

## **A manual of pathology / by Joseph Coats.**

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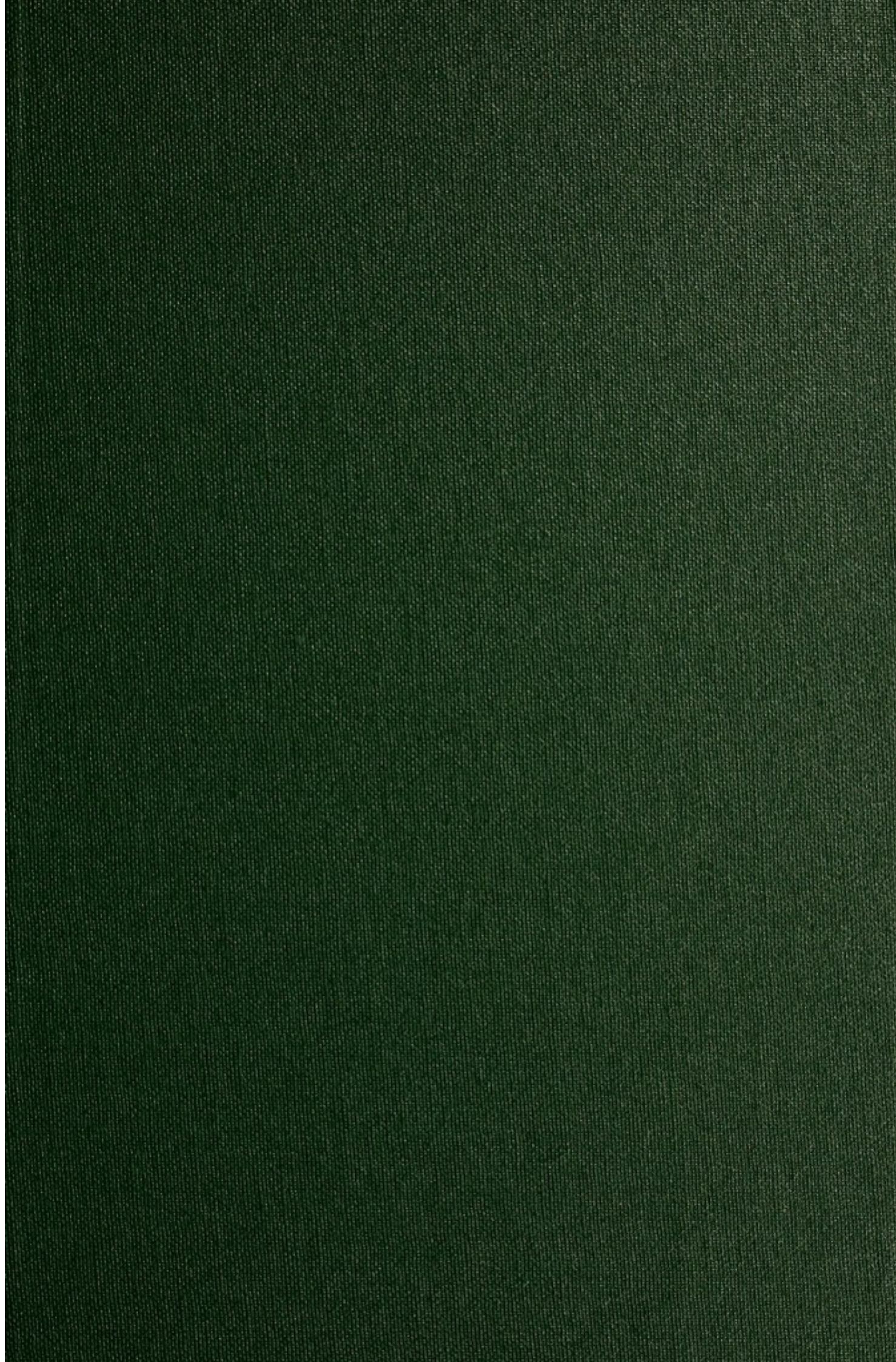
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James D. Cochran

Glasgow University

April 25<sup>th</sup> 189

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A MANUAL OF PATHOLOGY.



A MANUAL  
OF  
PATHOLOGY.

BY  
JOSEPH COATS, M.D.,

AUTHOR OF "LECTURES TO PRACTITIONERS ON THE PATHOLOGY OF PHTHISIS PULMONALIS ;"  
PATHOLOGIST TO THE WESTERN INFIRMARY AND THE SICK CHILDREN'S HOSPITAL, GLASGOW ;  
LECTURER ON PATHOLOGY IN THE WESTERN INFIRMARY ; EXAMINER IN PATHOLOGY  
IN THE UNIVERSITY OF GLASGOW ; FORMERLY PATHOLOGIST TO THE ROYAL  
INFIRMARY, AND PRESIDENT OF THE PATHOLOGICAL AND  
CLINICAL SOCIETY OF GLASGOW.

SECOND EDITION, REVISED AND MOSTLY REWRITTEN.

WITH THREE HUNDRED AND SIXTY-FOUR ILLUSTRATIONS.

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TO

SIR JOSEPH LISTER, BART., M.B., F.R.S.,

WHOSE METHOD OF REGARDING PHYSIOLOGICAL AND PATHOLOGICAL  
PROCESSES AFFORDED TO THE AUTHOR, DURING THE EARLIER YEARS  
OF HIS CAREER, AN EXAMPLE OF SCIENTIFIC INSIGHT FOR WHICH HE  
HAS NEVER CEASED TO BE GRATEFUL,

AND TO

W. T. GAIRDNER, LL.D., M.D.,

WHOSE ENTHUSIASM IN THE STUDY OF PATHOLOGY, IN THE POST-MORTEM  
ROOM AND AT THE BEDSIDE, HAS BEEN TO THE AUTHOR, DURING THESE  
MANY YEARS, A PERPETUAL STIMULUS, WHILE HIS GENEROUS FRIEND-  
SHIP HAS BEEN A CONSTANT ENCOURAGEMENT,

This Work is,

WITH GREAT RESPECT AND ESTEEM,

DEDICATED.



## PREFACE TO THE SECOND EDITION.

THIS edition, although greatly altered from the first, will be found to be similar in its objects and scope. In the preface to the first edition, issued in 1883, in referring to the scope of the work, I stated that "the whole aim of Pathological Anatomy is to elucidate the actual vital processes of disease, and it has been felt by the author in lecturing on Pathology, that the subject gains in interest and intelligibility when the Etiology, Anatomy, and General Pathology are all taken into account in forming a general conception of each morbid condition." The work, therefore, is not merely a treatise on Morbid Anatomy, but aims at the elucidation of Morbid Processes, on the basis of Experimental Pathology, Pathological Anatomy and Histology, and Clinical Observation.

While the objects and scope remain the same, I have found it necessary to make so many alterations and additions that the book has been almost entirely rewritten. Although not really more systematic, the arrangement is, in this edition, rendered more obvious to the eye by the adoption of prominent numbered headings which are intended to make the work more easy of reference, and also to aid the student in a systematic perusal. The summaries of contents introduced at the heads of the sections have a similar object.

The additions have been considerable in both the text and the illustrations. In the General Part three new sections have been added,

namely—Section I. Nature, Causation, and Terminations of Disease ; Section II. Teratology ; and Section XII. Fever, while in the Special Part there are two additional sections—V. The Thymus and the Thyroid, and XI. The Eye and the Ear. The new illustrations number over 70, nearly all of them prepared from my own specimens. I have also introduced brief notices of the Literature which are intended to guide the student in following out any particular subject. Notwithstanding these additions the number of pages is not greatly increased.

In the preparation of the section on the Eye and the Ear I have been much indebted to Dr. Thos. Reid and Dr. Thos. Barr, who revised the manuscript of this section. I am further indebted to Dr. Barr for the loan of the woodcuts illustrating the diseases of the Ear. Dr. Lindsay Steven has also kindly given me the use of two cuts.

I am again under great obligation to Dr. Geo. S. Middleton for a very careful revisal of the proofs and for many practical hints on various points.

GLASGOW, *August*, 1889.



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A MANUAL OF PATHOLOGY.





# A MANUAL OF PATHOLOGY.

## INTRODUCTION.

**P**ATHOLOGY is the scientific study of disease. The term Disease, which strictly means nothing more than discomfort, has reference primarily to the subjective sensations of the person affected, but as these feelings have an objective basis, and as it is possible to distinguish various groups of phenomena each with its own mode of divergence from the healthy conditions, the term diseases has come to be applied to these groups of phenomena. It might perhaps be more correct to call them morbid conditions, but the more familiar word has obtained the authority of general use.

In Pathology we aim at getting close to the various diseases, so as to study their essential nature. It is a scientific study, and it is pursued by scientific methods. There are three principal methods by which the nature of disease is elucidated, as there are three main sources of our knowledge of disease. These may be designated respectively, Experimental Pathology, Pathological Anatomy and Histology, and Clinical Observation. To each of these is Pathology indebted in endeavouring to form a conception of each disease in its true characters.

John Hunter may be said to have laid the foundation of experimental pathology. So far as it can be prosecuted, this mode of study, which aims at the production of morbid conditions in animals so as to observe them more closely, is calculated to afford us the most directly trustworthy information as to the nature of diseases, and much of the advance in modern pathology is due to this line of study. If we can actually induce a given disease in an animal, we already know a great deal as to its causation, and are in the way



of finding out much as to its nature. In some cases we are able actually to observe the morbid processes in the living animal; in others we can kill the animal at various periods after the onset of the disease, and observe the results at various stages. We can vary the experiment in different ways so as to eliminate on one side and another the various elements, and so reduce the problem to its simple and necessary parts. Experimental pathology has thus carried us far towards understanding many of the conditions in disease. Reference need only be made to the reformations which this line of study has effected in our knowledge of Inflammation, Embolism, Tuberculosis.

The study of morbid anatomy may not effect such revolutions in Pathology as experimental pathology, but it is by this method that the scientific basis of Pathology has been laid, and in the training of the individual medical student it is calculated to play an even more important part in giving him an understanding of the nature of the diseases he has to deal with. Morbid anatomy deals with the physical changes which disease produces in the body. These changes are, as it were, the expression of the pathological processes, and are only to be regarded as indications of the existence of these processes. It is probable that all morbid processes produce changes in structure, although the changes may sometimes elude our observation. For the most part, however, we are able to associate the change of structure with the morbid process, and the observed change leads us up to the actual nature of the disease. In fact, the nature of disease might be studied on the basis of the morbid structure, advantage being taken also of all information afforded by experimental pathology and clinical observation.

Morbid Anatomy is the study of the structural changes in all periods and stages of the disease. It is sometimes objected to this mode of study that in observing the state of the tissues after death we see only the last stages of disease, and in some cases only the results. This, however, is not the case. Every clinical observer will admit that in the same disease the fatal issue occurs in different cases under the most diverse circumstances. Death ensues, as a rule, when by reason of defective nutrition, or a deficiency in the supply of oxygenated blood mainly by interference with the respiration, the functions of the body can no longer be carried on. As there is in different individuals an infinite variety in the vigour of the vital processes, so a condition which in one will very slightly affect the essential phenomena of life, will, in another, seriously compromise them. This is an experience of our every-day life. And thus it happens that patients die in all stages of



disease. Even in the same case we are often able to study the various changes in their different stages.

As an illustration of what is meant, take the case of pneumonia or acute inflammation of the lungs. Patients die in this disease, as a rule, from failure of the heart and consequent cessation of the circulation; but the paralysis of the heart is not related solely to the condition of the lungs. It is often much more closely related to the elevation of temperature which is to be traced to changes in the nervous system, while the power of resistance of the heart in each person is an important element in the matter. And so it happens that one man will die with a limited and advancing pneumonia, while another will survive through a very extensive condensation. The lung at death in the one case is in the same stage of the disease as in the other at a certain date during life. We can study the nature of the disease in the former and we are surely warranted in assuming that, in the patient who survived, the state of the lung is virtually identical with that in the patient who died.

Again, in a carcinoma growing up to death, we have, at the growing margins of the tumour, changes which are the same as those by which the tumour has all along been extending, and we can study the methods of its growth as well as its relations to the structures in which it has been growing.

Thus it is that, although morbid anatomy by itself may in some cases throw little light on the nature of the disease, yet, taken as an expression of the morbid process, it is of the highest value. It may be added that, as similar processes must have similar changes as their correlatives, we may take these changes as a kind of test of the existence of the processes.

In referring to processes and changes it must be understood that the finer structure of the body is taken into account. In Pathological Anatomy, we include Pathological Histology, and it may indeed be said that, as the coarser, naked-eye appearances are merely the expression of the finer structural details, so we shall not understand the former unless we are able to refer them to the latter. We can have no accurate idea of the proper structure of the normal liver by simply looking at it with the naked eye, and we can have little insight into its morbid conditions by the same method of observation. The morbid conditions, however, do generally cause changes in the naked-eye appearances, and this is of great consequence as enabling us, when roughly examining an organ, to infer that its structure is changed. But these macroscopic appearances are only of value when they are used to throw light on the microscopic changes or to direct attention to them. The really important processes are those which concern the finer elements of the



tissues, and so it is to these that we shall chiefly attend in studying the various forms of disease.

Clinical observation is of great importance in the study of pathology, and that in various ways. If what has been stated above be correct, then it is the processes occurring during life that are the real subject of study in pathology, and it is only by careful clinical observation that we are able to get, as it were, into close relations with these processes during their actual currency. Clinical Observation and Pathological Anatomy should thus be taken as mutually elucidating each other, and as working together towards just conceptions as to the nature of the conditions concerned.

There is another aspect in which clinical observation is of importance in relation to pathology. The pathologist, if conversant only with morbid anatomy and experimental pathology, is apt to form a too mechanical and simple conception of the processes involved, and is likely to theorize on the basis of a too limited view of the subject. Clinical Observation is a needed corrective to this, bringing the pathologist back to the practical aspects of the matter, and checking the tendency to theorize by the discipline of facts.

In the systematic treatment of the various forms of disease, the plan adopted has been to take up first the diseases in their general aspects, and afterwards the diseases of the special organs and systems. The work, therefore, divides itself naturally into a General Part and a Special Part.

In the General Part will be taken up those diseases which are not confined to any organ or tissue; most of them, indeed, may affect any organ of the body. This is preceded by a Section on the Nature, Causation, and Terminations of Disease.

In the Special Part, the diseases as they manifest themselves in the individual organs and tissues will be considered. It is clear that the diseases described in the General Part will again come under review in each division of the Special Part, and they will be taken up very much in the same order as in the General Part. But each disease presents various specialities according to its locality, and, from a practical point of view, it is important to understand the modifications thus occurring. The Special Part will be, in a certain sense, an expansion and amplified illustration of the general principles educed in the General Part.



## PART FIRST.—GENERAL DISEASES.

### SECTION I.

#### NATURE, CAUSATION, AND TERMINATIONS OF DISEASE IN GENERAL.

NATURE OF DISEASE.—*Normal activities of tissues concerned with self-preservation ; in disease their efficiency impaired.* CAUSATION OF DISEASE.—*Action of external forces nearly always present, but powers of resistance in the tissues an important element.* A. *Influence of external forces: (1) direct action of the physical forces, poisons and morbid poisons ; (2) indirect action in producing susceptibility to disease.* B. *Influence of internal conditions : (1) congenital diseases and susceptibilities ; (2) influence of inheritance, in producing racial and individual characteristics, structural and physiological abnormalities, and susceptibilities or predispositions to disease ; (3) influence of age and sex.* TERMINATIONS OF DISEASE.—*Recovery implies vis medicatrix naturæ. Death by failure of respiration or of heart's action.*

**P**ATHOLOGY, as we have seen, is concerned with morbid conditions, taking account of the causes of disease, the nature of the changes induced in the tissues, and the resulting alterations in function. This implies a knowledge of the condition of the body in health and a comparison between the healthy and morbid states. In this preliminary section it will be proper to lay down some general principles in regard to the difference between normal and pathological actions.

In our conception of the living body we are not to look upon it merely as a whole, but also in its finer elements. John Hunter laid it down as one of his most important primary principles that “every individual particle of animal matter is possessed of life, and the least imaginable part is as much alive as the whole.” This principle of John



Hunter's may be regarded as the basis of the modern **Cell theory**, which, in the hands of Goodsir and Virchow, has been applied to Pathology with very great effect. According to this theory, which may now perhaps take rank as a definitely established principle, the tissues of the body contain multitudes of quasi-independent bodies, which have their own actions and their own life. We do not properly appreciate any action of the body, whether normal or pathological, unless we take into account the finest details of the tissues and have regard to the active cell-life which goes on there.

The primary actions of the cells will be found to be remarkably similar to those of the larger structures of the body and of the body as a whole. Each of the cells is engaged, in the first instance, in **self-preservation**. It has its own substance to preserve, and it has also in many cases a certain portion of intercellular substance to look after. Its first duty is to resist any disintegrating action and to make up for any waste which may occur in the exercise of its ordinary functions. In like manner the body as a whole has, for its primary action, the preservation of its own existence, opposing its resistance to forces external to itself, and making up for the ordinary waste of life. To this are subservient all the constant organic functions, digestion, respiration, circulation, etc. The individual organs of the body have also, in many cases, special arrangements for self-preservation, which come into play when any interference from without incommodes them in their functions. Thus resistance to forces external to themselves in order to self-preservation enters into the primary actions of all living animal structures, and it will be found that living bodies, whether in mass or in detail, have an inherent power of resisting external forces and restoring any injury such as dead matter does not possess. This may be illustrated before going further.

A living egg consists of a single cell, surrounded by a mass of nutritious matter destined to afford material out of which the future animal will be built up. But the egg, by virtue of the living principle in it, possesses, as John Hunter showed, a remarkable power of resisting external influences. It resists the influence of cold and of the agents of putrefaction to a degree which contrasts very strikingly with an egg which is not alive. When a dead egg (*i.e.*, one that has been killed by previous freezing) is placed in a freezing mixture its temperature soon falls to 32° F., and it begins to swell and congeal. In the case of a fresh egg, however, the temperature falls to 29½°, as does that of living animals, without freezing, and it takes 25 minutes longer than the dead one to freeze. Before freezing the temperature rises to 32°. Again, all through the process of hatching, the yolk, and what re-



mains of the albumen, continue perfectly sweet ; but if a dead egg be in the nest it becomes putrid like an ordinary piece of dead matter under similar circumstances. These observations of John Hunter show that a small piece of living matter has a special power of resisting the influences of external forces such as dead matter does not possess.

The body as a whole is also engaged in resisting the action of forces external to itself. An example of this is afforded by the preservation of the bodily temperature in warm-blooded animals. In such animals there is a continuous loss of heat by radiation and convection, and it is an important function of the body to compensate for the loss. Again, locomotion and even the retention of the erect posture are effected by the opposition of the various organs of locomotion to the action of gravitation. The resistance to this force involves the functions of bones, ligaments, muscles, and nervous system.

In the actual living body, then, the cells are engaged in preserving themselves from influences which may exist inside the body but are external to themselves, the regular wear and tear of life, and the body as a whole is similarly engaged. Health is preserved so long as the individual elements and the tissues as a whole are completely successful in resisting all disturbing influences, so long as they preserve themselves in a perfect state. In that case all the functions are performed with such ease that they do not obtrude themselves on the consciousness, and the body as a whole is preserved from decay and injury. On the other hand, when the forces external to the body as a whole or to its individual elements successfully obtrude themselves and impair the due efficiency of the living structures, then disease supervenes. In disease, therefore, we have to take into account the vital activity of the tissues, chiefly of the cellular elements, and the alterations wrought by forces external to the structures concerned.

## CAUSATION OF DISEASE.

In the causation of disease we have to look chiefly to external forces as the immediate agents, but we have also to take into account tendencies and susceptibilities existing in the tissues, which tendencies and susceptibilities may sometimes amount to actual determining causes. Disease may be induced in two ways : on the one hand external agents may attack the tissues with such energy as to break down the resistance of the living structures, or, on the other hand, the tissues may be so weakened as to be unable to resist the impact of the ordinary forces external to themselves. This may be made clearer by illustration.



At the mucous and cutaneous surfaces of the body there are innumerable microbes or bacteria capable of preying on animal matter of exactly the same chemical constitution as the tissues on which they lie. When the body dies these begin their action on the tissues, and the process of putrefaction leads to the disintegration of the tissues. During healthy life, however, these microbes are restrained by the living epithelial cells of the mucous and cutaneous surfaces. It seems, indeed, an important function of these cells to prevent the inroads of such living organisms. But let an injury of some kind destroy the protecting layer of cells, then the agents of putrefaction may find entrance, and by their action produce serious inflammations. Again, some of these bacteria, even when they find entrance to the tissues, produce no effect in most persons, while in others, by reason of some weakness in the powers of resistance, they lead to the processes of disease. It is so in regard to Tuberculosis, for instance. Most people are exposed to the inroads of the tubercular bacteria, inhaling them into their lungs, and taking them with their food into the alimentary canal, but only a small percentage of those exposed acquire the disease. On the other hand, bacteria may be of such vigour that they readily overcome the resistance of the living barriers, even in healthy persons, and find entrance to the body. Thus, in an epidemic of cholera most people who are exposed to the contagion—that is, most people who take into their stomachs the microbes concerned—become affected with the disease.

As another illustration, we may cite the ordinary external mechanical forces. Our tissues are formed to resist all ordinary mechanical forces to which they are exposed. They will succumb, however, to exceptionally vigorous impacts of the mechanical forces, and so arise such morbid conditions as wounds, fractures of bones, etc. But, on the other hand, the tissues may be primarily altered so as to expose a less degree of resistance than usual to the impact of the mechanical forces. There is a condition, for example, in which the blood-vessels are readily ruptured and the resulting hæmorrhage is with difficulty stayed. Persons so affected may bleed to death from a trivial injury. Again, there are persons whose bones are unable to support the weight of the body efficiently; they bend under it. These are examples of inherent morbid conditions which, indeed, do not come into evidence till the external forces act on the body, but which are present as definite changes in the tissues, and are hence designated diseases or morbid conditions.

From these illustrations it will appear that, while in the production of the actual manifestations of disease external forces are, in most cases at least, the determining causes, yet that in many cases there are pre-



existing conditions in the tissues, which form an important element in the causation, sometimes even more important than the other. This pre-existing element may be some manifest and definite imperfection in the structure of the body rendering it inefficient in relation to some of the ordinary actions of life, or it may be some obscure peculiarity which renders the tissues less able to resist certain external agents. In the one case the pre-existing morbid condition is described as the **Disease**, in the other, we speak of **Susceptibility** or **Predisposition** as existing. It is to be noted that such susceptibility or predisposition is usually to a particular form of disease and that it may exist in persons who are otherwise perfectly normal. It is clear also that the susceptibility may exist throughout life without the person having been exposed to the external influence which is necessary to the production of the disease.

The term **Diathesis** is frequently used to designate a peculiar constitutional state, predisposing persons to certain forms of disease. Thus the rheumatic, scrofulous, and gouty diatheses are spoken of. These conditions are, however, too vague to be of much real scientific value, and, as a matter of fact, the term is often used to designate not merely a predisposition but an actually existing condition of disease.

#### A.—INFLUENCE OF EXTERNAL FORCES IN THE CAUSATION OF DISEASE.

External influences may have a direct or an indirect influence in the causation of disease. That is to say, they may act directly on the tissues so as to bring about certain phenomena which we term diseases, or else they may so influence the tissues as to increase their liability or susceptibility to certain diseases.

##### I.—DIRECT INFLUENCE OF EXTERNAL FORCES.

1. **The physical forces**, mechanical, chemical, thermal, and electric, often directly affect the tissues. Their effect is to damage the parts on which they act. The mechanical forces cause wounds, bruises, fractures, etc.; the chemical forces produce disintegration or less severe injury to the tissues; excessive heat or cold in their extreme degrees kill the tissues, and, acting less strongly, do a certain amount of damage. In all these cases the immediate effects are local ones, the living structures acted upon are injured. But as the tissues are living tissues, certain phenomena ensue which are the expression of the physiological processes, altered by the damage done to the tissue. We have a series of phenomena generally grouped under the term **Inflammation**, which



arise more or less directly from the damage done to the living structures, and more particularly to the blood-vessels. The phenomena of inflammation are essentially local, having their seat in the damaged part; they are the expression of its own action. To some extent these phenomena are also the result of an attempt in the part to recover from the damage inflicted, or, as it is called, of a **reaction of the living tissues** against the injury done.

But even in the case of purely physical causes, the phenomena do not always remain purely local. As the blood-vessels of the injured part are in communication with the general circulation, the products of the changes occurring in the tissues may be carried throughout the body, and produce changes in the blood and tissues as a whole. An example of this is furnished by some cases of extensive burns of the skin. The blood in the parts exposed to the excessive heat may be damaged; the red corpuscles are sometimes killed. The dead corpuscles are disintegrated and their products are carried with the general circulation, and may do damage, especially while in process of elimination by the kidneys. Again, in almost all inflammations the fluid of the blood passes out, to some extent, amongst the tissues or on to surfaces. In these situations it may undergo putrefactive or other changes, the products of which may pass back into the blood by way of the lymphatics or veins, and so be carried throughout the body. The result of this is a different series of phenomena, included under the term **Fever** or **Pyrexia**. Hence fever is a frequent accompaniment of inflammation, even when produced by physical causes.

2. **Poisons**, in the ordinary sense of the word, can scarcely be said to produce diseases. Some of them, indeed, are chemical agents, such as alkalies and acids, which act locally on the tissues, and produce inflammations in the usual way. Others, such as arsenic and cantharides, produce local inflammations, not by any obvious or gross chemical action, but by some more intimate influence on the vitality of the structures. These two agents, like many other poisons, whatever the portal by which they find entrance to the body, being carried by the blood, select certain localities on which to act. Arsenic acts as an irritant on the mucous membrane of the stomach and intestines, producing the phenomena of inflammation there; cantharides acts similarly on the kidneys and bladder. Most other poisons resemble these two in respect that they select a certain locality for their action, and that they produce their effects by some intimate influence on the vital processes of the living tissues, this influence probably having to do with the finer chemical processes concerned in the vital phenomena. In the case of most poisons, the nervous system is more or less affected, many of them



having special kinds of actions, and some of them selecting special parts of the nervous system. Thus, strychnia acts as an irritant upon the motor centres of the cord and medulla oblongata. The term **Intoxication** (which literally means poisoning) is used by Continental writers to designate the effects produced on the nervous system by poisons in general. In this country it has become appropriated to the effects on the nervous system of poisoning by alcohol.

3. **Morbid poisons** stand in a totally different position from ordinary poisons. The term is used for agents, which, when introduced into the body, increase by self-multiplication, and produce effects altogether disproportionate to their original amount. They may be introduced in exceedingly minute quantities, but as a result they may have very severe effects, resulting even in death. Each of these morbid poisons produces a specific form of disease.

It may now be confidently asserted that these morbid poisons are, at least in the great majority of cases, related to minute vegetable organisms, belonging to the class of **Bacteria**, but often called somewhat generally **Micro-organisms** or **Microbes**. They propagate chiefly by simple division of their cells, and do so, generally with great rapidity, so that they multiply very greatly in a limited time. They do not produce their effects directly, but by means of poisons eliminated by them in the course of their vital processes.

Some of these microbes produce in the first instance purely local effects, these effects having as a rule the characters of inflammations, sometimes with special characters which serve to distinguish the various kinds of agents. Thus syphilis and tuberculosis always begin by the local application of the morbid poison, and their primary phenomena are more or less modified inflammations. But even those which have a primary local seat very readily extend their influence beyond it to the circulating blood, while many forms appear to have no primary local seat, but immediately pass into the blood. It is so with the morbid poisons of typhus fever, yellow fever, scarlet fever, and others. This extension to the blood may be of the morbid poison itself, the bacteria propagating into the blood and multiplying there, or it may be a mere filtration of the poisonous products of the micro-organisms, the latter remaining local. Thus tuberculosis may remain entirely local, and yet fever is a nearly constant result from the products passing on into the blood. Indeed there are some forms which apparently never pass beyond their local seats and yet may produce general phenomena. Thus the microbe of cholera appears to confine itself to the intestinal canal, but its products are absorbed and produce effects upon the nervous system and otherwise.



The terms **Infective** and **Zymotic Diseases** have been applied to affections of this class with reference to the fact that the morbid poison is self-propagating. The term **infective** is to be carefully distinguished from **infectious**. The former has reference to the extension of the morbid poison in the person affected. In multiplying the agent infects neighbouring parts, and so an infective disease is a spreading one, extending the territory affected. Infectious, on the other hand, refers to the extension from person to person. A disease may be infective, yet not infectious, although many diseases are both. Thus tuberculosis is a typically infective disease, having a characteristically spreading character, while it is doubtfully infectious. Syphilis, on the other hand, is both infective and infectious.

The mode of extension of this class of diseases amongst the community is somewhat various, and in some cases somewhat difficult to determine. There are many in which the disease is simply passed from person to person, either directly or by clothing, excreta, or otherwise. Such diseases are called distinctively infectious or contagious. This latter term is sometimes confined to the propagation by actual contact, but it is often used more generally to distinguish an indirect transmission. The term **contagium** or **contagium vivum** is equivalent to morbid poison.

But the mode of extension is in many instances not so simple as this. There are some morbid poisons which appear to reside in special localities, being apparently fostered by the special conditions of temperature, moisture, and other factors in these localities. In order to acquire such affections a person must visit the locality, and the morbid poison must, as it were, rise from the ground and pass into his body. The term **Miasma** is applied to such morbid poisons, and the diseases are called miasmatic. The most typical instance of it is the malarial fevers, although there is reason to believe that acute pneumonia, acute rheumatism, and some other affections are miasmatic.

There are some diseases which may be propagated either by miasma or infection. Examples of these are cholera, typhoid fever, dysentery, yellow fever.

To the class of diseases here under consideration the terms epidemic and endemic are frequently used. A disease is **epidemic** when it occurs in large numbers within a limited time in a place where it is not generally prevalent. The occurrence of an epidemic implies the introduction in considerable quantity of a morbid poison, and in such a case the disease will mostly be infectious. On the other hand, a disease is **endemic** when it is habitually prevalent in a particular locality. In that case the disease is most likely to be miasmatic in its origin.



## II.—INFLUENCE OF EXTERNAL CIRCUMSTANCES IN PRODUCING SUSCEPTIBILITY TO DISEASE.

External circumstances doubtless produce definite effects on the living body, but these effects are frequently very difficult to determine. Climate, temperature, condition of the atmosphere as regards purity, or moisture, the character of the soil, the dwellings, the food and drink, the clothing, the occupations of men, all have influence on their susceptibility to disease. The geographical distribution of disease and the effects of occupations have been made especially the subject of observation, but a more particular examination of these points scarcely lies within our province.

External circumstances will affect the living tissues chiefly by rendering them less vigorous in their resistance to morbid influences. As an illustration of this we may cite the effect of **exposure to cold** in the causation of disease. It is an admitted fact that exposure to cold, especially when the body is fatigued and the general vitality lowered, is frequently followed by attacks of disease. But the exposure can scarcely be the cause of the disease as it produces in different persons most diverse results. Thus a similar exposure will lead in one person to acute catarrh of the air-passages, in another to acute rheumatism, in a third to acute inflammation of the kidneys, in another to diarrhoea, and so forth. It is clear that the effect of the exposure is, in most cases by reflex nervous influences, to render some part unduly susceptible to the exciting causes of disease, the locality which is thus rendered susceptible being determined often by the individual characteristics of the person. In many cases the disease, indeed, is due to a morbid poison, which is able to make good its entrance when the structures are weakened. In what is specifically designated a common cold, the exposure to a low temperature is merely one of the predisposing conditions. Colds frequently arise without any such exposure. The fact that such colds are often obviously infectious is an indication that they are due to morbid poisons, which must be very prevalent in the community.

**Occupations** will influence the production of disease according as they tend to weaken particular tissues. Employments which imply severe muscular exertion, for instance, cause considerable strain on the vascular system, as every strong muscular effort is accompanied by a closure of the glottis, with suspension of the respiratory movements, and a rise in the blood-pressure. Such efforts frequently repeated will render the heart and large vessels peculiarly susceptible to disease. Hence we find that affections of the heart and arteries, more particularly aneurysm,



are much more common in men than in women, and in men who are engaged in occupations which require strong and sustained muscular exertion, such as blacksmiths, engineers, etc. Again, the frequent respiration of impure air, while it leads to a general weakness of the body, produces a special susceptibility to disease in the lungs. On the other hand, the inhalation of dust often leads directly to disease of the lungs, by the particles of dust actually injuring the lung tissue.

## B.—INFLUENCE OF INTERNAL CONDITIONS ON THE CAUSATION OF DISEASE.

From what has gone before it will appear that the condition of the tissues will form frequently a strong predisposing element in the causation of disease. It will also appear, however, that there are certain conditions of the tissues which in themselves are counted diseases, and which arise without any apparent external cause having acted; they are in this sense **spontaneous**, although this expression as well as the synonymous one **idiopathic**, merely indicate that the cause is obscure.

Looking at the tissues as living, we recognize that in their activity they are capable of morbid change in the direction of defect on the one hand and excess on the other. External causes may produce either of these morbid changes, damaging the tissues on the one hand or stimulating them to undue activity on the other. But the tissues in themselves may be defective or excessive, and may be so either in actual structure or in their tendencies and susceptibilities. These defects or excesses may be obviously the result of the action of external forces, but internal conditions have frequently a strong and sometimes a paramount influence in their causation. It will be proper to consider several different kinds of influence.

## I.—CONGENITAL DISEASES AND SUSCEPTIBILITIES.

A condition is congenital when the person is born with it. There are some diseases which are manifestly present at birth, and for the causation of which it is necessary to go back to the period of life *in utero*. Some of these are obviously due to the action of external forces, mechanical or other, but there are others in which the causation is quite obscure. Among those whose cause is clear may be cited congenital syphilis, which is always a manifestation of hereditary syphilis. In this case a morbid poison is propagated from the parent to the offspring and acts on the latter while still unborn.



Many of the congenital diseases consist in **Malformations** of the body as a whole or some part of it. These occur during the course of development and growth of the body, and consist in errors of excess or defect in the formation or disposition of the tissues. Some of them can, without difficulty, be traced to the action of external forces in the foetus. For example, a limb or part of one may be amputated in the uterus, and the person may be born with a corresponding defect, perhaps partly remedied by subsequent growth of the injured part. There are probably many congenital defects, which, if they could be traced back to their origin, would be found due to mechanical interference, but it is only in regard to a few of them that this can as yet be done. Such mechanical interference may be due to inflammation, producing adhesion of parts, and thus hindering their due expansion. It seems probable that inflammations play a considerable part in the causation of diseases in the uterus, and these inflammations will be due to the action of external forces just as are those of extra-uterine life.

In regard to congenital susceptibilities, these, again, may be divided into those tending towards defect on the one hand and excess on the other. Persons are born, it may be, with certain of their tissues unduly weak, and tending to decay. These tissues may give way and become the seat of disease as a result of the action of causes which in most persons would produce no such effect. It may even be that at a certain period of life they may degenerate without any apparent determining cause. Then, as opposed to this, people may be born with a tendency to undue excess in the growth of certain parts of their tissues. The causation of **Tumours** or **Morbid growths** is very obscure, but we may say at least that obvious congenital conditions, such as soft warts or moles, are not infrequently the starting point of tumours in later life. We may also infer that unseen conditions, which were equally congenital, may be the starting point of other tumours, the susceptibility to which has existed during life.

It will appear from what has gone before that congenital diseases are due to a variety of causes, and that they owe comparatively little to inheritance. All that the term means is that the child comes into the world with the disease existent. Congenital susceptibility, on the other hand, will be for the most part due to inheritance.



## II.—INFLUENCE OF INHERITANCE IN THE CAUSATION OF DISEASE.

At the outset it is necessary to distinguish two very different classes in relation to the inheritance of disease.

**Hereditary Diseases.**—The term hereditary disease is usually applied to cases in which a definite morbid condition is transmitted from parent to offspring. The commonest case of the kind is Hereditary Syphilis. Here a morbid poison, presumably related to a form of microbe, is transmitted by the parent to the offspring. It is a case in which an external force acting on the parent is directly passed on to the offspring, and becomes active in the latter as it has been in the former. Many of the acute fevers have been similarly transmitted to the child *in utero*, the foetus either passing through the fever successfully or dying from it. Thus small-pox, intermittent fever, measles, relapsing fever may be communicated to the child *in utero*. Children have been born with the eruption of small-pox or the cicatrices visible on their skins; others have presented the large spleens and cachectic appearances of ague.

It is obvious that in all these cases the hereditary form of the disease does not differ essentially in its causation from the ordinary forms of the same disease, and that inheritance is in no sense a cause, but merely a mode of transmission of the cause. We may therefore entirely eliminate this class of cases in considering the influence of inheritance in the causation of disease.

Most cases of hereditary disease in this sense are congenital, but they are not necessarily so. In the case of syphilis, for instance, the manifestations of the disease may not be present at birth, although later on lesions occur which are due to hereditary transmission. It may also be remarked that the use of the term hereditary disease, as distinguished from inherited disease, is to some extent arbitrary, but it is justified by convenience and also by the fact that true inheritance does not usually produce disease, but rather a susceptibility; hence the term hereditary disease is hardly correct, except in the sense in which it is used above.

**1. The General Principles of Inheritance.**—In order to determine the domain of inheritance in the causation of disease it is necessary first to consider the nature of its influence in the normal anatomical and physiological conditions of our bodies.

**Race the Result of Inheritance.**—We see the influence of inheritance not only in the fact that human parents produce human offspring, but also in that the offspring conform to the **race** of the parents. It is



important in connection with what follows to consider what constitute the differences of the races of men. There is more or less pigment in the deeper layers of the epidermis; the hair, in transverse section, is round or oval in outline, or more or less oval; the nasal bones are articulated at an acute or obtuse angle; the lower jaw is more or less massive; the shape and details of the bones of the skull are various; the eyes are horizontal or are directed upwards at their outer extremities; and so on. It is these differences in structure that make the chief distinctions of race, and, if we compare two such dissimilar races as the negro and the Englishman, we shall see that the difference lies in a multitude of details, each of which if taken apart might seem trivial. And yet the fertilized ovum inherits the power of modelling the bodily frame down to these minute details, and it practically never goes wrong.

**Individual Characters the Result of Inheritance.**—But, besides the characters which distinguish men as men, and the men of one race from those of another, we know that **men are distinguished** from each other **individually**. When we consider the general likeness of the men of one race to each other, it seems marvellous that, among the multitudes with whom we come in contact, we rarely mistake one for another. The only explanation of this is that there are innumerable shades of difference—in colour of skin, hair, and eyes; in shape of nose, mouth, and eyebrows—an infinite number of small peculiarities, which make up the whole portrait. We unconsciously blend the whole of them in one mental picture of the individual, and if one of them be altered we at once recognize that our portrait is out in some way and has to be re-adjusted, as when we change the arrangement or length of the hair or beard, or when the mouth is altered by the loss of teeth or the addition of an artificial set. This may be further illustrated by the well-known fact of the insufficiency of photographs. In them we have an exact picture but without the colour, and the absence of this one element in the portrait has often the effect of almost destroying the likeness.

It need hardly be said that most of these fine shades of difference between man and man are due to inheritance. The ovum enters on its career with these already inherent in it, and the various minute points of difference are but repetitions variously compounded of points in its progenitors. We are so accustomed to look for resemblances between children and parents that we are apt to forget what these imply. It is a particular colour of the iris, a special tone in the voice, a trick of manner, such as a twitching of the upper eyelid, or a way of fidgeting with the hand, which recalls the parent or grandparent. These minute points of resemblance become the more remarkable when we remember that they



are all inherent in the fertilized ovum. Without any external assistance it goes on modelling the tissues and endowing them with their functions according to a prearranged pattern, and this power continues throughout life ; so that it often happens that, on to middle life or old age, points of resemblance come out which had not been previously visible. Sometimes a child will resemble one parent most up to puberty, and will then, in undergoing the extraordinary changes of that period, come to resemble the other more. The child is not a mere reproduction of its parents, but it is as if the mother and father transmitted to the ovum certain forces inherent in themselves and derived from their ancestors. These forces will be variously proportioned in each case, and the product will be a very complex one. It is quite impossible to tell how, in any particular case, these forces are mingled, but we can often identify the individual items as coming from this or that parent.

A highly important illustration of the truth of what has been adduced is afforded by the facts connected with **Twins**. We have seen that each ovum which leaves the ovary and becomes fertilized is thereby endowed with the forces which mould it into the future human being. These forces seem variously distributed in the ova and the spermatozoa, so that each ovum has them in such varying proportions that even the brothers and sisters of the same family are often very different. But if a single ovum produces two embryos which are simultaneously fertilized, then we might expect that the forces inherent in these two embryos would be much more nearly alike than usual, and that in consequence the two individuals would resemble each other more closely than is usually the case. And so it is. There are two kinds of twins. In the one the individuals are derived from separate ova, just as the progeny of animals which bear several young at the same time are ; while in the other kind there is only one ovum, which probably produces the twins by fission. In the case of twins derived from the same ovum we may look for close resemblance, and they will always be of the same sex. In the other case, where they are derived from separate ova, they may or may not be of the same sex, and they will not necessarily resemble each other more closely than the ordinary brothers and sisters of a family. In a well-known work by Francis Galton we have a collection of interesting facts in illustration of the life-history of twins. He gives many anecdotes supplied by twins as to the mistakes made on account of close likeness, from which it will be gathered that the "Comedy of Errors" is scarcely a burlesque. The resemblance in many cases lasted throughout life, and extended to the smallest matters, even of feeling and thought, and, indeed to their illnesses. From these facts we have



demonstrative proof that in the fertilized ovum we have already inherent virtually all the finer details of structure and function, although many of these only come into effect in after-life. It will be observed that the sex is determined in the ovum.

**2. Inheritance of Structural and Physiological Abnormalities.**—From what has gone before we have seen that minute differences in local structure are the usual subject of inheritance. Coming to more considerable differences in structure, such as constitute more or less definite divergences from the normal, it is not surprising to find that they are also inherited.

A common structural peculiarity is webbing of the fingers or toes, and this comparatively slight divergence is very distinctly matter of inheritance. The author knows of a case in which the second and third toes of both feet are imperfectly separated, so that, although the bones are complete, the toes are united nearly to their tips by webs of skin. This peculiarity existed in a man who transmitted it to a son, who died in boyhood, to a grandson, the son of one of his daughters, and, although in a diminishing degree, to several great-grandchildren. Sedgwick, in his very elaborate series of papers in the *Medico-Chirurgical Review* for 1861 and 1863, adduces many cases of inheritance of supernumerary digits, crooked fingers, cleft iris, squinting, etc. He gives, for instance, a case in which a supernumerary finger was attached to the outer side of the first phalanx of the little finger. The deformity had occurred for five generations, and the person under observation was the fourth child of a family, all of whom except the second were born with this deformity.

There is also the well-established inheritance of **Ichthyosis**, the most striking instance of which is that shown by the family of Lamberts, the so-called "Porcupine family," adduced by Darwin in his "Animals and Plants under Domestication." In the original Lambert the skin was covered by warty projections which were periodically moulted. He had six sons and two grandsons similarly affected, and it is remarkable that the inheritance was confined to the male sex, the two grandsons having seven sisters who were free of it. A still more striking instance of the inheritance of ichthyosis is related by Sedgwick. The disease was observed in a boy fourteen years of age, and on tracing his family history it was found that he derived it from his grandfather. This man had been affected with the disease, and it is a striking example of atavism or latency in one generation that none of his children showed the disease, although there were three sons and three daughters, while of seven grandchildren (five males and two females) four of the males were affected and none of the females. The affected males were the children of daughters of the original case.



**Hæmophilia** or hæmorrhagic diathesis depends on an unknown structural peculiarity either in the blood or vessels, by virtue of which bleeding readily occurs, and when it takes place is stopped with difficulty. This condition is in the highest degree inherited, and it is so in a very peculiar way. Many cases have been traced in their connections, and it appears that it is never directly transmitted. It occurs almost alone in the male sex, but a person affected does not transmit it to his own sons but to those of his daughters. It always in this way misses a generation, and appears in the grandsons of the person transmitting.

**Daltonism** or **Colour-blindness** is also demonstrably inherited, and it is usually transmitted in the same peculiar fashion as hæmophilia. In nine tenths of the cases Daltonism occurs in the male sex, and it is usually transmitted to the grandsons of the original case through his daughters. (Sedgwick, Wickham Legg.)

**Diabetes insipidus** is another condition, which is apparently inherited in a remarkable degree. In this affection the person drinks excessively and passes a large excess of urine. We may presumably infer that the kidneys are unduly large, and we know that the bladder is unusually capacious. This was proved in a case recorded by Dr. Finlayson, and was also determined by Weil in the cases to be presently mentioned. Gee has recorded cases showing that this condition is hereditary, but Weil records a most remarkable instance of its inheritance. Weil met with a case, and he set himself, by hunting up and down the country, to find all about the relatives of his patient, with the result of producing a most elaborate genealogical history of the family. He traced ninety members of the family extending through four generations, seventy of whom were living at the time of the observation. He personally investigated most of these seventy cases. The disease was traced to a man called Johann Peter Schwartz, who was the common ancestor of the family, and was born in the year 1772. His descendants were five children, twenty-nine grandchildren, and fifty-six great-grandchildren, in all ninety-one persons. Of these no less than twenty-three were certainly diabetic, and thirteen doubtfully so. Omitting the doubtful cases altogether, there remain seventy-eight persons, of whom twenty-three were affected, or 30 per cent. According to Weil this condition was perfectly consistent with good health. They were evidently prolific enough and generally long lived. The original Schwartz lived to be eighty-three, and of his three daughters one died at seventy-four, and the other two were still alive at the time of observation, being respectively seventy-six and sixty-seven years of age.

The facts hitherto adduced suffice to show that, in accordance with



what we find in regard to normal differences in structure, abnormal peculiarities in structure or function are also subject to inheritance. It now remains to show on a similar basis of fact that susceptibilities to disease are also matters of inheritance, and that the varying susceptibilities of different persons depend on peculiarities, which cannot be, perhaps, demonstrated like those given above, but which frequently coexist at least with definite peculiarities of structure.

### 3. Inheritance of Susceptibilities or Predispositions to Disease.—

We have seen that certain forms of disease are due to the action of morbid poisons, which, obtaining an entrance into the body, multiply there and give rise to certain phenomena. In the case of some of these diseases microbes have been demonstrated as the immediate vehicles or sources of the morbid poisons. It is matter of frequent observation that different persons are very variously susceptible to diseases of this class, and it is an important fact that inheritance plays an important part in determining the varying degrees of susceptibility.

**Race** is an important factor in determining the susceptibility to this class of diseases. A striking experimental illustration of this has been afforded by Koch. He found that by inoculating mice with putrefying blood he could induce a specific acute fever, which was exceedingly fatal, the animals dying in from forty to sixty hours after inoculation. The disease thus induced is exceedingly infectious, so that the most minute portion of blood from a mouse which has died of it suffices to produce the disease when inoculated into a healthy mouse. It is a fact of great importance that, while this is the case in the ordinary mouse, yet the field mouse, so similar that it is scarcely distinguishable at a first glance, is absolutely insusceptible. Koch ascribes this to differences in the blood of the two animals, the existence of marked differences being shown by the fact that the blood of the field mouse readily produces, from the red corpuscles, hexagonal crystals, while that of the ordinary mouse does not. Whatever the difference may be, we have here an example of comparatively slight peculiarities producing the most profound difference in susceptibility.

A similar but less extreme difference is noted by Chauveau in the susceptibility of sheep to anthrax. This disease may be produced readily by inoculation in most animals, and in sheep among the rest. But it appears that it is much more difficult to produce it in Algerian sheep than in ordinary breeds, a larger dose being required. The racial peculiarities of this breed determine a striking difference in susceptibility. Pasteur found a similar difference in relation to the poison of chicken cholera amongst the different breeds of fowls. Ordinary fowls are highly susceptible, but those of the Cochin China breed are hardly at all so.



In man similar racial differences of susceptibility are observable. In a certain sense the "plague of lice" is a serious disease, and it is highly infectious. According to Murray (quoted by Darwin) the different races of men present great differences in the species of pediculi which infest them, and when those of one race stray on to the bodies of persons of a different one they generally survive only a few days. A structural or physiological peculiarity of the skin connected with the differences of race determines the susceptibility to the attacks of the parasite.

Race has much to do with the susceptibility to **Yellow fever**. There is abundant evidence that negroes are almost entirely insusceptible to this disease, which is so virulent in white men. The immunity of the negro is not due to his having lived for generations in countries where yellow fever prevails, because it is manifest when negroes are brought from parts of Africa where yellow fever is unknown. The absence of susceptibility extends to persons partly of negro descent, generally in proportion to the amount of the negro in their constitution. Negroes seem also to have considerably less susceptibility to **Malarial fevers** than the white races.

On the other hand, negroes are more susceptible to **Small-pox** than Europeans, and it generally attacks them in a more virulent form. They are also more susceptible to **Cholera**.

We may infer, therefore, that along with differences of race there go peculiarities of structure and function which determine differences of susceptibility.

**Family peculiarities** are of the same nature as racial ones, the latter being, probably, in their origin derived from the former. There are great and manifest differences among families in susceptibility to morbid poisons. There are many facts in existence which show that scarlet fever and diphtheria are special scourges in particular families. Again Tuberculosis of the lungs is acknowledged by almost all authorities to prevail in families, so much so that authors use the expression "family phthisis." It is difficult in the case of families to trace them through a sufficient number of generations, but such facts as those just indicated show that the peculiarities of structure which make family and individual distinctions are co-related to differences in susceptibility to the class of diseases under consideration.

### III.—INFLUENCE OF AGE AND SEX IN THE CAUSATION OF DISEASE.

It will be understood from what has gone before that age and sex will affect the causation of disease only by increasing or diminishing the susceptibility to certain diseases.



The first weeks and months of life are the most fatal, so much so that about a fourth of the children born die within the first year, while in many towns more than half the deaths are of children under a year old. There are various explanations of this, perhaps the chief being that the helpless infant can scarcely let its wants be known, and, being dependent on the care of others, is more susceptible to all sorts of accidental external influences. Hence it is that the children of the well-to-do are much less affected by disease than those of poorer parents, while illegitimate children who receive least care are much more liable to disease and death than those born in wedlock. Besides this, however, there are other conditions of susceptibility. The child, emerging from the protection and warmth of the uterus, undergoes a sudden transformation in its circulation and nutrition, and at the same time has to cope with physical conditions, chiefly mechanical and thermal, which are new to it. The first days and weeks of life are therefore the most trying to the child. In the child, again, the tissues are growing, and have the double duty of nutrition and growth thrown upon them. They are therefore more liable to derangement than those of adults, although they possess greater powers of restoration when damage is done. The smaller bulk of the child's body, also, renders it more susceptible to changes of temperature, so that cold will penetrate more deeply into its tissues than into those of adults, a fact which is not always remembered in the clothing of children. Besides all this the bodies of children present, as it were, a virgin soil for the propagation of the various morbid poisons. The children of the poor can scarcely escape exposure to the ordinary infectious diseases, and their bodies are apparently very susceptible to these influences. A striking illustration of this statement lies in the fact that, in Glasgow, whooping-cough is the cause of more deaths in children than is any other communicable disease (Russell). Even tuberculosis, which is not in the proper sense infectious, is exceedingly prevalent in children, and is much more liable to extend beyond its primary local seat than in adults; as if the tissues of children afforded a better nidus for the infective agent.

The greatest liability to disease is in the first year of life, and it diminishes from the first year to the eighth. From the eighth till the eighteenth year the liability to disease increases, but diminishes after the latter age, reaching a minimum between twenty-four and thirty, to increase from that point onwards. The mortality by no means follows the liability to disease. It is, indeed, greatest in the first year, diminishing from that period onwards. The diminution, however, does not cease at the eighth year, but goes on till adult life, the minimum being



between the ages of twenty and forty-five. After the latter age the death-rate rises regularly but slowly.

The increase of disease from the eighth to the eighteenth years is related to the fact that these are, in general, the years of school-life and include the time of puberty. School not only, by exposure in going and coming and sometimes by too prolonged confinement, renders the body less resistant, but also exposes children to infection. The period of puberty, involving serious changes in the bodily functions, is liable to induce a general weakness of the body, and also exposes females to the special affections of menstruation.

In advanced life, the body is liable to diseases which imply decadence or senility of the tissues. The bones are more brittle by reason of a diminution in their animal constituents, and hence are liable to give way under a less severe mechanical strain than ordinary bones are. The blood-vessels are more subject to senile changes affecting their liability to disease than any other tissues, and age is more potent in the causation of these diseases than any other factor. Affections of the brain are frequent in old persons, but nearly all of these depend on the condition of the blood-vessels and are not primarily diseases of the brain. Cancer is very specially a disease of middle and advanced life, but not of old age. It is characterized by an excessive growth of the epithelial tissues, and it is difficult to explain how this should occur preponderatingly at an age beyond that of greatest vigour unless on the assumption that, so long as the surrounding tissues remain vigorous, they restrain any tendency in the epithelial tissues to grow to excess.

**Sex** influences the liability to disease in several ways. Diseases of the generative organs are of course distinctive in the two sexes. The tissues of the male being, as a rule, larger and firmer, they might be expected to offer more resistance to the external forces, but this is very doubtfully the case, and at nearly all periods of life the female is less susceptible to disease than the male. It is a remarkable fact that, apparently in all lands, there are more male than female children born (in the proportion of 103-105 boys to 100 girls). But this disproportion begins to be reduced at the very point of birth, by the fact that more males are still-born than females (in the proportion of 14:10). This is apparently due to the fact that male children are on the average more bulky—especially have larger heads, than females, and are therefore more exposed to the accidents of parturition. Even after birth the females show less liability to disease, so that by the end of the first year the sexes are already equal in numbers. Throughout the remaining years of life the females are, at all ages, usually in the majority. Even the dangers of childbearing do not reverse the balance, as the risks are



counterbalanced by the greater exposure of men during the corresponding years of life, to accident and injury in the pursuance of their occupations.

### TERMINATIONS OF DISEASE.

The termination of disease is in recovery or death. The recovery may be much prolonged and only partial, or death may ensue only after a long struggle.

**Recovery from disease.**—It has been indicated above that disease is due to an interference from without with the normal physiological processes, or to an abnormal construction or tendency in the living tissues. The existence of such interference or tendency does not abolish the normal functions, but rather, in many cases, stimulates them to increased activity. The healthy tissues are continually engaged in dealing with external forces, and are generally successful in so doing; and, in cases where the external forces have for the moment obtained the supremacy, the tissues are frequently stimulated to exceptional vigour, so as to overcome the interfering agent. As already indicated, many of the phenomena of disease are really due to the reaction of the tissues against the agent producing the disease. There is therefore on the part of nature an attempt to get rid of the disease, a natural tendency towards recovery. This tendency is expressed in the aphorism **vis medicatrix naturæ**, this expression implying that nature possesses a power of healing.

It will follow from this that recovery is more apt to occur when the disease is due to the direct action of the external forces. In all diseases due to the physical forces, such as wounds, fractures, burns, etc., the tissues at once set to work to repair the injury, and in most cases with considerable success. In the treatment of such diseases the endeavour is made to place the parts under as favourable circumstances as possible for nature to do its part. In the case of diseases due to morbid poisons, again, we see usually an effort on the part of nature to overcome the morbid agent. According to one theory the elevation of temperature, which is characteristic of fever, is a means of destroying the morbid agent, and at any rate, in most cases, the normal forces, after a time reassert themselves, and the patient recovers. In some cases, on the other hand, the disease keeps on advancing with very little check on the part of the tissues. It will be found that when this is the case there has generally existed a special susceptibility to the disease, and that the attack of the morbid agent has, as it were, been invited by the special



condition of the person. If, after the morbid process has been established, the susceptibility remains, the disease is likely to remain and extend. In tuberculosis this seems peculiarly the case, and it is only by profoundly altering the conditions of life that we are able, as a general rule, to bring about an arrest of the process, without the actual removal of the morbid agent. It may, perhaps, be hoped, that in the case of infective diseases, medicinal agents may be discovered which will directly attack the morbid agent, but except in the case of syphilis and malarial fevers, such a discovery has not yet been made.

The tendency to recovery is much less in diseases due to causes in the tissues themselves. Malformations may be partially rectified during the processes of development and growth, but they will rarely be completely recovered from. Tumours generally continue their growth without reference to the activities of the tissues. The diseases due to decadence of the tissues are seldom the subjects of spontaneous recovery.

**Death** is the unfortunate termination of many diseases, but the tendency to death is frequently more an accidental circumstance than a necessary part of the phenomena of the disease.

Life may persist along with the abolition of many of the functions of the body. If the respiration and circulation persist, the remaining functions, even those of the brain, may be in abeyance and yet the person may survive. If we look at the actual process of dying we shall find that in most cases the heart or the respiratory movements first give way, although in some instances it may be difficult to determine which has first ceased. According to Bichat there are three modes of death, namely, by the brain, by the lungs, and by the heart, but from what has been noted above it is clear that these may be included in the two mentioned.

**Death by the lungs** will be due to failure in the respiratory movements. This may occur suddenly by paralysis of the respiratory centre in the medulla oblongata, as in hæmorrhages in the brain, but for the most part it occurs gradually from exhaustion of the respiratory centre, and is then called death by **Asphyxia**. Such exhaustion will ensue when, from obstruction of the air passages or vitiation of the air, the respiration has been for a time carried on with great difficulty.

**Death by failure of the heart's action** is probably a more frequent mode of death. It may be the result of irritation of the vagus centre in the medulla oblongata from injury to the brain, though in this case the failure of the heart may be simultaneous with cessation of respiration. This form of paralysis of the heart, however, is of rare occurrence. Again, a sudden reflex paralysis of the heart from shock may occur. There may also be death from failure of the heart due to



disease in the organ itself, and here also the death is very often sudden. A frequent cause of sudden death, for instance, is obstruction of a branch of the coronary artery. This may lead to such derangement of the nutrition of the heart's muscle as to cause paralysis and cessation of its contractions.

But cessation of the heart's action is often the mode of death when the seat of disease is distant from that organ or its nervous apparatus. In that case the heart is affected secondarily. It is weakened it may be by the condition of the blood being altered, so that the requisite amount of nutriment is no longer afforded to the heart; or it may be injured by the blood being increased in temperature or contaminated by abnormal products. In these cases the heart partakes in the general weakening of the body, and gradually ceases to contract.

Failure of the heart has an immediate effect on the circulation in the lungs, and we shall see afterwards that oedema of the lungs is one of the most constant effects of this condition. (Edema of the lungs, on the other hand, with exudation of fluid into the lung alveoli, which passes on into the air passages, obstructs the respiration. The laboured breathing and rattle in the throat, which are so frequent in the last stages of disease, are the usual signs of oedema of the lungs, so that although these signs call attention to the respiratory organs, the real primary failure may be in the heart.

It will be seen that the proclivity to death in the various forms of disease will depend on the degree to which the disease affects the respiration or the heart's action, but more particularly the latter. In this view of it the actual question of death or survival will frequently depend on the staying power of the heart. The ability of the heart to continue its contractions during the most severe period of an illness will frequently determine whether the person is to die or recover. This is an important point to keep in mind in actual practice. There are great differences in the staying power of the heart in different persons, and so there are great differences in the fatality of the same diseases. The ability of the heart, like other local conditions, is largely determined by inheritance. It is probably one of the main factors in determining the duration of life. Longevity depends probably more on the character of the heart than on any other single factor.

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## SECTION II.

## TERATOLOGY.

## GENERAL MALFORMATIONS. MONSTROSITIES.

INTRODUCTION.—*Definition of terms and grouping. Causation of monstrosities.*

I. MONSTROSITIES BY EXCESS, in size or number. A. *By excess in size, giants, local hypertrophies.* B. *By excess in number—Fission of embryo.* I. *Complete fission, twins, with subsequent partial union* (1) *of xiphoid, (2) of thorax, (3) of thorax and head, (4) of crania, (5) of pelvis.* II. *Abcaudal fission:* (1) *head single, and bodies equal; (2) head single, one body undeveloped and parasitic; (3) fission of pelvis and legs.* III. *Abranal fission:* (1) *partial fission of head—double-face; (2) complete division of head—double-head; (3) one division undeveloped and parasitic; (4) supernumerary arms.* IV. *Fission both abcaudal and abcranial.* V. *Double fission, triple monsters.* VI. *Reduplication of parts.* II. MONSTROSITIES BY DEFECT in size or formation. A. *Defect in size—Dwarfs.* B. *Defects in formation.* I. *From imperfect closure of cerebro-spinal canal:* (1) *Brain absent, Anencephalus; (2) brain displaced—Encephalocele, Hernia cerebri; (3) Brain defective—Cyclopia; (4) vertebral canal and cord imperfect—Spina bifida.* II. *Imperfect closure of visceral arches:* (1) *facial clefts; (2) congenital fistula of neck; (3) defective closure of thorax; (4) of abdomen.* III. *Defect of orifices and canals.* IV. *Absence or defect of extremities.* III. ABERRANT MONSTROSITIES, chiefly transposition of viscera.

## INTRODUCTION.

THE conditions to be considered here are all referrible to errors in the development of the embryo. They are to be traced to causes acting on the embryo and causing it to deviate in its formation either as a whole or in part from the normal. Malformations may consist merely in slight local deviations from the regular type, as where a muscle or a blood-vessel has abnormal relations, in which case the term **Anomaly** is frequently used. On the other hand, they may affect the body as a whole and may be such as to produce the most serious deformities, many of them quite incompatible with life. In that case the term **Monstrosity** is frequently used. It is with these latter that we are chiefly concerned here. The malformations of individual organs will be referred to in their places, but in this general section we have



chiefly to consider those which affect the body as a whole or the more important parts of it. The science of monstrosities or **Teratology** is a very wide one : it will only be possible here to give a general outline of it.

The congenital malformations form only a part of the congenital diseases (see p. 15). In some of the latter we have simply the ordinary pathological changes such as are met with in extra-uterine life, but they have happened to occur in the fœtus. In the case of malformations, on the other hand, the structure of the tissues in themselves is normal, but there is an erroneous arrangement owing to some interference occurring during the period of development.

The malformations are divisible into three great groups. These are malformations by excess, by defect, and by peculiarity of form.

**Malformations by excess** are divisible according as the size or the number of parts is in excess. There may be a general excessive size of the whole body or of some of its parts, but such gigantic formation is uncommon. Much more common is numerical excess, consisting in a doubling or even in a trebling of the body as a whole or of its parts. A large group of malformations is included under the name of **double monsters**, in which a considerable portion of the body is doubled, or there are two complete bodies joined together. **Malformations by defect** occur either in the form of actual deficiency or smallness of parts, or else of hindered or checked development. In the latter case the parts may be of full size, but they represent some period of embryonic life and not that of full development ; thus, structures which are formed separately in the embryo in order afterwards to unite and form single structures may fail to do so, and clefts or fissures may be the result. **Malformations by peculiarity of form**, or Aberrant malformations, are local anomalies in which without any actual defect the development has taken a different course from the normal, structures which usually atrophy as development proceeds perhaps continuing to grow, while those which usually develop completely are atrophied. Many malformations of the heart and vessels and of the generative organs belong to this class, which also includes many local anomalies. These latter do not fall to be described in this section.

**Causation of Malformations.**—As the malformations take origin in the embryo at comparatively early stages of its development, it must be generally difficult to trace the exact nature of the cause, but there are at least some indications of the kinds of causative agents.

In the case of the malformations by excess we must suppose an excessive stimulation of the embryo at an early period of development, but the nature of the stimulation is difficult to determine. It may here be pointed out, that in certain of the lower animals which are capable of



restoring lost parts there is sometimes a reduplication of the parts reproduced. Thus in the case of the lizard, when the tail is broken off, there may be two or three tails instead of one reproduced. In the salamander also, when the foot or hand is amputated or divided longitudinally, there may be a new formation of one or more supernumerary fingers. In these cases the stimulation of the wound, in which the germinal tissue is forming a new member, results in the reduplication of the member. In a similar fashion the over-stimulation of the embryo may induce it to a reduplication, which will affect the whole body or parts of it, according to the period and locality of its application. The stimulus may in some cases be related to the activity of the ovum and spermatozoa respectively, but there is reason to believe that external stimulation may in some cases have important influences. Thus Gerlach asserts that by varnishing an egg so as to leave only a certain selected part free for the penetration of air he succeeded in certain cases in producing a doubling of the anterior extremity of the chick. Panum, Dareste, and others have also produced malformations by varnishing, by variations of temperature, and by placing the eggs vertically.

In the causation of malformations by defect, it may be that deficiency in the ordinary stimulus inherent in the ovum takes part in the production of some forms, but many of them can be traced to disease of the embryo or its coverings. Thus important malformations result from accumulations of fluid or dropsies in the cavities of the embryo, more especially the cerebro-spinal canal. This leads to defects and other changes in the central nervous system and its bony case. Abnormal adhesions, usually produced by inflammations, also produce malformations, partly by holding back the expanding structures, partly by causing obstruction of orifices and otherwise. There are also strangulations and even amputations of parts by the umbilical cord or by threads accidentally formed. In these various forms of disease the starting point may be injuries to the foetus, sometimes from blows on the uterus or violent movements on the part of the mother.

**Inheritance** has little influence in the causation of malformations. In the case of the slighter forms or anomalies, such as supernumerary digits, inheritance has a most marked influence, but in the monstrosities proper it has little or no effect. There may possibly be a hereditary excess or defect in the inherent stimulus of the embryo, and the ultimate size of the individual is determined by its inherited endowments. The more considerable malformations, however, seem to be more dependent on accidental or local circumstances.

**Sudden frights** or shocks are often assigned popularly as the causes of malformations. The sight of a person with harelip or with an



amputated limb, by the mother, is supposed to give rise to a like malformation in the foetus, or even a simple fright is stated as the cause. There is, however, little basis for this opinion, and the assigned cause often happens at a period of pregnancy when the malformation must have been already present. Most of the malformations, as we have seen, are referrible to the early periods when perhaps the mother is as yet ignorant of the existence of pregnancy.

**Classification.**—The malformations form a regular series, and subject themselves to a classification on comparatively simple lines. Most of the systems are based on that laid down by Foerster, which has been already indicated above. Cleland has suggested a classification which follows somewhat on Foerster's lines, taking more account, however, of the causation of the lesions. This classification will be in general followed.

The author is indebted to Professor Cleland for many valuable suggestions in regard especially to the causation and classification of monstrosities. These suggestions are contained in several monographs on specific examples of malformation (see Literature), and in a volume of *Memoirs and Memoranda*, by Professor Cleland and his assistants. In his classification the author has made great use of a paper in the latter volume in which a suggested classification is tabulated.

## I.—MONSTROSITIES BY EXCESS.

The excess may be either in size or in number, and these again may affect the body as a whole or individual parts.

### A.—MONSTROSITIES BY EXCESS IN SIZE.

1. **General hypertrophy of the body: Giants.**—We include here persons who greatly exceed in height the stature of ordinary men. As a rule a man who measures 7 feet and upwards is counted a giant; there are cases on record from 7 to 8 feet in height, and even up to 9 feet, although many of the accounts are doubtful. The excess in growth is chiefly of the osseous system, and especially in the legs, but the other bodily structures are also in excess. In some cases sexual maturity is delayed or remains absent, and most giants become prematurely old and do not live long.

2. **Local hypertrophies.**—Excessive growth of one half of the body is a rare form of malformation. We have also unilateral hypertrophy of a limb, or of individual fingers or toes. The bones of the skull sometimes grow excessively, especially the lower jaw. There are also



hypertrophies of the larynx and tongue, and more rarely of the internal organs, which as they have no explanation in any acquired disease, are regarded as malformations.

#### B.—MONSTROSITIES BY EXCESS IN NUMBER—FISSION OF THE EMBRYO.

These form a very important group, and include a large proportion of the more considerable malformations of the body. They present great variety in form and degree. At the one end of the scale we have the monstrosity consisting of two complete individuals, united by a narrow band, or indeed disunited. At the other end we have simple duplicity or reduplication of parts.

In all cases of double monstrosity there has been to begin with one ovum. Opinions have been divided on the question whether in the single ovum there have been originally two embryos which have partially united, or whether there has been but one embryo which has given rise to the doubled parts by fission. The latter view is supported by many very forcible facts.

Multiplication by fission is common in the vegetable kingdom, and is not unknown, as we have seen, in animals. In the embryo an excess of formative material is apt to express itself in a more or less complete attempt at fission in the longitudinal direction. At the earliest period of all, when there is over-stimulation of the ovum soon after impregnation, there may be fission of the whole germinal mass, with the ultimate production of two separate individuals or twins. At later periods, when the central part of the cerebro-spinal axis has been already laid down, the fission may still affect the upper or lower end of the embryo, and we may have doubling of these either together or singly, while a portion of the cerebro-spinal axis is single. Then at still later periods when the appendages are being formed, the germinal material may be in excess, and by fission lead to multiplication of parts. There are even cases in which the fission has been repeated more than once, so that there is not only a doubling but a trebling of parts, and the monster contains portions of three individuals.

An interesting confirmation of the view that double monstrosity arises by fission of one embryo and not by union of previously separate embryos has been furnished by Cleland in his observations on supernumerary legs. According to the old theory supernumerary legs are parts of an undeveloped fœtus which has become adherent to, and partly included in that which has attained full development. The developed legs, however, do not belong to one pelvis and the supernumerary ones to another, as the ordinary theory would presuppose. On the contrary, the two developed legs belong to two different pelves, as do also the supernumerary ones. That is to



say, let there be two pelves and four legs, two of which are the proper legs of the individual and two are supernumerary and probably ill-developed, then the developed legs are the right leg of the right pelvis and the left leg of the left pelvis, while the supernumerary ones are the left leg of the right pelvis and the right leg of the left pelvis. The only possible explanation of this is that there has been a partial fission of the hinder part of the embryo, and that of the two pelves and four legs those parts have fully developed which have had most room to expand. As expansion takes place laterally it is the extreme right and left halves which develop most fully, the adjacent left and right halves being more or less crushed, and perhaps even included, forming in some cases no more than a sacral tumour (teratoma).

The doubled parts frequently attain varying degrees of development. The one half may grow much more quickly than the other, and when that is the case the lesser half may undergo various degrees of displacement and defective development. As the well-developed half grows in length it may even tear away parts of the less developed half from their relations, and so produce complex arrangements. The more quickly growing part may even come to include the less developed one, and the latter may form a kind of **Parasite** on the former.

Another law has to be remembered in considering double monstrosities, namely, that by which symmetrical parts tend to adhere. This law has a wide application in normal development, lateral parts adhering and coalescing in the middle line in front. It has also applicability in cases of doubling, the symmetrical parts of one half sometimes adhering to the symmetrical parts of the other. This is an adhesion strictly comparable with that occurring in normal development. There is, however, not infrequently an adhesion and partial coalescence of parts of double monstrosities if they meet in the middle line. Thus supernumerary arms or legs where the doubling is imperfect will often meet in the middle line, and they may coalesce more or less, so that instead of two extra legs or arms there may be but one, but this generally a compound one. This is an adhesion quite distinct from the normal adhesion of symmetrical parts of the embryo.

The large majority of double monstrosities are of the female sex, and the two bodies are always of the same sex.

The double monstrosities are divisible into four groups according to the extent and situation of the fission. The fission is (1) complete, involving the whole cerebro-spinal axis, or (2) of the hinder extremity—abcaudal fission, or (3) of the anterior extremity—abcranial fission, or (4) of both extremities while a portion of the cerebro-spinal axis is undivided.



## I.—COMPLETE FISSION. TWINS.

Most twins are developed from two ova, and are just as distinct as children who are born one at a birth. But some twins are developed from one ovum, are contained in one chorion, and have a common placenta. In that case the two individuals have arisen by complete fission of the original embryo, and the fission has remained permanent. It is consistent with this view that such twins are always of the same sex and closely resemble each other, so closely that their individual identity is often mistaken. In a certain sense, therefore, such twins are examples of double monstrosities.

There are, on the other hand, twins in whom the cerebro-spinal axis is complete in each, there having been originally a complete fission, but union has taken place, so that the two are connected by living tissue. For the most part the two bodies are placed parallel to each other, and with the anterior surfaces turned towards each other, and they are usually united by their anterior parts. The explanation of this may be that, as the anterior parts are the last to close, adhesion is more likely to take place here. This rule is, however, not without exceptions.

Although the cerebro-spinal axis is complete, there is in this form of monstrosity, not infrequently a suppression of certain parts, the suppressed parts being symmetrical, as if, being in contact, portions had been crowded out by the growing structures. There are thus several forms.

1. **Union confined to the neighbourhood of a common umbilicus** (*Xiphopagus*).—This is the least degree of union, and is illustrated by the well-known Siamese twins. There is a common umbilicus and umbilical cord, and there is also a cartilaginous bridge between the two xiphoid processes.

2. **Union from umbilicus upwards, to form a single thoracic cavity** (*Sternopagus*).—In this case as the union is very close some of the adjacent structures may be suppressed or ill-developed. The two intestines may be united at the jejunum, but double above and below. The liver is usually double, but the two livers are generally coalesced. The lungs are always double, and the heart usually so, but the two hearts may have coalesced at their borders and be externally single. There is sometimes a partial or complete coalescence of two of the arms, so that there are only three arms. The coalesced arms will be the left of the right twin, and the right of the left.

3. **Union of thorax and head** (*Prosopo-thoracopagus* or *Syncephalus*).—The spinal column and base of the cranium are separate, but the faces



have come in contact and partly coalesced. There may be two faces, but they are often partially undeveloped. The union here, as in the two preceding classes, is anterior, so that it is the faces which come in contact and coalesce. Symmetrical parts of the two faces may thus unite, especially those in the middle line, as the mouths and noses, while the ears are brought close together. There is in some cases a peculiar coalescence of the two faces, as if while facing one another they had become flattened out against each other and the parts carried to either side. There are thus two faces looking to the right and left, but each face really belongs half to one body and half to the other. This form of Mr. Facing-both-ways is called **Janiceps**. Sometimes one of the faces is only a rudiment. In all these forms the mouth and tongue are single in the posterior part; the oesophagus, stomach, and duodenum are single. The lungs, urinary, and sexual organs are double, but the heart single. The arms are nearly always completely double.

These three forms are usually included in the genus **Thoracopagus**, and they constitute a very common form of double monster. Amongst them the second or **Sternopagus** is the most frequent. In some cases one of the twins is ill developed, and exists as an appendage to the other, forming a parasite attached to the abdomen or thorax. This form is called **Parasitic thoracopagus**, and is illustrated by the case of Lazarus Colloredo, who was born in 1716, and lived to an adult age. The smaller twin had most of the external parts, with the exception that there was only one leg.

It is customary to include under the thoracopagus parasiticus cases in which the head of the ill-developed fœtus is absent (acephalic parasites), but this is more probably due to incomplete or abcaudal fission. (See further on.)

4. **Union of cranial vaults** (*Kraniopagus*).—The twins are complete, and separate except that the cranial vaults have united. The union may be frontal, parietal, or occipital (*Kraniopagus frontalis*, *parietalis*, *occipitalis*).

5. **Union pelvic** (*Ischiopagus*).—This is the converse of the preceding form. The two bodies are united below, and diverge from one another so that the heads are at the opposite poles. The two pelves are united, and sometimes the two sacra in such a way as that the spinal canals are continuous. All the organs of the chest and abdomen are doubled, but one set of the external sexual organs may be imperfectly formed or they may have coalesced. There are generally four legs, which are thrown to the sides, but there may be three or only two.

Ahlfeld divides the cases of complete fission into those in which the two parts are homogeneously developed and those in which they are unequally. Under this latter are included chiefly very ill-formed monstrosities, which occur along with, and frequently as appendages to, fully-developed fœtus. Most of them have the heart either rudimentary or absent (hence **Acardiacus**), and they are usually parasitic on



the more fully-developed fœtus, whose heart carries on the circulation in the parasite. As Cleland has indicated, most of these are really cases of abcaudal or aberanial fission in which various distortions have occurred owing to one half growing more quickly than the other, and they will be mentioned further on.

## II.—ABCAUDAL FISSION.

The fission in this case may be more or less extensive, from below upwards, and it may involve the whole vertebral column, but the head will always be single. One of the halves may develop excessively, and the other may be very imperfect.

1. **Head single, bodies double and equally developed** (*Dipygus*).—There is here one head and two bodies, but the bodies and arms may be to a variable extent coalesced. There may be the regular four arms, or there may be only two; hence dipygus tetrabrachius and dibrachius. This form of double monstrosity is rare in man, but common in animals.

2. **Head single, one body much larger than the other** (*Parasitic forms*).—We have here a great variety of malformations according to the degree of defect of the smaller portion. The fully-developed fœtus is continuous with the head, while the imperfect one seems a mere appendage, or is even broken off from the former. We are able to recognize various modifications.

(a) **Smaller part appended to front of larger** (*Acephalic parasites*).—In this form we have the fully-developed fœtus with part of an ill-developed one hanging from the front of the thorax and abdomen in the form of two legs, or two legs with part of a body and two arms. This form is sometimes called **Epigastrius**. It is the commonest form of the parasitic monstrosities, and is illustrated in the case of a Hindu girl, called Lalloo, who is now about 18 years of age.

(b) **Smaller part included in the other** (*Inclusio foetalis, Fœtus in fœtu*).—The included portion is within the abdominal cavity of the larger part (**Engastrius**), and is very ill developed. It lies in a sac composed of connective tissue, and the parts are rather confusedly mixed, but there are usually recognizable the bones of legs and arms with hands and feet—sometimes also portions of the vertebral column.

(c) **Smaller part completely separated** (*Acephalus. Amorphus*).—In this case the smaller division has got broken off from the larger, and has developed separately, being, however, still dependent for its blood supply on the cardiac contractions of the completed fœtus. As these forms have only a rudimentary heart, or none at all, they are sometimes called *Acardiaci*. The separate monster has its own umbilical cord, whose vessels, instead of passing to the placenta, open into those of the fully-formed fœtus. Hence the circulation in the monster is peculiar



in respect that blood is carried to it by the umbilical artery and leaves it by the umbilical vein, and that it receives blood which has already passed through the body of the well-formed fœtus.

The *acephalus* presents various degrees of development of the body and limbs. There may be only pelvis and legs, or there may be part of the vertebral column, and even thorax and arms. The legs and arms, if present, are often coalesced. There may even be some trace of cranial bones. The lungs and heart are always absent.

The *amorphus* is a rounded mass covered with skin, and showing externally no indication of parts of the body. It contains internally fat and connective tissue, with a rudimentary vertebral column. Sometimes there is more of an approach to human form, and the mass contains distinguishable rudimentary parts of the body.

3. **Fission affecting pelvis and appendages** (*Supernumerary legs, sacral teratoma*).—There is in this case primarily a fission which affects the inferior part of the embryo; the primordial pelvis, and perhaps also the lower part of the vertebral column are divided. If the latter is at all divided, then one half disappears, and besides the one fully developed trunk there is ultimately nothing but the two pelves with their appendages.

From the observations of Cleland, it appears that the two halves present peculiar and interesting relations. The fission is in the middle line, so that a right and a left pelvis result. As the two pelves are closely connected behind (by the spinal column), there is much more room for expansion in front, and so while the proximate posterior parts are crushed together and liable to imperfect development, the anterior halves attain full development. The result is two perfect limbs, which, however, belong properly to two different pelves, being the right leg of the right pelvis and the left leg of the left, and in addition a more or less imperfectly formed composite pelvis with supernumerary legs, variously coalesced or dwarfed, behind.

This subject has been worked out by Cleland in his memoir on "Birds with supernumerary legs," etc. He there refers to cases in the human subject, especially to that of Dos Santos, a monstrosity which has been described by several writers. In this case there was a supernumerary leg attached posteriorly. It had the knee turned backwards, and ended in a composite foot with the great toes joined and the fifth toes at the outer sides. This composite limb, therefore, had not arisen by the fission of two limbs of the same pelvis, but by the coalescence of the left leg of a right pelvis and the right leg of a left pelvis. In front there were two penes between the fully-developed limbs, another indication that the anterior parts and the two developed limbs belonged to different pelves.

The condition of the supernumerary parts behind varies considerably.



There is usually one coalesced limb, like that mentioned above; but there may be two, or the indications of a second.

On the other hand, the whole ill-developed part may be included in the well-developed parts, and the abnormality appear as a tumour in the sacral region. The **sacral teratoma** is a tumour in which portions of bone, muscle, fat, etc., are found, and they arise by an inclusion such as this. The contained bones may sometimes be recognized as belonging to the pelvis or lower limb.

### III.—ABCRANIAL FISSION.

As in the case of abcaudal fission, the division may be more or less complete, and the lateral halves may be unequally developed, so that in the extreme cases one is parasitic.

1. **Partial fission of head** (*Diprosopus*, *double-face*).—In this there are indications of fission in the middle line, but the corresponding parts have coalesced more or less, so that while the head is single, the parts of the face are at least partly doubled. In the lowest degree there is apparent externally only a broadening of the head, and there is no doubling of external parts, although the mouth and nasal cavities show a certain amount of duplicity. From this there are successive degrees. the double parts, as it were, emerging as successive stages are reached. As the two heads are applied laterally the eyes and nose first emerge, and then the ears. Hence, beginning with the lowest, we have cases with two eyes (*Diprosopus diophthalmus*), with three eyes (*triophthalmus*), with four eyes (*tetrophthalmus*), with three ears (*triotus*), and with four ears (*tetrotus*).

This form of monstrosity is rare, and as the brain and fauces are usually defective, it is not capable of life.

2. **Complete division of head** (*Dicephalus*, *double-head*).—In this case also we have degrees of fission varying from cases of two heads on one neck to those in which the body is also to a large extent doubled. The condition of the arms indicates approximately the extent of division of the spinal column; hence we have as varieties, in different stages, cases with two arms (*Dicephalus dibrachius*), with three arms (*tribrachius*), and with four arms (*tetrabrachius*). There are even cases in which an additional lower limb has been present in a rudimentary state (*Dicephalus tripus*), the fission having extended almost throughout.

These forms are very common, forming indeed the most frequent double monsters. The malformation causes great difficulty in parturition, but, if safely delivered, the monster is quite capable of life, and may even attain to old age.

3. **One division much larger than the other** (*Parasitic forms*).—As in



the case of abcaudal fission, so here one of the halves may develop fully, while the other is dwarfed, and remains as an appendage or parasite. It may be considerably displaced or even separated from its connection with the fully developed portion.

(a) **Smaller part appended in front of larger.**—This form is much rarer than the corresponding one in abcaudal fission. There is a head and part of the body appended to the thorax and abdomen of a fully developed foetus (*Epigastrius*), but the parasitic portion is dwarfed. The monstrosity is quite consistent with life.

(b) **Smaller part included in the other.**—To what extent the occasional occurrence of teratoma inside the skull or in the mediastinum may be due to aberanial fission with imperfect development of one half, is not known.

(c) **Smaller part completely separated** (*Acormus*).—In this, as in the corresponding form of abcaudal fission, the separated part is connected by the umbilical vessels with a fully developed foetus, from which it receives its blood. The separated portion consists of a head either with no spinal column or a short piece. The head itself is ill formed, the parts frequently much altered. The umbilical cord proceeds from the neck. There is, of course, no heart, as there is no trunk, and the monstrosity is included amongst the *acardiæ*.

4. **Supernumerary arms.**—As in the case of supernumerary legs in abcaudal fission, so by a somewhat similar cause we may have supernumerary arms produced in aberanial fission. In this case the spinal column is single, one of the original divisions having disappeared, but the primordial limbs of both divisions remain. As in the former case also, the outer limbs of each division develop into the proper limbs of the animal, while the internal or proximate ones become dwarfed appendages. The developed arms are therefore the right arm of the right division and the left arm of the left, while the appended ones are the left of the right and the right of the left. (Cleland.)

#### IV.—FISSION SIMULTANEOUSLY ABCRANIAL AND ABCAUDAL.

This will present many of the features of complete fission, but the spinal column will be in part at least undivided. There will be two heads or the indication of such a division, and four legs or indications of them. The "Two-headed Nightingale" is a monstrosity of this kind.

#### V.—DOUBLE FISSION. TRIPLE MONSTERS.

It is not very uncommon to have triplets in one chorion, arising by double fission of one embryo. It is also not very rare to meet with a



partially divided foetus along with a fully developed one. On the other hand, a double fission in a foetus which remains united is excessively rare. In one authentic case there were three heads (**Tricephalus**), two of which were on a single vertebral column, the cervical vertebrae being alone divided, while the other had a separate column to itself. There had been a nearly complete abcranial fission followed by a partial abcranial fission of one of the divisions.

#### VI.—REDUPLICATION OF PARTS.

The tendency to fission does not confine itself to the cerebro-spinal axis, but extends to the individual parts of the body, more especially to those accessory parts which as it were bud out from the main body.

**Polydactylism** is the reduplication of the fingers and toes, a somewhat common malformation. The lowest degree is that in which a small appendage is attached by a narrow neck to the outer aspect of the hand or foot. This may or may not have a bony phalanx. In a higher degree the finger has an additional metacarpal bone, and even an added carpal one. In the highest degree the hand or foot has nine or ten fingers or toes. The multiplication may affect one hand or one foot, or both hands or both feet, or all the four members simultaneously. When individual fingers or toes are divided it is most frequently the little one; next to that the thumb or great toe, and very seldom one of the intermediate ones.

These malformations are remarkably subject to hereditary transmission. It is also remarkable that supernumerary digits when amputated are liable to grow again. (See Darwin, "Animals and Plants under Domestication," vol. ii., p. 14.)

**Multiplication of bones and muscles.**—It is not uncommon to find additional vertebrae in the various regions of the column, and they are sometimes rudimentary. Sometimes there is a prolongation of the column so as to produce a **Tail**. There is, however, sometimes no bone in such caudal appendages, the vertebrae being only prolonged as membrane, and sometimes the appendage is not a proper tail at all, but merely a cutaneous projection.

The ribs are somewhat frequently reduplicated, so that we may have **cervical or lumbar ribs**.

Reduplication of muscles is frequent, and some anomalies of this kind are so common as to be regarded as mere varieties.

**Supernumerary mammae** are perhaps doubtfully to be regarded as instances of reduplication of parts, as they may be rather due to reversion. There are cases of three, four, and five mammae. The extra mammae are usually near the proper ones, and generally under them,



but there are cases of very considerable removal, as in the inguinal region, or on the back. The mammae may be represented only by nipples.

**Internal organs** are sometimes reduplicated, most commonly the spleen, but also the pancreas and other organs.

## II.—MONSTROSITIES BY DEFECT.

It will be understood that we have here to do with the more general defects, and that the more distinctly local ones fall under their respective special sections. In studying the malformations by defect we have to go back to the developing embryo, and to see, in interferences with the expansion and development of parts, the causes of the defects.

### A.—DEFECT IN SIZE, DWARFING.

Defect in size of the body as a whole results in the production of a dwarf. **Dwarfs** may be well formed although diminutive in size. Generally, however, the head is disproportionately large, and sometimes it is so excessive in comparison with the body as to give the body as a whole a deformed appearance. The body is also sometimes deformed, and the extremities crooked. Dwarfs have frequently good health, and may live to a considerable age. The usual height of dwarfs in the adult state is about  $2\frac{1}{2}$  feet, never under 2 feet, and sometimes as much as  $3\frac{1}{2}$  or  $3\frac{3}{4}$  feet.

**Partial dwarfing** occurs in various parts of the body, but especially in the extremities. The extremities may be well formed, but defective in size. This may affect all four limbs (*micromelus*), or the arms (*microbrachius*), or the legs (*micropus*). The brain may be defective in size as a whole (in idiots and cretins) or in parts, such as the cerebellum. The face is sometimes at birth dwarfed on one side (*Hemiatrophy*), this apparently depending on some affection of the brain. The intestine may be abnormally short.

### B.—DEFECTS IN THE FORMATION OF PARTS.

These vary considerably in form and causation, and it is difficult to divide them into perfectly definite groups. Although the causation is obscure in many cases, yet the constancy of the resulting forms shows that common influences have been at work.



## I.—DEFECTS FROM IMPERFECT CLOSURE OR DROPSY OF THE CEREBRO-SPINAL CANAL.

In the early embryo the cerebro-spinal axis is represented by the medullary plates, beneath or in front of which is the notochord. The medullary plates, rising up at the sides and folding round dorsally, come in contact and form the medullary canal which is the representative of the ventricles of the brain and the central canal of the spinal cord. It has been shown by Lebedeff and Cleland that this closure of the medullary canal dorsally may fail to occur, and it has been ascribed to an over-growth of the medullary plates causing them to flatten out and go into folds or curvatures, while the notochord prevents proper longitudinal expansion. On the other hand, after the closure of the medullary canal there may be, perhaps from over-stimulation, an accumulation of fluid or dropsy, and this again may lead to a secondary rupture of the canal or else to a bulging of it backwards. We have therefore these three possibilities, non-closure of the canal, its rupture by dropsy, and over-distension with bulging. According to Cleland the first of these, occurring at a very early period, is likely to produce such disturbance in development as to cause the death of the embryo, and in consequence the more permanent malformations are due to accumulations of fluid producing rupture or distension.

The condition of the parts formed by the epiblast and mesoblast will depend on whether there has been rupture or not. When there is rupture the bone and cutaneous tissue which normally cover the dorsal aspect over the cerebro-spinal axis will be defective, and the latter will be from the first **open** or exposed. On the other hand, if there is mere distension of the medullary canal, the cutaneous structures may be continued over it, forming a tumour, but the bony structures are likely to be pressed aside by the bulging tumour and imperfectly formed.

The changes mentioned may affect chiefly certain parts of the medullary canal, and may occur at different periods of development, the ultimate result varying accordingly. Thus the part affected may be the cerebral ventricles either in whole or in part, or it may be the central canal of the cord, or it may be both of them in various degrees. Certain well-marked forms are described.

1. **Anencephalus**.—This name implies that the brain is absent. It arises by an early rupture of the medullary canal or, according to Lebedeff, from a non-closure of it.

The child is often born at the full time, and the trunk and limbs are usually well developed. But the vault of the cranium is absent, and the base of the skull exposed. The base is occupied by some loose



membrane in which there may be some cysts. Occasionally there is a vestige of brain, with perhaps a small cavity communicating with the surface, in which case the early rupture has been limited and has allowed of some development of the brain. The membrane at the base represents arachnoid and pia mater turned aside, and with the ventricles exposed. It may be in some cases that there is a bulky sac occupying the place of the brain, but this is of doubtful occurrence. If it did occur it would represent the distended but unruptured ventricles. The absence of the cranial vault renders the eyes unduly prominent as they project at the edge of the open skull, giving the head the appearance of a toad, from which the malformation is sometimes called popularly *Toad's head*.

The cranium shows almost complete absence of the flat bones of the vault, a shortening of the base and an angular curvature between the sphenoid and occipital bones. In view of the open state of the bones anencephalus is sometimes designated **Cranioschisis** or **Acrania**. The malformation is often combined with a similar lesion of the spine (*spina bifida*).

2. **Encephalocele, Hernia cerebri**.—In this case the brain or a portion of it projects outside the skull. There may be distension of the ventricles in the extruded part (*Hydrencephalocele*) or there may not. It arises by dropsy of the ventricles without rupture.

In the more extreme cases the appearance of the skull is at first sight like that of anencephalus, but instead of the cranial bones being absent they are flattened down so as to form a very diminutive cranial cavity, while the brain, sometimes nearly of full size, lies outside, communicating with the interior through an aperture in the bones. In this case there has been at an early period a partial dropsy, which has caused displacement of the brain, and there is often evidence of this in the presence of two or three vesicles in the diminutive cranial cavity. Cleland has shown that these vesicles sometimes represent dropsical olfactory lobes with infundibula, which have pushed backwards the cerebral hemispheres, the latter at the period concerned being of small size. In other cases the extruded brain is hydrocephalic, and sometimes there is little more than a sac containing fluid (sometimes called *Meningocele*).

The position of the protrusion is most commonly behind the occiput or at the root of the nose.

3. **Cyclopia** (*Monophthalmia*).—In this condition there is a single orbital cavity in the middle line, containing, in some cases, only a rudiment of the eyeball, in others a fully-developed globe, or it may be two, close together. There is no retina in the eyeball, which consists



alone of parts developed from without. The cerebrum consists of one mesial portion containing a single ventricle.

According to Cleland there is here a dropsy of the roof of the thalamencephalon, including the pineal body. The enlargement of these parts causes defect of the anterior cerebral vesicle, and by its pressure also interferes with the development of the face. There may be, from a similar cause, a still further defect of the face, such as the condition named **Agnathia**, in which the lower jaw is deficient and the ears are brought close together so as to touch one another. This may be associated with Cyclopia, in which case the brain is defective as in that form of lesion, although otherwise the upper part of the face and the brain are unaffected.

4. **Spina bifida**.—In this condition the spines of the vertebræ are usually more or less incomplete, and there may be a tumour projecting whose internal cavity communicates with the spinal canal. But this tumour is absent in certain forms, and it is possible to divide the cases into those in which the vertebral canal is open without tumour, and those in which there is a tumour.

(a) **Open spina bifida** (*Rhachischisis*).—This form is strictly comparable with anencephalus, with which it is often associated. The medullary canal has never been completed or an early rupture has occurred. Accordingly the integuments have not been carried to the middle line behind, and the arches of the vertebræ are wanting, so that the vertebral canal is exposed, covered only with a membrane. This membrane is continuous laterally with the skin. The exposed canal does not, however, appear as a gutter; it is flattened out and shows at most a slight groove, but is frequently even convex posteriorly. The surface of the membrane represents the internal surface of the medullary canal, that is to say, the central canal of the spinal cord, and the cord itself, to the extent of the lesion of the spine, is absent or present as a mere trace, like the brain in anencephalus. In a case described by Cleland the membranous surface was continuous with a dilated central canal, thus proving that the former is really the open central canal. Although the cord is absent, the spinal nerves are present, arising from the membrane in an inner and an outer series, representing the anterior and posterior roots.

Rhachischisis is often associated with anencephalus, in which case it affects the whole length of the column or its upper part, the condition being inconsistent with life. On the other hand, it may affect a limited area usually at the lower part of the column, being produced by a local rupture or defect of the medullary canal. In that case we may have a persistent dilatation of the central canal of the cord, and perhaps a hydrocephalus.



(b) **Spina bifida with tumour.**—In this case there is a persistent dropsy with protrusion of portion of the sac, a condition comparable with hydrencephalocoele. The dropsy may be of the central canal of the cord (*syringocele*) or merely of the meninges (*meningocele*), but the cord is usually carried outwards with the protrusion (*myelomeningocele*). In the syringocele the central canal is expanded so as to form the internal lining of the sac, and the spinal cord may be atrophied or partially preserved. The nerve roots arise in front of the sac. In the meningocele and myelomeningocele the fluid is in the subarachnoid or arachnoid space, usually the latter, and, according to its seat in relation to the circumference of the cord, will be the condition of the cord itself and the nerve roots. The nerves may lie in front of the sac, or may arise within it and course in its walls.

Spina bifida with tumour may occur in any part of the spine but is most common in the lumbo-sacral region. (See under Affections of the Nervous System.)

## II.—DEFECTIVE COALESCENCE OR CLOSURE OF PARTS IN FRONT.

In the completion of the parts in front, the visceral arches grow forward, and, for the most part, coalesce in the middle line in a fashion similar to that of the neural arches on the dorsal surface. The face and neck are partly formed by the subcranial and branchial arches, which variously unite with each other, and with the fronto-nasal plate. In all these there are possibilities of non-union, and so we have various forms of clefts.

The incomplete closure may arise, as in the conditions considered above, by dropsical accumulations, especially in the case of the thorax and abdomen. It may be, also, that protrusion of viscera, from various causes, may prevent closure. But there are many cases in which no mechanical explanation is apparent, and the defect has no obvious cause.

1. **Facial clefts.**—The most extreme case is that in which the fronto-nasal and the superior maxillary plates are defective, and the face presents in its middle part a large opening, which to a greater or less extent replaces nose and upper jaw, and in some cases also involves orbits and lower jaw. This condition is called **Aprosopus** or **Schistoprosopus** ( $\pi\rho\acute{o}\sigma\omega\pi\omicron\nu$  = a face).

Less degrees of it are shown in **Cleft palate** and **Harelip**. The naso-frontal plate forms the central part of the upper lip and of the alveolar process of the upper jaw. Hence, in these parts, the line of union is on either side of the middle line, while in the palate it is mesial. The cleft in the lip and alveolar process is therefore lateral and that in the palate mesial. All degrees of non-union exist, from complete fissure



of the palate, with cleft of the alveolus and lip on each side, to the slightest notch on one side of the upper lip. These defects date back to the third month of foetal life, at which period the closure ought to occur.

2. **Congenital fistula in the neck** (*Fistula colli congenita*).—This arises by imperfect closure of the lateral aspects of the branchial arches. There is a small aperture, only sufficient to admit a small probe or bristle, usually situated half an inch to an inch above the sterno-clavicular articulation, more rarely further upwards and outwards, or in the middle line. The aperture leads into a canal which is directed towards the larynx, trachea, or pharynx, but does not usually communicate with any of these canals. The fistula is lined with mucous membrane, and mucus may be discharged at the orifice. There is usually one fistula which is most frequently on the right side or in the middle, but occasionally there are two symmetrically placed.

Occasionally, from a similar origin, we have **Cysts in the neck** without external aperture. These may be filled with serous fluid (*Congenital hydrocele of the neck*), or may have a more epidermic structure and contents.

3. **Defective closure of thorax** (*Fissura sterni*).—There are various degrees of this, and various combinations with defect of the anterior abdominal wall, the highest degree being a complete cleft of thorax and abdomen. In these extensive clefts the organs are generally extruded, but even when there is a limited cleft of the sternum, the heart is liable to lie outside the chest. In this **ectopia cordis** the heart generally lies in the middle line, communicating with the inside of the chest by a narrow peduncle composed of the great vessels. The heart itself is usually malformed. Sometimes the heart lies at the root of the neck, the manubrium being cleft. Sometimes ectopia cordis occurs without any cleft in the sternum, in which case the heart lies in the neck or in the epigastrium, in the latter case projecting through an aperture in the diaphragm. These facts indicate a displacement by pressure from behind, and probably the whole phenomena are due to dropsy of the pleural cavity at an early period.

4. **Defective closure of abdomen** (*Fissura abdominalis*).—As noted above, this is frequently associated with defect of the thorax. In complete fissure of thorax and abdomen, as there is no umbilicus, there is usually no proper umbilical cord, and the vessels pass directly from the placenta by the amnion to the cleft in the abdomen. The proper abdominal fissure, in its highest degree, extends from the ensiform cartilage to the pubes. In that case the viscera are extruded, the urinary bladder is cleft, and the genital organs are absent or defective.



Sometimes the defect is less, and the abdominal contents lie outside the abdomen in a large hernial sac. The umbilical cord is usually absent, and the placenta is directly in contact with the sac, the vessels traversing the latter to reach the abdomen. Two special forms merit more particular attention.

(a) **Hernia of the umbilical cord** (*Hernia funiculi umbilicalis*).—This is really a minor degree of the condition last mentioned. There is, at the seat of the umbilicus, a rounded tumour, from the distal extremity of which the cord passes off, there being no proper umbilicus. The tumour consists of a peritoneal pouch which is protruded through the abdominal wall. There is an aperture in the wall of the abdomen, and the peritoneal pouch is covered with amnion, which is continuous on the one hand with the surface of the abdomen, and, on the other, with the surface of the umbilical cord. The hernia is one, therefore, of the umbilical cord, and not simply of the umbilicus. When the cord drops off, the amnion goes with it and the peritoneal sac is exposed. If the tumour be large it will become gangrenous, and the patient will rapidly succumb. Even if the tumour be small the exposure generally results in fatal peritonitis, and it is only exceptionally that the gap in the abdomen is closed by healing.

(b) **Fissure of the bladder**.—(*Inversio s. extroversio vesicæ*). In this condition the abdominal wall between umbilicus and pubes is incomplete. The allantois has failed to close completely inside the abdomen so as to form the urinary bladder, and the lateral borders of the latter are adherent to the sides of the cleft, while the anterior wall is entirely absent. The posterior wall of the bladder therefore fills the cleft, and as it is pushed forward by the abdominal viscera it protrudes as a soft red area of mucous membrane, on the surface of which the ureters open, and which readily bleeds. At the upper end of the cleft is the umbilicus which is frequently displaced downwards.

The defect generally extends to the parts below. The pubic bones do not meet in the middle line, the rami ending in rounded lateral prominences. In the female the urethra is usually absent as well as the clitoris, and sometimes the vagina is defective or absent. In the male there is a rudimentary penis, which is not traversed by a urethra, but presents on its upper surface a deep groove which represents the cleft urethra.

Minor degrees of the defect occur. The cleft may confine itself to the upper part of the bladder, which presents itself as a red protrusion just beneath the umbilicus, the parts below being perfect. A still less degree is a simple **persistence of the urachus**, which forms an open communication between the bladder and the surface at the umbilicus.



On the other hand it may be the lower part which is defective, the urinary bladder being well formed, but the urethra cleft and defective. In the male this constitutes **Epispadias**. The penis is short and cleft on its dorsum, the urethra forming here a deep groove. The groove ends posteriorly in an aperture which communicates with the bladder beneath the pubes.

The causation of these malformations is obscure. It may be that there is a simple failure in the formative material of these parts, or that an over-distension of the allantois has prevented a proper closure.

### III.—ABNORMAL CLOSURE OR DEFECT OF ORIFICES AND CANALS.

Some of these are local and fall under their special sections, but there is one form which has more general results.

**Imperforate anus, Persistence of cloaca.**—Up to the fifth week of embryonic life there are no external openings for the intestine and urino-genital organs. The rectum is still closed below, but communicates with the allantois, which forms a common opening for the intestinal, urinary and genital passages, and is itself still open through the imperfect abdominal wall in front. This condition may persist in a greater or less degree.

The most extreme form is where, along with imperforate anus, there is fissure of the abdomen and bladder, a combination of the conditions described in last section with that now under consideration. In some cases the colon is deficient, and the ileum may communicate with the extroverted bladder. Such conditions are hardly compatible with life.

In the simpler forms there is imperforate anus and the intestine communicates with urinary or genital passages. The communication may be with the vagina (*anus vaginalis*), or with the urethra (*anus urethralis*), or with the urinary bladder (*anus vesicalis*). The communications may be very small, so that continuance of life is impossible from accumulation of feces, but there may be fuller communication, and sometimes the condition is surgically remediable.

**Simple imperforate anus**, may be a simple absence of the aperture, but in many cases there is also a defect of a portion of the rectum. The latter condition may exist while the anus is perfectly formed.

### IV.—ABSENCE OR DEFECT OF THE EXTREMITIES.

Defective formation of the extremities may be due to a failure in the inherent powers of the germinal matter which forms these parts, or to mechanical interference. Where there is a symmetrical defect, then we may presume that there has been a failure in the material destined



for the limbs, whereas a non-symmetrical deformity is probably due to a local interference. As the limbs form by lateral expansion, they are more likely than other parts to come in contact with the amnion and the umbilical cord, or with any band or bridge which may have accidentally formed. There may even be an amputation of a limb by the cord or a band of fibrine. Various forms are described.

1. **Defect of all the four limbs.**—(a) *Amelus*, or absence of all the extremities. The body is usually well formed, but at the shoulders and hips there are short rounded or warty projections, at the ends of which there may be horny or nail-like appendages. Sometimes there are ill-formed bones present.

(b) *Peromelus*. The whole four extremities are defective or deformed. The body may be well formed, but sometimes the deformity is associated with other malformations which prevent the persistence of life.

(c) *Phocomelus*. This is a variety of the former, in which the long bones are absent or very defective, and the hands and feet are seated immediately on the shoulders and hips and are well formed.

2. **Defect of the arms.**—(a) *Abrachius*. The upper extremities absent, while the lower are developed.

(b) *Perobrachius*. Both arms are defective. The long bones may fail and the hands be planted on the shoulders, or the arms may be short and deformed, and the hands have only two or three fingers. In most cases, however, the upper arm is little deformed, but the fore-arm and hand are defective.

(c) *Monobrachius*. Absence of one limb. This sometimes occurs as part of a greater defect of the thorax and abdomen, but the body may be fully developed.

3. **Defects of the legs.**—Defects of the lower extremity are similarly classified to those of the upper, and have similar characters. We have *Apus*, *Peropus*, *Monopus*. In the last mentioned the absence of the leg may be associated with defect of the pelvis and protrusion of the abdominal contents. In addition to this we have the following special form.

**Defect and coalescence of lower limbs** (*Sympus*, *Siren*).—This arises by an imperfect development and peculiar distortion of the posterior extremity of the embryo, and it involves not only the limbs, but also the pelvis and its organs. The osseous pelvis is rudimentary, the external organs of generation are seldom present, and the urethra and anus are closed. These conditions render the continuance of life impossible. The lower limbs are united in varying degrees, so that the body is prolonged downwards as a conical process (*Siren*). The legs are revolved so that the anterior surfaces present backwards, and the coales-



cence is by their outer aspects, the external condyles of the femur, the fibulæ and the little toes being the parts united. There are various degrees of union and consequently of defect. The foot may be absent (*Sympus apus*), or there may be one foot at the extremity of the combined leg (*Sympus monopus*), the single foot being sometimes defective, sometimes normal, and sometimes with redundant toes, or there may be two feet, which also may present defects (*Sympus dipus*).

**4. Defects of the fingers and toes.**—These present themselves as *absence* or *defect of the digits*, and as *coalescence*. These are very frequent malformations, and they may affect all four members, or be limited to two or one. In cases of coalescence it is usually only the skin which unites the fingers or toes, but sometimes the muscles and tendons are united, and more rarely the bones.

These defects of the fingers and toes are, like polydactylism, in a high degree inherited. This applies to the symmetrical forms, and not to such defects as may be due to mechanical interference, as by amputation.

### III.—ABERRANT MONSTROSITIES.

In this division we include malformations in which there is little or no quantitative defect, but there is a qualitative difference from the normal, an error in the development; they are the forms which Foerster has classified as *Monstra per fabricam alienam*. The malformations affect the thoracic and abdominal viscera and the external organs which are in immediate connection with them. The heart and blood-vessels and the generative organs are most frequently affected. The malformations of these will be described in their special sections, we have here to deal with general monstrosities.

**Transposition of the viscera** (*Situs transversus*).—This malformation is not very rare. The entire viscera of chest and abdomen are transposed, so that the aorta comes off from a right ventricle, the venæ cavæ are on the left side, the liver is on the left, the spleen on the right and so on. The organs are properly formed and their function is normal. The existence of the malformation may altogether escape observation, or may be accidentally discovered when the person is being examined medically.

Transposition is constant in double monsters, the left twin having the normal arrangement, and the right having the viscera transposed. This seems to indicate that in single embryos the usual situation in the umbilical vesicle is left, but that occasionally it may be right, in which case the viscera are transposed.



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## SECTION III.

AFFECTIONS OF THE CIRCULATION AND DISTRIBUTION  
OF THE BLOOD.

PHYSIOLOGY OF THE CIRCULATION.—*Innervation of Arteries, etc.* LOCAL HYPERÆMIA.—*I. Active hyperæmia ; caused chiefly by dilatation of arteries ; phenomena. II. Passive hyperæmia ; caused chiefly by weakness of heart or venous obstruction. Phenomena, including diapedesis, œdema, etc.* LOCAL ANÆMIA or ISCHÆMIA ; *chiefly from obstruction of arteries.* THROMBOSIS AND EMBOLISM.—*I. Thrombosis. Coagulation of the blood. The process of thrombosis shown by experiment ; characters and forms of thrombi. Causation, from stagnation of blood ; from alteration of wall. Growth of thrombi. (Absence of thrombosis from capillaries.) Changes in thrombi, chiefly softening and organization. Results of thrombosis. II. Embolism, Causation, chiefly by thrombosis. Phenomena, when anastomosis free are trivial ; when anastomosis imperfect (End arteries) frequently the hæmorrhagic infarction ; various occurrence of engorgement of vessels, hæmorrhage and necrosis ; disposal of infarction. Special forms of embolism, malignant tumours, fat, air, and infective.* HÆMORRHAGE ; *by rupture or by diapedesis ; causation various ; stilling of hæmorrhage ; seats of effusion and disposal of the blood.* ŒDEMA AND DROPSY, *depend on the lymphatic circulation. Causation and nature of process ; rarely from lymphatic obstruction ; usually from passive hyperæmia ; hydræmia and hydræmic plethora as causes, especially in Bright's disease ; nervous influences in causation. Position and character of exudation.*

**Physiology of the Circulation.**—In the distribution of the blood the capillaries and veins may be regarded as virtually passive channels. They are able to accommodate more or less blood according to circumstances, but of themselves they have probably little to do with the variations in the supply. The supply of blood to the tissues varies, chiefly according to their needs; the brain or muscle while working requires and gets a fuller and more rapid supply of blood than when at rest. This supply is regulated by the arteries; when they dilate more blood passes into the capillaries and on to the veins, and when they contract less blood reaches the capillaries and veins. We may regard the arteries as regulating the supply of blood, the capillaries as distributing the blood, bringing it within reach of the elements of the tissues, and the veins and lymphatics as carrying it off. The amount of blood



admitted by the arteries depends on the state of contraction or relaxation of the muscular fibre-cells in the middle coat, and this, like all other muscular actions, is under the command of the nervous system. When the vaso-motor nerves are stimulated the muscular coat contracts, when these nerves are paralysed it relaxes, and when a moderate amount of stimulus is supplied, then we have that state of moderate contraction to which the name of **Tonicity** is applied.

**Innervation of Arteries.**—Hunter pointed out that the walls of arteries possess not only elasticity but muscular contractility, and this contractility is due to the existence of the muscular fibre-cells of the middle coat. Constriction of vessels is brought about by stimulation of these cells, and this is effected by stimulation of a special set of nerve-fibres, the *vaso-constrictors*. These have been shown to exist in the sympathetic nerves, and also in the cerebro-spinal nerves of the extremities, where they are perhaps derived from the sympathetic system. Division of constrictor nerve-fibres will induce relaxation of arteries, because it will remove the stimulation which keeps up their tonicity.

Bernard's discovery of the fact that electric stimulation of the chorda tympani causes dilatation of the vessels of the submaxillary gland, proved the existence of *vaso-dilator* fibres, and further observation seems to prove that both kinds of fibres pass to the vessels in all parts of the body. Both kinds probably belong to the sympathetic system.

These facts have suggested that the regulation of the calibre of arteries is effected by the nerves through the intermediate action of local nervous ganglia in or near the walls of the arteries. Arteries may be compared in their innervation as well as in their structure to the heart. The heart has intrinsic ganglia and it has accelerator fibres derived from the sympathetic which stimulate these ganglia, and inhibitory fibres from the vagus which depress them. Nervous ganglia have been discovered in the neighbourhood of very few arteries, chiefly those of the submaxillary gland and the penis, but the nerves are known to form a close plexus around the arteries, and the known facts can only be explained on the assumption of local centres. A further proof of this is to be found in the fact that when arteries are cut off from all central connections they are still capable of varying their calibre. If the sciatic nerve in a frog is divided, the arteries indeed dilate, but after a time they resume, at least partially, their state of tonicity and are capable of still further variations. Again, an excess of carbonic acid in the blood will cause arteries to contract after all their central connections have been severed, and they may be made to dilate by means of chloral or atropine.



The vaso-motor nerves are known to run, to a large extent at least, in the sympathetic system, and they are under the influence of the sympathetic ganglia. The sympathetic system, however, is not an independent one, and the proper vaso-motor centres are in the spinal cord and medulla oblongata, with which the sympathetic has numerous communications. Most of the physiological variations in the calibre of the vessels is brought about by reflex action, and the reflex centres are situated in the cord and medulla. From these centres there seems to pass a continuous slight stimulation inducing that moderate contraction which we name tonicity. If the connection be severed the tonicity ceases, the arteries dilate. There is a general centre for the whole vaso-motor system in the medulla oblongata. When this is stimulated all the arteries appear to contract. When it is destroyed all the arteries in the body dilate. But without being destroyed the centre may be paralysed or inhibited. It is inhibited by the inhalation of nitrite of amyl, and the arteries dilate. It may also be inhibited by the stimulation of a nervous branch which passes upwards from the heart, the depressor nerve. This general centre, therefore, has nervous connections of a similar kind to the local ganglia.

**Literature.**—A very good historical account of the physiology of the vessels is that in VON RECKLINGHAUSEN's *Handbuch der allgemeiner Pathologie*, 1883, p. 4 *et seq.* See also HUNTER, On the blood, inflammation, and gunshot wounds, 1793, Palmer's ed. vol. iii., p. 145; HENLE, *Pathologische Untersuchungen*, 1840; *Handbuch d. rationelle Pathologie*, 1846; BERNARD, several series of *Leçons*, 1858-1876; LISTER (who inferred the existence of peripheral ganglia), *Phil. Trans.* for 1858, vol. cxlviii; WALLER, *Proceedings of Roy. Soc.*, 1862. CYON & LUDWIG, *Ludwig's Arbeiten*, 1866 and 1867; VULPIAN, *Leçons sur l'appar. vasomot.*, 1875; STILLING, *Jenaische Annalen*, 1851. SAVIOTTI, *Virchow's Archiv*, l. 1870; GOLZ, *Virchow's Archiv*, vols. xxvi., xxviii., and xxix.

## LOCAL HYPERÆMIA.

This term is applied to conditions in which the vessels, and especially the capillaries, contain an excess of blood. Two forms are distinguished according as, on the one hand, the blood is too freely admitted by the arteries, or, on the other, meets with some obstacle to its passage by the veins. These are called respectively active or arterial and passive or venous hyperæmia.

### I.—ACTIVE OR ARTERIAL HYPERÆMIA. ACTIVE CONGESTION.

This occurs when an excess of blood is admitted by the arteries into a part. The terms *atonic* and *arterial hyperæmia* are synonymous with active hyperæmia, as are also *fluxion* and *determination of blood*.



**Causation.**—Active hyperæmia is brought about chiefly by causes which induce dilatation of the arteries, although a local increase of blood-pressure, by forcing more blood into the arteries, will have a similar effect.

**Local increase of blood-pressure** is not of frequent occurrence, because as a general rule the vessels are so completely under the regulation of the nervous system that a rapid accommodation is readily effected, by contraction of the arteries. If the arteries be diseased, however, so that their walls are rendered rigid (as by atheroma) then variations in blood-pressure will not be readily compensated. Hence in old people, whose cerebral arteries are atheromatous, temporary increase of blood-pressure may cause congestion of the brain.

**Collateral hyperæmia** might be supposed to be due to a local increase of blood-pressure. It occurs when in consequence of obstruction of an artery or otherwise, the blood supply is diminished in one locality, with the result that there is an increase in the amount of blood in another, generally a neighbouring locality. Doubtless an obstruction in an artery leads to increase of pressure in the arteries proximal to the obstruction, and there may be a resulting hyperæmia of neighbouring parts which will assist in establishing an anastomotic circulation. But the problem is not always so simple as this. The arteries are so completely under the command of the nervous system that the site of the collateral hyperæmia is frequently at a certain distance from that of the anæmia, and is determined not by increase of blood-pressure but by relaxation of arteries. Thus obstruction of one renal artery will induce collateral hyperæmia in the kidney of the opposite side.

**Dilatation of arteries** leads more directly to active hyperæmia in several different ways. It may be due to purely local causes, or to an action on the vaso-motor nerves or the vaso-motor nerve-centres.

As examples of dilatation of arteries from **Local causes** may be cited cases in which the sudden withdrawal of pressure induces local hyperæmia. When the abdomen, for instance, is the seat of a collection of fluid, pressure is exercised on the vessels, and the arteries will relax as completely as possible to allow the blood to overcome the pressure from without. But if the fluid in the abdomen be suddenly removed, the pressure outside the vessels will be greatly reduced and, till the arteries have time to contract, there will be an active hyperæmia. This effect of the sudden removal of pressure is generally, at least in part, obviated by the application of a bandage. On similar principles removal of large ovarian tumours which have pressed on the kidneys and renal arteries sometimes leads to active hyperæmia of the kidneys, which may be accompanied by albuminuria.



Other examples of active hyperæmia due to local causes are afforded by the application of warmth to the surface of the body as by poultices, the result being a direct dilatation of the cutaneous arteries. Mechanical irritation of the skin, as by stroking it, usually induces, first a contraction of the arteries, evidenced by paleness, and then a dilatation, shown by a red streak. Similarly chemical irritants produce dilatation, sometimes preceded by contraction, the dilatation leading to active hyperæmia, which, in this case, may be the first phenomenon of inflammation. Again a mere temporary deprivation of blood may induce dilatation of arteries and a local hyperæmia. Thus in surgical operations when the circulation has been suspended by the application of an elastic bandage, the removal of the latter is often followed by an active hyperæmia.

Active hyperæmia due to causes influencing the **Vaso-motor nerves** may be produced by paralysis of the vaso-constrictors or by irritation of the vaso-dilators. Injury or disease of the sympathetic nerve in the neck is sometimes followed by unilateral congestion of the face, and unilateral sweating together with narrowing of the pupil, smallness of the eyeball and ptosis, these all being signs of paralysis of the sympathetic.

Such cases have been described by Weir Mitchell and others as following gunshot wounds, etc., and by Hutchinson as the consequences of fractures of the clavicle; but some of the manifestations, at least, may ensue from the pressure of tumours, or aneurysms, or the extension of abscesses, tuberculosis, etc., to the neighbourhood of the sympathetic in the neck.

In certain forms of neuralgia, especially in hemicrania, the phenomena indicate first an irritation and then a paralysis of the vaso-constrictors. Du Bois-Reymond by analysis of his own symptoms in such attacks came to this conclusion. In his case there was redness of the external ear; in other cases redness of the retina has been observed. The temperature as tested by the thermometer, in the external meatus, is usually raised, both in the neuralgic cases and those due to injury.

There are comparatively few cases of active hyperæmia which can be directly referred to irritation of the vaso-dilators, but after wounds of nerves, and during the process of healing there sometimes occur severe attacks of pain accompanied by redness and elevation of temperature, which seem only explicable on the supposition that the vaso-dilators are irritated. These phenomena are sometimes followed by trophic disturbances, more particularly the "glossy skin" of Paget.

Paralysis of **Vaso-motor centres** is calculated to induce active hyperæmia if the dilatation of the arteries be localized. Thus in a case observed by the author there was a traumatic lesion of the medulla oblongata which injured the vaso-motor centres of the kidneys. The result was an intense hyperæmia of both kidneys, visible after death, and manifested during life by an excessive secretion of watery urine.



During the few hours of life a large quantity of urine was twice removed by catheter, and after death the bladder was found greatly distended. Some authors ascribe diabetes mellitus to paralysis of the centres in the medulla oblongata, or in the semilunar ganglion. Such a paralysis would induce congestion in the abdominal organs, and more especially in the liver. (See section on Diabetes.)

**Phenomena of active congestion.**—When a local dilatation of the finer arteries of a part occurs, the most direct result is that the blood is admitted more freely, and at an accelerated rate. It meets with less resistance in the arteries, and reaches the capillaries and veins at a higher pressure than normal. The part so affected is of a bright red colour, and, if it be an external part, its temperature is raised. In many cases the active congestion is of short duration, but if it continue it leads commonly to more definite changes. There is increased transudation from the vessels, and consequent swelling. The secretion of the part is increased. This has been observed more particularly in the case of the secretion of sweat; unilateral hyperæmia and sweating have been ascribed to pressure on the sympathetic by an aneurysm (Gairdner) or a tumour (Ogle and Verneuil). There may also be considerable hypertrophy in consequence of active hyperæmia. (See Hypertrophy.)

It has been said that active hyperæmia may lead to hæmorrhage, but experiment seems to prove that even a very great rise in the blood-pressure in the capillaries does not lead to hæmorrhage unless the vessels are badly supported, or else defective in some way. By obstructing the respiration in a dog the arterial pressure may be raised enormously, but there is no rupture of the capillaries, or only in such delicate structures as the retina, brain, or conjunctiva. The increase in pressure in a pure active hyperæmia is, of course, greatly less than this.

**Literature.**—WEIR MITCHELL, MOREHOUSE, and KEEN, Gunshot wounds and other injuries of nerves, 1864; HUTCHINSON, Lond. Hosp. Reports, 1866; WEIR MITCHELL, Injuries of nerves and their consequences, 1872; DENMARK, Med. chir. trans. iv.; PAGET, Med. Times and Gaz., 1864; BERGER (Hemicrania), Virchow's Archiv, vol. liv., 1874; GAIRDNER, Clin. Med., 1862, p. 557; VERNEUIL, Gaz. des hôpit. 1864.

## II.—PASSIVE OR VENOUS HYPERÆMIA OR PASSIVE CONGESTION.

This is a condition in which the blood stagnates in the vessels; they are overfilled with blood which, as it remains too long in the vessels, has a venous character, hence passive hyperæmia is also called Venous Hyperæmia.



**Causation.**—Passive hyperæmia frequently occurs as a consequence of **Weakness of the heart**. In the normal state the forces of the circulation, in order to remove the blood from depending parts, have to overcome gravitation. The force of the heart propagated through arteries and capillaries to the veins is generally sufficient to do this. It is, however, assisted by the muscular movements, which, in conjunction with the valves in the veins, materially assist the venous current; also by the aspiration effected by the inspiratory movements. But if the heart is weak, the blood is apt to linger in depending parts or in parts far removed from the centre; hence the name **Hypostasis** or **Hypostatic hyperæmia** applied to such conditions. Weakening of the heart occurs in many forms of disease. It is often very marked in fevers, such as typhus and typhoid, or in long-continued debilitating diseases, which produce anæmia. In these cases the blood often stagnates in the dependent parts of the lungs, or in depending parts of the skin, over the sacrum and shoulder blades in persons lying on their backs, over the trochanters in persons lying on their sides. In these latter situations the weakness of the circulation along with the mechanical effects of the weight of the body and the irritation of decomposing material, leads frequently to the sloughing of the skin and the formation of bed-sores. In fevers there may be hyperæmia of the extremities of the fingers or toes, resulting even in necrosis or gangrene.

Again, there may be difficulty in overcoming gravitation on account of the force of the heart being partly lost by reason of **Obstruction of arteries**. Thus atheroma, by producing a thickening of the internal coat may cause a partial obstruction, which is often increased by the formation of thrombi on the affected surface. In consequence of this, the force of the heart may be insufficient and the blood stagnate in the parts supplied. Complete occlusion of an artery will under certain circumstances produce extreme passive hyperæmia. (Hæmorrhagic infarction, see under Embolism).

**Obstruction of veins**, however, is the most direct cause of passive hyperæmia. This may be produced by pressure of tumours, exudations, bandages, the pregnant uterus, even hard masses of fæces, by coagulation of blood within the veins, or by the bursting of tumours into veins, or their growth through their walls.

Lastly, **Disease of the valves of the heart** produces in a large proportion of cases a general venous hyperæmia.

**Phenomena of passive hyperæmia.**—Taking the simplest case, that of obstruction of a venous stem, the first result is an increase of blood-pressure in the veins behind the point of obstruction, and an accumulation of blood in the part. If the veins have abundant anastomoses, then



the blood will soon to a considerable extent find its way by other channels and the normal conditions be restored. But from the list of causes of passive hyperæmia it will be seen that most of these involve sets of veins, or whole regions of the body; even in the case of thrombosis, the coagulation usually extends to a number of veins; hence, relief by anastomosis cannot be obtained. Apart from relief by anastomosis, however, it might be supposed that when obstruction in a vein occurs the increase of pressure would, as in the case of obstruction of an artery, be compensated by changes in the state of contraction of the arteries. But a local relaxation of the arteries would only intensify the result by superinducing an active hyperæmia, and the desired compensation could only be effected by a reduction of the blood-pressure throughout the system, which will scarcely happen. The increase of pressure re-

mains, and it affects not only the veins but, more particularly, the capillaries.

The further effects are to be traced to the excessive pressure in the capillaries. The blood accumulates in excessive quantity in these vessels and at an excessive pressure. In consequence of this an increase in the natural transudation of fluid through the capillaries occurs, and the blood corpuscles, especially the red ones, escape from the vessels. Each of these phenomena merits more special consideration.



Fig. 1.—Normal capillary, with endothelium mapped out by treatment with nitrate of silver.



Fig. 2.—Capillaries after passive hyperæmia. Apertures between the cells greatly enlarged; the so-called stomata. (ARNOLD.)

**Hæmorrhage by diapedesis** is the escape of the red corpuscles from the blood-vessels without rupture of their walls. This process occurs mainly, if not entirely, in the capillaries, and it can be shown by experiment that it does not involve rupture of these vessels. If the tongue of a frog is ligatured at its base so as to include all the veins, but excluding the artery, there will be the phenomena of hyperæmia greatly intensified, and among these phenomena diapedesis. But if the



ligature be loosened within a moderate period, the circulation is restored, and the phenomena disappear. If the escape of blood corpuscles had been by rupture, it would have continued after resumption of the circulation. As to the manner in which corpuscles escape, it is probable that they pass between the endothelial cells. Fig. 1 shows the appearance of the endothelium of a capillary mapped out by treatment with nitrate of silver, and Fig. 2 shows capillaries similarly treated after passive hyperæmia had existed. The latter illustration is taken from Arnold, who asserts that while in the normal condition there are minute apertures between the endothelial cells, chiefly at the angles where two or three meet, these are found much enlarged in passive hyperæmia. The small apertures may be called *stigmata*, and the larger ones *stomata*. The excessive pressure in the capillaries seems to be the chief agent in causing the escape of the corpuscles as well as the increased transudation of fluid. The hæmorrhage is usually inconsiderable, but in some cases it attains to considerable amounts. (See under Thrombosis of cerebral sinuses.)

That there is an increased **Transudation of fluid** can be directly proved by experiment. The flow in the lymphatic vessels has been proved to be excessive. If the lymphatics are not capable of disposing of the entire excess, then the fluid accumulates in the serous spaces and cavities of the body, giving rise to **Œdema and Dropsy**, which are to be more fully considered afterwards. This accumulation will occur if, on the one hand, the transudation be too excessive for the lymphatics to dispose of it, or if on the other, for some reason, the lymphatics do not take it up sufficiently. The current in the lymphatics depends on the blood-pressure, and we have just seen that the cause of the increased transudation is excess of blood-pressure, and so the same condition which determines the increase will, to a certain extent, cause it to be more rapidly disposed of. In cases of hypostasis, however, there is a special tendency to œdema. The hyperæmia here is due as we have seen, to weakness of the heart associated with the action of gravitation. Both of these causes will equally act on the lymphatic circulation and induce the transuded fluid to linger and accumulate.

The exuded fluid, as may be inferred, contains red corpuscles, but it does so to a much less extent in actual pathological processes than might be supposed from experiments in animals. In the latter there is a sudden obstruction, with exaggerated results; in actual disease in man the processes usually develop slowly, and there is some accommodation of the vessels. It should be added that the white corpuscles pass out of the vessels as well as the red, but not to such an extent, and that the



corpuscles, both red and white, escape from the small veins as well as from the capillaries.

The part which is the seat of passive hyperæmia will be unduly red and the colour will be dark or livid. It will be swollen, both from the overfilling of the vessels and from the œdema, and it will usually be lowered in temperature. In organs readily capable of increase in bulk, there may be considerable **Enlargement** as a result of passive hyperæmia. This is especially the case in the spleen.

Two other consequences frequently follow in prolonged passive hyperæmia. The distended capillaries exerting pressure on the surrounding structures may through time produce **Atrophy**. This is seen especially in passive hyperæmia of the **liver**, where the central parts of the lobules often show a striking absence of hepatic cells. It is also seen in the kidney and the retina. An opposite result is effected by prolonged passive congestion in the connective tissue, namely **Hypertrophy**, induced by the increased transudation which bathes the connective tissue. The hypertrophied connective tissue is dense and it gives increased density to the organs affected, hence the term **Cyanotic Induration**, applied to it. This is very common as a result of valvular disease of the heart and is seen in the heart itself, the lungs, kidneys, liver, etc. In the lungs the induration is associated with pigmentation, due to the hæmorrhage by diapedesis, hence the term **Brown Induration**.

**Necrosis** or death of parts is not a common result of passive hyperæmia. It occurs when the conditions are such as to produce complete stagnation of blood. If a loop of intestine be caught in a hernia in such a way as to obstruct all the veins, then passive hyperæmia may be followed by gangrene. Similarly ligature of the femoral vein or its obstruction by a tumour may produce gangrene, as this vein has few and insufficient anastomoses. This only occurs if the vein be suddenly obstructed.

**Literature.**—For good account of passive hyperæmia, from experimental side, see COHNHEIM's *Allgem. Pathologie*, 2nd ed., vol. i., p. 138. See also RECKLINGHAUSEN, *Allgem. Path.*, p. 28; ARNOLD, *Virchow's Archiv.* lviii., 1873; and ZIELONKO, *ibid.* lvii., 1873.

### LOCAL ANÆMIA OR ISCHÆMIA.

In this condition the vessels are more or less empty of blood, and the part is correspondingly pale.

**Causation.**—The vessels, and especially the capillaries, may be directly emptied by pressure from without. We have an artificial anæmia produced in this way by Esmarch's elastic bandage. A tumour



or an abscess may by pressure empty the vessels, and, if long continued, this may lead to necrosis of the structure concerned. In most cases, however, anæmia is due to obstruction of arteries and this may occur in various ways.

**Spasm of arteries**, that is to say, violent contraction of their middle coat, may cause such obstruction as to produce an extreme anæmia. The application of cold to the skin causes contraction of the cutaneous arteries—ether spray causes a striking anæmia by this means. The rigor at the beginning of many fevers, accompanied as it is by paleness and coldness of the surface, is due to spasm of the cutaneous arteries. In some persons the vaso-motor system is peculiarly sensitive and slight causes are sufficient to induce spasm of the arteries.

The series of phenomena which may be thus produced is grouped under the name of **Raynaud's disease**, from the writer who first gave a full description of these phenomena. In predisposed persons an ordinary exposure to cold, as in washing, will induce such a spasm as to render the fingers bloodless and anæsthetic; they are said to be "dead" (*local syncope*). In more severe cases the skin becomes dark blue and various eruptions may form (*local asphyxia*). In very severe cases there may be actual necrosis of the ends of the fingers and toes. This form of disease is always symmetrical. Some neuralgias are associated with spasm of arteries. (Du Bois-Reymond.)

**Disease of the walls of arteries** may narrow their calibre, more especially atheroma and endarteritis obliterans (which see). In both these conditions there is thickening of the internal coat, but it is only when they occur in small arteries that serious obstruction results. In the case of atheroma, thrombosis or coagulation of the blood is often superadded and materially increases the obstruction. Lastly, arteries may be obstructed by ligature or by plugging. (See Embolism).

**Phenomena and results.**—The case of complete obstruction of arteries will be considered more fully under embolism. Where there is incomplete obstruction some blood will pass by the arteries, but the capillaries will be imperfectly filled. The part will be pale, reduced in temperature, flaccid. Its nutrition will be diminished, and its elements prone to undergo atrophy and degeneration, or even, as we have already seen, necrosis. The function will be interfered with if the nutrition is depreciated, and if the anæmia affect an important organ, the results may be serious. Thus, obstruction of the coronary arteries may cause death by paralysis of the heart. Lastly, a local anæmia may produce a hyperæmia elsewhere—a collateral hyperæmia.

**Literature.**—RECKLINGHAUSEN's Handbuch, p. 35; VIRCHOW, Handbuch d. spec. Path. und Therapie, vol. i.; RAYNAUD, De l'asphyxie local et de la gangrène symétrique, 1862; and Arch. gén. de méd. xxiii., 1874; DU BOIS-REYMOND, Arch. f. Anat. und Physiol., 1860.



## THROMBOSIS AND EMBOLISM.

These two conditions are often associated, but must be carefully distinguished. Thrombosis is the coagulation of blood within the vessels or heart. Embolism is the obstruction of a vessel by a plug brought from a distance. The coagulum which forms in thrombosis is a **thrombus**, the plug which obstructs in embolism is an **embolus**. A thrombus detached from its place becomes an embolus, and an embolus, whether consisting of coagulum or not, may grow by successive deposition of clot or thrombosis.

## I.—THROMBOSIS.

In considering the mode of occurrence of thrombosis, it is necessary to refer to the **Coagulation of blood**. According to the views which are identified with the name of Alexander Schmidt (although Andrew Buchanan anticipated many of his results) three agents are necessary for coagulation, the fibrinogen, the fibrino-plastic substance or paraglobulin, and the fibrine ferment. The two former unite to form fibrine in the presence of the latter. The fibrinogen is dissolved in the blood-plasma; the paraglobulin, at least chiefly, and the ferment, entirely, reside in the white corpuscles. It is only by destruction of the white corpuscles that the paraglobulin and ferment are set free; so long as the white corpuscles circulate in the blood and remain alive fibrine cannot form. Fibrine will form when the conditions are such that the white corpuscles are no longer preserved alive. Fibrine, it will thus be seen, is the result of a chemical combination, and the resulting albuminous substance, the fibrine, is not a vital structure but a dead chemical compound. In order to the preservation of the white corpuscles they must not be exposed to the contact of dead matter. You may keep blood fluid for a long time if you simply ensure that it is in contact with living tissue. Lister long ago showed that if an artery be ligatured in two places, and cut out while full of blood, it may be hung up and the blood will remain fluid for some days. Within the body if a vessel be ligatured carefully in two places, the middle portion remaining in connection with the living tissues, then the blood may be kept fluid from twelve to fifteen days. When blood is exposed to a perfectly smooth surface, even if it be of dead matter, it does not coagulate readily. Thus a piece of glass in the circulating blood does not induce coagulation, and blood in a vessel whose internal surface is smeared with oil does not readily coagulate.

It appears from the observations of Rauschenbach that the cells of lymphatic glands when treated with water yield, like leucocytes, the ferment necessary for



coagulation. Most other cells probably possess a similar power. Foa and Pellacani assert that when fresh brain substance is treated with water and filtered, the filtrate injected into the jugular vein of rabbits induces rapid coagulation.

Hayem and Bizzozero have recently alleged the presence of a third form of corpuscle in the blood. Hayem gives the name *hæmatoblast*, and Bizzozero that of *Blutplättchen* to this corpuscle. Hayem's name implies the view that these form the red corpuscles, and as this view is probably incorrect the name is inadmissible. We shall refer to them under the designation blood-disklets. They are small colourless bodies, oval or round in outline and disc-shaped. In cold-blooded animals there are spindle-shaped bodies which are regarded as of a similar nature. The disklets undergo rapid changes when once removed from the body, running together into indefinite granular masses. The existence of the disklets as definite corpuscles has been questioned by several observers (Löwit, Weigert), and it is acknowledged that they are difficult of detection on account of the readiness with which they change when the blood is shed. Bizzozero, however, asserts that he has seen them in the circulating blood of the mesentery of guinea-pigs and rabbits. They may be prevented from changing if the blood be rapidly dried on a cover glass and stained with methylviolet (Schimmelbusch). Bizzozero asserts that it is the disklets and not the leucocytes which have to do with coagulation. He would adopt the view given above of this process, substituting the disklets for leucocytes. This view, however, has not been supported by subsequent observers. (See Eberth and Schimmelbusch.)

**The Process of thrombosis.**—If a portion of blood inside a vessel be cut off from the circulation, then it will by and by coagulate just as it does outside the body, the main agent in the coagulation being the disintegrating white corpuscles. In this way a **Red thrombus** is produced. But vessels are rarely so situated as to allow of a complete coagulation such as this; much more frequently the thrombus is formed from the blood which is still moving, although, it may be, slowly, and it is of gradual growth. The formation of thrombi in the living vessels has been carefully studied by Zahn, whose experiments throw much light on the process.

The mesentery of a frog is exposed and subjected to microscopic examination. A vessel of some size, an artery or vein, is chosen, and its wall in some way injured, as by twitching it slightly with the forceps, or placing a small crystal of common salt near it. Very soon white blood-corpuscles begin to adhere at the injured part. As the blood passes over it successive layers of white corpuscles adhere, and a growing clump of them is formed. Along with the white corpuscles a stray red one may be insinuated, or there may be several red ones. The clump so formed, be it wholly white or partly mixed with red corpuscles, may be carried off, in which case a new one begins to form; but the clump may remain fixed and be continuously enlarged by successive depositions of corpuscles from the circulating blood. In course of time a change occurs in the appearance of the clump, the white



corpuscles lose their individual outline to a great extent, and the clump gathers itself together into a grey granular mass in which neither by acetic acid nor by staining are the majority of the white corpuscles to be discovered. It has, indeed, very much the characters of fibrine which has been obtained by whipping the blood outside the body. The clump of white corpuscles, in fact, by the disintegration of the corpuscles and the attraction of the fibrinogen from the blood plasma has converted itself into a fibrinous coagulum. All the white corpuscles, however, are not disintegrated, some are still recognizable in the mass. A similar mode of formation is observed when a vessel is injured by pricking with a needle, or by cutting it. The presence of foreign bodies if they are rough on the surface produces adhesion of white corpuscles and their conversion into thrombi in similar fashion. The thrombus formed in these various ways is called by Zahn the **White thrombus**.

Eberth and Schimmelbusch, from a very elaborate series of observations, have come to several very important conclusions. They assert that Zahn's observations are correct with the exception that it is the disklets and not the leucocytes which go to form the white thrombus. They do not regard the process as one of coagulation, but rather of conglutination; that is to say, the clumps of disklets run together and form white masses, in which no proper fibrine is present. When a foreign body, such as a thread, is introduced into the circulation then true fibrine is deposited, but not in the simple white thrombus. Leucocytes are often caught in the thrombi, in which they may be present in larger or smaller numbers.

In actual pathological processes in the human subject the pure white thrombus is frequently seen, especially in the heart, but it is more frequently mixed. As a general rule the thrombus enlarges by fresh deposition, and it not infrequently happens that, as corpuscles accumulate, sufficient ferment is produced as to allow of the coagulation of a layer of entire blood. Hence strata of red coagulum may alternate with strata of white. Again the thrombus after its formation may become compacted into a dense white structureless layer having a hyaline appearance, and composed of fibrine which adheres to the wall, and is so united that its boundary is indefinite; the fibrine is sometimes traversed by canals (channelled fibrine). This condition is often seen in the external layers of clot in aneurysms. We may thus distinguish four forms of thrombus according to structure. (1) The **red thrombus**, composed of the entire blood; (2) the **white thrombus**; (3) the **stratified and mixed thrombus**; and (4) the **hyaline thrombus**.

The main cause of coagulation, then, is the contact of the white corpuscles with dead matter or altered living tissue. In the living body anything which interferes with the integrity of the vessel-wall or the endocardium is likely to predispose to coagulation. Stagnation of the blood is often set down as a cause of coagulation, but this will act



chiefly by altering the vessel-wall, and by keeping the white corpuscles removed from contact with the living tissue. As it is the endothelium of the vessels and endocardium which is in most immediate contact with the blood, it will be interferences with it that will conduce to coagulation.

**Causation.**—In most cases of thrombosis there is either a **stagnation of the blood** or else some **palpable injury to the vessel** compromising its endothelium, and the various thrombi may be studied according as they are formed in one of these ways or the other, while in some cases the action of both may be traced. As we have already seen, the stagnation is rarely complete, and accordingly the thrombi are mostly white or mixed, being formed in the way just described.

(a) **Thrombosis from stagnation of blood.**—It will be found that wherever the circulation is so altered as that, while the blood moves more or less freely in the general current, there are subsidiary currents or eddies, thrombosis is liable to begin in the situation of the latter. **Local dilatations** produce this very directly. In cases of aneurysm or varix there is a pouch containing blood which is out of the regular current and thrombosis is liable to occur, although in the case of aneurysm there is the additional fact that the wall is considerably altered. **Weakness of the heart**, especially when associated with dilatation of its cavities, leads to imperfect emptying of its cavities, and the blood forms eddies in the parts most removed from the general current. White thrombi often form in this way behind the columnæ carneæ, in the auricular appendages, and, if there is much dilatation, near the apices of the ventricles. These thrombi often assume a globular shape (Globular vegetations of Lænnec), and may grow to a large size. **General weakness or marasmus**, occurring in consequence of debilitating diseases, such as typhoid fever, phthisis, etc., renders the current in the veins sluggish. There is under these circumstances not only weakness of the heart, but the muscles being weak do not assist in emptying the veins, and respiration also fails to give due assistance. While there is a general sluggishness, the localities in which the thrombosis begins usually correspond with bye currents in the circulation. The blood in the veins moving slowly and at a low pressure does not press back the valves against the walls of the veins. Hence as the valves are half open, the blood will stagnate **in the sinuses** of the valves, and thrombosis is liable to occur there. Coagula so formed grow up the veins, but even when they are continuous they usually present a knotted character from the parts corresponding with the valves being bulkier. This knotted condition is sometimes detectable through the skin. Again marasmic thrombi often begin in the **longitudinal sinus** of the



dura mater. This vessel has somewhat rigid walls, and the tributary veins from the pia mater are small. If the current in the latter be slow then there will be eddies in the sinus, especially at the parts lying highest. The most common starting point is indeed the highest part of the sinus (Recklinghausen). Thrombosis in the **uterine veins** after delivery is partly due to stagnation of blood in cases where the uterus contracts imperfectly, but the injury to the veins in the removal of the placenta contributes.

(b) **Thrombosis from alteration of the wall.**—Wounds of vessels induce thrombosis, and this process bears an important part in stilling hæmorrhage. If the blood is stagnating in the veins, as is the case in passive hyperæmia from heart disease, a trivial wound may start a thrombosis. In cases of valvular disease the legs are not infrequently punctured to relieve the œdema which is so common in such cases, and the punctures may be the starting points of thrombosis. Ligation of vessels causes thrombosis as there is rupture of the internal and middle coats. Acute inflammation of the walls induces coagulation, as one sees so frequently in acute endocarditis. Chronic endocarditis and atheroma, by producing palpable alterations in the endocardium or internal coat of arteries, are frequent causes of thrombosis, more particularly when calcareous matter is deposited and becomes exposed to the blood. An occasional cause is the protrusion of tumours through the walls of vessels, but this scarcely ever occurs in arteries and is rare in veins.

(c) There are cases in which coagulation occurs apparently in consequence of the sudden setting free of the ferment and fibrino-plastic substance in the general circulation. This has been mostly in cases of transfusion, where blood from one of the lower animals has been used. (See case by Recklinghausen).

It is important to distinguish **Thrombi** which have formed during life from mere **Post-mortem Coagula**. It will be clear from what has gone before that the thrombi are mostly either white or grey, but being formed largely of white corpuscles they are of a dead or **opaque white** or **grey** appearance. Post-mortem clots are sometimes pale from the sinking of the red corpuscles or otherwise, but they are gelatinous and smooth on the surface and mostly transparent, whereas the thrombi are firmer, drier, more opaque and granular on the surface. The thrombi also adhere to the wall to some extent, whereas the clots do not, although in the heart, from getting entangled among the columnæ carneæ, they may have an appearance of adhesion. Lastly, the thrombi are often stratified, indicating their deposition in successive layers.

**Growth of Thrombi.**—Thrombi begin as local depositions on the internal surface, but they are liable to grow till they may fill the vessel, obstructing it and extending to further portions of the vessel. The



thrombus is composed of dead matter, and its surface is rough, so that the conditions are given for further deposition, unless the current is very rapid. In the case of the veins the blood will be for the most part stagnant above and very often below the thrombus, and it is in these vessels particularly that they are apt to grow very extensively. Thus a coagulation starting in the uterus may travel along to the internal and common iliac veins, and thence pass into the veins of the legs. In a case observed by the author, coagulation had its origin in a cancer of the kidney which involved the wall of the renal vein; it extended thence to the inferior vena cava, and to the veins of both legs which were distended with old thrombi. Thrombi in the veins will often be found growing on till they reach a vessel where the circulation is very free, and the globular thrombi in the heart often grow so as to project into the cavity. Growing thrombi will be mostly white or mixed, but as the blood above and beneath an obstructing thrombus is usually at a stand-still or nearly so, they may be almost purely red in parts.

Before leaving this subject, it may be well to consider **Whether coagulation occurs in the living capillaries.** It is to be remembered that, so long as the living endothelium exercises its due influence on the blood, coagulation will not occur. As the capillary wall is entirely composed of flat cells, it is only when necrosis takes place that the conditions requisite for coagulation will be fulfilled. Moreover, as the tissues depend on the capillaries for their nutrition, obstruction by thrombosis will involve the death of the tissues themselves. Hence thrombosis of the capillaries, or even complete stagnation in them, may be put out of the question except in connection with necrosis. If this were not so, the thrombi in veins would grow into the capillaries, and, through them, into the arteries. The capillaries, not admitting of this, form a barrier to the extension of the coagulation. It thus happens that, after extensive coagulation in the veins, the circulation is largely maintained by arteries, capillaries, and lymphatics, and necrosis rarely occurs. It is worthy of note in connection with this matter that the blood found in the capillaries after death is nearly always fluid.

**Changes in thrombi.**—If a thrombus contains red corpuscles, then their colouring matter is soon dissolved out and stains the coagulum, giving it frequently a deep brown tint. The pigment sometimes deposits crystals of hæmatoidin, which may remain long unaltered. (See Fig. 3.) In course of time further changes occur, of which the more important are softening and organization.

**Softening** is a frequent result, especially in the coagula in the heart and veins. The globular thrombi in the heart frequently exhibit this process. It begins in the central parts of the thrombus, and the



coagulum breaks down into a turbid brownish juice, the softening extending gradually outwards. The juice is often like a mixture of pus and blood, or in very white thrombi it may be like pure pus. It consists of the debris of the thrombus, and no well preserved corpuscles are to be found in it. The softening may extend outwards till a mere rind is left, and this may give way and cause the juice to be launched into the circulation.

Fig. 3.—Crystals of hematoïdin found closely aggregated in the midst of an old thrombus in a vein. The crystals have a deep red colour.  $\times 350$ .

The **Organization and Absorption** of a thrombus is a process of some interest. As the coagulum is dead, it is clear that it cannot take part in the process of organization. In studying inflammation this subject will come up for fuller discussion, but it may here be stated that when

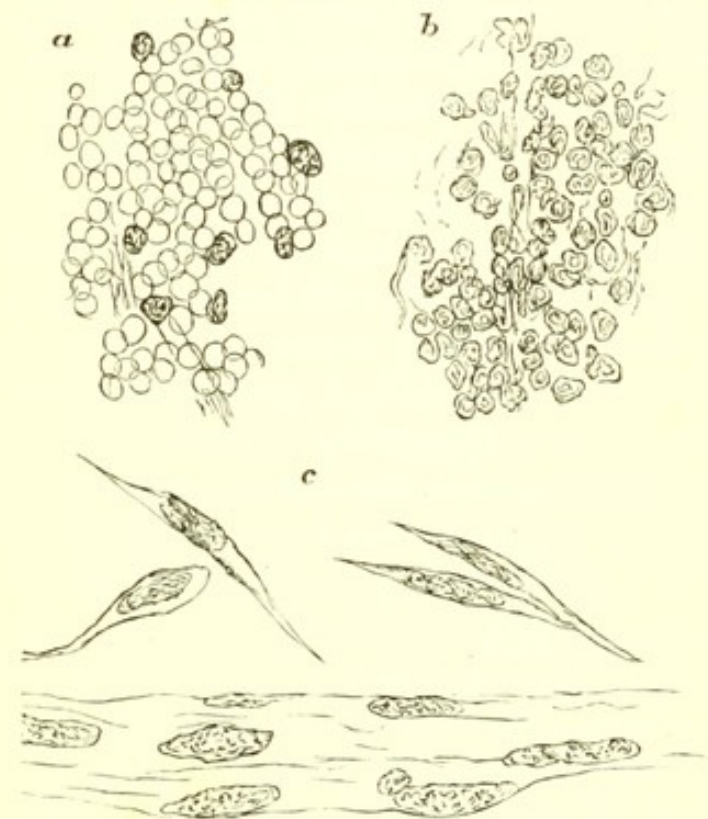


Fig. 4.—Organization of blood-clot. *a* represents the clot little altered; red corpuscles are chiefly seen, with a few solid round cells, which are like white corpuscles, but more numerous than in normal blood. In *b* the red corpuscles are entirely replaced by round cells. In *c* the round cells have given place to large spindles.  $\times 350$ .

tissue which subsequently develops into connective tissue. Like other new-formed connective tissue of this kind, this tissue tends

Thrombi, instead of softening, may dry in and shrivel. In such case we may have lime salts deposited, and even through time the formation of little stony masses — **Vein-stones or Phleboliths.**

a piece of dead animal substance is present among the living tissues, the first step towards its absorption is usually its replacement by an elementary tissue, the dead structure forming, as it were, a mould on which the new tissue forms itself. The constituents of the thrombus are, therefore, pushed aside or disintegrated by masses of round cells (as shown in Fig. 4 *b*); and at the same time blood-vessels springing from the vasa vasorum extend into the substance of the new-formed tissue. The thrombus is thus converted into, or replaced by, a vascular



to contract. The usual result is that the vessel is permanently obliterated and replaced by a solid cord of connective tissue.

There are some cases in which the process is somewhat modified. The contracting connective tissue acts on the new-formed vessels of the tissue which has replaced the thrombus rather than on the walls of the original vessel, and it has the effect of widening these thin-walled vessels. In this way a sponge-like or cavernous tissue may form, and the result may be the re-establishment of the circulation, the wide spaces of the sponge-like tissue coming to communicate with the regular calibre of the vessel. A favourite place for the formation of this cavernous tissue is the place of union of the iliac veins to form the vena cava. Here the vessels are sometimes found filled with a spongy tissue through which the circulation is carried on. By a still further contraction of the new-formed tissue, the calibre of the vessel may be completely re-established, the sinus-like blood-channels expanding into the calibre of the vessel.

In studying thrombosis, we have left out of account all cases of what is called **septic thrombosis**, where the coagulation arises in connection with the introduction of decomposing material into the veins. Such processes will receive consideration in another part.

**Results of thrombosis.**—Embolism is a frequent consequence of thrombosis in any situation. (See under Embolism).

Thrombosis in veins leads most directly to **passive hyperæmia** with its consequent **œdema** and dropsy. The occurrence of serious œdema depends largely on the extent of the anastomosis of the obstructed veins, and also somewhat on the rapidity with which the thrombosis has occurred. If the thrombus forms slowly the functions of the plugged veins may be largely taken up by the lymphatics and by other veins which remain unaffected. In the case already referred to, where a cancerous tumour in the kidney had burst into the renal vein, and thrombosis had extended to the vena cava and down the veins of both legs, there was comparatively little œdema at any time, although all the main venous trunks of both legs were plugged. The fact that the thrombosis does not extend into the capillaries allows the circulation to be carried on by arteries, capillaries, and lymphatics.

At the seat of thrombosis there is usually an **inflammation**, which may be of considerable intensity, and is sometimes accompanied by considerable pain. Even when the thrombus does not include specially irritating material, there is inflammation of the wall, leading, especially in the case of veins, to adhesion to parts around with induration (*Periphlebitis*). A periphlebitis, accompanied by considerable œdema and by severe pain, occurs in **Phlegmasia alba dolens**.

It is important to know that these secondary effects of thrombosis, and especially œdema, will not usually show themselves for some time after the onset of the process. A coagulation beginning in a vein will



take some time before it completely obstructs it, and even when it has done so, it may be necessary for it to grow into other veins before any pronounced œdema will develop. Hence it is that a thrombosis may lead to embolism by detachment of portions of the thrombi before it has manifested its presence by hyperæmia and œdema.

Thrombosis in veins seldom leads to **Gangrene**, and in the few cases in which gangrene actually occurs some additional interference with the circulation will usually be discoverable. Weakening of the heart, in conjunction with thrombosis, may cause it, and so may disease of the arteries diminishing the force of the blood.

Thrombosis in arteries, occurring, as it does, chiefly along with atheroma, may have as its result, ischæmia and its consequences. (See above.)

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## II.—EMBOLISM.

**Causation.**—Any solid material in the circulating blood may obstruct a blood-vessel which it finds too small to give it passage. It may be a foreign body, such as a parasitic animal or vegetable, or a piece of a tumour or a piece of cretaceous matter broken off from a degenerated valve or vessel. But the most frequent source of embolism is a **pre-existing thrombus**. Thrombi most readily become detached from the heart or veins, and in either case the thrombus is most dangerous when it has grown into the current so as to be exposed to the force of the blood. In a vein a thrombus which completely obstructs the vessel will not be readily carried away. The actual displacement of the thrombus will often take place in consequence of some compression or movement of the body affecting the part where it is seated. This is sometimes seen in the case of thrombosis in the uterine veins where the first considerable movement after delivery may displace the thrombus and lead to embolism of the lungs.

The embolus, whatever its source, will obstruct, for the most part, an artery or capillary. The only practical exception to this is the case of the portal vein, which in its ramifications in the liver has the distribution of an artery. The possibility of a proper venous embolism has



been asserted. Foreign bodies, especially if heavy, may fall backwards in the venous system, but this is not an occurrence of any practical moment, if, indeed, it actually occurs in human pathology (Recklinghausen). A thrombus or other loose solid body in the circulating blood will usually in its course be caught at a place where an artery is dividing; it often rides on the bifurcation. It sometimes becomes broken against the projecting bifurcation, and its fragments may be dispersed to the smaller branches, producing numerous embolisms in them. The embolus, acting as a foreign body, will usually induce thrombosis on its surface, so that it may get covered in by an **encapsuling thrombus**. As the embolus may be derived from a thrombus, and may induce thrombosis afterwards, it may be difficult to distinguish the one process from the other.

The **Diagnosis of embolism** from thrombosis rests on a survey of the existing conditions. In veins and in the heart any existing coagulum must be a thrombus, as embolism does not occur in them. In arteries we may have either, but the situation and circumstances will often guide us. Thrombosis in arteries almost necessarily implies disease of their walls, and the thrombus is firmly adherent. The embolus, on the other hand, will often be found riding over a bifurcation and non-adherent, or there may be part adherent (the encapsuling thrombus) and part non-adherent. Assistance may often be obtained by finding a source of embolism, such as thrombosis, elsewhere. Again, there are some arteries much more prone to embolism than others. In the arteries of the lungs, intestines, kidneys, and spleen, thrombi are comparatively rare, but in those of the brain, in the coronary arteries of the heart, and in those of the extremities both thrombosis and embolism are common (Cohnheim).

**Phenomena of embolism.**—These phenomena vary greatly in different cases, the variations depending chiefly on the circumstances of the vessels in regard to anastomosing communications. In this respect we have all degrees of difference.

If the **anastomosis be very free** then embolism will at once cause increase of pressure on the proximal side of the plug, and this increase of pressure may be sufficient to restore the circulation by means of the anastomosing connections. At the seat of the obstruction thrombosis will occur on each side, and by the organization of the thrombus the artery will be permanently converted into a solid cord from the nearest proximal to the nearest distal branch, the circulation being completely carried on by the anastomosing branches. The arteries with freest anastomoses are, in general, those whose branches are most liable from their situations to temporary obstructions from external pressure or



otherwise. Hence in the arteries of muscle, of the skin, of the intestine, of the circle of Willis in the brain (the common carotids are liable to pressure), and of acinous glands, embolism is usually of trivial consequence. A similar result will follow **capillary embolism**. As the capillaries are in the freest possible inter-communication, embolism produces little disturbance of the circulation, unless, as sometimes happens, there be many obstructed simultaneously.

The opposite extreme is where the **anastomosis is very imperfect**. Cohnheim has endeavoured to distinguish certain arteries of the body as being entirely devoid of anastomosing connections—they are like the branches of a tree, each division having no distal communications with its fellows. Such arteries he calls **End arteries**. It is impossible to have carry out this distinction absolutely, inasmuch as nearly all arteries at least fine communications, and all communicate by their capillaries. It may be said, however, that the following arteries have at least exceedingly imperfect anastomoses, and are practically end arteries, namely, the pulmonary, renal, and splenic arteries, the coronary artery of the heart, the central artery of the retina, and the nutrient arteries of the brain, to which may be added the portal vein. In addition, the superior mesenteric artery and the external iliac arteries have anastomosing communications, which, in relation to the size of these arteries, are small. In such arteries as those mentioned the results of embolism are not strictly uniform, but are nearly always serious. The typical result is the formation of the hæmorrhagic infarction, which will be first considered.

The **Hæmorrhagic infarction** is seen most characteristically in the lung as a result of plugging of the pulmonary artery. A piece of lung tissue, generally in the form of a wedge with its base at the surface, is solidified and presents a deep red colour, as if a solid blood clot had replaced the lung tissue. The entire vessels in the affected area are distended with blood, and blood has escaped from the vessels so as to fill up the lung alveoli, which are entirely destitute of air.

The mode of formation of the hæmorrhagic infarction has been carefully observed by Cohnheim, using the tongue of the frog as a subject of observation. The two main lateral arteries of the tongue are not end arteries, but their lateral branches are. Cohnheim introduced little blackened pellets of wax into the division of the aorta which communicates with the arteries of the tongue, and these were sometimes carried to the tongue, obstructing the arteries there. Sometimes one pellet obstructed the anastomosing branch between the main arteries while another stuck in one of the main arteries, so that the latter was converted into an end artery. Whether a main artery thus conditioned



or a plugged lateral branch was observed, the process was virtually the same. The blood was, immediately after the plugging, at a standstill, in artery, vein, and capillaries. These vessels might be moderately full of blood or might contain only blood plasma. Soon, however, a backward flow of the blood from the veins was observed (diagrammatically shown in Fig. 5), and by and by this produced an engorgement of the entire vessels, veins, capillaries, and arteries. After a time a remarkable diapedesis began to manifest itself through the capillaries into the surrounding tissue, which became engorged with blood. The portion of tissue concerned would, in accordance with the area of distribution of the artery, be wedge-shaped, and the result was a wedge-

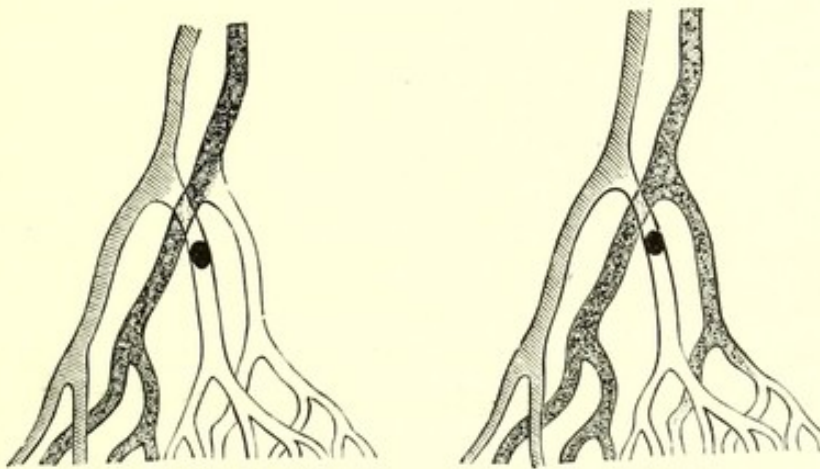


Fig. 5.—Diagram of conditions following embolism of an end artery. In the figure to the left the state of anemia immediately after the embolism is shown. In the other figure the regurgitant current from the vein is indicated. (After COHNHEIM.)

shaped red pulpy piece of tissue, not unlike a blood clot—the **hæmorrhagic infarction**. If the frog survived, the piece of tissue by and by sloughed off.

The explanation of these phenomena is not very difficult. The obstruction of the artery reduces the pressure in the vessels supplied by it to nothing. The veins are still in open communication with the veins of the part, in which, although the blood-pressure is low, it is still something. Accordingly, the backward flow occurs from the veins to the capillaries, and so to the obstructed artery. Thus the engorgement of the vessels is not difficult to explain. The diapedesis is to be referred to the fact that the capillaries, being deprived of blood, have the integrity of their walls impaired. We have already seen that passive hyperæmia produces diapedesis, owing to interference with the capillary wall. Much more serious is the nutritive deterioration in this case, and so the diapedesis is very great, leading to a marked infiltration of blood into the tissues around. The deprivation of fresh blood causes death of the tissue, which here, as it is an external part, sloughs.



It will be seen that in this occurrence three processes are involved, namely, engorgement of the vessels or hyperæmia, hæmorrhage, and necrosis. These results are by no means uniform, and in their application to human pathology they require further elucidation.

**Engorgement of the vessels** is ascribed by Cohnheim to the regurgitant current from the veins, but further observation has shown that when an artery is obstructed blood flows into the capillaries from all surrounding communications, from arteries and capillaries as well as veins, and the engorgement may be due to the flow from these as much as from the veins. Moreover, the current from the communicating arteries and capillaries may be sufficiently strong to carry on the circulation and prevent any considerable engorgement or diapedesis. This is frequently the case in the lungs, where the hæmorrhagic infarction often fails to develop after embolism. The circulation in the lungs is somewhat special. The capillaries are wide and very abundant, and the bronchial artery not only supplies nutrient branches to the lung tissue but forms anastomosing communications with the pulmonary artery. In consequence there may be none of the phenomena of the hæmorrhagic infarction, and even if the engorgement and hæmorrhage occur, it may stop short of an actual necrosis of the tissue.

On the other hand, it is common in some situations to have **Necrosis** with very little engorgement or none at all. In order to engorge the vessels must remain alive, otherwise the blood will coagulate and obstruct any backward current. But necrosis may occur before time

has been given for engorgement, or the engorgement may be limited to the peripheral parts. In the **spleen** and **kidneys** this is frequently the case. The tissue, especially if the artery be of large size, dies without any considerable engorgement, and in these organs the dead tissue undergoes a peculiar process of coagulation (see Coagulation Necrosis), the result being the formation of a dense pale wedge, the **white infarction**. (See Fig. 6.) In the spleen and kidney we have all gradations between the white and the

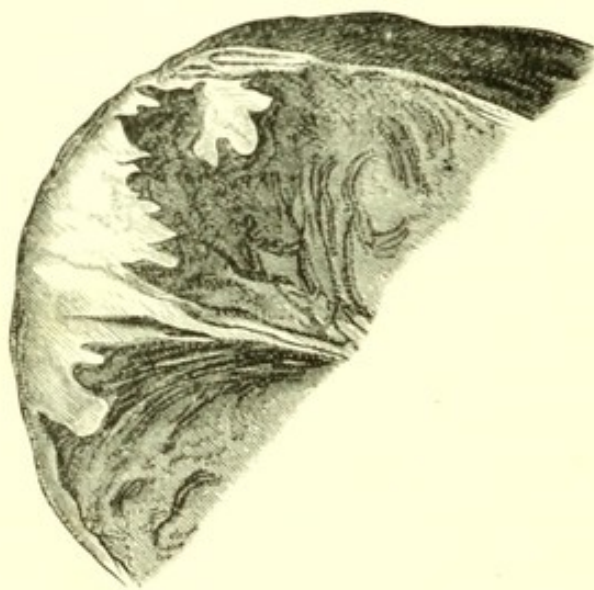


Fig. 6.—White infarction of kidney, showing the wedge-shaped piece of tissue affected. (After RAYER.)

hæmorrhagic infarctions, and a white infarction is often surrounded by



a red zone in which hæmorrhage has occurred. In the **brain** and **retina**, necrosis occurs in the form not of coagulation but of softening, and there is usually little hæmorrhage, hence a white softening.

The **superior mesenteric artery**, although it has many anastomosing branches, is so large in comparison with these vessels that its obstruction may lead to engorgement and necrosis in the portion of intestine to which it is distributed. The infarction is not co-extensive with the distribution of the artery, the peripheral parts of the district being sufficiently supplied to escape. Just as in the case of the lung the plug is usually some distance to the proximal side of the apex of the infarction, and the wedge is appreciably smaller than the area of distribution of the artery.

The **portal vein** is in its distribution an end artery, but infarction does not occur as a result of plugging. The explanation is that, not only is the liver supplied by the hepatic artery in addition to the portal vein, but the blood of the former, after passing through its own proper capillaries in the walls of the vessels and connective tissue of the liver, is carried into the inter-lobular veins which are the terminals of the portal vein. Obstruction of the latter will not therefore stop the circulation.

It will be obvious that **arteries possessing free anastomoses may be reduced to the condition of end-arteries** if their anastomoses are no longer available. If, as sometimes happens, an embolus passing to the leg breaks up, say, by being propelled against a bifurcation, and is scattered to a number of stems simultaneously, then the circulation will be re-established very slowly or not at all, and necrosis is liable to occur, especially if the circulation is already feeble. Thus, we may have gangrene of the toes occurring in this way. It is to be added that in old people obstruction of a number of arteries sometimes occurs from thrombosis as a result of atheroma, and this may likewise lead to necrosis.

**Disposal of the infarction.**—We have seen that the infarction whether hæmorrhagic or not, generally implies necrosis of the piece of tissue involved. This is not always the case in the lung, however. The engorgement and hæmorrhage imply a certain delay in the occurrence of necrosis, and in the case of the lungs the bronchial artery may sufficiently nourish the tissue as to prevent actual necrosis, and even to allow of ultimate restoration by absorption or discharge of the blood exuded. In some cases, however, necrosis does occur, so that, if the patient survive, symptoms like those of phthisis develop, and lung tissue has been known to be spit up. As the necrosed lung tissue is disposed of a cicatrix will take its place.

The solid infarctions in the spleen and kidneys, whether red or



white, are treated like dead animal matter in any protected situation among the living tissues. They are gradually absorbed (see Fig. 7) and their place is taken by cicatrices, in the midst of which, even at a late date, little pieces of cheesy-looking material may be seen. In the case of red infarctions, the blood-pigment is first dissolved out of the corpuscles, and then partly absorbed and partly deposited in the form of solid granules, the result being that the infarction becomes pale.

The softened brain tissue is also absorbed, and a cicatrix or a cyst takes its place. So is it with the retina, the piece of tissue is lost and absorbed. In the case of the superior mesenteric artery, the slough of

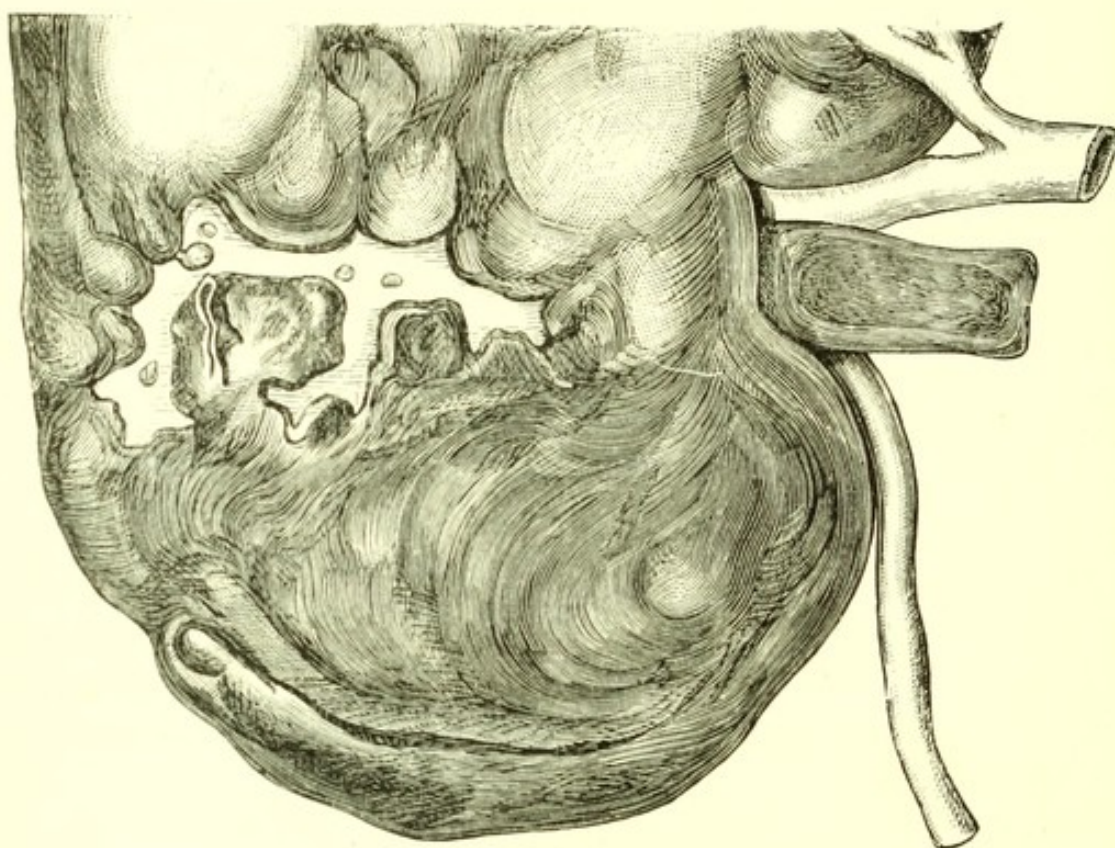


Fig. 7.—Infarction of kidney undergoing absorption. The infarction is white and depressed below the level while the tissue is puckered around it. (After RAYER.)

the bowel and hæmorrhage lead on mostly to a fatal result, but cases do occur in which, after the separation of a slough, an ulcer is left which may ultimately cicatrize.

**Special forms of embolism.**—(1) **Malignant tumours** not infrequently extend into the blood and form secondary tumours by embolism. (See Metastasis of Tumours.)

(2) **Fat-embolism.**—Oil or fluid fat not infrequently gets into the blood, and it may obstruct especially the capillaries, forming **capillary embolism**. This occurs mostly in consequence of fracture of bones, but also in consequence of injuries to the subcutaneous adipose tissue, as



in operations, or to the liver when it contains fat in excess. In these cases the fat finds entrance to the blood by the veins which are ruptured; but there are cases in which apparently it reaches the blood circuitously by the lymphatics; in suppurations occurring in parts rich in fat, such as the uterus after delivery (endometritis) this may occur (Cohnheim). Oil also occurs in the blood in cases of diabetes. (See Lipæmia.)

The oil finds its way, carried by the blood, to the right side of the heart and on into the lungs. If small in quantity it obstructs a few capillaries and small arteries, and there is no appreciable effect. But, if a considerable number of small arteries near each other be obstructed or even a large number of capillaries, we may have a condi-



Fig. 8.—Oil or fat embolism of the lung. The fat occupies the smaller arteries and extends into the capillaries, which, towards the right, form a network injected with oil.  $\times 90$ . (THIERFELDER.)

tion resembling the hæmorrhagic infarction. In some cases the oil may, to some extent, pass through the lung capillaries, and it is then found mostly in the Malpighian tufts of the kidney, one or two loops being here and there filled. There are a few rare cases on record in which extensive embolism has occurred in the smallest vessels of the brain, conjunctiva, skin, muscles, heart, intestines, liver, and kidneys, and has apparently been the cause of death.

The oil may be detected in the capillaries or small arteries in the lung or elsewhere (see Fig. 8), especially in sections which have been treated with osmic acid.

(3) **Air-embolism.**—The admission of a considerable quantity of air



into veins near the heart often leads to a fatal result, and some have supposed that embolism in the lungs is the cause. As the pressure in the veins near the heart is very low, and there is even a degree of suction during inspiration, there is great danger in opening such a vein as the jugular. Air admitted at a distance from the heart, as by the uterine veins after delivery, may also cause death (see case referred to by Birch-Hirschfeld). In regard to the cause of death in these cases it is important to notice that a repeated admission of small quantities of air does not produce serious results, whereas a single large admission is rapidly fatal. This hardly looks as if embolism were the cause, and as a matter of fact, the air is found after death mainly churned up with the blood in the right auricle and ventricle, which are usually enormously dilated. It seems that the right ventricle, in contracting, merely squeezes together the elastic air which again expands during the diastole. The force of the heart is thus fruitlessly expended and the blood is not sent into the pulmonary artery. Moreover, the over-distension of the right auricle with air prevents the admission of blood from the venæ cavæ. Thus the circulation is at a standstill and death results.

(4) **Infective embolism.**—Hitherto the phenomena of embolism have been referred to on the supposition that the obstructing plug is in itself of an indifferent nature. If this is not so, and especially if the plug contains specific micro-organisms, especially those of inflammation (pyogenic), then the results are very different. In some cases the veins are in communication with wounds which are the seat of decomposition, and the thrombi which form contain pyogenic micrococci. If embolism occurs, then each embolus, whether in an artery or capillary, is likely to form a centre of inflammation. Hence arise multiple abscesses, chiefly in the lungs, but also in the kidneys, liver, and elsewhere. A similar infective embolism may occur in consequence of the existence of a focus of infection in the heart. (See Ulcerative Endocarditis.)

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## HÆMORRHAGE.

Two forms of hæmorrhage are to be distinguished, namely, by rupture (*per rhexin*), and by diapedesis (*per diapedesin*). In the latter case the blood escapes through the walls without solution of their continuity. Hæmorrhage by rupture may take place from any size or kind of vessel, and when it occurs there will be an escape of the whole blood. Hæmorrhage by diapedesis occurs only in the finest vessels, mainly the capillaries; it is chiefly the red corpuscles which escape, and with them, of course, some fluid, but the fluid has not the constitution of the entire blood-plasma.

1. **Causation.**—The escape of blood is produced for the most part by injury or disease of the walls of the heart or blood-vessels, or by an increase in the blood-pressure, or by both combined. Some would add to these, alteration in the constitution of the blood, but this will act in most cases by injuring the walls of the vessels. It is not possible to separate accurately in their causation hæmorrhage by rupture and by diapedesis. It will be more useful to enumerate the various conditions which may lead in one way or other to hæmorrhage.

Hæmorrhage is most simply produced by **Direct injury** (traumatic hæmorrhage) as in wounds, bruises, fracture of bones, etc. It may be produced by **Reduction of pressure outside** the vessels, so that the blood-pressure is relatively increased. In this way we may have hæmorrhage as a result of reducing the atmospheric pressure in cupping, in ascending hills, or in balloons. To a similar cause may be ascribed hæmorrhage on sudden removal of tumours or exudations.

**Venous obstruction** is one of the most important causes of hæmorrhage. A frequent example of this is afforded by hæmorrhage from the stomach and intestine in consequence of obstruction of the portal vein by thrombosis or by cirrhosis of the liver. Thrombosis of the longitudinal sinus of the dura mater may lead to hæmorrhage in the brain substance, which is usually capillary, but may be in the form of a large extravasation. Violent contraction of muscles is not a common cause, but in severe vomiting the obstruction of the veins by the contraction of the muscular coat of the stomach frequently produces ecchymosis of the mucous membrane and erosions. Obstruction to the respiration may lead to hæmorrhage by obstructing the venous circulation. Thus, in death by asphyxia there are subpleural and subpericardial hæmorrhages. In all these cases the venous obstruction induces passive hyperæmia, and the hæmorrhage is mostly by diapedesis.

**Disease of the walls** of the heart or vessels is another cause of hæmorrhage. In this category are included conditions of the heart due



to disease of the coronary arteries (which see), aneurysm and atheroma of arteries, and varicose dilatation of veins. In all of these a rise in the blood-pressure is frequently the determining cause of the hæmorrhage, and hypertrophy of the left ventricle, implying a more or less permanent elevation of the blood-pressure, forms a constant source of danger, where the vessels are diseased. New-formed vessels, whether in inflammations or in tumours, are prone to rupture, presumably on account of thinness of their walls. **Obstruction of arteries and capillaries** produces hæmorrhage, as we have already seen in the case of the hæmorrhagic infarction. Fat-embolism even when the obstruction is entirely capillary may have this effect, and according to Busch and Recklinghausen the hæmorrhages may be in the brain and heart as well as in the lungs.

**Hæmophilia** or **Hæmorrhagic diathesis** is a condition in which hæmorrhage is apt to occur from very trivial causes, and it is very difficult to get the hæmorrhage stopped. It is an eminently hereditary condition—probably the most hereditary of all diseases—and is probably due to faulty construction of the vessels. Virchow has observed thinness and smallness of the aorta in one case, and we may perhaps infer that the whole system of blood-vessels is imperfectly formed. Recklinghausen seems to regard hæmophilia as in some way related in its causation to affection of the vaso-motor nerves.

**Alteration of the blood** is another cause of hæmorrhage. It may be that the blood has been altered as regards one or more of its essential constituents, as in scurvy, anæmia, leukæmia, or that a morbid poison is present in the blood, as in typhus fever (petechiæ in skin), small-pox, yellow fever, snake-bite, poisoning by phosphorus. Watson Cheyne, and Russell have recently ascribed the hæmorrhage in **purpura** to obstruction of capillaries by colonies of micro-organisms. Weigert has described a similar condition in hæmorrhagic small-pox.

**Nervous influences** may lead to hæmorrhage. The physiological hæmorrhage of menstruation is effected by means of the nervous system. The same applies to **vicarious menstruation**, in which hæmorrhage occurs in another situation, as the nose, mouth, or lungs, in consequence of interference with the ordinary process in the uterus. **Bloody-sweating** or **Hæmatidrosis** is another instance of bleeding in consequence of nervous influences.

2. **Stilling of hæmorrhage.**—In the case of hæmorrhage by diapedesis, and more particularly in hæmorrhage due to passive hyperæmia and alterations in the blood, the hæmorrhage will cease when the cause is removed. If the vessel has been actually ruptured, then the stoppage will occur by **thrombosis**. The torn edge of the aperture will itself



induce thrombosis, and as the blood is in motion the thrombus will be a white or mixed one. It often happens that as the current is too strong to allow of a thrombus readily forming at the aperture, the coagulation may begin in the blood outside, and the clot thus formed may materially assist in stopping the aperture. Thus in the case of a divided artery the contraction of the muscular coat causes narrowing of the calibre and a withdrawal of the vessel within its sheath. There is thus left a certain extent of empty sheath in which the blood may coagulate. This contraction of the muscular coat is brought about in the first instance by the stimulus of the agent which caused the rupture, and it will be kept up by the irritation of the blood on the exposed and torn structures. Other circumstances may favour the stilling of the hæmorrhage. If the bleeding is considerable the anæmia, by weakening the heart and reducing the blood-pressure, will more readily allow of thrombosis. On the other hand, circumstances may be unfavourable to the stilling of the hæmorrhage, more especially when they hinder contraction of arteries. Amongst these may be mentioned disease of the walls of arteries (atheroma, aneurysm), the application of warmth, and a longitudinal wound. In the last-mentioned case the contraction of the artery will cause the wound to gape, and the hæmorrhage is with difficulty stopped unless the surgeon cuts the vessel right across.

3. **Seat of the effused blood.**—When bleeding occurs at the free surface of the body, the blood usually passes away and is lost. On the other hand, it may take place into an existing cavity, or the blood may make a cavity for itself by tearing. Sometimes the blood in such a cavity forms a distinct tumour-like mass, which may be somewhat permanent; this condition is called **Hæmatoma**. If the blood permeates the interstices of the tissues without tearing, the condition is called **Hæmorrhagic infiltration**. Infiltration of the skin or subcutaneous tissue is called **Ecchymosis** or **Sugillation**. Small patches of hæmorrhage in the skin arising chiefly in purpura and acute fevers are called **Petechiæ** or **Vibices**.

Certain hæmorrhages have received special names, thus, **Epistaxis** is bleeding from the nose, **Hæmatemesis** from the stomach, **Hæmoptysis** from the respiratory organs, **Metrorrhagia** from the uterus, **Hæmaturia** from the urinary organs. When blood accumulates in cavities, special names are sometimes given, thus in the case of the uterus, **Hæmatometra**; of the pleura, **Hæmothorax**; of the pericardium, **Hæmopericardium**; of the tunica vaginalis of the testicle, **Hæmatocoele**.

4. **Disposal of the blood.**—In the case of hæmorrhage by diapedesis and in most hæmorrhagic infiltrations, the blood, being in the serous spaces and in communication with the lymphatics, is readily carried off. Before it is finally disposed of in this way, however, it may have under-



gone changes, especially in the red corpuscles, such as result in a greater or lesser degree of pigmentation. These changes are, however, similar to those about to be mentioned as occurring in larger hæmorrhages.

In larger hæmorrhages the blood forms a mass and coagulates, while at the periphery there is a certain amount of infiltration into the

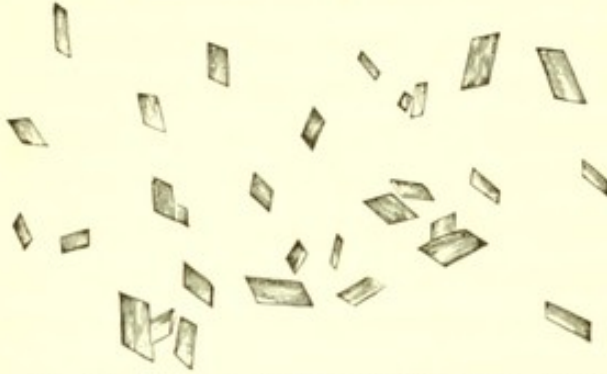


Fig. 9.—Crystals of hæmatoidin from an old hæmorrhage in the brain. Their colour is reddish brown.  $\times 350$ .

surrounding tissue. An early phenomenon is the solution of the hæmoglobin of the red corpuscles. A red solution is thus formed which may stain the tissues for some distance around. The peculiar bright red colour which one sees in the neighbourhood of a hæmorrhage in the brain, or the various colours seen in the skin after an effusion of blood into it, are due to staining of the tissues with the dissolved colouring matter of the blood, but neither of these is permanent. As the pigment is dissolved it will pass off with the fluid into the lymphatics, and the staining, while it may be somewhat extensive, will be evanescent. A more durable **Pigmentation** may result in two different ways. In the first place, the colouring matter after being dissolved out of the corpuscles is often, after a time, deposited in the solid form, appearing as crystals of hæmatoidin (see Fig. 9), or as granules. In the second place, the red blood corpuscles may be taken into the substance of other cells, and so disposed of. (See

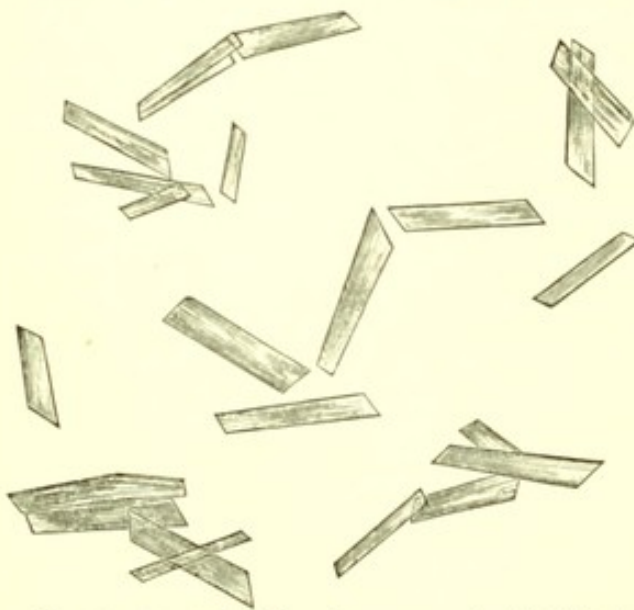


Fig. 10.—Crystals of hæmin prepared artificially by adding glacial acetic acid to a drop of blood, heating and evaporating to dryness.  $\times 350$ .

Pathological pigmentation.) If the blood is considerable in amount it comes to be treated as a piece of dead animal matter in the midst of the living tissues. Inflammation occurs around it, and this leads to what is called **Organization of the clot**. In this case the process is very similar to that already referred to in the organization of the thrombus (which see), and the result is usually the formation of a cicatrix. Sometimes a process

in some respects similar to the formation of cavernous tissue in place of the thrombus occurs, but instead of the meshes being filled with blood

in some respects similar to the formation of cavernous tissue in place of the thrombus occurs, but instead of the meshes being filled with blood



they are filled with serous fluid. Thus in the case of a hæmorrhage in the substance of the brain the new-formed connective tissue which replaces the effused blood, is unable, from the brittle nature of the brain tissue, and the fact that it is contained in a closed rigid cavity, to form a cicatrix by its contraction. Instead of that, by its contraction it leaves spaces which are filled with serous fluid, and so we have the **apoplectic cyst**.

An old clot which is not in a position to be readily disposed of in any of the ways described may dry in and finally become impregnated with lime salts. It need hardly be said that when exposed to septic influences, a clot is liable—perhaps very liable—to undergo decomposition.

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#### CEDEMA AND DROPSY.

In order to understand these conditions it is necessary to refer to certain points in the normal relations of **the lymphatic circulation**. The connective tissue throughout the body is as it were permeated with spaces of various shapes in which the connective tissue corpuscles lie. These lacunæ or serous spaces are lined with endothelium like the blood and lymphatic vessels themselves. The serous spaces are provided (as shown in Fig. 11) with numerous anastomosing processes, and they communicate on the one hand with the interior of the blood-vessels, and on the other with the lymphatics. Arnold and others (see Fig. 12) have succeeded in injecting the serous spaces from the blood-vessels, especially when, as in passive hyperæmia, the channels of communication between blood-vessels and spaces have been widened. We are to suppose that a circulation is continually proceeding from the capillary blood-vessels into these spaces, and so into the capillary lymphatics. The serous cavities of the body are to be regarded as large serous spaces. They also are lined with endothelium, and are continuous with the lymphatic capillaries. To a considerable extent also the lung alveoli are similar to the serous spaces. We know that fine dust inhaled into the alveoli readily passes into the lymphatics, and in œdema of the lung the fluid is in the alveoli.



(Edema is over-filling of the serous spaces with fluid, dropsy is over-filling of the serous cavities, but the latter term is often used in a general sense to include both. When œdema affects the surface of the body generally it is often called general dropsy, or **Anasarca** or **Hyposarca**. Opposed to this is local œdema and local dropsy.

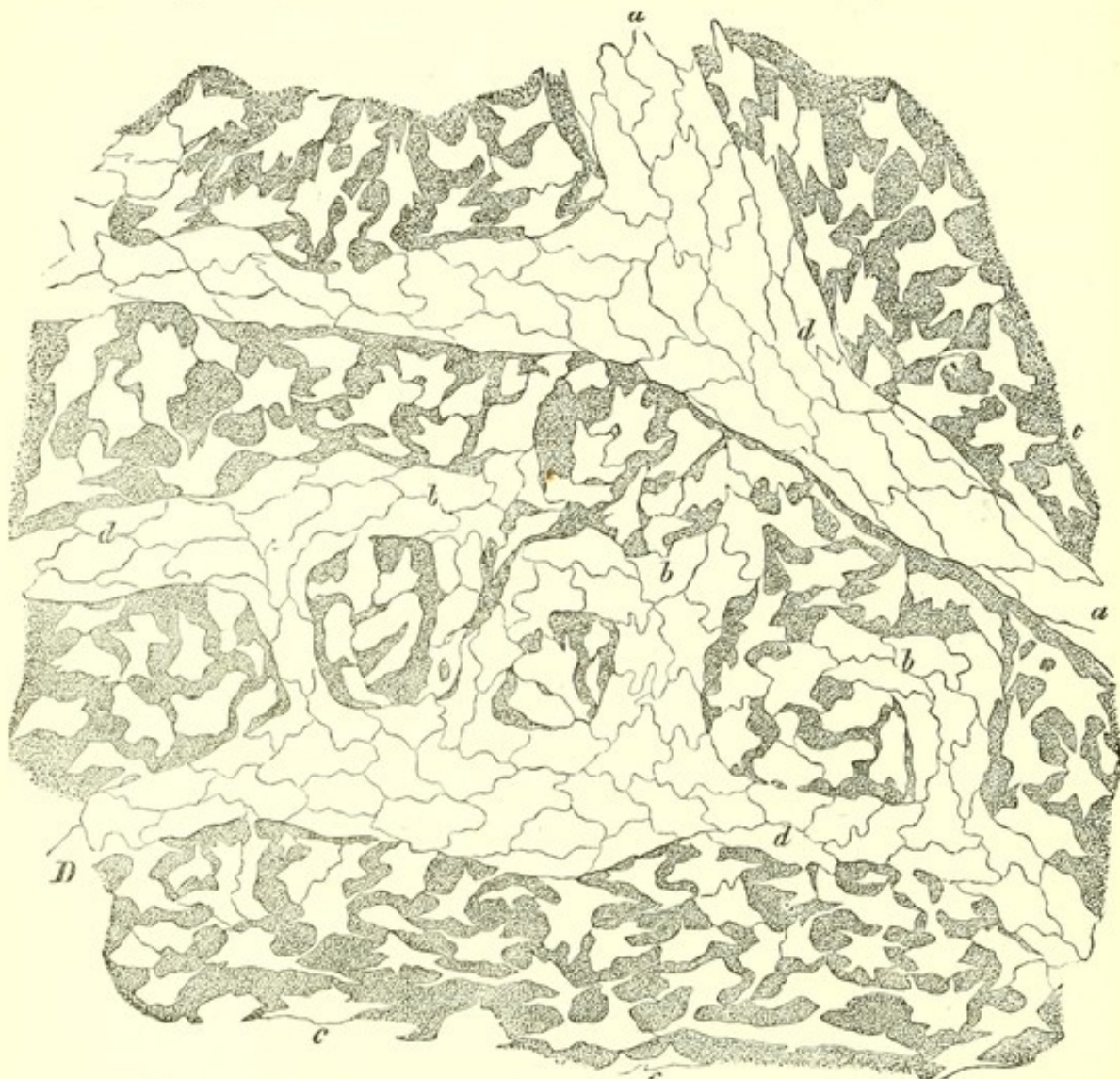


Fig. 11.—Serous spaces (*ce*) and lymphatic capillaries (*dd*) on pleural surface of diaphragm in preparation treated with nitrate of silver. The connection of the branched spaces with the commencing lymphatic vessels is shown. (RECKLINGHAUSEN.)

**Causation and nature of the process.**—Anything which causes an excess of fluid in these spaces or cavities will produce œdema or dropsy, and the first condition which suggests itself is obstruction of the lymphatics. The force which carries on the lymphatic circulation is largely the blood-pressure propagated from the capillaries, and it may be expected that **obstruction to a lymphatic stem** would produce accumulation of fluid in the spaces. But it is to be remembered that the serous spaces are in as close relation to the blood-capillaries as they are to the lymph-capillaries, and any obstruction to the lymphatics will probably have the effect of causing the transuded fluid to return to the blood-



capillaries. Majendie demonstrated that the power of absorption possessed by the veins is very great, and experiment has shown that the whole lymphatics of a limb may be ligatured, or, what is equal to that the entire lymphatic glands excised without producing œdema. The blood-vessels take up the functions till new lymphatic channels are formed.

When the **Thoracic duct** is obstructed there is not necessarily any œdema or dropsy, although lesions which obstruct the duct generally interfere also with the veins and predispose to œdema. On the other hand, when the duct is obstructed there is distension distal to the seat

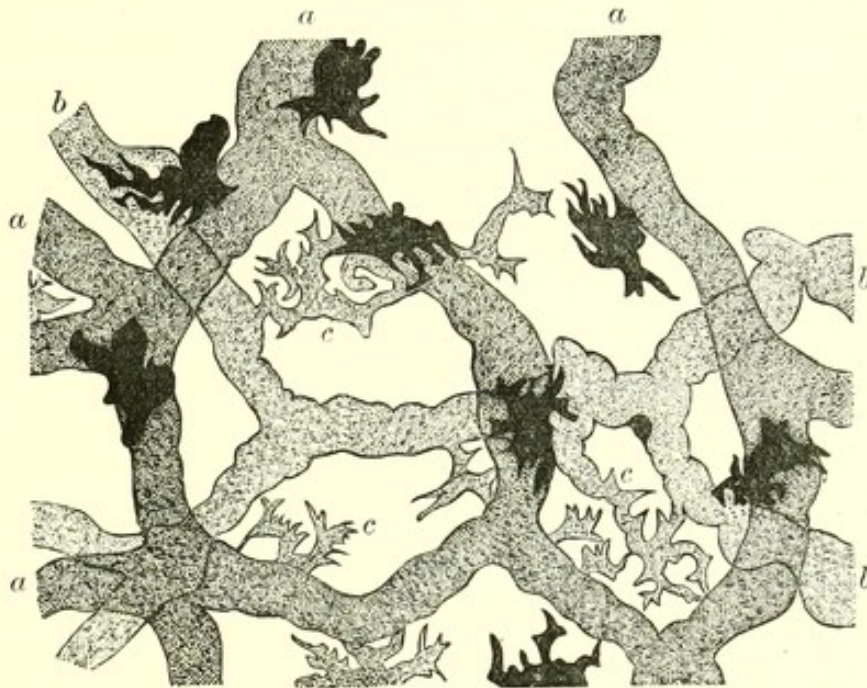


Fig. 12.—Capillary blood-vessels, serous spaces and capillary lymphatics of the frog's swimming web, filled with (blue) injection material thrown into the blood-vessels. (The black branched bodies are the normal pigment cells.) *a, a*, Capillary blood-vessels filled with injection material. *b, b*, Lymphatic capillaries also injected but less full than the blood capillaries. *c, c*, Serous spaces injected from the blood-vessels. The injection was made after passive hyperæmia had been produced by ligaturing the vein. (ARNOLD.)  $\times 250$ .

of obstruction, and this may lead to rupture of the chyle vessels in the abdomen or of the duct itself. In this way occur **Chylous ascites** and chylous pleural exudation. Recklinghausen has also seen a case of chylous ascites arising from obstruction of the chyle vessels of the mesentery with cancerous growth. Manson, again, accounts for the occurrence of chylous urine and lymph scrotum by obstruction of the lymphatic glands with the ova of the *filaria sanguinis* (see under Parasites). We may say, however, that with these exceptions, obstruction of lymphatics does not of itself lead to œdema or dropsy, although it is clear that it may aggravate an œdema whose cause is to be found in some other condition.

**Passive hyperæmia** is the most frequent cause of œdema and dropsy. Here there is increased pressure in the capillaries and veins, and a



greatly increased transudation. From what has gone before, it will appear probable that the increase of pressure is not the only element in the case, but that the capillary wall is so altered as to be abnormally permeable. This view appears more probable when we find that **Active hyperæmia** rarely produces œdema unless complicated with inflammation, in which case the agent which causes the inflammation presumably acts on the capillary wall. (See under Inflammation.) In passive hyperæmia the fluid exuded will be a watery serum, and as there is diapedesis it will contain red corpuscles in considerable numbers. The red corpuscles will be taken up by the leucocytes as already described, and carried to the lymphatic glands where they will disintegrate.

It should be added that it requires considerable increase of pressure in the capillaries to produce an œdema by passive hyperæmia when acting alone. It has even been stated that a simple passive hyperæmia, apart from vaso-motor paralysis (producing also active hyperæmia), does not lead to œdema. It has been shown that if the iliac vein of a dog be ligatured there is no œdema till the sciatic nerve is cut. The explanation of this, however, may be that this vein has such free anastomoses that its obstruction by ligature does not raise the blood-pressure sufficiently to induce œdema, whereas a coincident vaso-motor paralysis producing dilatation of the arteries raises the blood-pressure sufficiently. Experiment proves that if a sufficient number of veins be obstructed œdema will result without vaso-motor paralysis. Sotnitchewsky, by introducing plaster of Paris into a peripheral vein of the leg of a dog while the thigh was constricted by an elastic band, succeeded in obstructing a number of veins by the gypsum hardening in them. In this case œdema followed without the induction of vaso-motor paralysis. In passive hyperæmia from thrombosis of veins there are usually many veins obstructed before œdema occurs. In the case of passive hyperæmia from heart-disease it may well be supposed that the prolonged venous engorgement will produce a more than usual deterioration of the capillary wall, besides a depreciation of the blood itself. Hence valvular disease of the heart is the most frequent cause of this form of œdema.

A watery state of the blood, or **Hydræmia**, is frequently asserted as a cause of the so-called **cachectic œdemas**. According to Cohnheim and his assistants a simple anæmia induced in animals by repeated blood-lettings and the replacement of the blood by a solution of common salt, is not sufficient to induce œdema. In the human subject, however, the conditions are different from those of such an acute hydræmia, in respect that in chronic emaciating diseases we have a prolonged deterioration of the blood, during which the nutrition of the tissues generally will suffer. Under these circumstances the blood-vessels become more permeable. There is usually also in these diseases weakness of the heart, and consequently a tendency to passive hyperæmia in dependent parts. A slight amount of passive hyperæmia will, in the existing state of the blood and tissues, induce œdema. In accordance with these facts, the œdema is usually in the lower extremities.



The causation of the characteristic **Œdema of Bright's disease** is much more difficult to explain. There is, indeed, in acute Bright's disease a serious reduction in the specific gravity of the blood serum. It has been found reduced to 1020, or even 1013, instead of standing at 1030, which is the normal. This hydræmia is induced in two different ways. There is the loss of albumen, which renders the blood directly more watery, but there is also an abnormal retention of the water in the blood. In acute Bright's disease, especially, there is a great reduction in the amount of urine secreted, and this implies an increase in the water of the blood. It is in these cases, it will be observed, that the occurrence of œdema is most frequent. These facts naturally suggest that if the œdema in Bright's disease is not due simply to the hydræmia, it may be due to this in conjunction with the increased bulk of the blood, to a **Hydræmic plethora**. This view has obtained much support. It has been supposed that, the entire bulk of the blood being increased, there will be increased pressure in the capillaries and increased transudation from the watery blood. This explanation, however, is not a sufficient one, as it is opposed by observations in the human subject and by experiments on animals. In the first place there are cases of absolute suppression of urine, as in hysterical females, etc., in which we must presume an exaggerated hydræmic plethora, but without a trace of œdema, and, in the second place, an exaggerated hydræmic plethora may be induced in animals without leading to œdema of the skin.

In some experiments by Cohnheim and Lichtheim, a hydræmic plethora was induced by injecting a solution of common salt into the vessels of animals. Enormous quantities, in some cases more than three times the bulk of the blood, could be injected without killing the animal. As in other cases of transfusion, these injections did not produce any permanent rise in the blood-pressure. The most direct result was a great acceleration in the speed of the blood current, presumably from the diminished friction of the watery blood. There was also a greatly increased transudation, and consequently, increased flow of lymph. But this was remarkably localized. There was a greatly increased flow of lymph in the thoracic duct, showing that in the viscera the transudation was increased; while the lymphatics of the limbs showed no increase, indicating that in the skin and muscles the transudation was not increased. There was also greatly increased secretion from the salivary glands, stomach and intestine, liver, lachrymal glands, and kidneys. In these organs an œdema was developed, that is to say, in the mucous membrane and submucous tissue of the stomach and intestines, in the lymphatic glands of the mesentery, in the pancreas, kidneys, gall bladder, salivary glands; there was also ascites, but no œdema of the subcutaneous tissue. The explanation suggested for this localization of the increased transudation and the œdema is, that these organs are all normally concerned in the removal of water from the blood, and that probably their blood-vessels are unusually permeable to water. The experiments produce a great increase in the water of the blood, and the vessels give ready passage to it.



These observations and experiments may seem to throw little light on the œdema of Bright's disease, except in a negative way. But looking closely at the experiments, they show that hydræmic plethora induces increased transudation and œdema in parts predisposed. In Bright's disease we have two elements which are wanting in these experiments. On the one hand the kidneys are inactive, and on the other there is in the blood an irritant, which, acting on the kidneys, has induced the disease in them. This irritant may be presumed to act on the skin as well as on the kidneys. As Cohnheim has pointed out, there are many facts which indicate a close relation between the skin and kidneys in Bright's disease. In scarlatina, for instance, we have an acute inflammatory hyperæmia of the skin during the febrile stage. But there are, even in this stage, inflammatory changes in the kidneys, and when these phenomena advance sufficiently to induce a hydræmic plethora, it is perhaps not remarkable that œdema occurs in the previously damaged skin. In the experiments above referred to, it was found that although the normal skin did not become œdematous, yet when it had been previously inflamed, as by pencilling with iodine solution, or by exposing to a strong sun, then œdema showed itself. The œdema of Bright's disease hence approaches in its pathology to inflammatory œdema.

**Nervous influences** sometimes play a part in the production of œdema. The most familiar examples of this are afforded by erythema nodosum, and urticaria, where nervous irritation, usually reflex, induces localized swellings, presumably œdematous, in the skin. The bites of insects, especially fleas, produce similar effects in some persons, and, in this case as in the others there are sensations indicating irritation of the nerves. In injuries to nerve-stems, some authors have observed local œdema, and œdema sometimes follows paralysis, both paraplegic and hemiplegic, in the latter case forming a hemi-anasarca. Probably in all these cases the nervous lesion merely induces a condition of the vessels which predisposes to œdema, and its actual occurrence is determined by other causes. Thus in the case of urticaria and flea bites, the parts are usually subjected to rubbing, while in paralysis the position of the limbs, conspiring with vaso-motor paralysis may determine œdema by inducing passive hyperæmia.

**Position and character of the transudation.**—The fluid transuded accumulates where there is room for it, and especially in places where the tissue is loose and easily stretched. The elasticity of the tissue will therefore have a considerable influence on the locality of the œdema. The œdema of Bright's disease, for instance, frequently



begins in the loose subcutaneous tissue about the eyelids, and œdema in the larynx is in the loose ary-epiglottic folds.

The dropsical fluids are deficient in albumen as compared with the liquor sanguinis or inflammatory exudations, but they contain soluble salts in proportions nearly equal to these fluids.

The amount of albumen varies in different cases, being determined to some extent by the state of the blood, and the condition of the blood-pressure. It varies also very markedly according to locality. Thus the percentage of albumen is stated by Reuss to be, in the case of the pleura, 22·51; pericardium, 18·33; peritoneum, 11·14; subcutaneous tissue, 5·79; brain and cord, 1·14. Recklinghausen accounts for the larger amount of albumen in the pleura and peritoneum, as compared with the subcutaneous tissue and pia mater, by the greater richness of the former in capillaries. Some authors assert that in the pia mater there are only veins and arteries. The fluid according to its circumstances may contain bile pigment, biliary acids, fat, etc., and in Bright's disease it regularly contains urea.

**Myxœdema** is a term of recent introduction, which designates a condition characterized by certain alterations in the skin and elsewhere; the condition seems to be induced by atrophy or removal of the thyroid gland. (See further on.)

**Literature.**—VIRCHOW, *Hanb. d. spec. Path.*, i.; ARNOLD, *Virch., Arch.* lxii. 157, and lxiv. 120; RECKLINGHAUSEN, *Allg. Path.*, p. 94, etc.; SOTNITSCHESKY, *Virch. Arch.* lxxvii. 85; COHNHEIM und LICHTHEIM, *Virch. Arch.*, lxix. 106, and COHNHEIM, *Allg. Path.*, vol. i.; HILTON FAGGE, *Principles and Practice of Med.*, ii. 442; RNEBERG (Filtration of solution of Albumen), *Arch. d. Heilk.*, xviii. 1; HEIDENHAIN, *Hermann's Physiol.* v.; WEIR MITCHELL, MOREHOUSE, and KEEN, *Gunshot Wounds*, etc.; REUSS, *Deutsch. Arch. f. Klin. Med.*, xxiv. 183.



## SECTION IV.

## ALTERATIONS OF THE BLOOD AND ITS CONSTITUENTS.

GENERAL HYPERÆMIA, *Plethora* ; produced artificially by Transfusion ; *Plethora* as a pathological condition. GENERAL ANÆMIA, *Spanæmia* ; Causation ; hæmorrhage, hæmoglobinuria ; Character of lesion ; in oligæmia ; in pernicious anæmia ; in chlorosis ; in secondary anæmias. Secondary changes in anæmias. LEUKÆMIA, Causation obscure ; changes in blood ; spleen ; bone-marrow ; lymphatic glands ; and in liver, kidneys, etc. ABNORMAL CONSTITUENTS IN THE BLOOD. (1) *Melanæmia* ; pigment in blood and organs ; (2) *Lipæmia* ; fat in blood, chiefly in diabetes ; (3) *Uræmia* ; (4) *Diabetes mellitus*, arising from conversion of glycogen into sugar ; relation between food and sugar production ; experimental production of glycosuria ; relation to lesions of pancreas and semilunar ganglion.

THE blood is to be regarded as a tissue which is being continually changed by loss and renewal of its constituents. It is a fluid in which are suspended red and white cells or corpuscles. It is a vehicle by which the nutritious material required by the tissues, and the oxygen necessary for their respiratory processes are conveyed ; the former function being performed by the liquor sanguinis, and the latter by the red corpuscles. It also carries off to their proper places of excretion the products of tissue change, and the carbonic acid resulting from the respiratory process in the tissues.

The mode of formation of the corpuscles of the blood is still obscure, but it is generally believed that the bone-marrow, spleen, and lymphatic glands have essentially to do with their renewal.

## GENERAL HYPERÆMIA. PLETHORA.

These terms designate the condition in which the total quantity of blood in the body is in excess. The first question that arises here is, whether the vascular system is capable of accommodating more blood than it contains normally. There is no difficulty in answering this question in the affirmative. If the vessels as a whole could contain no more blood than they normally hold, then there could be no local variations, no blushing, no increase in the quantity of blood during



activity of organs. The capillaries present great varieties in the amount of blood they contain at different times, and if we take the whole capillaries of the body into account, it will appear that in them there is a great reserve space which may possibly on occasion be used for the accommodation of an excess of blood. This matter has been brought to the test of experiment, blood being transfused into the circulating apparatus in animals.

**Transfusion of blood** is the artificial introduction into the vascular system of blood from another person or animal.

In the experiments of Worm Müller, which are chiefly of importance here, the defibrinated blood of dogs was injected, with precautions, into the vessels of animals of the same kind. It is not remarkable that the vascular system can accommodate a large additional quantity of blood, but it is remarkable that it should do so with little disturbance to the health of the animal. A quantity equal to a half or three fourths of the blood normally present in the animal may be injected without injuring the health of the animal, and the quantity needs to be double the normal, or over it, before the animal's health is seriously impaired. It might be expected that when the vascular system is thus overfilled the blood-pressure would be greatly raised, but registration of the pressure, by a cannula introduced into the carotid and connected with a kymographion, shows that though during the actual operation there is a rise of pressure, yet it soon falls to within normal limits. If a quantity is injected more than equal to the normal bulk of the blood, then the pressure begins to show remarkable elevations and depressions, and the animal usually dies in the course of the same day or the next. Life is immediately endangered when one and a half times the normal bulk is injected.

When blood is transfused then in large quantity it finds accommodation chiefly in the capillaries and veins, the arteries being relaxed in order to admit of its passing into these, and the blood-pressure is not raised. It appears that the capillaries and veins of the abdominal organs form a great reservoir for the accommodation of excessive blood, and that the blood after transfusion lodges here chiefly.

The excess of blood thus supplied to an animal does not, however, remain permanently as a part of its organism. There seem to be arrangements in the body for disposing of it. Worm Müller endeavoured to determine what time it took to **dispose of a large excess of blood**. After the transfusion the animals were starved, and the blood and urine examined at intervals. It was concluded that the fluid of the blood is rapidly disposed of, being excreted by the urine. A few hours after transfusion much of the excess of liquor sanguinis had already gone, and even when 60 to 80 per cent. of the entire blood was injected, the whole excess had disappeared in the course of two or three days. The rapid disposal of the liquor sanguinis leaves the blood corpuscles greatly in excess, and the rate of disappearance was determined by counting them by means of Malassez' method. If the quantity transfused is small, the corpuscles may be disposed of in a few days; but in the case of large transfusions it may take two or even three weeks. In this comparatively short period of time, however, the whole excess of blood, both plasma and corpuscles is removed, and we may infer that there are arrangements in the body for the regulation of the amount of blood.

These experiments all refer to the transfusion of defibrinated blood. If blood is



injected when just **about to coagulate**, or if when it has just begun to coagulate some of the serum is pressed out and injected, then in some cases the animal dies rapidly with symptoms referrible to pulmonary embolism. The blood in the right side of the heart and pulmonary artery is found after death to be coagulated. The explanation of this is that, at the time of coagulation of the blood, the fibrinoplastic substance and the ferment, which Schmidt's observations have shown to be so directly concerned in coagulation, are both set free. It appears that even a small quantity of these substances introduced in the free state into the blood of a living animal induces coagulation of it. The importance of these facts in regard to transfusion in human beings will not be overlooked.

Hitherto the transfusion of blood from an animal of the same kind has been referred to, dog's blood injected into a dog. But when blood **from a different species** of animal is used, there are frequently symptoms of poisoning developed. The urine is blood-coloured, and it contains albumen and tube casts; there is vomiting of bloody material, diarrhœa, etc. It is as if the foreign blood had acted as a poison, and we may inquire what is the nature of the poison. The observations of Ponfick and others have thrown considerable light on this matter. The urine is blood-coloured, but microscopic examination shows that this is due, not to the presence of red blood corpuscles, but to that of the colouring matter of the blood in solution. It is not a bloody, but a blood-coloured urine, not a hæmaturia, but a **Hæmoglobinuria**. The fact seems to be that there is no poison in the foreign blood plasma, but that the corpuscles are unable to survive in the alien blood; they dissolve, and their hæmoglobin being dissolved in the plasma acts as a poison, and is eliminated by the kidneys. It seems to exercise its deleterious influence chiefly on the kidneys themselves, the tube casts and albumen in the urine being evidences of the irritation of these organs. The general symptoms, in fact, are largely referrible to irritation of the kidneys.

If hæmoglobin be introduced into the blood in other ways it acts equally as a poison. The red corpuscles of animals may be destroyed by repeatedly freezing and thawing the blood, and when that has been done, the blood of the same kind of animal will produce symptoms such as those referred to.

It is to be observed that all animals are not exactly in the same relation to each other in this respect. It is found, for instance, that a much smaller quantity of lamb's blood than of hen's blood produces hæmoglobinuria in a dog. The importance of these facts in regard to transfusion in the human species will be apparent.

In this place it may be mentioned that the blood transfused does not, according to Panum, act as food or take its place. If an animal is undergoing starvation, transfusion does not avert or delay the process of inanition, it rather hastens it by increasing the waste. In using transfusion as a means of treatment the principal object is to make up the bulk of the blood, and this can be effected by the injection of serum or solution of common salt, with a small proportion of alkaline salt, making the solution near the specific gravity of the blood.

**Plethora, as a pathological condition**, is not of great importance. It may be presumed that in some persons the blood-forming organs, probably along with the other structures of the body, are unduly active in their nutritive processes. In that case there may be an excess of blood in the vascular system, which will manifest itself in an overfulness of the capillaries and veins throughout the body, but especially



in those of the abdominal viscera. Persons of vigorous digestion and active habits have often a florid appearance, as if the vessels, of the skin at least, were overfilled. The excess of blood is used, to a considerable extent, in the formation of fat, and we know that the subcutaneous adipose tissue and that of the abdomen are often much increased. But the observations mentioned above show that any excess of blood is disposed of with considerable rapidity, and we may infer that in the human subject a moderate tendency to plethora will be overcome. It will develop when the formation of blood keeps in advance of its destruction by the arrangements provided for that purpose.

There may be a certain amount of plethora resulting from the stoppage of customary bleedings, as from piles, excessive menstruation, etc., and there will be a definite increase in the amount of blood relatively to the capacity of the vessels when, before amputation of a limb, its vessels are emptied into the circulation by the application of an elastic bandage.

We have again an increase in the total volume of the blood in cases where the capacity of the vascular system is increased. This occurs in many cases of disease of the heart, where not only the cavities of the heart are often greatly enlarged, but the capillaries and veins in the liver, kidneys, etc., are much dilated. Some of the symptoms in heart diseases may be due to difficulty in dealing with the increase of the mass of blood.

In **new-born children** there will frequently be a sudden increase of the mass of the blood. Before birth the foetal blood is divided between the child and the placenta. After the birth of the child, if the umbilical cord be not immediately ligatured, the contraction of the uterus will empty the foetal part of the placenta into the child. It has been estimated that this will sometimes amount to 100 grammes, or equal to one half the previous bulk of the blood. A portion of this blood will be accommodated in the lungs, which, on their inflation, increase the capacity of their vessels, but these will not always take up the whole excess. It is asserted by some that the common icterus of the new-born (see *Icterus*) is due to the disposal of the excess of red corpuscles.

**Literature.**—WORM MÜLLER, *Transfusion u. Plethora*, 1875; PANUM, *Virch. Archiv*, vols. xxvii., xxix., and lxiii.; PONFICK, *ibid.* lxii.; BUDIN, *Comptes Rendus*, 1876; GOLTZ, *Virch. Archiv*, vol. xxix; KRONECKER und ZANDER, *Berl. Klin. Wochensh.*, 1879, No. 52; JENNINGS, *Transfusion of blood and saline fluids*, 1888.

#### GENERAL ANÆMIA. SPANÆMIA.

The term **Anæmia** would mean literally want of blood, but it is used in a wider sense to indicate defect in any of the essential constituents



of the blood. **Spanæmia**, which means poverty of blood, would more strictly designate these conditions, but this term is seldom used.

As the blood may be defective in its various constituents different terms are used to indicate the character of the defect. **Oligæmia** is a defect in the bulk of the blood as a whole, such as results from a severe hæmorrhage, and it is virtually equivalent to **Acute traumatic anæmia**. **Oligocythæmia** is a defect in the number of the red corpuscles. This condition is sometimes called **Aglobulism**. When the red corpuscles, although, perhaps, normal in number, are deficient in hæmoglobin, the condition is called **Achromatosis**. A watery or dilute condition of the liquor sanguinis is called **Hydræmia**. As the essential constituent of the blood plasma is albumen, the term **Hypalbuminosis** is used instead of hydræmia, especially in cases where there is a direct drain on the albumen.

**Causation.**—As the conditions included are various, so the causes differ considerably.

The most immediate cause of anæmia is direct loss of blood either by a single large hæmorrhage or by successive small ones. A continuous anæmia may be kept up by bleeding piles, excessive menstruation, etc.

An occasional cause of anæmia is the **destruction of the red corpuscles** in the blood. It is mentioned above that the transfusion of alien blood leads to the solution of the corpuscles and a **Hæmoglobinuria**. This condition may be brought about, however, by the solution of the normal corpuscles. It is so to some extent in burns which involve extensive tracts of skin, also in pyæmia, in certain fevers, and, above all, in **Paroxysmal hæmoglobinuria**.

**Hæmoglobinuria** (*Hæmoglobinæmia*).—We have already considered at p. 94 the mode of occurrence of hæmoglobinuria when alien blood is transfused. It has been also produced artificially by the injection of distilled water, of biliary acids and diluted glycerine. Extensive burns of the skin, by destroying the vitality of the corpuscles in the vessels exposed to the high temperature, induce their ultimate solution. Certain poisons also partially dissolve the corpuscles, especially arsenic and arseniuretted hydrogen and nitric acid. It occurs in pyæmia, and occasionally in the specific fevers, such as typhus and typhoid fever. It occurs also in new-born children. (See under Icterus.)

**Paroxysmal hæmoglobinuria** is a condition in which the red corpuscles are peculiarly susceptible of solution, which is induced by slight causes. The commonest cause is exposure to cold, but a case is recorded in which it always occurred in a soldier after a long march. As exposure of the surface to cold is the usual cause, the affection occurs mostly in winter, but Rosenbach produced it in summer by a cold foot bath. The hæmoglobin is set free by the solution of the corpuscles. Ehrlich, in a case of this kind, put an elastic ligature round the finger and immersed the latter in ice-cold water. A drop of blood from the finger showed the corpuscles in various stages of decolorization, and also altered in size and shape (microcytes and poikilocytes). The urine in hæmoglobinuria gives a precipitate on heating like that of albumen, but this is hæmoglobin. The hæmoglobin is found in the urine in the form of little beads, often in rows, and sometimes forming casts of the tubules.



The free hæmoglobin may stain the tissues, but this does not usually occur unless the hæmoglobin is transformed into hæmatoidin. The so-called hæmatogenous icterus, observed in pyæmia and other conditions, has this origin. (See under Icterus.) Recklinghausen found crystals of hæmatoidin in the blood in a case in which lamb's blood had been used for transfusion.

There is also sometimes a destruction of red corpuscles in ague, but the colouring matter here is not in solution but in solid granules, constituting the condition of melanæmia. (See under Pigmentary Infiltration.)

In contradistinction to these cases of direct loss or destruction of the blood, we have cases in which the cause is very obscure, and which are designated by the term **Spontaneous anæmia**. In one of these, chlorosis, some have asserted that the organs of generation are defective, but Virchow has found in some cases structural lesions of the vascular system (see below). In another, namely essential anæmia or pernicious anæmia, we have no doubt several different conditions grouped together, of which the causation is obscure, although, according to Hunter and others, there is in this disease a destruction of red corpuscles chiefly in the portal vessels, probably induced by a poison absorbed from the intestinal canal.

**Secondary anæmias** are those which depend on some defect in the general conditions of life or on some grave disease which either causes a waste in the constituents of the blood, or, by lowering the nutrition as a whole, leads to defect in the blood.

**Character of the lesions.**—From what has gone before it will be apparent that in the different forms of anæmia the state of the blood will vary considerably, and it will be proper to refer to each individually. The condition of the red corpuscles is the most important point. Their numbers may be estimated by one of the forms of Hæmacytometer founded on the method of Malassez, and the amount of their hæmoglobin may be determined by the Hæmoglobinometer, in which the depth of colour is compared with a standard solution.

In **Anæmias due to hæmorrhage**, or in those in which a direct destruction of the red corpuscles has occurred, the number of the corpuscles is reduced, and the plasma is watery.

Immediately after a severe hæmorrhage we have an actual **Oligæmia**, but this simple reduction in the bulk of the blood does not long remain, as various processes ensue which modify the condition of the blood. In the first place, the bulk of the blood is rapidly made up by absorption from the tissues and alimentary canal, and by diminution of the excretion of water by the kidneys. The result will be that, while the bulk of the blood is made up, the plasma is watery and the red corpuscles deficient, a condition of **Oligocythæmia** and **Hydræmia**. In the blood, soon after a hæmorrhage, there is a certain proportional excess of white corpuscles, a



**Leucocytosis.** This arises partly from the fact that the leucocytes, being more adhesive than the red corpuscles, do not escape so readily as these, and also because the leucocytes are more rapidly renewed than the red corpuscles.

If there has been a single hæmorrhage the blood is gradually, although slowly, restored. The plasma comparatively soon recovers its due concentration, but the red corpuscles are very slowly replenished, and it may be weeks or months before their numbers are made up. The condition of the blood itself will interfere with the activity of the blood forming as well as other organs. Where there are repeated, although small, hæmorrhages, a more or less permanent hydræmia and oligocythæmia result.

**Pernicious anæmia**, or essential anæmia, is a condition in which, without apparent cause, the blood becomes progressively deteriorated. In some cases which have during life presented the features of this disease, post-mortem examination has revealed organic disease of the intestinal canal, such as a deep-seated cancerous tumour, or (according to Nothnagel and Fenwick) an induration of the stomach with atrophy of the gastric glands. There remains a considerable number of cases in which the cause is quite obscure, the blood becoming progressively more defective in red corpuscles, till a fatal issue results.

The observations of Quinke, Peters, Russell, and others, which have been confirmed and elaborated by Hunter, show that in pernicious anæmia there is a great excess of pigment, containing iron, in the liver. The iron is demonstrable by micro-chemical tests, sulphide of ammonium giving a dark colour, and ferrocyanide of potassium, with dilute hydrochloric acid, a beautiful blue (see Hunter). The pigment is present chiefly in the outer two thirds of the hepatic lobules, and is contained in the hepatic cells. The cells in the central parts of the lobules usually show fatty degeneration. A similar pigment is sometimes present in the kidney, where it is mostly in the epithelium of the convoluted tubules. The bone-marrow also contains an excess of pigment in which iron is present. On the other hand, the spleen does not contain an excess of iron, although usually enlarged.

From these observations Hunter infers that in pernicious anæmia there is a great destruction of red corpuscles mainly in the blood of the portal circulation, the hæmoglobin not passing unchanged into the urine as in hæmoglobinuria, but being caught and stored, in an altered form, in the liver. He has found that a poison, toluylen-diamin, when injected into the blood of animals, induces a similar destruction of red corpuscles in the portal circulation and a similar accumulation of iron in the liver. In these experiments also there was no hæmoglobinuria, this being accounted for partly by the fact that the destruction of corpuscles takes place mainly in the portal circulation, and partly by the inference that the hæmoglobin is altered by the poison. On the analogy of this poison, it is inferred that pernicious anæmia is due to a poison absorbed from the intestine. The occasional co-existence of cancer of the intestine, or atrophy of the intestinal glands, agrees with this view, as these lesions may facilitate the absorption of the poison. The frequent co-existence of gastrointestinal symptoms is also in accordance with this view.



In this disease there is a marked deficiency in number in the red corpuscles; they have been found to number as few as 500,000, instead of 5,000,000 in the cubic millimetre. Although deficient in number, they are not usually defective in hæmoglobin. Besides this the red corpuscles present varieties in size and shape. In most cases there are to be detected in the blood during life, small, globular, deeply-coloured red corpuscles, which are evidently altered red corpuscles. These have been named **Microcytes**. (See Fig. 13.) There have also been observed red corpuscles of various forms (*Poikilocytes*). Nucleated red corpuscles are also occasionally present.

An almost constant concomitant of this disease is an alteration of the bone-marrow. In the long bones we have instead of the yellow adipose tissue, a semi-fluid red marrow, occupying the medullary cavity of the shaft. Under the microscope the normal adipose tissue is seen to be replaced by large granular cells, not unlike epithelial cells (*b* in Fig. 14), along with which nucleated red corpuscles are present, as well as red corpuscles of very various sizes.

This condition of the bone-marrow is by many regarded as secondary, a similar condition being present in other forms of anæmia, such as leukæmia. In leukæmia, however, the bone-marrow resembles lymphatic gland tissue, while here the cells are large and granular, not at all like lymph-corpuscles. There is not as some assert, merely an over-growth of lymphoid tissue, as in leukæmia, but a fundamental alteration of the bone-marrow whether primary or secondary.

The condition of the spleen varies considerably; sometimes it is enlarged, it may be greatly so, sometimes it is scarcely at all altered. There may be also in it variously altered red corpuscles, as in the bone-marrow. In some cases the lymphatic glands are enlarged.

**Chlorosis** is a form of anæmia which occurs in females usually about the time of puberty. The condition here is rather that of deficiency of hæmoglobin than of corpuscles, as the latter, although usually deficient in number, may not be so. The **achromatosis** may be such as to indicate that the hæmoglobin is reduced to a half or a fourth of the normal. The rapid recovery of cases of chlorosis under treatment with

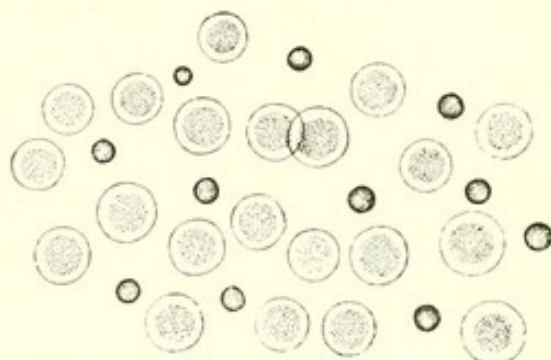


Fig. 13.—Blood in Pernicious Anæmia. The larger bodies are the normal red corpuscles. The smaller are the round more deeply coloured ones usually found—so called Microcytes. (EICHHORST.)

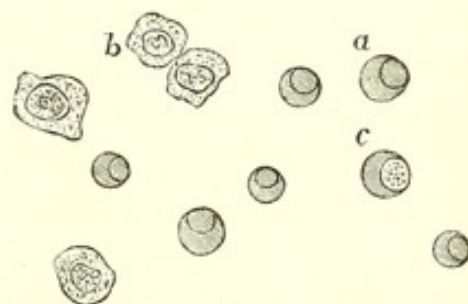


Fig. 14.—From Red Medulla of bone in Pernicious Anæmia. *a*, Nucleated red corpuscles. *c*, A red corpuscle with granular nucleus. *b*, Large nucleated cells, forming the bulk of the altered marrow.



iron is consistent with the fact that it is not so much replenishment of red corpuscles as of hæmoglobin that is needed.

The occurrence of chlorosis at puberty has suggested as its cause some defect in the sexual organs, and there are cases in which the uterus or ovaries have been imperfect (Rokitansky). But in the majority of cases this is not so, and it is more probable that the changes at puberty develop the condition by throwing an extra strain on the blood-forming organs. Virchow has observed in cases of chlorosis certain congenital defects in the vascular apparatus, the chief of which are narrowness and thinness of the aorta, irregularity in the origin of the branches from the aorta and smallness of the heart. The narrowness of the aorta may lead to compensatory hypertrophy of the heart in the same way as a narrowness of the aortic orifice. These lesions do not explain the chlorosis, but they suggest the existence of a congenital defect in the vascular system which may extend to the blood-forming organs, and may cause them to give way to the strain of puberty.

**Secondary anæmias** are generally due to grave exhausting diseases, such as phthisis pulmonalis, ulcerating cancers, Hodgkin's disease (which is also called **anæmia lymphatica**) leukaemia, fevers, albuminuria, etc. Malaria without manifesting an actual melanæmia often leads to anæmia.

The condition of the blood in secondary anæmia is usually that of hydræmia or hypalbuminosis with oligocythæmia; the blood is watery and deficient in red corpuscles, but the corpuscles remaining are generally of normal colour.

**Secondary organic changes in general anæmias.**—The altered condition of the blood commonly induces secondary changes, which are not peculiar to any form of anæmia but are liable to be more pronounced the greater the degree and the longer the duration of the disease. The condition of the blood interferes with nutrition and function, and the most definite changes are degenerative, and more particularly **fatty degeneration**. This manifests itself chiefly in the heart, but it also occurs in the walls of the blood-vessels. In chlorosis and other anæmias a yellow figuring of the intima of the aorta due to fatty degeneration is frequently observed. There may also be fatty degeneration of the epithelium of the kidneys (frequent in phthisis pulmonalis) and of the hepatic cells in the liver. There is also in some cases the change in the bone-marrow already referred to.

Experiments on dogs show that successive large hæmorrhages rapidly produce fatty degeneration of the muscle of the heart. The author has met with a case in man in which a single very severe hæmorrhage had led to a very pronounced fatty degeneration of the heart, having the usual characteristic naked-eye appearances of that condition (which see).

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Phys. chemie, 1881; GAMGEE, *Physiol. chemistry*, vol. i. *Hæmoglobinuria*—LICHTHEIM, Volkmann's Vorträge, No. 134; LESSER (Burns), *Virch. Arch.* vol. lxxix.; WINKEL, *Deutsch. Med. Wochenschr.*, 1879, No. 24; WICKHAM LEGG, *St. Barth. Hosp. Rep.*, vol. x. 1874; FORREST and FINLAYSON, *Glas. Med. Jour.*, 1879; PONFICK, *Berl. Klin. Woch.*, 1883, No. 25. *Pernicious Anæmia*—WILKS, *Guy's Hosp. Rep.*, 1857; *Lect. on path. anat.*, 1859; IMMERMAN, *Ziemssen's Handb.*; ZENKER, *D. Arch. f. Klin. Med.*, vol. xii.; EICHHORST, *Die progressive perniciöse Anämie*, 1878; BROADBENT, *Practitioner*, 1875; COHNHEIM, *Virch. Arch.*, vol. lxxviii.; WALDSTEIN, *Virch. Arch.*, vol. xci.; MUSSER, (*Pernic. Anæm. in America*) *Philadelphia Med. Times*, 1885; DICKINSON, *Path. Soc. trans.* vol. xxix., 1878; QUINCKE, *Deutsch. Arch. Klin. Med.* xxv., xxvii., xxxiii.; PETERS, *ibid.*, 1878, xxxii.; HUNTER, (*literature fully*) *Lancet*, 1888, vol. ii.; MACKENZIE, *ibid.*, 1878, vol. ii.; COUPLAND, *ibid.*, 1881, vol. i. *Chlorosis*—BECQUEREL, *Gaz. des hôpit.* 1856; SÉE, *Du Sang.*, etc., 1886; VIRCHOW, *Ueber die Chlorose*, etc., 1872; WILLOCKS, *Lancet*, 1881; ROKITANSKY, *Lehrb. d. path. Anat.* vol. ii.

#### LEUKÆMIA. LEUCOCYTHÆMIA.

These terms mean literally white blood and white-cell blood, and they express a condition in which the blood is light in colour, from the fact that the white corpuscles are in great excess. The excess of leucocytes is expressed by the term *Leucocythæmia*, originally applied by Bennett, but now generally discarded for Virchow's term *Leukæmia*.

**Causation.**—This is entirely obscure. There have been a few cases in which an injury to the spleen is supposed to have been the starting-point, while others have been ascribed to syphilis, rickets, malaria. The disease presents many features which suggest analogies with infective diseases, and the analogy of the tissue lesions with those in Hodgkin's disease, which belongs to that class, is consistent with this view. Cases have been recorded in which pernicious anæmia developed into leukæmia (Waldstein). The disease is twice as frequent in males as in females.

**Character of the morbid changes.**—The normal proportion of white corpuscles to red in the blood is stated as about 1 in 300 to 1 in 450, but it may vary within normal limits. There is after hæmorrhage, as we have seen, a slight increase in the proportion of white corpuscles, a *leucocytosis*, but it is of no special significance. There is also an excess of white corpuscles in many infectious and infective diseases, as pyæmia, erysipelas, relapsing, typhoid, and intermittent fevers, etc. In leukæmia however, the relative proportion of white to red corpuscles is greatly altered. A case is not a very severe one in which the corpuscles are as 1 to 10, and they may be as 1 to 2 or even equal.

This alteration in the proportion of white and red corpuscles, is not the only alteration in the blood. The white are more varied in size than normal, some larger and some smaller, and they are sometimes more distinctly nucleated. Then the red corpuscles are not only



proportionately few in number, but they are absolutely reduced, and that to a very high degree. Occasionally nucleated red corpuscles have been found in the blood, these corpuscles being of larger size than the normal red corpuscles. The specific gravity of the blood is considerably reduced; it may fall from 1055, which is the average normal, to 1040 or 1035. The blood has thus **the characters of anæmia** along with a great increase of the white blood corpuscles.

In leukæmic blood, after death, there are found small, colourless, glancing crystals of an octahedral shape, which are usually called from their discoverer **Charcot's crystals** (Fig. 16). There are also certain other abnormal chemical constituents, chiefly glutine and hypoxanthine.

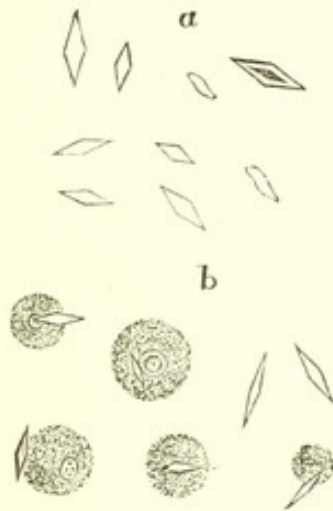


Fig. 15.—Charcot's crystals from the blood in leukemia after death. In *b* the crystals are partly inside the white corpuscles. (ZENKER.)  $\times 500$ .

With regard to the probable explanation of this peculiar condition of the blood, it must always be remembered that there is a very great diminution in the red corpuscles as well as a great increase in the white. There must be either a greatly increased newformation of white corpuscles and destruction of red ones, or a serious interference with the conversion of the white into the red corpuscles. This latter is the view originally taken by Virchow, and is the more probable. There may be an increased formation

of white corpuscles, but the diminution in the number of the red is probably due to a reduced conversion of white into red. In the existing obscurity as to the origin of the red corpuscles, this statement cannot be made unequivocally. The nucleated red corpuscles have been regarded by some as transition forms, their presence indicating a delay in the conversion.

The blood in persons affected with this disease is unduly pale, but it has not a watery appearance. It is even a thicker fluid than usual and distinctly more opaque, resembling a mixture of pus and blood.

**Tissue changes in leukæmia.**—The changes in the blood are associated with lesions in the tissues, some of which are generally regarded as primary, and some as secondary, although authors are not wanting who regard all the tissue changes as secondary to those in the blood. The primary changes affect the organs which are generally believed to have to do with the formation of the blood corpuscles, namely, the spleen, the lymphatic glands, and the bone-marrow. Attempts have been made to divide the cases, according as the lesions have been chiefly in one or other of these organs, into a splenic, a



lymphatic, and a myelogenic form, but this division can hardly be carried out, especially as the bone-marrow seems to be affected in nearly all cases.

The **Spleen** is affected in most cases, and when affected it is in all stages enlarged. The enlargement in the earliest period appears to be due to an active congestion, and is accordingly of rapid development. Rindfleisch mentions a case which he saw at Virchow's demonstrations in which the enlargement had been so rapid as to cause a rupture of the capsule of the spleen. This enlargement is merely from overfilling of the vessels, but it is succeeded by, or develops into, a solid enlargement. The Malpighian bodies of the spleen have the structure of lymphatic follicles, and it is mainly by the enlargement of these that the permanent solid enlargement occurs, a great increase, therefore, of lymphoid tissue. The spleen thus becomes converted into a hard, dense, bulky organ, sometimes like a piece of wood. It is also paler than normal, and we can often see the enlarged Malpighian bodies as whiter areas on the cut surface. In addition, there are frequently hæmorrhages in the spleen, and these may take the form of the regular wedge-shaped infarction. The capsule of the spleen is greatly thickened, and there are not infrequently dense localized thickenings of a cartilaginous consistence. The capsule is often firmly adherent to the diaphragm and other neighbouring structures. The greatly enlarged, dense, and heavy organ is frequently dislocated downwards by its own weight.

In the **Medulla of bone** the changes are very similar. There is a great newformation of round-cell tissue which occurs at the expense of the adipose tissue and also of the bone. This new tissue consists mainly of cells as large as white blood-corpuscles, and has the structure of lymphatic tissue, showing the reticulum and round cells as in Fig. 16. Sometimes there are larger cells than these, and there may be also nucleated red corpuscles. There is thus a great newformation in the medulla and that mainly of lymphatic tissue. As the medulla of bone, especially the red medulla of spongy bone, normally contains lymphatic tissue, the change in leukæmia may be regarded as a hypertrophy of this, and the amount of hypertrophy is sometimes very great. Thus the whole yellow marrow of the hollow shaft of the femur may be replaced by lymphatic tissue. The cavity of the shaft may also be widened at



Fig. 16.—Medulla of bone in leukæmia, from the cavity of the shaft of the femur. Adenoid reticulum and lymphoid cells are shown.  $\times 350$ .



the expense of the bone, and many smaller trabeculae here and in the spongy bone may be destroyed. In this respect as well as in intimate structure the leukæmic bone-marrow differs from the red marrow of pernicious anæmia, as the latter does not cause a proper atrophy of the bony tissue. The occurrence of hæmorrhages or even infarctions has been observed in the bone-marrow just as in the spleen.

The appearance of the medulla is, of course, greatly altered. It has generally a greyish red colour without any of the normal greasy appearance, but it may be pale so as to look like solidified pus, or dark red, these differences depending on the degree of the newformation and the state of fulness of the vessels. It has, however, the character of a tolerably solid tissue, not a semi-fluid material as in anæmias. These changes may occur in all the bones in the body, and in both spongy and hollow bones, the affection being, however, variously distributed.

The lesion in the **Lymphatic glands** consists in an enlargement of them. This begins mostly in a particular set of glands usually situated externally, as in the axilla, groin, neck, etc., and spreads to other sets, generally first to those nearest. The enlarged glands may be three, five, or even ten times their normal size, but there is no tendency to any degeneration of their tissue.

It will be observed that in all these primary lesions there is a newformation, or a hyperplasia of existing lymphatic tissue. In the **Secondary lesions** there is sometimes also a newformation of lymphatic tissue, partaking more of the characters of tumour-formation, as it may occur in places where there is normally no such tissue present, in which case the term lymphoma is sometimes applied to the newformation. More frequently, however, there is merely an infiltration of the connective tissue of organs with leucocytes. These may have been derived from the blood, but they appear to multiply *in situ* as Bizzozero has observed evidences of division (karyokinesis) in the cells.

The **Liver** is nearly always enlarged, and its connective tissue is occupied by myriads of collections of round cells. These are often in the form of a general infiltration of the capsule of Glisson, attaining to greater intensity at certain points, but it may be in such definitely localized areas as to suggest minute tumours. The collections of cells resemble, on the one hand, miliary tubercles, and, on the other, inflammatory infiltrations (as in cirrhosis), but they differ from the former in respect that they have no tendency to degeneration, and from the latter in that they do not form connective tissue.

The **Kidneys** are not infrequently affected, and here the appearance



to the naked eye is often as if the organs were greatly enlarged by the presence of large pale tumours in the cortical substance. On microscopic examination the lesion is seen to consist in an enormous infiltration of the stroma of the kidney with round cells, the proper secreting tissue remaining, but, of course, greatly pressed on. This infiltration occurs in definite areas, as if some agent had addressed itself to certain defined portions of the organ.

The **Closed follicles of the intestines**, both the solitary ones and those aggregated in Peyer's patches, may be enlarged, and as they are of lymphatic structure, their enlargement is a simple hyperplasia. This is not of frequent occurrence, and still less frequent is the formation of leukæmic tumours in the skin, these tumours consisting of infiltrations of round cells.

The connective tissue in other regions may also be infiltrated. The author met with a case in which the connective tissue of the mediastinum was enormously infiltrated, so that the tissue formed a bulky tumour. The infiltration extended to the pericardium much in the fashion of Hodgkin's disease.

It is proper here to mention that in **Hodgkin's disease** the organic lesions somewhat resemble those of leukæmia, but without the increase of white blood corpuscles, the blood being simply anæmic. There is great enlargement of the spleen and of the lymphatic glands, consisting as in leukæmia in a newformation of lymphatic tissue.

**Literature.**—VIRCHOW, *Froriep's Notizen*, 1845, *Gesam. Abhandl.* p. 190; BENNETT, *Ed. Monthly Journal*, 1850-51, and *Leucocythæmia*, 1852; TROUSSEAU, *Gaz. des hôp.*, 1858; MOSLER, *Die Path. d. Leukämie*, 1872, and *Virch. Arch.*, vol. lxxv; NEUMANN, *Myelogene Leukämie*; *Berl. Klin. Wochenschr.*, 1878, No. 6; WALDSTEIN, *Virch. Arch.*, vol. xci.; BIESIADECKI, *Wien Med. Jahrb.*, 1876, p. 234; PONFICK, *Virch. Arch.*, vol. lxxvii; BIZZAZERO, *Virch. Arch.*, vol. xcix. ZENKER, *Arch. f. Klin. Med.* xviii.

#### ABNORMAL CONSTITUENTS IN THE BLOOD.

We include here conditions in which, with or without alterations in the corpuscles or blood plasma, substances are present in the blood other than its normal constituents. These substances are either suspended or dissolved in the plasma.

1. **Melanæmia.**—By this name is meant, literally, black blood. It is used to designate a condition in which pigment occurs abnormally in the blood and is deposited in the tissues. The pigment is in the form of solid granules, and, although black in colour, it is derived from the blood pigment, and contains iron. It is met with in cases of malarial fever, but not in every case, as a rule only in the more severe forms. The view usually entertained is that during the acute paroxysm of an ague, blood-corpuscles are destroyed in the spleen, and their pigment is set



free in the granular form. Lately, Arnstein has brought forward the view that the destruction of the blood-corpuscles occurs not in the spleen, but in the blood generally. Whether the spleen be the seat of its formation or not, the pigment is certainly stored up there for years, and may pass into the blood by degrees, although it is more abundant immediately after the acute paroxysm.

When the pigment gets into the blood it is rapidly taken up by the white blood-corpuscles, so that these appear in the blood as pigmented

cells. These pigmented cells are often larger than normal white corpuscles, being enlarged by their abnormal contents. It will thus be seen that very little free pigment will be present in the blood except immediately after the acute paroxysm, the white corpuscles picking up the solid granules just as they do when a solid pigment, such as vermilion, is artificially introduced into the blood of an animal (see Fig.

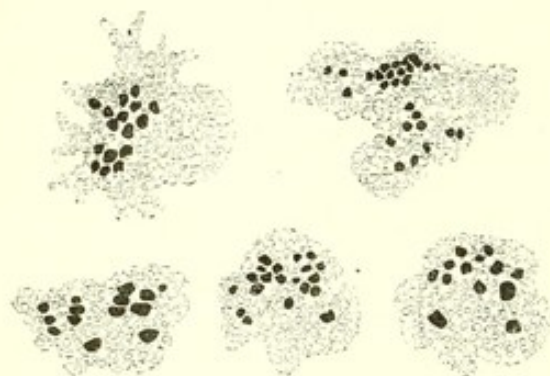


Fig. 17.—White blood-corpuscles of the frog, containing granules of vermilion, and showing amoeboid movement. (After KLEIN.)

17). The white corpuscles containing the pigment accumulate in certain organs, especially the spleen, liver, and bone-marrow. In the spleen and bone-marrow the pigmented cells appear to leave the capillaries readily, and pass into the tissues, producing an actual pigmentation of them, but in the liver they linger longer in the capillaries, which may be seen with many such corpuscles in them (see Fig. 18), although here

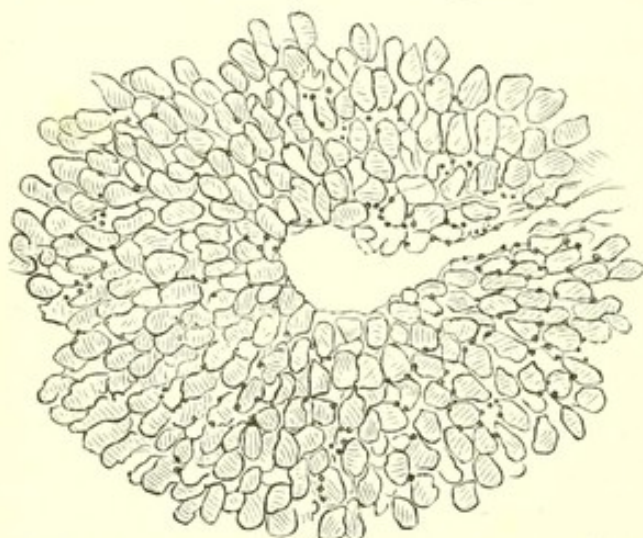


Fig. 18.—A lobule of the liver in melanæmia. The black pigment granules are seen between the hepatic cells, being really in the capillaries.  $\times 75$ .

also the latter may pass out into the surrounding tissue. In very severe cases, other organs and tissues may be pigmented in this way. The brain is very often pigmented, especially in cases where there has been much cerebral excitement. This will be in very acute cases, but it is doubtful whether the pigment has much to do with the cerebral symptoms, which may be merely a consequence of the acute attack. The kidneys are

sometimes the seat of it as well as other tissues, such as pancreas, intestine, etc.



This abnormal pigmentation of course produces changes in the colour of the organs affected. The spleen is slaty-grey or almost black, the liver is steel-grey or blackish, the grey substance of the brain is of a dark chocolate or graphite colour, and the kidneys present greyish spots.

**Literature.**—ARNSTEIN, *Virch. Arch.*, vol. lxi., p. 494; MOSLER, *Virch. Arch.*, vol. lxi., p. 369.

**2. Lipæmia. Piarrhæmia.**—These names designate a condition in which fat is abnormally present in the blood.

There is a certain quantity of free fat in the blood normally, and after a meal it may be somewhat abundant. In lipæmia, however, there is so much fat present as to give the blood a peculiar milky appearance to the naked eye. Under the microscope the fat is visible as fine fat drops,

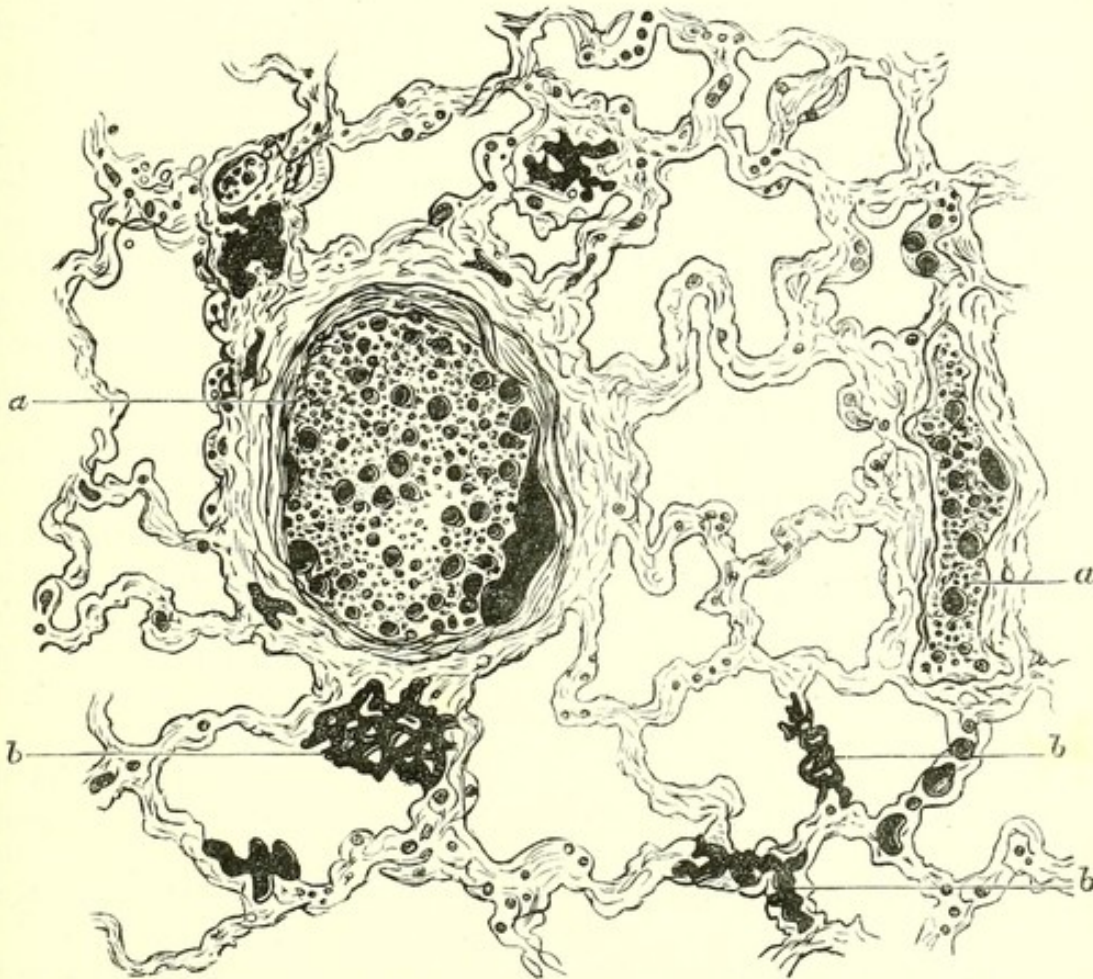


Fig. 19.—Lung in Lipæmia, the fat stained black with osmic acid. *aa*, Fat in pulmonary artery, and *bb*, in capillaries. (SAUNDERS and HAMILTON.)

which are free or else enclosed in cells. The fat may be present (in some cases of diabetes) in such quantity as to alter the entire appearance of the blood. Thus the blood in the vessels will present opaque white portions where the fat has come to the surface like cream. The fat is generally finely divided, but it may be caught in the fine arteries and capillaries of the lungs (see Fig. 19), the brain, and the kidney. It is



readily detected in the glomeruli of the kidney, and it may pass thence into the urine, constituting a form of chyluria.

Lipæmia has been met with in its most aggravated form in diabetes, but it also occurs as a consequence of alcoholism, in some cases of dyspnoea, and in relapsing fever. The source of the fat is obscure. By some it is believed that it is the fat of the food which is not oxidized and so accumulates, but this is not a probable explanation. A more probable one is that it arises by fatty degeneration of the endothelium of the blood-vessels in the spleen and elsewhere, and perhaps it does so in some forms of lipæmia. The enormous amount of fat found in some cases of diabetes, however, precludes the idea that it can be derived from a fatty degeneration of the endothelium. In some such cases, almost the entire blood has a milky appearance. The only probable explanation of this seems to be that the sugar which is present in the blood in diabetes in large quantities has been replaced by or converted into fat.

Ponfick found, in cases of relapsing fever, cells in a complete state of fatty degeneration in the blood, especially in that of the splenic and portal veins. These cells were endothelial cells of blood-vessels, which in consequence of becoming fatty were readily separated and carried off by the blood. The author observed in a case of diabetes with extreme lipæmia a very striking fatty degeneration of the endothelium of the splenic pulp, so that the organ as a whole presented an opaque pinkish appearance suggestive of salmon roe. The fatty degeneration was observed in the endothelium of the capillaries of the liver and elsewhere, but not to such an extent as in the spleen. In another case these conditions were present but less marked. In these cases, however, there was no appearance in the blood of free cells presenting fatty degeneration. Looking to the fact that in kidneys, spleen, brain, liver, lungs, and elsewhere, the blood was occupied by very finely divided fat, the only probable conclusion seems to be that a constituent of the liquor sanguinis had become converted into fat; the only constituent capable of such a conversion is the grape sugar.

Hamilton and Saunders suggested that the coma which frequently marks the fatal termination of diabetes, may be due to the accumulation of fat in the blood. One cannot but believe that the lesion must have an important effect on the function of the blood, but there is sufficient evidence to show that diabetic coma often occurs independently of lipæmia.

**Literature.**—SAUNDERS and HAMILTON, *Edin. Med. Jour.*, July, 1879; CHRISTISON, *Edin. Med. and Surg. Jour.*, vol. xxxii., 1830; HUSS, *Der chron. Alkoholismus*, 1852; LANCEREAUX, *Traité d'anat. path.*, vol. ii.; PONFICK, *Virch. Arch.*, vol. lx., p. 166, 169; COOTE, *Lancet*, Sept., 1860. See also under Diabetes.

**3. Uræmia and Lithæmia.**—These terms designate conditions in which two constituents of the urine, namely, urea and uric acid or its salts, are present in excess in the blood. Uræmia is more a clinical than a pathological term, however, and is used to distinguish a group of symptoms whose relation to disease of the kidneys will be considered further on.



Lithæmia, or an excess of uric acid or urates in the blood is believed to have to do with the causation of gout, in which disease there is a deposition of these salts in the tissues, chiefly of the joints and kidneys.

4. **Diabetes mellitus.**—This term properly designates an excessive flow of urine containing sugar, but as the sugar passes into the urine because it is present in the blood the disease is really one in which the blood contains sugar in excess. The terms **Melituria** and **Glycosuria** are equivalent in their meaning, but are generally used to designate a temporary condition, generally symptomatic of some other disease, while diabetes is a disease by itself. The term **Glycosæmia** is sometimes used to designate the condition of the blood. The substance thus present abnormally is grape sugar or **glucose**.

The blood and urine contain normally a small amount of sugar, which is believed to have its chief source in the glycogen of the liver. This latter substance is formed by the hepatic cells, this being regarded by many as one of the most important functions of these cells. The uses of glycogen, which is found in the muscles and other tissues besides the liver, both in the foetus and adult, are quite obscure. It can scarcely be believed that its only use is to supply sugar for burning, and its presence in muscle suggests that it may have to do with the production of muscular force. It obviously subserves some important function, whatever that may be. In diabetes the glycogen in the liver is converted to an abnormal extent into sugar. The sugar being a crystalloid while the glycogen is a colloid substance, the former readily diffuses into the blood and passes into the urine. But this is not all. There is an excessive production of glycogen which, becoming transformed into sugar, gives rise to an excessive discharge of sugar, which may amount to 25 ounces in the 24 hours.

The whole process is probably traceable to the abnormal conversion of glycogen into sugar. If we believe that glycogen itself subserves an important function, then this function will suffer by the conversion of the glycogen into sugar; there will be a deficiency of glycogen. To supply this deficiency the liver will be stimulated to increased action, and there will be increased formation of glycogen whose transformation leads to excessive production of sugar. We may thus explain the progressive nature of the disease and the extraordinary quantities of sugar which are produced.

In the normal condition the sugar in the blood is at a constant minimum, unaffected by the amount or kind of food taken. In diabetes there is commonly a very obvious **relation between the food taken and the sugar**. There are some diabetics who cease to excrete any excess of sugar so long as they abstain from starch or sugar, but whenever they take any such food it is mostly converted into grape sugar and so



excreted in the urine. On the other hand, there are diabetics who excrete an excess of sugar whatever the kind of food they take, and though the sugar may be diminished by the use of a mainly nitrogenous diet, yet it cannot be thus made to disappear. In that case it is obvious that sugar is formed not only from carbo-hydrates, but also from the albuminous principles of the food. According to Traube, diabetes is divisible into a slighter and a more severe form according as the sugar is formed from the carbo-hydrates alone, or from nitrogenous principles as well, and it is probable that these forms represent an earlier and a later stage in the disease. Donkin, in view of the relation of the sugar production to the food, distinguishes three stages; first, that in which only the starch or sugar of the food furnishes the grape sugar; second, that in which fats as well as these are converted; and third, that in which albuminous as well as starchy and oily foods undergo conversion into grape sugar.

It is obvious that in this latter stage there is a great **Consumption of albumen**, and, in the process of formation of sugar, urea is formed as a waste product in the chemical transformation. We find, therefore, that the **Urea** is also greatly increased in the urine, amounting sometimes to two or three times the normal quantity in the twenty-four hours. In this stage of the disease there is no doubt that the fatty and **Albuminous constituents of the tissues** are also used for the formation of sugar, and that the albuminous principles of the tissues, like the albumen of the food, furnish sugar and urea which appear in the urine. We may believe that these various constituents are carried to the liver, there to undergo a conversion, first into glycogen and then into sugar.

In the normal living body the liver, apparently, contains glycogen and not sugar. If the liver of an animal be cut out immediately after death and without delay placed in boiling water after being cut into small pieces, then it will be chiefly glycogen that will be found, and any sugar that exists has probably formed after death by the transformation of the glycogen. It is difficult indeed to avoid the occurrence of traces of sugar in this experiment, and if the removal of the liver be delayed a large amount of sugar will be found. These facts show that glycogen is always just ready to be converted into sugar; its conversion is, as it were, every moment imminent. During life this conversion does not take place to any considerable extent, and it is even doubtful whether the small amount of sugar existing normally in the body is due to a conversion of the glycogen in the liver, a comparison of the blood in the hepatic vein with that in the portal giving doubtful results.

The **Experimental production of glycosuria** throws some light on the pathology of diabetes. Bock and Hoffmann succeeded in the production of an **Artificial diabetes mellitus** by injecting into the blood of rabbits large quantities of a watery solution of common salt. The animals, soon after the injection was made, began to secrete a large quantity of urine, and this urine soon became saccharine. If the injection of salt solution was persisted in, the sugar by and by diminished in the urine and ultimately disappeared. It was made clear by examination that the sugar in the urine came from the liver and was caused by the conversion of the glycogen into sugar. In all cases where the animal was killed after the diabetes had passed off the liver was found free both of the glycogen which exists normally and of sugar. If, on the other hand, the animal was killed while the melituria existed, then glycogen and sugar were both present in the liver. The inference from these observations is perfectly obvious. The abnormal condition of the blood causes the transformation of the liver-glycogen into sugar, and the latter being a crystalloid and readily



diffusible, it is at once washed out of the hepatic cells and passes into the circulation and on into the urine. The diabetes ceases because all the glycogen in the liver has undergone conversion into sugar and the source of supply is exhausted, the liver in these cases being found free of both glycogen and sugar.

It may be said that the diabetes thus produced is temporary and not comparable with the permanent and progressive disease in the human subject. But if there is a permanent cause leading to the conversion of the glycogen into sugar, then the consequences of increased formation would follow in due course. The liver will be stimulated to an ever-increasing formation of glycogen which will immediately pass into sugar. We can understand how in the earlier stages all foods which are near to glycogen in their chemical composition will be at once utilized, and how as the disease goes on and the demand becomes more urgent, the other kinds of foods and even the tissues of the body will be used.

So long therefore as the hepatic cells retain their activity we may presume that they will react to the hunger for glycogen, and the production of sugar will be the result. If the hepatic cells be destroyed or weakened the probability is that the diabetes will diminish, the disease requiring that the hepatic cells retain their energy.

It is difficult to account for the abnormal conversion of the glycogen into sugar. The experiments of Bock and Hoffmann seem to indicate that an alteration in the character of the blood circulating in the liver has this effect. In other experiments in which glycosuria is produced artificially there is also presumably an alteration in the circulation in the liver. The nature of the alteration may be summed up in the statement that the blood reaches the hepatic capillaries in a condition approaching that of arterial blood instead of venous blood, as under normal conditions.

The inhalation of nitrite of amyl induces a temporary glycosuria. We know that this agent produces a general vaso-motor paralysis, and a general dilatation of the systemic arteries. With the other arteries those of the abdomen will dilate, and the blood passing through the capillaries at an increased rate will reach the portal vein at a higher pressure than normal. The circulation in the liver will therefore be accelerated and the blood will be less venous in character, having passed rapidly through the capillaries of the intestines and other organs.

Bernard produced diabetes by puncturing the medulla oblongata in the floor of the fourth ventricle, and we know that in this part of the nervous system are situated the principal vaso-motor centres of the body. There is therefore here also a paralysis of the arteries just as after the inhalation of nitrite of amyl. Injuries to the brain, spinal cord, and sympathetic sometimes produce a temporary glycosuria, and they also may cause vaso-motor paralysis.

The observations of Pavy are important in this connection. He produced glycosuria by the injection of defibrinated arterial blood into the portal vein, and here again there is an excessive supply of blood to the liver. But then Pavy also produced glycosuria by ligaturing the portal vein, and so cutting off the blood supply except through the hepatic artery; at first this result seems a very contradictory one.

The effect of that experiment would be to cause the liver to be supplied with



arterial blood alone, and we may presume that the hepatic artery would dilate and so allow of an additional flow of blood. According to the researches of Cohnheim and Litten the blood of the hepatic artery, after supplying the connective tissue, gall ducts, and walls of the large blood-vessels, passes into the inter-lobular veins and on into the proper hepatic capillaries, so that the blood of the hepatic artery is finally distributed with that of the portal vein. When the portal vein is closed, the circulation in the liver will be kept up, but only by the blood of the hepatic artery.

From these observations Pavy concluded that the cause of the glycosuria when the portal vein was ligatured was the circulation of arterial blood in the liver, and he proceeded to determine whether it was possible in animals to produce diabetes by supersaturating the whole blood with oxygen, so that the blood in the portal vein would be virtually arterial. He effected this in various ways; by causing the animals to inhale oxygen, or by using artificial respiration till the fact that the animals ceased to make any spontaneous respiratory movements showed a deficiency of carbonic acid. In these ways glycosuria was produced. It would appear therefore that when diabetes is artificially produced in animals, the one essential condition is that the liver should be supplied with unduly oxygenated blood, such blood causing the glycogen to be transformed into sugar in the liver itself.

Many endeavours have been made to connect experimental glycosuria with actual cases of diabetes. The central nervous system has been scrutinized with very uncertain results, and evidences have been sought of vaso-motor disturbances in the liver. It is to be remembered that the portal vein, having little tonicity, is scarcely liable to great variations under different vaso-motor impulses. The circulation in the portal vein depends on that in the coeliac and mesenteric arteries. When these are dilated the blood will pass more rapidly through the capillaries of the organs which these arteries supply, and will reach the portal vein at a higher pressure and in a condition more nearly approaching to arterial blood than usual. If diabetes be owing to vaso-motor disturbances, therefore, we must look for these in the arteries and capillaries of the alimentary canal rather than in the liver itself.

The most direct evidence of the dependence of diabetes on a nervous lesion is afforded by the fact that a considerable number of cases are now on record in which diabetes was associated with disease of the head of the pancreas. The immediate connection of the head of the pancreas with the coeliac plexus and semilunar ganglion at once suggests the extension of disease to the latter. Hale White again has recorded several cases of diabetes with disease of the sympathetic, and especially of the semilunar ganglion. Pavy has also described a case in which injury to the superior cervical ganglion has led to diabetes.

The forms of disease of the pancreas with which diabetes has been found associated are—cysts, formation of calculi, cancer, atrophy. In cases observed by Klebs there was, along with the atrophy of the pancreas, a very marked change in the coeliac plexus consisting of a great destruction of ganglion cells. It seems very probable, considering the nearness of the pancreas to the coeliac plexus, that a



cancer, or a large cyst, or the changes resulting from the formation of calculi in the ducts of the gland, might exercise pressure on the cœliac plexus, and Klebs accounts for the diabetes in his cases on the view that there was vaso-motor paralysis produced by the destruction of the ganglion cells of this plexus. In one of his cases, indeed, there was observed after death an extraordinary dilatation of the hepatic and splenic arteries and the gastric branches of the cœliac axis, the last named branches attaining the size of a goose quill and presenting a highly convoluted course. These facts seem to confirm the view that in man diabetes may be produced by paralysis of the arteries in the domain of the cœliac plexus.

There have been also cases in which diabetes seemed to follow injury to the spinal cord or medulla oblongata, but these have been cases of temporary glycosuria comparable to that following Bernard's diabetic puncture. On the other hand there have been a few instances of softening of the brain and intra-cranial tumours in cases of diabetes, but these have been too rare to be regarded as of much importance in the pathology of diabetes. For the most part there is little or no alteration visible to the naked eye in the central nervous system.

Recently Dickinson has described minute changes in the central nervous system, consisting mainly in excavations around the arteries, produced apparently by exudation from these vessels with disintegration of the brain substance around, but his observations have not been confirmed, and the changes are probably to be regarded rather as secondary than primary.

In the abdominal organs themselves, there have been in many cases indications of congestion observed. We have already referred to the enlargement of the arteries in one of the cases described by Klebs. The liver itself is generally described as congested, and in a few cases there has been thrombosis of the portal vein.

The characteristic red tongue of advanced diabetes has been cited by Pavy as evidence of a progressive congestion of the alimentary canal. The congestion at first is confined to the abdominal organs, but in the later periods extends to the mouth, the vaso-motor paralysis being progressive.

Most of the known facts may be held as indicating that diabetes depends on the liver being supplied with blood of an abnormal character and at an accelerated rate of speed. The congestion of the liver depends on abnormal nervous arrangements, and these may be local, affecting the cœliac plexus, or possibly situated in the central nervous system.

In regard to the other changes in diabetes mellitus, little has to be said. The tissues appear to be peculiarly vulnerable; wounds heal badly, inflammations are generally very severe, often going on to supuration and gangrene, phthisis pulmonalis often closes the scene, and is frequently very acute. General emaciation is a striking feature in advanced cases. The kidneys may be found enlarged and their epithelium fatty, but this has no direct connection with the essential



pathology of the affection. Lipæmia, as we have seen, sometimes occurs, but only in a small percentage of cases.

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## SECTION V.

## INFLAMMATION.

**ETIOLOGY.** *Produced by irritants, whose character is to injure the tissues ; influences of nervous system, etc. ; mode of entrance of irritants.* I. **PRINCIPAL PHENOMENA,** *exhibited in an experiment ; (1) State of vessels—(contraction not essential) dilatation with active hyperæmia, followed by retardation ; explanation of these ; paving of veins. Increase of temperature, not due to heat production but to excess of blood.* II. **THE INFLAMMATORY EXUDATION.** (1) *the serous and (2) the fibrinous exudation, (3) exudation of white corpuscles by emigration and red by diapedesis, (4) cells from other sources, (5) the purulent exudation, suppuration due to pyogenic microbes ; the abscess ; (croupous and diphtheritic exudations).* III. **CHANGES IN THE TISSUES.** (1) *Parenchymatous changes ; (2) Newformation of tissue, chiefly after the type of the granulating wound ; the formative cells and their origin ; newformation of blood-vessels and of epithelium ; newformation around foreign bodies.* IV. **THE ISSUES OF INFLAMMATION.** V. **THE FORMS OF INFLAMMATION.**

**T**HIS is a subject whose importance in pathology can hardly be overestimated. From the time of John Hunter its phenomena have been taken as the basis of various speculations on the nature of pathological phenomena in general. It is impossible to give a strictly accurate and complete **Definition** of inflammation. The oldest and still most generally received definition distinguishes it as characterized by the four cardinal signs, redness, swelling, heat, and pain (*rubor, tumor, calor, dolor*). These signs, however, by no means apply universally, and they are typical rather of the early stages of inflammation than of the later ones.

**Etiology of Inflammation.**—In many cases the causes of inflammation are obscure, and this applies mainly to internal organs whose actions are greatly hidden from our view. Inflammation is **producible artificially**; and if we study the modes of production we shall find that agents are used to which the general name of **Irritants** is applied. The inflammation is produced, it is said, by irritating the part in some way. This name irritant is apt to be misleading, as it embodies the conception of a stimulating action ; whereas the so-called irritants, in their nature and action, possess characters rather of a deadening



than of a stimulating kind. They are agents which, when acting strongly, are calculated to kill the tissues, and acting less strongly they may be supposed to damage them; it is in this latter case that they produce inflammation. A certain degree of heat or cold will kill, a less degree will lead, if sufficiently prolonged, to inflammation. Such agents as croton oil, nitrate of silver, chloride of zinc, have a damaging action on the tissues, and they are some of the commonest in use when inflammation is to be produced. Again, traumatic causes produce inflammation; a direct injury inflicted may kill, but acting less vigorously it may produce inflammation. The agents, however, although their direct action is deleterious, may yet induce a reaction in the tissues and in this sense stimulate them.

It may be said, perhaps, that in inflammation of internal organs there is no such definite cause to be discovered. In pneumonia it might be supposed that there is no direct irritant or deleterious agent attacking the lung tissue. But even here there is reason to believe that there is some such agent at work. In one of the hospitals in Glasgow a medical man prescribed a table-spoonful of cough-mixture; the nurse gave a table-spoonful of carbolic acid instead. The result was an acute pneumonia presenting the usual physical signs and symptoms. In this case there was a deleterious agent introduced into the stomach and so into the circulation, and it acted on the tissue of the lungs (perhaps mainly on the blood-vessels), possibly during the process of excretion by these organs; the result was an acute inflammation.

Again, when large blisters are applied, it sometimes happens that the active chemical principle is absorbed, and, passing into the blood, produces inflammation of the kidneys probably during the process of excretion. And so in all inflammations we are to infer the existence of some deleterious agent, although it is very often difficult to tell whence it has come and what is its nature.

Many inflammations are produced by morbid poisons, which, again, are related, as we have seen, to specific microbes. These may be applied directly to the tissues or may be carried by the blood. We have already seen that in this latter case they generally select some special locality for their action. This applies also to ordinary poisons as the instances given above indicate.

At this point it may properly be inquired, whether **the Nervous system** has anything to do with the causation of inflammation. There are not a few facts which suggest, at least, some connection. In some of the instances the connection is clearly an indirect one. For instance, when inflammation of the eye results from division of the fifth nerve,



the inflammation is really traumatic in its origin. By the severance of the nerve the reflex action of winking is abolished as well as the secretion of tears, and so the organ is exposed to injuries and unable to get rid of foreign bodies. The inflammation is the result of irritation from without, and, if the eye be carefully protected (say by stitching the lids together) the inflammation does not occur.

But there are cases where nervous action has a very important and apparently direct relation to the cause of the inflammation. A man with a stone impacted in his urethra may have an inflammation of the testicle. Or, after the passage of a bougie through a urethra with a partially healed wound, there may be a rigor with immediate suppression of urine and an acute inflammation of the kidney. These results can only be effected by reflex action, and there are various ways in which we may suppose them to occur. Reflex irritation may produce, through the vaso-motor nerves, a contraction of the arteries of an organ, and in the case of the kidney this contraction may conceivably be sufficient virtually to bring the circulation to a standstill. The immediate suppression of urine resulting on the passage of a bougie can scarcely be accounted for except on the supposition of such a spasmodic contraction amounting almost to occlusion of the renal arteries. If such a stagnation of the blood continues long enough the vessels will be so damaged that when the circulation is restored inflammation must result.

But there are cases of internal inflammations which are greatly under nervous influences, but in which the connection cannot be explained merely on the supposition of a reflex stimulation of the vaso-motor nerves. The manifest relief which many inflammations undergo in consequence of counter-irritation has been pointed out by Lister as indicating the influence of the nervous system in producing inflammation. The application of the actual cautery to the skin over an inflamed and painful joint will often relieve the pain and inflammation instantaneously. It is as if there was in the nerves of the part a state of over-action which affected injuriously the tissues. This over-action being relieved by a still greater stimulation of nerves connected reflexly, the inflammation subsides.

It is probable that even in such cases nervous action alone is not sufficient to produce inflammation. It is more likely that under certain conditions of the local nervous arrangements the tissues are specially weak, and are liable to be damaged by agents which, under normal conditions, they would resist.

As an example of how local conditions may determine the occurrence of inflammation, the fact may be cited that a mild inflammation of the bone-marrow produced by caustics may be converted into an intense inflammation by causing the



animal to eat putrid food. Here an existing damaged state makes the tissues unable to resist the attack of a further damaging agent which normally they are able to withstand. It may be similar in the case of many internal inflammations produced apparently by exposure of the surface of the body to cold. Such exposure may produce, by reflex action, a state of the nervous system rendering the organ peculiarly liable to damage from conditions of the blood which otherwise would not produce any such effect. In all these cases, also, it is to be borne in mind that the reflex action may affect specially the vascular system.

It may be added that besides conditions brought about through the nervous system there may be **Individual peculiarities, hereditary or acquired**, rendering different persons variously liable to the action of irritants, and even in the same person the various organs of the body may show different degrees of resistance. The character of the inflammation will also, to some extent, be influenced not merely by the nature of the agent causing it, but also by the state of the individual.

In studying individual cases of inflammation it will be important to consider by **what path the irritant has reached the part** which has become inflamed. In many cases it reaches it by the blood, and as the blood is distributed in every part of the organ, the inflammation will not probably show any special localization. And so, when we find two symmetrical organs both of which are attacked in every region, we generally infer that the agent causing the inflammation has come by the blood or, at least, by a path common to both. Of course, there may be local differences in the organ itself of such a character that every part will not be equally susceptible to the action of an agent calculated to produce inflammation, and so the disease may develop more in one part of the organ than in another, although the general character of its distribution will generally still suggest the path by which the agent has come.

#### I.—THE PRINCIPAL PHENOMENA OF INFLAMMATION.

Looking to the cardinal signs of inflammation already enumerated, it will appear that two of them, namely, redness and heat, are intimately related to the condition of the circulation, while the swelling may possibly have similar relations. John Hunter, in his conceptions of inflammation, regarded the phenomena as essentially connected with the state of the blood and blood-vessels. On the other hand, Virchow in his great work on cellular pathology emphasized the importance of the tissues, and regarded the phenomena of inflammation as due to a stimulation of the cells. Under the influence of the discovery, by Cohnheim, of the emigration of leucocytes from the vessels there was a revulsion towards the vessels, and the attempt was made by this author



to limit the term inflammation essentially to the phenomena connected with the vessels, and to regard it as a disturbance of the circulation. There is no doubt that at the outset of acute inflammations the alterations in the circulation are the main factors, but even in the early stages the reaction of the cells must not be ignored, while in the later stages and in chronic inflammations the tissue changes are the more important.

A simple **Experiment** may be performed to illustrate the principal phenomena at the outset of an acute inflammation. If a frog be paralysed with curare, the web of the foot may be spread out and observed under the microscope. With a pair of scissors a superficial longitudinal wound may be made, taking care to remove little more than the epithelium. By this operation the connective tissue of the web, with its vessels, is exposed; the action of the scissors in cutting, and the unusual exposure to the air, affect these structures, and the various phenomena of inflammation soon begin to manifest themselves, care being taken to keep a moist atmosphere around the web so as to prevent the wound, deprived of its epidermis, from drying. At first the circulation goes on in the bottom of the wound as before, the area is merely more transparent from the absence of the epithelium. But very soon, if an artery is near or in the wound, it dilates, and there is an acceleration of the stream in the capillaries and veins. But this soon disappears, and the circulation, especially in the capillaries, becomes slower and slower, till here and there the blood-corpuscles now and then stand still for a moment or two. By and by a peculiar condition becomes visible in the veins. Normally, the blood-corpuscles flow down the middle of the vein, and the peripheral zone contains plasma with a few white corpuscles rolling along. As the inflammation proceeds, the white corpuscles come to occupy this zone, and to adhere to the inner surface of the vessel. The individual corpuscles may not be all absolutely stagnant, they adhere for a time and then depart, but the result of the process is that there is a nearly complete filling up of the zone with white blood-corpuscles, so that the vein seems paved internally with these cells. This is seen not only in any vein which may happen to be in the bottom of the wound, but also in those for a short distance outside it.

If the attention be now directed to the surface of the wound, it soon becomes manifest that certain peculiar bodies are appearing there. These are at first seen mostly towards the edge of the wound, and are especially numerous in the neighbourhood of veins where the white corpuscles are adherent. They are of various shapes, and present a



transparent gelatinous appearance. If observed carefully they are seen



Fig. 20.—A leucocyte from human blood showing amoeboid movement. (KLEIN.)

to be altering their shapes, presenting the well-known amoeboid movement, as shown in Fig. 20. These cells gradually increase in numbers, and by their contractile power, they move from the periphery towards the centre of the wound till they may come to cover it entirely.

These bodies may be removed from the wound by placing the end of a capillary glass tube on the surface. A fluid runs up into the tube, and in this fluid are these free cells. The fluid may now be blown on to a glass slide and examined under a higher power of the microscope, when the slow amoeboid movement will be still more manifest. If, in the drop of fluid, these bodies are allowed to die or are killed by the addition of a reagent, they become globular and granular, in fact, have the characters we recognize as those of white blood-corpuscles, lymph corpuscles, pus corpuscles—of leucocytes in general. The addition of acetic acid to the living cells will first cause them to assume the globular form, and then will bring out the nucleus or nuclei as in an ordinary white corpuscle. The fluid in which these corpuscles are found is coagulable, and if it be kept till the corpuscles have died strings of fibrine will be found in it. If this experiment be made in summer, the whole of these phenomena will manifest themselves in a few hours, and in five or six hours the entire wound may be plastered over with amoeboid cells.

In the course of a few more hours the wound begins to be covered in with new-formed flat epithelium. This begins at the margins, and if the wound is small it may be wholly covered within twenty-four hours. The amoeboid cells are covered in, but they very soon disappear, and the connective tissue of the web remains with a thin transparent epithelium covering it.

Without following out this experiment further, we may now proceed to consider the principal phenomena of inflammation, which have been partly illustrated. We shall consider, first, the state of the vessels; secondly, the condition which is illustrated in this experiment by the fluid on the surface of the wound containing amoeboid cells; and lastly, the condition of the tissues in inflamed parts.

#### 1. THE STATE OF THE VESSELS IN INFLAMMATION.

When an irritant is applied to a transparent vascular tissue, such as the tongue or web of the frog, it produces effects which vary slightly,



according to its nature. If croton oil be applied, there is first a contraction of the arteries, extending to their whole length, followed by dilatation. If ammonia be used, there is dilatation without previous contraction. The dilatation affects chiefly arteries and veins, but also, though to a less extent, capillaries. The **dilatation of the arteries** leads to an **Active hyperæmia**, the current is accelerated in the arteries, capillaries, and veins, and these vessels are overfilled; there is a great excess in the quantity of blood passing through the vessels. The acceleration of the current does not persist, however, in the most affected parts; on the contrary the blood corpuscles begin to lag, especially in the capillaries and veins, although there is still acceleration in the arteries, and in the capillaries and veins of the less inflamed parts. This stagnation in the capillaries and veins may assume a high degree, especially in the part most acted on by the irritant. Although the current is slow in these vessels they remain overfilled, a **Passive hyperæmia** supervenes on the active hyperæmia; at the same time the white corpuscles accumulate along the internal wall of the veins in the manner already described, and they also adhere at intervals in the capillaries. The circulation may come almost to a standstill in the capillaries of the parts most affected, while at various distances out from this there will be manifest a less and less amount of retardation till a zone is reached where the retardation disappears, and by and by gives place to acceleration.

We have now to consider what may be the explanation of these various phenomena which the vessels manifest. The observations of Lister present us with a view of this subject which has been largely confirmed by other observers. Saviotti's researches are for the most part confirmatory of Lister's views.

**Contraction of the arteries** is not, in any proper sense, a part of the inflammatory phenomena. It is simply the result of stimulation of the nerves by the irritant, and while in the case of many irritants it does not occur at all, it is always transitory. Contraction of arteries may be produced either by reflex or direct irritation of nerves. It is produced reflexly by irritation of sensory nerves, as when the arteries in the web of the frog's foot are seen to contract when the skin is tapped or twitched with the forceps. This contraction does not occur if the nerves be first divided. On the other hand the irritant may cause contraction directly, by stimulating the vaso-constrictor nerves (as cold does), and so produce a temporary contraction of the arteries.

**Dilatation of the arteries**, leading to **active hyperæmia** (also called



*determination of blood*), is induced by **paralysis of the arteries**, just as in other cases of active hyperæmia.

It is important here to recall the fact that **irritants act injuriously** on the tissues, and, in a certain sense, paralyse them. Lister in his important researches on inflammation brought this fact into prominence. The skin of the frog is supplied with pigment cells. These cells are contractile bodies (they are shown in Fig. 12, p. 87). In the state of rest they are extended into numerous branches, which make a fine pigmented reticulum under the skin; in the active state they are drawn together so as to make a dark clump. They are under the command of the nervous system, and by their means the animal is capable of changing its colour, presenting a dark hue when the cells are relaxed, and a lighter colour according to the degree of concentration. Some irritants have the immediate effect of relaxing the pigment cells, and this itself is so far an evidence of paralysis, as the dispersed condition is the state of rest of the cells; but whether the pigment is dispersed or not, the animal loses control of its pigment in the affected area, which does not change its colour with the rest of the skin, and may be found dark while the animal is pale, etc.

In a similar manner the dilatation of the arteries is effected by a paralytic influence of the irritant. It is not easy to determine whether this paralytic influence is exercised through the peripheral ganglia, whose existence we have already seen reason to infer, or directly on the wall of the vessel, but as these ganglia are in or near the vessel-wall, we may infer that both are influenced. The view has been held that the dilatation is reflex, but this is excluded by the observations of Cohnheim. He found that if the sciatic nerve has been divided in a frog's leg the arteries dilate, but the application of an irritant produces a further dilatation. He found also that, after destruction of the brain and spinal cord, irritation of the tongue still produces dilatation of the arteries. The acceleration of the blood-current in the arteries, capillaries, and veins, will be understood from what has gone before to be a direct result of the dilatation of the arteries; we have, in fact, an active hyperæmia.

**The Retardation** is to be referred to an **increased adhesiveness** of the blood-corpuscles. This is a matter of direct observation. The corpuscles in the inflamed area can be seen to move sluggishly along the wall as if attracted to it, and the **Pavementing of the veins** with white corpuscles is clearly due to increased adhesiveness. It has been pointed out by Lister that, when the blood is removed from the vessels and comes in contact with dead matter, the blood corpuscles acquire an adhesiveness which they do not possess inside



the normal vessels. The red corpuscles stick together by their flat surfaces and form the well-known *rouleaux*. The adhesiveness of the white corpuscles is not so obvious when a drop of blood is examined outside the body, but there is reason to believe that it is even greater than that of the red. Now the irritant damages the walls of the vessels, with the result that the corpuscles behave as if in the presence of dead matter—they become adhesive.

This is excellently shown in an experiment of Lister's. He ligatured the leg of a frog, producing thereby stagnation of the blood in the vessels, but on examining the web it could be seen that the corpuscles were able to move freely among one another—there was obviously no adhesiveness. But now, when a piece of mustard was applied to the web, this free movement ceased in the area affected; the corpuscles became adherent among themselves and to the walls of the vessels. The result of this was an accumulation of the corpuscles in the irritated area; any corpuscles which happened to glide into the area remained adherent there, and so, by degrees, the vessels became overfilled—a state of hyperæmia superinduced on stagnation. If any corpuscle happened to escape from the affected area, it ceased to be adhesive, and moved freely about.

We may therefore infer that the retardation of the current and the paving of the veins with white corpuscles are the result of the injury to the vessel wall, and it may be added that, in connection with inflammations, all degrees of stagnation up to absolute stoppage or **Stasis** may be produced, and are often manifested together in the same case.

We may now **sum up** the conditions presented by the blood-vessels in the early periods of inflammation as follows. The arteries dilate by a relaxation of their muscular coats, due to a paralysis of the peripheral ganglia. In some cases this dilatation is preceded by an evanescent contraction. The immediate result of the dilatation of the arteries is active hyperæmia, or determination of blood, involving overfilling of the arteries, capillaries, and veins, with acceleration of the current. This is followed by retardation of the current, the vessels remaining dilated and hyperæmic, and this retardation may go on to almost complete stasis in the capillaries. The retardation is due to adhesiveness of the corpuscles, and to the same cause is to be traced the paving of the veins with white corpuscles.

It is sometimes possible in acute inflammation of the skin, as in the case of a boil, to observe conditions directly traceable to the state of the vessels here indicated. Thus at the peripheral parts of such a focus of inflammation the skin presents a fiery-red appearance due to determination of blood; the red colour may be pressed away with the finger, but it immediately returns. Inside this zone there is an area in which the redness is not so vivid, and when the red colour is pressed away



it returns sluggishly; the corpuscles are here already adherent, and the current retarded. Then in the more central parts a dusky red appearance is presented, and on pressure it may be impossible or very difficult to remove the redness; here a condition of extreme stagnation exists.

**Increase of temperature in inflammation.**—**Calor** is one of the cardinal signs of inflammation, and a feeling of heat is usually experienced when external parts are the seat of acute inflammation. In regard to internal parts, their nerves are not capable of conveying impressions of differences of temperature; a hot substance swallowed gives a sensation of pain. Many experiments have been made with a view to determining whether a part which is in a state of acute inflammation is the seat of increased production of heat.

There is no doubt that the temperature of inflamed external parts is increased in inflammation. John Hunter determined this by actual observation. He had a case of hydrocele to deal with, and undertook its treatment by the old operation of laying open the sac and inserting lint dipped in an irritating salve with a view to producing inflammation. At the time of the operation he found that the temperature in the tunica vaginalis was  $92^{\circ}$ , but next day, inflammation having been induced, the temperature had risen to  $98\frac{3}{4}^{\circ}$ . The question here arises whether this increase of temperature is due to an actual production of heat in the inflamed part, or to an increased supply of hot blood, due to dilatation of the arteries.

That there is a **greatly increased supply of blood** to inflamed external parts has been proved by experiment and observation.

Cohnheim found that in a recent inflammation produced by scalding the fore leg of a dog in hot water, or by painting with croton oil, the amount of blood issuing from a cannula inserted into a vein was greatly increased, sometimes reaching nearly double that issuing from a corresponding vein on the other side. It may even be as great as when the arteries have been relaxed in a limb by dividing the axillary plexus of nerves. This extreme degree of difference may not continue long, but even for some days it is frequently very considerable.

It is an every-day observation of surgeons that an incision in an inflamed part is accompanied by a much greater escape of blood than in a normal part, and if an artery be incised it spouts more blood and to a greater distance than a similar artery in a non-inflamed part. It is similar with the veins; Laurence performed venesection in both arms in a person who had an acute inflammation of one hand, and he found that the blood flowed two or three times more rapidly from the vein on the side affected. The relaxation of the arteries in acute inflammation evidently produces a determination of blood which extends



beyond the immediate focus of the inflammation, and is not counter-balanced by the retardation.

John Hunter, having determined the great increase in temperature above referred to, yet came to the conclusion that it was due to the determination of blood, for he found that the temperature of an inflamed external part never exceeded or even quite reached that of a normal internal part, or, in other words, of the blood in internal organs. Since Hunter's time, Simon has asserted that there is some development of heat in inflamed parts. His experiments seemed to show that the arterial blood passing to an inflamed external part is not so warm as the focus of inflammation, and that the venous blood returning from the part is warmer than the arterial blood, although not so warm as the focus of inflammation. But the experiments of Jacobsen, made with more exact instruments, entirely confirm the views of Hunter. It appears that the most intense inflammations of the skin or of muscles never cause an elevation of temperature sufficient to reach that of the rectum, vagina, or abdomen, the difference being generally  $1^{\circ}$  to  $2^{\circ}$  C. Again, in inflammations induced in internal parts, as the peritoneum or the pleura, the temperature was never raised to that of the blood in the left ventricle, being always from  $.2^{\circ}$  to  $.5^{\circ}$  C. under it. This means that in these parts, where the temperature is already near that of the blood in the left ventricle of the heart, inflammation, leading to an increased supply of blood of a similar temperature, causes virtually no elevation above the normal heat. Again, Cohnheim has found that, when in one fore-paw of a dog a state of acute inflammation is induced, while in the other active hyperæmia is produced by dividing the axillary plexus of nerves, the temperature in the inflamed foot is always slightly less than that in the other. We may safely infer, then, that in inflammation there is no local production of heat.

**Fever** often accompanies inflammation, but it only does so when the blood is affected. This may happen in one of two ways. The agent which caused the inflammation may be primarily in the blood, and it may produce both the general manifestations of fever and the local manifestations of inflammation. This applies to many morbid poisons. On the other hand, a local inflammation may give rise to products whose absorption into the blood will cause fever. (See under Fever.)

## II.—THE INFLAMMATORY EXUDATION.

In the experiment sketched at the outset the inflammatory exudation was the fluid which collected on the surface of the wound. We saw that this fluid contained amœboid cells, and that it was coagulable. We



may thus consider the inflammatory exudation as consisting of cells, serous fluid, and fibrine, and we shall in the first place pass each of these under review.

1. **The serous exudation.**—Nearly all inflammations are accompanied by a transudation of the fluid part of the blood from the blood-vessels, and this passes, in the direction of least resistance, wherever it finds room. In the case of a wound it flows from the surface, forming a serous **discharge**; in that of a mucous membrane it also flows from the surface, forming a **catarrh**; in serous cavities it accumulates, forming an inflammatory **dropsy**; in the tissues it passes into the serous spaces constituting an inflammatory **œdema**; in the lungs it is situated in the lung alveoli, where also it produces œdema. The exuded fluid will in many cases find its way into the lymphatics, and experiment has proved that the current in the lymphatics is much increased in inflammation.

The serous exudation is somewhat different in constitution from an ordinary transudation fluid. It is much more concentrated, approaching more nearly to the liquor sanguinis, and containing more leucocytes than the exudation in simple œdema. It is also coagulable, so that when shed it may deposit fibrine, and sometimes does so in the living body. When this occurs we have the fibrinous exudation. We may associate these characters of the serous exudation with the fact that in acute inflammations the tissues including the walls of the vessels are seriously damaged, and the latter allow more readily of the escape of the fluid of the blood.

2. **The fibrinous exudation.**—This is seen most typically in acute inflammations of serous cavities such as the pleura or pericardium. In such cases there is, in addition to the serous exudation occupying the cavity, a deposition on the surface of a soft yellow layer of coagulated fibrine. There is frequently a similar deposition on the surface of a freshly-inflicted wound, the fibrine forming a glaze on the surface, while the serous fluid passes off as a discharge. Fibrine is seldom deposited in the meshes of the tissues, unless there be an actual necrosis, as in the case of a boil or a carbuncle, where the slough which forms in the skin is composed partly of dead tissue and partly of fibrine. In acute pneumonia, however, the exudation in the lung alveoli is fibrinous.

The term **Lymph**, or coagulable lymph, is often applied to the fibrinous exudation as seen on serous surfaces, but the use of this term is not to be commended. From its use by John Hunter the term has interesting historical relations, yet, as it implies a theory which is not now held, namely, that the so-called lymph has the power of developing into organized tissue, its use is apt to lead to confusion. This is all the more true, because the term lymph is frequently used in a very loose way to



designate connective tissue formed as a result of inflammation. According to Hunter's view, the connective tissue developed directly from the fibrine, and so was legitimately called lymph; but as his views are now departed from, this use of the term is quite unwarranted.

**The fibrine** of the exudation has all the characters of that in an ordinary blood-clot. This is shown in Fig. 21, where a network of fibres

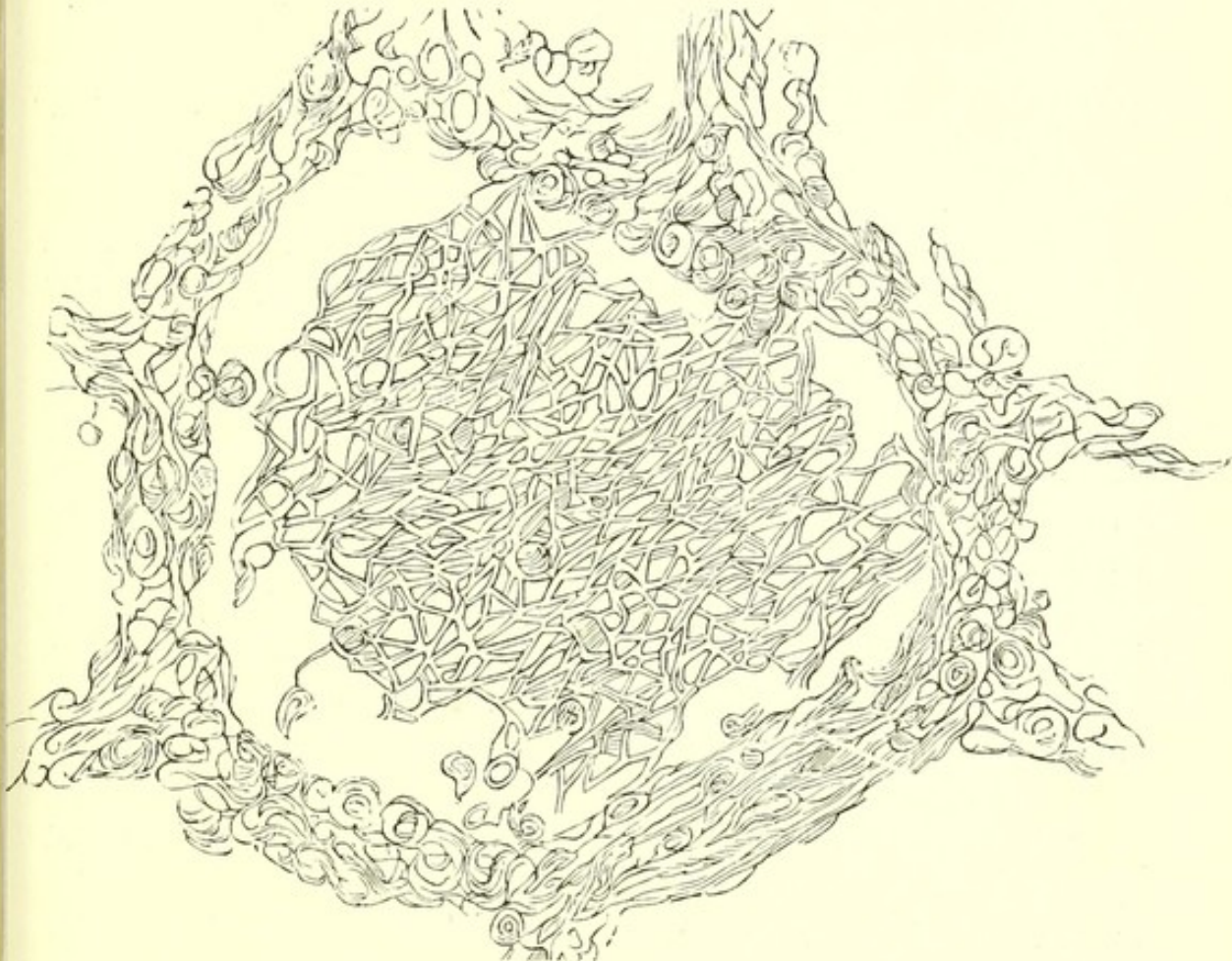


Fig. 21.—Lung alveolus in acute pneumonia. A plug of fibrine, with a few leucocytes in it, fills the alveolus.  $\times 350$ .

is seen filling a lung alveolus. It has also a similar origin to that in the blood-clot. We have seen (under Thrombosis, p. 64) that coagulation of blood occurs when the necessary constituents are present, and that in order to this there must be a disintegration of the leucocytes so as to yield the paraglobulin and the ferment. Leucocytes are always present in the serous exudation, and in order to their disintegration they must be sufficiently removed from the influence of the living tissue. This can scarcely occur except in the case of an extended surface as of a membrane. Just as thrombosis does not occur in the capillaries, because the blood in these narrow vessels is in intimate contact with the living cells forming the capillary walls, so in inflamma-



tion the fibrinous exudation scarcely occurs in the serous spaces unless there be actual necrosis of the tissue. In some very acute inflammations of connective tissue (acute phlegmon) there may be a deposition of fibrine, forming a kind of **fibrinous œdema**, such as sometimes occurs in the skin in erysipelas, but in these cases necrosis is a frequent if not a constant concomitant. It is where the exudation is on a surface, and is to a great extent removed from contact with the living endothelium that the fibrinous exudation is most constantly found.

Epithelium, like endothelium, has the power of preventing the disintegration of the leucocytes, and hence a fibrinous exudation seldom occurs on a mucous membrane or the skin. Its occurrence implies that the epithelium has undergone necrosis or has been shed.

3. **Exudation of white and red blood-corpuscles.**—The resemblance of the cells met with in acute inflammations to the leucocytes in the blood long ago suggested the idea that they are white blood-corpuscles. In the year 1846 Waller observed the pavementing of the internal coat of the veins with white blood-corpuscles in the inflamed tongue of the frog, and as he saw similar cells outside the vessel he inferred that the white corpuscles had got through the wall. It was, however, difficult to believe that the solid globular white corpuscles could pass through the intact wall of a blood-vessel, and Waller's views, although supported by William Addison, were lost sight of. The discovery by Recklinghausen that pus corpuscles and white blood-corpuscles possess contractile power by virtue of which they are able to move from place to place, and to alter their shapes in the most diverse fashion, paved the way for the actual observation of their passage through the walls of the vessels made by Cohnheim.

**Emigration of the white corpuscles.**—This was observed by Cohnheim in the mesentery of the frog. When this exceedingly delicate and transparent structure is drawn out of the body through a wound in the lateral aspect of the abdomen, the mere exposure to the air is sufficient to set up an acute inflammation, the phenomena of which can be readily observed under the microscope. Let us suppose that the pavementing of the veins has occurred, and that there is an occasional white corpuscle adherent in the capillaries, and the following surprising phenomena show themselves, as described by Cohnheim himself. "One sees, as a rule, first in a vein which presents the pavementing with white corpuscles, but sometimes in a capillary, a pointed projection in the external contour of the vessel; it pushes itself farther and farther outwards, it increases in thickness, and the pointed projection develops into a colourless rounded knob; this increases in length and thickness, sends out fresh points, and draws itself gradually outwards from the



vessel-wall, with which it comes to be connected only by a long thin stem. Finally, this also lets go the vessel, and there lies outside a colourless, dull, glancing, contractile corpuscle, with several short processes and a long one, of the size of a white blood cell, with one or several nuclei; in a word, a white blood corpuscle."

In the investigation of inflamed tissues in the usual way after death, there are often indications to be met with of this process of emigration. The leucocytes in the tissues, for instance, are frequently aggregated especially around the blood-vessels. In the accompanying Fig. 22, the

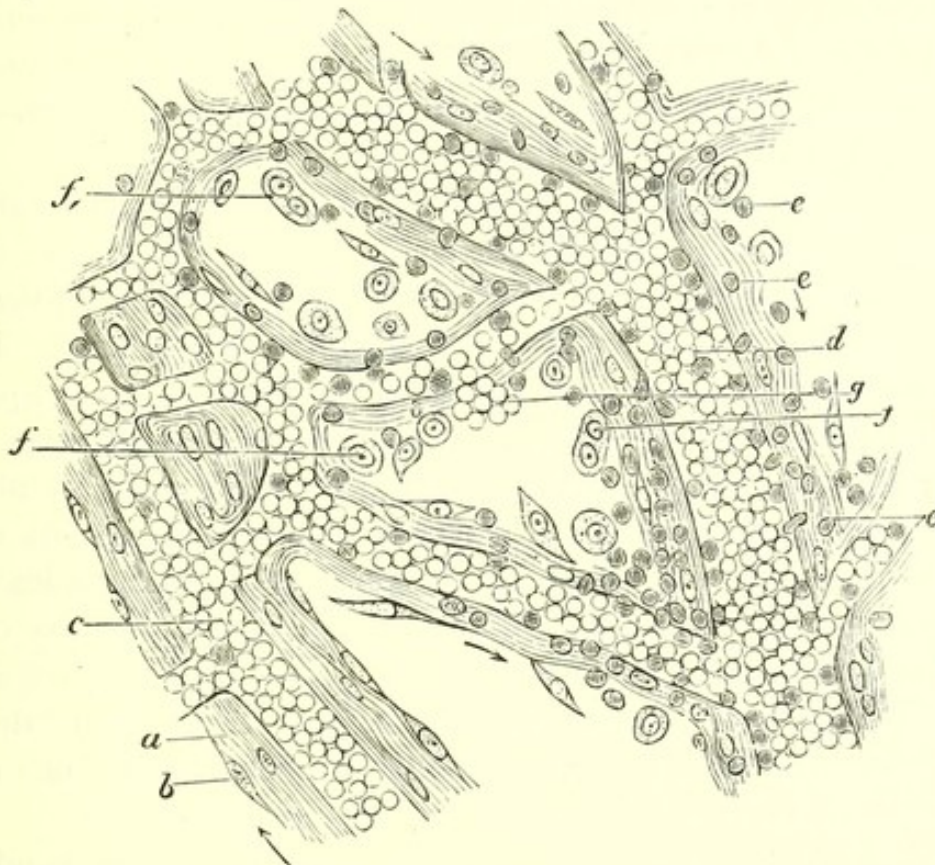


Fig. 22.—Inflamed human omentum. The phenomena of inflammation are seen in the veins and capillaries. The condition is normal at the artery (*c*) where *b* represents endothelium covering the trabecula (*a*). In the vein (*d*) there are many white corpuscles along the wall; some of these are emigrating (*e*). *f*, desquamated endothelium; *g*, extravasated red corpuscles. (ZIEGLER.)

conditions in the human omentum are shown in a somewhat diagrammatic form, and the appearances there are sufficiently suggestive.

**Diapedesis of the red corpuscles.**—This is a frequent accompaniment of acute inflammations. The white corpuscles are active contractile cells, and they generally pass through the vessels to a larger extent than the red ones, but there are some inflammations in which the red corpuscles also pass through in large numbers. They are not active, but are passively pushed through the walls, just as they are in the diapedesis of passive hyperæmia.

This passage of the white and red corpuscles through the walls of the vessels is to be accounted for by changes in the vessels induced by



the agent producing the inflammation. These changes, as we have seen, induce an increased adhesiveness of both white and red corpuscles, and lead to the pavementing of the veins. The altered vessel-wall may also permit of freer exercise by the leucocytes of their power of amœboid movement. The living wall of the vessel seems to restrain this movement, but in inflammation the altered wall allows itself to be penetrated by the contractile leucocytes. Increase of pressure has been suggested as a cause of the extravasation of the corpuscles, but its influence is in any case subordinate. Even in the case of the red corpuscles the diapedesis is related to the alteration in the vessel-wall, which permits of the corpuscles being pushed through. It occurs to the largest extent in inflammations of a particularly virulent kind, in which the vessel-wall is presumably most seriously compromised.

The passage of the corpuscles takes place, presumably, through existing apertures in the walls of the vessels. The capillaries are not composed of membrane, but of flattened cells, which have places where they join, as well as apertures between them (*stomata*, see under Passive Hyperæmia). Both red and white corpuscles pass through apertures much narrower than their own proper breadth. They undergo alterations in shape so that the part actually in the wall is very attenuated.

It is customary to call all the leucocytes which are met with outside the vessels in inflamed tissues by the name of **Pus corpuscles**. This term is hardly warranted unless there is the actual presence of fluid pus. But as the characteristic corpuscles of pus are leucocytes, and are identical with those met with in other acute inflammations, the term may be retained in this wider significance as meaning cells of a similar character to those in pus.

Besides leucocytes we often meet with other cells in exudations, which are obviously not derived from the blood, and even in regard to the leucocytes themselves it cannot be taken for granted that they have no other source than the blood. Hence it is necessary to discuss the extravascular origin of certain cells in the exudation.

**4. Cells from other sources than the blood.**—Although Cohnheim and his followers have asserted that pus corpuscles have no other origin than the leucocytes of the blood, there are many facts which render this conclusion very doubtful. In the first place, pus corpuscles outside the vessels may multiply by division, a process which has been actually observed by Stricker. Then, also, observations on non-vascular structures seem to show that the connective-tissue corpuscles may give origin to leucocytes. These observations have been made chiefly on the cornea and on cartilage.

Hoffmann and Recklinghausen produced inflammation in the cornea and then



excised the eye or the head of the animal and preserved it in a moist chamber. They found in two or three days that at the seat of irritation groups of amœboid cells (pus corpuscles) had formed, as in ordinary keratitis. Then, also, several observers have described and figured the cornea corpuscles in inflammation, as drawing in their processes, and in doing so becoming amœboid. They also leave detached portions of their protoplasm which become pus corpuscles. It has also been asserted that the epithelium on the posterior surface of Descemet's membrane, as well as that on the surface of the cornea and elsewhere becomes, at the outset of inflammations, amœboid.

The so-called endogenous production of pus corpuscles is more doubtful. By this is meant the formation of these cells inside other cells. It has been asserted by Buhl, Rindfleisch, and others, that epithelial cells give origin in this way to pus corpuscles. No doubt the latter forms are sometimes found inside the former, but as both forms may be contractile, this does not imply that the one has given rise to the other, more especially as red corpuscles are found inside cells under similar circumstances.

It may be said in the meantime that while pus corpuscles undoubtedly arise by emigration of the blood corpuscles, there is considerable probability that as the inflammation proceeds the existing tissues, and more especially the connective tissue, give origin to these cells. It is clear that in the course of a suppurative inflammation, in which there is an enormous production of pus corpuscles, there must be a very great new formation of leucocytes somewhere. In such cases the lymphatic glands and spleen sometimes undergo enlargement, and Cohnheim asserts that the number of leucocytes in the blood may undergo a great increase.

The cells met with in inflammatory exudations are not all of the character of pus corpuscles. They are sometimes larger and partake of the character of **derivatives of the epithelial cells**. This is more particularly the case in certain inflammations of the lung, where the alveoli are frequently occupied by cells which are evidently produced from the epithelium of the alveoli (so-called *catarrhal cells*). A similar formation of cells is frequently seen in the uriniferous tubules in inflammations of the kidneys. (See further on under Parenchymatous Changes.) Again, in suppuration in epithelial structures generally, amœboid cells are often found which are much larger than pus corpuscles and are derived from the epithelium. Neumann asserts that ciliated epithelium sometimes becomes amœboid and can be recognized, even after it has become detached, by the persistence of some of its cilia.

**5. The purulent exudation. Suppuration. Pus.**—Pus is a fluid usually of a yellow colour. It consists of a fluid portion, the liquor puris, and of pus corpuscles, which are identical with ordinary leucocytes. Pus is met with under a variety of circumstances, but in nearly all cases it is produced by the indirect action of microbes belonging to the class of micrococci. The association of suppuration with the presence of certain forms of microbes has given rise to some interesting



discussions as to the explanation of this association. It is asserted by some that the leucocytes are designed to destroy the microbes by taking them into their substance and digesting them. (See under Phagocytes in section on Bacteriology.)

The purulent exudation frequently follows the fibrinous exudation, especially in serous cavities, and it is not uncommon to find an intermediate condition in which the fibrine is, as it were, infiltrated with pus. This implies that the cause of the inflammation (micrococci) continues to act intensely. The emigration of leucocytes goes on vigorously, and these, crowding into the fibrinous coagulum, the latter disintegrates and liquefies.

Many experiments have been made in order to determine whether suppuration can be produced without the agency of microbes. From some of the most recent of these, such as those of Grawitz and De Bary, it appears that various substances, such as turpentine, may alone induce suppuration, and that cadaverin, obtained from decomposing matters, but free from microbes, may do so. We may infer that while suppuration may be produced by different active agents, yet in actual cases micrococci are always present. We may also infer that it is by evolving irritating chemical principles (animal alkaloids—ptomaines) that they produce this effect.

The solution of the fibrine is not easy to explain, nor is it easy to understand why pus does not coagulate. There are abundant leucocytes, and there is the fluid presumably containing the fibrinogen. A few pus corpuscles added to fresh liquor sanguinis induce coagulation. Its absence in pus has been ascribed to the existence of some chemical agent evolved by the micro-organisms which inhibits the process of coagulation, or to a conversion of the fibrinogen into peptones, which latter are known to be present in pus.

In the case of serous membranes the occurrence of suppuration implies a very intense inflammation, but in the case of mucous membranes pus is often produced when the inflammation is comparatively trivial. The ordinary catarrh of the nares and bronchi is first accompanied by a serous exudation, and this often gives place to an exudation of pus.

**Purulent infiltration** is a term applied when the suppuration occurs not at a surface but in the midst of the tissue, and the pus fills up the spaces, the condition being comparable, in respect of the locality of the exudation, with œdema. Purulent infiltration implies an intense inflammation, and it is frequently associated with necrosis.

**Abscess** frequently follows purulent infiltration. The name abscess or **Apostema** is applied to a collection of pus in a cavity generally formed to accommodate it, although it is sometimes used in cases where the pus has accumulated in a pre-existing cavity. Purulent infiltration usually precedes, and the abscess is formed by the necrosis or liquefaction of the tissue, so that a cavity results. Hence abscesses often contain shreds of tissue or sloughs.



The abscess has often a distinct membrane forming its boundaries. This membrane is composed of granulation tissue, and it has often been regarded in the light of a secreting surface by which the pus is produced; hence the name **Pyogenic membrane** applied to it. The membrane, however, is by no means necessary to the formation of pus, and it is, in fact, a secondary product after the abscess has actually formed. It may, indeed, supply pus from its vessels, and so cause enlargement of the abscess, but it more probably rather limits the enlargement, as granulation tissue is less sensitive to irritants than other tissues. The membrane may be regarded as **a protective layer**.

As abscesses usually contain the agents of inflammation which have produced the suppuration, they usually enlarge. This mostly occurs in the direction of least resistance, but gravitation often plays a part in the advance of an abscess. There is a gradual liquefaction of the tissue before the advancing pus, and this goes on, as a rule, till a surface is reached and the pus discharged.

A **cold abscess** is not really a collection of pus, but only of matter resembling pus. This is usually softened caseous matter, produced, in connection with tuberculosis of bone, but pus corpuscles are often mixed with this debris. Cold abscesses frequently extend long distances, assisted by gravitation, before they reach the surface. Their opening, by admitting septic micrococci, may lead to acute suppuration within the cavity.

**Pus**, whether in abscesses or elsewhere, frequently **undergoes changes**. The pus corpuscles often undergo fatty degeneration, or they may swell up and disintegrate. The pus may in this way become absorbed. It sometimes, by absorption of its fluid, thickens into a pasty matter, in which lime salts may be deposited. Again, pus, or indeed a serous exudation, may be mixed with other matters, the secretion of glands, such as mucus, urine, or bile, or with oil where suppuration occurs in a tissue rich in fat.

[**Croupous and diphtheritic exudations**.—These terms have got into common use, but their employment is not to be recommended.

The **croupous exudation** is primarily that which occurs in the larynx in Croup (a Scotch word derived from the peculiar sound made during inspiration in certain diseases of the larynx). The exudation in the larynx is a whitish layer resembling fibrine, and by many regarded as a fibrinous exudation. Wagner believed that it arose by transformation of the epithelium, and Weigert has asserted that it is due to a coagulation-necrosis of the epithelium.

The **diphtheritic exudation** implies a necrosis not only of the epithelium but of the mucous membrane. A fibrinous exudation occupies the meshes of the necrosed tissue as well as its surface, and it is consequently adherent to and involved in the membrane.



These terms, although originally applied to croup and diphtheria, are now frequently used in a purely anatomical sense of exudations on mucous membranes. A croupous exudation is one which lies on the surface and is not associated with necrosis of the mucous membrane, while a diphtheritic exudation is both on the surface and in the substance of the membrane and implies necrosis.]

### III.—CHANGES IN THE TISSUES AND THE INFLAMMATORY NEWFORMATION.

1. **Parenchymatous changes.**—In what has gone before, the changes referred to have mainly concerned the blood-vessels and the connective tissue with its lymph-spaces and surfaces. Besides these, however, we have to consider the proper parenchyma of the tissues, the structures which perform the special functions of parts, and which the connective tissue, as a general rule, only supports. We have the numerous epithelial tissues, some of them covering surfaces, and others forming the secreting structures of glands; and we have, besides, the parenchyma of muscle, of nerve-tissue, of bone, etc.

Cohnheim and his followers would limit the proper phenomena of inflammation to those already considered, and would regard the parenchymatous changes as secondary and unessential. It is true that in many acute inflammations the only obvious changes, apart from the blood-vessels, are degenerative, consisting mainly of necrosis and fatty degeneration, but there are many inflammations in which active changes are visible in the parenchymatous cells, and these may or may not be associated with degeneration of these cells.

The active parenchymatous changes consist mainly of enlargement of the protoplasm, due to an infiltration with albuminous material. The cells acquire, in consequence, a granular clouded appearance, which is expressed in the name **Cloudy swelling**. (See under Albuminous Infiltration.) At the same time the cells frequently multiply, and they do this by the process of karyokinesis. The new-formed cells frequently depart from their position, especially in inflammations of epithelium. This process of **Desquamation** causes the cells to form sometimes a material part of exudations. (See above.) These phenomena of cloudy swelling and desquamation are frequently associated with fatty degeneration. Another occasional sign of increased activity is an increase to the secretion of glands involved in inflammations. Thus the secretion of mucus is frequently increased in inflammations involving mucous membranes.

There are some inflammations in which these phenomena are so characteristic that Virchow's name of **Parenchymatous Inflammation** is properly used for them. It may be presumed that the irritant in these



cases, usually a morbid poison, has special affinities for the parenchymatous structures. It is not to be inferred, however, that the phenomena in the blood-vessels already described are absent in these cases. They are probably present in all cases, and they may be much more prominent during life than one would infer from the appearances after death, when the parenchymatous changes are perhaps unduly obtrusive.

**2. Newformation of tissue.**—In what has gone before the phenomena described have been chiefly those of the earlier periods of inflammations and hence of the acute stage. If an inflammation persists, it usually becomes less intense, and in that case changes ensue which result in newformation of tissue. It may be said, indeed, that most **Chronic inflammations** are characterized by newformation. The new-formed tissue sometimes exercises an important function in serving as a bond of union between surfaces which have been separated, but it is very frequently of no such beneficial character.

**The Granulating Wound**, with its subsequent development into the **cicatrix**, may be taken as a type of the inflammatory newformation. A granulating wound forms a red, somewhat irregular surface, and consists of a highly vascular tissue. Under the microscope, the tissue consists principally of round cells embedded in a fine reticulum, and permeated by abundant blood-vessels which form loops towards the surface. (See Fig. 23.) If the wound be of some duration, there are, besides round



Fig. 23.—Vessels of a granulating wound injected. (BILLROTH.)

cells, spindle-shaped ones, and sometimes large giant cells. Indeed, in the deeper layers, one commonly observes layers of spindle-cells. (See Fig. 24.)

On careful examination the round cells of the granulating wound are seen to be of two kinds, namely, leucocytes (Fig. 25, *a* and *a*), many of which show the usual divided nucleus, and larger cells with single large oval nuclei (*b b*). These latter somewhat resemble epithelial cells and are hence called **Epithelioid cells**. As they are concerned in the further development into connective tissue they are sometimes called **Fibroblasts** or **Formative cells**. Sometimes the formative cells enlarge



greatly and acquire many nuclei, thus taking the characters of giant cells ( $c_1$ ).

According to the observations of Ziegler, which were made by introducing into the living tissues small glass slides cemented together, so as to leave capillary spaces between, into which the new-formed tissue grew, the epithelioid cells are concerned in the further development of the tissue. The foreign bodies introduced set up inflammation. In about five days the formative cells made their appearance. They are, like pus-corpuscles, contractile, but their movements are slow. These cells elongate and become spindle-shaped (see Fig. 25), forming the large spindles seen in granulating wounds.



Fig. 24. — Semidiagrammatic section of a granulating wound.  $\times 85$ .

The formation of the fibrous intercellular substance is rather difficult to follow. According to Ziegler, the protoplasm of the formative cells becomes fibrillated, but in many cases the appearances are suggestive rather of a process of secretion or formation around the cells. The conditions, in fact, sometimes closely resemble

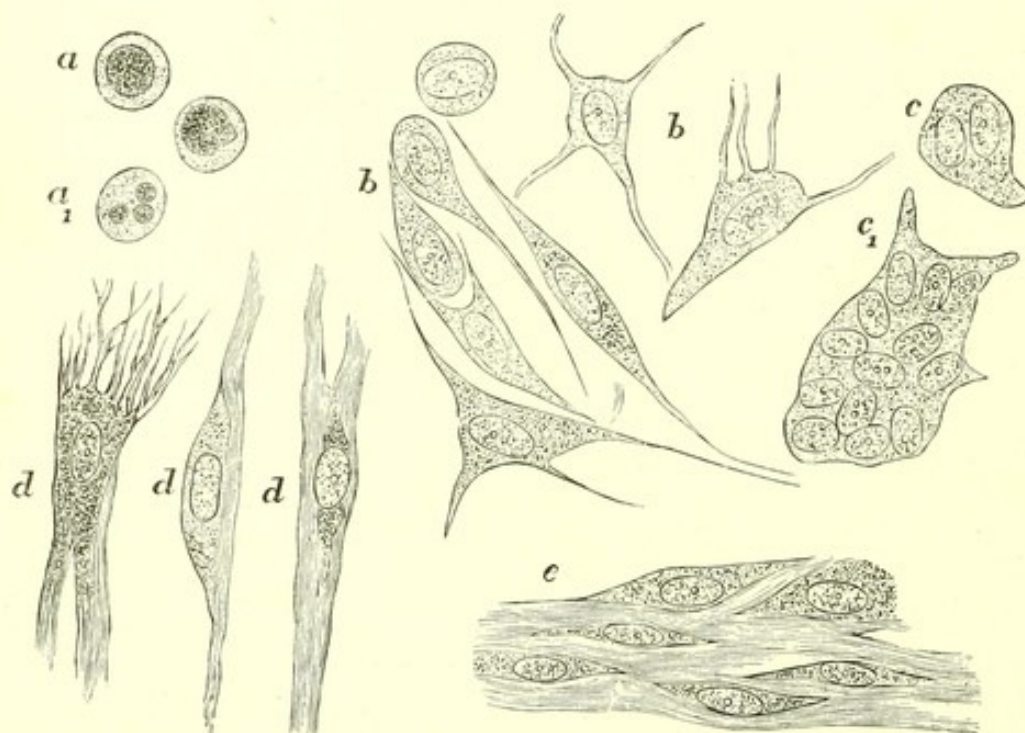


Fig. 25. — Granulation cells.  $a$  and  $a_1$ , leucocytes;  $b$ ,  $b_1$ , various formative cells;  $c$ , formative cell with two nuclei;  $c_1$ , with many nuclei;  $d$ ,  $d_1$ ,  $d_2$ , Formative cells developing connective tissue;  $e$ , complete connective tissue.  $\times 500$ , Picrocarmine preparation. (ZIEGLER)

what is seen in the normal development of bone, in which the osteoblasts surround themselves with zones of osseous matrix. The result



is the formation of a properly organized connective tissue, and this, when the process has involved the entire granulation wound, forms the cicatrix.

**The origin of the fibroblasts** was asserted by Ziegler and others to be from leucocytes. It was stated that these enlarged, or two or more coalesced, and so formed the epithelioid cells. It may be that to some extent they have such an origin, but there is evidence that the fixed cells of the connective tissue are engaged in the process of newformation, and it is probable that they alone are the essential agents. These cells have been observed in inflammatory newformations to be engaged in the process of **karyomitosis** or **karyokinesis**, which is observed in cells which are undergoing multiplication, and is evidence of active newformation. From what has already been mentioned at page 131 it will appear that fixed cells of various kinds may give origin to contractile cells, which are indistinguishable from leucocytes. It may also be regarded as supporting the view that the existing connective tissue is the essential source of that which is new-formed, that undoubtedly the blood-vessels are always produced from existing vessels.

**The Newformation of blood-vessels** takes place by a process of sprouting or **budding** from the existing vessels. The protoplasm of one of the cells in the wall of a small artery or capillary protrudes outward, like a bud, and extends to a neighbouring vessel, or to a corresponding process from another vessel. In this way solid arches (see Fig. 26) are formed which increase in thickness. The vessels are formed

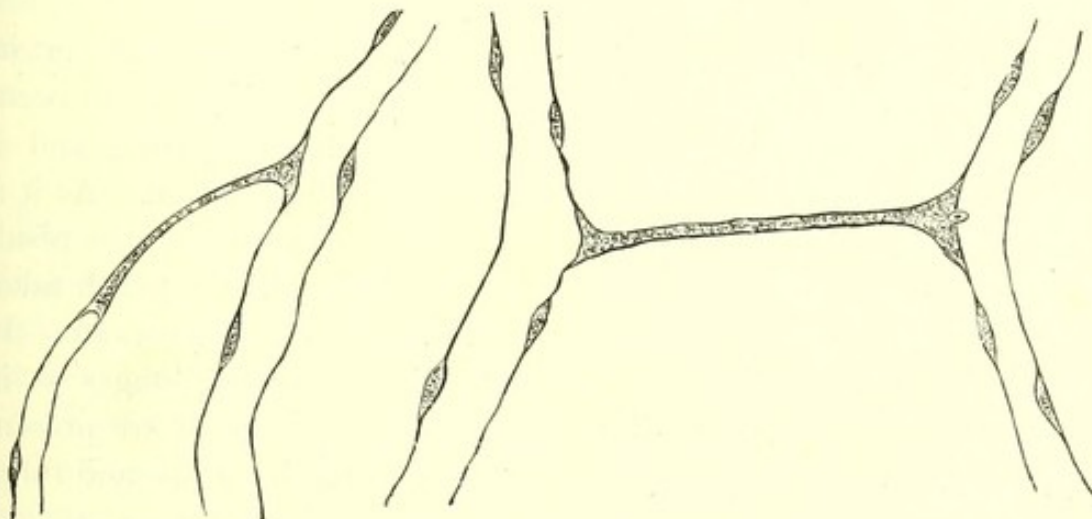


Fig. 26.—Newformation of blood-vessels in a granulating wound. (After ARNOLD.)

by the tunnelling of these. It has been asserted that the tunnelling takes place by the conversion of some of the protoplasm into red corpuscles in a manner similar to that which occurs in the original formation of vessels in the embryo, but this is very doubtful. According to Thiersch blood-vessels may be formed by a process of channelling among the cells. The



wall of the vessel, infiltrated and softened by the inflammation, allows the blood plasma to pass out, and this forms canals which widen and admit the blood-corpuscles.

**Bone and cartilage** are, histologically, forms of connective tissue, and they are, in pathological processes, to some extent interchangeable with ordinary connective tissue. Hence in inflammatory newformations we may have, according to circumstances, these tissues produced along with or instead of connective tissue. This applies more particularly to inflammations in connection with bone. In fractures, for example, the so-called callus is composed, in many cases, of all three tissues.

**The Newformation of epithelium** is an essential part of the process of the conversion of a granulating wound into a cicatrix. This occurs by the process of karyomitosis, the existing epithelium furnishing the new cells. Hence this part of the process occurs usually at the margins of the wound, and will only take place in the midst of the wound if epithelial structures, such as hair follicles or glands, survive there. A certain limited amœboid movement is asserted by which the young epithelium passes into its proper place. The newly-formed epithelium is translucent and delicate, so that the granulations beneath show their colour through it. The epithelium remains somewhat translucent, like that of the stratum lucidum of the epidermis, and there is not a formation in the cicatrix of a proper malpighian layer. Living epithelium may be transplanted from a distant part or from another person, to the surface of a wound, where it may multiply. (See further on, under Transplantation.)

**In all other inflammatory newformations** the process is, in its elements, similar to that just described. For newformation to occur the inflammation must be prolonged for a considerable time, and it must not be too severe, hence usually a chronic inflammation. As it is connective tissue which is chiefly concerned in this process, the result is, in most cases, an increase of existing connective tissue, which takes the form of thickenings of membranes, induration of organs, etc. In such cases the process does not go through such definite stages as in the case of the granulating wound, but the various steps are present side by side. The tissue shows leucocytes, formative cells, and fully-formed connective tissue.

**Foreign bodies, or dead pieces of tissue**, frequently give rise to inflammatory newformation. If a foreign body be introduced among the living tissues, if not very virulent in its own nature, it sets up a mild chronic inflammation, with the result of producing a vascular rudimentary tissue like granulations. If the foreign body be permeable by this tissue, then the granulations will grow into it, and, as it were,



devour it, replacing it first by their own rudimentary tissue, which afterwards gives place to connective tissue. As this connective tissue is comparatively small in bulk, and tends to contract more and more, the result of the whole process is an **absorption of the foreign or dead substance** and the gradual disappearance both of it and of the tissue which has replaced it. But if the foreign body is not permeable, or only partially so, then the inflammation results in the production of a layer of connective tissue around it, and so the body becomes encapsuled.

Many instances of this might be given. If a piece of dead animal tissue be introduced into the body, as, for instance, a piece of liver previously hardened in chromic acid solution, or a piece of prepared catgut used to ligature a vessel, then the dead tissue is first replaced by rudimentary tissue which gradually shrinks away. Again, if a piece of a tissue or organ dies, then if severe inflammation is kept off, it is replaced by rudimentary tissue and absorbed. In fractures of bones it often happens that a piece is entirely separated and dies. Such a piece of bone may lie exposed in the wound in a compound fracture, and it has frequently been seen how it has been eaten into by the granulations and absorbed by them.

**The encapsuling of foreign bodies** is frequently seen. A parasite such as the trichina or echinococcus obtains a connective tissue capsule. Dead material in the body which is not permeable by the granulations is similarly treated, such as dried-in inflammatory products, which have formed first a caseous and then a calcareous mass. We frequently find such calcareous material surrounded by a fibrous capsule in the lungs and elsewhere.

#### IV.—THE ISSUES OF INFLAMMATION.

The manner in which inflammations conduct themselves till their conclusion, and the results which remain, have, to a large extent, been explained in what has gone before. We have here briefly to gather up the facts and bring them into relation.

1. **Resolution, or Restitutio ad integrum.**—These terms are applied to the subsidence of the inflammation and a restoration of the parts to their previous normal condition. This can scarcely take place except in acute inflammations, as the occurrence of newformation constitutes a more or less permanent lesion. In the case of acute inflammations, as soon as the irritant ceases to act, the phenomena, so far as the blood-vessels are concerned, will soon cease, and there will remain to be disposed of the exudation and the altered tissues. The exudation, so far as it consists of serous fluid and leucocytes, is, as a rule, readily



absorbed. It is generally placed so as to be in immediate communication with the lymphatics, and the leucocytes, by their own movement may pass into these. Where absorption is not so readily effected, the cells frequently undergo fatty degeneration, and this applies to other cells besides leucocytes, which may be in the exudation. Cells which have undergone fatty degeneration readily disintegrate, and the resulting fatty matter is easily absorbed as a general rule. In like manner, the fibrinous exudation may undergo fatty degeneration, or otherwise disintegrate and undergo absorption.

2. **Adhesion of inflamed surfaces** is a frequent result of newformation. In the case of the granulating wound, if two granulating surfaces, such as the flaps of a stump, be brought together, they coalesce, and the two layers ultimately form a single bond of union, the cicatrix. It is similar elsewhere. The inflamed surfaces of pleura, pericardium, peritoneum, etc., commonly coalesce and form permanent adhesions, composed of connective tissue. By means of these adhesions the blood-vessels of the two surfaces will communicate, and the adhesion will be a permanent one, the cavity being obliterated so far as adhesion has occurred. There may be, in some cases, a regeneration of tissue which may have been lost, but this is generally imperfect. (See under Regeneration.)

3. **Induration** is a frequent result when the connective tissue of an organ or considerable piece of tissue is affected by chronic inflammation. The new-formed connective tissue, like that of the cicatrix, is dense and tends to shrink. Hence the structures are rendered more dense and processes result which are sometimes designated by the term **Sclerosis**.

4. **Necrosis**, or death of a portion of tissue, is a frequent issue of acute inflammation. The dead piece of tissue may be absorbed in the manner already referred to, or it may be discharged as a slough.

#### V.—THE FORMS OF INFLAMMATION.

We have had occasion already to notice that the various phenomena of inflammation are not always present in every case, and one or other of them may be so pronounced as to give its character to the inflammation. Hence, names have been given according as the inflammation shows certain special characters. It may, however, be remarked that **hyperæmia** is a nearly constant accompaniment of inflammation, and in slight acute inflammations it may be almost the only phenomenon. **Exudation** is also of almost constant occurrence. Indeed, in the examination of the tissues after death, the presence of leucocytes or



round cells is often the most definite evidence of the existence of an inflammation.

In the various forms of inflammation now to be mentioned, it is not to be supposed that a classification of inflammations is attempted. These are merely names in current use which call for explanation.

1. **Parenchymatous inflammation** is a term introduced by Virchow to indicate that the inflammation affects mainly the tissue elements. It is characterized by cloudy swelling and fatty degeneration of the structures. It has been referred to at p. 134.

2. **Interstitial inflammation** is almost the converse of the former. In it the inflammation affects chiefly the connective tissue, which forms the supporting stroma of organs. In acute inflammations there will be an infiltration of the connective tissue with leucocytes, and in some cases this may be so intense as to give rise to infiltration of pus. In chronic inflammations there will be newformation of tissue, usually resulting in **induration**. A frequent consequence of chronic interstitial inflammation is atrophy of the proper parenchyma of the organ, produced by the shrinking of the new-formed tissue, partly acting directly on the parenchyma and partly compressing the blood-vessels.

3. **Suppurative inflammation**.—From what has gone before it will appear that, for the most part, it is in intense inflammations produced by virulent microbes that suppuration will occur. **Phlegmonous inflammation** is a very acute inflammation accompanied by suppuration and necrosis. It is usually applied to such inflammations of the skin and subcutaneous tissue, of which erysipelas affords the best example.

4. **Infective inflammations** imply the presence of microbes, which by their multiplication, produce an extension of the inflammation from its original seat. The infection usually extends locally by means of the serous spaces and lymphatics (as in erysipelas), or it may be carried by the blood to distant parts, and produce inflammations by **metastasis**. (See under Infective Embolism.) The infective inflammations are mostly suppurative.

5. **Ulcerative inflammation, Ulceration**.—These terms imply necrosis or loss of tissue, generally of a progressive character. The loss of tissue may be rapid, and involve the formation of visible sloughs, or it may be gradual, a molecular necrosis, the dead particles being absorbed or removed in the exudation. These results are produced by very intense irritants, and the inflammations are usually of an infective and suppurative character. While ulceration expresses the process of progressive destruction of tissue, the ulcer is the open exposed gap in the tissue. If the ulceration pauses, then the ulcer becomes covered with granulations and is converted into a granulating wound.



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## SECTION VI.

## RETROGRADE CHANGES.

NECROSIS *caused by direct injury, by obstruction of arteries, spasm of arteries, obstruction of veins, nervous influences, assisted by weakness of the heart. Forms of necrosis determined chiefly by inflammation and putrefactive changes. Various forms of gangrene; coagulation-necrosis and caseation. Issues of necrosis. SIMPLE ATROPHY, its physiological type. General and local atrophy. ALBUMINOUS INFILTRATION, also in general and local forms. FATTY DEGENERATION, a transformation of nitrogenous principles. General and local forms. Character of lesion and results. FATTY INFILTRATION, in connective tissue; in the liver. PATHOLOGICAL PIGMENTATION—Origin of pigment. Alterations of physiological pigmentation; pigmentation from inflammation and hæmorrhage; melanæmia; icterus, hepatogenous and hæmatogenous; pigmentation in tumours and from without; pigmentary atrophy. AMYLOID DEGENERATION—Causation and nature of process: changes in tissues and seat; local amyloid degeneration; corpora amylacea. MUCOUS, COLLOID, AND HYALINE DEGENERATIONS. CALCAREOUS INFILTRATION, mainly in dead or obsolete structures; characters and effects.*

UNDER this designation are included a number of conditions, all of which imply a degradation of the nutritive processes in the tissues. As the cells are concerned in the nutrition of the tissues, it is for the most part these which are at fault, although it may be that in some cases the most manifest visible changes are not in them. The most extreme case is where the nutrition ceases altogether, and the structure dies. Short of that, we have various lesions manifesting themselves. Thus there is a simple diminution of the vitality, and the structures dwindle. Again, the chemical constituents of the structures change, splitting up, it may be, into more elementary principles. Or the tissues are unable to prevent the deposition in them of extraneous material, which is thus infiltrated into them. These last are called infiltrations, whereas the conditions in which there is a degradation of the normal constituents into lower chemical substances are designated degenerations.



## NECROSIS : GANGRENE : MORTIFICATION.

The term **Necrosis** is equivalent to local death of tissue, and includes all forms of lesion in which a part of the body loses its vitality. **Gangrene** has a more limited significance, being applied chiefly to cases in which the necrosis is accompanied by putrid decomposition of the part, and especially to death of considerable portions of the external parts of the body. **Mortification** has a similar meaning. **Sloughing** is the death and separation by ulceration of smaller parts of the soft external tissues. **Sphacelus** has a similar significance. Necrosis is sometimes used in the limited sense of death of bone or cartilage.

I. **Causes of necrosis.**—An agent may cause necrosis by acting directly on the tissues, or may do so indirectly by interfering with their blood-supply or innervation.

Instances of **Direct action** are afforded by chemical agents, by traumatic action, and by extremes of temperature.

**Chemical agents**, such as strong acids or alkalies, destroy the vitality of the tissues by their direct action, and they sometimes dissolve the tissues at the same time. Necrosis is a very frequent result of the action of **microbes**, and this effect is produced by the chemical principles evolved by these. The products of ordinary putrid decomposition may produce necrosis, as, where decomposing urine is extravasated into the tissues. We see also in the cases of pyæmia, erysipelas, tuberculosis, syphilis, diphtheria, that when microbes settle in the tissues and multiply, they commonly give rise to necrosis when acting in a concentrated form. The chemical products concerned are presumably animal alkaloids. (See under Ptomaines, further on.)

**Traumatic action**, besides its direct effect in injuring the tissues, will produce necrosis more frequently by obstructing or rupturing the vessels.

**Extremes of heat or cold** produce necrosis, partly by affecting the living structures directly, and partly by their effect on the vessels. The ear of a rabbit immersed in water heated to 130° to 136° F., or in a freezing mixture reduced to about 0° F., suffered necrosis even when the immersion was for a very short time. If the temperature was less extreme a short immersion produced intense inflammation, and a longer immersion produced necrosis. (Cohnheim.)

**Pressure** on a part produces necrosis when long continued, and it does so chiefly by emptying the vessels, and especially the capillaries. Pressure from within is exemplified in the case of abscesses and tumours which advance to the surface and cause sloughing of the



skin. External pressure is seen to produce necrosis in the case of bed-sores, or where bandages or splints, by pressing on a bony prominence, cause sloughing of the skin over it.

**Obstruction of arteries** is a frequent cause of necrosis. As the nutrition of the tissues depends on the capillary circulation, obstruction in an artery will scarcely produce necrosis, unless it brings about a complete stasis in the capillaries. (See under Embolism.)

Besides obstruction of arteries by embolism, there is very frequently a partial interference with their calibre as a result of atheroma, especially when thrombosis is superadded. In **senile gangrene** there is always atheroma of the arteries of the lower limb, and this even without complete obstruction may be the cause of the gangrene, although weakness of the heart and of the tissues resulting from old age may contribute. **Softening of the brain** in old people is similarly produced from atheroma. These softenings are frequently in the cortex of the brain, and lead as much to weakness of mind as to motor paralysis, whereas softening from embolism is usually central, and leads to more definite paralysis.

**Spasm of arteries** is an occasional cause of gangrene as in Raynaud's disease (see p. 63). It has also been assigned as the cause of gangrene in poisoning with **Ergot of Rye**. In former days there used to be epidemics of what is now recognised as Ergotism from eating bread made with grain in which ergot was present. The symptoms consisted of lesions of sensation and violent cramps, followed by redness of the skin, and sometimes culminating in necrosis of the tips of the fingers and toes, or of the nose and ears. This result is ascribed by some to spasm of the arteries, a view which is confirmed by an observation of Recklinghausen, who found by experiment in fowls that under the influence of ergot the arterioles of the cock's comb and of the tongue showed a violent and persistent contraction, during which thrombosis occurred, obstructing or occluding their calibre. Besides this action on the vessels, the poison may have a directly poisonous effect on the tissues, and the necrosis may be further assisted by the injury to which an anæsthetic part is exposed.

**Obstruction of veins** seldom produces necrosis, as these vessels anastomose so freely that stasis in the capillaries will rarely occur. It takes place, however, when a piece of intestine is incarcerated in a sac with a narrow neck, as in a strangulated hernia, in which case gangrene is frequent. Even extensive thrombosis may lead to gangrene.

**Nervous influences** frequently contribute to the production of necrosis. Lesions of the peripheral nerves, the spinal cord and brain, are sometimes followed so rapidly by the formation of sloughs in parts of the



body exposed to pressure in lying in bed, that the term **Acute bed sore**, or **Acute decubitus**, has been applied. This has been ascribed to the primary lesions affecting the trophic nerves, but it remains doubtful to what extent the immobility of the patient, his constant retention of the same posture, and the alteration in the circulation, may account for the necrosis without calling in the action of trophic nerves.

A similar difficulty exists as to the necrosis in **Anæsthetic leprosy**. Here the affected parts are devoid of feeling and the vaso-motor nerves are implicated, hence the parts are more exposed to injury and to variations in temperature, which are not compensated by alterations in the circulation.

In all forms of necrosis **Weakness of the heart** may exercise an influence, and in some cases it may even be the main element. In extreme cases of general weakness, and in some cases of specific fever, there may be necrosis of the extremities partly due to weakness of the heart and partly to alteration in the constitution of the blood.

The various tissues comport themselves somewhat differently in relation to the causes of necrosis, or in other words they are in different degrees able to survive a deprivation of blood. For example Litten found that if he ligatured the renal artery in a rabbit for  $1\frac{1}{2}$  to 2 hours and then removed the ligature, the circulation was perfectly restored, and the blood-vessels and connective tissue survived; while the epithelium of most of the convoluted tubules underwent necrosis. Ehrlich and Brieger found that a suspension for one hour of the circulation in the lumbar part of the spinal cord, caused necrosis of the grey substance, while the white substance was not affected. Muscle seems also peculiarly sensitive to deprivation of blood. Thus in embolism of the coronary artery of the heart the muscular fibres will die before the circulation can be re-established, while the connective tissue survives. In this relation skin, bone, and connective tissue possess great powers of resistance, while nervous tissue, muscle and the secreting tissue of glands are more vulnerable.

The tissues again may be rendered unduly susceptible of necrosis. Anæmia and passive hyperæmia render the tissues more vulnerable. Diabetes has a similar effect; boils and carbuncles are common in that disease, and slight injuries are liable to go on to necrosis. Again children who are in extreme states of inanition, especially after acute fevers, are liable to **Cancrum oris** or **Noma** in which extensive necrosis of the soft parts in the neighbourhood of the mouth occurs.

**II. Forms of necrosis, and changes in the tissues.**—The changes which the dead tissues undergo vary considerably according to circumstances, and the resulting appearances are so different that special names are given and forms described according to the appearances presented. From what has gone before, it will appear that **inflammation** frequently goes along with necrosis; the two are sometimes produced simultaneously by the same cause, or the dead structures may in themselves, or by the products evolved by them, produce inflammation.

The circumstances which determine the form which the necrosis will



assume are chiefly these—the position of the dead structure, whether internal or external, whether protected from the access of air or not;



Fig. 27.—Formation of a pyæmic abscess in the kidney. Section stained with aniline brown. The necrosed portion *a* shows no structure and is not stained. The surrounding inflamed area (*b, b, b*) is deeply stained.

the presence or absence of acute inflammation; the bulk of the dead piece; its structure and chemical constitution. The most important circumstance is whether **putrefactive changes** occur or not. Dead pieces of tissue, like all dead animal matter, are liable to decomposition under the influence of microbes. In most cases of necrosis in external parts and in the lungs, the air finding access carries with it the microbes concerned in these processes, and putrefaction is the result. In this case the term gangrene is usually applied.



Fig. 28.—Capillary blood-vessel (*b*) in kidney filled with micrococci, from a case of pyæmia. There is a tubule on either side (*a, a*), the nuclei of which are visible except in the neighbourhood of the capillary.  $\times 650$ .



In most cases of necrosis the process is characterized by the **disappearance of the nuclei of the cells**. In Fig. 27, for example, necrosis of a portion of the kidney has occurred by embolism of a small artery. The section has been stained so as to bring out the nuclei of cells, but the dead portion in the centre has no nuclei. In Fig. 28 also, necrosis of the renal epithelium has occurred in consequence of the presence of microbes in a capillary, and it is seen that the nuclei have disappeared.

1. **Dry gangrene or Mummification**.—These terms are applied in the case of external parts, when the circumstances are such that there is little moisture in the parts. Hence it will not occur where acute inflammation precedes or accompanies the process. It is mostly met with in cases of embolism or other obstruction of the arteries, and is characteristic of senile gangrene and of Raynaud's disease. The part undergoes a gradual deepening of colour; at first merely livid, it passes into purple, deep blue, and even black. This is due to the fact that the blood-pigment is dissolved out and stains the tissues, which deepen in colour as the pigment becomes concentrated by drying. The cuticle generally gets raised by the accumulation of a red fluid beneath it, and bullæ are formed. If the cuticle separates, evaporation is accelerated. The part, which is usually the extremity of the lower limb, gradually shrinks, and is converted into a hard black mass, often with a mouldy smell. In dry gangrene there is putrid decomposition, but as there is a deficiency of fluid this occurs to a subordinate degree.

2. **Moist gangrene, Sphacelus, Sloughing**.—In the conditions designated by these names, putrid decomposition plays a prominent part, and the tissues are separated in a softened condition. Moist gangrene is the condition where a considerable portion of the body has died and is undergoing separation. It is well seen in cases where a portion of the leg has died in consequence of injury. The parts which were at first hot, red, and painful, become mottled with brown, blue, and black, and the surface often presents blisters. The part becomes cold and darker except at the margin, where a dusky red line of demarcating inflammation appears. The tissues are universally stained with blood-pigment, the cuticle gives way and putrescence advances, causing breaking down of the tissues, which are separated in all stages of softening, the more resistant tissues such as bone and cartilage retaining their form. The part exhales a strong odour, and its juices contain fluid fat, phosphates, extractives, as well as multitudes of microbes.

3. **Desiccation without putrescence**—This is somewhat similar to dry gangrene, except that, occurring in internal parts, there is no decomposition. The part simply dries in and shrivels. The most



striking example of this is afforded by extra-uterine pregnancy, in which the foetus after its death is retained in the abdomen of the mother. It may remain for many years simply drying in and getting encased in a capsule, which becomes impregnated with lime salts. Virchow in a case of this kind found muscle, connective tissue, and vessels still recognizable after twenty years. In twin pregnancies also, one of the foetuses may die at an early period, but, being retained, it is born along with the other in the form of a dry flattened object.

4. **Softening without putrescence, Colliquifaction.**—This is almost peculiar to necrosis of the nervous system in consequence of obstruction of arteries. The process is accompanied by fatty degeneration.

5. **Coagulation-necrosis.**—This term was introduced by Cohnheim, and the subject has been elaborated by Weigert. When necrosis occurs in internal parts which are rich in cells, then the tissue frequently becomes converted into a solid, firmly compacted mass. Weigert compares the process to the coagulation of the blood, and asserts that the cells in dying enter into combination with the fibrinogen contained in the fluid which permeates the tissue. In order to this process the tissue must be a very cellular one, and an abundant supply of fluid must be present. The embolic infarction in the spleen and kidney forms the most typical example. The infarction forms a stiff, firm wedge which may be pale or may contain blood. The process is characterized by the disappearance of the nuclei from the affected structures.

A peculiar change which occurs in voluntary muscle is also regarded by Weigert and others as a form of coagulation-necrosis. This is the process described by Zenker as **waxy degeneration of muscle**, which is also designated colloid and **hyaline degeneration**. It is brought about by direct injury to the living muscle. It also occurs in certain febrile states where the temperature runs high, especially in typhoid fever and

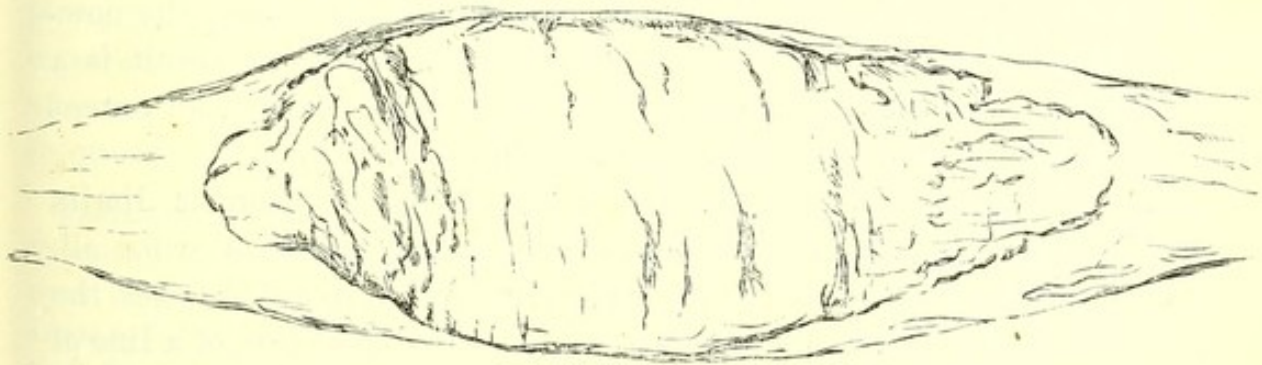


Fig. 29.—Coagulation-necrosis in muscle. The sarcolemma surrounds the coagulated clump, drawing together at the ends.  $\times 350$ .

phthisis pulmonalis, and is sometimes seen in paralysed muscles. It may be produced artificially by freezing the muscle of a living animal, or by injuring it before the occurrence of post-mortem rigidity. The



change is met with, in the human subject, chiefly in the diaphragm, the rectus abdominis, and the adductors of the arms. It consists in a coagulation of the contractile substance, the fibre being converted into a homogeneous translucent cylinder. The affected fibres are brittle, and the cylinders get broken up, often presenting transverse cleavages, or giving origin to oval clumps as in Fig. 29.

In **atheroma** of arteries there are often translucent 'structureless' pieces in the walls, which Weigert also claims as examples of coagulation-necrosis. Similarly some tube casts in the kidney are supposed to originate by necrosis and coagulation of the epithelium, and the membranous exudation in diphtheria is asserted to have a similar origin.

**Hyaline degeneration**, as used by Recklinghausen, to a large extent covers the same ground as the coagulation-necrosis of Cohnheim and Weigert. (See further on.)

6. **Caseous necrosis**.—This is a condition closely allied to coagulation-necrosis. In the latter the necrosis occurs for the most part suddenly in structures hitherto unaltered, whereas caseous necrosis is always part of another pathological process. It is highly characteristic of tuberculosis, so much so that the term is nearly equivalent to the older designation tuberculization, and caseous matter is virtually the same as crude or yellow tubercle. It is also frequent in syphilitic newformations. In all cases there is a great abundance of cells, which are not the normal cells, but are produced by exudation or newformation. The necrosed tissue undergoes a process of condensation or coagulation, accompanied by fatty degeneration, the result being a somewhat dense brittle matter, which has been compared to cheese.

In its microscopic details the process involves a complete obscuration of structure. The nuclei disappear as in coagulation-necrosis, and the presence of abundant fine fat granules renders the structure very opaque and homogeneous. As the necrosis involves not only the new-formed cells, but the tissue in which they are seated, the result is a disappearance of the details of the tissue, sometimes to such an extent as to render the structure difficult of identification.

III. **The issues of necrosis**.—In many cases the necrosis limits itself at once, the agent which produced it having acted once for all. It only remains to dispose of the dead structures. In other cases the limitation does not take place so directly, and the formation of a **line of demarcation** is anxiously looked for. This is frequently the case in traumatic necrosis where it may for some time be doubtful to what extent the tissues have been injured beyond recovery. The co-existence of inflammation, especially when this is associated with decomposition, often renders the limitation of the necrosis more difficult. In



infective processes also, such as tuberculosis, the necrosis follows the advance of the lesion. Again in senile gangrene where the arteries are seriously obstructed, the starting point of the necrosis may be a trivial injury, and its progressive extension may go on without any signs of limitation for a considerable time.

In the **disposal of the dead tissue** inflammation plays a most important part. We have seen that violent inflammation is often produced by the same cause as the necrosis, or may supervene on it. This will mostly be the case in external parts where decomposition occurs. The inflammation is characterized by hyperæmia and exudation, and commonly goes on to suppuration. A layer of pus comes thus to divide the dead tissue from the living, and the dead is cast off as a slough. There remains a suppurating wound or ulcer. This loss of tissue, whether molecular or as a massive slough, is the essential feature of an ulcer of whatever kind, and a suppurating surface will usually be left.

In internal parts, if the necrosis be accompanied or followed by the production of irritating chemical substances, then a violent inflammation will be produced around. It is so in the case of pyæmia, where there is septic embolism. In this case the inflammation will usually be violent enough to produce suppuration, and the result will be the formation of an abscess. (See Fig. 27, p. 147).

In the case of internal parts where there is no disturbing decomposition, or in external parts which are protected from septic contamination, the inflammation is of a much milder character. The necrosed portion now comes to act as a foreign body or dead piece of tissue; and is subject to the changes already described at p. 138. The dead tissue is often eaten into and replaced by vascular granulations, which finally contract and leave a small residue of connective tissue; or the dead piece is encapsuled and may lie quiescent. Not infrequently the encapsuled tissue undergoes infiltration with lime salts, as we shall see afterwards. In the case of necrosis in bone the external capsule is frequently composed of new-formed bone.

**Literature.**—CARSWELL, *Elementary forms of disease*, Art. Mortification, 1834; VIRCHOW, *Handb. d. spec. Path.*, vol. i.; PAGET, *Lect. on surg. path.*, p. 340; COHNHEIM, *Allg. Path.*, vol. i., p. 526, and *Die embol. Proc.*; RECKLINGHAUSEN, *Allg. Path.*; LITTEN, *Zeitsch. f. klin. Med.*, vol. i.; KOCH, *Traumatic infective diseases* (Syd. Soc. transl.); BURDON SANDERSON, *Path. trans.*, xxiii., *Brit. Med. Jour.*, 1877; WEIGERT (Coagulation-necrosis), *Virch. Arch.* lxxix, p. 89; CHAUVEAU, *Nékbiose et Gangrène*, *Bullet. de l'Acad. de Méd.*, 1873; BRIEGER, *Ueber Ptomaine*, 1885; AITKEN, *On animal alk.*, 1887; BROWN, *A treatise on animal alk.*, 1887 (Introduction by Gautier).



## SIMPLE ATROPHY.

By this term is meant a simple diminution in the nutritive activity of the structures, and a consequent diminution in size without further change. Strictly speaking, we should distinguish from this a smallness due to defective growth, to which the terms **Hypoplasia** and **Aplasia** are applied. But atrophy is frequently used so as to include these conditions.

**Physiological atrophy.**—There are certain normal processes of decay which occur in the body. At certain periods, for instance, the milk teeth drop off, and this is effected by an atrophy of the fang so that the crown is shed. At a still earlier period the thymus gland atrophies. Then again throughout life there is a continual shedding of the hair; if a cast-off hair from the eyelash be examined under the microscope, it will be seen that its bulb is atrophied, and this is the cause of its being shed. In some persons the hair of the scalp is largely shed at a comparatively early age, without being properly reproduced, there being here an atrophy of the hair sheath and papilla. Then there is the normal atrophy of the tissues generally, which occurs in old age. The atrophy of old people may in many cases be due to some organic disease whose symptoms are not manifest; but we are all familiar with the healthy old person with shrivelled hands and face and plicated skin.

In all these cases there is a kind of intention in the tissues, so to speak, according to which they live a certain period and then decay. As Paget has pointed out, such atrophies may almost be regarded as active processes. The fall of the leaf is due to an active absorption or atrophy of the fibres uniting it to the stem; if the leaf dies before its time, or is killed, it remains hanging, but in the natural course it drops when its time is come. So with our tissues and the whole organism, there is a period to their activity. The period varies in different persons, and in this respect hereditary influences have an important bearing. Just as these largely determine the period of growth of the body and its rapidity, so do they influence the duration of activity of the tissues. This is plainly seen in the case of the hair; baldness runs in families, just as longevity does.

**Causation and Forms of atrophy.**—Atrophy in many cases depends on some interference with the supply or alteration in the quality of the nutritious material supplied to the tissues. It is also related frequently to diminution in the function of the parts, while in some cases it depends more directly on interference with the nervous arrangements.

**General emaciation** indicates that the tissues generally have been affected in such a way as that their nutrition is diminished. This will occur as the result of an alteration in the blood. The blood, being the



vehicle for the conveyance of nutriment to the tissues, may be impoverished because of a direct interference with the food-supply, as in starvation, in stricture of the œsophagus, excessive vomiting, diarrhœa, etc.; or, there may be an excessive consumption of the nutritious material, in cases of excessive discharges, as in phthisis pulmonalis, or ulcerating cancers, or there may be, as in fevers, an increased consumption of the nitrogenous elements of the tissues.

In general emaciation the various constituents of the tissues do not atrophy in an equal degree. According to experiments by Chossat, in which animals were deprived of food, the fat and blood diminished most, next to them came the muscles and the abdominal glands, while the bones and central nervous system diminished least.

**Senile atrophy** is closely related to the physiological atrophies already mentioned. The atrophy of some tissues in old age renders them more liable to pathological processes. Thus the bones, being diminished in size, and having proportionately less animal matrix, are more liable to fracture; and the lungs, having lost their supporting tissue, are more liable to emphysema (senile emphysema). The brain also suffers atrophy in old people, and the kidneys frequently do so.

**Atrophy from disuse.**—This manifests itself chiefly in the muscles and glands. When a limb is disused from being set up in a rigid apparatus, or from paralysis, the muscles undergo atrophy, and even the bones diminish if the condition be prolonged.

**Atrophy from pressure** is exemplified in the atrophy which occurs as a consequence of the advance of tumours and aneurysms, or external pressure from stays, etc. In the last mentioned case the liver frequently suffers considerable atrophy. Atrophy is also not uncommon in organs which are the seat of interstitial inflammation, the new-formed connective tissue, by its direct pressure on the proper tissue, or by obstructing the blood-vessels, causing atrophy.

**Atrophy from nervous lesions.**—This is a somewhat wide subject and will be more fully considered in the special part of this work. In ordinary motor paralysis there is atrophy from disuse. But there are atrophies of a more active kind following lesions of nerves and of the spinal cord which are referred to interference with the trophic nerves or centres. These include atrophy of the nerve-fibres and of the muscles. According to Charcot a muscle or nerve atrophies when cut off from its trophic centre, and this may be effected by interruption of the conductivity of nerve-fibres, or by destruction of the centre. The atrophy of the muscles in **lead-palsy** belongs to this class.

**Hemiatrophy of the face** belongs to the class of neurotic atrophies. It affects the soft parts of the one half of the face and of the tongue,



while the bones are not affected unless the disease has occurred in early life, and even then the bones are unequally affected. The lesion is probably due to interference with the nerves, perhaps in their passage through the cranium or at their ganglia. A **hemiatrophy of the body** sometimes occurs from cerebral lesions in the fœtus or young child. This may be in the form of a crossed hemiatrophy, the face and extremities being affected on opposite sides.

In the various forms of atrophy, with the exception of general emaciation, the proper functioning tissue is that which chiefly suffers diminution. Thus in muscles it is the contractile substance, in glands the secreting cells, in nerves the nerve-fibres, which are specially affected. The atrophy of the proper tissue is often accompanied by **newformation in the accessory structures**. Thus, atrophy of muscle is often associated with increase of the interstitial connective tissue. This connective tissue frequently becomes the seat of fatty infiltration, so that adipose tissue largely replaces the muscle, bringing about a pseudo-hypertrophy. (See under Hypertrophy and Fatty Infiltration.) A similar process frequently occurs in and around disused glands.

**Literature.**—PAGET, *Surgical Path.*, 3rd ed., p. 69; RECKLINGHAUSEN (*Neurotic atrophy*), *Allg. Path.*, p. 326; CHARCOT, *Senile diseases* (*Syd. Soc. trans.*), 1877, *Dis. of nerv. syst.* (*Syd. Soc. trans.*), 1881; CHOSSAT, *Rech. exp. sur l'inanition*, 1843.

#### ALBUMINOUS INFILTRATION—CLOUDY SWELLING.

The condition designated by these terms was first described by Virchow, and regarded by him as characteristic of **Parenchymatous inflammation**. The term **Parenchymatous degeneration** has a similar signification.

It is possible to distinguish a local from a general cloudy swelling. The **local form** occurs in parenchymatous inflammation, more especially of the kidneys, and is sometimes the most pronounced evidence obtainable post mortem of the existence of that condition. The **general form** occurs in most febrile diseases and is very characteristic of some.

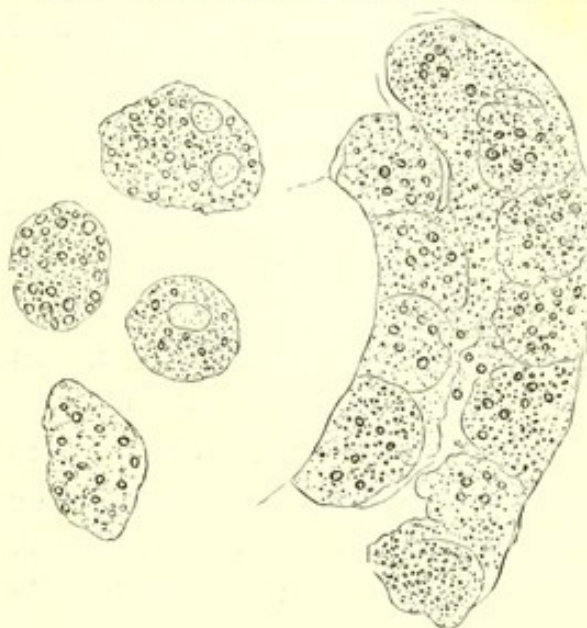


Fig. 30.—Cloudy swelling of renal epithelium with slight fatty degeneration. A portion of a tubule is shown, and some isolated cells. From a case of acute phthisis pulmonalis.  $\times 350$ .

It is met with in the specific fevers, in erysipelas, diphtheria, acute phthisis pulmonalis, etc., and seems to be related in these cases to the



high temperature and the altered state of the blood. It also occurs as one of the results of acute poisoning with phosphorus, arsenic, mineral acids. In the general form the lesion is diffused through various organs, but affects especially the liver, kidneys, heart, and voluntary muscles.

In these various cases the condition seems to be due to an irritation of the cells, which are induced to absorb more albumen than they can assimilate. The cells are enlarged, and they are clouded with albuminous granules (see Fig. 30), which obscure the nuclei. The condition implies a qualitative defect in the cells, although quantitatively there is excess. This is nearly always shown by the co-existence of a minor degree of fatty degeneration. (See Fig. 30.) The fine fat granules may be obscured by the albuminous granules, but if the albumen be dissolved by adding liquor potassæ, or a dilute mineral acid, the fat comes out very prominently.

The organs affected have, to the naked eye, a characteristic appearance. They are enlarged, sometimes to a high degree, and have, on section, a bulky appearance, while the tissue has a grey, opaque, dull character. In acute fevers the enlargement of liver and kidneys is sometimes very great.

**Literature.**—VIRCHOW, Cellular Pathology, transl. by Chance, also various papers in his Archiv.

### FATTY DEGENERATION.

In this condition, which affects the cells of tissues, we have a change in the chemical composition of the cell-contents; the albuminous constituents split up and yield fat.

**Causation.**—Fat may be formed in the animal body either from the carbo-hydrates or albuminous substances of the food. In the case of fatty degeneration the fat is derived from the albuminous constituents of the tissues. This implies that these constituents break up, yielding their nitrogen in some lower form of combination which is usually carried off, and leaving the fat in the tissue. In accordance with this we find that, where general and extensive fatty degeneration occurs, there is simultaneously an excess of urea or other extractives in the urine. The splitting up of the albuminous constituents of cells implies a most serious alteration in their chemical constitution. In all such cases, therefore, it is to be inferred that the vitality of the cells is greatly reduced, and in some cases the condition approaches to, or is associated with, necrosis.

The proof that fat may be formed from carbo-hydrates lies in the fact that bees will produce wax when fed on honey alone, this being a solution of sugar (Gund-



lach). The formation of fat from nitrogenous substances is proved by various facts. Thus the tissues of the body, including the nitrogenous constituents, are sometimes changed after death into **Adipocere**, a peculiar waxy substance, composed of fatty acids combined with ammonia and lime instead of glycerine, and therefore more strictly a soap than a fat. This substance is occasionally found in graves, and it has been produced by leaving the dead body in running water for a time. Again, it has been shown that, in lactating animals fed with animal food carefully deprived of fat, the milk is even more abundant and rich in fat than in animals fed on diet containing much fat. Fresh milk becomes richer in fat and poorer in caseine (which is nitrogenous) during the first day after its withdrawal from the mammæ (Hoppe-Seyler).

The most conclusive proof that albuminous tissues yield fat is afforded by the effects of poisoning by phosphorus. When a dog has been deprived of food till all its spare fat has been exhausted and the nitrogen in the urine has reached a constant minimum of 8 grammes in the twenty-four hours, the administration of small doses of phosphorus causes a marked increase of the nitrogen, which may reach nearly 24 grammes. This is coincident with a very large increase of fat in many internal organs.

The causes of fatty degeneration may be divided into those which, depending on some morbid condition of the blood, act on many organs, and those which have simply a local influence. So we may speak of a general and a local fatty degeneration.

**General fatty degeneration** is produced by certain poisons, pre-eminently by phosphorus, but also by arsenic, antimony, iodoform, chloroform, and others. It occurs also in some general diseases in which the blood is greatly altered, in acute yellow atrophy of the liver, in some fevers, in pernicious anæmia, and in some other forms of anæmia. It has been produced artificially by confining animals in an over-heated space for thirty-six hours (Cohnheim). In these cases the altered blood has acted on the cells of the tissues and caused them to alter their chemical constitution. The change occurs mostly in the parenchyma of organs, as in the hepatic cells, the renal epithelium, the striated muscular fibres, especially those of the heart, but is also seen in some cases in other structures, such as the intima of arteries.

**Local fatty degeneration** is frequently the result of deprivation of blood, as where an artery is occluded. In the brain, occlusion of arteries is followed by softening of the cerebral tissue, a species of necrosis, but this is associated with the appearance of cells filled with finely divided fat (the so-called compound granular corpuscles of Gluge). It is true that the fat here may be partly derived from the myelin of the nerve-fibres, but these corpuscles are present in the grey substance where there is no myelin, and fat is also visible in the walls of the blood-vessels (as in Fig. 31). Inflammation is a frequent cause of local fatty degeneration, especially in parenchymatous organs, where the



cloudy swelling often goes on to fatty degeneration. (See Fig. 30.) In quickly growing tumours, and even in slowly advancing cancers, the

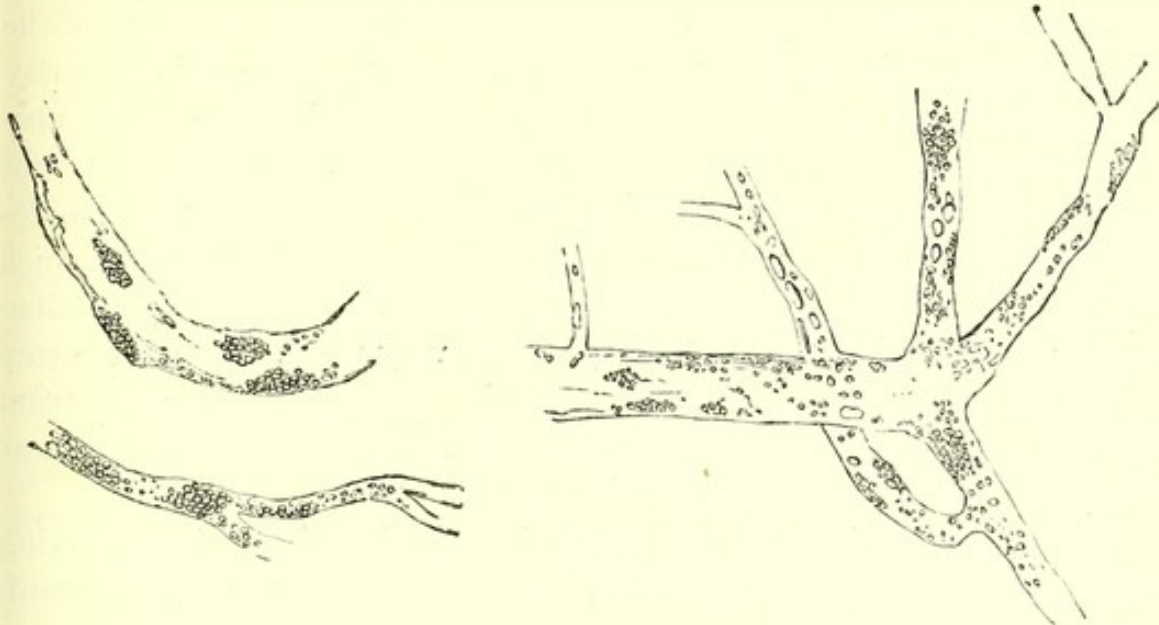


Fig. 31.—Fatty degeneration in the cerebral vessels in softening of the brain. (PAGET.)

cells frequently undergo fatty degeneration. Lastly, nerve-fibres which have been divided show not only atrophy, but also a fatty degeneration.

Some authors have endeavoured to account for fatty degeneration on the supposition that it is due to a deficiency of oxygen. It is said that in general fatty degeneration the blood is deficient in oxygen, and in the local form the tissues are deprived of oxygen. In the former case, however, there is usually an obvious alteration of the blood apart from simple anæmia, while in many local fatty degenerations there is no deprivation of oxygen, as in inflammations, in tumours, and after section of nerves.

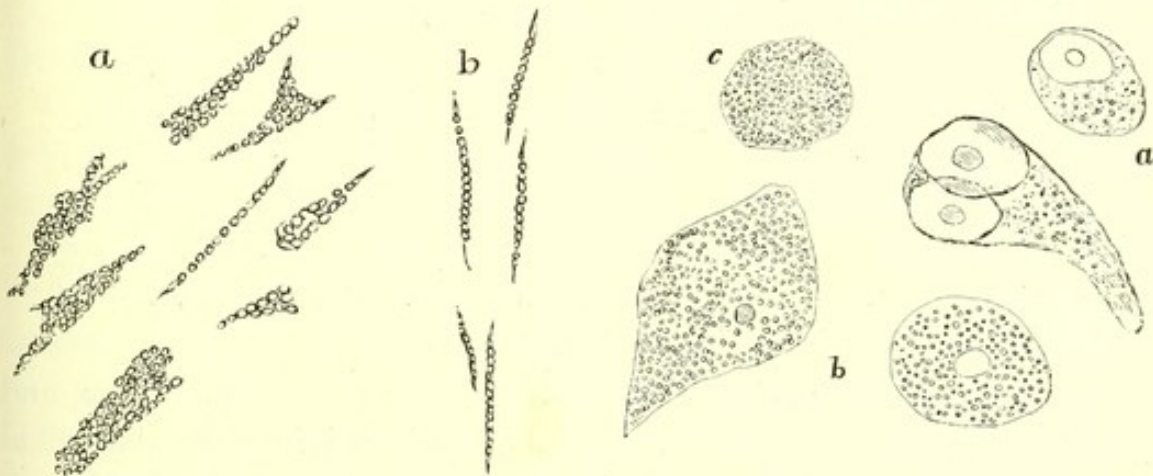


Fig. 32.—Fatty degeneration in an atheromatous aorta. The shapes of the cells brought out by the fat; *a*, from internal coat; *b*, muscle cells from middle coat.  $\times 350$ .

Fig. 33.—Fatty degeneration of cells in a cancer of the mamma; *a*, slightly affected; *b*, more so; *c*, completely fatty—the compound granular corpuscle.  $\times 350$ .

**Characters of the lesion.**—The degeneration occurs mainly in the cells of the tissues. The fat, arising as it does by the chemical decom-



position of the protoplasm of the cell, appears in the form of fine drops or granules, which are strongly refracting. (See Figs. 31, 32, 33.) These granules are still separated from each other by the remains of the cell contents and are therefore isolated. It may happen in this way that, as in Fig. 32, a fatty degeneration occurring in a structure may render its constituent cells unusually distinct, their form being brought prominently out by the fat in them. As time goes on, the fat granules increase till the whole cell is filled with fine refracting oil drops, which still remain isolated (see Fig. 33), being each surrounded by an albuminous envelope. The process is, in fact, very much like what occurs in the cells of the mammary gland in the secretion of milk, the colostrum cells being like the fully degenerated cells, which are often described as compound granular corpuscles.

When finely divided fat suspended in fluid is present in the living tissues, it is very readily absorbed. We know how readily the emul-

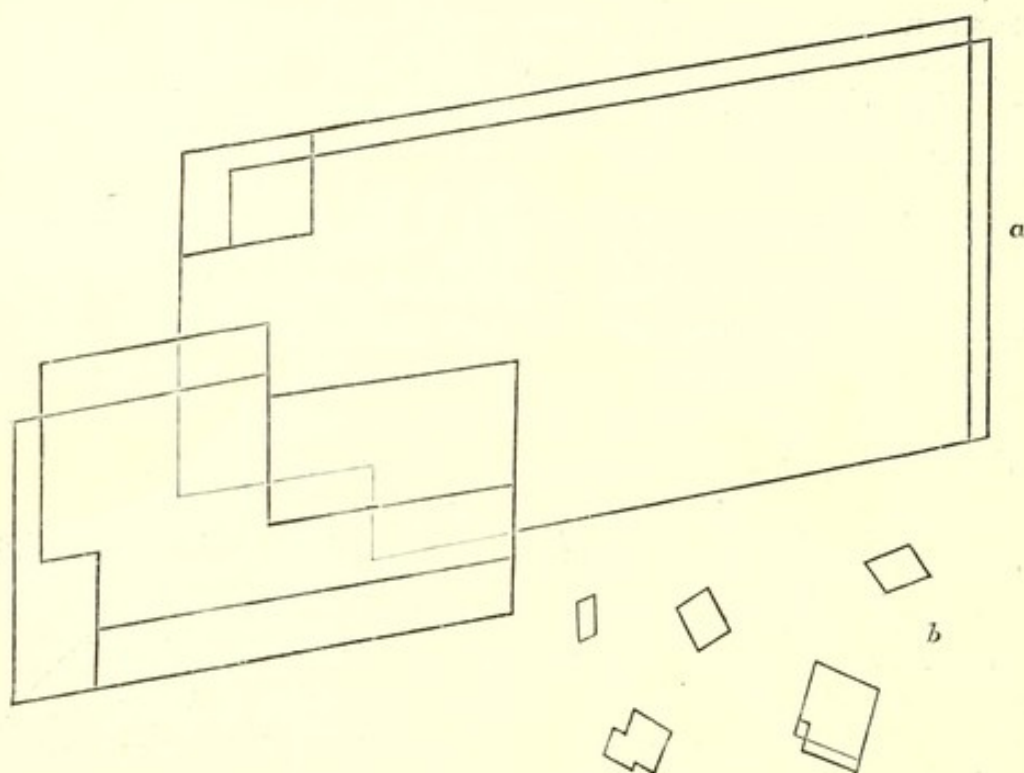


Fig. 34.—Crystals of cholestearine; *a*, large ones from an old hydrocele; *b*, from stagnant bile in gall bladder.  $\times 350$ .

sionized fat in the alimentary canal is taken up by the epithelium, and passed on into the lacteals. When milk is injected into the abdominal cavity of a living animal, or even laid on the surface of the diaphragm after death, it very quickly passes into the lymphatics. In the case of fatty degeneration of cells, if fluid be present, the cells disintegrate, an emulsion is formed, and absorption occurs just as in the case of milk.

But sometimes the fatty degeneration occurs in connection with structures not adapted to absorption, as in a hydrocele, where the



tendency is rather to transudation, or in an ovarian tumour. In that case, the fat undergoes further changes, resulting usually in the production of crystals of **cholestearine** (Fig. 34) or **margarine**.

**Cholestearine** occurs as a normal constituent of the central nervous tissue and bile, in which latter it is dissolved. Its crystals are rhombic tables whose angles measure  $79^{\circ} 30'$  and  $100^{\circ} 70'$ . On adding strong sulphuric acid to a crystal of cholestearine, letting it gradually run in and act on it, the crystal appears to melt from the edge inwards, and take on a fatty appearance, and by and by it gathers into a brown drop. On adding iodine and sulphuric acid to a crystal, there is at first a beautiful display of colours. **Margarine** occurs in the form of radiating needles such as one sees frequently inside the fat cells in adipose tissue.

Local fatty degeneration is not infrequently followed by **Calcareous infiltration**, where, from deficiency of fluid or otherwise, the fat is not absorbed.

We have already seen that in **Caseation**, fatty degeneration is associated with necrosis.

#### FATTY INFILTRATION.

By this term is meant the infiltration of free fat into the tissues. This condition is only in a restricted sense pathological, especially when it is general. It is necessary to distinguish fatty infiltration of the liver from other forms.

1. **Fatty infiltration in connective tissue.**—Adipose tissue is a form of connective tissue, and is, to a considerable extent, interchangeable with loose connective tissue. Adipose tissue is formed by the infiltration of fat into the connective tissue cells, and at different times it may be variously abundant.

In **Obesity** an excess of fat is present in the body, and it is laid down in store chiefly in the subcutaneous connective tissue and the omentum, but also in other situations where loose connective tissue is present.

**Around or in disused or atrophied organs**, it is common to find a fatty infiltration. A most typical example of this is afforded by **muscles** which have become fixed at their two ends by stiffening of joints. The muscle can no longer produce any movement, and its fibres gradually atrophy as we have already seen. At the same time, in the connective tissue around the muscle and in that which supports it, there is a great infiltration of fat, so that adipose tissue appears between and around the fibres. (See Fig. 35.) Then, again, in pseudo-hypertrophic paralysis—a disease chiefly of children—there is a similar process. The muscular tissue atrophies, but there is at the same time an excessive transformation of the connective tissue into



adipose tissue, so that the wasting of the muscle is more than counterbalanced by the excess of adipose tissue, and there is thus a pseudo-hypertrophy.

A similar fatty infiltration occurs in **the heart** and may seriously incommode it in its action (see further on).

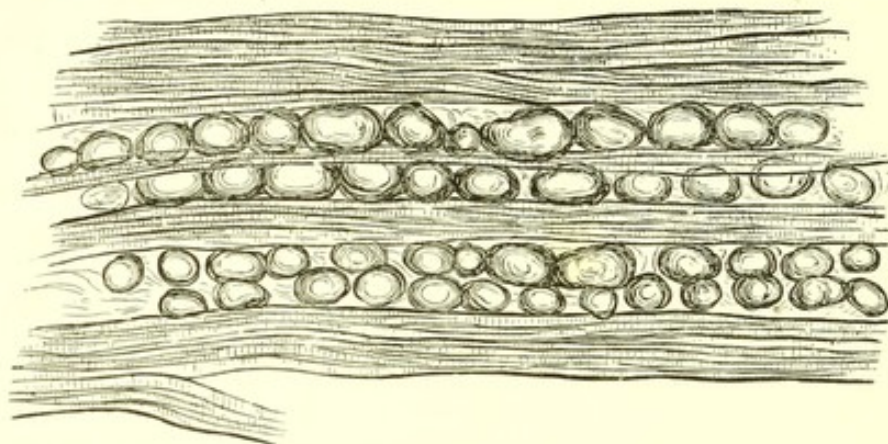


Fig. 35.—Fatty infiltration of muscle from a case of stiffening of the ankle by an epithelioma of the skin over it. The muscular fibres are narrowed, and adipose tissue appears between them.  $\times 80$ .

It is difficult to account for this local deposition of fat in connection with muscle. It may be that as, normally, fat is largely oxidized in muscle, or largely produced there in connection with the active process of contraction, when the demand is gone it is laid down by the vessels in the neighbouring connective tissue.

Fat is often deposited in excessive quantity **around diseased and useless glands**, such as the kidney, pancreas, etc. In the contracted kidney of chronic nephritis there is often an excess of fat at the hilus, which may make the kidney appear much less reduced in bulk than it really is. In hydronephrosis there is often an enormous increase of the fat which normally surrounds the kidney. This is also difficult to account for.

2. **Fatty infiltration of the liver**, which we meet with very frequently, is also difficult to explain. Fat is often found in large quantities in the liver in cases where, in the subcutaneous tissue or elsewhere, there is an actual deficiency of it. The fat in the liver is in the peripheral parts of the lobules, and from this it is to be inferred that it has been brought by the portal blood. This fatty infiltration occurs most frequently in phthisis pulmonalis.

Its accumulation under these circumstances may, in part, be accounted for by supposing that the fat which is normally used for the formation of the fatty acids and the cholestearine of the bile is not so used and is therefore stored in the hepatic cells. It is known that the secretion of bile is greatly diminished in such cases, and that the bile is watery. In that case the fatty infiltration here would be like that in muscle, due to diminished activity of the organ. Another view, and one



having some appearance of probability, is based on the theory that one of the functions of the liver is to prepare fat for oxidation. Naumann (Reichert and Du Bois Reymond's *Archiv*, 1871, p. 41) has shown that the liver fat is much more oxidizable than ordinary fat, and that in the vertebrata the size of the liver is in inverse proportion to the activity of the respiration, being largest in fishes and smallest in birds. It is therefore suggested that in phthisis and cachectic diseases the liver may produce an excess of easily oxidizable fat and store it up ready for use. Hence, perhaps, the utility of liver oils in cases of phthisis, etc.

We have seen that in fatty degeneration the fat appears in the form of fine granules or drops, and that as these increase they remain isolated.

In fatty infiltration there are, of course, first fine fat drops, but as more fat is added the drops grow in size. In the case of the conversion of connective tissue into adipose tissue, there is a single fat drop in each cell, as in Fig. 35. In the case of fatty liver the fat drops are of various sizes (see Fig. 36), but, as a rule, much larger than in fatty degeneration. The size is by no means

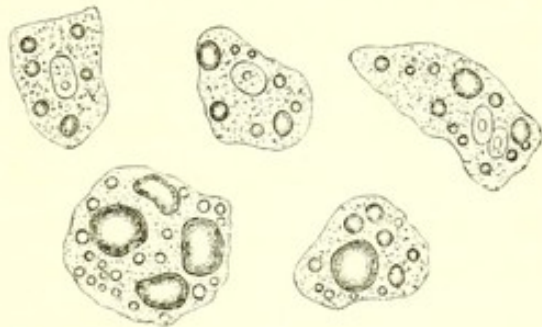


Fig. 36. — Fatty infiltration of the liver. Isolated hepatic cells with drops of fat of various sizes.  $\times 350$ .

an absolute criterion, but it is an important practical indication. It is only in the liver that there can be much difficulty in distinguishing between fatty degeneration and fatty infiltration, and here the fact that in the latter the fat is deposited at first in the cells at the peripheral parts of the lobules, and always continues more abundant there, is sufficiently distinctive.

#### PATHOLOGICAL PIGMENTATION.

The pathological variations in colour in the tissues come under a considerable number of different categories.

The endeavour has been frequently made to refer all pigments found in the body to the blood-pigment, but this relation has been definitely disproved for some (as in the case of pigments introduced from without), and rendered doubtful in others. The normal pigments in the body, those of the blood-corpuscles, of the skin, eyeball, etc., arise in cells and are probably formed by the cells elaborating them from the nutrient material afforded them. It is an assumption to suppose that the pigment of the skin is derived from the blood pigment and not elaborated by the cells. The bile-pigment (*bilirubin*) is very closely related, if not identical chemically, to hæmatoidin, which is the immediate derivative of hæmatin, and is probably produced from the blood-pigment, but the black pigment of the eyeball and rete malpighii (*melanin*) is different, and in particular is devoid of iron.

#### 1. Alterations of physiological pigmentation.—The pigment of the



skin varies much in different persons within physiological limits. There is a pathological absence of pigment (*albinism*), and excess (*negrism*). The pigment also varies in the same person at different times; it is often increased by exposure to the sun, by pregnancy (*chloasma*), etc. A peculiar form of pigmentation is that in **Addison's disease**, in which certain parts of the skin assume a bronzed colour. The excess of pigment here is not only in the rete malpighii but in the papillæ of the cutis and around the veins. This disease is associated with tuberculosis of the suprarenal capsules, but the connection is not very clear.

It is usually supposed that the suprarenal capsules contain nerve centres, and that the bronzing is due to nervous derangements, but some assert that pigment is produced in excess in the capsules and deposited in the skin.

**2. Pigmentation from inflammation and hæmorrhage.**—Many chronic inflammations are associated with pigmentation which may remain long after the inflammation has subsided. This is doubtless due to the escape of red corpuscles by diapedesis, and the pigmentation of the tissue by the products of their pigment. This pigmentation is met with chiefly in the skin and intestine. In passive hyperæmia a similar pigmentation occurs, such as that seen in the lung in brown induration.

Pigmentation as a result of hæmorrhage is of frequent occurrence. When blood is effused in quantity the tissues around usually become stained and may retain their colour to a large extent for years. The colouring matter is hæmatoidin, which is a derivative of hæmatin, and very near to it in elementary constitution, but much more stable. It occurs as granules and crystals, the latter in the form of orange-

coloured rhombic prisms which resemble in shape those of hæmin. (See Figs. 9 and 10, p. 84.)

There are two ways in which the pigment from the effused blood is dealt with. In the first place it may be simply dissolved out of the corpuscles and this solution may afterwards deposit hæmatoidin in granules or crystals. On the other hand the blood-corpuscles may be taken into the substance of amœboid leucocytes and there undergo changes.

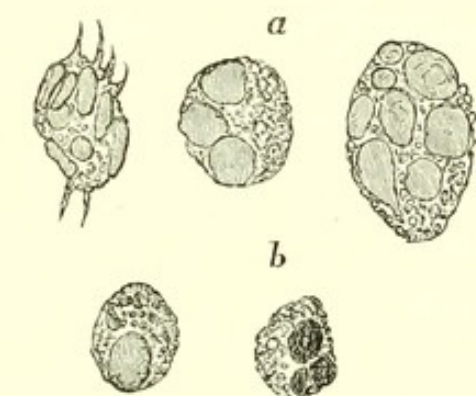


Fig. 37.—Cells containing blood-corpuscles from the neighbourhood of a hæmorrhage: *a*, with fresh corpuscles; *b*, with dark granules from disintegration of red corpuscles.

The blood in the tissues acting as an irritant induces the exudation of leucocytes, which have the power of taking up solid bodies. We have thus cells containing red corpuscles. (See Fig. 37.) These cells



may pass off into the lymphatics, but it appears that they may remain permanently in the part. The red corpuscles inside the cells shrink and become converted into irregular pigment granules.

3. **Melanæmia**.—This name is given to cases in which free pigment is present in the blood. The pigment is derived from that of the red corpuscles, and it implies a great destruction of these. The pigment is in the form of granules and flakes and being deposited in certain organs leads to a coloration of them. (See p. 106.)

4. **Icterus or Jaundice**, is a condition in which the blood contains a yellow pigment which stains the tissues of the body generally. The pigment is usually that of bile, namely, bilirubin. This substance is elaborated by the hepatic cells and secreted as a constituent of the bile. When the exit of the bile is hindered by obstruction of the ducts the pigment is re-absorbed and passes into the blood. Icterus which is thus due to the bile pigment is called **Hepatogenous**, that is, derived from the liver.

We have already seen that hæmatoidin is of similar constitution to bilirubin, and we may presumably have icterus from the formation of this pigment in the blood by destruction of the red corpuscles independently of the bile. This form of icterus is called **Hæmatogenous**. The exact domain of this form is not determined. The icterus of some acute diseases such as pyæmia and typhus fever is hæmatogenous. More doubtful is the **Icterus neonatorum**. This form of jaundice occurs in a large number of new-born children, and is usually regarded as due to a destruction of red corpuscles occurring in consequence of the changes in the circulation consequent on birth.

It is asserted by Birch-Hirschfeld that this icterus is hepatogenous. He says that the sudden change in the circulation at birth causes œdema of the interstitial connective tissue of the liver, and that this produces obstruction of the ducts. Cohnheim agrees with Birch-Hirschfeld that the icterus is probably hepatogenous but does not see sufficient evidence of œdema. He suggests that as there is at birth a sudden increase in the secretion of bile, the ducts may take some time to accommodate themselves. Recklinghausen believes the icterus to be hæmatogenous on the apparently sufficient ground that the fæces are coloured with bile and that therefore the bile ducts are not obstructed.

A peculiar feature in icterus neonatorum is the occurrence of Crystals of hæmatoidin or bilirubin in the kidneys, and also in the tissues and blood. That is to say the pigment not only stains the tissues but is deposited in the crystalline form. It is probable, however, that this crystallization is a post-mortem phenomenon. Hæmatoidin crystals may be found even where there is not enough pigment present to produce jaundice.

5. **Pigmentation in tumours**.—In certain cancers and sarcomas a brown pigment is present in the cells of the tumours, giving to the tissue a brown or black colour. The pigment here has the chemical



characters rather of melanin than of blood-pigments and is presumably elaborated by the cells. These tumours mostly take origin in structures where pigment cells normally exist, as in the choroid of the eye, the superficial layers of the cutis, the pia mater (Virchow), and the conjunctiva of the eye, where it passes into the cornea. Such tumours also originate in pigmented *nævi*, in the rete malpighii and in the cornea (especially in horses). In some of these cases the melanin passes into the urine, where it may deeply colour that secretion (Tennent and Coats).

**6. Pigmentation from without.**—Pigmented substances introduced may lodge in the tissues and even permanently colour them. **Salts of silver**, when long administered, or when taken in excess in one dose, may cause a bluish staining of the skin (*argyria*). The oxide of silver is deposited also in the tissues of internal organs.

The **Dust of the air** which is inhaled, passes to some extent into the substance of the lungs, and gives a dark colour to them. (See under Lung Diseases.) Workers whose trade exposes them to variously coloured dust, present similar pigmentations of the lungs. **Tattooing** is an operation by which granular pigments are introduced and lodge in the skin and lymphatic glands.

**7. Pigmentary Atrophy.**—When coloured tissues atrophy there is usually a concentration of their pigment. Thus, we have a brown atrophy of muscle, especially of the heart, and a deepened coloration of the fat in emaciated persons and in old age.

**Literature.**—VIRCHOW, Virch. Archiv, vols. i., ii., iv., vi., Geschwülste, vol. ii.; ADDISON, Dis. of suprarenal capsules, 1855; LANGANS, Virch. Arch., vol. xlix.; RINDFLEISCH, Path. Histology; KEHRER, Stud. üb. Ikterus neonatorum; BIRCH-HIRSCHFELD, Virch. Arch., vol. lxxxvii.; COHNHEIM, Allg. Path., vol. ii., p. 75; RECKLINGHAUSEN, Allg. Path., p. 437; GUSSENBAUER, Virch. Arch., vol. lxiii.; KUNKEL, Virch. Arch., vol. lxxxi.; RINDFLEISCH and HARRIS, Virch. Arch., vol. ciii.; TENNENT and COATS, Glasg. Med. Jour., xxiv., 1885.

#### AMYLOID DEGENERATION.

This name is applied to a condition in which the constituents of the tissues are converted into a substance whose chemical characters are different to those of any normal principle in the body. The degeneration is also called **waxy** and **lardaceous** from the physical characters of the substance produced. This may be called for convenience **amyloid substance**, and, as the name suggests, it was originally supposed to be allied to starch. It has really no chemical relation to starch, being a nitrogenous substance and a modified form of albumen. It resembles starch, however, in respect that it gives a colour reaction with iodine.



The presence of amyloid substance is determined by its physical characters or by certain colour tests. The earliest known of these latter is the reaction with iodine. The iodine reaction is useful for roughly testing macroscopically at the time of the post-mortem. For this purpose a watery solution, consisting of iodine 10 grains, iodide of potassium 20 grains, and water 4 ounces, is poured on the surface of the structure to be tested. A mahogany-red colour indicates the presence of amyloid matter. The further addition of dilute sulphuric acid sometimes produces a deeper red or a bluish colour.

For microscopic purposes watery solutions of methylviolet or gentianviolet as introduced by Cornil are most suitable. These dyes produce a rose-pink colour with the amyloid substance, while normal tissues are stained blue. In testing by iodine microscopically a solution half the strength of that mentioned above is to be used.

The substance itself has a peculiar bright translucent glancing appearance (see Fig. 38), and, as the structures in which it occurs are enlarged, they are often remarkably prominent under the microscope. It is a very dense heavy material, and after death, at least, is somewhat brittle, but the absence of hæmorrhage in amyloid organs would seem to indicate that it is not so during life.

**Causation.**—Looking to the history of the cases in which amyloid degeneration occurs it is clear that it is to be referred primarily to an alteration in the blood, probably an impoverishment of it in albumen. The disease is not an independent one, but comes on in certain cachectic states due to chronic tuberculosis, syphilis, diseases of bone involving prolonged suppuration, chronic dysentery, etc.

As more unusual causes of amyloid degeneration may be mentioned, leukæmia, Hodgkin's disease (malignant lymphoma), very rarely cancer or sarcoma. It is frequently associated in the kidney with chronic inflammation of that organ, but it is doubtful whether the latter is to be regarded as its cause, for, on the one hand, amyloid disease may lead to nephritis, and, on the other, both conditions may be the result of syphilis.

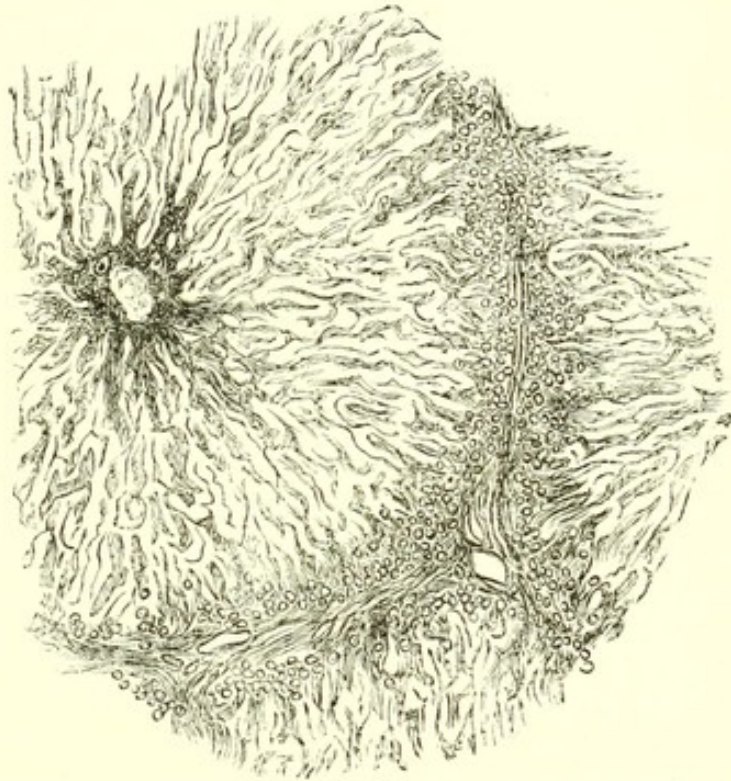


Fig. 38.—Advanced amyloid disease of the liver. The arrangement of the transparent amyloid material suggests its formation in the capillaries. The round bodies at the peripheral parts of lobules are fat drops, there being slight fatty infiltration.  $\times 70$ . (THIERFELDER.)



It is more difficult to determine the nature of the connection between the vice in the blood and the disease in the tissues. By some it is supposed that the amyloid substance arises in the blood by modification of the albumen and is then infiltrated into the structures. But this view cannot be accepted, for various reasons. In the first place, the substance is eminently insoluble, and it is difficult to understand how it can be carried by the blood; besides this it does not displace the normal structures simply, but replaces them, these structures being converted into the amyloid substance. It is more consistent to suppose that the tissues are reduced in vitality by the altered condition of the blood, and that the albumen of the blood enters into combination with the protoplasm in such a way as to produce this peculiar substance. The process may perhaps be compared to the coagulation of the tissues, which, as we have seen, sometimes occurs when they undergo necrosis, the tissues entering apparently into a chemical union with the fibrinogen in the fluid exuded from the blood-vessels, so as to form fibrine or some substance allied to it. Amyloid matter has frequently been compared to fibrine, and Dickinson has suggested its affinity with de-alkalized fibrine. The existence of localized amyloid disease is strongly confirmatory of some such view as this. In this condition abnormal structures enter into this peculiar chemical combination with the albumen of the blood, while normal structures do not. In this connection also, the fact that amyloid disease affects the connective structures of the body is not to be forgotten. It is as if the chemical basis of these structures had a special relation to the amyloid substance. Amyloid disease is therefore essentially a degeneration, although, in order to the formation of the amyloid substance, it is necessary to have, added to the tissue, material from without, and this adds greatly to the bulk and weight of the structures.

**Character of the lesions.**—Amyloid disease may affect any organ of the body, but it is particularly frequent in the spleen, liver, kidneys, intestine, and lymphatic glands.

In all these organs it begins in the **walls of the blood-vessels** more especially the walls of the capillaries and smaller arteries, or in **the connective tissue**. In advanced cases the amyloid substance is in such quantity and the proper tissue of the organs is in many cases so much atrophied that it is often difficult to determine its precise seat. In early cases, however, it will be found that the arteries and capillaries are nearly always the primary seat. It is readily seen in the liver, for instance, that the capillaries are affected, the hepatic cells undergoing atrophy. Even in advanced cases the arrangement is often suggestive



of radiating capillary tubes, as in Fig. 38. In the kidneys, again, it is always the vessels which are first affected, although extension may occur to the basement membrane of the tubules. In the spleen the arteries are mostly affected, and in addition to these either the walls of the sinuses in the pulp or the reticulum of the malpighian bodies. (See under Spleen.)

The distribution of amyloid disease varies greatly in different cases, both in regard to the organs chiefly affected and the parts of the organs. Thus in phthisis pulmonalis it may be chiefly present in liver, spleen, or kidney, and it may be absent in one of these organs while present in the others. Of the two forms of amyloid disease of the spleen one (the sago spleen) is characteristic of phthisis pulmonalis, while the other is probably the form mostly met with in syphilis.

Considerable discussion has occurred as to the existence of amyloid disease in epithelial structures. It is now generally admitted that these are rarely if ever involved, and if they are it is in advanced cases. In the liver it is admitted that the arteries and capillaries are chiefly affected, but Kyber and others have asserted that the hepatic cells are involved. The author has not been able to detect any amyloid change in the hepatic cells.

The amyloid substance is a very inert matter. It is insoluble in water and alcohol and even in gastric juice. During life it renders the structures involved passive, so that they are incapable of vital changes. This material is insoluble in the juices of the body, but consistently with its character as inert matter it gives ready passage to fluids, so that during life the prominent symptom of amyloid disease in the intestine is diarrhoea and in the kidneys an excessive discharge of watery urine.

A frequent result of amyloid disease is diminution in the calibre of the blood-vessels, and this must lead to **anæmia** of the organs. To this may be partly ascribed the **fatty degeneration and atrophy** which so frequently accompany the process, although these are also due to the pressure of the swollen structures.

Structures which have undergone amyloid degeneration are greatly increased in bulk and weight and this tells on the organ as a whole. The liver, spleen, and kidneys are often greatly enlarged, and they present a peculiar dense translucent appearance which has given rise to the names waxy and lardaceous disease often applied to amyloid degeneration.

**Localized amyloid disease.**—This does not occur in tissues previously unaltered, there is always some preceding local lesion. It is met with chiefly in new-formed inflammatory tissue and cicatrices, especially when of syphilitic origin, and also in tumours. It has been seen in syphilitic cicatrices in the liver, tongue, and larynx, in degenerating



cartilage, etc. In some cases the amyloid piece of tissue is of considerable size, and as it differs in its hard translucent character from the tissues around, it may itself look like a tumour.

**Amyloid concretions. Corpora amylacea.**—In old extravasations of blood in the lungs we sometimes meet with round or oval stratified bodies of small size (see Fig. 39 *b*), which somewhat resemble starch granules, and give with iodine the amyloid reaction. Sometimes they contain in their central parts a foreign body, such as a blood crystal. Again in the prostate gland (*a*) we meet with concretions of considerable size, it may

be visible, as brown granules, to the naked eye, with all the characters of stratified amyloid concretions. They are also met with in the tissues of the central nervous system (*c*); they are present in the normal brain, especially in the ependyma of the ventricles, but in cases of sclerosis they may be present in enormous numbers.

The reaction of these bodies is not quite the same as that of ordinary amyloid matter. They give with iodine more of a blue tint, and that without adding sulphuric acid. They sometimes fail to give a red colour with methylviolet. Their significance is not usually very great in a practical point of view, but their presence under these

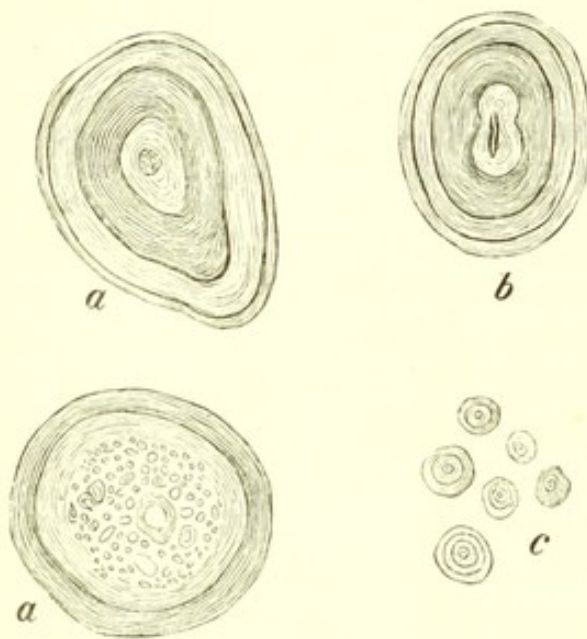


Fig. 39.—Corpora amylacea: *a*, from the prostate; *b*, from a hemorrhagic infarction of the lung; *c*, from the spinal cord.  $\times 400$ . (ZIEGLER.)

various conditions seems to prove that various albuminous substances may undergo conversion into the so-called amyloid substance.

In the nervous system there are frequently developed artificially clear glancing bodies which somewhat resemble amyloid bodies. They occur as a result of the action of alcohol in hardening the tissue, and this agent should therefore, as a general rule, be avoided in preparing the brain and spinal cord for histological investigation. The bodies give with iodine a pale yellow, and with methylviolet a reddish colour. They are devoid of pathological significance, although some writers have referred to them as true morbid lesions. (See a review of the subject by Middleton, who regards them as formed by the action of alcohol on the myeline of the medullated nerve-fibres.)

**Literature.**—VIRCHOW, Virch. Arch., vi.; WILKS, Guy's Hosp. Rep., 1856; KEKULE, Heidelberg Jahrb., 1858; KÜHNE und RUDNEFF, Virch. Arch., vol. xxxiii.; CORNIL, Arch. d. phys., 1875; KYBER, Virch. Arch., vol. lxxxii.; BUDD, Lancet, 1880; Report on lardaceous disease, Path. Soc. trans., vol. xxii.; FAGGE, Path. Soc. trans., vol. xxvii.; DICKINSON, Discussion on lardaceous disease, Path. Soc. trans., vol. xxx.; see also Greenfield, Goodhart and others in this discussion; COATS (Amyloid dis. in



Phthisis), in Gairdner and Coats, Lect. to practitioners, 1888; ZAHN (Local amyloid dis.), Virch. Arch., vol. lxxiii.; MIDDLETON, Glas. Med. Jour. xxii., 1884.

### MUCOUS, COLLOID AND HYALINE DEGENERATIONS.

There are many pathological conditions in which translucent, glancing substances appear in the tissues, and it is frequently difficult or impossible to determine the chemical and other relations of these substances, which were all at one time called colloid substances. In regard to one of them, namely, amyloid substance, the reactions are so definite that it can readily be detected even in small quantities. It has, therefore, been separated from this group. Mucin is also a tolerably definite substance, whose reactions generally allow of its detection. But even in regard to it there are cases in which its presence is doubtful, and there remain many conditions in which the colloid or hyaline appearance is visible, but the nature of the change is obscure.

Most authors use the term colloid degeneration to cover the more indefinite forms, using a term which formerly had a wider significance. Recklinghausen has introduced the term **Hyaline** to indicate a substance having a clear translucent appearance. This author includes under this name both solid and semi-fluid substances having the optical characters mentioned. As the term means glassy, it seems hardly consistent to call by this name tenacious fluids, such as that found in the thyroid gland in some cases of goitre. Perhaps it may be convenient to retain the term colloid for the semifluid matters, and hyaline for the more solid.

1. **Mucous degeneration.**—This is characterized by the presence in the tissues of **Mucin**. This is a normal secretion of certain glands, and is a body of definite chemical reactions. It is closely allied to albumen, but it is precipitated by dilute mineral acids, and by organic acids (acetic acid), and is not re-dissolved by excess of acid. With alcohol it gives a membranous and fibrous coagulum, which is partly re-dissolved in excess of water. Albumen, on the contrary, is not precipitated by organic acids, and its precipitate with alcohol is flocculent, and not re-dissolved by water. The physical characters of mucin are notable in that, in even small amounts, it gives fluids a sticky, tenacious character. Thus a fluid containing 5 per cent. of mucin will be tenacious, while the blood serum which contains 9 per cent. of albumen is quite liquid. **Paralbumin** is closely allied to mucin, if not the same substance.

Mucin is present pathologically either in cells or in the intercellular substance. In cells it has its physiological type in the secretion of mucus. This takes place by a transformation of epithelial cells, which may be either in proper mucous glands or else on the surface of mucous membranes. The cells show in their protoplasm a clear substance which



gradually distends them, and they become goblet cells. The mucin is discharged, the cell being either destroyed or returning to the normal condition. An exaggeration of this process occurs in catarrhs of mucous membranes, but this can scarcely be called mucous degeneration. There may also be an accumulation of mucus in a cavity or cyst, but this also is to be distinguished from degeneration.

A definite mucous degeneration occurs in tumours, notably in ovarian tumours, where the result is rather paralbumin than mucin, and in certain cancers. In the colloid ovarian cystoma, the mucous (or colloid) matter is produced by a process of secretion in glandular structures, goblet cells being characteristically present. (See Fig. 40.) In colloid or

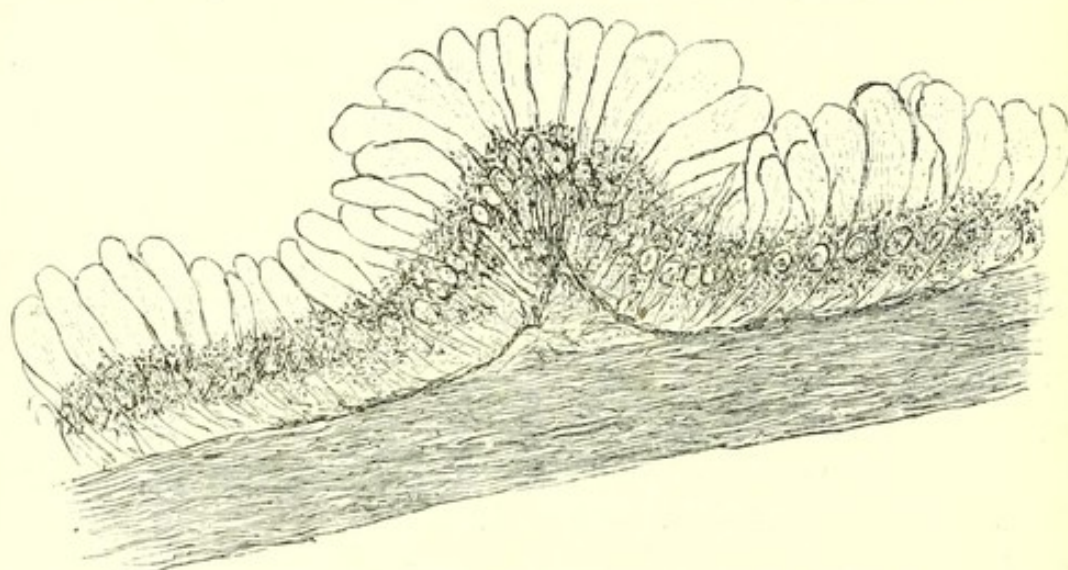


Fig. 40.—From internal surface of a colloid ovarian cyst. The lining epithelium is mostly in the form of goblet cells, the superficial parts having clear transparent contents.  $\times 350$ .

mucous cancers the epithelial cells of the tumours undergo a mucous transformation. The tumours which present this change all have their seat where cylindrical epithelium is a normal constituent.

Mucin differs from albumen in respect that it has no tendency to become absorbed, so that a cavity containing a mucous fluid is not likely to have its contents reduced by absorption.

Mucin is present in the **intercellular substance** in the tissue of the umbilical cord, which may be taken as the type of **Mucous tissue**. In this tissue the intercellular substance is composed of a soft jelly. Tumours occur which are composed of a similar tissue. (See Myxoma). A true mucous degeneration occurs when the dense matrix of cartilage or bone becomes transformed into a jelly containing mucin, or when the adipose tissue becomes like mucous tissue.

2. **Colloid degeneration.**—Under this name are included conditions in which the cells of structures secrete or become converted into a clear homogeneous substance, this transformation implying the destruction of



the cells. It occurs almost as a physiological process in the **Thyroid gland**, as it is always present in later life in that gland. It forms an important element in the commonest form of **Goitre**, in which there is an enlargement of the gland. The thyroid gland consists of saccules lined with epithelium. The epithelial cells become colloid and the saccule comes to be occupied by a translucent clump of colloid matter, which by swelling up causes enlargement of the saccule. In the **Kidneys** we frequently find cysts occupied by colloid matter, which has arisen by transformation of the epithelium of the tubules or glomeruli.

**Myxœdema** is a condition in which mucin is present in the skin, subcutaneous tissue, etc. It is associated with changes in the thyroid gland under which it will be discussed.

3. **Hyaline degeneration**.—Recklinghausen has introduced this term to designate conditions in which a clear homogeneous substance of a vitreous appearance is present, which does not yield the reactions of amyloid substance or mucin. This **Hyaline** has special reactions to some staining agents, being deeply coloured by most of the acid dyes. Thus carmine, picro-carmine, and to a less extent, hæmatoxyline, eosine, and acid fuchsine stain it deeply. It is an inert substance and its presence implies that the structures involved are obsolete if not dead. The substance is insoluble in water and alcohol, and is unaffected by acids and alkalies. It is insoluble in the juices of the tissues; at most, swells up when acted on by them and remains as an inert substance. Like amyloid matter and mucus, it arises chiefly by transformation of cells, but there are some conditions included by Recklinghausen, in which this can hardly be said.

It is not asserted by Recklinghausen that hyaline has a determined chemical constitution, and there are probably many different conditions included in the name. Some of these have already been referred to. (1) Colloid degeneration is included in hyaline degeneration; (2) Some instances of coagulation-necrosis are included, and more particularly the waxy degeneration of muscle; (3) Tube casts in the kidneys (hyaline cylinders) and similar structures met with in inflammations in the ducts of the sweat glands, and in the ovarian follicles; (4) In many tumours, as in lymphomas, sarcomas, and cancers, as well as in tubercles; (5) On mucous membranes, forming the main constituent of diphtheritic membranes; (6) In thrombi and fibrinous exudations, where the fibrine, at first forming a net-work, becomes converted into a hyaline homogeneous material; the thrombi in aneurysms often assume this character; (7) In the eye as prominences in the hyaloid membrane.

**Literature**.—RECKLINGHAUSEN, *Allg. Path.*, p. 404.

#### CALCAREOUS INFILTRATION AND CONCRETIONS.

By these terms is meant the pathological deposition of lime-salts. In **Ossification** we have the salts of lime united with an organic matrix, and the tissue has a definite structure in which living, active cells are present. In **Calcareous infiltration** the same salts, chiefly the carbonate and phosphate of lime are deposited in tissues without entering into any proper union with them, and, in fact, the deposition of the lime is in



itself evidence that the tissue has virtually lost its vitality. A **Concretion** is a solid body, formed generally by deposition from a fluid. Such solid bodies consist in many cases of lime-salts.

**Causation.**—Lime-salts or other soluble matters may be deposited because they are present in excess. In cases of rapid destruction of bone, as by an advancing cancer, the absorbed lime-salts are present in the blood in excess, and we may have a **metastatic calcification** of the lung or intestine. There may be thus an incrustation such as to make the tissue like pumice-stone. In some cases of this kind there has been a co-existent disease of the kidneys hindering the due excretion of the lime-salts.

In most instances, however, the presence of **dead or obsolete matter** or a foreign body is the chief determining cause of deposition. The blood and principal fluids contain lime-salts in solution, and, under certain circumstances, these are liable to precipitation. In the living

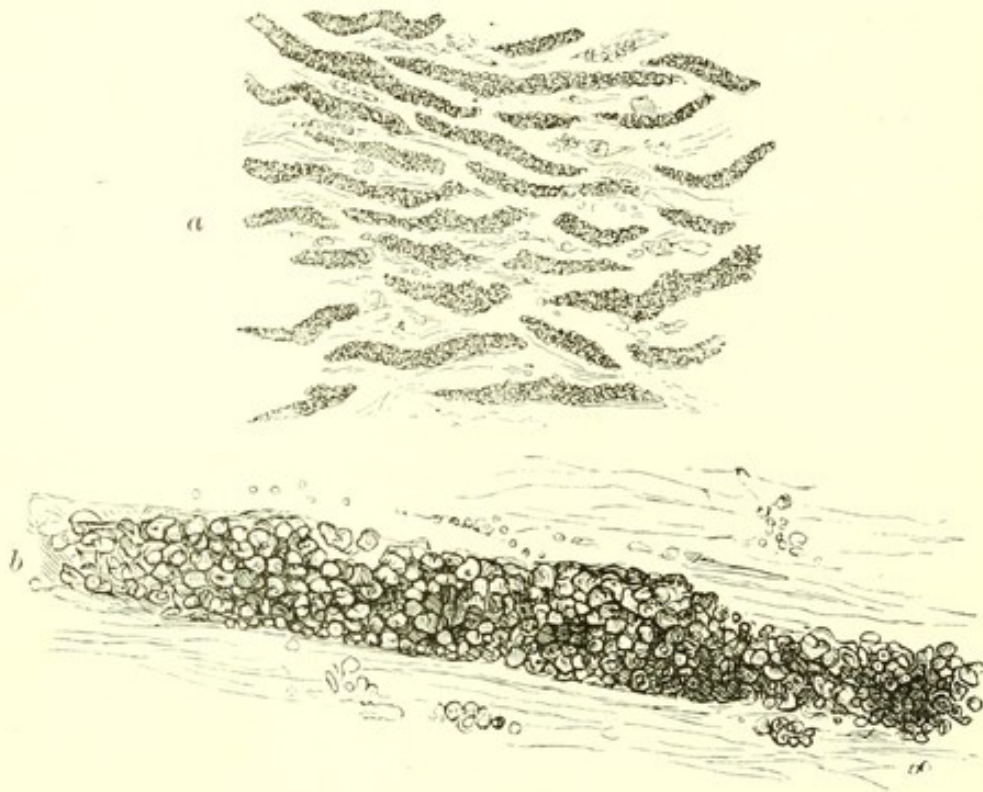


Fig. 41.—Calcareous infiltration in a tumour; *a*, cells of smooth muscle filled with lime granules; *b*, a blood-vessel converted into a solid rod.  $\times 350$ .

tissues there is a continual circulation of the fluids, and these latter do not linger long enough to undergo any serious chemical change. In dead or obsolete structures, on the other hand, the juices will lie stagnant, and are liable to undergo chemical changes. Both in the case of concretions and infiltrations there is usually a foreign body or piece of dead matter as the centre of deposition. In addition, the circumstances are frequently such as to cause stagnation of the fluid.



**Characters of the lesions.**—The lime-salts are deposited in the first instance in the form of fine globular granules, either in the protoplasm of cells or in the intercellular substance. The structure is as if dusted with refracting granules, and the appearances in many respects resemble those of fatty degeneration. (Fig. 41 *a*.) As the salts accumulate, the appearance of granules is somewhat lost and a more continuous petrification results. (Fig. 42 *b*.) Sometimes the structure becomes in con-

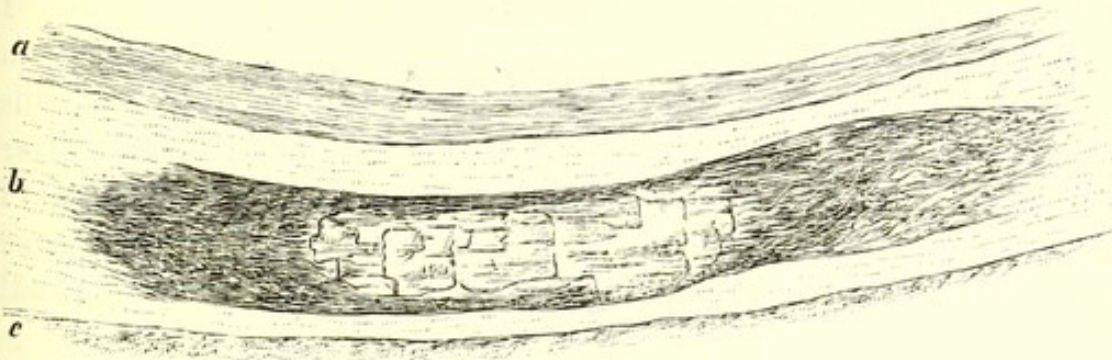


Fig. 42.—Calcareous infiltration of the middle coat in an artery. The lime salts have aggregated together so as to produce a crystalline appearance.  $\times 22$ .

sequence homogeneous and somewhat transparent as in Fig. 42. The addition of a dilute mineral acid causes the salts to dissolve, and, as carbonates are nearly always present, solution occurs with evolution of gas.

Examples of this process are very numerous. A minute parasite, the *trichina spiralis*, occurs in the embryo form in the muscle of man and

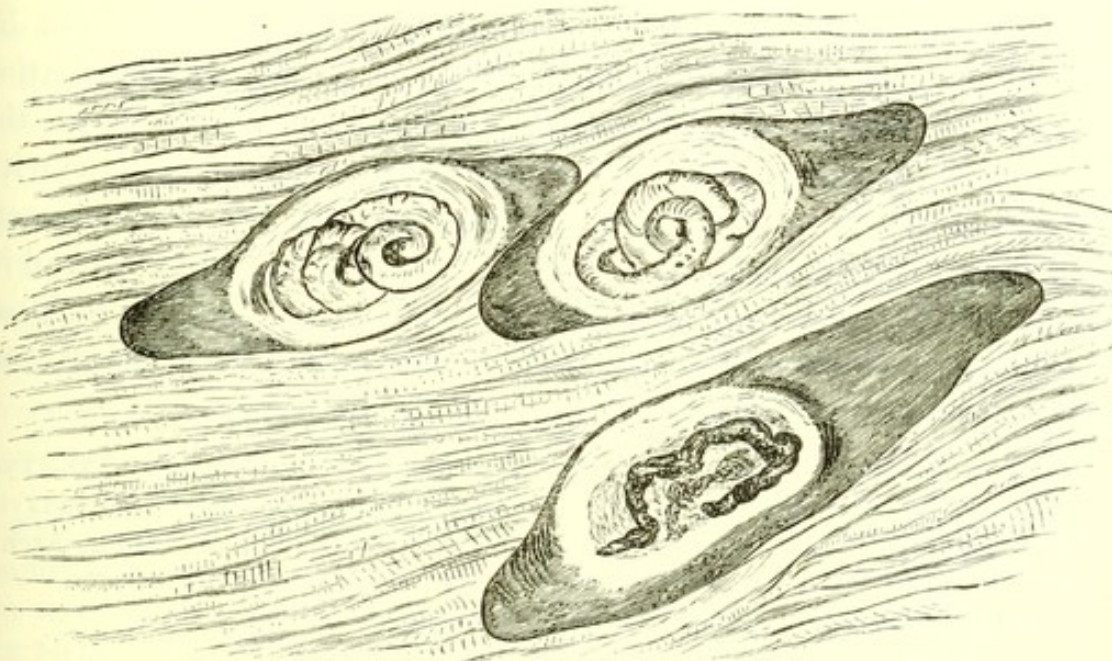


Fig. 43.—*Trichina spiralis* in muscle. The capsules infiltrated with lime, and in one case a dead worm shrivelled and impregnated with lime.

animals; it lies there quiescent, coiled up spirally and surrounded by a capsule. It is virtually a foreign body, and the capsule is by degrees



impregnated with lime, assuming an opaque appearance at its poles (Fig. 43). If the embryo itself dies, it also may become impregnated with lime (see Fig. 43, lower part). Sometimes an extra-uterine foetus dies and remains inside the abdomen as a foreign body. It becomes surrounded by adhesions and partially encapsuled. The capsule and superficial parts of the foetus become through time encrusted with lime, forming the so-called **Lithopædion**. Again, an inflammatory exudation in the pericardium may dry in and become impregnated with lime. In phthisis pulmonalis, if healing occurs, the contents of cavities and inflammatory products may dry in and become surrounded by a capsule; impregnation with lime results, leading to a pultaceous or mortary material, which may ultimately condense into a stony mass. Again, in valvular disease of the heart, due to chronic endocarditis, the new-formed connective tissue, by its contraction, becomes hard and dry and virtually obsolete, and deposition of lime-salts occurs.

These are examples of calcification of foreign bodies or pathological products, but we may have the process occurring **in the ordinary tissues**, when they have become obsolete. In the middle coat of arteries of old people, calcareous infiltration, beginning in the muscular fibre-cells of the middle coat, frequently leads to massive petrification of the arteries (see Fig. 42), so that they form rigid tubes. The cartilages of old people are also liable to impregnation with lime. The crystalline lens of the eye may undergo a similar deposition in certain forms of cataract.

In some cases the calcification is followed by a **true ossification**. Thus calcareous infiltration of the middle coat of arteries is not infrequently associated with ossification (Paul, Coats), and the calcification of the ribs of old people frequently passes into ossification. Again the author found in an old hydatid cyst in the liver true bone associated with calcification. In these cases the rigid calcified structure probably acts as a foreign body, inducing the formation of granulation tissue around it. The granulation tissue may eat into the calcareous mass, and it looks as if the presence of lime-salts induced it to develop into bone rather than into ordinary connective tissue.

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## SECTION VII.

## NEWFORMATION.

HYPERTROPHY, REPAIR, REGENERATION, TRANS-  
PLANTATION.

NEWFORMATION OF TISSUE *by Karyomitosis.* *Genesis of newformations, from indifferent cells; Metaplasia.* HYPERTROPHY, *compared with normal growth; (1) Hypertrophy from congenital proclivity; (2) compensatory; (3) from increased blood-supply; (4) from direct stimulation; (5) from friction.* REPAIR OF INJURIES, *limited powers of restoration in man; healing of wounds.* REGENERATION OF TISSUE, *chiefly blood, epithelium, and connective tissues, also nerve and muscle.* TRANSPLANTATION, *effected by experiment in animals, spontaneously and by operation in man.*

## I.—NEWFORMATION OF TISSUE.

THE pathological newformation of tissue occurs by processes analogous to those concerned in the physiological formation of tissue in the process of growth.

**Karyomitosis** (*καρῖον* = nucleus, *μίτος* = a thread or fibre).—The newformation of tissue, whether physiological or pathological, implies cell-division. According to the views of Remak this process consisted in a direct division of the nucleus and cell. The recent observations of Flemming and Strassburger show that in the growth of both animal and vegetable tissues the process is not so direct, but involves certain changes in the nucleus of a striking and peculiar nature. To this process the names **Indirect division**, **Karyokinesis**, and **Karyomitosis** have been applied.

This process occupies a comparatively short time, seconds or minutes, and is not readily seen in the products of post-mortem examinations. In order to observe it the tissues should be obtained from the living body and immediately subjected to the proper hardening processes, preparatory to microscopic examination.

The **Nucleus** in a state of rest is not a homogeneous body. It has a limiting membrane, inside which the contents are composed of two substances. One of these is deeply stained by certain re-agents and is



hence called **Chromatic substance**, while the other is less stained and is called **Achromatic**. The chromatic substance forms a finely **fibrous stroma** (see *a*, Fig. 44), between which the achromatic substance and

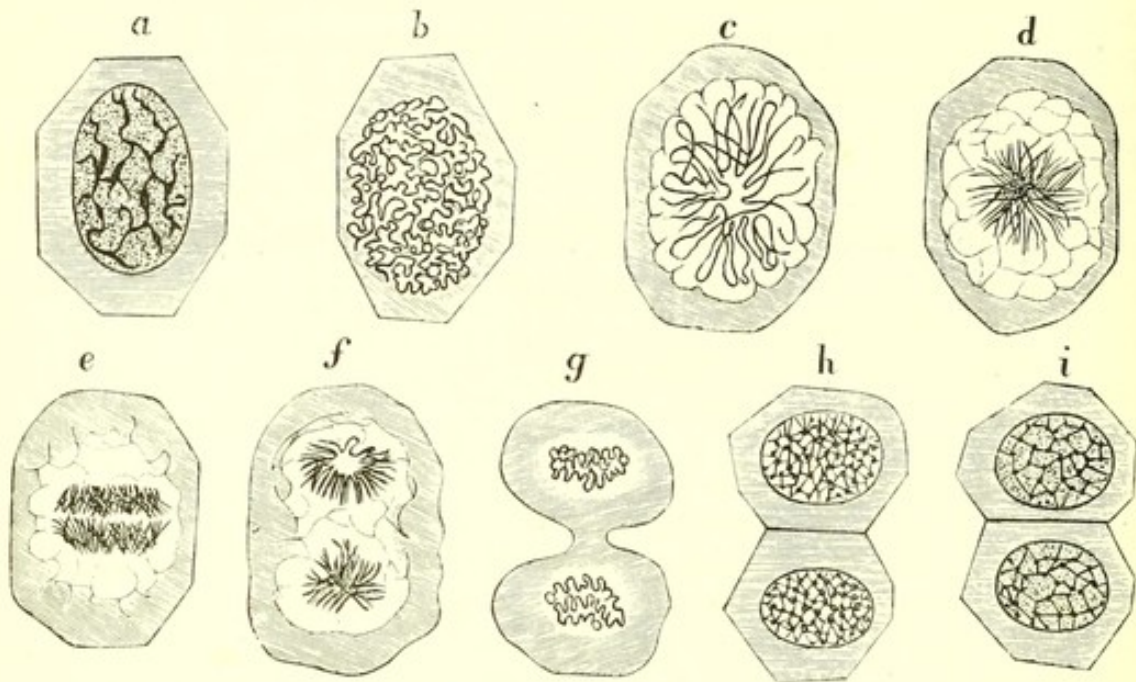


Fig. 44.—Division of cells. Explanation in text. (After FLEMMING.)

the nucleoli lie. It is the fibrous stroma which is mainly concerned in the processes which precede division. The membrane and nucleoli disappear and the fibres become thicker and stain more deeply than before (see *b*, Fig. 44), being converted into a convoluted fibre, or system of fibres. These fibres seem to have a power of movement according to which they alter their arrangement as the process proceeds (hence the name karyokinesis, from *καρύον* = nucleus, and *κίνησις* = movement). From this convoluted form develops the stellate form of the fibres (*c*), the fibres by longitudinal division becoming finer (*d*). The fibres now arrange themselves in the equator of the nucleus (at *d* and *e*), and here they divide into two. The two halves now diverge from one another towards the poles of the cell, forming there the fibres of the new nuclei, and having at first a stellate arrangement (*f*). This by degrees gives place to a convoluted arrangement and the nucleus subsides into the state of rest, the fibres to a great extent disappearing, and the cell-membrane being formed. After the polar separation of the daughter nuclei the cell itself shows signs of division, and so the process is completed.

The process of karyomitosis is to be observed in all kinds of pathological newformation. It is abundantly evident in growing tumours, in tubercles (according to Baumgarten), in inflammatory newformations,



and it is probably the chief, if not the only method, of newformation of permanent cells. In fact, the activity of newformation may in general be estimated by the number of cells in which karyomitosis is visible.

According to Arnold the process is not so uniform as that described above, but follows two types which he designates segmentation and fragmentation, but he suggests that the two may be simply modifications of the same type. There is probably, in addition to this, a direct division of cells, which occurs chiefly in leucocytes.

**Genesis of newformations.**—An important question in regard to newformations is, as to their origin and their relation to the existing tissues. It will be found that the new-formed tissue always conforms in the details of its structure to one or other of the normal tissues. No newformation is therefore foreign to the organism, there is no heterologous or heteroplastic newformation in this sense. For the most part, also, the new-formed tissue obviously springs from and is in close relation with tissue of its own kind. It sometimes happens, however, in the case of tumours that their tissue extends to structures of a different nature from their own, and in this sense the term heterologous is sometimes used.

In its earlier periods, after the cleavage of the germinal vesicle, the embryo consists of a mass of round cells which, from the fact that they present no visible differentiation of structure, may be called **Indifferent cells**. These cells form the three germinal layers, the epiblast, hypoblast, and mesoblast. When once this differentiation has occurred these layers remain distinct, and each produces its own special tissues. The same distinction is presumably carried out in pathological newformations. It was, indeed, suggested by Virchow that the connective tissue which exists in every part of the body, penetrating amongst other tissues, may be regarded as the remains of the undifferentiated embryonic tissue. In this sense he regarded connective tissue as the essential agent in newformation of tissue, and as capable of producing epithelial tissues as well as those of its own kind. In more recent times there was a tendency to ascribe to the leucocytes which emigrate from the blood-vessels a somewhat similar position, and to ascribe at least some newformations to the development of these cells. The general belief now is that the differentiation of the germinal layers holds for pathological as well as for physiological newformation, and that tissues do not originate except from tissues of their own nature. The greatest difficulty in this respect is in regard to cancers, under which heading the subject will recur.

**Metaplasia.**—A certain interchangeability is manifested among the different tissues belonging to the connective tissue series. These



really form a single tissue having certain modifications, so that fibrous tissue, bone, cartilage, mucous tissue, adipose tissue, may not only be developed to a certain extent from each other, but may even be converted, when mature, into each other. Here we have a true metaplasia. Thus, adipose tissue, by absorption of the fat, becomes loose connective tissue, connective tissue by attraction of lime salts becomes osseous tissue, cartilage also develops into bone, or, by a different change in its matrix, forms mucous tissue.

**Classification of newformations.**—The newformation of tissue occurs under three different circumstances. It may be virtually a continuation of normal growth, the new tissue being produced to subserve the normal functions of the body. Secondly, it may be produced in consequence of the application of an irritant which directly stimulates the tissues. Inflammatory newformation is an example of this, and we have a further example in the specific or infective newformations. Lastly, there is a group in which no cause is apparent for the newformation, the tissue simply grows, without any apparent stimulus and without any purpose in the economy. This comprises the group of tumours proper.

In the present section the first of these groups will be considered.

**Literature.**—*Karyomitosis*—FLEMMING, Virch. Arch., vol. lxxvii., and Zellsubstanz, Kern- und Zelltheilung, 1882; STRASSBURGER, Zellbildung und Zelltheilung, 1876, Kerntheilung, 1884; EBERTH, Virch. Arch., vol. lxxvii.; ARNOLD, Virch. Arch., xciii., xcv., xcvi.; BAUMGARTEN, Tuberkel und Tuberkulose, 1885; MARTIN, Virch. Arch., lxxxvi.; KLEIN, Quarterly Jour. of Micr. Science, xvii. and xix.; BIZZAZZO, Virch. Arch., cx. (here a very good method of staining which the author has found very efficient).

## II.—HYPERTROPHY.

The term hypertrophy means overgrowth or excessive growth. Looking to the cellular constituents of the tissues, Virchow has drawn a distinction between an increase of tissue due to an enlargement of the cells, and that due to a numerical increase, applying to the latter the term **Hyperplasia**. This distinction, however, cannot be carried out, as in many cases both substantial and numerical increase may be present. The term hyperplasia may, however, be used where it is intended to convey the meaning that cell-division or proliferation is present.

**Normal growth**, as seen in the tissues during the period of adolescence, is determined by impulses inherent in the impregnated embryo. Already, at this early period, the sex, and details of structure are implicitly inherent in the embryo, which enters on its career of development with a pre-determined plan. The tissues cease to grow at the period of maturity not because their powers of newformation have



been exhausted, but because this plan has been fulfilled. Hypertrophy may occur because of some error in the embryonic arrangements, or it may be due to some stimulus acting on the tissues after birth. In the latter case the effect will be greater when the stimulus is applied during the period of normal growth than after the state of maturity has been reached. In all cases the new-formed tissue is in structure and function essentially similar to the normal tissue of its kind and forms an addition to the existing active tissue of that kind.

1. **Hypertrophy from congenital proclivity.**—There may be an excessive growth of the whole body, so that the person becomes a **giant**; or there may be a localized hypertrophy, as of the fingers, one side of the face, etc. There are also cases in which at birth there is a hypertrophy of the tongue, penis, neck, or a lower extremity.

2. **Compensatory hypertrophy.**—This form of hypertrophy implies that, as a result of some defect in the organism, some function has been called into unusual exercise. As a result of the continued excessive exercise, the tissue is increased. The necessity for this increased exercise may arise in one of two ways; there may be from atrophy or destruction, an actual loss of tissue, and the remaining tissue enlarges to bring it up to the normal amount; or the circumstances may be such as to call for the exercise of a particular function in excess of what is usual, so that, in relation to the increased need, the existing tissue is defective in amount. In this case the new-formed tissue constitutes by so much an absolute excess over the average normal.

The most striking instance of compensatory **hypertrophy from loss of tissue** is that afforded by the enlargement of one kidney as a result of destruction or disease of the other. The hypertrophied kidney will sometimes attain to the bulk of the two normal ones, especially if the lesion has occurred in a young person. A similar hypertrophy occurs in the lung in cases of congenital non-inflation of one lung; also in the testicle, where one is wanting or defective in its development, and in the liver, where, from destruction of a large portion of the right lobe, the left may attain to the size which the right normally presents. (See under the affections of these various organs.)

It is necessary to distinguish **Pseudo-hypertrophy** from the proper compensatory hypertrophy illustrated above. Atrophy of a tissue may be accompanied by an excessive growth of tissue different from that which has been lost, and the new tissue may not only make up the normal bulk but exceed it, so as to give an appearance of hypertrophy. Atrophy of muscle is frequently accompanied by development of adipose tissue between the ultimate fibres, and in pseudo-hypertrophic paralysis a deceptive appearance of hypertrophy of the muscles is produced. A similar excessive production of adipose tissue sometimes occurs around effete and disused glands, such as the mamma (see p. 159).



The other form of compensatory hypertrophy, that characterized by **absolute excess of tissue**, is exhibited chiefly in muscular organs. In the case of canals with muscular walls, constrictions of the canals or orifices and defect in the valves frequently occur, and these may necessitate increased muscular effort to compensate for the defect. Thus the walls of the heart frequently hypertrophy from disease at the orifices, in the valves, or in the general vascular system. The urinary bladder shows hypertrophy of its muscular coat in consequence of obstruction at its neck (enlarged prostate) or in the urethra. The muscular coat of the stomach frequently hypertrophies from obstruction of the pylorus, and that of the intestine from obstruction of its calibre. (See under the organs named.)

3. **Hypertrophy from increased blood-supply.**—In ordinary growth

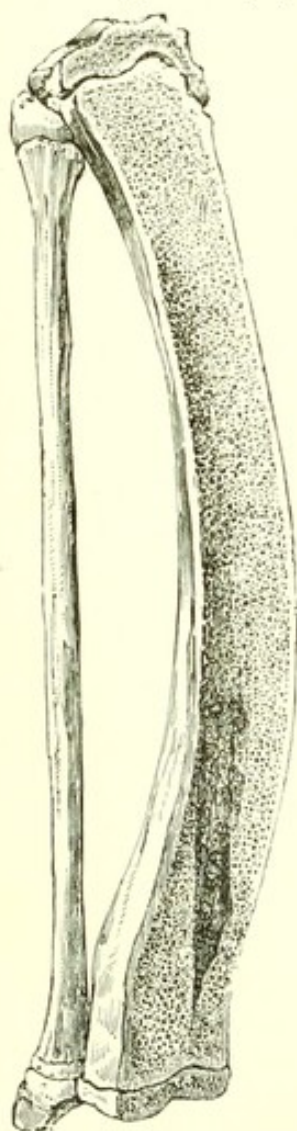


Fig. 45.—Elongation and curvature of tibia, the result of necrosis during period of growth. (PAGET.)

of tissue, whether normal or pathological, the blood-vessels strictly follow the growing tissue, and are formed according to its needs. But if, from some accidental circumstance or by artificial interference, the blood-supply be greatly increased, then excessive growth may result. If the spur of the cock be removed from the leg and successfully transplanted into the comb, it will grow with excessive vigour, forming a prominent horn-like structure. Here the excessive supply of blood, the comb having a much more active circulation than the leg, induces an excessive growth of the epidermis forming the spur.

In human pathology we have numerous instances of hypertrophy analogous to this. Increased activity of the circulation will often lead to hypertrophies. We see it **in the neighbourhood of inflammations**. Determination of blood exists outside the immediately inflamed area, and if this is prolonged it may lead to overgrowth of the tissues. In this way we may account for the excessive growth of hair sometimes seen in the neighbourhood of ulcers, near diseased joints, and at the ends of stumps which have remained long inflamed.

A very striking instance of hypertrophy of this kind is sometimes afforded **in bones**. In the neighbourhood of inflamed joints the surface of the bones is often nodulated, and the bones



greatly thickened by newformation under the periosteum. Again, if a boy has a necrosis of the femur, the whole bone may be more richly supplied with blood, and the normal growth accelerated. The necrosis may be recovered from, and the person be left with a permanently elongated femur which may be as much as two inches longer than the other, and this may lead to considerable lameness. The tibia is differently situated to the femur. Its two extremities are tied to the ends of the fibula by firm ligaments, and so the bone cannot freely elongate. If overgrowth occurs the bone must curve so as to accommodate itself. An example of this is shown in the accompanying Fig. 45, from a preparation in St. Bartholomew's Museum (quoted by Paget), in which the bone, measured over its curve, was two inches longer than the healthy one.

4. **Hypertrophy from direct stimulation.**—From the observations of Gies it appears that small doses of arsenic administered to growing rabbits cause increased growth in the bones. When administered to pregnant rabbits the young were born with both soft parts and bones larger than those of the young of rabbits of similar size. Wagner found that small doses of phosphorus given to young animals caused the epiphyseal layer of cartilage to produce dense bone instead of spongy. The periosteum also produced bone in excess, so that increased thickness resulted. In these instances the arsenic and phosphorus seem to act as direct stimulants to the bone and other tissues.

5. **Hypertrophy from pressure or friction.**—Thickenings of the epidermis occur in places where the skin is exposed to unusual pressure or friction. Continued pressure, as by a splint or bandage, causes atrophy, but intermittent pressure, by allowing the parts to recover, and by affording time for increased nutrition, gives rise to hypertrophy. We have thus the **horny hands** of workmen, and **corns** which consist of concentric thickenings of the epidermis. The same law applies to internal parts, but as pressure from within, produced by tumours, aneurysms, etc., is usually constant, atrophy is much more frequently the result. Hence the original statement of John Hunter is justified, that pressure from without produces thickening, while that from within causes atrophy, although it is not to be taken without reservation.

**Literature.**—HUNTER, Palmer's ed., vol. i. pp. 421 and 560; VIRCHOW, Cellular Path.; PAGET, Lect. on Surg. Path.; STANLEY, Diseases of the bones, 1839; COATS, Compensatory hypert., Proc. Lond. Med. Soc., vol. vii.; GIES, Arch. f. exp. Path. u. Pharm., vol. viii.; WEGNER, Virch. Arch., lv.



## III.—REPAIR OF INJURIES. REGENERATION. TRANSPLANTATION.

The term **Regeneration** is applied to the restoration of portions of the body which have been lost by injury or disease. The regeneration of a part is to be carefully distinguished from mere growth or hypertrophy. A tissue may be able under suitable stimulation to reproduce its elements, and increase in size; but for the replacement of a lost part, if at all considerable, there must be, virtually, a renewal of the process of development.

The **reproduction of lost parts** in their entirety occurs readily in some of the **lowest forms of animals**. In the hydra, if the creature be cut in two, each half will develop into a complete animal, and the process may be repeated indefinitely. This power of reproduction of the whole animal from a part seems confined to those creatures which can propagate by spontaneous fission or gemmation. When we come to animals higher in the scale the power of reproduction seems to be limited to the restoration of lost limbs, antennæ, etc.

Without going into details, which will be found by reference to Paget's "Lectures on Surgical Pathology," it may be said that there are indications which seem to show that there is some kind of **law** according to which the reparative power in each perfect species is in inverse proportion to the amount of change which the animal has passed through in its development from the embryonic to the perfect state. It is as if, in the process of development, the formative power as distinguished from mere growth were gradually exhausted, and the process of reproduction, which, we have seen to be, as it were, a renewal of that of development, only occurs when this power has been comparatively little expended. It appears, for instance, that in insects the power of reproducing antennæ or limbs is limited to those species which have attained the perfect state through a comparatively simple and direct course of development. It is consistent with this view that in the larval state insects show a much greater power of reproduction than when perfect. The larva of one of the higher insects will be able to reproduce its limbs, while the perfect insect is not.

In man, and in the vertebrata in general, the long course of development seems largely to exhaust the reproductive power of the body, and, in the adult state at least, the power of restoration of lost parts is very limited, and the processes concerned are almost as much related to growth of tissue as to development. In the embryonic state it is probable that the power of restoring lost parts is much greater than in the adult. Some children are born with a short arm at the extremity of which are imperfectly developed fingers; it is probable that in these cases amputation of the arm has occurred *in utero*, and an attempt at restoration has followed.

In the adult it may be said that restoration of lost structures is almost confined to the epithelial and connective tissues and to the blood. Along with the connective tissues we have, of course, blood-vessels which are readily reproduced, and we may also, to a limited extent,



include nerve fibres, which, as we shall see afterwards, are sometimes restored.

1. **Repair of injuries.**—While the absolute restoration of complete and considerable portions of the body is scarcely possible in man, yet the body is by no means unprovided with powers by which injuries are repaired and loss of structures is made good. It may be said that the higher and more complex animals are endowed with greater ability to protect themselves from injury, and that their tissues possess the necessary powers of restoration in the case of those injuries to which they are specially liable. In the various processes here to be considered, it will be seen that what may be called the definite intention to attain a certain result which is shown in the process of development, is distinctly visible, and that the tissues have a remarkable power of meeting adverse conditions.

**Healing of wounds.**—The power of repair is well seen in the various processes concerned in the healing of wounds. There are some wounds which heal by a process fitly designated **Immediate union**. That is to say, the surfaces are brought together and coalesce without any new-formed material being produced to serve as a bond of union. This occurs mostly in clean-cut wounds, which, shortly after their infliction, are closed so as to bring the cut surfaces into close contact. It is necessary for this process that all inflammation be avoided either at the time of infliction of the wound or afterwards. We are to think of the living tissues not as mere mechanical pieces of texture, but as possessed, by virtue of their vitality, of wonderful powers of adaptability to circumstances. When two living surfaces are brought into contact and all disturbing conditions are averted, then the blood-vessels will form communications, the nerves will by and by unite and become continuous, and the connective tissue coalesce. In this way it frequently happens that a wound in the skin, or even in muscle, unites, and no trace of a bond of union or even of the line of union can be found after a few days. The epidermis probably does not unite so directly, and the wound is covered by new-formed epidermis.

**Primary adhesion** is a process of a more complicated kind. In it inflammation plays a part. When a wound is inflicted the mere mechanical injury, or exposure afterwards, frequently leads to a trivial but acute inflammation, resulting, as we have seen, in the coating of the cut surface with a fibrinous exudation, the so-called glaze. If two surfaces thus coated with fibrine be brought in contact they unite, the fibrine acting as a glue or cement. But the fibrine does not form a permanent bond of union, and if union is not effected by other methods, then the wound will subsequently gape by the breaking down of the



fibrine. In order to effect a permanent union we must have formative cells produced, such as are concerned in the formation of connective tissue out of granulation tissue, and we must have a newformation of blood-vessels. The uniting tissue is very trivial in amount; there is merely a limited production of formative cells which replace the thin layer of fibrine, and a budding of the blood-vessels till communications are formed between the two surfaces. This whole process may occupy only a day or two, and the permanent new-formed tissue forming the cicatrix is usually very small in amount.

The term **Union by the first intention** is commonly used so as to include both of the conditions described above, any case of union occurring without separation of the surfaces from the first application being so designated. (See more fully in Paget's "Surgical Pathology.")

**Union by the second intention** or **by granulation** is a name given to the closure of a wound by the adhesion and coalescence of two layers of granulations. We have already seen how this comes about. The granulations ultimately develop into connective tissue which forms the **cicatrix** or permanent bond of union. In this case the cicatrix will be a much more considerable one than in union by the first intention.

2. **Regeneration.**—As we have seen, this process is somewhat limited in its range in human pathology. **The blood** is gradually regenerated when in consequence of hæmorrhage its bulk is reduced. The fluid portion is rapidly restored, the white corpuscles are also soon replaced, but the red corpuscles somewhat more slowly. (See Anæmia.)

**The epithelial structures** of the body are to a large extent continually undergoing a physiological process of loss and regeneration. The hairs of men and animals fall out at intervals and are restored; the feathers of birds undergo a similar process; the nails and horny layer of the epidermis are continually lost and replaced by newformation. The plucking out of hairs or feathers, or the removal of nails is followed by their restoration, so long as the papillæ are not destroyed. It is an interesting fact that when the whole distal phalanx of the finger is removed, or even the two terminal phalanges, there may be a partial restoration of the nail in the remaining terminal phalanx. There is also a case recorded in which a boy, apparently affected with ichthyosis, regularly shed his nails. (See references in Recklinghausen.)

On the general surface of the skin and mucous membranes there is normally a continuous shedding of the surface epithelium, and a newformation in the deeper layers to replace that which is lost, a kind of physiological regeneration therefore. When, by accident or otherwise, a superficial portion of epithelium is shed before its time, it will be



replaced by the normal growth of the deeper layers, probably accelerated by the requirements of the body. When the whole thickness of the epithelium is destroyed the gap is by degrees filled by the proliferation of the epithelium at the edges of the wound, as we have already seen in the case of the cicatrization of a granulating wound. According to the observations of Klebs the new-formed epithelium acquires a slight power of amœboid movement, so that it can proceed to the spot which it is to occupy.

The proliferation of the epithelium proceeds by the process of karyomitosis. In the accompanying illustration (Fig. 46, Eberth) the

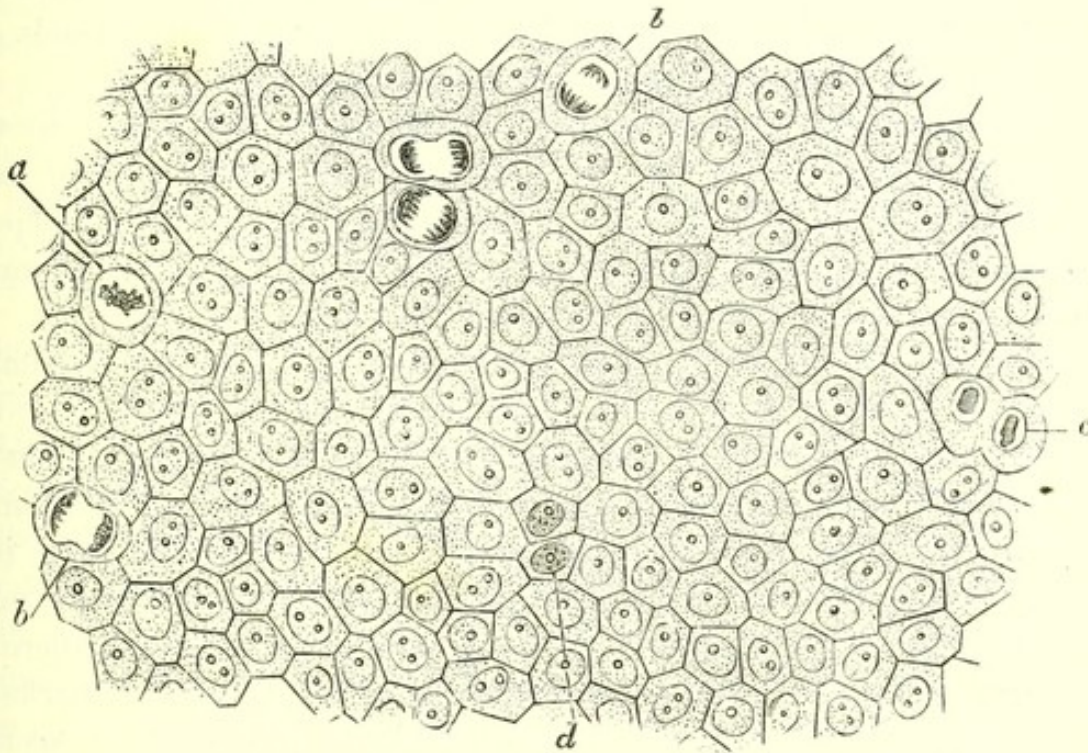


Fig. 46.—Regeneration of epithelium in cornea of a rabbit. (a) Fibrous transformation of nucleus. (b) Partial separation of the fibres and hour-glass change of nucleus. (c) Complete division of nucleus. (d) Complete division of cell. (EBERTH.)

fibrous transformation of the nucleus and the other changes, as seen in the cornea of the rabbit, some days after a portion of the epithelium had been removed, are shown. In the normal cornea and in the malpighian layer of the epidermis there are evidences of a similar process by which, we may presume, the physiological regeneration occurs.

**Gland epithelium** also to some extent undergoes a physiological loss and regeneration. The secretion of some glands implies a destruction of cells and their restoration. According to Bizzozero, who examined the various glands in respect to the activity of karyomitosis, the sebaceous glands, the mucous glands of the stomach, Lieberkühn's glands in the intestine, and the uterine glands, are actively engaged in regenerating their epithelium and consequently lose it in the process of



secretion. In some diseases there is a great loss of epithelium, and a restoration by karyomitosis. Thus in acute nephritis the epithelium is to a large extent shed and we may find desquamated epithelium lying in the tubules, while young epithelium lines them. In acute yellow atrophy of the liver also, there is great destruction of the hepatic cells, and there is often visible along with that a newformation of cells as if there were an attempt at restoration.

**The lens of the eye**, which in its development is an epithelial structure, may be in part or in whole regenerated, after its removal on account of cataract.

**Connective tissue**, as we have seen, is frequently regenerated, and the new-formed tissue is the means of union of wounds. **Blood-vessels** are similarly restored, forming really part of the connective tissue.

**Cartilage** seems scarcely capable of regeneration, at least in the adult. Fractures of the cartilaginous ribs are united by bone, and wounds of cartilage are replaced by connective tissue or bone. Experiments on young animals show that in them there may be considerable regeneration, and it is probably so also in the human subject.

**Nerve-fibres** are regenerated after division of nerve-stems. A simple section of a nerve may be followed by immediate union, and the function may be restored in a few days. Even when portions of nerves have been removed (as much as 2 inches) there is a restoration of function, but at a much longer interval. In order to this there is a newformation of nerve-fibres from the central extremity, and these meet those of the peripheral end. The division of a nerve, unless there be immediate union, implies, as we shall see further on, a remarkable change in the whole peripheral portion from the point of section onwards. The power of regeneration and accommodation of nerves is shown by the fact that, after transplantation of skin, when the parts are separated from their nervous connections, there is a restoration of sensation.

**Muscle** is, to a limited extent, liable to regeneration. Wounds of muscles are usually united by connective tissue, although subcutaneous wounds, as proved by experiment, will often heal without cicatrix. Weber found also that in the neighbourhood of fractures no cicatrices existed in the muscles, although they had undoubtedly been torn by the ends of the bones. In the healing of wounds in muscles by granulation, it is believed by some that a newformation of muscle takes place to some extent, the muscle corpuscles taking part in the process. There is also a restoration of muscle after atrophy and degeneration. In emaciating fevers there is a great atrophy of the voluntary muscles and a restoration as convalescence advances. Fatty degeneration of muscle



is probably followed, as in other cases of fatty degeneration, by absorption of the affected structures, and this again involves a regeneration. In these cases the muscle corpuscles are not lost, and they seem to be the agents in regeneration.

**3. Transplantation.**—By this term is meant the removal of parts of the living tissues from their normal position, and their implantation in another situation. This process has been frequently effected by experiment in animals; it is not of infrequent occurrence as a pathological phenomenon in man, and it is sometimes made use of for therapeutic purposes as a surgical operation.

The best known experiment is the transplantation of the spur of the cock to other parts of the skin or to the comb. Zahn implanted a whole foetal femur into the kidney of an adult rabbit, and found that it survived and grew there. Bert performed many experiments, in which he removed the tails of rats and implanted them on their backs. A remarkable fact brought out in these experiments was, that if the implantation was made with inversion of the tail, so that the tip was in the back and the root projected out, yet the tail survived, and even sensation was restored, conduction occurring in the nerves in the reverse direction.

Several practical results arise from these experiments. For one thing, the larger the surface by which the transplanted piece was in contact with the living tissue, the greater the likelihood of success. Hence, small pieces of tissue and those which were completely buried in the living tissues were the most successful. In experiments with rats' tails, the latter required to be denuded of skin for some distance, so as to bring a considerable raw surface into contact with the subcutaneous tissue. Another fact was, that tissues from young animals were more successfully transplanted than those of adults. Again the transplanted piece commonly grew in its new situation, sometimes very markedly, as in the case of the cock's spur on the comb. This growth, however, was generally temporary, and in many cases was succeeded by diminution and complete absorption of the transplanted piece. If the transplanted piece, however, was so placed as to restore a lost part, then it remained permanently. Lastly, the tissues of animals of different species did not seem congenial, so that when, for instance, the tissues of rats were transplanted to birds they gave rise to severe inflammations (Ollier).

**In man**, transplantation sometimes occurs **spontaneously**. The greatest example of this is furnished in some cases of **Tubal pregnancy** in which, after rupture of the tube, the ovum may be transplanted to the peritoneum, acquiring adhesions there, and developing its placenta in connection with the vessels of the peritoneum. Then **Tumours** of the uterus or ovary (myomas and cysts) sometimes separate from their



seats and acquire connections with other parts. It is probable that in these cases there is a gradual transplantation, the new connections being formed before the old are completely severed. Again, **pieces** of tissue are sometimes **broken off**, such as the appendices epiploicæ in the peritoneum, or pieces of synovial membrane, cartilage or bone in joints. These may become free bodies, retaining their vitality without any vascular connections, or they may become attached in new positions.

Transplantation, as a **Surgical operation**, has long been practised. In plastic operations, involving the surface of the body or the buccal cavity, the transplantation is usually partial, the transplanted piece being left, at least for a time, in partial connection with its original seat. A complete transplantation is effected in **skin-grafting**, in which portions of the living epidermis are transplanted to the surface of granulating wounds. The granulating surface, being exceedingly vascular and composed of cells, very readily coalesces with any living structure placed on it.

**Transplantation of bone** is an interesting achievement of modern surgery. Macewen has succeeded, by successive transplantations, in restoring almost the whole shaft of the humerus, which had been lost by necrosis, and this surgeon has also shown that, after trephining the skull, the piece removed may be restored, and it will retain its vitality and acquire fresh connections. (See further under Affections of Bones).

**Literature.**—PAGET, Lect. on surg. path., ed. by Turner, 1870; HUNTER, l. c.; RANVIER, Le develop. du tissu osseux, 1865, and in Cornil et Ranvier, Manuel d'hist. path., 1881, i. *Transplantation*—RECKLINGHAUSEN (very fully), Allg. Path., 1883; HUNTER, Works by Palmer, iii., 273; ZAHN, Congrès périod. internat. Genève, 1877; BERT, Annal. de science nat., v., 1866; REVERDIN, Gaz. d. hôpit., 1870-71, Arch. gén. de Méd., xix., 1872; MACEWEN, Phil. trans. of Royal Soc., 1881, and Annals of surgery, Oct. and Nov. 1887.



## SECTION VIII.

## SPECIFIC NEWFORMATIONS. INFECTIVE TUMOURS.

*Newformations due to specific morbid poisons, which in most have been determined.*

**SYPHILIS.**—*The primary lesion, or Hunterian chancre; affection of lymphatic glands. The secondary lesion, the virus in the blood. The tertiary lesion, gummata; amyloid disease; affection of arteries; hereditary and congenital syphilis.* **TUBERCULOSIS.**—*Causation, the bacillus; tuberculosis by inoculation; contagiousness; inheritance. Character of the lesion, the miliary tubercle; how produced; caseous necrosis; fibrous transformation; softening and ulceration. Local tuberculosis; mode of access of bacillus; extension of local process, by lymphatics and along surfaces. Effects of local tuberculosis, emaciation, fever, etc. General tuberculosis, chronic and acute. Tuberculosis in animals.* **LEPROSY,** *the bacillus; character of lesions in tubercular and anæsthetic forms.* **GLANDERS;** *causation and character of lesion.* **ELEPHANTIASIS;** *causation from unknown morbid poison; character of lesion.* **ACTINOMYCOSIS;** *causation; character of lesion and locality.* **MALIGNANT LYMPHOMA or Hodgkin's disease;** *character of lesion.*

THE affections in this group show in their structure and general relations considerable analogies to inflammation on the one hand and tumours proper on the other. As already indicated, the expression 'infective' means that the disease is 'spreading,' that it depends on some virus which propagates itself, and tends to reproduce the same kind of lesion outside its original seat. In most of the forms of disease included here, the exact nature of the virus has been made out, and in all of these except one it is found to be a microbe in the form of a short rod-shaped bacillus. It may perhaps be legitimately inferred that in the rest specific microbes are the infective agents. Several of these diseases are not only infective but infectious, communicable, that is to say, from person to person; some are capable of being inoculated into animals.

The virus has, to begin with, a local seat, where it acts in a concentrated form, inducing newformation of tissue. The newformation has many features in common with that of inflammation, but it has also certain distinctive or specific features. It consists chiefly of structures analogous to **granulation tissue**, not simply as in inflammations, replac-



ing a part of the normal tissue, but forming more or less independent masses which resemble tumours. Hence, the group of diseases included here is often designated **Granulation-tissue tumours**. Outside these more specific newformations, there are usually the ordinary lesions of inflammation, the virus having acted in a less concentrated form.

But the tissue of the tumours presents certain differences from ordinary granulation tissue, chiefly in its tendencies. The granulation cells tend to undergo fatty degeneration, and so the tissue may become caseous or break down. At the same time there is the more normal tendency to undergo development into connective tissue, and this may go on in an imperfect way alongside the other change. Hence, the tumours frequently present great varieties in structure, and it is sometimes difficult fully to unravel their relations.

#### SYPHILIS.

**Causation.**—This disease is due to a morbid poison, which is asserted to be related to a bacillus, discovered by Lustgarten. This bacillus has special reactions to staining agents, which will be more fully referred to in the section on Bacteriology. The disease occurs only in the human subject, being transmitted from one person to another by contact.

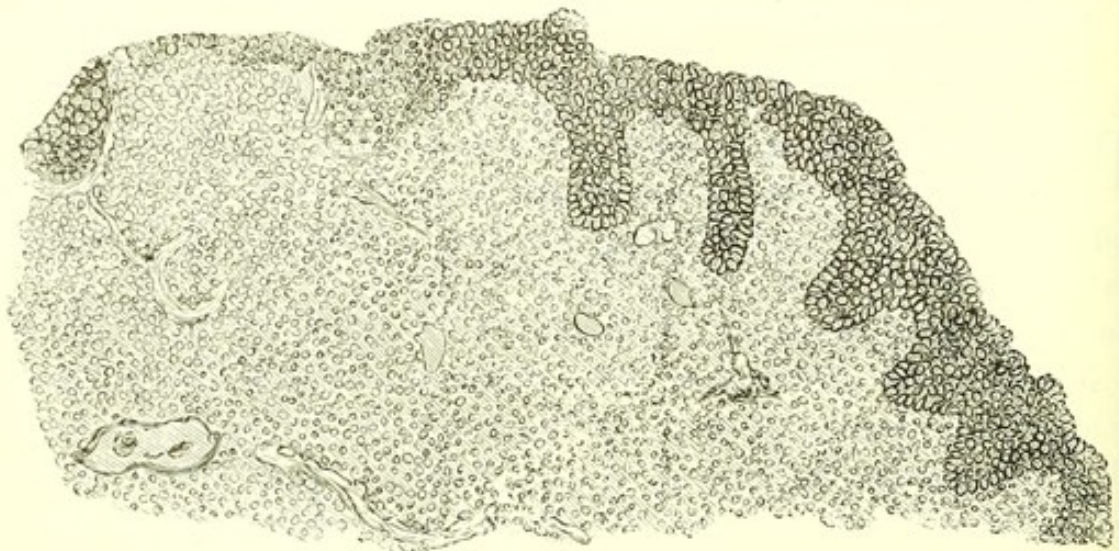


Fig. 47.—Section of a hard chancre of the lip at its marginal part. The granulation tissue occupies the skin under the epidermis.  $\times 75$ .

**Primary lesion.**—The contagium is usually applied to some part of the external generative organs. It may be applied, however, to some other part, as to the finger (in the case of a surgeon examining a part), to the lip (as in the case from which Fig. 47 is taken), to the eyelid or other external part. After a period of incubation, usually extending to three weeks or more, it begins to show signs of local action. This is in



the form first of a papule or vesicle, which acquires a hard or indurated base, and so takes the characters of the **Indurated** or **Hunterian chancre**. This consists of a raised surface, whose base has a hard, almost cartilaginous feeling. Examined microscopically, it is found that the epidermal covering of the skin is not necessarily lost, but that beneath it, there is a layer of granulation tissue, which in the central parts may to some extent replace the epidermis and come to the surface (Fig. 47). The true skin is entirely replaced by this granulation tissue, which forms a bulky mass of round cells. This structure may be regarded as inflammatory, but it may also be regarded as a kind of tumour, and may be taken as the type of the syphilitic tumour.

In its less common seats it may very closely resemble another form of tumour. Thus a chancre of the lip may be mistaken for an epithelioma, and be excised under that impression. This actually occurred in the case from which Fig. 47 was taken.

The tissue, although like in structure to granulation tissue, does not readily form connective tissue, but remains long in the same rudimentary condition, and when it disappears leaves comparatively little of a cicatrix. On the other hand it sometimes becomes caseous, but this does not so readily occur in the primary lesion.

The virus is carried from the indurated chancre by the lymphatics, and lodges in the neighbouring **lymphatic glands**, where it again produces similar results, namely a great production of ill-formed granulation tissue which has little tendency to develop into proper connective tissue, but readily undergoes an irregular caseous metamorphosis. The glands undergo a slow enlargement and become hard.

**Secondary lesions.**—These are due to the fact that the virus passes from its local seat into the blood. This it does after an interval of some weeks from the time of the primary lesion.

The virus probably reaches the blood directly from the primary chancre, as well as by the lymphatics, but a certain period of time is necessary before it reaches the blood in sufficient quantity to produce any effect. Lang distinguishes in syphilis two periods of incubation, a **First incubation** extending from the time of infection to the appearance of the primary lesion, and a **Second incubation** between the primary and the secondary lesions. The first incubation has, according to the statements in recorded cases, a minimum duration of ten days, a maximum of forty-two days, and an average of twenty-four days. The second incubation is longer, having a minimum of eight to fourteen days, a maximum of one hundred and fifty-nine days, and an average of six to twelve weeks.

By some it is believed that the induration of the primary lesion is itself due to a constitutional infection, and they cite in proof of this the fact that auto-inoculation, from an indurated chancre, is rarely successful, that is to say, the person is already protected by the primary indurated chancre from the production of another chancre by inoculation. It is to be observed, however, that in all such cases the original



chancre has the start of the inoculation, and before the period of incubation of the inoculation has elapsed there may be time for the blood to have been saturated with the virus. It is admitted that auto-inoculation is possible when done early enough. It seems scarcely possible to explain the second incubation except on the supposition that the primary lesion is, at least relatively, a local manifestation.

When an agent exists in the blood in a finely divided state it will be carried to all parts of the body, and if it produces lesions they will probably be symmetrical, as the corresponding parts in each lateral half of the body are for the most part in similar circumstances, and are similarly affected by any agent acting equally on them. The existence of symmetrical lesions is presumptive evidence that a disease is due to something in the blood. In the secondary stage of syphilis then we have the virus in the blood, and the result is symmetrical lesions of the skin, mucous membranes, bones, etc.

These secondary lesions are inflammatory in character, and have generally a resemblance to those of ordinary inflammations. They are most frequent in the skin, and so we have the syphilitic Roseola, Eczema, etc., but other parts may be affected, and we have syphilitic Periostitis, Pharyngitis, etc. It is a question to what extent inflammations occur in internal organs in this stage; according to Hutchinson they are more frequent than is generally supposed, but are rarely seen because persons seldom die in the secondary stage.

The secondary stage has been aptly compared with the eruptive stage of specific fevers, it is like a fever long drawn out. There is in both cases a virus in the blood, and in syphilis there is frequently elevation of temperature. The analogy between the rash of secondary syphilis and that of measles, scarlet fever, small-pox, etc., is also suggestive, the skin affections in both classes of cases being inflammatory. During this stage then the virus is active in the blood, and the blood and secretions are contagious. The person is also in the position of transmitting the disease to his offspring, the virus apparently passing into the germ and sperm cells. Just as in the specific fevers, the virus dies out of the blood spontaneously, and the various secondary lesions disappear, generally in six to eighteen months.

In this secondary stage it is not common to meet with tumours like the indurated chancre. They are characteristic rather of the next stage, and when they do occur in this stage they are small and accompanied by more pronounced inflammatory manifestations.

A certain approach to the formation of granulation tissue, however, is often seen in the skin and mucous membranes during the secondary stage in the form of **mucous tubercles** or **flat condylomata**. These are flat superficial elevations of the skin or mucous membranes, usually met with near the external organs of generation, the anus, or the mouth and pharynx.



**Tertiary lesions.**—These are chiefly characterized by the formation of tumours to which the name **Gummata** is applied. They are composed similarly to the indurated chancre, of granulation tissue, but in them this tissue has a much greater tendency to undergo caseous necrosis, and is accompanied by a greater newformation of connective tissue.

To the naked eye the tumour is a whitish or greyish body, commonly with a yellow caseous appearance in its central parts, or irregularly distributed. It varies in size ; it is sometimes as small as a millet seed, in which case it is usually multiple, but it is generally much larger, and may attain the size of an apple. The tumour is not generally sharply defined, but its periphery merges in a firm connective tissue which usually extends outwards into neighbouring structures, so that the tumour appears planted in the midst of a cicatrix.

Under the microscope the tumour will be found, as in Figs. 48 and 49, to replace a certain portion of the normal tissue. The central caseous part will be opaque as in *d*, Fig. 48, and *c*, Fig. 49. Externally the tissue is more transparent (*b*, Fig. 49), while around and in neighbouring parts of the organ there is new-formed connective tissue, as at *d* in the figure.

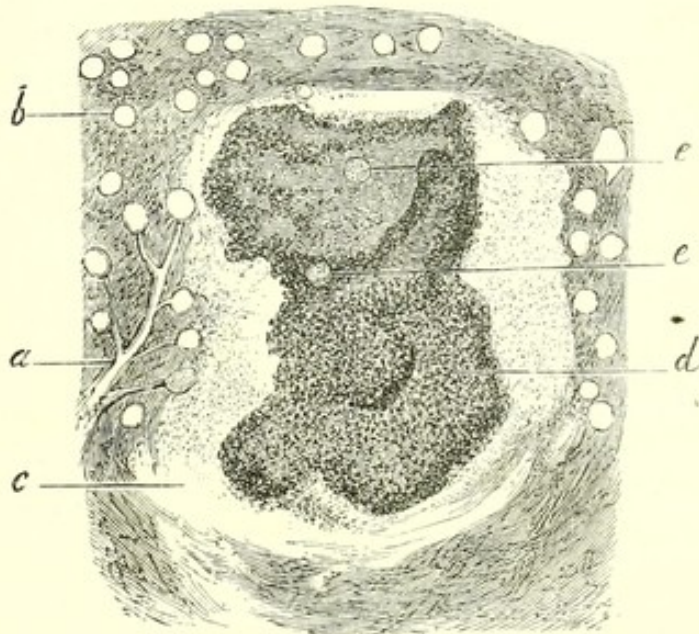


Fig. 48.—Gumma of kidney. The opaque central part (*d*) is caseous ; around this there is fibrous and granulation tissue (*c*).  $\times 20$ . (CORNIL and RANVIER.)

In Fig. 50 the appearances seen under a higher power are shown, the parts taken being from *b* and *c*

in Fig. 49. It is seen that the peripheral portions (Fig. 50 *a*) of the tumour present innumerable round cells mixed with fibrous tissue, which latter is often very pronounced. The caseous parts are opaque and present fine fat granules with shrunken cells and nuclei (Fig. 50 *b*).

The tumours are met with in almost all the tissues of the body, skin, mucous membranes, subcutaneous tissue, in the substance of muscles (as in the tongue), periosteum, liver, dura mater, soft membranes of the brain, cerebral nerves, etc. The name gumma does not express their usual consistence, and is stated to have been first applied to the periosteal form.



The caseous necrosis leads to various results, according to the situation and circumstances of the gumma. If the tumours have a superficial situation, then **Ulceration** results, and we have a deep excavated ulcer with swollen infiltrated walls, consisting of tissue like that of the gumma, and with the same tendency to degeneration, so that the ulceration extends. As the tumour involves neighbouring structures which undergo necrosis along with the caseous process in the tumour, there may be great destruction of tissue brought about. In internal organs the caseous material may long lie apparently unaltered. The gumma may be virtually healed, its granulation tissue absorbed or converted into connective tissue, while the caseous matter remains, and is finally left in the midst of a cicatrix where it may become calcified.

We have still to inquire what may be the relation of these tertiary lesions to the virus. The virus no longer exists in the blood, and in accordance with this the lesions are characteristically unsymmetrical. The most probable supposition is that, after the close of the secondary stage some of the virus has remained lying in a particular part. Perhaps a small gumma has formed, and the virus has lain in it quiescent but still surviving. It may be waked up by some accidental circumstance in the life of the patient, at any period afterwards, perhaps as long as twenty years. The virus propagates itself, but its effects are local. It may produce a tumour of large size, but it does not pass into the blood, and does not produce the lesions of the secondary stage. It has been matter of dispute whether a tertiary gumma is an infectious lesion, and the fact that the blood of the patient himself does not become infected might seem to answer the question in the negative. It is to be remembered, however, that the affected person already possesses an immunity by having passed through the secondary stage. The gumma is probably capable of producing syphilis when its juices are brought into contact with the tissues of a susceptible person.

While the conditions described as characteristic of the tertiary stage usually succeed those of the secondary period, it should be understood that there are great variations both in degree and order of occurrence. The secondary manifestations may be greatly prolonged, and the tertiary may develop to some extent coincidently. The seat of the tertiary lesions varies very greatly.

The tertiary stage of syphilis is often in its later period associated with **Amyloid disease**. This may be due in some cases to chronic suppurations induced by the specific lesions, but the observations of Fagge show that it is not always so. Of 76 cases of amyloid disease associated with syphilis there was evidence of former or present bone disease only in 34. In the total autopsies in cases of syphilis over a period of years,



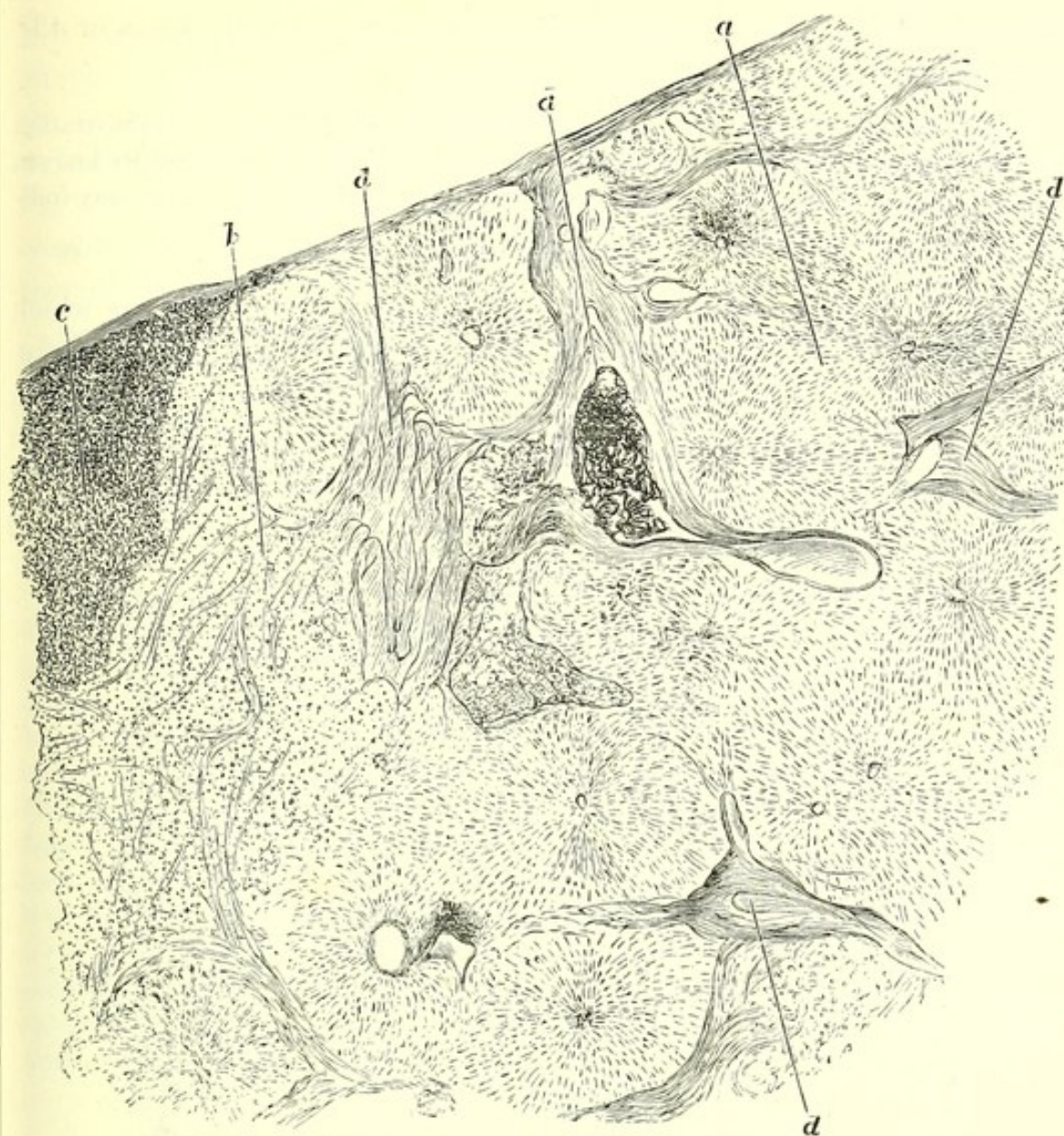


Fig. 49.—Gumma of liver. Explanation in text.  $\times 16$ .



Fig. 50.—From gumma of liver, same section as Fig. 49; *a*, taken from recent part (*b*, in other figure); *b*, from caseous (*c*).  $\times 350$ .



amounting to 177 cases, amyloid disease was present in 76 cases or  $43\frac{1}{2}$  per cent.

It appears that the **absorption of a gumma** may be promoted by the administration of remedies. The exact process by which this is brought about is hardly known, but there seems to be a simple fatty degeneration with absorption, in the way indicated in the section on fatty degeneration.

Syphilis is often associated with a **condition of the arteries** which will come up for discussion further on. Wherever there is, as so frequently happens, a considerable formation of granulation tissue passing into connective tissue, the arteries take part in the inflammation, and we have, especially, thickening of the internal coat, sometimes going on to complete obliteration of the calibre of the vessel. (See Fig. 51.)

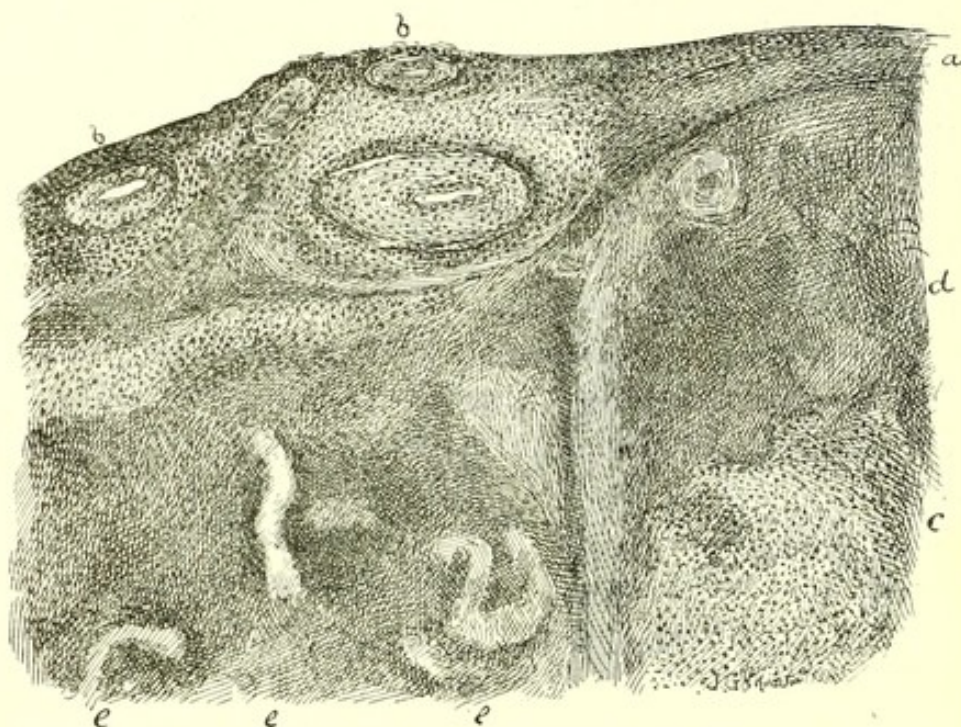


Fig. 51.—Syphilitic affection of meninges and brain. *b, b*, Arteries showing thickening of internal coats.

This is sometimes very strikingly seen in the neighbourhood of gummata, and by diminishing the blood supply, it may contribute to the degeneration of the gumma. It may also lead to degenerations in parts around, as where softening of the brain occurs in connection with gummata.

**Hereditary syphilis.**—We have already seen that syphilis, in the secondary stage at least, may be transmitted to the offspring. In the acute period death often occurs *in utero*, or the child sickens soon after birth and dies within a few weeks. But it often happens that the children do not show any evidence of syphilis for months or years. In



this way we may distinguish cases of congenital syphilis from cases of simple hereditary syphilis, the former being born with syphilitic lesions, the latter acquiring them afterwards.

In **Congenital syphilis** the most constant and unequivocal lesion is the affection of the bones, which will be considered afterwards. In this condition there is an error in the process of ossification, with inflammatory conditions.

In **Hereditary syphilis** the lesions are, like those of the secondary stage, mainly inflammatory. There are inflammations of the skin, mucous membranes, cornea, etc. The characteristic malformation of the teeth which Hutchinson has pointed out seems related to inflammation of the mucous membrane of the gums during the development of the teeth.

**Literature.**—The author, in describing the general pathology of syphilis, has followed chiefly Hutchinson and Virchow. HUTCHINSON, *On Syphilis*, 1887, and *Debate on Syphilis*, *Path. trans.*, vol. xxvii., 1876; see also other speakers in this debate; VIRCHOW, *Krankhafte Geschwülste*, ii., p. 393; LANG, *Path. und Therap. der Syphilis*, 1884; RICORD, *Traité prat. des malad. vén.*, 1838; LANCEREAUX, *Traité hist. et pratique de la syphilis*, 1866; HILTON FAGGE, *Medicine*, vol. i., p. 109, and *Path. trans.*, vol. xxvii.; KASSOWITZ, *Die Vererbung der Syphilis in Stricker's Med. Jahrb.*, 1875, p. 359; FOURNIER, *Leçons sur la syphilis*, 1881; VAN HARLINGEN, *International Encycl. of Surgery*, vol. ii., 1882. *Bacillus of Syphilis*—LUSTGARTEN, *Med. Jahrb. d. Wien Gesellsch. der Aerzte*, 1885. Since the publication of Lustgarten's observations, many have observed the bacillus, but some authors have stated that it exists in the normal smegma of the prepuce. This controversy is chiefly in the pages of the *Deutsche Med. Wochensch.* See also BITTER, *Virchow's Arch.*, cvi., 1886.

## TUBERCULOSIS.

Tuberculosis is an infective disease in which the tissue-changes are due to the action of a specific virus or morbid poison. As in the case of syphilis there is always a local or primary lesion, but it is only exceptionally that the virus extends to the blood and infects distant parts of the body. On the other hand the local lesion is, for the most part, a constantly extending one, infecting neighbouring parts till the death of the person, which is usually brought about, directly or indirectly by the tubercular process.

**Causation.**—While tuberculosis is due to a specific virus, there are undoubtedly conditions of the body which predispose certain persons to its action. Considering that this virus is very largely present in nearly all inhabited places, it must be apparent that most people are exposed to its action, and yet only a limited number of persons become affected. For practical purposes, therefore, the predisposition may be regarded as perhaps of greater importance than the actual infection.



This predisposition is frequently the result of inheritance. It has been abundantly proved to be so in the case of the commonest form of tuberculosis, namely that of the lungs, but not so obviously in other forms. Inheritance determines a local predisposition usually in one organ or system (see *ante*), but when a tuberculosis is once established in the body, it is not uncommon to find it extend by one path or other to other organs, even without there being evidence of a general extension by the blood.

The **Tubercular bacillus** is the essential agent of infection. This is a micro-organism, consisting of minute threads, each measuring in length about two thirds of the diameter of a red blood-corpuscle.

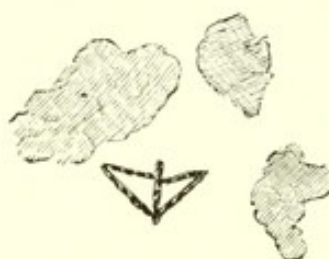
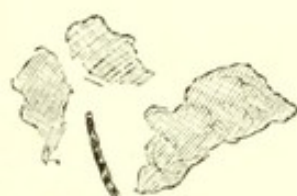


Fig. 52.—Tubercular bacillus prepared from sputum in phthisis pulmonalis.  $\times 1500$ .

The thread contains small rounded bodies like spores. (See Fig. 52.) The bacillus is present in somewhat varying degrees of abundance in different tubercular lesions, but it has been met with in all forms of tuberculosis.

The connection of the bacillus with tuberculosis and the essential identity of the various forms of the disease have been proved

by the cultivation of the microbe outside the body, and by the results obtained by inoculating the cultivated products.

The methods of cultivation will be referred to in the section dealing with micro-organisms. The result of inoculation in suitable animals is the production of local or general tuberculosis according to circumstances. The bacillus is recognized both by its form and by its reaction when treated with staining agents. It is not stained with ordinary watery solutions of the aniline dyes, as most microbes are, but it is readily stained by solutions of fuchsine or gentianviolet containing free aniline or carbolic acid, and when so stained it does not yield its colour when treated with nitric or sulphuric acid. (See section on Bacteriology.)

The tubercular bacillus grows slowly and only at a temperature approaching to that of the living body. We may therefore infer that it does not to any appreciable extent multiply except in the body, as the conditions will seldom be favourable as regards soil and temperature, unless these are carefully arranged for experimental purposes. On the other hand the bacilli and especially the spores are very resistant to external influences, so that they retain their vitality under adverse circum-



stances, and are ready to begin growth when they obtain a fitting locale. The bacilli and spores survive even when dried, and they may be suspended in the air and carried as dust, ready to deposit on surfaces or to be inhaled during respiration. Tuberculosis is not commonly produced by direct communication by contact between person and person, in this respect contrasting with syphilis.

The bacilli in themselves, consisting of minute threads, cannot be supposed to have much effect upon the tissues. They produce their results by means of irritant products which they evolve. These products will be most concentrated in the immediate vicinity of the bacilli, but being in solution they will have a wider action in various states of dilution. Hence it is possible to recognize a more specific lesion presumably from the concentrated action, and certain less specialized conditions from the more diluted action. Both of these somewhat resemble inflammatory processes, but the former is characterized by small local formations called tubercles, while the latter has more of the characters of ordinary inflammation.

The infective nature of tuberculosis does not rest alone on the facts connected with the tubercular bacillus, it was inferred long before the discovery of this micro-organism. The most direct proof consisted in the production of tuberculosis by the inoculation of tubercular material in animals. This was first demonstrated by Villemin in 1865, and was repeatedly confirmed by many observers, among whom may be mentioned Lebert, Waldenburg, Klebs, Cohnheim, Tappeiner, Bollinger, before the discovery of the tubercular bacillus in 1882. The artificial production of tuberculosis is now regularly performed in experimental laboratories, and the material used may be tubercular products obtained from post-mortem examinations or otherwise, the sputum of patients suffering from phthisis pulmonalis, or artificial cultures of the bacillus. The disease has been induced by inoculation (Cohnheim, Koch, Baumgarten), by inhalation (Tappeiner), and by feeding (Wesener, Bollinger).

As an example of the mode of inoculation and its results we quote the following from Koch's work on the Etiology of Tuberculosis (Sydenham Society's translation, 1886). It recounts the effects of inoculating fragments of tissue from various organs in cases of human tuberculosis, such as from the lungs in phthisis, from strumous joints, scrofulous glands, lupus, and from tuberculosis in various animals.

"The inoculation was effected by making a small incision in the abdominal wall of a guinea-pig with the scissors, inserting the point of the scissors to form a pocket-like subcutaneous wound, about half a centimetre deep. Into this little pocket a fragment of the inoculation substance about the size of a millet or mustard-seed was pushed as deeply as possible. On the following day the inoculation wound was always united, glued together, and showed no reaction. Generally it was not till after a couple of weeks that a visible swelling of the lymphatic glands next the seat of inoculation occurred, usually the inguinal glands on one side, and at the same time induration and the development of a nodule took place in the inoculation wound, which up till then had remained perfectly healed. After this the lymphatic glands enlarged rapidly, frequently to the size of a hazel-nut.



The nodule at the seat of inoculation then generally broke and became covered with a dry crust, beneath which was a flat ulcer with a cheesy floor discharging very slightly. The animals began to lose flesh about this time, their coat became bristly, dyspnoea set in, and they died generally between the fourth and eighth weeks, or they were killed within the same space of time."

Tuberculosis has been induced or observed in **nearly all the commoner warm-blooded animals**, and according to Koch no bird or mammal is capable of permanently resisting infection. Animals show, however, very different degrees of susceptibility, and in similar degrees they are variously liable to spontaneous tuberculosis. Rabbits, guinea-pigs, cattle, and apes are peculiarly susceptible. Carnivorous animals are much less so, spontaneous tuberculosis being very rare, for instance, in cats or dogs. In the same class of animals the individuals show varying susceptibility just as in the case of man.

The infective character of tuberculosis is also to be inferred from the spreading character of the lesion, to be now described.

**Contagiousness of tuberculosis.**—From the inoculability of the tubercular bacillus it might be supposed that tuberculosis would be eminently contagious. In this respect, however, it differs from syphilis and most other analogous conditions. It appears as if the element of predisposition were so important that mere contact is seldom sufficient to induce the disease in persons not predisposed. It seems necessary in order to direct communication that the virus be applied in concentrated form. Thus phthisis pulmonalis may be communicated to a person who habitually sleeps with one subject to this disease. The chronic thickenings or warts to which the fingers of pathologists are liable, and which are sometimes called post-mortem tubercles, are asserted to be due to the inoculation of tubercular bacilli.

**Inheritance of tuberculosis.**—While inheritance plays a very important part in the predisposition to tuberculosis, it has not yet been proved that, in man, tuberculosis is ever communicated from parent to offspring, although there are some cases recorded in infants which may have been congenital. This is consistent with the fact that the bacilli do not readily pass through membranes. Johnes has recorded a case of congenital tuberculosis in a calf whose mother was affected with severe tuberculosis of the lungs. In the calf the tuberculosis was situated in the liver, portal glands, and bronchial glands, the infection having been by the umbilical veins in the placenta. (Referred to by Birch-Hirschfeld.)

**Character of the lesion.**—The typical lesion in tuberculosis is the so-called miliary tubercle, and it is this which was referred to above as being related to the concentrated action of the virus.

**The miliary tubercle**, is a minute rounded body (Fig. 53) composed of cells and devoid of blood-vessels. A single tubercle is scarcely visible to the naked eye, but, by the confluence of several there may be



larger nodules formed, and even, by the concurrence of vast numbers in successive generations, large masses. At first sight the tubercle seems composed simply of round cells, but closer inspection shows usually three forms to be present. The most peculiar and typical is the **Giant-cell**, which in the early stages is nearly a constant constituent. It is a large body (seen in the middle in Fig. 53) presenting at its margin radiating processes, especially at its poles, and containing numerous oval nuclei, usually arranged in a row near the periphery of the cell. The giant-cell is generally near the centre of the tubercle, but may be considerably removed from this position. There are frequently several in one tubercle. The second constituents, the **Epithelioid cells**, are smaller than the giant-cells and possess one nucleus, but are still of considerable size. They resemble epithelial cells, and are called, like the similar cells in the inflammatory newformation, epithelioid cells. They surround the giant-cell and in their general appearance somewhat resemble it. **Ordinary round cells** or leucocytes form the third constituents and they are variously abundant according to circumstances. They are present at first at the periphery, and may be regarded as representing the ordinary inflammatory products, which are commonly seen in the tissue around. The giant-cells and epithelioid cells seem to be the essential constituents of tubercles, and in perfectly recent examples they may be almost the only ones. They are, according to Baumgarten, the immediate products of the action of the tubercular bacilli, the leucocytes being the ordinary result of inflammation.

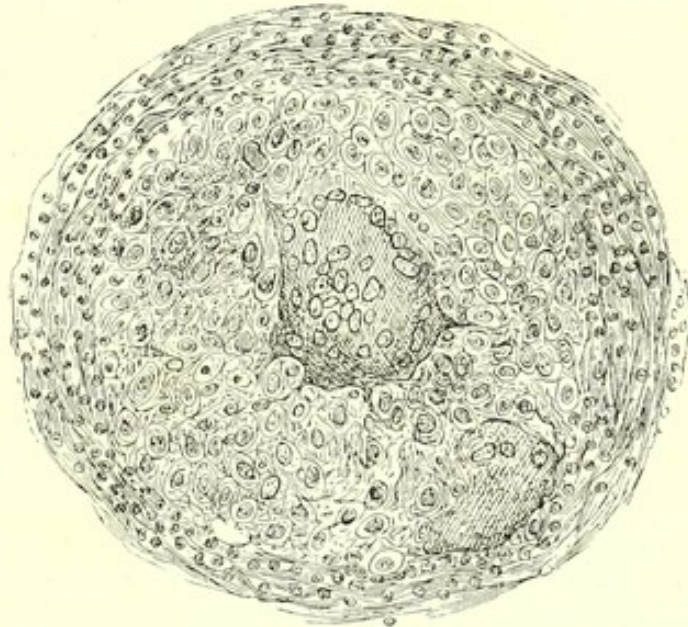


Fig. 53.—Tubercle showing constituents. A giant-cell is in the middle, and a smaller one lower down. Epithelioid cells form the rest of the tubercle except at the margins, where leucocytes are present.

Baumgarten, from a very elaborate series of experimental studies, has described the **Origin of the various constituents of the tubercle**, and their relation to the bacilli. When bacilli invade a tissue, the fixed cells of the tissue, those of epithelium as well as of connective tissue, first show evidences of change. They are induced to proliferate by the bacilli which may be in their substance or in their neighbourhood. The nuclei of these cells show karyokinetic figures, and both giant-cells and epithelioid cells result from the changes in the fixed cells of the tissues. In



the case of the giant-cell the nuclear division goes on, but the cell division does not ensue, there being thus a kind of arrest of development. By this process a rounded tubercle arises which is composed entirely of giant-cells and epithelioid cells. But the neighbouring blood-vessels are affected by the virus, and from them an emigration of leucocytes occurs. The leucocytes are at first outside the **large-celled tubercle**, but penetrate into it, and may convert it into a **small-celled or lymphoid tubercle**. This transformation may be rapid or slow, its course depending on the number and energy of the bacilli. If the latter are abundant and vigorously propagating, or if they are accompanied by other foreign micro-organisms, then the production of leucocytes may be very great, and the tubercles rapidly pass into the small-celled condition. This may occur so rapidly that the large-celled condition may be almost omitted. Hence it is where the bacilli are comparatively few and the cultivation pure that the typical large-celled non-inflammatory tubercle is most liable to occur.

Bodies having a structure similar to that of tubercles may be produced by the presence of small foreign bodies (rabbit hairs, Baumgarten; particles of stone, Hamilton), but in all such cases the lesions have not an infective character. A more near approach to the true tubercle is met with in Actinomycosis, where the presence of a micro-organism has the effect of producing results not unlike those of tuberculosis (see Actinomycosis).

The typical tubercle, as here described, is not infrequently difficult of recognition in actual cases. Inflammatory processes occurring around and invading it, obscure its structure, and the tubercle itself is prone to degenerative changes.

**Caseous necrosis or Caseation** is the most characteristic change in tuberculosis. As already described this change consists in necrosis with fatty degeneration of the structures. It manifests itself in a granular condition of the cells, whose nuclei disappear. The result is an obscuration of the structure, and the replacement of it by a homogeneous structureless and somewhat opaque material. This change usually begins in the central parts of the tubercles and produces an opacity there (see in Fig. 55, p. 207), but it is liable to overtake the whole structure. Indeed, it extends to the parts around, so that the tissue which is merely infiltrated with inflammatory products is frequently involved in the caseation. Hence, when the process has overtaken a considerable portion of tissue, the structure may be much obscured. This process in the rapidity and completeness of its occurrence depends on the abundance and vigour of the bacilli. Where they are numerous and where consequently there is much infiltration with leucocytes, caseation is rapid (Baumgarten). The caseous material so produced is a yellow, brittle substance, which resembles cheese in its appearance. So characteristic is this process of tuberculosis that caseation is almost equivalent to the older term **Tuberculization**, and caseous matter to **Crude or Yellow tubercle**.

In very acute cases we may have an approach to ordinary necrosis or sloughing, or at least an acute softening.



The process of caseation is to be related to the action of the virus. Where it acts most vigorously caseation is most characteristic. Some have endeavoured to account for the caseation by the absence of vessels in the tubercles. But much larger pieces of matter may be kept alive in the body without the intervention of vessels, such as free bodies in the joints or abdomen. Besides, the caseation extends beyond the tubercles to the tissue around, which is vascular.

**Fibrous transformation** is a much more unusual change in tubercles, and occurs only where the process is very chronic and the bacilli very few. While some tubercles undergo caseation, others, and along with them the surrounding tissue, develop fibrous tissue like that in chronic inflammation.

**Softening** is the usual result of caseation. The caseous matter, although it may remain for a time unaltered, in most cases ultimately



Fig. 54.—Tubercular ulcer of intestine. The rounded form of the tubercles around the ulcer (*a*) is seen, some of them breaking up the muscular coat (*b*); one (*d*) outside this coat, and beneath the peritoneum.  $\times 16$ .

liquefies or breaks down. The result is the formation of a **Cavity or Ulcer**. (See Fig. 54.) It is to be remembered that, as the caseation involves not only the tubercles but the surrounding tissue, the cavity or ulcer is associated with an absolute loss of tissue. The walls of the ulcer will be tubercular, and the disease will usually continue advancing, so that the ulcer or cavity will enlarge.

On the other hand, the **caseous matter** may accumulate without softening, and we may have considerable masses of it, such as we find in the brain and in the lymphatic glands. In course of time the caseous matter may be partly absorbed, or may undergo **Calcareous infiltration**.



**Scrofula and Struma.**—These terms are used to designate conditions of the lymphatic glands, which are now generally recognized as tubercular. They will be described in their proper places. The terms have been somewhat indefinitely extended so as to include affections of joints, bones, skin, and elsewhere, but such affections are nearly all really tubercular.

**Local Tuberculosis.**—We have already seen that tuberculosis is always, to begin with, a local affection, due to the implantation of bacilli in the living tissue, and it may be added that, even when it becomes generalized, the secondary lesions are due to the multiplication of bacilli transplanted from the primary seat, the growth of the micro-organism occurring not in the blood but in the local seat. Hence the description already given applies to the local process however originating.

The bacillus finds access to the body by various channels. The most common seats of tuberculosis are in direct communication with the surfaces of the body. The commonest seat of all is the **Lungs**, and entrance is here obtained by the inspired air. Next to the lungs the **Lymphatic glands** are most frequently affected, and in the case of children they are perhaps more frequently the seat of tuberculosis than the lungs. The lymphatic glands which are attacked, are in communication either with the skin or a mucous membrane, and there are frequently catarrhs or other forms of inflammation in the tissue with which the affected glands are related. The bacilli may, however, find their way from surfaces which are unaltered. The glands most frequently affected are those of the neck, communicating with the mouth, of the mesentery, communicating with the intestines, and of the bronchi, communicating with the lungs. The skin may be directly attacked by tuberculosis, the resulting conditions being described as Lupus or Scrofuloderma.

While these more direct modes of entrance constitute the majority of cases, there remain a large number in which the access is more circuitous. In some of these the tuberculosis is due to a secondary extension from a tubercular lymphatic gland. It is thus that many cases of tuberculosis of **Serous membranes** arise. But in a considerable number of cases, the bacilli can only have found access by the blood. This applies to tuberculosis of the **Bones**, which is very frequent in children, and which often begins in the vertebræ, or in the cancellous tissue at the extremities of the long bones. The **Brain** is also liable to primary tuberculosis, especially in children. Tuberculosis of the **Urino-genital** system begins very frequently in the testicle, and the bacilli are carried thither by the blood.

In some of the cases mentioned, tuberculosis exists in some primary seat, and the bacilli may be carried from it, but in many cases we can



only suppose that a few bacilli are accidentally conveyed, and, finding seats predisposed to their reception, multiply and produce their effects. The blood is not infrequently the vehicle of small solid particles, whether micro-organisms or not.

**Extension of local tuberculosis.**—A tuberculosis once established in a locality presents usually a tendency to indefinite extension. Tuberculosis is generally a slow process, and the extension is also chronic. The extension is to the immediately neighbouring parts, by direct infiltration, or, more commonly, by the lymphatics or along surfaces and canals. The process of extension is usually stopped by the intervention of a membrane, as the bacilli seem not to be possessed of the power of penetrating membranes unless there is first a necrosis of them. Thus, tuberculosis of lymphatic glands does not pass through the capsules unless the latter have been perforated by necrosis; and tuberculosis of the lungs rarely extends to the pleura unless there be actual perforation of that membrane.

**Extension by the lymphatics** is very frequent and characteristic. Tubercles often form in the course of the lymphatic vessels connected with tubercular organs, as well as in the lymphatic glands. In phthisis pulmonalis, for example, there are tubercles in the substance of the lung seated in the lymphatic vessels, while the bronchial glands are also the seat of tuberculosis.

**The extension along surfaces** and canals is exemplified chiefly in serous and mucous membranes. A tuberculosis occurring in the pleura, pericardium, or peritoneum extends over the entire surface of these membranes. Tuberculosis, in communication with mucous canals, frequently travels considerable distances, involving the surface more or less continuously, but penetrating deeply to a very slight extent. Thus, tuberculosis of the lungs is frequently associated with tuberculosis of the bronchial mucous membrane, of the mucous membrane of the trachea and larynx, and of the intestine (from swallowing the expectoration). Again, tuberculosis of the urino-genital organs frequently begins in the testicle. From this it will extend the whole length of the vas deferens to the vesiculæ seminales and urinary bladder, and sometimes up the ureters to the kidneys.

It may here be remarked that in artificial cultures, the tubercular bacillus grows on the surface of the medium, and does not penetrate into its substance. (See section on Bacteriology.)

**Effects of local tuberculosis.**—The process of tuberculosis, as already described, involves destruction of tissue, and in many cases ulcers and cavities are the result. The destruction of tissue may itself involve serious consequences, as in the case of the bones and of the brain, in



which latter position the tubercular mass also acts as a tumour, pressing on the brain substance around. Most of the evil consequences, however, arise in connection with the ulceration and formation of cavities. The surfaces thus produced discharge and use up the available nutritious material of the body. Such processes are, therefore, accompanied by **Emaciation**. Even more potent in the production of general emaciation is **Fever**, which commonly accompanies tuberculosis. Fever in general is produced by the presence of foreign matter in the blood (see further on). In tuberculosis the bacilli do not produce fever directly, as they are not present in the blood, but their products are. We have seen that these products in their most concentrated form lead to the formation of tubercles, and, when less concentrated, produce inflammations around. They find their way into the blood in a still more dilute state, and there produce fever. In the case of discharging surfaces, again, there will commonly be putrid and other forms of decomposition in the matter discharged, and the absorption of the products will still further conduce to fever.

**General tuberculosis.**—This term is used in contradistinction to local tuberculosis, to designate a condition in which the tuberculosis has not one but many centres to which the virus has been carried by the blood. It has been pointed out above that tuberculosis has little tendency to penetrate deeply, preferring to extend along open channels or surfaces. Hence, the disease, for the most part, remains local, at least in the sense that it does not extend to the blood. It is also to be noted that the bacilli apparently do not multiply in the blood, this fluid only acting as their vehicle of conveyance. When carried by the blood, the bacilli may or may not settle down and multiply. This will depend on the number and vigour of the bacilli, and the resisting power of the tissues. If they do obtain a settlement they will slowly grow, and produce their usual effects, namely, the formation of tubercles and inflammation. Hence, in general tuberculosis we have what is equivalent to a number of local tubercles. For convenience of study, Weigert has divided general tuberculosis into chronic and acute.

1. **Chronic general tuberculosis.**—While tuberculosis does not readily penetrate deeply, still in vascular organs a few bacilli will occasionally reach the blood, passing either directly into the vessels of the part, or indirectly by the lymphatics. These may settle in predisposed situations, and give rise to secondary tubercular lesions in several different centres. When a few bacilli, thus at intervals, find entrance to the blood, the lesions will usually be comparatively few. These secondary lesions will run a chronic course and may sometimes rival the primary tuberculosis in size and effect on the body. Chronic general tuberculosis is, therefore,



characterized by the presence of a number of considerable tubercular lesions in different organs, and the case frequently looks like one in which several local tubercloses are simultaneously present. This form of disease is most frequently seen in children, whose tissues seem more susceptible to the tubercular bacilli in small numbers. In them we may find tubercular masses simultaneously in lungs, kidneys, brain, etc.

2. **Acute general tuberculosis, Acute miliary tuberculosis.**—In this disease there is a simultaneous and wide-spread eruption of tubercles in several organs. The tubercles have the usual structure, but they are present in vast numbers in lungs (see Fig. 55), liver, spleen, kidneys,

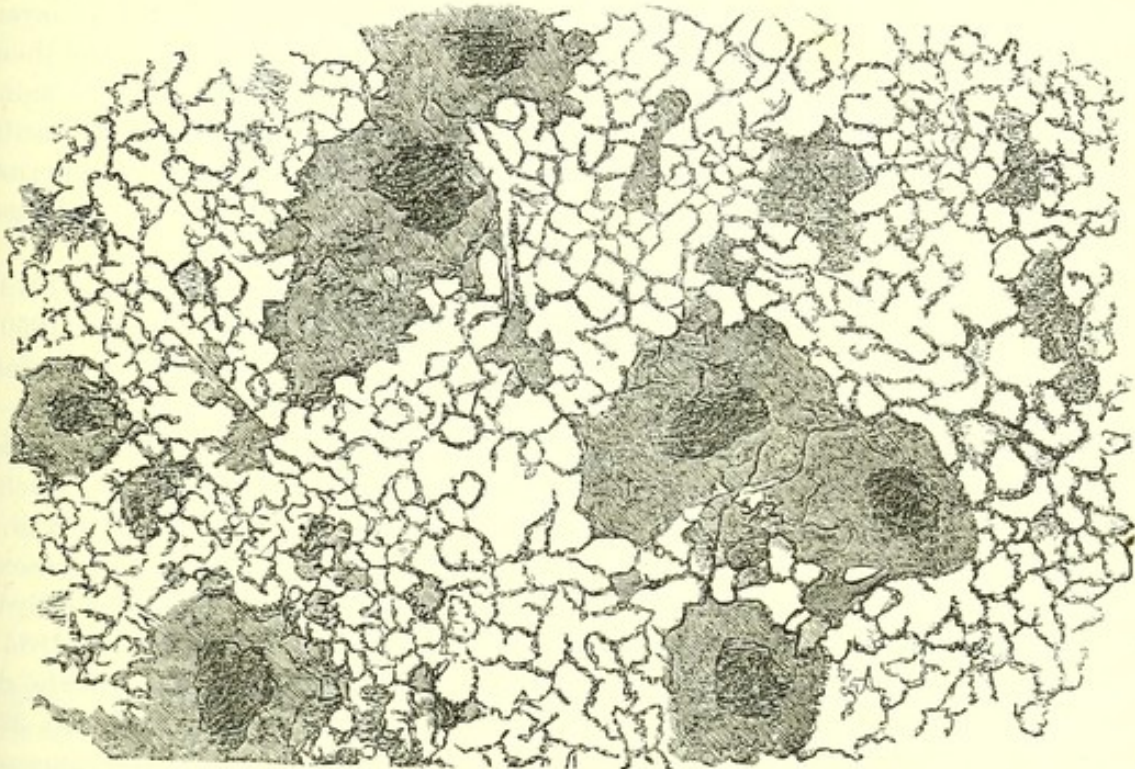


Fig. 55.—Miliary tubercles in lung. The opaque central parts are caseous.  $\times 16$ .

sometimes in the meninges, lymphatic glands, supra-renal capsules, etc. The author has seen them in the heart and endocardium. The disease is an acute febrile one, fatal in a few weeks, and the individual tubercles are small in size, each local formation being little more than a single miliary tubercle. They are often so small as to be only detectable by the aid of the microscope, especially in the liver where they are present in almost incredible numbers.

The simultaneous outbreak of numerous tubercular lesions implies that in a short interval the blood has received large numbers of bacilli. In some cases the bacilli have been detected in the blood during life (Weichselbaum), and not infrequently after death.

This overloading of the blood with bacilli can only occur when tuberculosis has extended so as to involve the wall of a still pervious vessel of some size, so that the bacilli may find free access to the blood.



Acute general tuberculosis has been found associated with tuberculosis of the thoracic duct, of the pulmonary, splenic, portal, hepatic and other veins. The extension may also occur from lymphatic glands directly into the blood-vessels of these glands (Koch).

This solution of the problem of acute general tuberculosis was first suggested by Ponfick, who observed tuberculosis of the thoracic duct in a case of this disease. Weigert has, in a large proportion of cases, found the source of the general infection in the extension to pervious veins, and he asserts that in most cases such a source will be found if diligent search be made. The pulmonary vein is that most frequently involved, and it is so in about half the cases. In a case observed by the author, in which the primary tuberculosis affected the lung, a branch of the pulmonary vein, the size of a crow quill, presented in its wall an elongated, yellow caseous layer, somewhat resembling a thin localized thrombus. The surface was smooth and there was no layer of fibrine between it and the calibre of the vessel which was here uninterrupted, except by the slight projection of the yellow mass. Tubercular bacilli were found in large numbers in this yellow structure, extending to its internal aspect. Bacilli were also found in the blood and in the miliary tubercles in lungs, liver, etc.

**Literature.**—*General Works*—BAYLE, *Recherches sur la phthisie pulmonaire*, 1810; LÆNNÉC, *Traité de l'auscult. med.*, etc., 1819; WALDENBURG, *Tuberkulose*, etc., 1869; VIRCHOW, *Geschwülste*, vol. ii., 1864; Discussion on pulmonary phthisis in relation to tubercle of lung, *Path. Soc. trans.*, 1873. *Inoculation and Tubercle bacillus*—VILLEMIN, *Du tubercle au point de vue de son siège*, etc., 1862; and *Etudes sur la tuberculose*, 1868; LANGANS, *Die Uebertragung der Tuberk.*, 1868; KLEBS, *Virch. Archiv*, vols. xlv., xlix., etc.; TAPPEINER, *Virch. Archiv*, vols. lxxiv. and lxxxii.; COHNHEIM, *Die Tuberkulose vom Standpunkt der Infectionslehre*, 1880; SANDERSON, *Researches on artificial tuberculosis*, 1869; KOCH, *The Etiology of tuberculosis*, *Syd. Soc. trans.*, 1886; BAUMGARTEN, Many papers, and *Tuberkel und Tuberkulose*, 1885; WEICHELBAUM, *Deutsche med. Wochenschr.*, 1884; JOHNE, *Die Geschichte d. Tuberkulose*, 1883; KARG (*Post-mortem tubercles*), *Centralbl. f. Chir.*, 1885, No. 32. *Anatomy and mode of extension*—VIRCHOW, *Geschwülste*, vol. ii., 1865; RINDFLEISCH, *Path. Histology*, *Syd. Soc. transl.*, 1872; LANGANS, *Virch. Archiv*, vol. xlii.; SCHÜPPEL, *Lymphdrüsentuberkulose*, 1871; BUHL, *Lungenentzündung*, etc., 1872; KÖSTER, *Virch. Arch.* vol. xlviii.; FRIEDLÄNDER, *Volkmann's Klin. Vort.*, No. 64; GÜTERBOCK (*Lupus*), *Virch. Arch.* vol. liii.; WEIGERT, *Virch. Arch.*, vol. lxxvii., and *Deutsch. Med. Woch.*, 1883 and 1885; BAUMGARTEN, *Tuberkel und Tuberkulose*, 1885; HAMILTON, *On the path. of bronchitis*, etc., 1883.

### TUBERCULOSIS IN ANIMALS.

It has been already mentioned that tuberculosis has been communicated to a large number of animals by inoculation, and that probably all warm-blooded animals are susceptible. As a spontaneous disease it occurs in many domestic animals. In these the lesions observed are virtually identical with those in man, consisting in tubercles formed by giant-cells and epithelioid cells with varying proportions of leucocytes. There are, however, certain minor variations in structure in certain animals. Bacilli presenting all the characters of the tubercular bacillus have been found in the affected structures. For practical purposes the most important form is that which occurs in cattle.



**Bovine tuberculosis.**—This is a common disease in cattle. The structures most frequently affected are the serous membranes—the pleura and peritoneum especially—and the lungs, but the disease often extends to lymphatic glands, alimentary canal, liver, spleen, nervous system, etc. In the **serous membranes** the tubercles are aggregated into considerable nodules, frequently as large as lentils. They are attached to the surface of the membrane or else supported on villous projections from the surface. (See Fig. 56.) There may be massive projections of these nodules

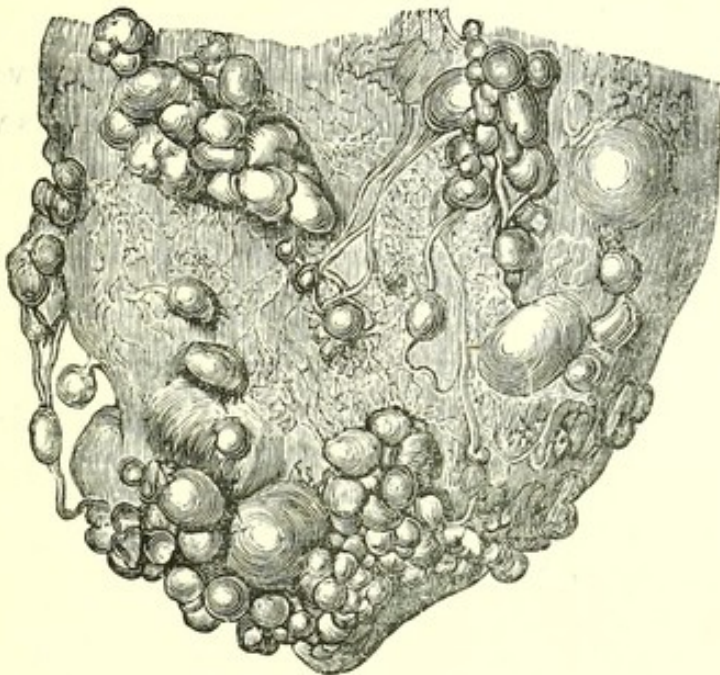


Fig. 56.—A piece of lung in bovine tuberculosis. On surface of lung many rounded tumours are seen, some pendulous. (VIRCHOW.)

hanging from the pleura or peritoneum. From the size of the nodules and their white appearance they have been compared to pearls, and the name **Perlsucht** has been used in Germany for the serous form of the disease. In **internal organs**, we have also comparatively large rounded nodules, and sometimes considerable infiltrations.

The nodules in bovine tuberculosis are peculiarly prone to **calcification**. The calcification is probably always preceded by caseation, but the latter may be so rapidly overtaken by the calcareous infiltration as to be masked. According to some authors, tuberculosis may be communicated to calves and to man by the milk of tuberculous cows. It is said also that bacilli have been detected in the milk.

**Tuberculosis in horses** somewhat resembles that in cattle, there being usually pearl nodules on the peritoneum. There may be also numerous nodules in internal organs, and these are said to be more like those in human miliary tuberculosis. Tuberculosis in horses is said to be sometimes mistaken for sarcoma; it is not of common occurrence.

**In swine** tuberculosis is a frequent disease. It resembles bovine tuberculosis in its general characters as well as in its tendency to calcification.

Tuberculosis is very common in **monkeys**, in which animals it usually begins in the lungs, but is apt to extend to other organs in the form of chronic general tuberculosis, so as to form numerous comparatively large foci, which usually break down so as to form cavities.



**Literature.**—For an account of the various forms of tuberculosis in animals see KOCH, *Etiology of Tuberculosis*, Syd. Soc. transl., 1886; and JOHNE, in Birch-Hirschfeld's *Lehrbuch d. path. Anat.*, where a full statement of the literature is given; see also BAUMGARTEN, l. c.; CREIGHTON, *Bovine tuberc. in man*, 1881.

### LEPROSY OR LEPRA.

This disease was at one time spread over the whole of Europe, but is now limited to certain localities in Norway, Russia, Iceland, and the coast of the Mediterranean. It is still somewhat prevalent in Asia, especially India, Japan, and China; in Africa, where it is very prevalent in Egypt, Abyssinia and the Islands of the East Coast; and in America, especially in the West India Islands and Mexico. (See Hirsch, "*Geograph. Path.*," vol. ii., p. 1.)

**Causation.**—The disease is generally regarded as hereditary, but a considerable number of cases of contagion have been observed.

The question of contagion as a cause of leprosy has been till lately an undecided one. But recently several cases of communication to Europeans, evidently by contagion, have been published, one in which it was communicated by vaccination. A final demonstration of contagion has been furnished by the inoculation of a condemned criminal at Honolulu by Dr. Arning. The inoculation was successful, but the period of incubation was very prolonged. It is probable that the incubation may extend over several years, and if this be so, it will be frequently very difficult to trace the source of contagion.

The leprosy bacillus somewhat resembles the tubercular bacillus, and it is stained by similar methods.



J.L.S.

Fig. 57.—Leprosy bacillus. Those in groups are inside cells.

It has been cultivated on solidified blood serum, but attempts at inoculation of the cultures, both in animals and man, have been unsuccessful, although direct inoculation has succeeded in the case referred to above. The bacillus is present in quite enormous numbers in the lesions in the skin in leprosy. It presents itself in the interior of round cells (the lepra cells of Virchow) which are considerably larger than leucocytes, but are probably these enlarged by the bacilli. (See Fig. 57.)

Some have denied that the bacilli are in cells, and assert that the rounded aggregations are simply free groups. The bacilli are so numerous as to suggest that the enlargements may be in part due to the actual bulk of the bacilli.



**Character of the lesions.**—The disease occurs in two forms which are designated *Lepra tuberculosa* and *Lepra anæsthetica*. In the first the newformation has its seat in the skin or mucous membrane; in the second it is the nerves which are affected. The tubercular form is sometimes called *Elephantiasis græcorum*.

When it attacks the skin, it is mostly the face and hands which are affected, and the legs if they are exposed to the weather. There appear large or smaller swellings, at first red or bluish in colour, which become firmer and harder. These tubercles may reach the size of a hazel nut or a walnut. They consist of granulation tissue in which cells of the size of leucocytes are most abundant. As in other granulation tumours, we often have ulceration, or, as in the case of lupus, there may be cicatrization without ulceration. By the formation of the swellings and cicatrization, great deformities frequently result, so that the patients have often a peculiarly hideous appearance, the face being knobbed and gnarled (Fig. 58).

In the anæsthetic form the nerve stems become the seat of granulation tissue, forming spindle-shaped swellings, sometimes of considerable length. The granulation tissue here is in the interstitial connective tissue, so that the nerve-fibres are separated and compressed. The newformation sometimes, but rarely, extends to the membranes of the spinal cord or brain.

Besides the anæsthesia there are usually other evidences of interference with the nerves, mostly connected with the nutrition of the parts, namely, vesicular eruptions of the skin, alterations in the pigmentation, deep ulceration, not infrequently leading to separation of the fingers or toes. No bacilli have been found in these peripheral lesions, they are simply the result of the interference with the nerves.

In leprosy there are sometimes tumours formed in internal organs, but this is very rare.



Fig. 58.—Leprosy. The face shows nodular swellings, especially on nose, eyebrows, lips, chin, and ears. Patient had also a large ulcer on right leg, and a small one on left. (VIRCHOW.)



**Literature.**—For a historical account see HIRSCH, Handbook of hist. and geograph. path., Syd. Soc. transl., vol. ii; see also VIRCHOW, Geschwülste, vol. ii.; GULL, Guy's Hosp. Rep., 1859; CARTER, Med. Chir. trans., lvi., and Path. trans., xxvii.; HANSEN, Virch. Arch., vol. lxxix., NEISSER, Ziemssen's Handb. d. Spec. Path. vol. xiv., and Virch. Arch., vols. lxxxiv. and ciii.; also NEISSER and CORNIL et SUCHARD, in translated essays, New Syd. Soc., 1886. *Contagion*—MONRO, Etiology and Hist. of leprosy; BESNIER, Acad. de Méd., Oct. 11, '87; Brit. Med. Jour., 1887, i. 1269, ii. 799, 1055, and 1119, and 1888, ii. 1171.

#### ELEPHANTIASIS ARABUM.

This disease, which is called elephantiasis arabum to distinguish it from *E. græcorum*, which is true leprosy, is often simply designated Elephantiasis, sometimes also Pachydermia.

**Causation.**—The disease is endemic in certain localities in India, China, Egypt, Arabia, the islands of the Pacific, etc., chiefly in places within a short distance of the sea. It obviously depends on some morbid poison, but the nature of it has not been discovered. The disease begins in an acute inflammation of the skin, accompanied by fever, and somewhat resembling erysipelas. These attacks pass off and recur, the disease ultimately subsiding into a more chronic condition. These are conditions which can only be produced by a morbid poison acting on the tissues locally and probably sending off products into the blood so as to induce fever. The fact that no secondary lesions occur at a distance is an indication that the virus itself does not reach the blood. The disease attacks chiefly the native races, but no race is exempt from it.

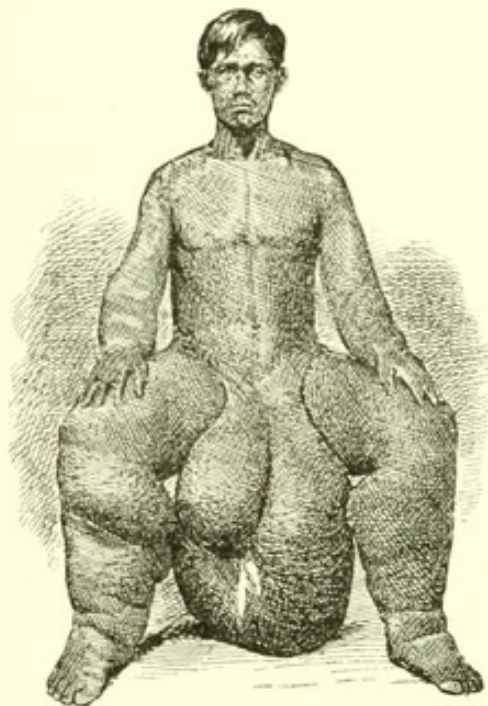


Fig. 59. — Elephantiasis affecting scrotum, legs and arms. (After photograph by TURNER.)

**Character of the lesion.**—The lesion consists of an enormous hypertrophy of the tissues of the skin. The cutis vera is thickened, the papillæ are enlarged, and the epidermis is thickened. In the earlier periods the new-formed tissue is somewhat cellular, and throughout it is succulent and oedematous. The disease occurs chiefly in the lower extremities and external genital organs, but sometimes attacks the upper extremities. In the lower extremities it produces an extraordinary irregular thickening of the skin, which has a folded and bagged appearance, like that of the legs of an elephant. (See Fig. 59.) In the genital organs it produces tumours



which sometimes grow to a massive size; tumours have been excised which weighed over 100 pounds, and which hung down so as to reach below the knees. (See Fig. 59.) Sometimes the tumour is not entirely made up of hypertrophied skin, but a proper isolated tumour is found internally. This was so in a case observed by the author. In the legs the newformation may extend inwards to the fasciæ, the intermuscular tissue, periosteum, and bone. The accompanying Fig. 60 shows how the bone may be the seat of newformation.

Sometimes there is great dilatation of the lymphatic vessels. It is asserted that the disease may develop from so-called lymph-serotum, and Manson has inferred that elephantiasis, like lymph-serotum, depends on the *filaria sanguinis*. (See under Parasites.) The author does not regard this inference as warranted.

**Literature.**—VIRCHOW, *Geschwülste*, vol. i.; ESMARCH und KULENKAMPFF, *Die elephantiasischen Formen*, 1885; CARTER *On Leprosy and Elephantiasis*, 1874; TURNER, *Amputation of scrotum for Elephantiasis*, *Glasg. Med. Jour.*, 1882, vol. i. (with photographs).

#### GLANDERS.

This disease is met with chiefly in horses, but is occasionally communicated to man. The name **Farcy** is sometimes given to the disease when the skin and lymphatic system are specially engaged.

**Causation.**—The disease has been shown by Löffler and Schütz to depend on a bacillus which resembles the tubercular bacillus in form, but does not contain spores and requires different methods of staining. (See in section on Bacteriology.)

**Character of lesion.**—The disease in horses manifests itself first in the formation of swellings of the mucous membrane or skin, consisting of granulation tissue. The nasal mucous membrane is generally first attacked, and there is either a diffuse infiltration of it or else a more localized series of swellings like those of lupus. From this seat the

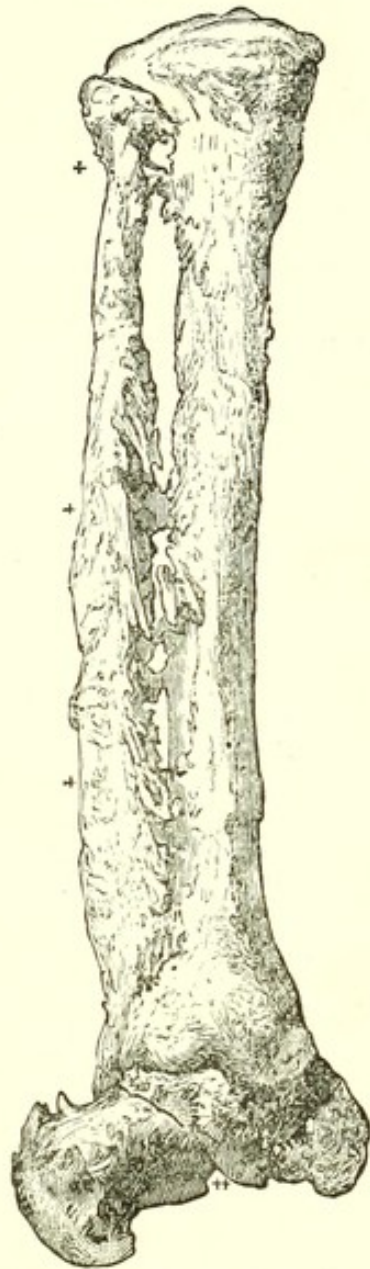


Fig. 60.—Hyperostosis and synostosis of the bones of the leg and foot in elephantiasis. The bones present everywhere flat or pointed projections. At + + + the tibia and fibula are united, as they are also above the ankle-joint. The astragalus and os calcis are also united. (VIRCHOW.)



disease spreads to neighbouring lymphatic glands, also along the mucous membrane to the lungs and intestinal tract. The skin also becomes the seat of lesions.

The granulation tissue here, even more than in the other infective tumours, tends to break down, so that, as a rule, ulcers soon form, which present a great tendency to spread. These ulcers arise by what is virtually a suppurative process, and if the tumours are situated deep in the skin or mucous membrane there may be actual abscesses, which, by bursting, form ulcers. If the ulcers heal, cicatrices are formed in the usual way.

The manifestations in internal parts are largely inflammatory in character. The lymphatic glands of the neck are the seat of inflammatory swelling. In the lungs there are nodules either consisting of granulation tissue with caeous central parts, or more distinctly pneumonic. There are also in the mucous membrane of the intestinal canal nodules which are sometimes more solid, at other times breaking down into pus.

The above description applies to the acute cases, which are the commoner, but the disease is sometimes chronic in its course. Chronic cases have usually the nodules in the skin, and there are secondary tumours in the muscles, etc.

In man, glanders, which occurs occasionally in persons engaged about horses, is usually an acute, rapidly fatal disease. There are nodules and ulcers in the mucous membranes and the skin, and frequently phlegmonous inflammation of the skin with abscesses among the muscles, etc. In fact, in man the disease takes the inflammatory character more distinctly than in the horse. We may have in man also nodules, ulcers, and abscesses in internal organs. Sometimes the abscesses in lungs, kidneys, etc., cause the disease to resemble pyæmia. At other times the juices from the ulcers in the nose or mouth being inhaled, may give rise to lesions like those of phthisis pulmonalis.

Chronic glanders in man is characterized by ulcers in the mucous membranes of fauces, bronchi, etc., and in the skin. There may be caseating nodules in internal organs.

**Literature.**—VIRCHOW, *Geschwülste*, vol. ii.; BOLLINGER, *Ziemssen's Handb.*, vol. iii.; LÖFFLER and SCHÜTZ, *Deutsche Med. Wochenschr.*, Dec. 1882.

#### ACTINOMYCOSIS.

This disease is prevalent in cattle in whom it was first recognized as parasitic by Bollinger. It has been observed in man in a considerable number of cases.



**Causation.**—As the name implies, the disease is due to a microbe, whose botanical position and life-history have not been fully made out. It consists of short threads, arranged in a radiating fashion many springing from a common centre. (See Fig. 61.) The threads are usually clubbed at their outer extremities, but they are not always so. They are united in the centre of the clump by a matted mass of fibres. The little masses thus form peculiar radiating bodies of a globular or oval form, having a somewhat glandular aspect. From the radiating arrangement the fungus is called the Ray-fungus or *Actinomyces* (*ἀκτίς*, a ray). The individual clumps are of very small size, not visible to the naked eye, but they may be united into larger masses as the heads in a cauliflower. These larger masses appear as small yellow sulphur-coloured grains in the discharges and newformations, giving the characteristic indications of the disease.

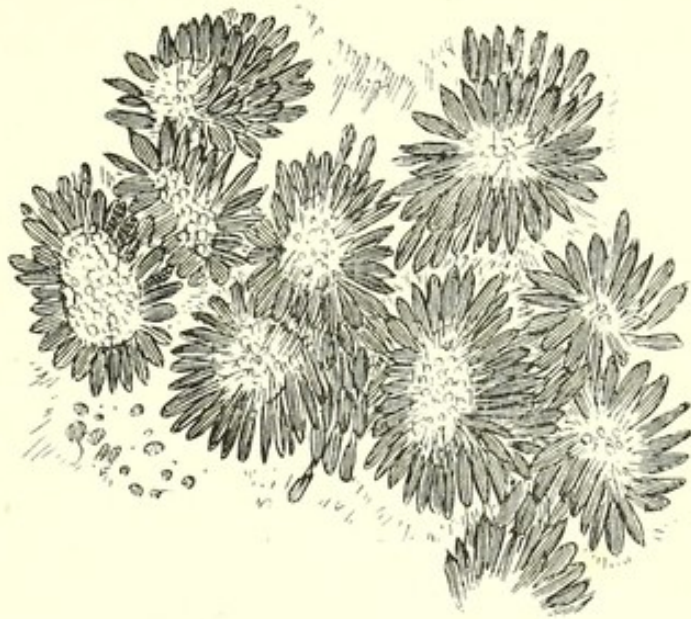


Fig. 61.—*Actinomyces*. A group of radiating heads.

The fungus has not been cultivated outside the body with unequivocal success, and even the direct inoculation of portions of newformation containing the fungus into the calf, have been doubtfully successful. The disease is not, apparently, contagious, and it seems probable that the fungus has some habitat outside the animal body. The occurrence of an epidemic in a locality where the inhabitants had eaten rye grown on land newly reclaimed from the sea suggests some such habitat. (Jensen.) This is supported by the fact that the disease occurs almost alone in herbivora and omnivora.

**Character of lesion.**—Whether in animals or man each little fungus head constitutes a centre of irritation, and there forms around it a little mass of granulation tissue which may contain giant-cells, in fact a structure closely resembling a tubercle, but with a different micro-organism in the centre. Besides these specific centres there is a more general formation of granulation tissue. There is no tendency to caseation, but rather to formation of pus on the one hand or cicatricial tissue on the other, but the new-formed tissue has more tendency to retain its undeveloped form than granulation tissue generally. In this way bulky masses, which have been taken for sarcomatous tumours, have been formed



about the jaws of cattle. In man there is more tendency to suppuration than in cattle.

The locality of the tumour and the seat of entrance of the fungus have been studied in the recorded cases. In cattle it has been met with chiefly in connection with the jaw bones, the tongue, and face. In man the seat of entrance has been distinguished as, the jaws, the respiratory tract, or the alimentary canal. Israel divides the cases according to this arrangement, and makes a fourth group in which the place of entrance is not determined.

In **Actinomycosis of the face and neck**, the fungus has obtained entrance by breach of surface in the mouth, usually in connection with carious teeth, so that the commencement is usually in connection with the jaw. There is a formation of granulation tissue around the jaw, usually the inferior maxilla, extending to parts around. The process is a chronic one, and by degrees the lesion extends to the surface. Abscesses form which discharge and leave fistulous openings. Sometimes the newformation extends to the vertebræ or base of the skull, especially when the upper jaw has been the starting point.

**The respiratory organs** may be the primary seat. There may be a superficial actinomycosis, producing simply a suppurative bronchitis. More usually the fungus is in the substance of the lung, giving rise to chronic condensations and indurations, in isolated patches. These enlarge and coalesce, and they often break down so as to form cavities. The process is a chronic one and somewhat resembles that in phthisis pulmonalis, but differs in respect that it does not usually attack the apices of the lungs. The disease is apt to involve the pleura and the neighbouring pericardium. It not infrequently extends to the bodies of the vertebræ, causing them to become carious. There is here a slow inflammatory process with exuberant granulations and suppuration. The abscess may reach the surface by a course similar to that of a tubercular abscess of bone.

**The alimentary canal** is more rarely the primary seat of actinomycosis. There are nodules and ulcers formed and the disease extends to the peritoneum, leading to the formation of granulation tissue and abscesses in the cavity of the peritoneum. There may be perforation of the bladder, intestine, or abdominal wall.

In some cases the fungus gets into the blood and is disseminated in various organs. In that case there are abscesses produced and the disease is like a **chronic pyæmia**.

**Literature.**—BOLLINGER, *Centralbl. f. d. med. Wissensch*, 1877, No. 27; PONFICK, *Die Actinomycose des Menschen*, 1882; ISRAEL, *Actinomycosis in man*, *Syd. Soc.*, 1886; SHATTOCK, *Path. trans.*, vol. xxxvi.; ACLAND, *ditto*, vol. xxxvii.



## MALIGNANT LYMPHOMA OR HODGKIN'S DISEASE.

The exact relations of this disease, or group of diseases, are somewhat obscure. In many respects it presents the characters of the infective tumours, and on the other hand it is related to leukæmia, this latter disease sometimes presenting most of the organic lesions of Hodgkin's disease. By some leukæmia is also included amongst the specific new-formations. The uncertainty as to the nature of the disease may be inferred from an enumeration of the names applied to it. Besides the above we have **Adenia**, **Pseudo-leukæmia**, **Lymphosarcoma**, **Malignant lymphadenoma**, **Malignant aleukæmic lymphadenoma**, and **Anæmia lymphatica**.

The disease begins in a particular group of lymphatic glands, usually those of the neck, but also frequently the mediastinal, bronchial, or mesenteric. The glands enlarge and the new-formed tissue is essentially like that of the normal gland, consisting of round cells in a reticulum. The disease shows local malignancy, extending not only from gland to gland, but infiltrating the structures around. In this way we may have bulky tumours produced, which are composed of glands compacted together and incorporated with the surrounding tissues. In its local advance the tumour will often mould itself on neighbouring structures. Thus, when originating in the mediastinal or bronchial glands it may incorporate the walls of the bronchi, the walls of the venæ cavæ, and other large veins (first causing obstruction by thrombosis), and even the pericardium and wall of the heart (as in a case observed by the author). Again, from the mesentery it may extend to the intestine, and the author has seen a considerable length of the small intestine replaced by tumour tissue, the general form, including the valvulæ conniventes, being repeated in an exaggerated form in tumour tissue. The local tumour not infrequently presents considerable induration due to development of fibrous tissue, and it may also present caseation.

Besides this local malignancy there is usually extension by metastasis to distant organs. The organ most frequently affected is the spleen, where round-celled tissue forms around the arteries. This may be a simple increase of the lymphatic tissue normally surrounding the splenic arteries. The tumour tissue is thus mixed with the tissue of the spleen, and the greatly enlarged organ appears infiltrated with a whitish substance which has been compared to lumps of suet in a pudding. The kidneys also not infrequently take part in the process. They are the seat of extensive infiltrations similar to those sometimes seen in leukæmia. The liver, follicles of the intestine, and tonsils are also not



infrequently affected. The liver may show an infiltration of the capsule of Glisson with round cells, just as in leukæmia. The enlargement of the solitary follicles of the intestine and of the tonsils is like the lesion in the spleen, an increase of existing lymphatic tissue.

The condition of the spleen attracted the attention of Hodgkin, whose original paper is entitled, "On some morbid appearances of the absorbent glands and spleen." It is probably always enlarged, and is commonly the seat of secondary tumours. Hodgkin's disease may give rise to amyloid degeneration, as in a case observed by the author. In this case a sago spleen was the seat of secondary lymphatic newformations. The newformation occupied the same structures as the amyloid disease, and it was interesting to observe how the round-cell infiltration caused absorption of the amyloid material, and how the arteries, whose walls were amyloid, recovered their structure under the influence of the newformation.

The blood in Hodgkin's disease is in the condition of anæmia, and the symptoms, apart from those due to the local enlargements, are those of progressive anæmia.

**Literature.**—HODGKIN, *Med. Chir. trans.*, 1832; WILKS (who suggested the name Hodgkin's Disease, and has chiefly brought it into prominence), *Guy's Hosp. Reports*, 3rd series, vols. ii. and xi., also cases in *Path. trans.*; MURCHISON, *Path. trans.*, xxi. 372; also papers by WILKS, GREENFIELD, and others in *Path. trans.*, xxix., 1878; HILTON FAGGE, *Medicine*, 1886, vol. ii. 334.



## SECTION IX.

## TUMOURS OR MORBID GROWTHS.

DEFINITION. STRUCTURE, *typical or atypical*. CAUSATION, *Cohnheim's theory of origin; inheritance; effect of injuries, etc.; parasitic microbes not a cause of tumours proper; influence of age*. GROWTH AND EXTENSION, *typical or atypical; local malignancy; metastasis and generalization of tumours; occasional malignancy of typical tumours*. CLASSIFICATION AND NOMENCLATURE.

## INTRODUCTION.

A TUMOUR means literally a swelling, and formerly any swelling was called a tumour. *Tumor* for example, is often named as one of the cardinal signs of inflammation, and it is still customary to speak of the inflammatory tumour. But the modern use of the word is limited to a class of newformations of which it is difficult to give a strict definition.

It may be said that tumours are pieces of tissue which have a life and growth of their own irrespective of the needs of the organism, and of the general local conditions around them. The structure of a tumour does not differ in its elements from that of normal tissues. It is nourished by the same blood, its blood-vessels are continuous with those of surrounding parts, its nerves are connected with neighbouring nerve-stems, and its tissue, if not always precisely the same as that near it, is frequently identical. But it grows independently and without apparent object. Thus a fatty tumour goes on increasing in size irrespective of the adipose tissue in which it has its seat, and the person may be reduced to the greatest emaciation, most of the ordinary fat being absorbed, while little or no impression is made upon the fatty tumour.

**Structure of tumours.**—While all tumours in their ultimate structure are analogous to the normal tissues, yet some of them vary considerably from the type of tissue to which they belong. Hence, it is possible to distinguish tumours whose structure is **Typical**, and others whose structure is **Atypical**. In the former, the details, including size and shape of cells, abundance of cells, relation to intercellular substance, and so on, are strictly like those of the corresponding normal tissue. In the latter the structure varies from the normal, chiefly in respect that the



cells become much more numerous, and that the intercellular substance passes into the background. We shall see afterwards that in their mode of growth, tumours show somewhat singular differences, some being typical and some atypical, and that in general the variations in mode of growth correspond with those in structure.

The terms **Homologous** and **Heterologous** were at one time employed to express the idea that the tissue of the tumour may, or may not, correspond with that of the tissues of the body, the heterologous tumours being looked upon as altogether different in structure and foreign to the body. In the modern use of the words a homologous tumour is one which exists in tissue of its own kind, as an osseous tumour growing from bone. A heterologous tumour on the other hand is one which is present in a situation where no normal tissue of that kind exists, as, for example, an osseous tumour in the brain.

**Causation of tumours.**—It is implied in the definition given above that the causation of tumours is exceedingly obscure. For the most part, without any apparent stimulus, they begin to grow, and go on without control.

**An explanation of the origin** of tumours has recently been suggested by **Cohnheim**, which may apply to some tumours but certainly does not account for all. He suggests that the primordial tumour is a piece of tissue, which in the process of development and growth has been, as it were, left over, has retained its embryonic powers of growth, and is not subject to the general laws which control the growth of the tissues.

In enforcing his argument Cohnheim points out that pieces of living tissue, especially if from young animals, will bear transplantation, and, if placed in a favourable situation, may grow to considerable dimensions. Thus, the spur of the cock, transplanted to the more vascular comb, will sometimes form a considerable tumour. Then Leopold has transplanted living cartilage into the anterior chamber of the eye, where it acquires vascular connections with the iris. If cartilage from an early fœtus be used it will grow so as to form a tumour of some size. The earlier the period of development of the fœtus the more likely is there to be a vigorous growth.

Certain facts in the pathology of tumours themselves lend some support to this view. Thus cartilaginous tumours not infrequently originate in bones, as if portions of the original fœtal cartilage had been left over and had afterwards grown. Then mucous-tissue tumours sometimes occur in the subcutaneous tissue, a situation in which this form of tissue exists in the embryo, the tissue being itself an embryonic form of tissue. Then also, tumours not infrequently develop from congenital moles or soft warts. These little outgrowths have commonly a rudimentary or embryonic structure (see section on Skin Affections), but they may lie quiescent throughout life, and only occasionally start into vigorous growth.

While Cohnheim's theory may apply to some tumours, more particu-



larly to the typical ones, it does not explain a large proportion of cases, and especially the cancers. In these the existing normal structures, usually in an increasing area, give rise to the tumour. There must be here some stimulus acting on the epithelial structures especially.

**Inheritance** is generally believed to play an important part in the causation of tumours. The embryonic tissue presupposed in Cohnheim's theory may, like an ordinary malformation, be transmitted by inheritance, or the predisposition to the abnormal growth in cancer may be so transmitted.

The whole subject of inheritance of tumours stands in need of elucidation. It is asserted on the one hand that "we cannot overestimate the importance of inheritance in the origination of cancer" (Paget), while on the other hand it is asserted that when the families of non-cancerous persons are compared with those of cancerous, there is little or no appreciable difference in the number of cancerous relatives (Snow, Cripps). We have already seen that so far as we are able to go on demonstrable facts, inheritance deals with local peculiarities of structure. It is stated by Paget that in the transmission of cancer, the disease often passes from one situation to another, as from the breast to the stomach or uterus. This author would even seem to indicate that in transmission a cancer may become a sarcoma. Such statements run so counter to the known facts of inheritance that they throw considerable doubt on the whole subject. The strongest arguments in favour of inheritance are based on the observation of individual families, such as those recorded by Paget, Broca, and others.

**Injuries, Irritations, and Chronic inflammations** play an important part in determining the occurrence of tumours. Thus, fractures of bones are sometimes the starting points of cartilaginous or bony tumours or of sarcomas. Cancers of the mamma are often referred to blows on the breast. The seat of election of cancers is often determined by the local conditions of exposure to irritation by friction or otherwise. For example, cancer of the lip and tongue are referred to irritation by a tobacco pipe or carious tooth; cancer of the stomach is often ascribed to prolonged irritation from alteration in the gastric juice. The frequency of cancer at the narrow parts of mucous canals, such as the pyloric orifice of the stomach, the ileo-cæcal valve, the os uteri, has also been ascribed to their special exposure to irritation.

**Parasitic microbes** have been suspected as forming the determining causes of tumours, but without any definite proof. We have seen that specific microbes are the exciting causes of the infective tumours, and the analogy between their mode of growth and that of malignant tumours, such as sarcomas and cancers, has led some to suppose a similar causation. There are some malignant tumours which closely resemble the infective tumours, and probably belong actually to this class. The condition designated Hodgkin's disease, for instance, is variously regarded as a malignant or as an infective tumour. An important dis-



inction must be drawn, however, between the infective characters of the tumours due to parasitic micro-organisms, and of tumours proper. In the former case, the infectiveness is due to the extension of a parasite, which wherever carried induces an effect comparable to inflammation, but with certain specific characters. The lesion is the result of irritation of the tissue to which the parasite is applied. In the case of a tumour, on the other hand, there is everywhere a proper newformation of tissue, which of its kind is completely organized, and not merely an effect of irritation. Even when produced at a distance from the primary tumour, the secondary one is, in the details of its structure, an exact reproduction of the tissue of the original growth. Thus, while cancers differ very greatly in structure according to their seat of origin, they always, so to speak, breed true; the secondary tumours, however various their seats, having the same structure as the primary one. A striking illustration of this argument is afforded by a case recorded by the author, in which an adenoma of the thyroid gland had led to the formation of secondary tumours in the bones of the skull. The primary tumour had the structure of the normal thyroid gland, and the tumours in the skull had the same structure. It is scarcely conceivable that any parasite could produce such results.

**Age** is a factor in the etiology of tumours. The frequency of tumours increases in proportion to the persons living up to the age of seventy, and is greatly increased from thirty upwards. This latter fact is due to the prevalence of cancer in the more advanced years of life. Cancer is a disease of decadence.

It seems strange that a disease characterized by undue activity of growth of certain elements should occur especially when the body generally is losing in vigour. An explanation of this was suggested by Thiersch, to the effect that, as cancer consists essentially in an exaggerated growth of epithelium, which invades the neighbouring structures, especially the connective tissue, the cause may lie rather in a falling away of the resistance of the other tissues than an extra vigour of the epithelium. The occurrence of cancers in old cicatrices, which consist of a very imperfect connective tissue, would lend some force to this view.

**Growth and extension of tumours.**—Most tumours enlarge by a newformation of tissue within themselves, just as the normal tissues increase during the period of growth. Their growth is **typical**. The tissue of the tumour is sometimes continuous with that of surrounding parts, sometimes it is separated by a capsule composed of connective tissue; sometimes it begins by being continuous and afterwards gets separated. In any case the ordinary tumour simply grows by newformation of its own tissue. It often produces effects on neighbouring parts, by pressure especially, but apart from these merely mechanical effects, it does not



prejudice neighbouring structures. Tumours possessing these characters are usually called **Simple** or **Innocent** growths.

But then there are tumours of which this is not true. They have a tendency to grow into and infiltrate neighbouring tissues, presenting characters of what may be designated **Local malignancy**: their mode of growth is **atypical**. This is peculiarly the case with sarcomas and cancers. In the case of sarcomas the tumour seems to penetrate into and develop as it were, on the mould of the existing tissue—apparently very much in the style that a thrombus or a piece of catgut in the tissue is replaced by granulation-tissue which moulds itself on it. In the case of cancers, on the other hand, their tissue, which is epithelial in character, penetrates among the tissues, largely destroying them and producing inflammatory disturbances.

Besides this local malignancy the same classes of tumours frequently show a still more atypical growth by extending beyond their local seat. This character is expressed in the term **Metastasis**, which is another feature of malignancy. In this case secondary growths spring up in parts removed from the primary tumour, and doubtless something is carried from the original tumour to the remote part. In the case of cancers the epithelial processes penetrating into the tissues readily find their way into the lymphatic spaces, and portions may be carried thence to lymphatic glands.

Besides this metastasis by the lymphatic system, there is sometimes a more distant metastasis by the blood, in which case the tumour is transplanted to various parts of the body, and the term **Generalization** is used. This occurs usually after the lymphatic glands have been involved, but in sarcomas it is usually direct, the tumour penetrating through the walls of the vessels. It is mostly the veins which are thus opened into, and the first extension is therefore to the lungs, but the pulmonary capillaries may be in part traversed, and implantation occur by the systemic arteries.

As to the nature of the material which passes from the tumour, it must be either something dissolved in the juices, or else solid particles of some kind. Looking to a secondary cancerous infection of the peritoneum, we can hardly escape the conclusion that it is a finely divided solid. The secondary tumours are not regularly distributed over the peritoneal surface, but occur here and there or in groups, just as if solid particles had been carried and produced their effects where they got leave to lie. Then also it must be solid particles which are arrested by the lymphatic glands and give rise to the secondary tumours there. The probability is that the actual cells of the tumour are carried off and deposited at a distance. Perhaps it is not the fully developed cells but small rudimentary ones.



In this connection it is important to note that the secondary tumour for the most part exactly reproduces the tissue of the original one, even to the smallest details, and it is natural under these circumstances to believe that pieces of the original tumour are actually transported.

**Occasional malignancy of typical tumours.**—It has already been stated that sarcomas and cancers are the tumours which regularly present a malignant tendency, but on rare occasions other tumours also do so. **Cartilaginous tumours** are not infrequently **malignant**. Next to them mucous-tissue tumours most frequently become malignant, but even fibrous tumours have been observed to do so, and lately Cohnheim and others have recorded cases of colloid goitre in which secondary tumours occurred. So that malignancy is not confined to sarcomas and cancers. On the other hand tumours having the structure of sarcomas may remain local to the end.

It has been usual to believe that malignancy is dependent on peculiarities of the tumour itself, and it is true that when a simple tumour, as sometimes happens, assumes malignant characters, it usually assumes the structure of a sarcoma or cancer. The view already referred to above as having originated with Thiersch has been amplified to account for the difference between a simple and a malignant tumour. In the case of a simple tumour the normal tissues are able to prevent the tumour-tissue from penetrating into them, but in the case of malignant tumours the tissues have become weakened and are unable to form a barrier to their extension. When a simple tumour becomes sarcomatous, it is because the tissues around have acquired a peculiar weakness. When pieces of a tumour are transported to a distance, a struggle, as it were, occurs between the tendency of the tissue of the tumour to grow, and that of the normal tissues to prevent its growth. In the case of a simple tumour the tissues assume the upper hand, and cause the absorption of the tumour-tissue. In the case of malignant tumours, however, the tissues are too weak to accomplish this, and the secondary tumour develops. This theory is not sufficient to explain all the facts, and we must suppose as well a special power of growth in the tumour. No doubt the tissues differ very greatly in their power of restraining the growth of secondary tumours. In the liver, for instance, cancers grow very freely, and attain large dimensions, but looking to a cancerous liver, it is difficult to believe that there is not a special activity in the growing tissue of the tumours.

**Secondary changes in tumours.**—Tumours are exposed to the same pathological processes as normal tissues, and in a higher degree. Thus we meet with fatty degeneration, which is common, especially in quickly growing tumours; calcareous infiltration in structures which are obsolete; hæmorrhages, principally in superficial rapidly growing tumours, where the blood-vessels are ill-formed; and necrosis mostly in tumours near the surface and thus exposed to mechanical irritation, the consequences of such necrosis being ulceration with, it may be, suppuration, hæmorrhage, decomposition.

Conditions like those last named will seriously affect the organism as



a whole, and they will occur most frequently in the case of malignant tumours, which grow quickly, and rapidly come to the surface. Malignant tumours also affect the organism as a whole by the readiness with which they extend to or produce secondary tumours in important organs, and secondary cancers frequently also induce inflammations as in the case of the peritoneum or pleura.

It is clear, therefore, that malignant tumours, especially by bleeding, by ulcerating and sloughing, by invading important parts, by producing inflammations, etc., have a tendency to deteriorate the system, producing anæmia and general weakness. If growing quickly they also tend to emaciate by using up the nutritive material of the body. A simple tumour, if it happens to be at the surface and exposed to mechanical violence, may also ulcerate and produce serious constitutional results, but it will be mainly in the case of malignant tumours that these effects, will follow. To these conditions of the body as a whole the name **Cachexia** is often given. Sometimes also the term **Dyscrasia** or **Dia-thesis** is used with the meaning that there is some peculiar condition of the system preceding and inducing the formation of malignant tumours, but of this there is no evidence, and the changes in the general condition are always secondary.

**Classification and nomenclature of tumours.**—Tumours have sometimes been classified and named according to their clinical characters, whether innocent or malignant, the term cancer being used to include malignant growths in general. A true system of classification will take into account the origin, structure, and mode of growth of tumours, and such a classification will be found ultimately to correspond with clinical facts.

In classifying tumours according to their origin we may assume that they take origin in tissues of their own nature. This is at least admitted in the case of simple tumours; a fatty tumour, for example, originates in adipose tissue. In the case of cancers, whose more important elements are epithelial, Virchow asserted that they originated from connective tissue. This was opposed by Thiersch, who showed that in the case of epithelial cancer the cells originate from existing epithelium. This view has been amplified by Waldeyer, who, in a very elaborate series of investigations, has shown that in their very various seats, cancers are in their origin connected with the epithelial structures. This has led to a reference to embryology in the classification of tumours. An attempt has been made to distinguish those arising from the various layers of the embryo, the tumours of mesoblastic being separated from those of hypoblastic or epiblastic origin. It is very difficult to carry out this distinction absolutely. The epiblast gives rise to much besides epithelium, as does also the hypoblast, and there seems even now consid-



erable doubt whether the serous cavities are hypoblastic or mesoblastic, and whether, in consequence, the layer of cells covering these cavities is endothelium or epithelium. The attempt to carry out rigidly this mode of classification has led to great difficulties in the nomenclature and placing of some tumours. Thus we have tumours of the pleura and peritoneum, which have the structure of cancers, but German authors regard the endothelium of these cavities as belonging to the connective tissue, and many of them would place such tumours amongst those arising from connective tissue, and separate them from the cancers.

The structure and mode of growth have to be taken into account as well as the origin of tumours. It has been indicated above that in respect to structure and mode of growth tumours may be typical or atypical. It will be convenient to divide tumours, in the first instance, on this basis into two great groups, the typical and the atypical. In the further subdivision of these groups the relation of the tissue to the corresponding normal tissue is taken into consideration, and we may have both typical and atypical tumours referrible to the same normal tissue.

The grouping of tumours into typical and atypical forms will nearly correspond with the clinical distinction of innocent and malignant, but it must not be forgotten that the distinction is not absolute, and that typical tumours sometimes assume the characters of the atypical.

In naming tumours the structure is chiefly taken into account. The typical tumours are named by adding the usual suffix *-oma* to the name of the tissue. The atypical tumours have special names which are variously derived. Thus, Sarcoma is a name originally used in a very indefinite way, but by Virchow applied to atypical connective-tissue tumours. Carcinoma also had at one time a wide and somewhat indefinite significance, but is now limited to atypical tumours whose more important structure is epithelium.

A very great advance has been made in our knowledge of tumours by the publication of Virchow's classical work, "Die krankhaften Geschwülste." From this work many of the following illustrations are borrowed by the kindness of the author.

**Literature.**—J. MÜLLER, *Bau und Form der Krankhaften Geschwülste*, 1838; VIRCHOW, *Die Krankhaften Geschwülste*, 1864-67; PAGET, *Lectures on Tumours*, 1852, *Surgical Pathology*, 3rd ed., 1870, *Path. trans.*, xxv., p. 319; LÜCKE, in *Pitha and Billroth's Handbuch*, vol. ii., part 2, 1869; COHNHEIM, *Allg. Path.*, 2nd ed., vol. i., 1882; BROCA, *Traité des Tumeurs*; BUTLIN, *Art. Tumors*, *Internat. Encycl. of Surg.*, vol. iv., 1884; ZAHN, *Congrès méd. internat. de Genève*, 1878; LEOPOLD, *Virch. Arch.*, lxxxv., p. 283; BAKER, *St. Barth. Hosp. Rep.*, vol. ii., p. 129; CRIPPS, *do.*, vol. xiv., p. 287; SNOW, *Clin. notes on cancer*, 1883; THIERSCH, *Epithelialkrebs*, 1865; WALDEYER, *Volkman's Sammlung* No. 33, 1872; VIRCHOW'S *Arch.*, xli. and lv.; COATS, *Path. trans.*, xxxviii., p. 399, 1887.



## SECTION IX.—CONTINUED.

## A.—TYPICAL OR HISTIOID TUMOURS.

1. FIBROMA, including *molluscum fibrosum* or *neuro-fibroma* and *hard fibroma*;
2. LIPOMA, *diffuse* or *encapsuled*, *pendulous*; 3. MYXOMA, *characters of mucous tissue, hydatid mole, proper tumour, varieties of*; 4. CHONDROMA, *as ecchondrosis or enchondroma, the latter usually in connection with bone*; 5. OSTEOMA, *chiefly as exostosis, Odontoma*; 6. MYOMA, *the rhabdomyoma rare, the leiomyoma common, structure and relations of latter*; 7. NEUROMA, *the true neuroma, Wood's painful subcutaneous tumour*; 8. ANGIOMA, *capillary, cavernous, lymphatic*;
9. GLIOMA; 10. PSAMMOMA; 11. LYMPHOMA; 12. PAPILLOMA; 13. ADENOMA, *of various structure according to gland*; 14. CYSTOMA, *Cysts arising from pre-existing cavities, including retention cysts; and cysts of independent origin, including dermoid cysts, adenoid cystoma, extravasation cysts, etc.*; 15. TERATOMA.

## 1.—THE FIBROMA OR CONNECTIVE TISSUE TUMOUR.

AS fibrous tissue is frequently the product of inflammation we may expect that the demarcation between inflammatory newformations and fibromas is not always easy to make. Elephantiasis arabum is sometimes regarded as a simple tumour, although more properly belonging to the infective tumours. There are cases, however, of local enlargement of the skin, which have not the regular course of elephantiasis, and which have more apparent analogies with tumours. To such cases Paget's name of "Cutaneous outgrowth" is applicable. There are cases also in which, with or without thickening of the cutis, there is a definite encapsuled tumour, composed of soft connective tissue, beneath the skin. Such tumours occur chiefly in the external organs of generation, and may attain large dimensions, one observed by the author weighing fifteen pounds. (See Elephantiasis.)

**Fibroma molluscum** or **Multiple fibroma of the skin** are terms applied to cases in which we have multiple isolated tumours beginning as little growths of connective tissue in diverse regions of the skin, and afterwards growing out and becoming pendulous. Thus we may have hundreds of more or less pendulous tumours in various parts of the



body. This purely fibrous molluscum must be carefully distinguished from molluscum contagiosum, a totally different disease, which will be considered in the section on Diseases of the Skin.

These multiple fibromata have been made the subject of special study by Recklinghausen, who found them in two cases associated with multiple tumours of the nerves. The tumours of the nerves formed usually oval swellings in their course, and were composed of comparatively soft connective tissue in which the nerve-fibres were embedded. These tumours are frequently described under the name of **Neuroma**, but as they are really fibrous tissue tumours they are to be regarded as false neuromata. The term **Neurofibroma** or fibro-neuroma is more correctly applied. In Recklinghausen's case the tumours of the skin were also found in their origin to be connected with nerve-stems, growing from the nerve sheath, although in growing they frequently involved the sheaths of neighbouring canals, such as blood-vessels and sweat-glands. The tumours of the nerves have frequently a plexiform character, being formed by enlargement of the sheaths of a plexus of nerves. A similar character is sometimes presented by the cutaneous tumours and may assist in their diagnosis. Whether Recklinghausen's view applies generally to the multiple fibroma of the skin or not, remains to be determined. It is confirmed by Kriege and Westphalen, while its universal applicability is questioned by Philippon.

**Hard Fibromas** or true Fibromas are exceedingly dense tumours composed of firm connective tissue tightly interlaced and resembling tendon in its structure. On section they show a brilliant white surface and often a concentric arrangement of the connective tissue bundles. We do not include here the uterine fibromas, which are really muscular tumours, and will be described as such. Fibromas are frequent in connection with periosteum and bone, especially on the jaws. Growing from the periosteum they are sometimes intimately connected with the bone which may be as if buried in the tumour. Sometimes the fibroma originates inside a bone. Fibromas also occur on fascias and membranes, as the dura mater, and are not uncommon in the mamma. Just as dense connective tissue formed in chronic inflammation may become calcified, so may dense fibromas be partially infiltrated with lime salts. They sometimes undergo partial ossification, or they may be mixed with bone, cartilage, or gland tissue.

**Literature.**—VIRCHOW's *Geschwülste*, i.; MORTON and COATS, *Glasg. Med. Jour.*, iii., 145, 1870; RECKLINGHAUSEN, *Die multiplen Fibrome der Haut*, 1882; KRIEGE, *Virch. Arch.*, cviii. 466; WESTPHALEN, *do.*, cx. 29; PHILIPPSON, *do.*, 602.

## 2.—THE LIPOMA OR FATTY TUMOUR.

This form of tumour consists of adipose tissue exactly like that of the body, as for instance the subcutaneous adipose tissue. Adipose tissue contains bands of fibrous connective tissue which carry the vessels and nerves, and so do lipomas, but in different tumours this is variously



abundant. If there is little connective tissue the tumour will be soft, and may even feel fluctuant. If there is much it will be hard, and we may even have an approach to the fibrous tumour, the **fibro-lipoma**. The fibrous character may be increased by irritation, as where a tumour is exposed to friction, producing a kind of indurative inflammation in the tumour.

Fatty tumours are mostly surrounded by a distinct capsule, but sometimes they are continuous with the surrounding fat. Thus the fat around the mamma or the kidney may undergo such an enlargement as to warrant the name of tumour (*Lipoma capsulare*). Billroth mentions a lipoma which had grown in between the muscles of the thigh in such a way as that it could not be removed completely. Lipomas which are not definitely circumscribed may be called **Diffuse lipomas**.

Lipomas are of common occurrence. Their most frequent seat is under the skin, especially of the trunk. They are rare under mucous membranes, as that of the stomach (Fig. 62), where there is normally a small quantity of fat corresponding with the subcutaneous fat. They are occasionally met with in connection with serous and synovial membranes, in the appendices epiploicæ of the intestines or the synovial fringes. They are rarely heterologous, occurring where fat is not present normally, as in the kidneys, brain, etc. The author met with a case in which an elongated piece of fat lay on the upper surface of the corpus callosum. Lipomas are for the most part single, but in some cases they are **multiple**, although rarely symmetrical. When multiple they do not grow simultaneously, and may appear in succession for months or years.

Fatty tumours sometimes become **pendulous**. Those in the appendices epiploicæ (Fig. 63) are so from the first, and the neck of the polypoid tumour may get severed, the tumour becoming a **loose body** in the peritoneum; it is similar with the lipomas of the synovial fringes. Subcutaneous lipomas may also become pendulous and polypoid, and may grow to great dimensions in this form. There is apt to be ulceration of the surface of such tumours, and even hæmorrhage. The tumours of the subcutaneous tissue may, by their weight, gradually

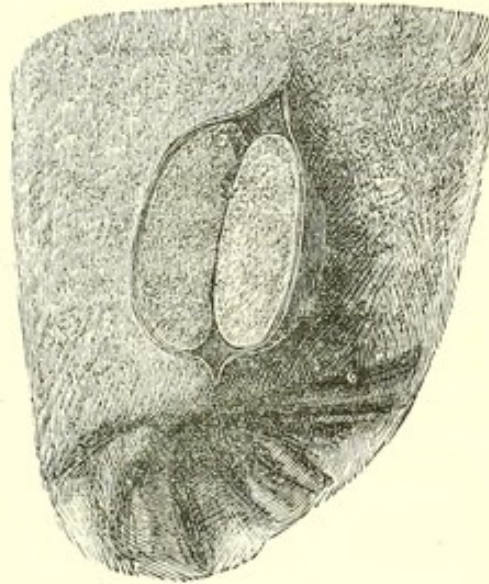


Fig. 62.—Lipoma in the wall of the stomach. It was the size of a hazel-nut, situated near the pylorus in the submucous tissue. Natural size. (VIRCHOW.)



slide downwards, leaving their old attachments and acquiring new ones. Besides these changes we have the occasional induration already men-

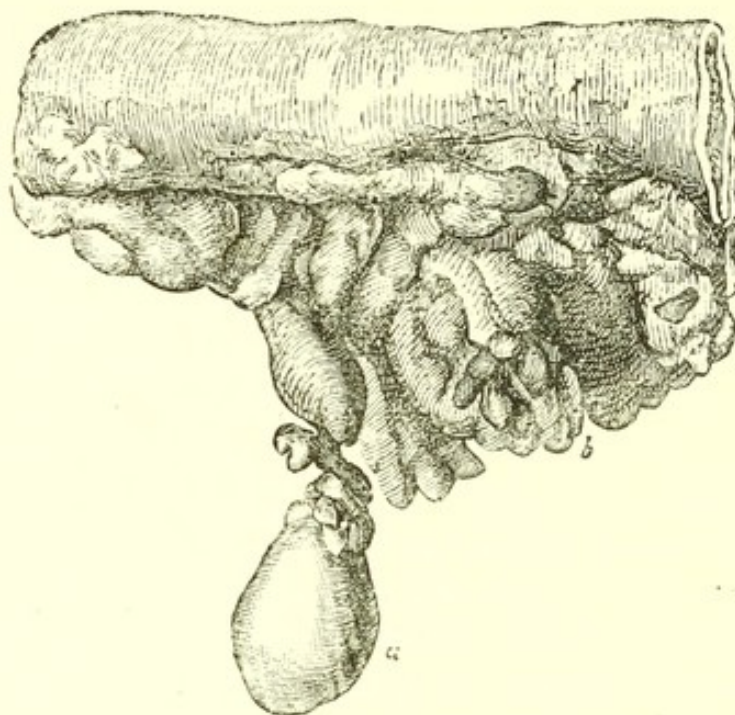


Fig. 63.—Pendulous lipoma of appendices epiploicae of colon. A pedunculated tumour (*a*) projects from the general mass of subserous adipose tissue (*b*). It is twisted twice on its axis, and the peduncle is very thin. (VIRCHOW.)

tioned, and sometimes calcareous infiltration follows. Softening, with the formation of a cyst inside the tumour is a rare occurrence.

### 3.—THE MYXOMA OR MUCOUS-TISSUE TUMOUR.

This form of tumour is composed of **mucous tissue**, and as this is not one of the physiological tissues of the adult, it will be proper to refer more specially to its characters. The blood-vessels of the umbilical cord are padded and protected from pressure by a gelatinous substance called Wharton's jelly. Under the microscope, this consists of variously shaped cells separated by a clear transparent intercellular substance. The intercellular substance is of gelatinous consistence, and owes this character to the fact that it consists of a watery solution of mucin.

Mucin is nearly allied to albumen, but when present in even small quantity in a solution, it gives the latter a sticky gelatinous character. Its chemical reactions also differ from those of albumen, in respect that though like it precipitated by alcohol, the precipitate is redissolved by water. Also, acetic acid and other organic acids precipitate mucin, but not usually albumen. The precipitate of mucin by alcohol or acetic acid is more membranous than that of albumen. The reaction with acetic acid can be readily brought out in Wharton's jelly, a microscopic section



showing, on adding acetic acid, a reticulated precipitate. These reactions can also be studied in the mucus from any mucous membrane, or in the bile, in which mucin is normally one of the dissolved constituents.

Besides in the umbilical cord, mucous tissue is present in the villi of the chorion of the foetus, the villi consisting of a covering layer of epithelium with mucous tissue internally. In the foetus, it is also present in early stages in the subcutaneous tissue where it has the place of the subcutaneous adipose tissue, being, in fact, related to fat very much as the temporary cartilage is to bone. Some remains of this tissue are met with in the adult. Thus, the vitreous humour of the eye is really composed of soft mucous tissue, and traces of it have been found in places where normally adipose tissue exists, as under the pericardium, at the hilus of the kidney, subcutaneously, and in the medulla of bone. It appears that in these positions there is sometimes a partial recurrence to the foetal condition. The connective substance of the brain, the neuroglia, is allied to mucous tissue, and seems to present a proneness to return to that form.

**The hydatid mole**, which arises by a great newformation of mucous tissue in the villi of the chorion, is regarded by some as a form of myxoma. (See further in Section on Generative Organs.)

**Mucous-tissue tumours** are met with principally in the subcutaneous tissue, where they may be regarded as perhaps due to a piece of embryonic tissue left over when the mucous tissue was converted into adipose, in this respect resembling the chondromas of bone. They form rounded or oval tumours generally soft, sometimes almost fluctuant in consistence and hyaline in appearance. These tumours are sometimes described as soft fibromas or fibro-cellular tumours. Examined under the microscope we have a very trans-

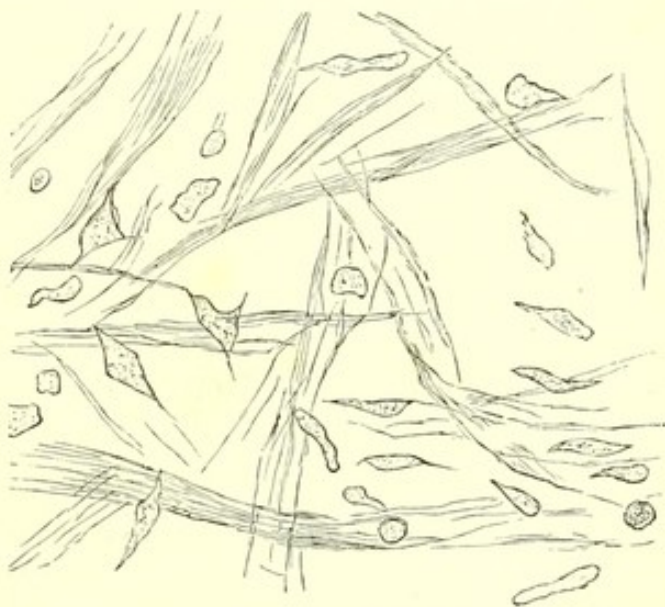


Fig. 64.—Microscopic section of a myxoma of the subcutaneous tissue. Isolated cells and strands of fibres are shown. Between them a clear gelatinous material was present.  $\times 350$ .

lucent tissue in the midst of which there are irregularly shaped cells. The tissue is intersected by more or less frequent bands of connective tissue (Fig. 64). The proportion of cells varies considerably in different cases, and even in different parts of the same tumour.



Myxomas are also of somewhat frequent occurrence in the brain as compared with other tumours in this organ. They arise in connection with the soft membranes of the brain and spinal cord, or in the ventricles, or in the substance of the brain. They sometimes grow to considerable size, and are liable by softening to take the form of cysts filled with a mucous fluid. They are also met with on peripheral nerves. They occur in the mamma, where not infrequently they appear to have the character of a diffuse formation of mucous tissue between the glandular acini, so that the gland as a whole is converted into a tumour. In the salivary glands they occur, but are usually of mixed structure, being partly formed of other kinds of tissue.

The simple pure hyaline myxoma is not very common, as the tissue is apt to be mixed with other forms. From the relation of mucous and adipose tissues it is not remarkable that a partial conversion of a lipoma into a myxoma or *vice versa*, is met with. There is also not infrequently a mixture with fibrous, cartilaginous, or gland tissue. Then, also, the myxomas vary very greatly in the proportion of cells, the very cellular ones being called medullary myxomas, and frequently graduating into sarcomas. In fact, with pathologists who take the embryonic nature of the tissue as the criterion of sarcomas, the fact that mucous tissue is the embryonic precursor of adipose tissue induces them to class this form of tumour among the sarcomas. Consistently with these facts there are some myxomas which show malignant characters, either local or general. In a case recorded by Virchow there were tumours on the nerves and in the dura mater of the cord and brain.

**Literature.**—MÜLLER, Arch. f. Anat. u. phys., 1836; VIRCHOW, Arch. xi. 286, and Geschwülste, i., p. 396.

#### 4.—THE CHONDROMA OR CARTILAGINOUS TUMOUR.

This tumour is composed of cartilaginous tissue. The cartilage may be hyaline or fibrous, generally the latter, and the matrix is not infrequently rather soft. The tumour is also intersected with fibrous bands which carry blood-vessels that nourish the tissue. If a cartilaginous tumour grows in connection with and out from cartilage it is called an **Ecchondrosis**, but if, as in the majority of cases, it grows in connection with other tissues, then it is called an **Enchondroma**. The term chondroma of course includes both.

The ecchondroses are usually small unimportant outgrowths chiefly of the cartilages of the septum nasi, larynx and trachea and ribs.

Virchow has described an interesting form of so-called ecchondrosis at the basilar portion of the occipital and sphenoid bones. The basilar parts of these bones are



formed from cartilage, and in adult life they are united, forming a single bone (os tribasillare). The junction takes place irregularly by a kind of toothed union. In this process a little bit of cartilage may be omitted, and this sometimes develops into a little tumour just under the basilar artery to which it may be adherent. It will be seen that we have here an actual instance of a little piece of embryonic tissue left over to develop afterwards into a tumour. This little tumour sometimes undergoes a remarkable change; the cells swell up and become like those of the chorda dorsalis, but in other cases it ossifies.

The cartilaginous loose bodies in joints are sometimes regarded as originally outgrowths from the cartilages or synovial fringes which have been broken off and grown after their separation.

**Enchondromas** are mostly met with in connection with Bones. As bones are developed out of cartilage for the most part, it may be supposed that little bits of the embryonic cartilage are left over and develop into tumours afterwards. The enchondromas of bone may be divided into central and peripheral, according as they originate in the medulla or at the surface. The former are met with chiefly in early life, and may be congenital. They occur especially in the fingers and toes, which may be the seat of multiple tumours as in Fig. 65. These tumours begin inside the phalanges or metacarpal bones, or, less frequently, the analogous bones of the foot. Growing inside the bones they may be for a time unperceived, but afterwards swell up the bones and may even burst through the external shell. These tumours often show a local malignancy, growing by the formation of new nodules in the tissue around. The peripheral chondromas are met with most frequently on the femur and pelvis, and most rarely on the bones of the face and skull.

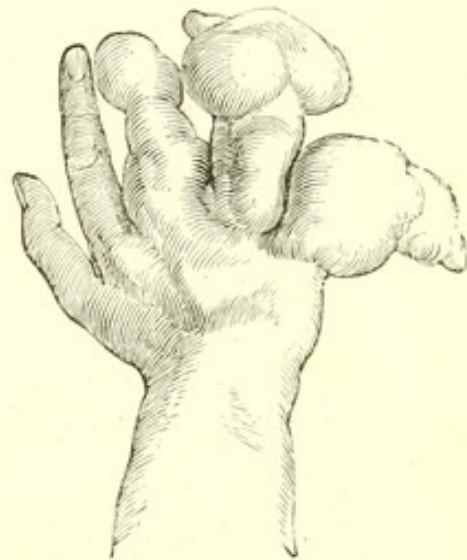


Fig. 65.—Multiple internal enchondromata of fingers. (CORNIL and RANVIER.)

In soft parts chondromas are found occasionally, but are commonly mixed with fibrous, mucous, glandular, sarcomatous, or cancerous tissue. They are particularly frequent in glands, as the testes, ovaries, mammae, and salivary glands. In these situations they are occasionally found to follow chronic irritations.

In their form chondromas are generally rounded tumours and distinctly encapsuled, but if large they are lobulated.

It has already been said that the chondromas are often mixed with other tissue, and it is to be added that secondary changes are not infrequent. They sometimes soften, and this is frequently due to partial



transformation into mucous tissue. This is particularly the case with the glandular enchondromas. On the other hand, those of bone are liable to ossify.

Lastly, chondromas are liable, as has already been said, to show a certain malignancy, forming secondary tumours, especially in the lungs. This is connected with the frequently mixed character of these tumours and their association especially with sarcomas and cancers.

**Literature.**—A. COOPER, *Surgical Essays*, 1818; VIRCHOW, *Geschwülste*, i. 435; MURCHISON, *Edin. Monthly Jour.*, 1852; SYME (Congenital enchondroma), *Lancet*, 1855, p. 116.

### 5.—THE OSTEOMA OR BONY TUMOUR.

In this class are included tumours composed of bone, not mere new-formations due to inflammation, nor tumours in which bone exists as a subordinate element with other tissue.

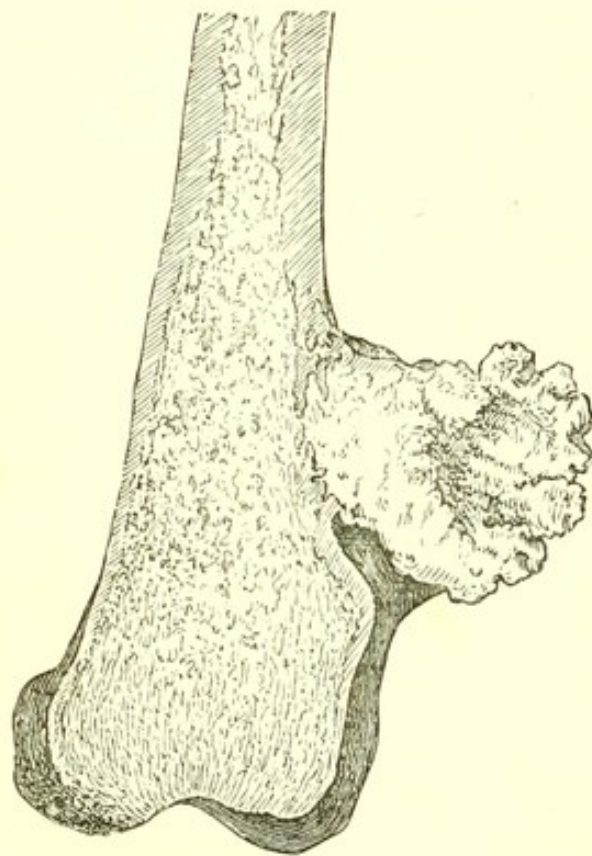


Fig. 66.—Cartilaginous spongy exostosis of femur. (VIRCHOW.)

The large majority of bony tumours grow from bone, and are hence called **Exostoses**. In regard to their structure, some are like spongy bone with the interstices filled with ordinary bone-marrow, and some are composed of dense bone such as forms the shaft of a long bone, and are called ivory exostoses.

There are also tumours which originate in the teeth. When these are composed of cement, they are properly called dental **osteomas**, but when formed of dentine, **odontomas**. The latter term, however, is often applied

to either form. The tumour may be little more than a local enlargement of the fang, rendering extraction difficult, or there may be a distinct tumour, growing sometimes to the size of a walnut. In the latter case it is usually a true odontoma, composed of a structure like dentine.

Of the proper exostoses several forms are distinguished.

(1) The **Spongy exostosis**, or the **Exostosis cartilaginea** (Fig. 66). These tumours occur mostly at the epiphyses of the long bones, and are



derived primarily from the epiphyseal cartilage. A rather favourite seat is the dorsal aspect of the last phalanx of the great toe, where they project beneath the nail and produce great pain and discomfort. They grow during childhood, and, just as the cartilage from which they originate ossifies, so do they, and the bony tumour formed is directly continuous with the bone beneath. The tumour begins as a small outgrowth, and so the first bone formed is a narrow piece. The cartilage, as it goes on growing, enlarges in every direction, and so overhangs its base, the tumour thus becoming larger as it grows outwards, and consequently pedunculated. The tumour consists of spongy bone with a thin layer of cartilage on its surface. It is enough to snip through the base in order to remove the tumour.

(2) **The Ivory exostosis** is mostly met with on the bones of the head, but also on the pelvis, scapula, great toe, etc. The tumours are usually rounded in form, and may be tuberculated on the surface. In their favourite seat on the head they may grow from the external table and project externally, or from the internal table and project internally (as in Fig. 67), in which case they may produce irritation of the brain substance beneath. It sometimes happens that an ivory exostosis grows

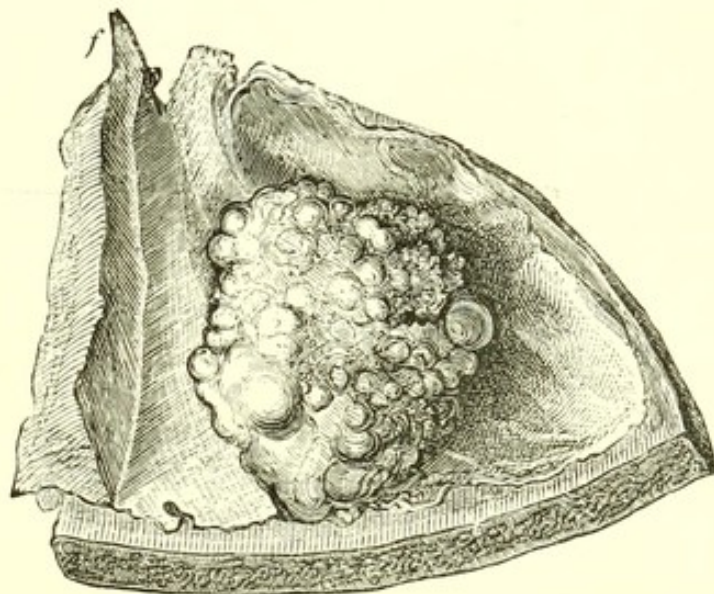


Fig. 67.- Internal ivory exostosis of the frontal bone. The surface is tuberculated, and the tumour has a narrow base. It was situated to the left of the falx (f). Natural size. (VIRCHOW.)

from corresponding parts of both external and internal tables. These tumours are sometimes multiple (as in Fig. 68).

(3) **Hyperostosis and Periostosis** are names applied to growths of bone which are not properly tumours. They are localized thickenings of bones or portions of bones. This growth of particular parts of the bones of the head may become so independent in its manner that the characters of a tumour are simulated.

**Osteomas of other parts** than bones are rather rare. It is remarkable, however, that bony masses occur sometimes in the central nervous system. They are met with in the arachnoid, where they used to be regarded as evidences of chronic irritation, and are hardly tumours.



Actual tumours occur in the dura mater, and even in the brain substance. They are also met with in the eyeball, in the lungs, and, as little bony granules, in the skin.

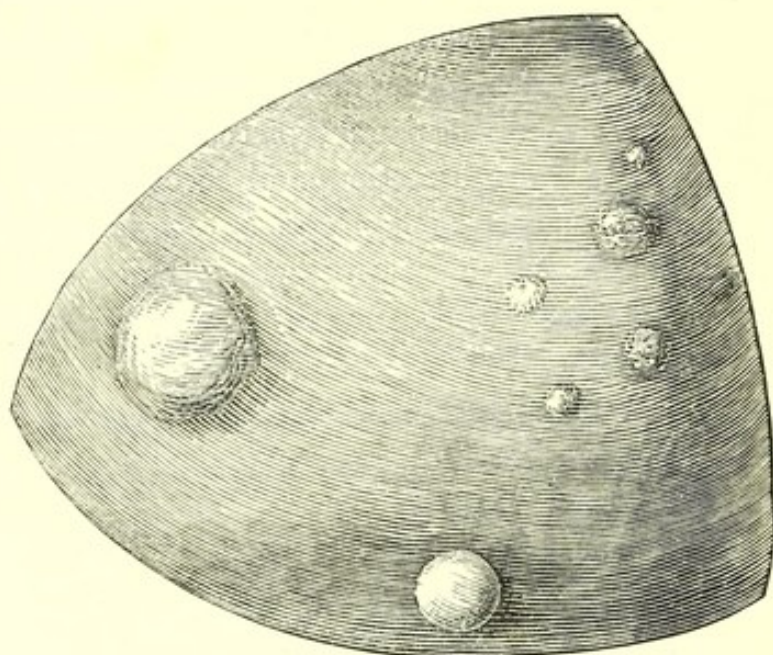


Fig. 68.—Multiple exostoses of the frontal bone. Natural size. (VIRCHOW.)

**Literature.**—WEBER, *Die Knochengeschwülste*, 1856; VIRCHOW, *Geschwülste*, ii. 1 and 53 (Odontoma).

#### 6.—THE MYOMA OR MUSCULAR-TISSUE TUMOUR.

These are tumours in which muscular tissue is the essential constituent, but just as all muscles have supporting connective tissue so have these, some more, some less. As there are two kinds of muscle, so are there two forms of muscular tumour, those composed of striated, and those of smooth muscle.

The **Myoma strio-cellulare** or **Rhabdomyoma** is very rare. Tumours of this structure are probably always congenital. They have been seen in the heart, kidneys, ovaries, and testicles. The tumour is not usually composed of ordinary striated muscle, but the muscular fibre is embryonic in character, consisting of spindle-shaped cells, which are transversely striated. Tumours of this kind have strong analogies with sarcomas, and are sometimes designated myo-sarcomas. Besides this, similar rudimentary muscle is occasionally met with in other forms of sarcoma, and in cystic tumours of the ovary and testicle.

The **Myoma lævi-cellulare** or **Leiomyoma** is an exceedingly common form of tumour, and is met with in almost every part where smooth muscle exists normally. According to the amount and density of the interstitial connective tissue is the consistence of the myoma—it may



be very dense and warrant the name Fibro-myoma, or it may be so hard as to resemble cartilage.



Fig. 69.—Section of a myoma of the uterus stained with carmine. The muscular nuclei are seen in longitudinal and transverse section.  $\times 350$ .

Myomas are often described as fibrous tumours, and in appearance they justify this designation. To the naked eye they appear fibrous on section, and even under the microscope they show a fibrous appearance. On adding acetic acid to a microscopic section, or on staining with carmine or other agent, the fibres are seen to be much more abundantly nucleated than ordinary connective tissue. In fact, rod-shaped nuclei (not spindles as in connective tissue) are so closely set as at once to

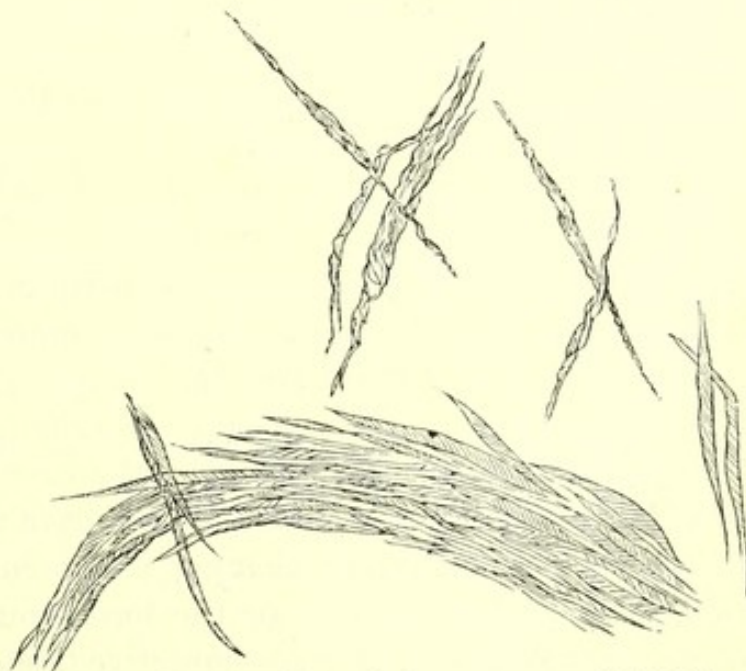


Fig. 70.—Muscular fibre cells from a myoma isolated by steeping in nitric acid.  $\times 350$ .



suggest a cellular tissue (see Fig. 69). It may here be remarked that in the unimpregnated uterus there is the same difficulty in distinguishing the individual spindle-cells. In both cases, however, the cells may be isolated by macerating the tissue for twenty-four hours in a 20 per cent. solution of nitric acid, or for twenty to thirty minutes in a 34 per cent. solution of caustic potash. This makes the tissue rotten, and separates the cells, which are recognised as spindles (as in Fig. 70).

Myomas always arise where muscle already exists, and as smooth muscle is most frequent in the walls of mucous canals and cavities, it is there that they usually originate. The tumour may be continuous with the muscular wall, forming an outgrowth from it, or it may be distinctly isolated and encapsuled. It may remain in the substance of the muscular wall (*intraparietal* or *intramural*), or it may slip inwards so as to bulge under the mucous membrane (*submucous*), or it may pass outward and present under the serous coat (*subserous*). In the two latter cases the tumours often become polypoid.

By far the most frequent seat of the myoma is the **female organs of generation**, generally the uterus, but also the ligaments and ovaries. The so-called uterine fibroids are myomas, and the most important of these are the submucous which so frequently become polypoid, and give rise to hæmorrhage, sloughing, etc.

In the **prostate** the hypertrophy frequently met with in old men is really from the formation of muscular tissue, and the third lobe which forms a prominent bulging projection at the neck of the bladder is an outgrowth from the muscle of the prostate. Sometimes there are even isolated muscular tumours in the midst of the prostate. This form of tumour is to be distinguished from the glandular tumour, which is a much rarer form of hypertrophy and occurs mostly in young men. Myomas of the **œsophagus, stomach, and intestine** are infrequent, and are usually submucous. They have also been seen in the urinary bladder. Lastly, they are rarely met with in the **skin**, one or two cases in connection with the nipple and the scrotum having been observed.

The myoma is of **slow growth**, but may go on as long as thirty or forty years, and may reach a very great weight, as much as 60 lbs. It is nearly always an innocent tumour, but a case is recorded by Brodowski in which a large myoma of the stomach gave rise to secondary tumours in the liver. **Retrograde changes** may occur, such as fatty degeneration, resulting in shrinking or the formation of cysts. If induration occurs from formation of hard connective tissue, this may calcify, leaving the muscular tissue in the spaces between the calcified trabeculæ. In some cases, from derangements of the circulation in large tumours, we



may have an actual necrosis of a portion of the muscular substance, resulting sometimes in absorption and the formation of a cyst. In other cases, the dead structures become calcified as in Fig. 71, where muscular elements, connective tissue, and walls of blood-vessels were all

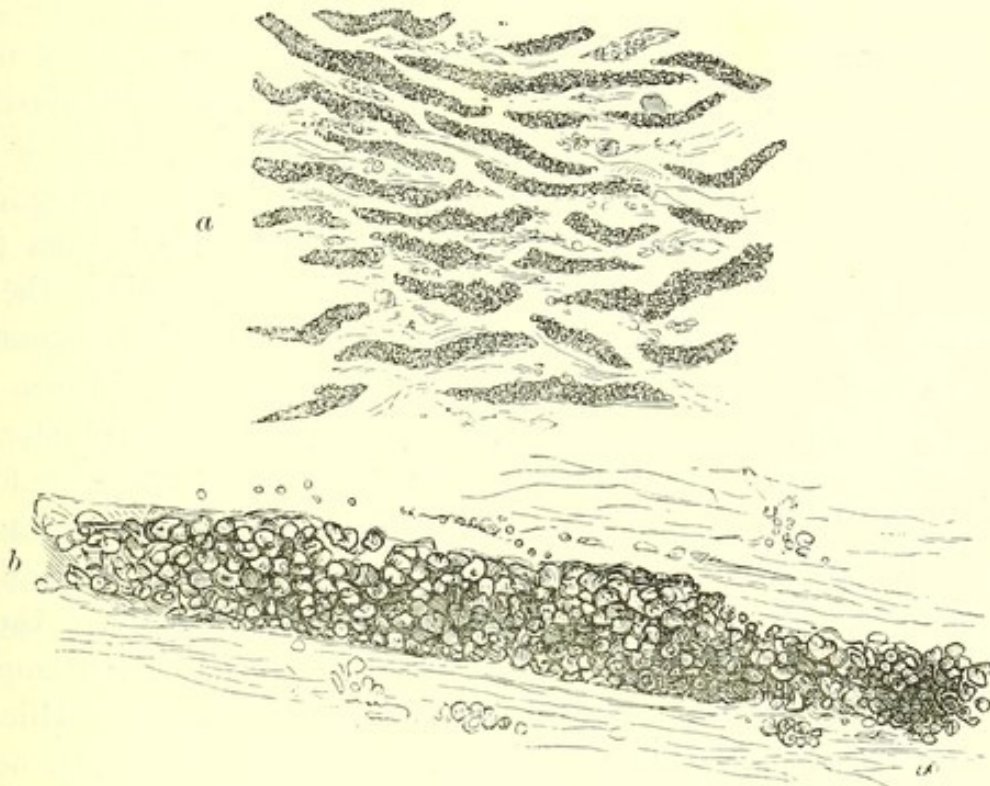


Fig. 71.—From a myoma, a portion of which had become calcified. *a*, muscular fibre-cells impregnated with lime salts; *b*, a blood-vessel with wall incrustated.  $\times 350$ .

found impregnated with lime. As the myomas so readily become poly-poid, they are liable to be insufficiently nourished, as the neck gets thinner; sloughing may even occur, especially if they present on a mucous surface, and become exposed to injury. Such tumours are also liable to bleed, especially as the vessels in them sometimes undergo great dilatation.

**Literature.**—A considerable number of isolated cases of Rhabdo-myoma are recorded. A list of these is given by KOLESSNIKOW in Virch. Arch., lxxviii. 554, and by HUBER and BOSTRÖM, D. Arch. f. klin. Med. xxiii., 208; see also COHNHEIM, Virch. Arch., lxvi.; MARCHAND, do., lxxiii. and c. (in last-mentioned paper, Glycogen found in muscle). *Leiomyoma*—see chiefly VIRCHOW'S Geschwülste, iii.

## 7.—THE NEUROMA OR NERVOUS-TISSUE TUMOUR.

A neuroma is properly a tumour composed of nerve tissue. As there are two kinds of nerve tissue so we may distinguish a ganglionic and a fibrous neuroma. There are some cases of small tumours projecting from the cortex of the brain which may be named ganglionic neuromas, but they are excessively rare, and the name neuroma is virtually reserved for tumours of nerve stems.



All tumours of nerve stems are usually designated neuromas, but as such tumours may really be composed of fatty, mucous, or fibrous tissue, it has become customary to distinguish those which actually contain new-formed nerve fibres as true neuromas, and the others as false neuromas.

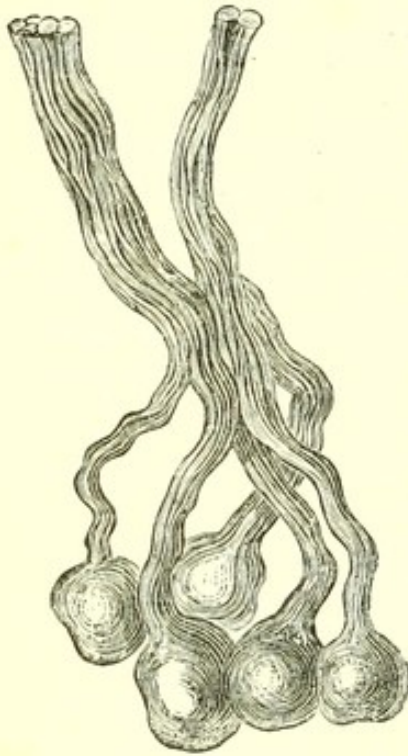


Fig. 72.—Amputation-neuroma. The nerves end in knobbed extremities. The case was one of amputation in the upper arm. Two thirds of the natural size. (VIRCHOW.)

The true neuroma is composed of nerve fibres, which may be either medullated, as in the ordinary cerebro-spinal nerve, or non-medullated. One of the most striking forms is the so-called **Amputation neuroma** (Fig. 72). Sometimes after amputations the cut ends of the nerves produce little knots or knobs which have a hard consistence and to the naked eye look fibrous. Under the microscope medullated nerve fibres are found running in bundles, but there are also many fine fibres which are probably non-medullated nerve fibres. It is in this case as if the cut end of the nerve had made an attempt at regeneration of the lost portion. Allied to this form is the traumatic neuroma, occurring as the result of injury in the course of a nerve.

But neuromas occur in the course of nerves spontaneously, and they are often multiple, forming oval swellings, hard and fibrous in appearance. They contain much fibrous tissue, in the midst of which there are medullated nerve-fibres (see Fig. 73) recognizable in the fresh

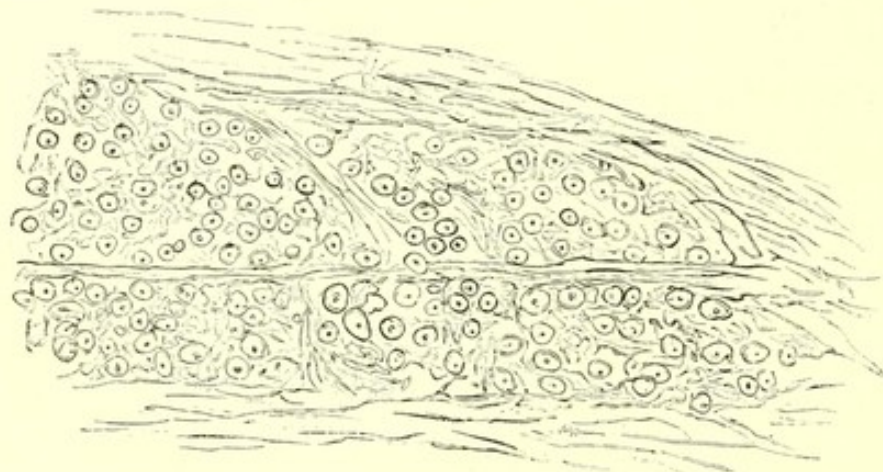


Fig. 73.—Transverse section of a neuroma. The medullated nerve-fibres are shown, the appearance being much like that of the section of a nerve.  $\times 80$ .

state by the double contour, especially when the connective tissue has



been rendered transparent by acetic acid or liquor potassæ. But usually in addition to these medullated fibres, there are fine nucleated fibres which Virchow regards as non-medullated nerve-fibres. In some neuromas these are very abundant.

There is no doubt that most tumours of nerves are really false neuromas. This applies, as we have seen at p. 228, even to the multiple tumours which have been usually regarded as the typical neuromas. It applies also, at least in part, to the amputation neuromas. It is obvious that as a nerve stem is composed of connective tissue and nerve-fibres, it must be often very difficult to determine whether there has been newformation of nerve-fibres or not. It seems questionable whether it is worth while, for the sake of consistency in nomenclature, to attempt to restrict the term neuroma to those actually composed of nerve tissue.

The **painful subcutaneous tumour, Tubercula dolorosa**, or Wood's tumour, is a somewhat difficult growth to classify. It occurs in the form of a small round tumour under the skin, and is commonly the seat of intense pain. The pain indicates some connection with the nerves, and it has been suggested that the tumour may even be a non-medullated neuroma. According to Virchow there are tumours of various structure in this category, myomas, angiomas, neuromas.

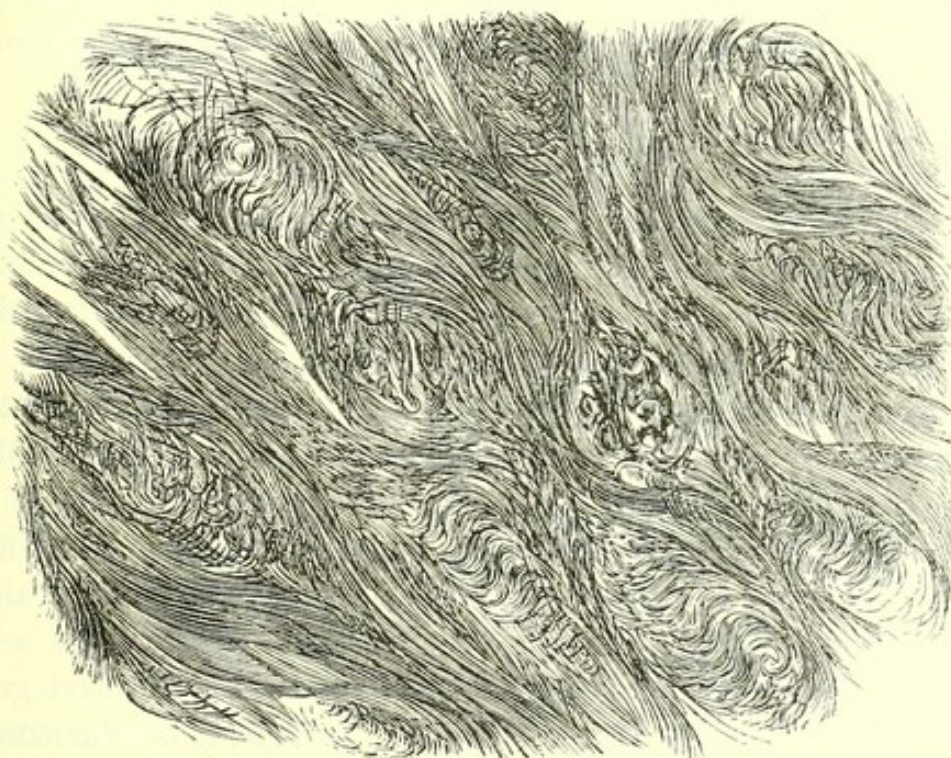


Fig. 74.—Section of a painful subcutaneous tumour (Wood's tumour). The appearance is somewhat like that of a myoma.  $\times 80$ .

Axmann has suggested, with some probability, that the little tumour may be an enlarged Pacinian body. Recently Hoggan has described one which he believes to be an adenoma of the sweat-glands, but which Virchow rather takes to be an angioma. In the case from which our figures are taken (Figs. 74 and 75) the tumour consisted of a dense



interlacing network of fibres, very suggestive of the appearance of a myoma (Fig. 74). On macerating portions in nitric acid the tissue broke up into large spindle cells, as shown in Fig. 75. This tumour is probably a myoma of the skin, and several others examined by the author had a similar structure.

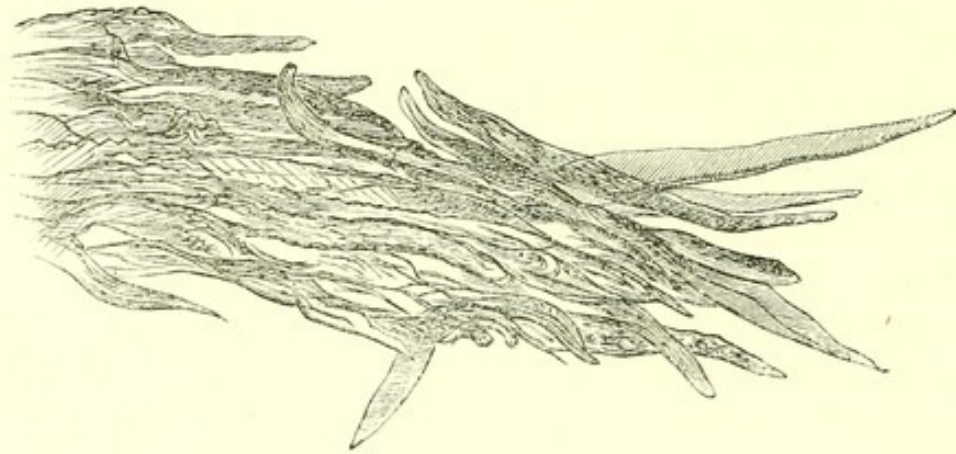


Fig. 75.—From same tumour as Fig. 74, after maceration in nitric acid. Large spindles are isolated, probably smooth muscle cells.  $\times 350$ .

**Literature.**—VIRCHOW, *Geschwülste*, iii. 233; R. W. SMITH, *On Neuroma*, 1849; WOOD, *On painful subcutaneous tum.*, 1812; PAGET, *Lect. on surg. path.*, 3rd ed., p. 490. *On false neuroma*, see especially RECKLINGHAUSEN, *On multiple fibroma*, 1882; KRIEGE, *Virch. Arch.*, cviii.; WESTPHALEN, *Virch. Arch.*, cx.; PHILIPPSON, *Virch. Arch.*, cx.

#### 8.—THE ANGIOMA OR VASCULAR TUMOUR.

We have here a tumour composed of blood or lymphatic vessels. The tumour varies in bulk according to the contents of the vessels, hence some of them are occasionally designated **Erectile tumours**. The term **Nævus** is almost synonymous with angioma of the surface.

The commonest form is (1) the **Plexiform** or **Capillary angioma**. This includes most of the vascular nævi, and consists of capillary and intermediate vessels forming a rich plexus. It is mostly a growth of the skin, and may be very small or cover a large area, forming a flat soft surface of dark or bright hue. It is nearly always congenital, although it may increase in size after birth. Minute capillary nævi are very common, and are frequently multiple. These capillary nævi graduate into those consisting of dilated veins (2) the **Venous** or **Varicose nævi** (Fig. 76). They have similar situations to the capillary form and similar appearances.

(3) The **Cavernous angioma** consists of tissue like that of the corpus cavernosum of the penis or clitoris, namely a network with meshes which communicate freely and are filled with blood. When empty they are seen to be composed of a pale tissue, in its texture resembling a sponge, with variously thick trabeculæ and variously wide spaces.



These trabeculae are all accurately lined with endothelium, and consist of connective tissue with some muscular fibre-cells in them. These

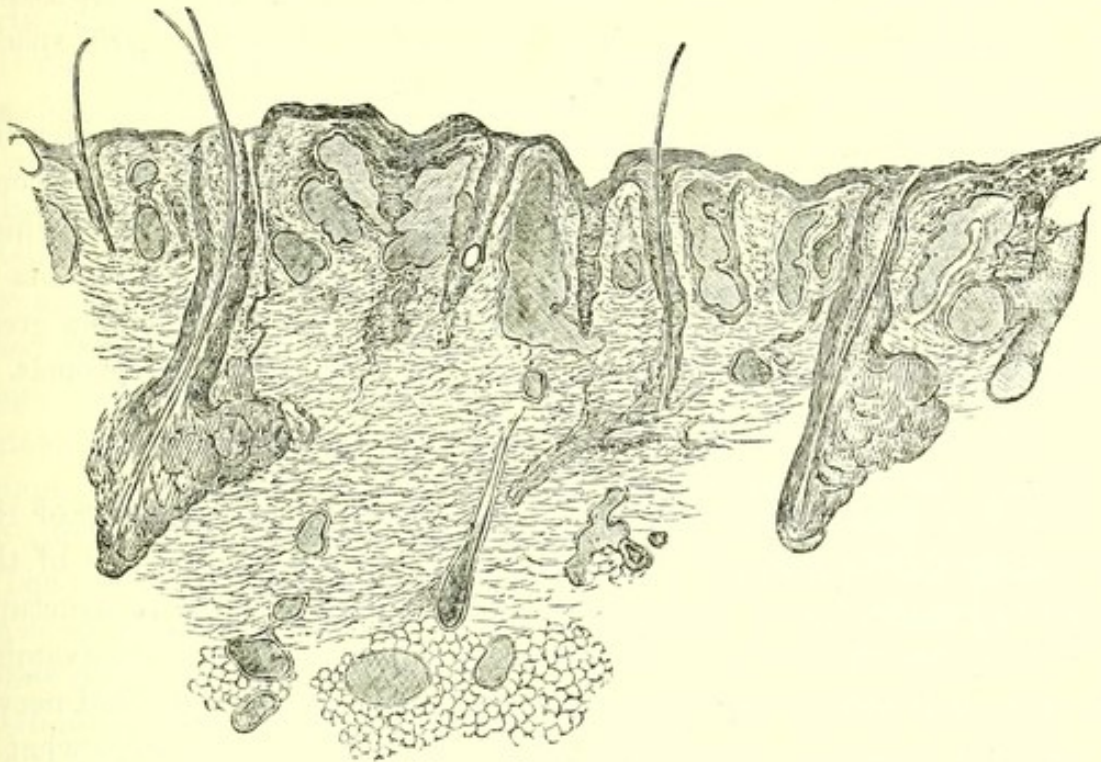


Fig. 76.—Section of skin in a case of diffuse venous naevus. The large sinuses (shaded) are seen to lie superficially between the hair-follicles and glands. (VIRCHOW.)

tumours are usually more or less prominent (*naevus prominens*); they are erectile and sometimes pulsatile. Sometimes the tumour merges into the neighbouring vessels without distinct boundary, but sometimes it has a distinct connective tissue capsule, which, however, appears to be a secondary formation. Sometimes also it is indurated in the centre, and the induration may gradually lead to the obliteration of the spaces and the destruction of the tumour. These tumours are not so usually congenital as the former kind, but they come on in childhood at latest, and they may develop out of the other form.

The skin is a frequent seat of the cavernous angioma, especially that of the face or head, but also sometimes of the trunk or limbs. They are also met with in the liver,

not forming prominent tumours, but simply replacing a piece of liver tissue by cavernous tissue. (See Fig. 77.)

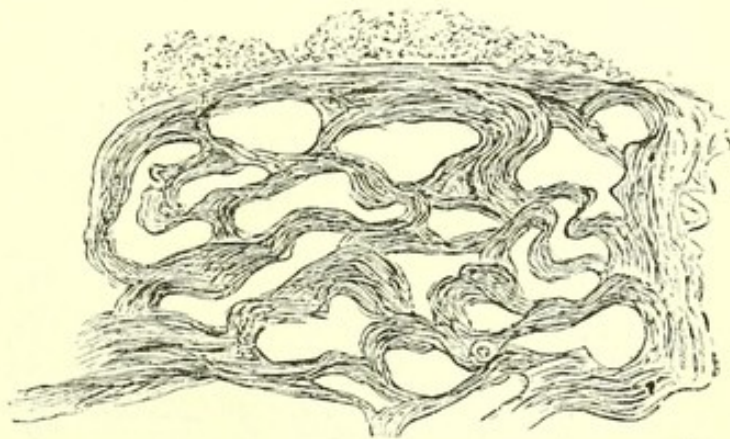


Fig. 77.—Cavernous angioma of liver. The fibrous septa are shown. The spaces between were occupied with blood during life.  $\times 90$ .



(4) The **Lymphangioma** is a vascular tumour composed of lymphatics. Two forms have been distinguished here also, namely, the **Plexiform** and **Cavernous**. The former consists of a congeries of dilated lymphatics, while the latter forms a more definite tumour. Sometimes the spaces dilate so as to form complex cysts.

These tumours are mostly congenital, and they form a considerable proportion of the congenital cystic tumours. The cavernous form seems to be the cause of the congenital enlargements of the tongue, to which the name **Macroglossia** is applied, as well as of similar enlargements of the lips and cheeks. In certain cases of elephantiasis there is a great dilatation of the lymphatics which some regard as forming angiomas.

#### 9.—THE GLIOMA.

This is a tumour with the structure of the connective tissue of the central nervous system, the **Neuroglia**. In examining a section of the brain substance, it is difficult to tell what is really nervous structure, and what the supporting connective substance, but when we examine the surfaces of the ventricles we find that the ganglion cells and nerve-fibres fall away, and just at the surface or ependyma we have what is presumably a purely connective substance. When hardened sections are examined, this is seen to consist of a finely reticulated network of fibres and round or slightly elongated cells. In the fresh state the fibres are not obvious, and we have a granular material. This connective substance has some of the characters of mucous tissue, and seems allied to it.

The glioma as it occurs in the brain does not usually form an

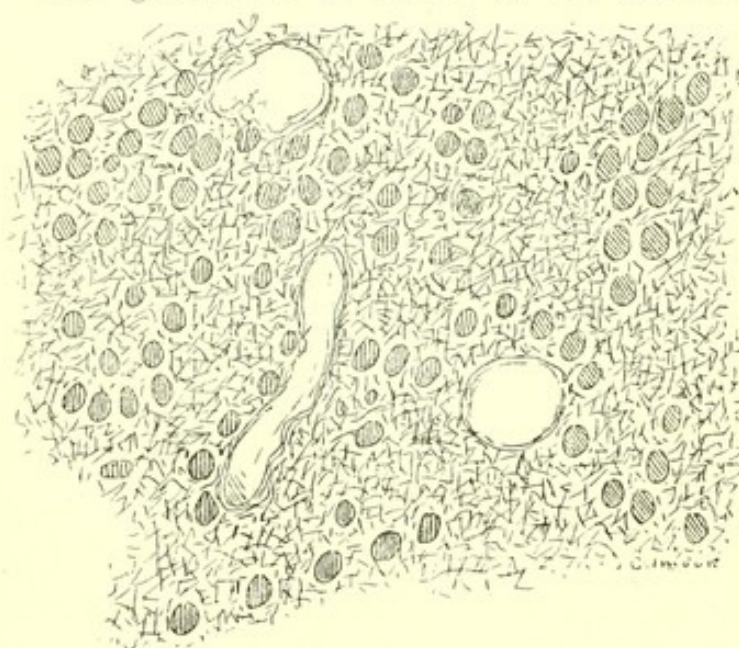


Fig. 78.—Glioma of brain.

isolated tumour, but, being continuous with the brain substance, has more the appearance of a swelling of part of the brain. It is seen also that the different shades of colour of different parts of the brain are lost when a glioma takes their place. Gliomas are usually soft in consistence and grey in colour, so as to resemble brain-sub-

stance, but they sometimes attain considerable density. They



sometimes occur as small granular or warty projections on the surface of the ventricles, but the more important ones involve considerable portions of the brain substance.

Under the microscope the glioma is seen to resemble the neuroglia, but the cells are much more abundant. There is a well-developed fine or coarse net-work and in it cells with oval nuclei. (See Fig 78.) The cells present considerable variety in size.

Gliomas being soft and somewhat cellular tumours, are liable to secondary changes. Hæmorrhage not infrequently occurs, and the blood causing pressure around, the case may end like one of hæmorrhagic apoplexy. The tumour may also undergo fatty or caseous metamorphosis, and if a limited hæmorrhage has occurred the clot may change in a similar way. In this manner a tumour which had originally the appearance of brain substance may change considerably.

The tumour is usually of slow growth and non-malignant, except in the sense that on account of its site it often affects important parts and causes death.

Gliomas occur also in the retina, forming soft tumours which fill up the eyeball. The true glioma is an innocent tumour, but sometimes it assumes a sarcomatous character and malignancy is developed.

#### 10.—THE PSAMMOMA OR BRAIN SAND TUMOUR.

The pineal gland contains calcareous particles like grains of sand, and tumours are met with in which similar particles are present. It is necessary, of course, to distinguish these from tumours in which simply a secondary calcareous infiltration has occurred. The psammoma is composed of soft connective tissue in the midst of which there are calcareous masses in the form of irregular globes, rods or spines. (See Fig. 79.) The commonest form is the globe, which has rounded projections on its surface like a berry. The origin and significance of these masses is obscure. The tumours are met with in the pineal gland, choroid plexus, and brain substance, and are usually small. They also occur in the dura mater, where they form half globular tumours, sometimes as large as a cherry and either smooth or irregular, on the surface.

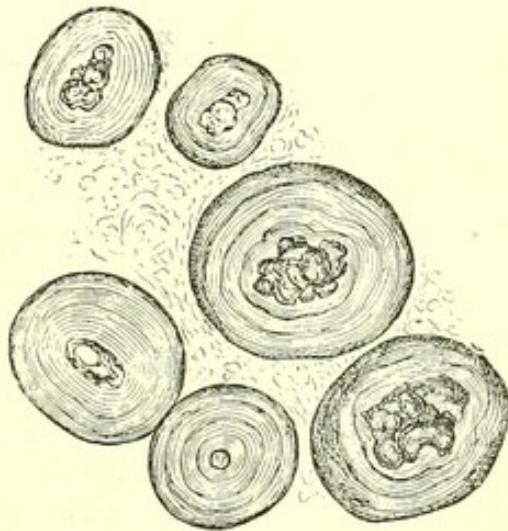


Fig. 79.—From a psammoma of the brain substance. Globular particles of brain sand are visible.



The calcareous particles probably arise by deposition around the new-formed vessels. Besides occurring in the simple psammoma, they are met with in sarcomas, myxomas, and other forms of tumour.

#### 11.—LYMPHOMA OR LYMPHATIC GLAND TUMOUR.

Under this designation are included tumours composed of typical lymphatic gland tissue. Such tumours originating in glands would be scarcely distinguishable from simple hypertrophies of these. It is doubtful whether such simple lymphomas exist. Enlargements of lymphatic glands occur as a result of tuberculosis and syphilis, and in leukæmia, but these all belong to distinct categories. The proper tumours arising in lymphatic glands and having their structure are nearly all infective in character and are described at p. 217.

#### 12.—THE PAPILLOMA OR PAPILLARY TUMOUR.

By this name is meant a tumour composed of a congeries of exaggerated papillæ like those of the skin, or like the villi of mucous membranes. A papilla or villus consists of a basis of connective tissue in which there is a loop of capillary blood-vessel, and a covering of epithelium. The epithelium is like that of the surface concerned, and may be stratified or in a single layer.

Their commonest situation is the skin, where they form the **Wart**, which is an overgrowth of a group of existing papillæ covered with hard epidermis. At the surface of the wart the papillæ may be covered over with a continuous layer of epidermis, or the individual papillæ may project independently. The **Horn** is also formed on the basis of a group of papillæ, but the hard horny epidermis is greatly developed, and forms a consistent outgrowth of considerable dimensions. The **Condyloma** is a syphilitic outgrowth due to exaggeration of the papillæ, with very soft epidermis. These occur near the genital organs mostly.

The ordinary wart is to be distinguished from the **Congenital soft warts** and **Moles**. These are often pigmented and sometimes covered with hairs. In their structure they not uncommonly contain pieces of tissue composed of round or spindle-shaped cells and so differ altogether from the true warts. It is these soft warts and moles which in after life are liable to give rise to sarcomas or cancers.

On **mucous membranes** papillomas may be gathered into local tumours, or cover a considerable surface, giving it a shaggy villous appearance. In the **larynx** (Fig. 80) they often form localized prominent tumours, especially on the vocal cords. They are not uncommon in the rectum. In the **urinary bladder** they are of considerable



importance on account of their tendency to hæmorrhage. In this situation they may form distinct tumours with long branched papillæ, or there may be a large surface which is simply villous in appearance. The papillæ are covered with delicate epithelium, and are liable to severe and frequent hæmorrhage.

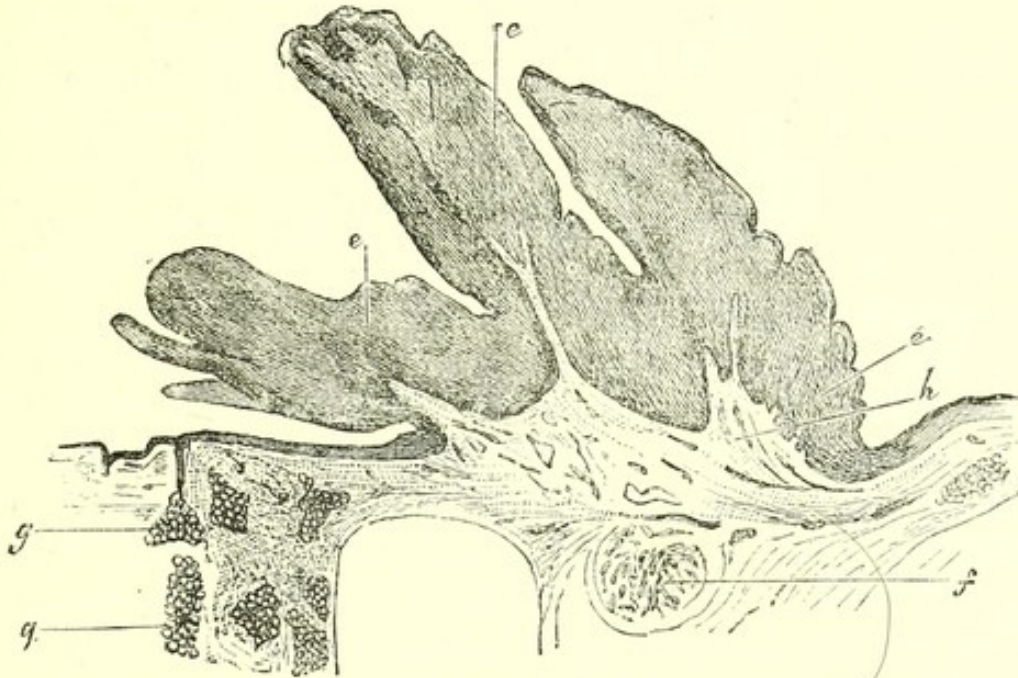


Fig 80.—Papilloma of larynx : *e*, epithelium ; *h*, connective tissue ; *g*, mucous glands ; *f*, an atrophied gland.  $\times 20$ . (CORNIL and RANVIER.)

The **pacchionian bodies** of the encephalon are really papillary formations, and Dr. Cleland has described tumours of this region which seemed to arise by extreme hyperplasia of these papillæ.

**Literature.**—CLELAND, Glas. Med. Jour., 1861.

### 13.—THE ADENOMA OR GLANDULAR TUMOUR.

As there are many glands of different structure, so are there various forms of adenoma. The several forms will be more particularly described under their respective organs, and it will only be necessary here to enumerate them. We have the mammary, the prostatic, the thyroid, and the hepatic adenomas. Besides these, the so-called mucous polypi, although often formed of hypertrophied mucous membrane, frequently contain glandular tissue which is apparently, in some cases, new-formed. Again, in ovarian tumours there is frequently a newformation of gland-tissue out of which cysts develop, so that the tumour is called an adenocystoma.

The term adenoma is not commonly used for tumours composed of lymphatic-gland tissue, although the expression lymph-adenoma is not infrequently employed. As the structure of lymphatic glands differs so markedly from that of the epithelial



glands it is perhaps better to reserve the term adenoma for the latter, and to use that of lymphoma for the former.

A rather unfortunate custom of calling lymphatic tissue "adenoid tissue," without specifying that it is lymphatic, has become somewhat prevalent. It would conduce to clearness if the term lymphoid tissue were used instead.

#### 14.—THE CYSTOMA OR CYST.

A cyst is a cavity having well-defined walls and containing material which is more or less fluid. Some cysts are formed simply by the transformation of existing structures, and it is clear that according to our definition of tumours these do not come under that designation, as they are not pieces of tissue which grow independently without regard to the needs of the organism. There are other cysts, however, which do owe their origin to the independent newformation of tissue, and therefore deserve the name of tumours. The two forms are so similar in appearance that it would be inconvenient to separate them rigidly, and without aiming at logical accuracy we shall here consider the cysts as a whole. If necessary the name Cystoma may be reserved for strictly cystic tumours. With such various origins cysts vary greatly in structure, contents, and size. The structure of the wall may be exceedingly simple or very complex. The cyst may be composed of a single cavity (unilocular) or of several sacs united (multilocular). The different sacs will often run together so that their number will be reduced, or a multilocular cyst may become unilocular. When this is the case there are usually remains of the former partitions visible inside the cyst.

In this general survey of cysts their mode of formation is the most important consideration. This will be more fully considered here, while the local varieties will be described under the special organs.

(1) **Cysts originating from pre-existing cavities or canals.**—Cysts belonging to this group will mostly arise by the accumulation of the normal contents in the existing cavity or canal. Most of them arise in connection with glands possessed of excretory ducts, and they originate by obstruction of the duct and consequent accumulation of the contents; but some are formed in glands without such ducts, and arise by reason of an abnormal accumulation of contents. The former have been named retention cysts by Virchow.

**Retention cysts** are formed as the result of obstruction of gland ducts, and the accumulation occurs either in the duct itself or in the acini of the gland. Examples of this on a large scale are furnished by cases in which a main duct is obstructed. Thus obstruction of the ureter may lead to the conversion of the kidney into a large cystic cavity (hydro-



nephrosis). (See Fig. 81.) Obstruction of the cystic duct may cause the gall bladder to form a large sac; and closure of the orifice of the vermiform appendage may lead to the formation of a large cyst.

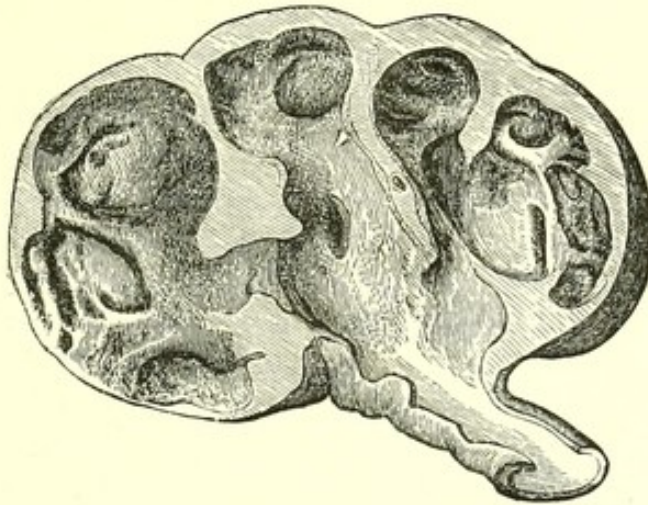


Fig. 81.—Hydronephrosis with granular atrophy of the kidney. The other kidney showed a marked compensatory hypertrophy. (VIRCHOW.)

Examples of retention cysts on a smaller scale are very frequent. **Cysts of the skin** frequently arise from sebaceous glands, and are called **Sebaceous** or **Atheromatous cysts**. They form chiefly on the scalp where they constitute the common **Wen**. Their contents consist of fat, cholestearine and epidermic scales, and their walls are formed of connective tissue with a lining of epidermis, which may form a very thick layer and may present papillary projections.

**Mucous cysts** also form an important group of retention cysts. They occur in situations where mucous glands are present, their chief sites being the nostrils and communicating cavities, the upper surface of the epiglottis, the larynx, the œsophagus, and in connection with the glands of Cowper and Bartolini.

The origin and mode of formation of the mucous cyst has been carefully studied by Recklinghausen. It has long been taught that these cysts arise by obstruction of ducts and accumulation of the contents behind them, but this scarcely explains the continuous enlargement of the cysts after they have been formed. It might be supposed that the epithelium lining the cysts would secrete mucus and so add to the contents, but there is the objection to this that the epithelium loses its glandular character, and there are no goblet cells to be found in it. The real state of matters appears to be that the cyst is formed, not out of the gland, but from its duct, and, as the gland persists, its secretion is thrown into the cyst. This is shown in Fig. 82, copied from Recklinghausen's paper. It would thus appear that the persistence of the gland is an essential element in the formation of the cyst. In the enlargement of the cyst the power which mucin has of swelling up and absorbing water is of some consequence. A small amount of mucin formed by the gland and discharged into the cyst will swell up and add considerably to its contents. Before a regular cyst forms, the orifice of the duct is obstructed, usually by an inflammation



around it. But on account of the peculiarity of mucin just noticed, a small temporary cyst may form without any considerable obstruction of the duct. If a quantity of mucin is discharged into the duct it may swell so much as to be unable at once to escape from the orifice and so form a small cyst, which afterwards discharges. In this way cysts often form in the mucous membrane of the mouth.

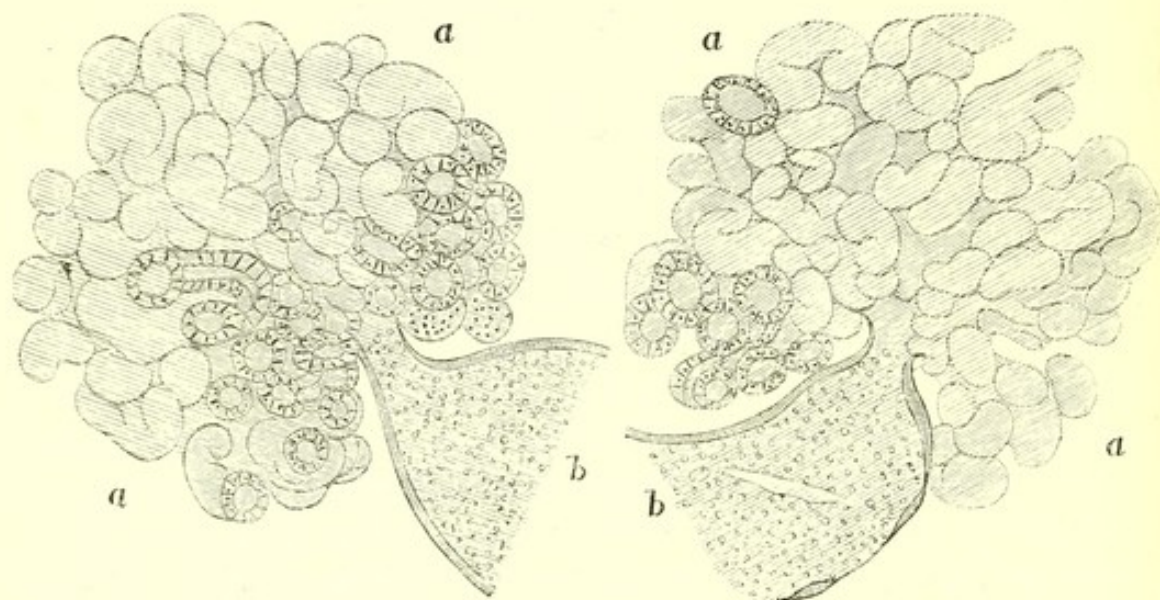


Fig. 82.—Formation of mucous cysts; *a*, the persistent mucous gland tissue; *b*, the dilated duct.  $\times 300$ . (RECKLINGHAUSEN.)

It has already been noted that cysts arise in mucous polypi, and they have a similar mode of formation to that just described. The obstruction of the orifices is here the more likely, as the polypus itself usually originates in connection with a chronic inflammation of the mucous membrane. Retention cysts are also of frequent occurrence in the larger glands, especially the mamma and kidney, but also the liver. In these there is obstruction of the finer ducts or tubules due usually to inflammation in the connective tissue around. As the inflammation affects the gland generally the resulting cysts are multiple, and usually of small size.

Besides the retention cysts we have cysts arising by **Dilatation of pre-existing cavities** where there is no excretory duct. A typical example of this is afforded by the ovary, in which cysts frequently originate by the enlargement of the Graafian vesicles. This cystic formation is often associated with chronic inflammation of the organ, the increase of connective tissue, especially in the capsule, preventing the usual rupture of the vesicles. In this sense there may be an analogy with the retention cysts. Another example is afforded by the thyroid gland, where the colloid degeneration in the saccules may lead to the formation of cysts, which enlarge both by accumulation of their contents and coalescence of neighbouring ones.

(2) **Cysts of independent origin.**—This group includes a considerable number of different forms.

**Dermoid cysts** are very complex structures, their walls consisting of tissues similar to those of the skin and its appendages, but frequently



associated with still more diverse tissues. The simplest of them are met with in the skin, where they resemble atheromatous cysts, but their wall is much nearer in structure to the skin, and usually contains growing hairs. They are usually congenital, arising by inclusion of portions of foetal skin. Their situations suggest this origin, being commonly at places where in the process of development fissures have united; thus the commonest seats are the outer corner of the eyebrow, the neck, and the mediastinum, where they apparently arise from imperfectly closed branchial fissures. The more complex dermoid cysts are found mostly in the ovary, where they arise from the ovum apparently by virtue of its formative power. They are frequently very complex in structure, their wall containing not only skin with hairs and sebaceous glands, but bone, cartilage, teeth, etc. Similar cysts have been met with in the testicle, brain, orbit, lung, peritoneum, and elsewhere. These complex dermoid cysts may be classed along with the Teratomas.

Cysts of the ovary frequently arise, as we have seen, from glandular tissue, so that they have their origin in **Adenomas**. These tumours will be fully described in the section on diseases of the ovaries. The gland tissue has no ducts, and as the epithelium produces a mucous or other secretion there is a great distension resulting in the formation of cysts. Sometimes the gland tissue is rather in the form of villous or papillary processes, the cysts forming by the union of these processes and accumulation in their recesses.

Cysts sometimes arise in connective tissue without apparent cause. Some of these may take origin from the lymphatics, and some from the remains of branchial fissures (see p. 47).

**Extravasation cysts** result from the changes which may occur after hæmorrhage. This is exemplified in the case of the **apoplectic cyst** where the blood-clot and softened brain substance are replaced by connective tissue containing fluid in its meshes. Such cysts may occur from softening of the brain without hæmorrhage.

**Parasitic cysts** are really connective tissue capsules formed around parasitic animals. The most important example of this is the so-called hydatid cyst which occurs in connection with the *tænia echinococcus* and sometimes attains large dimensions.

**Cysts in tumours** usually originate by dilatation of glandular structures and hence are most common in adenomas. They also occur not infrequently from softening of the tumour tissue. This is especially the case in large tumours and in those whose tissue is already comparatively soft. All forms of tumours when they grow to a large size are liable to have cavities in their central parts due to softening. On



the other hand the soft sarcomas often contain cysts without attaining to large dimensions. These latter cysts often contain blood, and indeed the tumour may present little more than the characters of a cyst filled with blood. **Blood-cysts** are apparently for the most part sarcomas which have undergone this process. (Godlee.)

**Literature.**—VIRCHOW, *Geschwülste* i., Lecture ix.; BUTLIN, *Internat. Ency. of Surg.*, iv. 655; RECKLINGHAUSEN, *Virch. Arch.*, lxxxiv. 425; GODLEE, *Path. Trans.*, xxvii. 270.

#### 15.—TERATOMA.

This name has been applied by Virchow to tumours into the structure of which a number of very different tissues enter. The name is derived from *τέρας*, a monster. The tumour contains various structures of the body as if from an ill-arranged foetus. Thus we may have in such tumours skin, bone, muscle, glands, nervous tissue.

Some of these tumours arise by the abnormal inclusion of a portion of a double monster (see p. 39). They are most frequently met with in positions where double monsters are usually attached to each other, namely, in the sacral region or further up the back, and in the head and neck. They are also somewhat frequent in the ovary where they may arise by an irregular developmental process, the ovum endeavouring to develop some of the perfect tissues of the body.

The **Sacral teratomas** and **those of the head and neck** are the more typical ones. They frequently contain pieces of bone which simulate the bones of the foetus, also brain substance and muscle.

The teratomas of **internal organs** do not usually represent such complete systems or so many different tissues as the sacral. They are most frequent in the ovaries, but also occur in the peritoneum, testes, lungs, and so on. The dermoid cysts already considered are the chief representatives of this group in internal organs.



## SECTION IX.—CONTINUED.

## B.—ATYPICAL TUMOURS.

- I. SARCOMA. *Definition. Structure; cells, intercellular substance, blood-vessels. Place of origin. Mode of growth; local malignancy and metastasis. Changes in structure; indurations, ossifications, cystic formation, ulceration, etc. Individual forms. 1. Round-celled, 2. Spindle-celled, 3. Giant-celled, 4. Pigmented, and 5. Plexiform sarcomas. Other forms described.* II. CARCINOMA OR CANCER. *Definition. Structure; cells, stroma, blood-vessels, lymphatics. Origin; from existing epithelium, shown in many cases, hence its localities where epithelium exists. Influence of age and sex. Growth and extension; frequently by infiltration; ulceration; secondary tumours in lymphatic glands; generalization by implantation of grafts; usual seats of secondary tumours; their large size; their mimicry of primary tumours. Retrograde changes; fatty degeneration, chiefly. The local nature of cancer. Individual forms. 1. Flat-celled epithelioma, including Rodent ulcer, 2. Cylinder-celled epithelioma, 3. Soft or medullary cancer, 4. Scirrhus, 5. Colloid cancer, 6. Melanotic cancer, 7. Mucous cancer, 8. Endothelioma.*

## I.—SARCOMA.

IN its literal meaning this term simply indicates a fleshy tumour, and it was formerly applied in a very indefinite way. Under the influence of Virchow, however, it has come to include a group of tumours, which, though in certain respects differing in structure, yet present such features in common that they form a consistent class of themselves.

**Definition.**—The sarcomas may be defined as tumours which originate in one or other of the forms of connective tissue, but differ in structure from their mother-tissue chiefly in respect that the cells greatly preponderate over the inter-cellular substance, and also that they frequently differ greatly in size and shape. This definition includes the origin of the tumour, and refers also to the atypical character of the structure. It may be added that in its mode of growth the sarcoma is also atypical, having the characters of malignancy already referred to.

**Structure.**—Sarcomas have been somewhat aptly compared in structure to inflammatory newformations. We have already seen that the



tendency of these, as exemplified in the granulating wound, is to develop into connective tissue. The round cells pass into spindle cells, and then the connective tissue develops out of the latter. The round and spindle cells may thus be regarded as the preparatory, or in a certain sense embryonic stage of connective tissue. In sarcomas we have tumours composed of round cells, and tumours composed of spindle cells, without any tendency to further development, as if the embryonic form had been stereotyped for the whole life of the tumour. Besides these forms sarcomas sometimes contain giant-cells (myeloplakes). We know that cells of this nature occur normally in growing bones where, as Wegener and Kölliker have shown, they exercise an important function (osteoclasts), and they are not unknown in granulation-tissue, even apart from bone. They also are to be regarded as connective tissue structures, and as belonging to a developmental stage of connective tissue.

From the definition given above of the sarcoma it will be apparent that the tumours included in this class will vary very greatly in structure and other characters. They will vary in the first place according to their tissue of origin, because, although in general following the type of granulation-tissue, yet they usually carry with them, especially in the characters of the intercellular substance, some indications of the mother-tissue, so that it might be possible to distinguish as many forms of sarcoma as there are typical tumours of the connective tissue series. But even when derived from the same tissue the tumours may vary according to shape, size, and abundance of cells, so that still greater complexity is thus introduced.

In the diagnosis of actual cases reference should be made to origin as well as to structure and mode of growth.

Sarcomas differ according to the form of cells and character of intercellular substance. So far as **the Cells** are concerned the chief forms are those already mentioned, namely, round, spindle-shaped, and giant-cells. The tumour is usually homogeneous in its structure, at least in its recent parts where not altered by degenerations or further developments, and so it is generally possible to distinguish sarcomas according as they are round-celled, spindle-celled, or giant-celled. There are, however, cases in which the cells are variously shaped, partly round and partly spindle-shaped.

As the characters of the individual forms of connective tissue are determined by their **Intercellular substance**, so in sarcomas the shape of the cell does not bear a constant relation to the character of the intercellular substance. Sarcomas are sometimes named according to their relations to the various connective tissues, thus **Fibrosarcomas**,



**Myxosarcomas, Chondrosarcomas,** and so on. We even have tumours of striated or smooth muscle which take on the sarcomatous mode of growth, and are hence called **Myosarcomas**.

The **Blood-vessels** of sarcomas are generally rather thin-walled, and they run usually in immediate contact with the tumour tissue. In some cases they are supported by complete or rudimentary connective tissue, which may divide the tissue into alveoli, and so give rise to an appearance resembling that of cancer (*alveolar sarcoma*). Sometimes the sheath of the vessels undergoes a peculiar transformation into a hyaline substance, which forms a mantle round the vessels, and gives a plexiform character to the tissue (*plexiform sarcoma, cylindroma*.)

**Pigmentation** is not infrequent in sarcomas, especially in those arising in pigmented situations such as the skin and eyeball. The pigment is usually in the cells but may be in the intercellular substance.

**Place of origin of sarcomas.**—As connective tissue is of nearly universal occurrence sarcomas may arise in almost any situation. They originate, however, for the most part where connective tissue in some form is abundant. Thus the bones, the skin, the mamma are frequent seats of origin. They are also common in the testicles and the brain, but are rare as primary tumours in other glands, in the muscles and in the lungs.

The place of origin has an important influence on the structure of the tumour and on the transformations and degenerations to which it is liable.

**Mode of growth and transformations of sarcomas.**—The sarcoma grows by multiplication of its own elements, and it is often surrounded by a capsule so as to be apparently self-contained. Even when so delimited, however, it generally presents the characters of local malignancy. The tumour grows along the existing connective tissue of the part, and outside the apparent boundaries, even outside the capsule, there are already the multiplying cells of the tumour.

This was very characteristically observed by the author in a case of pigmented sarcoma, where the pigmented cells were plainly visible in the connective tissue outside the mass of the tumour.

The tumour thus growing along the connective tissue sometimes moulds itself on the structure, causing atrophy of the special constituents of the tissue which it replaces.

Besides this local malignancy, sarcomas frequently give rise to **Secondary tumours** at a distance. As a general rule the lymphatic glands are not secondarily affected, but in sarcomas of the foot, the tonsil, the testicle, and probably the kidney, they are liable to be involved (Butlin.) This fact implies that either in their origin or in



process of growth the sarcomas of these localities come into relation with the lymphatic vessels. Either with or without an intermediate affection of the glands the infection is liable to extend by the blood first to the lungs, which are the most frequent seats of such metastatic growths, and then it may pass on by the systemic circulation to a large number of different situations. The secondary tumours repeat exactly in structure and mode of growth the primary one so that we may have, in the lungs or elsewhere, typical spindle-celled tumours, or even cartilaginous or ossifying sarcomas.

The sarcomas are somewhat liable to **metamorphoses and transformations**. The soft and quickly growing ones are specially prone to fatty degeneration and softening, so that cysts may form in this way. Then the tissue may show a tendency to develop into the mature tissue of its kind. Thus a fibrous development may occur in the sarcomas of membranes, or a partial formation of cartilage or bone may take place, and this tendency may be so marked as to give a distinctive character to the tumour, so that we may speak of an **Ossifying, Indurating, or Calcifying sarcoma**. This peculiarity may give rise to mistakes in diagnosis if a part of the tumour which has undergone transformation be examined. The tissue thus formed is not usually quite typical, the cells being as a rule more numerous and larger, hence it is always proper to seek for the growing margin of the tumour for examination.

Besides this the sarcomatous tissue may be **mixed** with other tissues. **Gland-tissue** is most frequently thus associated, especially in the mamma and testicle. It is sometimes difficult to determine whether the gland-tissue found in mammary tumours is simply the remains of the gland or new-formed. The presence of gland-tissue is important as it frequently, by dilatation, gives rise to **Cysts**. These cysts frequently give a special character to the sarcomas, especially in the mamma and testicle, the association being indicated by the term **Cystic sarcoma**. The tumour tissue often grows into the cysts, forming the so-called **Intracystic growth**, and often giving a peculiar character to the tumour when divided by the knife.

Sarcomas coming to the surface usually incorporate the skin, converting it into their own tissue, and then **Ulcerate**. The ulcer is thus formed of the tumour tissue, which may be excavated by softening, or may pout outwards into a fungating mass, liable to bleed (*Fungus hæmatodes*).

**The individual forms of sarcoma.**—The most convenient division of sarcomas is according to the form and other characters of their cells. While such a division is adopted, however, it must be remembered that tumours occur in which various forms of cells are present.



1. **The round-celled sarcoma** (Fig. 83) is also called the granulation-sarcoma and the encephaloid sarcoma. It is composed of round or slightly oval cells generally about the size of white blood-corpuscles but sometimes much larger. It is an exceedingly soft tumour, often half diffuent, and has usually a grey medullary appearance. Its blood-vessels are in the form mainly of large capillaries whose walls are embryonic in structure and often present varicose or aneurysmal dilatations. The vessels are liable to rupture, and so these tumours often present interstitial hæmorrhage. Between the cells there may be some intercellular substance. This is sometimes homogeneous and becomes opaque with acetic acid (Myxo-sarcoma), or it may be somewhat fibrous or reticulated.

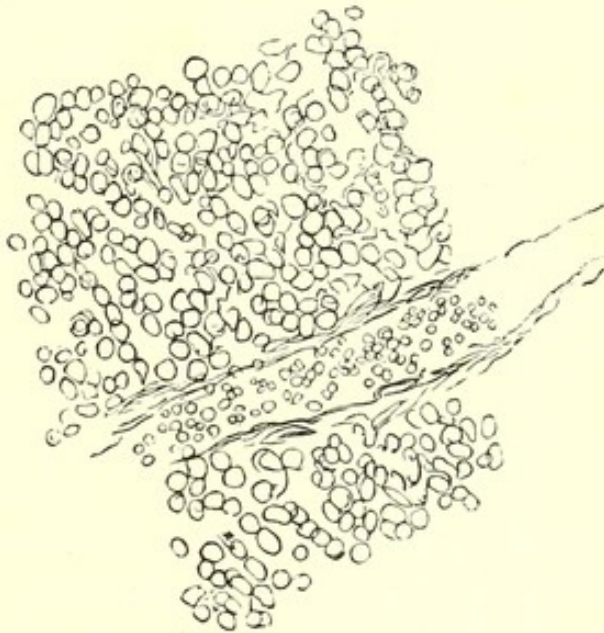


Fig. 83.—Section of a round-celled sarcoma of the brain. A thin-walled vein with blood-corpuscles in it, divides the section.  $\times 175$ .

This form of sarcoma is met with in the skin where it may originate in a congenital soft wart or mole; in the subcutaneous tissue; in the bones—forming the majority of the so-called medullary cancers of bone; in the muscles; in the glands—especially the mamma and testicle; in the brain and elsewhere. Being a soft tumour with delicate vessels it more readily produces secondary tumours by metastasis than other sarcomas. It is also usually a tumour of rapid growth and commonly imperfectly delimited from the surrounding tissue. The malignant lymphoma is sometimes described under the name of Lympho-sarcoma (see p. 217), as if it were a form of round-celled sarcoma.

2. **The spindle-celled sarcoma** (Fig. 84) is also called the fibro-

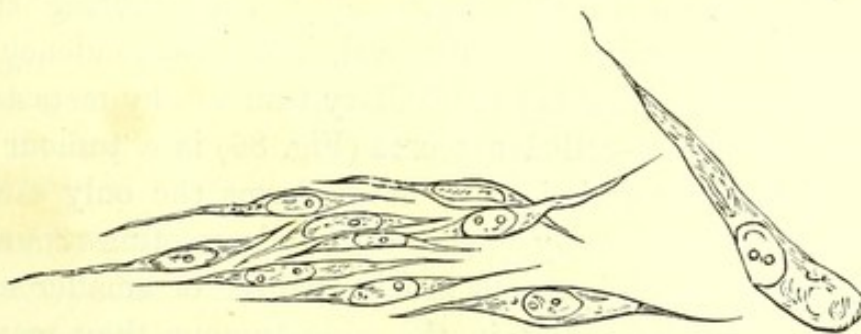


Fig. 84.—Large cells from a spindle-celled sarcoma.  $\times 300$ .

sarcoma and corresponds with Paget's class of recurrent fibroids, and



Lebert's group of fibro-plastic tumours. The cells are spindle-shaped like those in the deeper layers of a granulating wound, and there is comparatively little intercellular substance, the tumour being mainly composed of masses of spindle-cells. There is always, however, a certain amount of intercellular substance, which consists of fibres generally of some density. The cells are arranged in bundles which to a certain extent interlace like the bundles of fibre-cells in the myoma. There are great varieties in the size of the cells, some tumours being composed of very small cells, and these are usually soft, while at the opposite extreme are cases where the spindles are gigantic. When viewed in mass the individual spindles may not be apparent, but they are usually easily isolated, unlike the fibre-cells of the myoma. These tumours, except the small-celled forms, are usually

firmer than the round-celled sarcomas, and may even approach the fibroma in hardness. Many of them show a tendency to more complete organization into fibrous tissue, cartilage, or bone.

The spindle-celled sarcoma occurs frequently in the periosteum, and in that case is firmly attached to the bone. It is also met with in or under the skin (see Fig. 85), in muscles, in the testicle, etc. It is a frequent tumour in the mamma, and here it not infrequently forms the adenoid sarcoma and the cystic sarcoma.

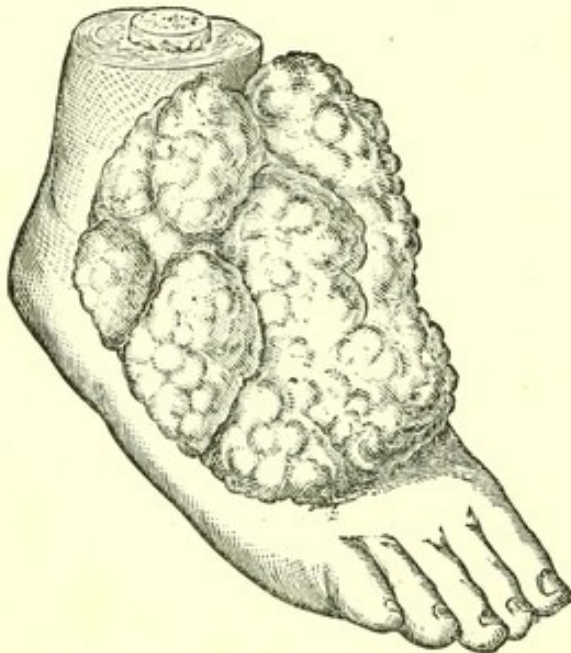


Fig. 85.—Large fungating spindle-celled sarcoma of the foot. (VIRCHOW.)

The spindle-celled sarcoma is usually a distinctly defined tumour but often, as in the case of the periosteal form, its boundaries are not defined, and it advances by incorporating neighbouring structures. Although prone to return after removal, it has less tendency than the round celled form to give rise to secondary tumours by metastasis.

**3. Myeloid or Giant-celled sarcoma** (Fig. 86) is a tumour in which the giant-cell is characteristic, but never forms the only sarcomatous element, there being generally spindle cells and sometimes round cells in great abundance. The giant-cells are in greater or smaller number in proportion to the others, and in the same tumour they may present various proportions in different parts. The tumour tissue is soft and very often of a brown colour. Cysts not infrequently develop by softening.



The myeloid sarcoma occurs in connection with bone, and most frequently grows from the medulla. This is especially the case with the long bones, where the tumours originate in the cancellated tissue at the extremity—the most usual situation being the lower end of femur or upper end of tibia. (See under Diseases of Bone.)

Myeloid sarcoma is also met with outside bones, growing from the periosteum, especially of the jaws. Many tumours, to which the name **Epulis** is given, are myeloid sarcomas.

The myeloid sarcoma is, for the most part, slow of growth and does not usually produce secondary tumours.



Fig. 86.—Giant-celled sarcoma.  $\times 175$ .

4. **Melanoid or Pigmented sarcoma** always originates in a situation where pigment already exists, the eye or skin. The cells of which it is composed are usually spindle-shaped, but may be round, and from the first they tend to the aggregation of brown or black pigment in their substance. The pigment aggregates first around the nucleus, but is very irregular in its distribution. In a melanotic tumour there may be bits unpigmented, and even in the pigmented parts some cells are free from pigment.

The melanotic sarcomas have a great tendency to metastasis, and as the material is conveyed by the blood there are pigmented tumours formed in a great variety of organs and tissues where they may grow to great dimensions, though the original tumour may be very small.

It is necessary to distinguish from these proper melanotic sarcomas those which become pigmented from blood. In the former the pigment is brown or black from the first, being obviously elaborated by the cells. In the latter the pigment is red or yellow, and the pigmentation may be related to a special weakness of the vessels allowing of hæmorrhage.

The term **Chloroma** has been applied to a form of round-celled sarcoma of a peculiar greenish yellow or grass-green colour. It occurs primarily in the periosteum of the face and head, and may lead to secondary tumours of similar colour in the liver, kidneys, etc. The colour is due to small refracting granules which appear to be composed of fat.

5. **Plexiform sarcoma or Cylindroma** is a name applied to a form of tumour whose relations are somewhat obscure, and it probably includes more than one kind. The peculiarity of the tumour is the existence of cylinders and rounded structures having a hyaline character, like mucous tissue. In the centre of the cylinder there is often a



blood-vessel, so that the hyaline material clothes it like a mantle. Then, between the cylinders of hyaline material there are frequently masses of cells which may form long processes so as to give a close resemblance to cancer, to whose cells these may also conform in general appearance.

The origin of these cylinders is not perfectly clear. In some cases it may be that we have a combination of sarcoma and myxoma, but this does not account for the peculiar form of the cylinders. A more probable explanation is that the cylinders arise by hyaline or mucous degeneration of the adventitia of the blood-vessels, and this is confirmed by the fact that they are often arranged around the vessels. If this be the proper view, then, in the tumour, the formation of blood-vessels is an important element. In this way we should have a sarcoma in which a peculiar transformation occurs in the external coats of the vessels. It is on this view that the name *plexiform angiosarcoma* is applied to this form of tumour.

The tumour as a whole is often of a gelatinous appearance, or it may be that the gelatinous material is seen to be in separate spaces throughout the tumour. It occurs in the orbit and its neighbourhood, or the upper and lower jaws; it may form part of the constituents of tumours of the parotid, and it is also found in the brain and its membranes and the peritoneum, where it may grow to a large size.

Besides these forms of sarcoma several others are sometimes distinguished and designated by special names. Thus we have **Alveolar sarcoma** in which the cells, which are generally round and frequently large, are arranged in loculi, so that both in the characters of the cells and their arrangement there may be a resemblance to cancer. The place of origin and the existence of a delicate reticulum, which is generally present between the cells, are generally sufficient to make the diagnosis clear.

The **Osteoid chondroma** and **Osteoid sarcoma** are closely allied forms of tumour of bone. In both there is a great tendency to the formation of osseous tissue, often of imperfect structure. The recent and growing parts of the tumours are composed of cells like those of ossifying cartilage, or of spindle-shaped or stellate cells with stiff fibrous intercellular substance.

The **Psammoma** is often regarded as a variety of sarcoma. No doubt spindle-celled sarcomas sometimes contain calcareous particles such as these already described. With less justification the **Glioma** is sometimes included with the sarcomas. The term **Endothelioma** is used by some to designate tumours which have a structure indistinguishable from carcinomas, but arise in presumably non-epithelial structures. We prefer to include them among the cancers where they will be described.

**Literature.**—VIRCHOW, *Geschwülste*, 1864-65, ii. 170; PAGET, *Lect. on surg. path.*, 3rd ed., 1870, p. 544; BILLROTH, *Lect. on surg. path.*, Syd. Soc. transl., 1878, ii. 401; BUTLIN, *Internat. Encycl. of Surg.*, 1884, iv. 600, and *Sarcoma and Carcinoma*, 1882; BIZZOZERO, *Wien. med. Jahrb.*, 1878, p. 4; ACKERMANN, *Volkman's Vorträge*, Nos. 233, 234, 1883; HUBER (*Chloroma*), *Arch. d. Heilk.*, xix., 1878, p. 129; CHIARI (*Chloroma*), *Zeitschr. f. Heilk.*, iv., 1883. *Cylindroma*—BILLROTH, *Die Entwick. der Blutgefässe*, 1856; *Virch. Arch.*, xvii. 364; SATTLER, *Ueber die sogenannte Cylindrome*, 1874; EWETZKY, *Virch. Arch.*, lxix.; WALDEYER, *do.*, lv.; FRIEDLÄNDER, *do.*, lxvii. . . . OPPENHEIMER (*Formation of pigment*), *Virch. Arch.*, cvi. 515.



## II.—CARCINOMA OR CANCER.

The term cancer is a clinical one, expressing the malignant characters of the tumour. Like sarcoma it was formerly applied in a general way, and included most sarcomas. The delimitation of the sarcomas by Virchow has led to a stricter definition of the cancers.

**Definition.**—The carcinoma is a tumour taking origin in epithelium and having an epithelial structure, but in the arrangement of the structure and in its mode of growth presenting atypical characters.

**Structure.**—The **Cells of cancers** are epithelial in origin and structure, but differ according to the form of epithelium from which they are derived. Thus we have flat or pavement cells, cylindrical cells, and glandular epithelial cells. Again, the cells do not always correspond strictly with the typical cells of the same kind, but as they are produced in great numbers, they often present great varieties in shape and size, yet they are distinguished usually by their large oval nuclei as epithelial cells. This applies especially to the glandular forms. The various forms of cells will be again referred to in describing the varieties of cancers. The cells always grow in larger or smaller masses, lying close together so as to form the so-called "cell nests," which are characteristic of cancers.

**The Stroma** encloses the cell masses, and supports the blood-vessels necessary to the nourishment of the tissue. This stroma may be merely the connective tissue of the part in which the carcinoma is growing, perhaps increased by a process analogous to that of chronic inflammation as indicated by the presence of round cells in large numbers. In other cases the stroma is as much a new-formed tissue as the epithelium, and it may occur in the form of a well-constructed net-work composed of connective tissue. (See Fig. 87.)

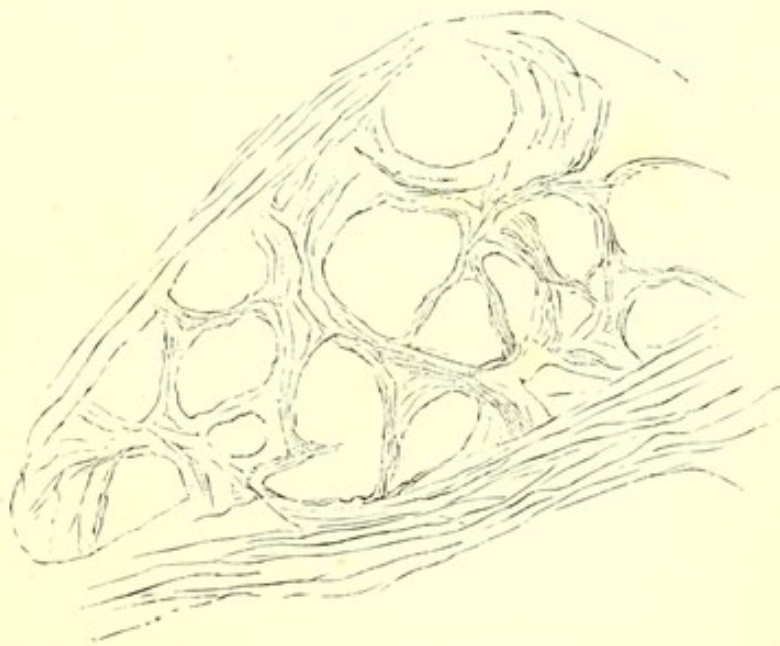


Fig. 87.—Stroma of a soft cancer of the mamma after the cells had been washed out.  $\times 82$ .



Where a cancer produces a well-formed stroma along with the epithelial masses, it will probably grow more readily into a distinct tumour than where the epithelial masses depend for their nourishment on the existing blood-vessels. In this latter case the cancer will commonly present more the characters of an infiltration of the tissues with epithelial structures, and this infiltration may be associated with such irritation as to lead to considerable newformation of connective tissue, giving sometimes a considerably fibrous character to the structure, as in scirrhus cancer.

The **Blood-vessels** in cancers run, as has been indicated, in the connective tissue stroma. They consist of wide capillaries with the usual arterial and venous connections. The great tendency which cancers present to extend by the lymphatic system suggests some special structural connection with the **Lymphatic vessels**. According to Cornil and Ranvier such a connection can be demonstrated by injection. If a cancerous tumour, before being laid open, be punctured with the needle of a hypodermic syringe and a watery solution of Prussian blue be injected, the material first runs into the alveoli around the puncture, mapping out, as it were, a series of cavities, and then passes on into the lymphatic vessels, issuing by their extremities divided in removing the tumour.

**Origin and Locality of carcinoma.**—In regard to their place of origin it may be said that cancers always arise where epithelium or endothelium is normally present, and there seems no doubt that the epithelium of the cancer takes origin in the similar cells of the normal tissue.

Virchow, although distinguishing sarcoma from cancer as a connective tissue tumour, asserted that the cancerous tissue takes origin in connective tissue. The great authority of Virchow has caused this view to be perpetuated more than it otherwise would have been. It was controverted first by Thiersch who showed that in epithelial cancer the cells can be seen to originate from the epidermic cells. Waldeyer, in a series of very elaborate papers, went over most of the seats of cancer and showed that in these the cancer tissue is directly derived from the existing epithelium.

In many situations the actual **Connection with the existing epithelial structures** can be traced, especially if the growing edge of the carcinoma be examined. Thus in a section of an epithelioma of the lip or tongue the cylinders of epithelium which form the essential constituents of the tumour, can be often traced into direct connection with the malpighian layer of the epidermis. (See Fig. 88.) Again, in some cases of primary cancer of the kidney it can be seen that the tumour is arising by direct transformation of the kidney tissue. The primary cancer of the kidney is not a tumour added on to the kidney, but it is generally a portion of, or sometimes the whole kidney, which has undergone an enormous enlargement while keeping its general shape ;



it is in fact the kidney or portion of kidney transformed. And when we examine the marginal parts of such a tumour we find the epithelium of the uriniferous tubules in an active state of germination, the

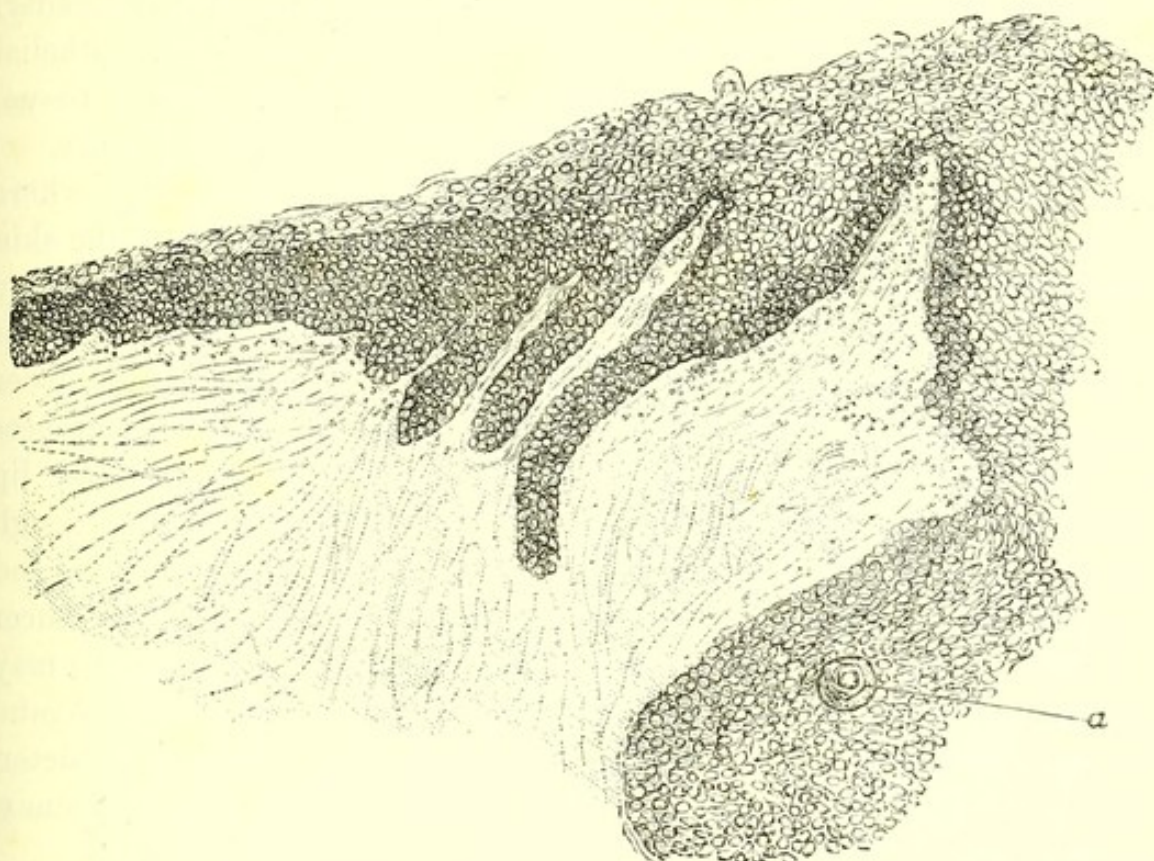


Fig. 88.—From the growing margin of an epithelioma of the skin. The interpapillary processes of epidermis penetrating inwards. *a*, a small laminated capsule.  $\times 100$ .

tubules getting distended with new-formed epithelium. The epithelium also in its state of activity, is altering its shape according to the mutual pressure of the cells, so that it sometimes gets elongated and tailed. (See Fig. 89.)

This activity of the normal epithelium seems to be the regular preliminary to the cancerous formation where it takes origin in glands. In the mamma, for example, there are some cancers in which the tumours have a special connection with the ducts (Duct-cancer; often co-existing with eczema of the nipple and areola). In these the epi-



Fig. 89.—From a cancer of the kidney. A tubule is represented in which the epithelium is undergoing alterations in shape.  $\times 300$ .

thelium of the ducts, from the nipple downwards, shows great activity, so that the ducts become distended with epithelium, which in accumulating loses its normal cylindrical form. The epithelium of the



acini of the gland also partakes in the active newformation. Similar processes have been observed in cancer of the uterus and elsewhere. We may therefore conclude that the first stage in the formation of a cancer is an abnormal activity in the epithelium of a particular locality.

The next stage in the development of the cancer is that its epithelial elements break bounds and extend out into the surrounding tissue. It is this atypical extension which is the most characteristic feature.

As already indicated, cancer may originate in any locality where epithelium or endothelium is normally present. It occurs in the skin and mucous membranes, in glands, in the lungs, in the brain, and (very rarely) on serous membranes. But it shows great preferences for certain localities. Thus the pre-eminent seats of cancer are the lower lip, tongue, mamma, uterus, and stomach. Many of these preferences can be accounted for by local peculiarities. Cancer of the lower lip and tongue have been ascribed to the irritation of short or rough tobacco-pipes and the jagged edges of carious teeth. The mamma and uterus suffer involution before other organs of the body, and as cancer is a disease of advanced life, the earlier decadence of these organs may determine the frequency of the occurrence of cancer in them. Again, exposure to injury and friction have been already alluded to as determining the localities of cancers. There remain, however, many unexplained facts in regard to the seat of origin of cancers.

**Age and sex** have important influences on the origin and locality of cancer. It is almost unknown during infancy and childhood, and is very rare under thirty years of age. It is frequent from thirty-five till sixty-five. The liability of the mamma and uterus to cancer causes a considerable preponderance in the female sex, which is only partly redressed by the frequency of cancer of the tongue, lip, and œsophagus in the male.

**Mode of growth and extension of cancer.**—We have seen that cancer is characterized by the atypical growth of the epithelium. The growing epithelium sends out buds or offshoots which penetrate into the underlying or surrounding tissue. In this way a primary cancer does not usually grow into a considerable tumour but rather insinuates itself amongst the tissues around. Hence we speak of **Cancerous infiltration** as characteristic of most tumours of the kind.

The question of the **Inoculation of cancer** is one which naturally suggests itself, and many experiments have been performed both on man and animals to determine the possibility of such inoculation. These experiments have all had a negative result. It is not uncommon, however, to find what may be called **Auto-inoculation**. Thus a cancer of the posterior wall of the urinary bladder will sometimes extend to a point on the anterior wall where the tumour comes in contact when the bladder is empty.



The growing cancerous processes frequently lead to much irritation in the connective tissue, so that we may have, in cancer of the lip, for instance, the extremities of the cancer cylinders buried in masses of round cells (see Fig. 90). Sometimes there is considerable cicatricial

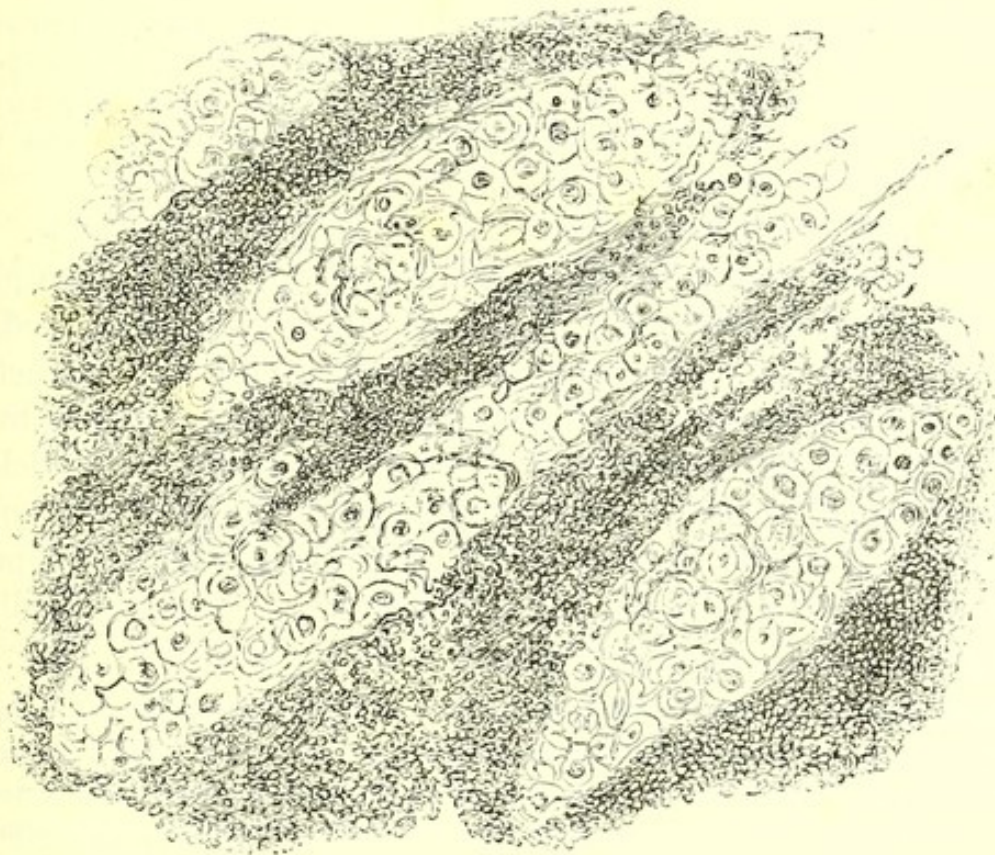


Fig. 90.—From an epithelioma of the skin. Cylinders of epithelium penetrating deeply and causing great irritation, as shown by the presence of abundant round cells.  $\times 200$ .

formation apparently induced by the insinuating cancerous processes. On the other hand the tumour may lead to little irritation, and the cancer with its characteristic stroma may grow amongst the normal tissues, merely acting on them by pressure.

The effect on neighbouring structures is to cause atrophy of their proper elements by pressure, so that the cancer takes the place of the normal tissue.

If the cancer originates at a surface or in its growth comes to present itself there, then its tissue, being less resistant than the normal structures, breaks down and we have ulcers forming. **Ulceration** is generally present in cancers of the skin and mucous membranes and in the later stages of glandular cancers.

Besides this local malignancy cancers frequently produce tumours at a distance. It is well known that the **Secondary tumours** mostly occur in the **lymphatic glands**. This may be explained partly by the anatomical connection already mentioned, and partly also by the fact



that the growing cancerous processes extending in the direction of least resistance will readily pass into the spaces in the tissues which are the radicles of the lymphatic vessels.

It seems not improbable on the other hand that cancer has a special affinity for lymphatics. The author has observed, for example, in cases of secondary cancer of the lung, that although brought to the lung by the pulmonary artery the cancerous growth has been chiefly in the lymphatic vessels, to begin with. In one such case it was quite common to find, in the neighbourhood of the pulmonary artery, the lymphatic spaces in the sheath filled out with cancerous masses. (See under Affections of the Lungs.)

The cancer forms a true newformation in the lymphatic glands. The tumour-formation mostly begins at the peripheral parts of the gland, these being the parts to which the afferent vessels are distributed, and here the characteristic epithelial structure is developed. It is sometimes very striking in the earliest periods of infection of the glands to find layers of cancerous tissue at the periphery enclosing the follicles of lymphatic tissue. The latter tissue undergoes atrophy, so that we may have only here and there groups of round cells representing the proper lymphatic gland-tissue in the midst of cancerous tissue. Finally all trace of the normal tissue disappears.

It is an important practical question whether the enlargement of lymphatic glands in connection with cancers is always a true secondary infection. There is no doubt that, just as in the primary tumour, the cancerous growth is often associated with irritation, sometimes leading to cicatricial conditions of the gland (especially in colloid cancer), but in that case there is the cancerous infection as well. In the case of ulcerating cancers, on the other hand, irritating products of decomposition may be carried to the glands and give rise to a simple non-cancerous enlargement. Hence enlargement of glands is of less significance in ulcerating cancers than in those which are not ulcerated.

The cancerous infection may for a time remain confined to the primary seat and the lymphatic glands, but it is liable to extend further and become generalized. This occurs by the material of infection reaching the blood and being carried by it to distant situations. For the most part this metastasis by the blood only occurs after the lymphatic glands have been for some time affected, and the infection takes place from the lymphatic glands. It may be that in some cases there is a direct extension from the primary tumour to the blood.

The infection may occur from the lymphatic glands after the complete removal of the primary tumour. This had happened in a case observed by the author in which, after excision of an epithelioma of the vulva, the lymphatic glands in the groin, having been affected, gave rise to multiple secondary tumours in various organs.

Having reached the blood the infection is carried throughout the body, and **Grafts are planted** in various organs. If the lymphatic



glands be in communication with the systemic veins then the infection will be carried to the lungs and on into the systemic arteries, but if they be in connection with the portal circulation, then the liver will be the organ to which they will be conveyed. This secondary (or tertiary) infection of distant organs will occur by **Embolism**, portions of cancerous tissue, perhaps only young cells, will be planted in various organs.

The metastatic growth does not occur so readily in some organs as in others. It is commonly said that the situations in which primary cancer occurs are comparatively seldom affected secondarily. Thus the mamma, uterus, and stomach are rarely the seat of secondary tumours. On the other hand the liver, lungs, kidneys, heart, skin, and bones are frequently the seat of such tumours. In some forms of cancer secondary tumours are peculiarly liable to form in the **Bone-marrow** in various parts of the skeleton. The brain is comparatively seldom affected either with primary or secondary cancers. Perhaps of all organs the **Liver** is most liable to secondary development of cancer. As it receives blood from the systemic as well as from the portal circulation, it may be infected whatever the seat of the primary tumour.

There are some apparent anomalies in the distribution of cancers to liver and lungs respectively. Thus a cancer of the lower end of the œsophagus will often give rise to secondary tumours in the liver, while cancer of the stomach may give rise to tumours in the lungs. The author believes that this depends chiefly on the relations of the lymphatic glands from which the infection of the blood occurs. In a case of cancer of the œsophagus observed by the author, he found that extension had occurred first to the lymphatic glands beneath the diaphragm and thence to the liver. On the other hand, in a case in which primary cancer of the stomach gave rise to tumours in the lungs, he found that the pre-vertebral glands were affected and that extension had occurred (as evidenced by the occurrence of thrombi) to the vena cava.

The secondary tumours are often **more favourably situated for growth** than the primary one. They may be better supplied with blood and less exposed to mechanical or other interference. Hence, they often grow to much larger size than the primary one, and may show the structure more fully developed. Thus the liver is often the seat of bulky tumours, while the primary tumour is quite insignificant.

In their structure the secondary tumours **imitate the primary one** even in the finer details. This applies not only to the shape and size of the epithelial cells, but to the abundance and arrangement of the stroma, and even of the vessels in the stroma. If the stroma be abundant and fibrous in the primary tumour, it will show, at least a tendency in the same direction in the secondary ones, although time may have failed to allow of the full manifestation of this.



This imitation of the primary growth produces very remarkable results, when one sees, for instance, gland-like tissue, lined with cylindrical cells, growing abundantly in the liver, or lung, or brain. A striking illustration of this mimicry was found by the author in a case where a cancer of the stomach showed a striking tendency to hæmorrhage; the patient actually died from the effects of a large hæmorrhage. There were numerous secondary tumours in the liver, which looked almost like masses of blood. The delicate character of the vessels had been repeated in the secondary tumours, and bleeding was characteristic of them as well as of the primary growth. Facts like these render it very difficult to believe that a micro-organism has anything to do with the causation of cancer.

**Retrograde changes in cancers.**—The cancerous tissue is much more prone to degenerations and secondary changes than is normal tissue.

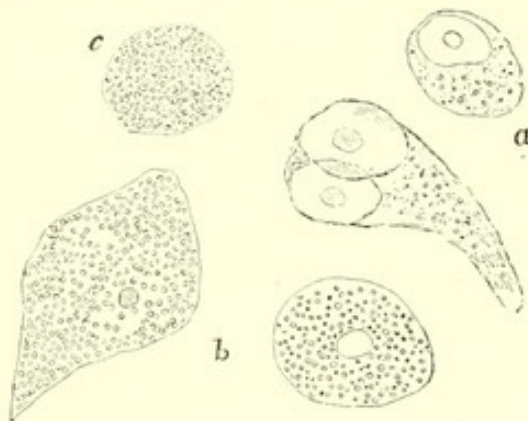


Fig. 91.—Fatty degeneration of cells in a cancer of the mamma; *a*, slightly affected; *b*, more so; *c*, completely fatty—the compound granular corpuscle.  $\times 350$ .

**Fatty degeneration** is very frequent. (See Fig. 91.) This may, in quickly growing tumours, affect considerable portions, so as to give rise to an appearance like caseation. In more chronic cases, the fatty degeneration affects more the individual cells. The degenerated cells are readily absorbed and this often leads to a relative preponderance of the stroma. Thus cancers frequently shrink and become cicatricial in their older or central parts.

This may lead to dimpling of the surface of a tumour, as we often see in cancers of the liver, producing the so-called **Umbilication**. **Mucous** and **Colloid degeneration** are not infrequent in cancers. They occur to a minor degree in many cancers of the intestine, but in a higher degree they are so characteristic as to give a special name to a form of cancer. (See Colloid Cancer.)

**Ulceration** is a regular result in most superficial cancers. As a general rule the cancerous ulcer is bounded by a prominent border, composed of tissue infiltrated with the growing tumour.

**The local nature of cancer.**—From what has been stated above, it will appear that cancer begins as a local growth of epithelium, accompanied by the formation of a connective-tissue stroma of varying complexity. It is in many cases a well-formed but atypical tissue. In the secondary extension the other tumours bear a definite material relation to the primary one. They arise by the implantation of grafts, first, as a rule, in the lymphatic glands, and secondly, it may be, in parts further removed. As the primary cancer sends offshoots amongst the tissues, and extends outwards to the lymphatic glands, it must usually be difficult to determine its limits, but if these limits can be determined,



and the whole growth removed, then we must infer that the disease will be eradicated. It is exceedingly rare to find two primary cancers in the same person. In the great majority of cases, all the existing tumours are direct descendants of a single primary growth.

**The individual forms of cancer.**—Various modes of classification have been adopted. That which we use here is not entirely satisfactory, but it is useful for practical purposes.

1. **Flat-celled epithelioma; Epithelioma proper.**—In English works the term Epithelioma is chiefly used to designate cancers of the

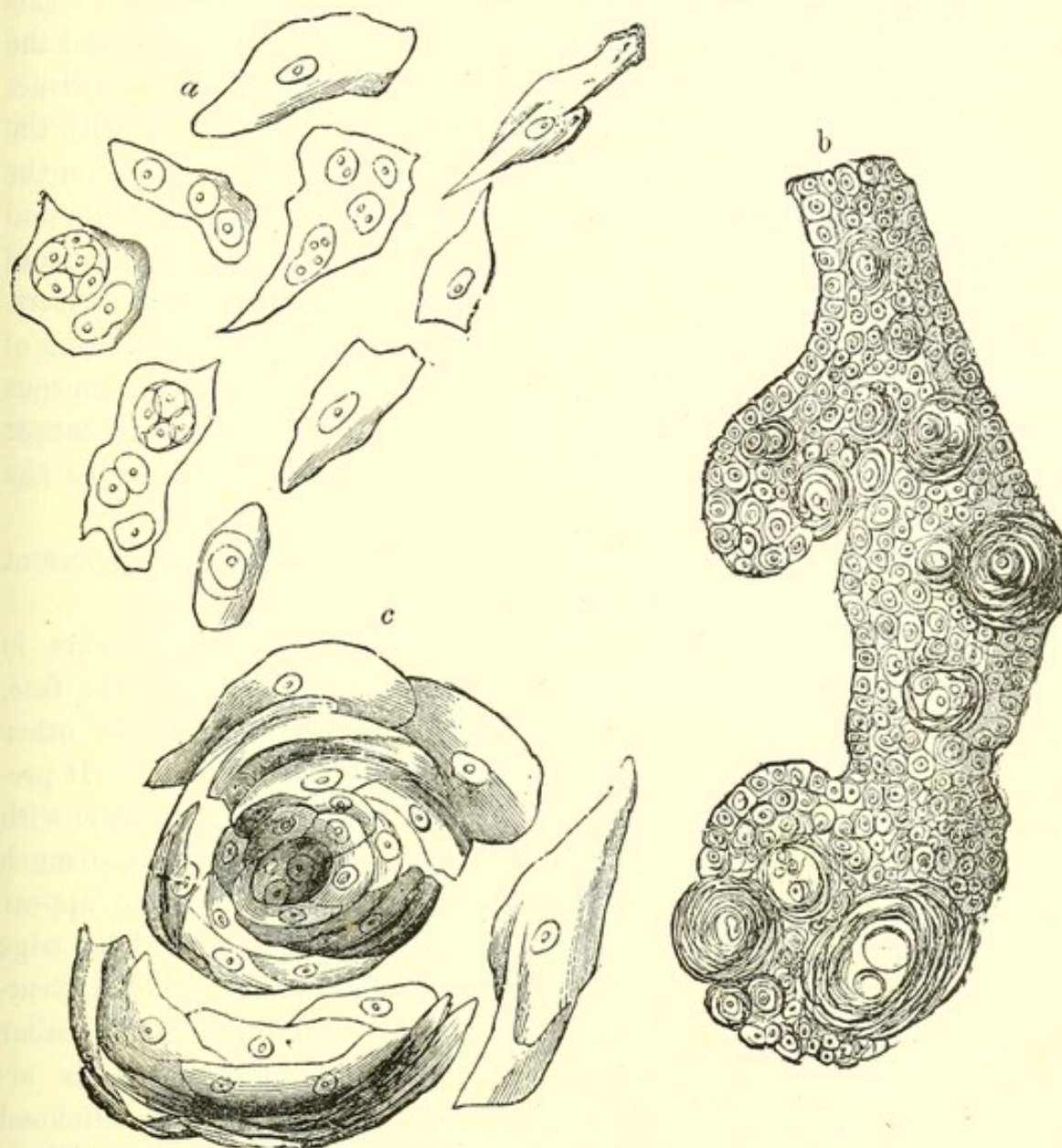


Fig. 92.—Flat-celled epithelioma: *a*, isolated cells; *b*, a cylinder with many laminated capsules; *c*, a laminated capsule opened out so as to show the cells composing it. (BILLROTH.)

cutaneous surface, tongue, and œsophagus, places where the surface is covered with flat epithelium. Similar tumours occur in the larynx, uterus, and vagina, and urinary bladder. It is possible to distinguish a penetrating or infiltrating form, and a more superficial flat form. This



latter form is almost equivalent to Rodent ulcer of English authors, and *Flache Krebs* of Thiersch and the German authors.

The common **Epithelioma of the lip** is the most familiar example of the infiltrating form. In it cylindrical processes of epidermis, taking origin in the surface epithelium, grow downwards into the true skin, infiltrating it, and destroying its connective tissue. These processes, as they grow, exercise a concentric pressure on their own cells, and so produce closely-packed rounded globular bodies, composed of epidermic cells, wrapped round each other. These bodies, variously called **Epidermic globes**, **Laminated capsules**, etc. (see Fig. 92), are very characteristic of this form of epithelioma. The cells in the globes are usually horny, and the consequent bright translucent appearance, as well as the shape, attract the eye in microscopic sections. In some parts there is, along with the production of these penetrating processes, a formation of papillæ on the surface. This is often manifest in the epitheliomas of the scrotum and certain other parts of the skin. It is also very pronounced in some of the vagino-uterine cancers, forming the so-called **Cauliflower cancers**. In the urinary bladder there is frequently such a marked production of elongated papillæ that the surface is quite shaggy, while the mucous membrane beneath is infiltrated. To this form the name **Villous cancer** is often given. The papillæ, like those of the simple papilloma of the bladder, are liable to hæmorrhage.

All these epitheliomas are prone to ulceration, and frequently present themselves as ulcers with infiltrated walls.

**Rodent ulcer** is a special form of epithelioma which occurs in



Fig. 93.—Rodent ulcer of face. The epithelial masses *b b* (the cells sometimes cylindrical) are beneath the surface epidermis (*a*). Towards the left a hair follicle (*c*).

the upper part of the face, and more rarely in other parts of the body. It presents itself as an ulcer with overhanging edges, so much overhanging as to appear "rolled over." The edge shows the epithelial structures in the form of peculiar small epithelial cells arranged in well-defined groups (see Fig. 93). These groups of cells are beneath the epidermis, and seem to have no connection even with the malpighian layer,

as Thin has pointed out. They often present a cylindrical arrangement,



and the peripheral cells are sometimes columnar. The tumour is a superficial one, there are no penetrating cylinders, no laminated capsules, and no tendency to invade the lymphatic glands.

The name **Cholesteatoma** or **Pearl tumour** has been given to a form which is variously regarded as an epithelioma or a cystic tumour. It contains bright, glancing, pearl-like structures, which are due to the presence of cholestearine and fat. Besides these, there are flat cells, epithelial in character, which are arranged into rounded bodies, inside which the cholestearine is contained. These tumours are sometimes surrounded by a capsule so as to resemble atheromatous cysts. They are most typically seen in the soft membranes and substance of the brain, but also occur in the subcutaneous tissue, testicle, ovary, parotid, and ear. (See Virchow, in *Virch. Arch.*, viii.; Eberth, *do.*, xlix.; Eppinger, *Prag. Vierteljahrschr.*, 1875; Chiari, *Cholesteatom des Rückenmarks*, *Prag. med. Wochenschr.*, 1883; Bristowe, *Path. trans.*, v. 24, 1854; Price, *do.*, xxxviii. 24, 1887.)

**2. Cylinder-celled epithelioma.**—This is a tumour of parts where cylindrical epithelium normally covers the surface, hence chiefly in the stomach and intestine, and more rarely in the uterus.

The tumour usually takes origin in the mucous membrane or its glands, but it may occur also in the brain or in the lungs. The tissue has a glandular appearance, forming a congeries of tubes and cavities, lined with cylindrical epithelium (see Fig. 94). Sometimes the glandular appearance is strictly preserved, but frequently the spaces enlarge,

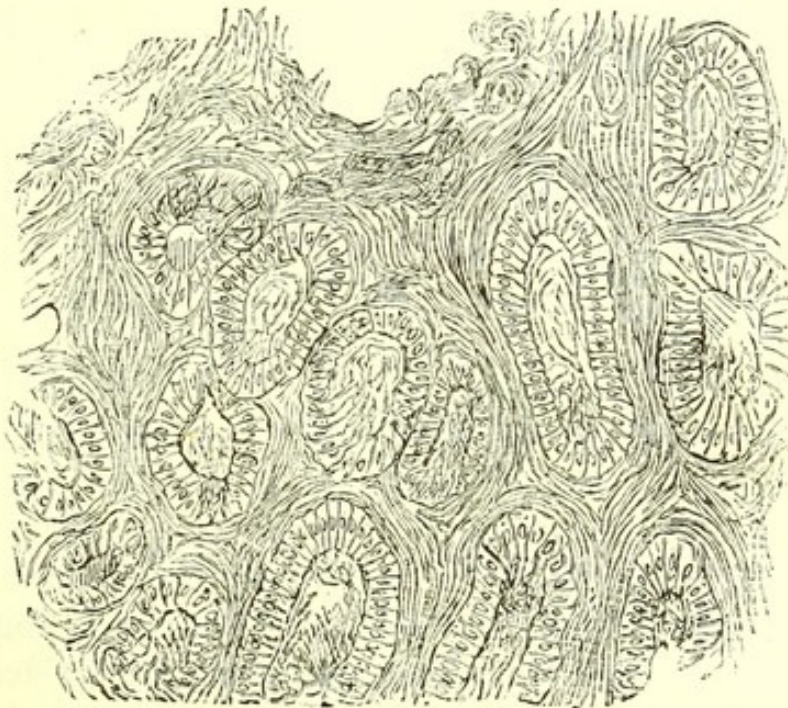


Fig. 94.—From a cylinder-celled epithelioma of the rectum.

and as the epithelial cells accumulate they lose their cylindrical form, although the outer layer of cells next the stroma may still preserve their shape (see figure). The glandular elements are con-



tained in a well-formed stroma, which is new-formed as well as the epithelial elements.

In growing, the tumour infiltrates neighbouring parts, insinuating itself among the muscular trabeculæ very often, and extending further. It may form a considerable tumour, projecting from the surface, and is frequently ulcerated, as it occurs mostly in situations where it is exposed to friction.

From the strikingly glandular character of these tumours, they are included by some writers among the adenomas. They are called by these writers **Malignant adenoma** or **Adenocarcinoma**. As they are so distinctly atypical in their mode of growth, there seems no reason to separate them from the cancers.

3. **Soft cancers** or **Medullary cancers** are characterized by the existence of a very delicate stroma in which are abundant cells, usually of small size, and loosely packed in the alveoli with a good deal of fluid. They occur chiefly on mucous membranes, in the ovaries, testicles, kidneys, less commonly in the mamma. Being soft, they tend to bleed, and if originating on a surface or coming to it in their growth, they may undergo ulceration. Sometimes the softened, ulcerating, bleeding tissue projects in a very striking manner from a surface, forming the **Fungus hæmatodes**, this condition also occurring, however, in soft sarcomas (p. 256). The cut surface of such tumours is grey in colour, and a somewhat fluid juice can be scraped from it. In this juice will be found cells and free nuclei, the latter large and mainly oval in shape. Many of the cells contain fat granules, and there may be some in an advanced state of fatty degeneration.

Cancers of this kind sometimes grow to a considerable size, and they are often of very rapid growth. They are, as a rule, very malignant; the young cells, being loosely attached, readily pass away and lead to metastasis.

4. **Hard cancer** or **Scirrhus** occurs most commonly in the mamma, but also in the stomach, the testicles, ovary, pleura, and peritoneum. It is characterized by the preponderance of connective tissue as compared with the epithelial structures.

The cancers of this kind have usually a very infiltrating character, the epithelial processes penetrating among the surrounding structures, and it looks as if these processes, by their irritation, produced an excessive amount of connective tissue. This view is borne out by the fact that the connective tissue is not usually in the form of a well-developed stroma as if planned to support the epithelial structures, but is irregular and even impinges on and destroys the cells. The cells readily undergo fatty degeneration, and they sometimes to a large extent entirely dis-



integrate, the stroma assuming the upper hand. It therefore happens that different parts of such tumours have often very different structures. The more recent parts will show well-marked epithelial masses with stroma (Fig. 95), while in the older parts the cells have almost disappeared, and there is nothing but dense connective tissue.

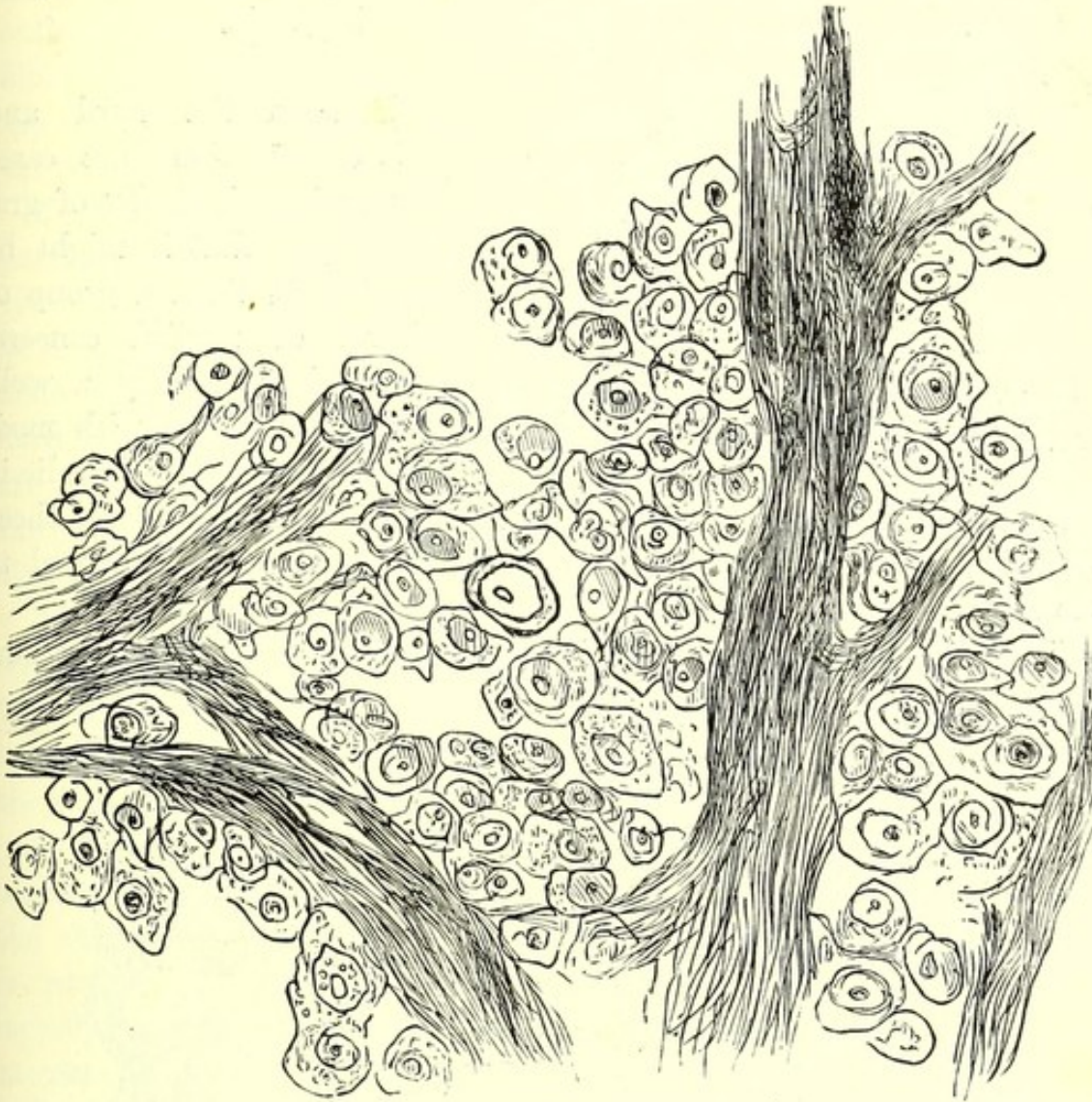


Fig. 95.—From a scirrhus of mamma, recent part showing stroma and epithelial cells.  $\times 350$ .

As scirrhus occurs more as an infiltration than a distinct tumour, the hardening and contraction of the connective tissue causes the organ in which it grows to be sometimes contracted rather than enlarged. It is frequently so in scirrhus of the mamma and of the stomach, its two most frequent seats. On cutting into the organ the tissue is felt to be dense and elastic. The cut surface is greyish and transparent, with opaque yellow markings indicating the existence of fatty degeneration in the cells. The juice to be obtained from the cut surface is scarce, and under the microscope it is seen to contain cells, often of large



size, and free nuclei (see Fig. 96). The cells vary greatly in size and shape, and they often contain two nuclei, or even a fully formed cell inside (mother and daughter cells).

These cancers are less malignant than the soft cancers, but they produce somewhat readily secondary tumours in lymphatic glands.

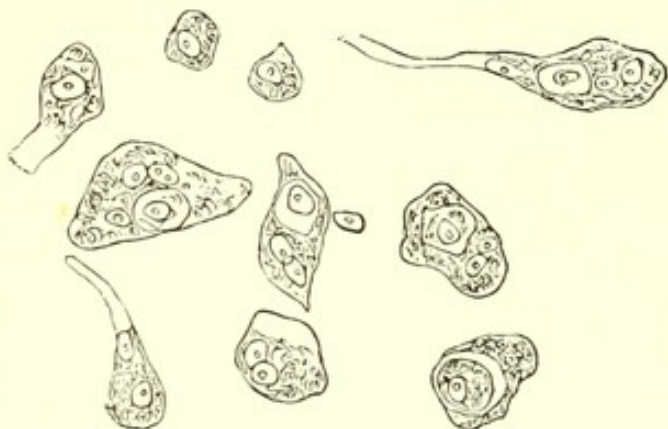


Fig. 96.—Cells from a cancer of the mamma. Most of them contain several nuclei, and some daughter cells.  $\times 200$ .

Between the hard and soft cancers there are cases presenting all shades of gradation; so that it might be possible to form a group of simple or normal cancers.

These have mostly a well-developed stroma with moderately sized cells which lie in the stroma in considerable spaces. The mamma presents these varieties, and they will be again referred to in the section devoted to the diseases of that organ.

5. **Colloid or Alveolar cancer** is a tumour characterized by the occur-



Fig. 97.—Colloid cancer. A finely reticulated stroma is seen with meshes full of colloid material, the cells having disappeared.  $\times 90$ .

rence of colloid degeneration of epithelial cells. It is met with chiefly in the stomach and intestines, and in the mamma, more rarely elsewhere.



There is here a definite newformation both of stroma and epithelial masses, and the stroma is often produced in the most beautiful and characteristic forms (see Fig. 97). As if it were in the plan of the growth, the cells regularly undergo colloid degeneration, and finally the masses of cells become converted into masses of colloid material which fill the spaces. In the bulk of the tumour therefore there may be nothing visible beyond the regularly formed stroma whose spaces are filled with clear transparent material. Occasionally there may be in the centres of the alveoli some remains of the cells visible, while the peripheral cells are already completely converted.

To the naked eye such tumours have a markedly gelatinous appearance, and as the fibrous stroma may in its coarser meshes be visible without the microscope, it may, even to the naked eye, look as if there were nothing but alveoli filled with gelatinous material, hence the name alveolar cancer. The tissue is frequently dense and hard to the feel. This arises apparently from the fact that the alveoli are tensely packed with the colloid material, and, the fibres being on the stretch, a dense resistance is offered, just as a tightly blown up bladder is hard.

These tumours mostly occur as infiltrations, frequently penetrating among the constituents of the tissues, and although they very often extend widely by continuity (as in the stomach), they show little tendency to metastasis; even when they attack the lymphatic glands secondarily they do not usually produce large tumours.

6. **Melanotic cancer** is a rare form of tumour compared with the melanotic sarcoma. It occurs primarily in similar situations, namely the skin and eyeball. It is really a soft cancer in which pigment is present in the cells and also sometimes in the stroma. It is usually a very malignant tumour, producing secondary growths by metastasis.

7. **Mucous cancer** includes tumours in which the stroma of the cancer assumes the characters of mucous tissue. The cells of the cancer may undergo a similar degeneration. The tumour as a whole is very gelatinous in appearance, and resembles the colloid cancer. These tumours are of rare occurrence, being met with in similar situations to colloid cancer, but attaining the highest development in the ovary.

8. **Endothelioma** is a name sometimes applied to cancers arising from the endothelium of serous membranes and elsewhere. (See above, p. 260.) Cancers of the pleura and peritoneum belong to this class. They will be described in connection with tumours of their own locality.



**Literature.**—HANNOVER, *Das Epithelioma*, 1852; LEBERT, *Traité des malad. cancéreuses*, 1851; FÖRSTER, *Handb. d. path. anat.*, vol. i. (a very full account); THIERSCH, *Der Epithelkrebs, namentlich der Haut*, 1865; WALDEYER, *Virch. Arch.*, xli. 470 and lv. 67, also Volkmann's *Sammlung Klin. Vort.*, No. 33; LÜCKE, in Pitha-Bilbroth's *Handb.*, 1869; PAGET, *Lect. on Surg. Path.*, 3rd edition, 1870; KÖSTER, *Die Entwicklung der Carcinome*, 1869; FRIEDLÄNDER, *Epithelwucherung und Krebs*, 1877; CORNIL et RANVIER, i. 208, 1881; Discussion on Cancer, *Glasg. Path. Soc.*, papers by MACEWEN, JONATHAN HUTCHINSON, Author, etc., *Glasg. Med. Jour.*, 1886; THIN, *Cancerous affections of Skin*, 1886 (full literature and good account of Rodent ulcer); PERLS, *Lehrb. d. allg. Path.*, i.; WOLFF, *Entstehung der Carcinomen aus traum. Einwirk.*, 1874.



## SECTION X.

## VEGETABLE PARASITES.

- A. BACTERIOLOGY.—I. GENERAL CONSIDERATIONS. *Definition and classification, structure and mode of growth of microbes; cell, cell-membrane, spores; methods of detection. Conditions of life, as to temperature, oxygen, etc. Action of disinfectants. Products of bacteria. Distribution, Cultivation, Polymorphism.* II. BACTERIA IN RELATION TO DISEASE—1. *Saprophytes produce ptomaines; origin of wound-fever, etc.; 2. Pathogenic forms produce local infection or may enter blood; lead to necrosis and inflammation; products may enter blood, or themselves.* III. THE INDIVIDUAL FORMS—A. *Some saprophytic forms; B. pathogenic forms—1. Micrococci, 2. Bacilli and spirilla, 3. Bacteria of specific newformations, 4. Some forms in animals.* IV. DISPOSAL OF BACTERIA; IMMUNITY; *natural and acquired; vaccination; causes of immunity; PHAGOCYTES.*
- B. PARASITIC FUNGI.—I. THE YEASTS. II. FILAMENTOUS FUNGI—1. *Saprophytes and occasional parasites; 2. Pathogenic fungi.*

## A.—BACTERIOLOGY. MICROBIOLOGY.

THE relation of minute vegetable organisms to disease has of late years assumed a high degree of importance, so much so that a special science has been created with its chairs and laboratories, and a literature which has already reached very large dimensions.

The minute organisms have received the names of **Bacteria** or **Microbes**, these terms being used in a general sense to include the whole group. The department of pathology which deals with them is designated **Bacteriology** or **Microbiology**. The reader is referred to special works on the subject for the details of methods of investigation, we are here chiefly concerned with the relation of the microbes to disease.

## I.—GENERAL CONSIDERATIONS.

**Definition and classification.**—The bacteria are the lowest forms of organic life. They consist of minute round, oval, or cylindrical cells, so small that they require high powers of the microscope for their detection. Like the fungi, they are, except in one or two doubtful cases, devoid of chlorophyll, and like them are incapable of evolving



their structure from inorganic matter. But they are even lower than the fungi, as they multiply almost entirely by division. It is from this that the name **Schizomycetes** is derived, a name equivalent in its meaning to **Fission-fungi**. There is an approach to reproductive organs only in a few forms in which spores are produced under certain conditions, but even in them the multiplication is mainly by fission.

It is not possible at present to make a proper scientific classification of the microbes. The working out of that may be left to the botanists. For our purposes the classification of Cohn is sufficient. He distinguished them according to their external form, into the globular, the short rods, the longer rods, and the spiral forms. The rod-shaped ones may be taken together, so that we have three forms, the globular or oval (cocci, micrococci), the rod-shaped (bacilli), and the spiral (spirilla).

**Structure and mode of growth.**—The bacterium, of whatever form, is a cell, consisting of cell-contents and membrane, but **without a nucleus**. The **Cell-contents** consist of an albuminous substance, which in its reaction to staining agents closely resembles the nuclei of ordinary animal and vegetable cells. This is more particularly the case with the aniline dyes, which the bacteria, as well as the nuclei of cells, usually absorb greedily, but the bacteria, in many cases, retain the colour more firmly than the nuclei, some of them even in the presence of acids. The contents sometimes present granules of starch, and in one form, of sulphur. The **Cell-membrane** is composed of a substance allied to cellulose. It is difficult to demonstrate, but by the use of iodine the contents may be made to shrink so as to display the membrane, which may be now stained of a different colour (Crookshank). The cell-membrane sometimes in its outer parts swells up or otherwise produces a **Gelatinous envelope**, which forms a cement between the individual bacteria, uniting them in various numbers. Some bacteria are possessed of a **Flagellum** or lash, by the active movement of which they move about. As the bacteria themselves are very small, and the flagellum is much narrower, the latter is very difficult of observation, but it may be rendered visible by staining with logwood or aniline dyes.

The bacteria multiply, as already mentioned, by fission, and they do so, under favourable circumstances, with enormous rapidity, so that, as has been calculated, a single individual will multiply a million-fold in twenty-four hours. In multiplying, the individual members may remain for a time united, and so give rise to specific appearances. Thus some of the cocci divide in succession in the same direction, and often appear first in twos, **diplococcus**, or elongated into **chains**. The bacilli also frequently form chains. The gelatinous envelope again may unite them into large groups or colonies, to which the name **Zooglœa** is given. The



zooglœa thus formed sometimes form membranous aggregations, such as we so often see as a scum on the surface of decomposing fluids. Seen thus in the mass in zooglœa, the bacteria give under the microscope a characteristic brownish clouded appearance, as shown in Fig. 98.

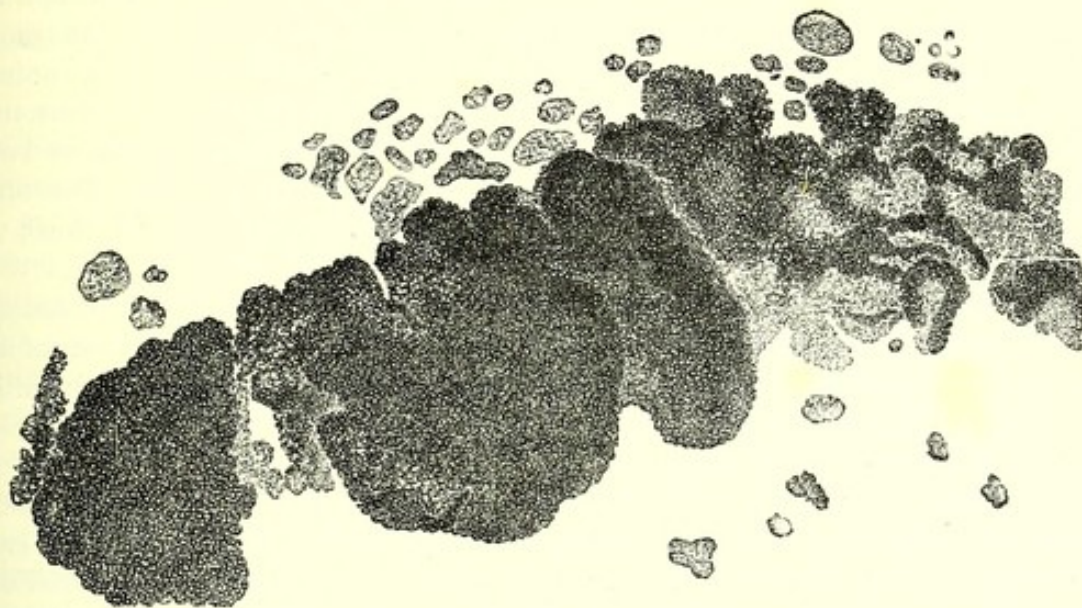


Fig. 98.—Swarms of bacteria growing in a nutritive material.  $\times 90$ .

Some bacteria produce **Spores**, and it is this faculty to which botanists pay special attention, with a view to classification. Two forms of spore-formation are described. In the one the spores form inside the bacteria, in which they are differentiated as round or oval bodies. These are called **Endospores**. In the other form, individual members of a group of cocci become larger and more prominent, having greater refractive power on light, and so assume the character of spores, while the remaining individuals undergo no such change. These are called **Arthrospores**. In whatever way produced the spore has a thick membrane and is much more resistant than the ordinary cell. The spores are more difficult to stain than the rest of the cell, but when stained they retain the colour more firmly. This fact may be made use of to effect a double-staining, the spores one colour and the rest of the bacterium another. They are stained brown with osmic acid, from which it has been inferred that they contain some form of fat. Bacteria sometimes present an appearance resembling endospores, but not truly of that character. The beads, so characteristic of the tubercle-bacillus for instance, are not universally regarded as spores, although by some viewed in that way. The great majority of bacteria show no spore-formation of any sort, and even those which bear spores do so only under special circumstances.

**Methods of detection.**—It will be obvious that these minute organisms are often very difficult of detection, especially when they are mixed up with other structures



or lie in the midst of the tissues. When they are in colonies their opaque clouded appearance and their frequent brown colour make them often very prominent objects. But when more isolated, and even when in sparsely distributed colonies, it may be difficult to distinguish them from the mere granular debris of the tissues. In order to render them more easy of detection, various methods have been devised. The simplest is that originally suggested by Recklinghausen, which is based on the fact that these organisms present a much greater resistance to alkalies and acids than the animal tissues do. The addition of solution of caustic potash to an animal tissue renders it transparent and obscures the structure. If bacteria be present, they will be rendered more prominent than before. Dilute acetic acid, which as we know clears up the connective tissue especially, may be used in a similar way. This often succeeds in bringing into prominence the zooglœa of bacteria, but is not of much use in detecting isolated ones.

Much advance has been made by the introduction of improved methods of staining bacteria, devised by Weigert and Koch. Bacteria absorb readily several kinds of dissolved pigments, but they show a peculiar tendency to become coloured with aniline dyes. Many such dyes have been used, but the most suitable seem to be methylviolet, gentianviolet, fuchsine, and methylblue.

Most bacteria stain readily with watery solutions of these dyes, but various methods have been devised either to render the stained bacteria more prominent, or to stain certain forms which are refractory to the ordinary dyes. One of the most useful of these is **Gram's method**, by means of which the bacteria are stained blue, while any other structures can receive a contrast stain of red or brown. There are also the special methods for the tubercular, syphilitic, typhoid, and other forms, in which also a double or contrast staining is aimed at.

In staining bacteria contained in fluids the co-called **Cover-glass method** is most convenient. A thin layer is spread on a cover-glass, and dried so as to form a thin film. If the fluid contains albumen, the film should be fixed by passing it three or four times slowly through the flame of a lamp, taking care not to scorch it. The film may now be stained by any of the methods referred to above, and then, after drying, mounted in Canada balsam. Sections of tissues are sometimes dyed and stained in a similar fashion, but as a rule they require to be kept moist. After removal from the staining fluid they are placed in alcohol, then in oil of cloves, or other clearing agent, and mounted in Canada balsam.

In all microscopic examinations for bacteria it is necessary to use an illuminating apparatus similar to that known as Abbe's condenser. This piece of apparatus has the effect that all coloured objects are rendered visible, but all uncoloured details are obscured. The bacteria, when stained with aniline dyes, are thus rendered very prominent.

The special methods will be found described in detail in works on Practical Pathology and on Bacteriology.

**Conditions of life.**—The bacteria require for their formation and growth the presence of organic matter. They differ from most plants in respect that they are unable, on account of the absence of chlorophyll, to eliminate their carbon from pure carbonic acid, and hence they derive it from higher compounds. The nitrogen may be similarly derived, but it may also be obtained from inorganic bodies, nitrates and ammonia salts. The bacteria grow best in a medium of an **alkaline**, or



at least **neutral**, reaction, in this respect contrasting with the fungi which grow vigorously in the presence of acids.

**Temperature** exercises considerable influence on most bacteria. Many of them grow best at the temperature of the blood, and some of them are only active at that temperature, but most of them grow readily at the ordinary temperature of the air.

As a general rule when the temperature falls to  $5^{\circ}$  C. ( $41^{\circ}$  F.) their growth and multiplication cease, but they are not killed even by extreme degrees of cold. Thus Pictet and Young exposed several forms of bacteria to a temperature of  $-70^{\circ}$  C. ( $-94^{\circ}$  F.) for 109 hours and then to that of  $-130^{\circ}$  C. ( $-202^{\circ}$  F.) for 20 hours, and found that many of them retained their vitality. This was the case with the anthrax bacillus which produced its effects when injected into animals after this exposure. Coleman and M'Kendrick exposed putrescible fluids to a temperature of  $-83^{\circ}$  C. for 100 hours, and found that the bacteria were not killed.

Bacteria have less power of resisting high temperatures. Activity is suspended usually at a temperature of  $45^{\circ}$  C. ( $113^{\circ}$  F.), and a prolonged exposure at a temperature of  $50^{\circ}$  to  $60^{\circ}$  C. suffices to kill most bacteria. Hence boiling generally suffices to destroy the vitality of bacteria. The spores are more resistant than the bacteria themselves, and some of them resist boiling unless it be prolonged. According to Pasteur, there are some spores which can withstand a temperature of  $130^{\circ}$  C.

**Water** is necessary to the growth of bacteria. The ordinary cells die when dried, but the spores survive and some of them may be preserved for years in the dry state.

**Oxygen** or atmospheric air is necessary for the growth of many bacteria, but for some the exclusion of air is a necessary condition. To the former Pasteur has applied the term **Aërobic**, and to the latter **Anaërobic**. There are very few which do not require at least a certain supply of oxygen, which they may derive from its presence in ordinary water.

Certain substances **inhibit** or **destroy** bacteria. It is not to be inferred that because an agent destroys one form that it will destroy all bacteria, for, as a matter of fact, the same agent will have very different effects on different forms. It is to be noted also that an agent may inhibit the growth of bacteria without destroying them. Carbolic acid, for example, stops the growth of many forms, but it may do so when too dilute to kill them. For surgical purposes it is often sufficient to inhibit the growth, although it is not surprising that even under carbolic acid dressings there may be a growth of bacteria when the agent has become greatly diluted.

Koch tested a series of **Disinfectants** as to their power of destroying the vitality of the anthrax bacillus and its spores, as well as of other forms of bacteria. Carbolic acid in a one per cent. solution destroyed the vitality of bacilli in two minutes, whereas the spores required exposure to a five per cent. solution for more than twenty-four



hours. A much more dilute solution, when present in a nutrient medium, inhibits the growth even of spores. The most active known agents are the salts of mercury, which in very dilute solutions kill even the spores of anthrax. A watery solution of chloride, nitrate, or sulphate of mercury 1:1000 destroyed the spores in ten minutes. Solutions of chloride of mercury in even greater degrees of dilution destroyed these spores. (See Koch On Disinfection, in Selected Essays, New Syd. Soc., 1886.)

**Products of Bacteria.**—In their growth bacteria are engaged for the most part in splitting up organic compounds, and in building up others. In this process they frequently eliminate chemical principles of greater or lesser complexity. In the souring of milk, for instance, the bacterium *lactis* changes milk sugar into lactic acid. Then in ordinary putrefactive processes the albuminous principles are decomposed, while various principles are produced, some of them alkaloids, and some gases, which present peculiar disagreeable odours. There are some which produce pigments; the micrococcus *prodigiosus*, for example, is characterized by the brilliant red appearance which it presents when growing in masses. Other bacteria produce blue, black, and yellow pigments. These pigments are, as it were, secretions of the bacteria, and in many cases, at least, the pigment is not in the substance of the bacteria but lies between them. It is now generally acknowledged that decompositions of organic matter and fermentations are the result of the action of bacteria, and it is recognized that each form of bacterium, when under similar conditions, produces the same chemical results. Certain of the alkaloids eliminated are called **Ptomaines** (see p. 285).

**Distribution.**—Bacteria are present almost everywhere in nature, but they are specially abundant where water and decomposable organic matter concur. They are present **in water**,—even the purest waters contain them in considerable abundance. They exist **in the air** provided it contains watery vapour. If the air is dry then spores will be present rather than the bacteria themselves. **In the earth** they are present, but, as they are here liable to drying, it is often chiefly spores which are present. There are not many below the level of three feet from the surface. They are present abundantly **on the human body**; on the skin wherever there is dirt, and on the mucous membranes.

**Cultivation.**—The greatest advances in our knowledge of bacteria have been made by their artificial cultivation. Following the example of Koch the bacteria are now cultivated not only in fluids, but on solid media, where they can be kept much more free from contamination and may be more completely observed in their growth. The cultivation takes place on the surface of nutrient substances, of which the chief are potatoes and meat-juice. Fluid media are rendered solid by the addi-



tion of gelatine or agar-agar. As the latter substance does not melt at the temperature of the body, it is exceedingly useful in cultures which require a higher temperature than that of the ordinary air.

**Pure cultivations** are made by first sterilizing the medium and inoculating afterwards, taking great precautions against contamination. It is possible to propagate many individual forms through successive generations and on different media, so as to observe their habits. The growing and multiplying bacteria are visible on the surface of the medium, and the individual forms present, as a rule, variations in mode of growth, colour, and otherwise, sufficient to enable a practised eye to discriminate the form which is under observation. In their growth in media which have been stiffened with gelatine, they often produce liquefaction. When one sees the visible characteristic layers of bacteria on the surface of potato or other medium one has a vivid conception of the reality of these forms which individually are mere microscopic objects.

The methods of cultivation will be found in text-books of Bacteriology, such as Klein's and Crookshank's. It may be noted here that by an ingenious application of Koch's gelatine method a quantitative estimate can be formed of the number of bacteria in a given amount of air, earth, or water. This is done chiefly by means of so-called **Plate cultivations**. The gelatine solution is heated gently so as to melt it, mixed with a determined quantity of the substance to be estimated, and then poured, in a thin layer, on a glass plate, where it solidifies. The bacteria present develop and form centres of growth which can be counted and so give an approximate estimate of the numbers in a given quantity of material. It has been estimated, for example, that in the water of the River Clyde at Glasgow there are 1,500 bacteria in every drop, while even in Loch Katrine water, which is exceedingly pure, there is about one in every drop (Maylard).

**Polymorphism.**—An important question has been raised as to whether the bacteria always retain their forms, and accordingly whether the genera can be formed on the basis of their outward configuration. It has been asserted that under varying circumstances not only the form but the physiological characters may vary. It is true that according to the stage of development, and the conditions of nutrition, the individual forms do undergo certain variations in size and shape. This is, to a certain extent, true of all living things. But it is also true that when the more perfect modern methods of cultivation are used, the identity of the different forms can be absolutely secured, and that in such case they have a definite configuration which is the constant expression of their complete state of development. As in the case of higher plants we can here recognize indications of genera and species, although the objects are too minute and the science too recent to permit of a permanent classification.



The alleged cases of alteration of form and characters arise largely from imperfections in the observations. It may be acknowledged, however, that in the case of some of the bacteria a very strong case has been made out for polymorphism. In the construction of genera and species here as in higher plants, attention will be paid to the whole life-history of the bacteria and not merely to their outward form. We are still far from a complete system on such a basis.

## II.—BACTERIA IN RELATION TO DISEASE.

It will be readily understood that the great majority of bacteria present no relation whatever to disease; they have their various functions in the economy of nature, and may never come into relation with the living body of an animal. In this connection bacteria are divisible into three groups, those which feed only on dead animal or vegetable matter, and which are called **Saprophytes** (*σαπρός* = putrid), those which feed only on the living body of higher organisms or **Obligate parasites**, and those which may live on either the one or the other and which may be called, according as they are viewed in one aspect or the other, **Facultative parasites** or **Facultative saprophytes**.

Another, and, for our purposes, perhaps a more suitable division of bacteria is according as they produce disease or not. Bacteria may exist in the living body, and live on its secretions or excretions and yet do no harm. Those which propagate in the living tissues and lead to definite morbid processes, such as inflammations and fevers, are called **Pathogenic**, while those which live simply on the excreta of the body are **Non-pathogenic** or **Saprophytic**.

As a general rule bacteria will produce disease only when they penetrate into the tissues of the living body, but it is not to be supposed that it is by their mere presence that they produce their effects. They are in themselves small particles of matter, often very insignificant in relation to the mass of the tissue. Nor is it by using up the material required by the tissues that they do harm. In a few cases, such as that of the anthrax bacillus, they may be present in enormous numbers in the blood, and it has been supposed that they do harm by using up, perhaps, the oxygen of the blood, or blocking the capillaries of the lungs so that respiration is impeded. But if at all true this view is only partial. Bacteria produce their effects essentially by the products which they evolve, and the effects vary according to the kind of bacteria and the nature of the products.

1. **Saprophytic bacteria** may lead to symptoms of disease if their products are absorbed by the living body. In the process of putrefaction in dead animal matter certain alkaloids are formed, which have been placed under the general designation of **Ptomaines**. The result of



the absorption or injection of these products into the living animal is a group of symptoms, which may be included in the term **Putrid or Septic poisoning**, the principal being hectic fever and diarrhoea. Such symptoms are producible in animals by the injection of such products even after the destruction of all living bacteria by boiling and filtering.

No doubt the so-called **Ptomaines** include many different alkaloids, some of which have been at least partially isolated. Panum isolated a virulent poison which was soluble in water but not in alcohol which he called the **Putrid poison**. Bergmann and Schmiedeberg isolated a crystalline substance which they called **Sepsin**, and several other crystalline alkaloids have been obtained, especially by Brieger. Amongst the latter may be mentioned **Neurine**, an alkaloid obtained from decomposing muscle. It possesses properties similar to those of muscarine, a poison obtained from some fungi, and peculiarly fatal to flies. Poisonous products evolved by the decomposition of animal foods sometimes lead to serious and even fatal results. There are not a few cases of poisoning by sausages, in some of which the symptoms have resembled those of poisoning by atropine (Brunton). There have been also cases where fish of various kinds, but especially crabs, have had this effect, and there are also cases in which tinned meats have apparently had poisonous properties. It appears as if the poisons might be evolved without the ordinary signs of putrefactive decomposition. In that case the bacteria concerned would not be associated with the ordinary septic kinds.

Somewhat more problematical are the so-called **Leucomaines**. It is asserted (Gautier) that in the physiological processes, such as muscular action, alkaloids are formed, which, like those arising from bacteria, are poisonous. To these physiological alkaloids the name leucomaines is given.

**Wound fever** is to be placed as a form of septic poisoning. This name is applied to the condition which occurs when a recent wound is left open so that putrefaction occurs in the discharges. In a recent wound the lymphatic channels and capillary blood-vessels are laid open, and are ready to absorb any products of putrefaction which may be present. Hence, in the first forty-eight hours we may have symptoms due to such absorption, which may cease when the absorbent tissues are covered up by granulations. A similar febrile condition often occurs when pus is confined in wounds or abscesses. If the pus be in a state of tension, the products may be absorbed in spite of the protecting layer of granulations, and we may have fever as a result, but it will disappear when the pus obtains free exit.

**In the intestinal canal** the decomposition of the contents will often evolve ptomaines of various sorts. It seems not improbable that many of the symptoms of dyspepsia and biliousness are due to the absorption of such poisons.

We know that odorous (often very mal-odorous) products are absorbed from the intestinal canal and exhaled by the lungs, and in a similar way poisonous ptomaines may be absorbed. So long as they are freely excreted they will not be in sufficient



quantity to do much harm, but when the secretions are interfered with they may do so. Bouchard ascribes the symptoms in some cases of uræmia to the retention of ptomaines which would be excreted by the kidneys, if these organs were healthy.

2. **Pathogenic bacteria** produce their effects in several different ways. All of them propagate at the expense of the tissues or their secretions, and they produce poisons which may have a violent local action or may pass into the circulation and produce general effects.

Those which propagate locally multiply and invade the tissues, irritating and destroying as they proceed. An example of this is afforded by septic infection, which is to be carefully distinguished from septic poisoning. The pathogenic septic bacteria (see further on under Micrococci) are capable of invading the tissues, extending along canals and spaces. Another example is afforded by tubercular infection, in

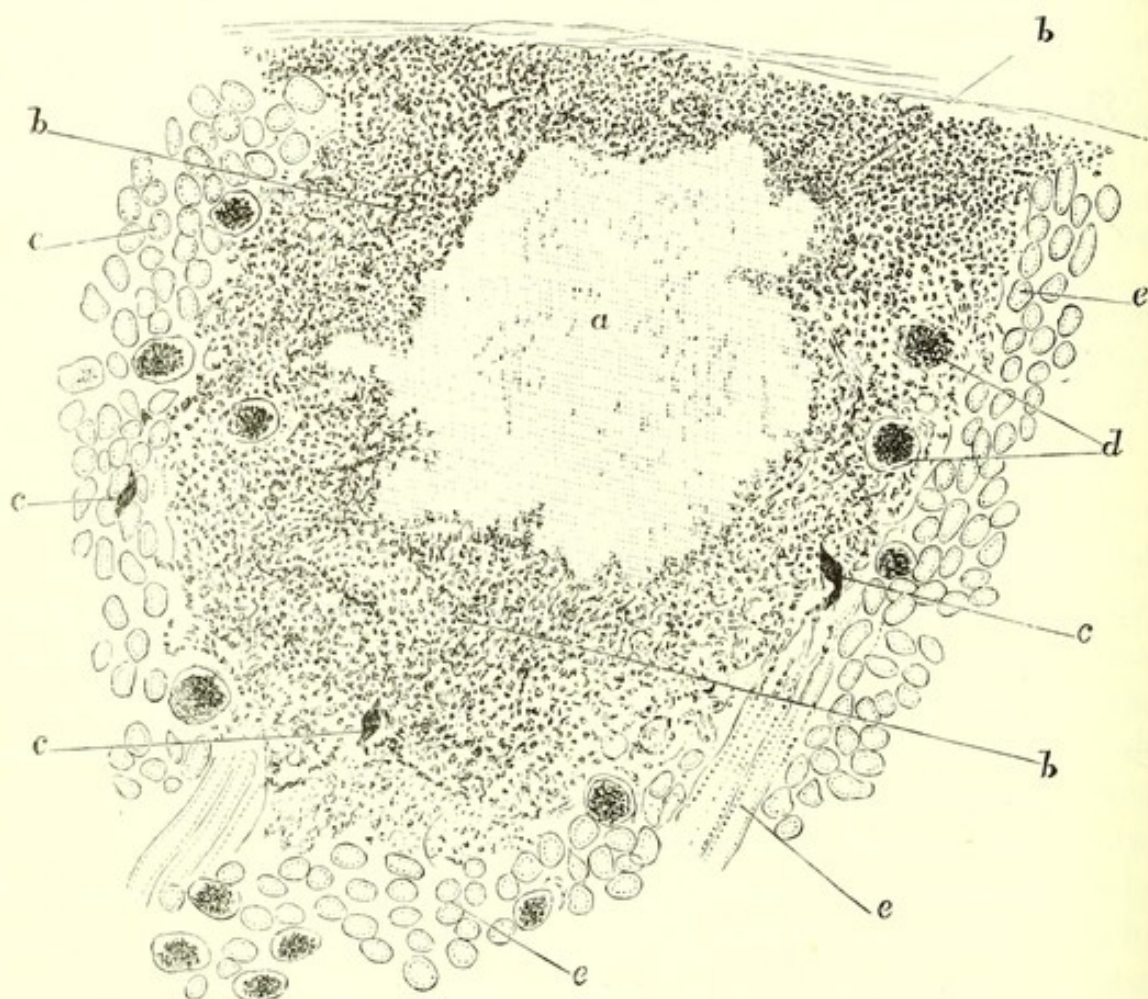


Fig. 99.—Pyæmic abscess of kidney; *a*, necrosed central part; *b*, area of inflammation, the tissue packed with pus corpuscles; *c*, capillaries filled with micrococci; *d*, Malpighian glomeruli; *e*, uriniferous tubules.  $\times 16$ .

which the tubercular bacillus propagates itself in the living tissues and advances. These locally-growing bacteria are capable of transportation, especially when they **enter the blood**. In the case of septic bacteria we may have them implanted in various regions, producing the



characteristics of **Pyæmia**. The tubercular bacteria are also sometimes disseminated by the blood leading to general tuberculosis. In these cases the transported bacteria do not generally propagate in the blood itself, but they settle in many localities where they multiply and invade the tissues as before. Diseases produced by these locally-growing bacteria will thus be **infective** in character.

The most important effects of most of the infective bacteria are **Necrosis and Inflammation**. This may be illustrated by the accompanying figure (Fig. 99) which represents a section of the kidney in a case of pyæmia. Septic bacteria have been transplanted to the kidney where they have settled and produced metastatic abscesses. The central part of the area is necrosed (*a* in figure). It shows no differentiation of tissue, and when stained with one of the ordinary nuclear staining agents it remains unstained or is homogeneously coloured. Around the necrosed part there are evidences of acute inflammation in the presence of myriads of pus-corpuscles. In this illustration the section has not encountered the actual embolism which caused the lesion, but there are at intervals two or three capillaries filled with bacteria. In Fig.

100 again we have septic bacteria in a capillary of the kidney, producing necrosis of the epithelium of the tubules. The specimen was stained with aniline brown, the effect of which is to stain the nuclei of cells and the bacteria, but if the cells are necrosed their nuclei are not stained; it will be observed that in the neighbourhood of the bacteria there are no nuclei visible.

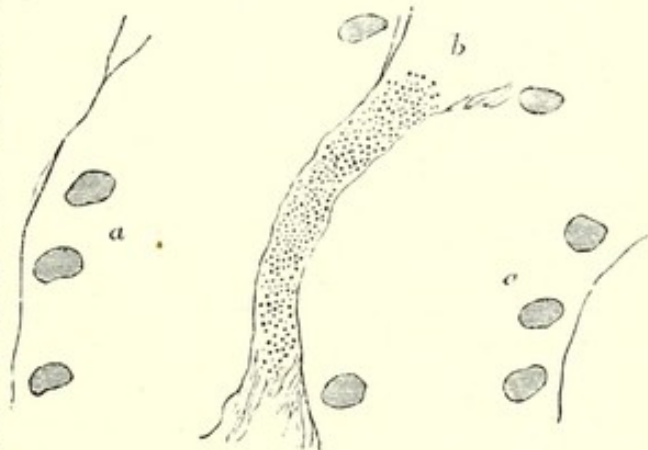


Fig. 100.—Capillary blood-vessel *b* in kidney filled with micrococci, from a case of pyæmia. There is a tubule on either side (*a*, *c*), the nuclei of which are visible except in the neighbourhood of the capillary.  $\times 650$ .

Many other examples of necrosis and inflammation produced by bacteria might be mentioned, as tuberculosis, where the necrosis is of a special kind (caseous), dysentery, diphtheria, erysipelas, etc.

The products of these bacteria pass beyond their actual seat, and the effects diminish in intensity as they pass outwards, being diluted in proportion as they depart from the centre. In Fig. 99 it is shown that in the centre of the affected area necrosis has occurred, while inflammation is visible in the peripheral parts. These products frequently pass into the blood, producing symptoms of poisoning such as those already referred to.



In the case of **Cholera**, the bacteria are, according to Koch, confined to the intestinal canal, where they evolve a violent poison, which not only irritates the mucous membrane, but, being absorbed, leads to symptoms of general poisoning.

Some of the pathogenic **bacteria propagate in the blood itself**. Some indeed are capable of general infection as well as local. The anthrax bacillus, for example, may produce a local infective disease

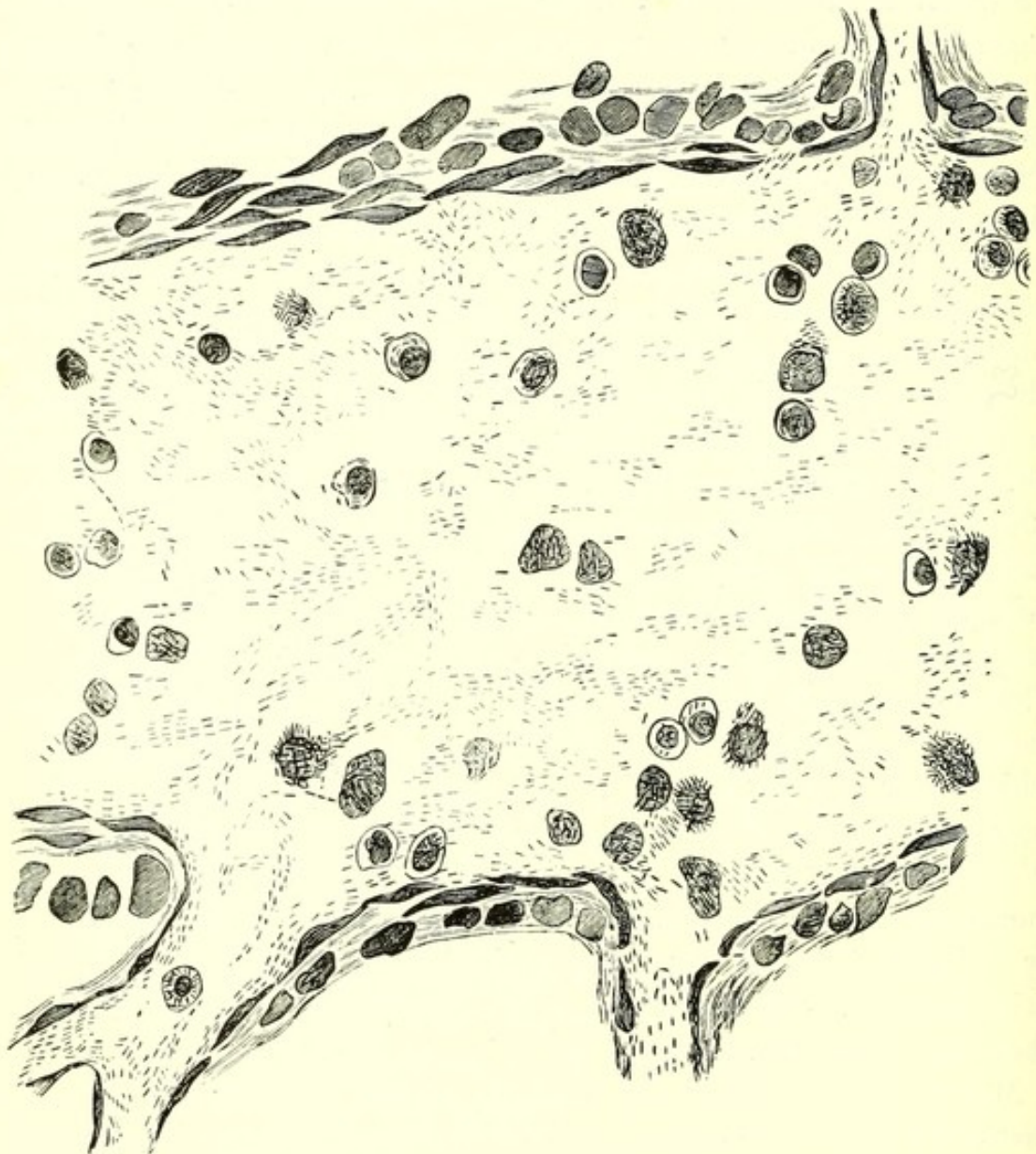


Fig. 101.—Vein of the diaphragm of a septicæmic mouse. The septicæmic bacilli are seen in great abundance, some of them attacking the white blood-corpuscles, which are occasionally reduced to masses of bacilli.  $\times 700$ . (Koch.)

(malignant pustule) as well as a general disease, in which it is present in the blood. In some cases the bacteria are present in enormous numbers in the blood. Fig. 101 for example shows a vein from the diaphragm of a mouse affected with a form of disease due to a minute bacillus (septicæmia of mice). It is seen that the bacteria are very numerous and that they are frequently attacking the white blood-corpuscles. It



is probable that in most of the acute specific fevers there are bacteria present in the blood, although in some of them they have a local habitat, and it is their products which induce the general symptoms. This is so in the case of typhoid fever where the bacteria are in the intestine, spleen, and lymphatic glands, and in small-pox where they are in the skin and mucous membranes.

### III.—THE INDIVIDUAL FORMS OF BACTERIA.

In what has gone before it has been indicated that there is not as yet a complete and satisfactory classification of the bacteria. Sufficient is known, however, to enable us to identify many of the forms although their generic relations may not be fully established. In order to determine the character and kind of a particular bacterium, it is necessary, not merely to prepare a specimen and observe it under the microscope, but also to discover its behaviour when grown as a pure culture. Hence the description of each form includes a statement of its mode of growth when so cultivated, as well as of its form. In the case of pathogenic bacteria the results of cultivation must be tested, where possible, by experiments on animals, and the position of any pathogenic form can hardly be regarded as established unless it has been found in persons affected by the disease, cultivated on nutrient media through several generations, and inoculated on animals with the result of producing the original form of disease (Koch).

In the following enumeration it is not intended to give a complete account of the bacteria. In regard to the saprophytes especially, we shall only refer to those which, from the frequency of their occurrence or from their association with man, are apparently of practical importance. In the nomenclature, the three terms descriptive of the forms of the individual cells are used as generic names, while the specific designation refers to some obvious characteristic. We have then, both in saprophytes and pathogenic forms, the three genera, micrococcus, bacillus, and spirillum.

The author has followed this simple arrangement after the example of Fraenkel, De Bary and others.

#### A.—SAPROPHYTIC BACTERIA.

1. *Micrococcus prodigiosus* is one of the commonest bacteria. It appears as blood-red stains on objects—sometimes on milk, bread, starch, etc., from which the names “blood-rain,” “bleeding host” were derived. As the blood-red spots have sometimes been regarded as portents or prodigies, the bacteria have received the name *M. prodigiosus*. The individuals are globular, oval, or short rods, and they grow vigorously on the surface of potatoes, or on nutrient jellies, and give a deep red colour. The colour is not resident in the bacteria but secreted by them. The



bacteria evidently exist abundantly in the air and readily infect any suitable medium which may be exposed. They absorb readily the aniline dyes. No pathological significance is attachable to this form.

2. **Sarcina** is a peculiar form of microbe which is now recognized as belonging to the group of micrococci. The old name is retained as it expresses the form of the bacteria. The sarcina is a micrococcus which in dividing shows lines of fission in two directions at right angles, so that it always divides into fours. The fours often remain adherent, so that we may have groups of four, eight, sixteen, or further

multiples (see Fig. 102). There are several forms of sarcinæ which may appear spontaneously in various media, their source being the air, and from these pure cultivations may be obtained. The forms are distinguished by the colour of the cultures, as white, yellow, and orange.

**Sarcina ventriculi** (Goodsir) is a form found in the stomach, especially when the organ is greatly dilated and processes of fermentation are proceeding. It occurs as cubical packets of micrococci which frequently have a brownish colour

Fig. 102.—(a) *Sarcinæ ventriculi*, (b) starch granules, and (c) fungus spores from vomited matter.  $\times 350$ .

when seen under the microscope (see Fig. 102). When cultivated on nutrient jellies, it grows in light yellow colonies. It has no special significance in the stomach, where it occurs along with other fermentative bacteria.

*Sarcinæ* have also been observed in other situations. If blood be taken fresh from the vessels into a capillary tube and preserved in a water-bath at a temperature near that of the body, then in almost every case *sarcinæ* will develop in a few days (Lostorfer and Ferrier). They first appear as globular glancing bodies and then form the regular cubical packets. They are smaller than those found in the stomach, but in nutrient media they grow to that size. *Sarcinæ* have also been found in gangrenous cavities in the lungs, and in the urine, being probably derived from the blood.

3. **Bacillus megaterium** has been fully described by De Bary, who made special observations as to spore-formation in it. It was first observed growing on boiled cabbage leaves and afterwards cultivated on jellies containing grape sugar, or meat extract. It is composed of cylindrical rods, which are often slightly bent, and contain



numerous granules. It forms spores in the granular substance, which ultimately become free.

4. *Bacillus subtilis*, Hay bacillus, is a very common form, and, as its cells are large as compared to those of most bacilli, it is readily seen, and was early discovered. From its name, hay bacillus, it will be inferred that it readily occurs in vegetable infusions, being carried by the air. It is composed of long rods, which are usually united into long threads. The bacilli are motile, possessing a flagellum at each end. Spore-formation occurs inside the rods, and as the spores enlarge the bacilli disintegrate so that the former are set free. If the spores are placed in a fitting medium they develop into rods. The membrane of the spore tears at one side and the young rod grows out of the cleft. This form grows readily on many nutrient media, forming on potatoes a whitish cream-like layer, and on other media a thick membranous layer. The hay bacillus is aërobie; deprivation of oxygen stops its growth.

The hay bacillus has a superficial resemblance to the anthrax bacillus, and it was at one time asserted by Buchner that the two forms were convertible. They present, however, several points of distinction, the anthrax bacillus for instance being motionless. The hay bacillus has been introduced into the blood of living animals, but has produced no morbid effects. It is soon removed from the blood, and deposited (according to Wyssokowitsch) in the liver and spleen, where it may remain for months, without producing any effect.

5. *Bacillus acidi lactici* is concerned in the souring of milk. When milk is withdrawn from the mammary glands and exposed to the air, it is liable to have several forms of bacteria and fungi developing in it. An almost constant contamination is that with the lactic acid bacillus, which has the double effect of converting the milk sugar into lactic acid, and precipitating the casein, the latter effect being the result of the presence of the free acid. The result occurs in pure cultivations in sterilized milk. The bacillus is a small short plump rod, mostly joined in twos, seldom in larger chains. It is motionless. The bacilli form endospores. The bacillus grows on nutrient jelly as a greyish white glittering layer, which does not liquefy the jelly.

There are probably several other bacteria which convert sugar into lactic acid, but this is the chief agent.

6. *Bacillus butyricus* (*clostridium butyricum*), the bacillus of butyric acid fermentation, is another of the forms which occur spontaneously in milk. In solutions of starch or milk sugar, it produces butyric acid, while carbonic acid and hydrogen are evolved. It is also capable of converting lactic acid into butyric acid. Coagulated casein is dissolved by this bacillus, and it is hence supposed to have to do with the ripening of cheese. The bacillus is an extremely active one, both in its movements and its growth. The rods are large and thick, resembling those of the hay bacillus. They are anaërobic, and not infrequently produce spores.

This bacillus presents, under certain circumstances, a peculiar reaction with watery solution of iodine, certain parts of the cells taking on a deep indigo blue colour, especially when cultivation has occurred in starchy media.

7. *Bacterium termo* is a name which, in view of the modern position of Bacteriology, can hardly be said to have a proper place. In putrid fluids every drop contains large numbers of bacteria, doubtless of various sorts. The commonest and most constant forms are moderately large, vigorously moving rods, and it is these which have received the name of *bacterium termo*. But attempts made to obtain pure cultivations of this form, seem to have shown that we have not here a single species, but several kinds which future research must be left to identify. The



ordinary bacterium of putrid decomposition is a short rod with a thick cell-membrane and a flagellum. No spore-formation has been observed. It grows actively on nutrient jellies, which it liquefies. On potatoes it forms a grey slimy layer.

8. *Leptothrix buccalis* is a form to which some ascribe pathogenic properties. It occurs

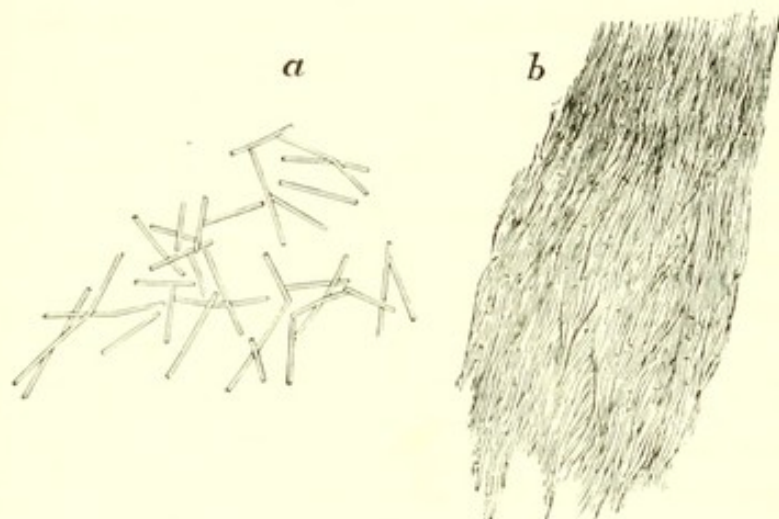


Fig. 103.—*Leptothrix buccalis* from the gums at edges of teeth. *a*, the filaments separated; *b*, masses of filaments.  $\times 350$ .

It occurs in the mouth and is constantly present in the soft white matter which lies in the corners between the teeth. It consists of long fine threads (see Fig. 103) which are deeply stained by the ordinary aniline dyes. By some this bacterium is believed to extract the lime from

the teeth and so induce caries, but it probably does not do this unless the teeth are already weakened. It is supposed to attract lime from the food and form the so-called tartar of the teeth.

## B.—PATHOGENIC BACTERIA.

### I.—PATHOGENIC MICROCOCCI.

Most of the pathogenic micrococci are characterized by the fact that they produce violent inflammations frequently resulting in suppurations. Hence they are mostly **Pyogenic**, that is, producers of pus. An endeavour has been made to form genera on the basis of the arrangement which the cocci assume in their growth. Some of them always divide in one direction, and so form more or less elongated chains, whence the generic term *Streptococcus*. Others do not divide in one determinate direction, and so form clumps or masses, from the resemblance of which to bunches of grapes, the term *Staphylococcus* has been applied. These terms must be regarded like the others as merely provisional and as expressing the present state of knowledge. All the septic micrococci stain readily with aniline dyes, especially by Gram's method.

1. *Streptococcus erysipelatis*, the micrococcus of erysipelas, is composed of perfectly globular cells of small size, which have a peculiar tendency to grow into long chains. It has been cultivated on various media, growing slowly at the ordinary temperature, and more quickly



when the temperature approaches that of the body. The cocci grow most abundantly in bouillon, and not at all on potato. In growing they form little white spots which have a characteristic appearance, and do not liquefy the gelatine.

In cases of erysipelas they are found in the lymphatic spaces and vessels, which they frequently fill out so as to form a kind of injection of them. They do not extend beyond these vessels, but their poisonous products not only produce intense local inflammation, but, passing to the blood, cause fever and other general symptoms. The bacteria are scarcely at all present in the inflamed area, but are abundant at the margins. Apparently the cocci produce a poison and subsequently perish, leaving the poison behind.

Erysipelas has been produced artificially by inoculation of cultures of this streptococcus. The most convenient experiment is to inoculate the ear of a rabbit, which leads to an acute inflammation, but without suppuration.

Fehleisen has practised inoculations in man with a view to the **treatment of certain tumours**. It has been observed that erysipelas accidentally acquired sometimes cures or stops the growth of malignant tumours. Acting on this idea Fehleisen inoculated the coccus from pure cultures in cases of sarcoma and cancer, with the result of inducing typical erysipelas. The tumours, in some cases at least, were benefited.

In ordinary cases of erysipelas the bacteria are usually derived from the air, but may be conveyed by contact. The point of entrance is probably in all cases a wound, but it may be a very trivial one.

**2. Staphylococcus pyogenes aureus.**—Micrococci are to be found virtually in all abscesses or suppurations, and the form now under consideration is the most important. It has been studied by many authors, of whom Ogston, who suggested the term staphylococcus (from *σταφυλή*, a bunch of grapes), and Rosenbach, who gave the full name, may be mentioned.

It is smaller than the coccus of erysipelas and in its growth tends to form little masses rather than chains (although Crookshank asserts that it forms chains, and hence calls it streptococcus). It is not known to produce spores, but it is peculiarly resistant to drying and heat. It may be kept dry on a cover-glass for ten days without losing its power, and it requires the temperature of boiling water for several minutes in order to kill it.

It grows at the ordinary temperature, but more vigorously at higher degrees. It grows on gelatine, agar-agar, and potato, forming in all of them bright orange-coloured masses, which have been aptly compared to layers of red oil paint. It rapidly liquefies gelatine and agar-agar.



Suppurative inflammations have been frequently produced by the artificial application of this bacterium. Garré applied it to his own person, once to a fissure at the finger-nail, and once by rubbing into his forearm. In the former case he produced a spreading suppuration, and in the latter a considerable carbuncle which took weeks to heal.

The relation of this bacterium to acute infective ostitis and to ulcerative endocarditis is very interesting. If large quantities of the culture be injected in animals they will die of poisoning in a short time, but if small quantities be used they will usually recover. However, an injury to the valves of the heart on the one hand, or to a bone on the other, will determine the settlement of the cocci in one or other of these situations, the result being a malignant infective inflammation. (See under Ulcerative Endocarditis, and Suppurative Ostitis.)

Abscesses may be produced in animals by the subcutaneous injection of cultures of this bacterium. Acute peritonitis is produced by injection into the abdomen, and acute synovitis by injection into the joints.

This micrococcus is very widely distributed, occurring in almost all suppurations, but also in the normal saliva, in water, and air. All open wounds are exposed to its inroads, and it will depend somewhat on circumstances whether it penetrates further inwards.

3. *Staphylococcus pyogenes albus* closely resembles that just described, almost the only difference being that when growing it does not produce a yellow pigment, but instead, a thick white layer like varnish. It is less common than the yellow form and apparently less malignant.

4. *Streptococcus pyogenes albus* is not infrequently found in pus, either alone or in conjunction with the *Staphylococcus aureus*. In all its characters it closely corresponds with the streptococcus of erysipelas, having similar modes of growth on nutrient media. In fact it seems doubtful whether it is not the same form, the principal point of difference being that while mice are not infected by inoculation with the micrococcus of erysipelas they are by this form.

In addition to these there are several other bacteria which have been met with in pus, but they have not been so fully worked out, and are much less common than those described above. These are the micrococcus *pyogenes tenuis* of Rosenbach, the bacillus *pyogenes foetidus* of Passet and others.

Along with the micrococci of pus may be mentioned the *Bacillus of green pus* (*Bacillus pyocyaneus*). Surgeons sometimes find that the dressings suddenly assume a bright green or blue colour, which may occur without disturbing the process of healing. This is due to a small bacillus which has found access to the secretions of the wound and has multiplied there. It is a slender rod, often united in small chains of four to six cells. It has very lively movements; no formation of spores has been observed. It has been cultivated in various nutrient media, the culture usually giving a greenish colour. The pigment has been extracted by



chloroform and separated as long needles. It has been observed also in the pus of cases in which the dressings have been stained. This pigment is named by Gessard pyocyanin. This form of bacterium has no pathological significance.

5. **Micrococcus gonorrhœæ, Gonococcus.**—This form, discovered by Neisser, is now acknowledged to be the active agent in the causation of gonorrhœa. It is a large micrococcus, which is generally found united in twos (diplococcus), the two surfaces facing each other being usually flattened.

The gonococcus is not to be detected by Gram's method, as the iodine decolorizes it. It may be stained by the ordinary aniline dyes, but it is best demonstrated as follows:—A cover-glass which has been smeared with the pus is placed for a few minutes in a concentrated alcoholic solution of eosine (best with slight heating). The eosine is then removed with blotting paper and the preparation treated for a very short time (about a quarter of a minute) with a concentrated alcoholic solution of methylblue, and then washed in water. The cocci are now seen to be blue against a red background, as the pus-corpuscles take up the eosine.

The principal peculiarity of the gonococcus is that it is found in the substance of the pus cells, often filling the protoplasm but leaving the nucleus free. It is exceedingly difficult to cultivate, success having only been obtained by the use of human blood serum, and even with this the growth has been comparatively slight. The identity of the coccus and its connection with gonorrhœa was proved by Bumm, who inoculated a culture of the third generation in the female urethra and produced a typical gonorrhœa in three days.

## II.—PATHOGENIC BACILLI AND SPIRILLA.

1. **Bacillus anthracis.**—This is one of the best known and most widely diffused pathogenic forms, occurring, as it does, both as a saprophyte and a parasite. As a parasite it gives rise to the conditions variously known as **Splenic fever, Anthrax, Charbon, Malignant pustule, and Wool-sorter's disease**, and it is sometimes described under the name *Bactérie du charbon*. It mostly occurs in animals, but is occasionally communicated to man.

The bacteria are rods of considerable size, generally united end to end so as to form longer threads. Each bacillus is about as long as the diameter of a red blood-corpuscle. The extremities of the bacilli are abrupt, and even

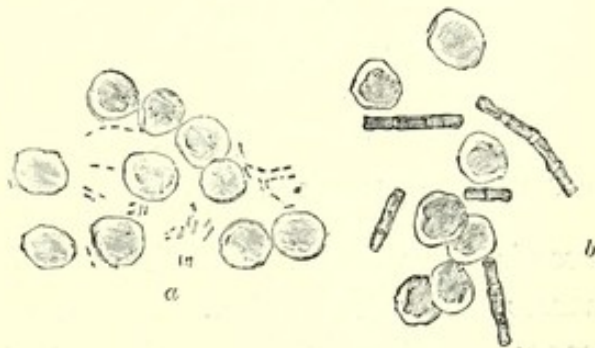


Fig. 104.—Bacillus of anthrax (*b*), and of the septicæmia of mice (*a*).  $\times 700$ . (KOCH.)



slightly concave, so that when two are united there is an oval space which is very characteristic. (See Fig. 104.)

The bacillus may be cultivated in many different nutrient media, nutrient gelatine, agar-agar, potato, etc. It grows readily in neutralized or weakly alkaline urine and in infusions of animal substances in general. On potato it grows very vigorously, forming a dry yellowish creamy layer.

While in the animal body the bacilli are either single or form short threads of two or three, but when grown on the surface of nutrient media so as to be exposed to the oxygen of the air they grow out into **long threads**, which at first sight appear to be continuous, but when properly displayed are seen to be composed of individual bacilli. The

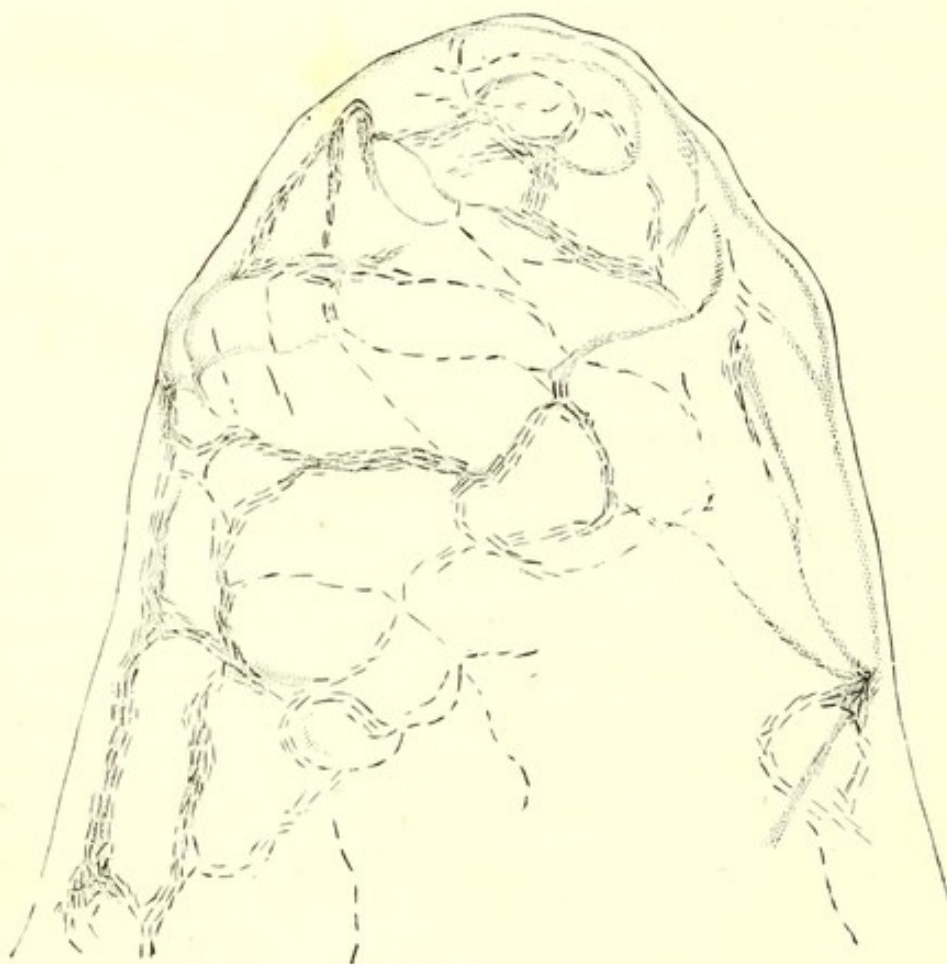


Fig. 105.—Villus of intestine in anthrax, the bacilli visible as minute threads.  $\times 250$ . (Koch.)

threads have a habit of interlacing so that interwoven or matted masses are thus formed. In these threads the **formation of spores** occurs when the circumstances are favourable. This requires a temperature not lower than  $18^{\circ}$ – $20^{\circ}$  C. and not above  $33^{\circ}$ – $34^{\circ}$ , while about  $30^{\circ}$  is the most favourable temperature. As a preliminary to spore-formation the bacilli become granular, and there afterwards appear glancing refractive bodies which gradually grow into oval spores. The spore



lies in the middle of the bacillus, and there is never more than one in each. The bacillus by and by disappears, leaving the spore free. If placed in a favourable position the spores germinate and grow into the regular rods. The spores contrast with the bacilli in being very permanent, as they resist drying and decomposition. Pieces of silk thread impregnated with spores from a culture on potato may be kept for years perfectly capable of germination.

The anthrax bacillus requires careful staining, but if due care be taken a watery solution of almost any aniline dye may be made to serve, or Gram's method may be used for double staining. The spores are more difficult to stain. If a cover-glass preparation be first made then it may be floated for about twenty minutes on the surface of a hot alcoholic solution of fuchsine, decolorized in weak hydrochloric acid and again stained with methylblue. The bacilli are stained blue and the spores red.

The anthrax bacillus is capable of transmission to a large number of animals, although considerable differences of susceptibility exist. Mice are the most readily infected, and next to them stand guinea-pigs, rabbits, sheep, and cattle. Rats are with difficulty affected, and dogs, birds, and frogs are scarcely susceptible. Frogs may be affected if after inoculation they are kept for a few days at the temperature of the brood-oven. Men are susceptible, but there is not infrequently after inoculation a local disease (malignant pustule), the bacilli being present in the exudation of the pustule.

The disease may be induced experimentally either by inoculation into the skin or mucous membranes or by introduction into the alimentary canal. Inoculation of the blood of an animal which has died of anthrax, induces the disease in a susceptible animal, or it may be induced from an artificial culture. In order to its production by the alimentary canal, cultures containing spores must be used, as the bacilli are destroyed by the gastric juice. When an animal is infected it shows the symptoms of an

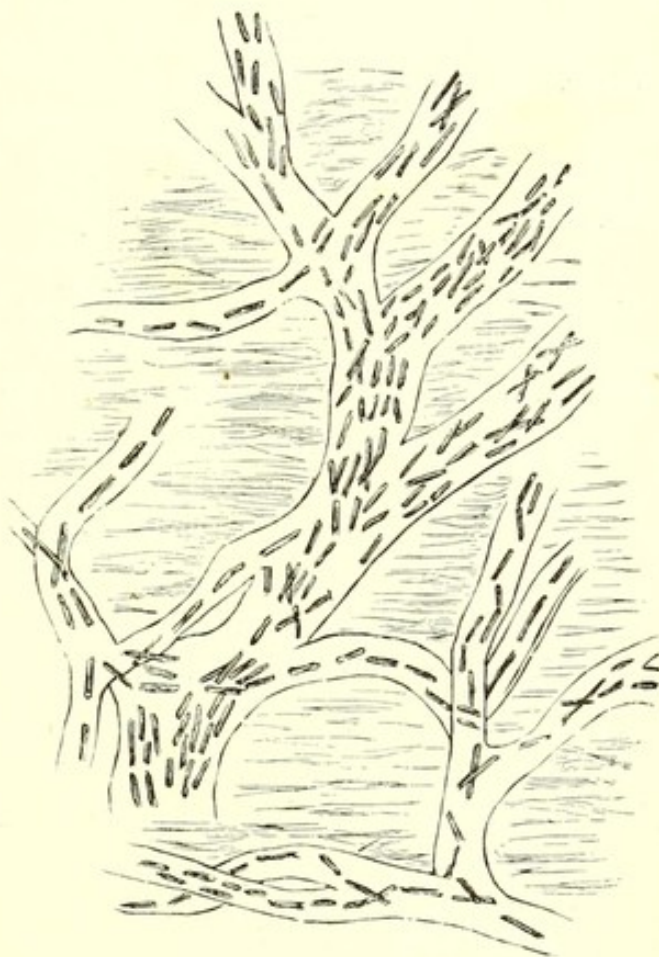


Fig. 106.—From same preparation as Fig. 105, more highly magnified. The bacilli visible as definite rods.  $\times 700$  (Koch.)



acute specific fever, which usually proves fatal in a few days. The bacilli are found in enormous numbers in the blood, and as a rule they are found only inside the blood-vessels. So abundant are they in some cases that a preparation stained in the proper way so as to render the bacilli prominent will appear as if the vessels were injected with a coloured material. (See Figs. 105 and 106.) It is chiefly in the capillaries that they are to be found, and they are most readily observed in the spleen, the villi of the intestine, and the glomeruli of the kidney. In the latter situation there is not infrequently rupture of the glomeruli and the passage of blood and bacilli into the uriniferous tubules. The bacilli, however, do not extend downwards further than the convoluted tubules.

**Spontaneous infection** with anthrax occurs in animals mostly with their food. It originates chiefly in cattle and sheep, and they acquire it in grazing. There are certain parts of the country where the disease is endemic, the grass apparently containing the microbe. As this infection is by the alimentary canal it is necessary that spores should be present. The bacillus grows on all sorts of vegetable and animal matter on which it may be planted by the blood or discharges of infected animals. Under suitable conditions of temperature it develops spores. It is important that such infection should be as much as possible prevented, and this should be borne in mind in the disposal of the bodies of affected animals. No spore-formation occurs without oxygen, and if the body is not opened the bacilli in the blood will perish. Hence such bodies should, where possible, be buried without opening, and at any rate, precautions taken against contaminating the ground with blood or tissues.

**In man** the disease has mostly been communicated by handling the tissues of animals which have been affected. Sometimes spore-formation will occur in such tissues, the bacilli growing on the tissue when exposed to the air and producing spores. Hence spores may adhere to hair, wool, or other material, and be conveyed long distances. Most cases in man have occurred amongst workers in hair or wool (hence the name of *wool-sorter's disease*). In several cases occurring in Glasgow, the hair had come from Russia. The infection usually occurs by the accidental inoculation of a wound or scratch, but it may be by the intestinal canal.

**Attenuation of the virus** was first attempted by Pasteur, and was attended by very striking results. By cultivating the bacillus at a temperature of 42°–43° C. for about twenty-four days, the infective power is destroyed, while a shorter period suffices to weaken it. When the virus, weakened by various periods of cultivation at a high temperature, is cultivated at the ordinary temperature, it remains weak in infective power, and so a permanent **Vaccine** is procurable. The bacilli have been also weakened when cultivated under the pressure of eight atmospheres (Chauveau), and also by passage through different species of animals. Thus the blood of white mice dead of anthrax produces when inoculated in sheep only a transitory illness, and the blood of guinea-pigs inoculated in cattle does not usually kill, although producing a serious illness.



When animals are vaccinated with the attenuated virus they have a mild form of the disease and acquire a certain degree of immunity to the more severe forms. The practical importance of this fact is seriously diminished because the immunity appears to be only from infection by inoculation, and does not extend to infection by taking the spores into the alimentary canal. The latter is, however, the most frequent mode of infection in spontaneous anthrax.

**Post-mortem appearances in anthrax.**—There is not usually much to be seen at the point of inoculation, but over the whole abdominal surface there is generally a marked jelly-like œdema. The subcutaneous tissue is friable and sometimes infiltrated with blood, but there is no development of gas. The superficial muscles are pale and almost like cooked muscle. The spleen is greatly enlarged, dark in colour, soft and brittle. The liver shows a moderate cloudy swelling.

**2. Bacillus of malignant œdema.**—This is a bacillus somewhat like the bacillus anthracis, and it is also probably identical with Pasteur's "vibrions septiques" which he found in his "septicæmie."

Malignant œdema has been observed in man in consequence of severe wounds, compound fractures, etc. There is a cutaneous emphysema, decomposition and œdematous swelling of the superficial muscles, and death generally results in a few days. The parts have been infected with the bacillus under consideration. The author met with bacilli closely resembling this form in a case of cancrum oris.

The bacterium is a slender rod, somewhat thinner than the anthrax bacillus and with rounded ends. It tends to grow into long threads even in the body, and it is motile. The bacilli are **strictly anaërobic** and they bear spores under circumstances which are not well understood. Their source in experimental observations is various kinds of decomposing matters, dust, garden earth, etc., and they are evidently widely dispersed. They are cultivated on various nutrient media in an atmosphere of carbonic acid or otherwise protected from the air.

The bacilli produce disease when inoculated in mice, guinea-pigs, rabbits, etc., and this occurs whether their source be such external matters as garden earth, or pure cultivations. Death usually results in twenty-four to forty-eight hours. Around the point of inoculation the subcutaneous tissue and superficial muscles are infiltrated with a dirty red stinking fluid in which are bubbles of gas. The bacilli are found in the œdematous fluid and to a very small extent in the fluids of the tissues; none are visible in the blood. The bacilli in fact remain in the subcutaneous tissue and on the surface of organs, not penetrating into the parenchyma and not extending to the blood-vessels. It is to be added that they grow vigorously after death, penetrating into the substance of organs and into the vessels. In the mouse the bacilli penetrate more



markedly during life into tissues and blood-vessels and the appearances approach to those of anthrax.

3. **Bacillus of typhoid fever.**—Eberth, and after him several other authors, have, in cases of typhoid fever, observed, in the closed follicles of the intestines, and in the lymphatic glands and spleen, peculiar bacilli, which were distinguished by their peculiar arrangement and defective reaction to aniline dyes. Gaffky has followed out the investigation and given a complete account of this form.

The bacteria are small rods with rounded ends, which are usually single or in pairs in the tissues, but may grow into longer threads. They possess very active mobility. Gaffky has described appearances which are probably those of spore-formation. The bacilli grow on various nutrient media, but especially on potato. They form on potato a thickish layer which to the naked eye scarcely differs from the surface of the potato, but a small portion is seen under the microscope to present vast numbers of the active bacteria. This growth on potato is very characteristic, and, occurring as it does at the ordinary temperature, is very important as showing that these bacteria are not obligate parasites, but that they can grow outside the body. It has also been shown that they grow vigorously in milk (Wolffhügel), and that they may be preserved and even grow in water.

The typhoid bacillus is difficult to stain, giving almost no colour with watery solutions of ordinary aniline dyes. With Löffler's alkaline methylblue, and with Ziehl's and Neelsen's carbolic acid and fuchsine solution, they are readily stained. Sections should be left twenty-four hours in the solutions. The preparations should be washed with water, and not with alcohol.

The bacilli have been transmitted to animals by Fraenkel and Simmonds and by Seitz. The former injected emulsions of cultures into the veins of the ears of rabbits; the latter introduced them into the alimentary canal. The simple administration of them with the food was not enough; but, by methods similar to those used by Koch to induce cholera in animals, Seitz produced a disease in animals similar to typhoid fever, of which most of the animals died. The bacilli were found abundantly in the contents of the intestine, and to a small extent in the organs, but the blood was free.

In man the bacilli have been found in the intestinal contents, in the Peyer's glands, the lymphatic glands, and the spleen, but only to a slight extent in the blood. In the tissues mentioned they grow in little masses with intervals between. They are not found in every case, diminishing apparently as the disease advances. Hence they are uniformly met with in recent cases, in which numerous clumps of bacilli will be found in the swollen Peyer's patches and glands. In later cases,



where ulceration has occurred, they will be found in the deeper layers of the patches, in the mucous membrane and muscular coat beneath the ulcer.

As the bacilli extend to the blood, it may be that the general symptoms of typhoid fever are due to this; but, as the extension is apparently not great, it may be that the bacilli form a poison in the intestine, glands, and spleen, which, reaching the blood, produces these symptoms.

The propagation of typhoid fever occurs by multiplication of the bacilli outside the body and their subsequent ingestion. There are abundant bacilli in the fæces of the patients, and they have been detected in the albuminous urine in severe cases. Considering how readily they propagate in milk and other media outside the body, it is not difficult to understand how they should frequently infect the ingesta and produce the disease in man. There are many epidemics which have been traced to contaminated milk.

4. **Spirillum of relapsing fever, *Spirillum Obermeieri*.**—This form is an active spiral bacterium which is found abundantly in the blood of persons affected with relapsing fever. (See Fig. 107.) It is present only during the acute attacks, disappearing in the interval, to return when the relapse occurs. It is readily stained by ordinary watery solutions of the aniline dyes. The disease has been communicated to man and to apes by the inoculation of blood containing the spirilla. The spirillum is found only in the blood, and Koch has observed it in the blood of the brain, liver, and kidneys of an ape, which was killed during the attack.

These spirilla have not been cultivated artificially.

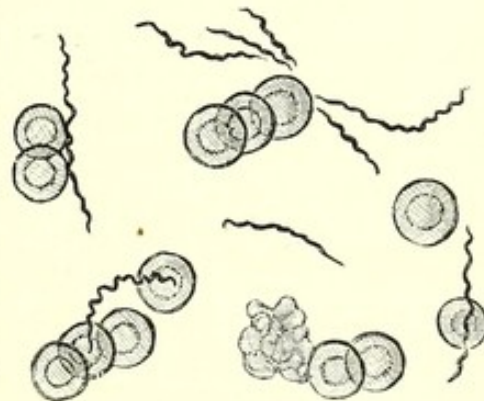


Fig. 107.—Blood from case of relapsing fever, showing corpuscles and spirillum *Obermeieri*.  $\times 750$ . (CARTER.)

5. **Bacillus of cholera, *Comma bacillus*.**—The observations of Koch have, in the case of this bacterium, afforded a strong basis for its connection with Asiatic cholera.

The bacillus is a short, thickish rod, about half the length of the tubercle bacillus, and somewhat broader. It is curved on its long axis so as to resemble a comma (but without the head of the printed comma). Sometimes there are two, end to end, with their curves in opposite directions, so as to resemble the letter S. They sometimes grow into longer spiral threads, from which they have been ranked as spirilla. As the spirals are not single cells, but made up of rows of commas, this conclusion is scarcely warranted, and the bacteria may still be



retained as curved bacilli. The bacilli are very actively mobile. In cultures an appearance has been observed which looks like formation of arthrospores, but as no permanent condition of the bacillus is known this is probably not so. The bacilli are readily stained by watery solutions of the aniline dyes, but the best is a strong watery solution of fuchsine. They require longer exposure than usual, not less than ten minutes for cover-glass preparations and twenty-four hours for sections. The fluid may be heated. Gram's method is not available, as the bacilli are decolorized.

Cultures of the bacilli in gelatine show very characteristic appearances. The growing bacilli liquefy the gelatine slowly, and form typical depressions and spaces in it. They also grow on potato, in milk, and even in water. They were found by Koch abundantly in the water of a tank in India.

Cholera does not occur in animals, but Koch has succeeded by a special method in inducing the bacilli to grow in the intestinal canal of guinea-pigs. In order to this he overcomes the acidity of the stomach by an alkaline carbonate, and at the same time controls the peristaltic action of the intestine by opium, and then finds that cultures introduced into the stomach survive and propagate. The animal dies in about two days, and the intestine is found to contain large quantities of fluid faeces which teem with comma bacilli. An accidental contamination of one of Koch's pupils during a course of bacteriology resulted in an attack of cholera, thus affording a proof of the identity of the bacillus.

In cases of Asiatic cholera Koch has always found the comma bacillus in the rice-water evacuations and intestinal contents. The mucous membrane of the intestine is red and swollen, especially in the region above the ileo-cæcal valve. The bacilli are present not only in the contents, but in the tubular glands, sometimes penetrating between the epithelium and the basement membrane. The bacillus does not penetrate beyond the intestine, but it apparently produces a violent poison, which irritates the intestine and, being absorbed, leads to weakening of the heart, lowering of the temperature, muscular cramps, etc.

The bacillus is destroyed by drying, and is apparently transmitted by means of water, milk, etc.

Klein, while acknowledging that the comma bacillus is always present in the intestine in the early stages of cholera, and agreeing in the opinion that it is an important diagnostic guide, holds that it is not proved to be the cause of cholera. (See Klein's "Bacteria in Asiatic Cholera," 1889.)

Other comma bacilli have been described, of which the most important is that of Finkler and Prior, who obtained it in cases of cholera nostras. The individual bacilli closely resemble those of Koch, but their mode of growth in cultures is quite distinctive. These bacilli have pathogenic characters, but are not so active as Koch's.



Daneke has obtained a comma bacillus from old cheese, which had also pathogenic characters in guinea-pigs.

6. **Bacillus of pneumonia.**—Pneumonia is now generally recognized as presenting the characters of a disease due to a morbid poison, and observers have sought for a special form of bacterium in it. It cannot be said that the existence of such a form has been absolutely demonstrated, but there are two candidates for the position. These are the "Pneumococcus" of Friedländer, and the "Pneumoniococcus" of Fraenkel, both of which although called cocci are really bacilli.

(a) **Friedländer's Pneumococcus** is a rod-shaped bacterium which is sometimes very short so as to be like a micrococcus, but sometimes grows to a considerable length. It has the peculiarity that, under certain circumstances within the living body, the cell-membrane swells so as to produce a capsule around the bacterium. This capsule does not stain by the ordinary methods and the result is that the bacterium is surrounded by a clear halo. There may be one or more cells in each capsule. A similar capsule has been observed in other bacteria, so that although distinctive it is not absolutely so. In cultivations outside the body the capsule is absent.

The capsule is not always visible even in specimens obtained from the lung or sputum. It is best displayed, according to Friedländer, by placing sections of the lung for twenty-four hours in a solution composed of fuchsine (or gentianviolet) 1 part, water 100, alcohol 5. After this they are rinsed in alcohol and then placed for a couple of minutes in a 2 per cent. solution of acetic acid, and afterwards treated with alcohol, oil of cloves and Canada balsam, in the usual way. Gram's method is not applicable to this bacterium as it decolorizes in the process.

When cultivated in a test tube in nutrient gelatine, the bacteria grow along the needle track, especially at the surface, so that the culture has the shape of a round headed nail. They also grow readily on agar-agar and potato.

Experiments on animals with these bacteria were unsuccessful in the case of rabbits. They were partially successful in guinea-pigs and very strikingly so in mice. In the latter an emulsion of the bacilli was injected through the thoracic wall into the lungs, with the result of producing a violent inflammation, in the products of which numerous bacilli were found. When the animals were made to inhale the emulsion, the result was less successful.

The relation of the pneumococcus to pneumonia is still matter of discussion and its significance is by many seriously questioned. On the one hand the operation by which the experiments were made in mice is a very serious one for such small animals, and on the other, the bacteria are not infrequently absent from the sputum and lungs of persons affected with pneumonia.

(b) **Fraenkel's Pneumonia-coccus** is a short bacillus, which apparently does not grow into long rods. It is commonly in twos, end to end, and occasionally forms chains of five or six. It differs from Friedländer's in being generally shorter, and in never forming elongated rods. It possesses a capsule when found in the body, and on the whole closely resembles Friedländer's pneumococcus, although behaving differently in cultures and when inoculated in animals. It is also strikingly different from Friedländer's in respect that by Gram's method it is readily demonstrated while the other is not.

These bacteria are readily cultivated on nutrient media. When cultivated in a



test tube the needle track shows numerous little round granules like those of the streptococcus of erysipelas.

Experiment shows that this form is very virulent in several animals, especially mice, guinea-pigs, and rabbits. If the bacteria be injected under the skin of a rabbit, they cause an acute febrile disease resulting in death in twenty-four to forty-eight hours. The bacilli have increased enormously in the blood so that in every organ the blood-vessels contain bacilli with their capsules.

This bacillus has a very limited existence when cultivated outside the body, so that in four or five days a culture in agar-agar, for instance, dies. It is therefore almost entirely parasitic. It appears also that in cultures it rapidly loses its virulence, and does so the sooner if the culture be at an elevated temperature.

It cannot be said that either of these bacilli yet holds the field. Fraenkel's form seems to be more virulent and it has been more fully investigated, but it is not known in what proportion of cases it may be present. It is not improbable that pneumonia depends on several different bacteria of which these may be two.

**7. Bacillus of diphtheria.**—Diphtheria presents all the characters of an infective disease with a local seat and also extension to the system generally. Many observers have sought for bacteria both in the local lesion and in the internal organs, but the results cannot yet be said to be satisfactory. Oertel in 1871 described a micrococcus which he believed to be the cause of the disease, but this is now generally recognized to be at most an accidental contamination. More recently Loeffler has described a bacillus, for which he does not claim more than a probability that it is the essential diphtheritic bacterium.

Loeffler's bacillus is a rod about the same length as the tubercle bacillus, but about twice as broad. It is found in the superficial layers of the exudation, in groups, and does not penetrate the mucous membrane. It is not mobile and has no spores. It does not stain well with the ordinary aniline dyes, but with Loeffler's alkaline methylblue solution it assumes a bright blue colour while the rest of the structures are faintly stained.

The bacillus has been isolated and cultivated on nutrient media. It grows best on blood-serum prepared in a special way, forming a thick white opaque layer. It does not grow on potato.

Inoculation of animals has succeeded in birds, such as sparrows, pigeons, and hens, and also in rabbits and guinea-pigs. Subcutaneous inoculation leads to local inflammation followed by general disturbance. Introduction into the trachea produced a formation of false membrane as in diphtheria.

This seems a pathogenic bacillus, but Loeffler himself questions if it be the real one of diphtheria, chiefly on the ground that he has not always found it in that disease, and that when applied to the uninjured mucous membrane it has no effect.

### III.—BACTERIA OF SPECIFIC NEWFORMATIONS.

**1. Bacillus of tubercle.**—The great frequency of tuberculosis both in man and animals renders the discovery of the bacterium by Koch one of the most important results of science in this century. The merit of



this discovery is the greater as the tubercular bacillus is peculiarly difficult both to observe in its usual seats and to cultivate.

The bacillus is a thin, rod-shaped cell, rather shorter than the diameter of a red corpuscle. It is often slightly curved or bent at an obtuse angle. The bacilli are mostly seen singly, but occasionally in pairs, more rarely in longer threads. They do not possess power of movement. They usually present beads (see Fig. 52, p. 198), and these have been generally regarded as spores, although Crookshank questions this inference. The remarkable persistence of the tubercular bacillus, and the manner in which it retains its infective powers when dried or when kept in putrid fluids, seem to indicate that there must be some more permanent condition, probably the spores mentioned.

The staining of the tubercular bacillus may be accomplished in various ways. The material to be stained may be either the discharges from tubercular lesions or the tissues affected with tuberculosis. Of the former the sputum from phthisis pulmonalis is most frequently the subject of examination.

In examining sputum it is important to choose a portion which has actually come from the lung, and not merely the clothing of mucus which the sputa obtain from the bronchial mucous glands. A portion of the sputum may be poured into a watch-glass and the latter placed on a black background. In the midst of the sputum will be found yellow rounded masses which have come from the lungs. A small portion of one of these should be separated with needles and placed on a cover-glass. Another cover-glass is placed on the top of it and the piece of sputum squeezed between the two till it forms a thin film. By gliding the two cover-glasses asunder we obtain two preparations, each consisting of a thin film of sputum. This should be dried in the air and then passed through the flame of a spirit lamp three or four times. It is then ready to be stained.

The primary staining fluid may be Ehrlich's or Ziehl-Neelsen's. The former consists of water shaken up with aniline oil (Aniline oil 5, water 100, shake for some minutes, then filter through moistened paper), to which a strong alcoholic solution of fuchsine or gentianviolet is added (about 5 parts to the 100). Ziehl-Neelsen's fluid is simpler, more permanent, and acts more quickly. It consists of water 100, crystalline carbolic acid 5, alcohol 10, and fuchsine 1. The sputum or section is placed in the solution and left, in the case of Ehrlich's fluid, for twelve to twenty-four hours, but the time may be curtailed by heating the solution till steam is seen to rise, so that for sputum a quarter of an hour will suffice. In Neelsen's fluid, even in the cold, a quarter to half an hour does for sputum, and an hour for sections. The deeply stained material is then treated with nitric acid (1 to 4 of water) or sulphuric acid (20 per cent.) till it has lost its colour and become greenish or brownish. It is then washed in 70 per cent. alcohol (ordinary methylated spirit) till it gives off no more colour. It may then be double stained with Bismarck brown (in the case of gentianviolet) or methyl-blue (where fuchsine has been used). A very useful modification of the process has been suggested in which the two last steps are combined. After removing from the fuchsine solution, the cover-glass is placed at once in Gabet's solution, consisting of sulphuric acid (25 per cent.) 100, methylblue, 2. In this solution the acid extracts the colour from all except the bacilli, while the blue dye stains the structures which are extracted by the acid.



In artificial cultures the bacillus is very slow of growth and difficult of cultivation. It grows scarcely at all on ordinary nutrient jelly, and the best medium is solidified blood serum. Nocard has recently introduced a medium of cultivation, consisting of ordinary bouillon to which is added glycerine in the proportion of 6 to 10 per cent., and which is solidified with gelatine or gelose. The bacilli require for their growth a temperature above  $30^{\circ}\text{C}$ . and under  $42^{\circ}$ , and the best is near the temperature of the body, namely,  $37.5^{\circ}$ . In about ten to fourteen days after implantation the growth first appears as dry whitish scales, which are entirely superficial. Under the microscope the scales are seen to be composed of colonies of bacilli, which, from their arrangement, form curved lines, some of them like the letter S (see Fig. 108). These

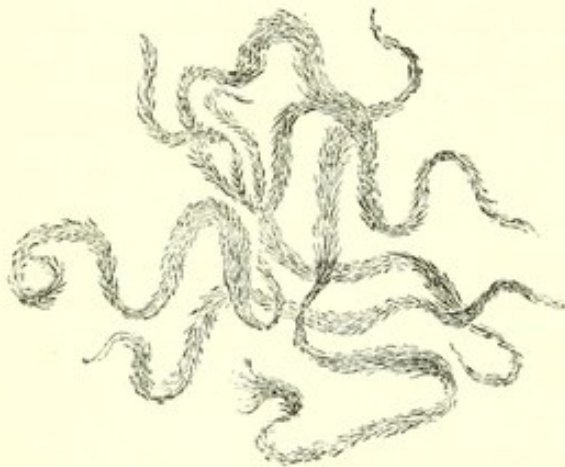


Fig. 108.—Tubercular bacilli, forming typical growths.  $\times 700$ .

bacilli give the characteristic reactions with the staining fluids mentioned above. The growth goes on for three or four weeks, and remains superficial all the time, the jelly not being liquefied. The culture may be propagated through many generations, the bacilli retaining their morphological and pathogenic characters.

Tuberculosis is producible in animals by administering either the products of disease (such as dry sputum or portions of caseous structures) or the results of cultivations. It has been induced by inoculation under the skin or into the anterior chamber of the eye, by injection into the serous cavities or into the veins, by inhalation and by ingestion with the food. Except when directly introduced into the blood, there is first a local tuberculosis, which may be followed by a generalization of the disease.

In man the bacilli are introduced accidentally by inhalation, by the food, or by inoculation. When they attack the skin they produce lupus, when they affect the lungs, phthisis pulmonalis, and when introduced with the food they may affect the intestinal canal, but are frequently carried to the lymphatic glands.

In multiplying in the living body they produce the results already described under Tuberculosis, but as these results are produced not directly by the bacilli, but by the poison evolved by them, the actual number of bacilli is not always in proportion to the effects. They are



to be found particularly in the advancing lesions, and they are present especially in the giant-cells, but also in other cells and between the cells.

In view of the fact that the tubercular bacillus is an obligate parasite, being only with difficulty cultivated under special arrangements, it will be obvious that tuberculosis can only be produced by communication, direct or indirect, from person to person, or from animal to animal. If the parasite could be got rid of the disease would cease. The bacillus is abundantly present in sputum, urine, pus, etc., in persons who are affected with the various forms of tuberculosis. The bacilli thus set free are scattered over the world, and retain their powers of growth when they find a fitting seat. We can only hope to stop the disease by a system of isolation of persons affected, coupled with disinfection of the discharges. The fact that about a seventh of the deaths in the community are due to tuberculosis ought surely to justify such isolation, with which indeed the efficient treatment of the disease would be combined. The isolation of lepers has not improbably had a good deal to do with the disappearance of this disease from most parts of Europe.

2. **Bacillus lepræ, Bacillus of leprosy.**—This form of bacterium was first observed by Hansen in Norway, and the observation confirmed by Neisser. It is a rod which closely resembles the tubercular bacillus, and, like it, is motionless. It presents clear spots, which remain uncoloured when the bacillus is stained, but it is not known whether these are spores or not. It is stained by the same process as the tubercular bacillus, but takes on the colour more readily, and it presents the important difference that it is stained by ordinary watery solutions of the aniline dyes, especially fuchsine and methylviolet.

The bacillus is found in all cases of leprosy in the lesions in the skin, nerves, and elsewhere. It occurs in very large numbers, so that when stained by Gram's method a section will have a decided blue colour from the stained bacilli alone. The bacilli are in cells, and so present themselves in little rounded clumps (see Fig. 57, p. 210).

The bacillus has not been cultivated artificially, but Melcher and Ortmann have succeeded in inoculating the anterior chamber of the eye in rabbits with pieces from a freshly excised leprosy nodule. Characteristic newformations appeared in almost all the internal organs, especially the cœcum, lymphatic glands, spleen, and lungs. In these the bacilli were abundantly present.

In man there is now sufficient evidence of direct communication of the disease from one person to another, but the period of inoculation is very long and the mode of communication often difficult to trace. The bacilli are found in the lesions, which affect the skin and the nerves chiefly, but may extend to almost all the organs of the body.

3. **Bacillus of syphilis.**—Lustgarten has described a bacillus whose connection with syphilis still stands in need of proof. He has found it



in the secretions and in the tissues affected with syphilis. It is a rod somewhat resembling the tubercle bacillus, and is always found inside cells. It presents peculiar relations to staining agents, requiring a complicated treatment with gentianviolet, permanganate of potassium, and sulphuric acid.

Giacomi ("Correspondenzbl. f. schweiz. Aerzte," 1885) describes a simpler method. A cover-glass preparation or section is placed for a few minutes in a heated fuchsine solution, afterwards washed in water containing a few drops of chloride of iron, and then decolorized in a concentrated solution of chloride of iron. The bacilli remain red, while all other bacteria are decolorized.

The position of this bacillus is very doubtful, on the one hand because a number of observers have sought for it in vain in syphilitic lesions, and on the other because a similar bacillus has been found in the normal præputial and vulvar smegma.

**4. *Bacillus mallei*, *Bacillus* of glanders.**—This form is abundantly present in the lesions of glanders, is readily cultivated, and easily communicated to animals.

The bacillus is a rod somewhat like the tubercular bacillus, but slightly thicker and shorter. It is generally slightly curved, and usually single or in pairs. It is not motile. It is not certain that it produces spores, but it preserves its virulence in the dry state for months.

The bacillus may be stained with the ordinary aniline dyes, especially with fuchsine, but better results are obtained with Loeffler's methylblue. Loeffler also gives a somewhat complicated method which yields very striking results.

The bacilli have been cultivated on nutrient agar-agar and blood serum, where they give a whitish or yellow culture. On potato there is a very characteristic growth, at first amber coloured, then getting darker in colour till it assumes a reddish brown or red tint. A temperature between 25° and 42° C. is necessary, the best results being obtained between 30° and 40°.

Animals are readily inoculated, but there are peculiar differences in susceptibility. As the disease is mainly one of horses, it is natural to find that these animals and asses are readily affected. Field mice and guinea-pigs are readily affected, but white mice, house mice, pigs, and cattle are hardly affected. There is always a local action at the point of inoculation, from which there may be a gradual extension, but not by the blood.

In man an accidental inoculation sometimes occurs, resulting in local abscesses, with secondary extension to the mucous membranes, joints, etc.

A remarkable fact is, that in cultivating the bacillus, it gradually loses its infective power, so that in the fourth or fifth generation it has become



so harmless that it is necessary to inoculate much larger quantities, and the effect is merely local. By further cultivation the bacilli lose their virulence. We may infer from these facts that the bacillus is an obligate parasite.

5. **Actinomyces or Ray fungus** has usually been regarded as belonging to the fungi proper, but recent attempts at cultivation, although only partially successful, seem rather to place it among the bacteria. It is said that cultures have been successfully carried out on nutrient gelatine and agar-agar, the best results being obtained at a temperature of 33°–37° C. The cultures showed threads, rods, and cocci, the ray form being regarded as only occurring when the soil is unsuitable to perfect growth. (See Fig. 61, p. 215.)

Besides the bacteria described above, there are a number of other forms, for which pathogenic characters have been asserted, but in regard to most of them further proof is required before their position can be admitted. Amongst these may be mentioned the *Bacillus scarlatinæ* of Edington, which this observer believes to be the cause of scarlet fever, the bacillus of Rhinoscleroma of Frisch, and the bacillus of acne contagiosa of Dieckerhoff and Grawitz.

**Micrococcus tetragenus** is a bacterium found by Koch in the contents of tubercular cavities in the lungs, and also in normal sputum. It has been investigated by Gaffky. It consists of round cells, which, when found in the living body, are arranged in fours, each group having a clear capsule around it. These groups somewhat resemble sarcinæ. The bacteria have been cultivated in nutrient media, but they lose the peculiar arrangement. They have been used for inoculation in animals, and are violently pathogenic in white mice and guinea-pigs, while house and field mice and rabbits are insusceptible. The cocci are found in the blood and tissues of all organs.

#### SOME PATHOGENIC BACTERIA IN ANIMALS.

A number of pathogenic bacteria have been very thoroughly investigated in animals, and of some of these a brief account may be given.

1. **Bacillus of septicæmia in mice** was described by Koch in his very important work on "Traumatic Infective Diseases." When one, or at most two drops of putrid fluid are inoculated in a house mouse, the animal, in a certain proportion of cases (about a third) becomes ill in about twenty-four hours, and dies in two or three days, with symptoms of an acute specific fever. The blood is then found loaded with small bacilli (shown in Fig. 101, p. 288), many of which are in the leucocytes of the blood. The bacilli are very small, and they are stained by the ordinary agents. They have been cultivated on nutrient media, and animals have been infected by inoculation. House mice and white mice, pigeons, sparrows, and rabbits are susceptible, but it is peculiar that field mice, although so like house mice, are not. Fowls and guinea-pigs are also insusceptible. In rabbits the result is not septicæmia, but an inflammation of the subcutaneous tissue, like erysipelas.

2. **Bacillus of fowl cholera.**—This form is found in fowls in a disease which presents some features in common with human cholera. The bacilli consist of short, thick rods, which stain chiefly at the ends, so that they may be mistaken for micrococci. They have been cultivated on nutrient gelatine and other media, and animals have



been infected. Besides poultry, geese, pigeons, sparrows, mice, and rabbits are susceptible both to inoculation and to ingestion with the food. The result is a disease characterized by weakness and torpor, which deepens into death. At the acme of the disease a fluid evacuation occurs from the bowels, which contains numerous bacilli. Unlike the human cholera bacillus, this one does not confine itself to the intestine, but extends to the blood and organs generally.

Pasteur has extracted from cultures of this bacillus a substance which is highly narcotic, inducing in fowls a condition of somnolence or coma like that of the disease. Pasteur has also attenuated the virus by cultivating it for some time with exposure to the oxygen of the air, at an elevated temperature.

Allied to this bacillus, if not identical with it, is the *Bacillus of rabbit septicæmia*.

3. *Bacillus of swine plague or Swine typhoid* has been described by Klein. The bacilli are like those of fowl cholera, and the cultures also resemble them. But the bacilli have a very different relation to animals. Fowls, pigeons, and guinea-pigs are not affected, while pigs, mice, and rabbits are. The inoculation is very fatal to pigs, which die in one or two days. The bacilli penetrate to the blood and all the organs.

4. *Bacillus of swine erysipelas (Rothlauf, Roget du porc)* has been investigated by Loeffler and Schutz. It is a very small bacillus, like that of the septicæmia of mice. It has been cultivated on nutrient media and thence communicated to animals. Inoculation is successful in pigs, rabbits, pigeons, house and white mice, while fowls and guinea-pigs are insusceptible. Feeding with the cultures did not succeed, even in pigs, although the disease seems to be spread by infection of the food. The disease, whether occurring naturally or produced by inoculation, is characterized by a red eruption of the skin, which is not painful or swollen. There are also signs of progressive weakness, and death follows in one or two days. The bacilli are found in all parts of the body, especially the lungs and spleen, but are not abundant in the blood.

Pasteur produced an attenuation of the activity of this bacillus, and by inoculation induced a mild form of the disease, which caused an immunity from the unweakened form. In order to this two vaccinations were made, first with a very weak form (*premier vaccin*), and then twelve days after with a much stronger (*deuxième vaccin*).

#### IV.—DISPOSAL OF BACTERIA. IMMUNITY. PHAGOCYTES.

In what has gone before we have seen that while there are many microbes which are entirely saprophytic, the truly pathogenic ones present great varieties in their relations to different kinds of animals. It may be said that all animals present an immunity from the pure saprophytes, and that animals present different degrees of susceptibility and immunity from the pathogenic forms. In this regard we may distinguish a natural and an acquired immunity.

**Natural immunity** is, to a large extent, matter of inheritance. Several illustrations of this have been furnished in the section on Causation of Disease (see pp. 21, 22). It may be stated here that not only the species of an animal, but the race and the family characters determine the degree of susceptibility to the pathogenic forms. It has already been pointed out that differences of race and amongst individuals depend on



peculiarities in the details of structure, some of them in appearance very insignificant, and we may associate the varying degrees of susceptibility with such fine differences in structure and function.

**Acquired immunity** is induced in many persons by the occurrence of an attack of the disease itself. Thus a single attack of small-pox generally renders a person insusceptible to a further attack, and the same applies to most of the specific fevers, although not to all.

**Vaccination** is an instance of immunity produced by the induction of a modified form of the disease. The virus of vaccinia is equivalent to an attenuated form of the virus of small-pox.

Pasteur has endeavoured to obtain material for similar protective inoculation, or **vaccines**, of various forms of pathogenic bacteria, and, as indicated above, he has succeeded in doing so in the case of anthrax, fowl cholera, and swine erysipelas. In the case of anthrax, this is done by cultivating the bacillus at a higher temperature than usual, and in the other two cases by prolonging the culture.

Pasteur has applied another method to **Hydrophobia**. Although the bacterium of hydrophobia has not been discovered, the morbid poison has been shown to be present in the central nervous system of animals affected by this disease. When a rabbit has become affected with hydrophobia by inoculation, then the spinal cord may be used for producing the disease in another animal by inoculation. Pasteur has found that by drying the cord the intensity of the virus may be diminished, so that, for instance, after drying for fourteen days, the inoculation will produce no effect. After inoculation with a virus thus attenuated, others may be used which have been treated for a shorter time, and thus, by a series of inoculations, we reach the fresh cord. After this preliminary series of inoculations the fresh cord, which otherwise is extremely virulent, produces no effect, and the animal is absolutely protected from the disease, when acquired in the usual way by the bite of a rabid animal.

This method of protective treatment has been applied in man, and Pasteur asserts that it may be begun after the person has been bitten, and yet prevent the development of the disease. The observations of this observer have been confirmed so far as animals are concerned by an English commission, but there seems still to be some doubt as regards the protective treatment.

An interesting fact in regard to immunity is that, to a certain extent, an **acquired immunity** may be **transmitted by inheritance**. This at least seems to be one explanation of the fact that in countries where specific fevers are prevalent the disease is milder than where it is introduced for the first time. The case of the extraordinary fatality of



measles in the Fiji islands may be cited in this connection. There is also an interesting series of facts recorded by Grawitz in regard to small-pox. A troupe of Esquimaux visited Berlin, and some were vaccinated with ordinary vaccine lymph, the result being that three died of an acute fever. Shortly afterwards the remainder were vaccinated in Paris, and they all died of what the Parisian physicians called severe small-pox. Of course in such cases there may be racial differences of susceptibility, which complicate the problem.

**Causes of immunity.**—From what has gone before in regard especially to natural immunity, it will appear that the living tissues of certain races and persons are more able to resist the attacks of pathogenic bacteria than those of others. This apparently depends on fine differences in the reaction of the tissues to the morbid agent.

In regard to acquired immunity the problem is much more difficult. It seems strange that having once succumbed to a certain form of microbe, recovery should establish a kind of protection against further attacks. Pasteur has suggested that, for the growth of each form, a certain chemical substance is necessary in the blood or tissues, and that as this is exhausted by the first attack it takes some time before, by its restoration, the parasite is supplied with a proper pabulum. But there are obvious objections to this. For one thing, the bacteria may be cultivated out of the body in the very blood in which they would not grow inside the body. Such an explanation breaks down entirely when we deal with natural immunity. We have cited, for example, the curious fact that while ordinary sheep are readily inoculated with anthrax, the Algerian sheep require a larger dose. There must be something very different here from a mere absence of a chemical principle, as, when the disease is established, the bacteria find sufficient pabulum. There must be here, as in similar cases, a special power of resistance in the living tissues, so that the bacteria are nullified unless they attack with greater force than usual.

This subject has been greatly elucidated by the very suggestive observations of Metschnikoff on what he calls the **Doctrine of the phagocytes**. This doctrine has received considerable support from the labours of this author and of others, but, although pointing in the direction of a true explanation of the facts under consideration, it cannot be regarded as fully established, especially in some of its details.

Metschnikoff's first observations were made by the introduction of fungus spores into the transparent bodies of a form of daphne. He found that when small quantities of spores were used, the amœboid leucocytes collecting around them, took them into their substance, destroyed their power of germination, and finally disintegrated them,



while, if many spores were introduced, some of them which were not taken up by the leucocytes, germinated and grew through the body.

In the further development of this doctrine Metschnikoff recognizes a difference between the leucocytes and the fixed cells of the tissues. Both may become amoeboid and both have the power of taking up solid particles and digesting them. In view of this power he names them **Phagocytes** (*φαγεῖν* = to devour). These two forms of cells present different reactions to the various bacteria, and for the sake of distinction the leucocytes which have divided nuclei are called **Microphags**, and the cells derived from the fixed cells of the tissues, which are larger and have large oval nuclei, are **Macrophags**. In all diseases of this class there is liable to be a struggle between the bacteria and the phagocytes, whether the microphags or the macrophags.

As an illustration of this we may cite, in the first place, Metschnikoff's observations on **Erysipelas**. As we have seen, in this disease the streptococcus erysipelatis gets into the lymphatics and multiplies there. It does so in spite of the presence of macrophags, which generally show little power of attacking micrococci. Then ensues the inflammation accompanied by a great invasion of leucocytes which, as microphags, eat up the micrococci. The latter are found inside the cells in various stages of degeneration; they are converted into irregular granules and digested. The leucocytes are not only capable of devouring bacteria, but any animal debris may be digested by them. Hence they play an important part in the resolution of inflammations.

Hess has also experimented with the staphylococcus pyogenes aureus. He introduced cultures of it into the cornea of rabbits, and found that it produces an acute inflammation, which by and by subsides and healing occurs. In this case the leucocytes take up the micrococci and make them disappear.

This same author has made some observations on anthrax which have important bearings on natural immunity. He used frogs which, as we have seen, are nearly insusceptible to anthrax. When the bacilli were injected into the blood of these animals they were rapidly taken up by the leucocytes, so that in a few hours there were none free. The leucocytes conveyed the bacteria to the liver and spleen where they were disposed of within the cells, presenting various stages of degeneration. In other animals it was found that the number of bacilli undergoing disintegration in the cells in the spleen was in inverse proportion to the susceptibility of the animal to anthrax. That is to say, in susceptible animals the leucocytes are unable to devour the bacilli, and the disease progresses. It is only the bacilli in the leucocytes which undergo degenerative changes, those free in the blood grow freely.



Metschnikoff has investigated the relation of the spirilla in relapsing fever to the phagocytes. The leucocytes in the general circulation do not take up the spirilla, and he inferred that when they disappeared from the blood they must be devoured by the macrophags of the splenic pulp. He found, however, that although they are disposed of in the spleen it is the microphags which there attack them.

This author has also made observations in regard to the tubercular bacillus. He has experimented in a small kind of marmot which is little susceptible to tuberculosis. He finds after inoculation that giant-cells here, as in other animals, take up the bacilli. There is then a struggle between the two living cells. In the marmot the giant-cells usually get the better of the bacilli, which undergo degenerative changes till they are converted into yellow clumps (yellow degradation). If the bacilli get the upper hand the giant-cells degenerate, the form of degeneration being caseous necrosis. In susceptible animals, such as the rabbit, this degeneration of the giant-cells is very marked, but the yellow degradation of the bacilli also occurs.

**Literature.**—The recent literature of Bacteriology is of enormous extent, as may be judged from the fact that in Crookshank's book it occupies 59 pages. The works more particularly referred to by the author are given here. *General works*—COHN, *Biol. der Pflanzen*, vol. iii., 1879; ZOPF, *Zur Morphologie der Spaltpilzen*, 1882 and 1885; DE BARY, *Vorlesungen über Bacteria*, 1885, and *Fungi, mycetozaa and bacteria* (transl.), 1887; HUEPPE, *Formen der Bacteria*, 1886; KLEIN, *Micro-organisms and disease*, 3rd. ed., 1886; CROOKSHANK, *Bacteriology*, 2nd ed., 1887; WOODHEAD and HARE, *Pathological Mycology*, 1885; CORNIL et BABES, *Les bactéries*, 1885; FRAENKEL, *Bakterienkunde*, 1887. *Limits of temperature*—PICTET and YOUNG, *Comptes Rendus*, xcvi., p. 747; COLEMAN and M'KENDRICK, *Proc. of Roy. Instit.*, 1885. *Ptomaines*—AITKEN, *Animal alkaloids*, 1887; BROWN, *Animal alkaloids*, 1887; BRUNTON, *On disorders of digestion*, 1886; BRIEGER, *Ueber Ptomaine*, 1885, and *Untersuchungen*, 1885. *Sarcinae*—LOSTORFER, *Wien. Med. Jahrb.*, 1871; FERRIER, *Brit. Med. Jour.*, i., 1872; FALKENHEIM, *Arch. f. experiment. Path. u. Pharmac.*, xix., 1885. *Pathogenic bacteria*—FEHLEISEN, *On erysipelas*, *Syd. Soc. transl.*, 1886; OGSTON, *Brit. Med. Jour.* i., 1881; ROSENBACH, *Microorganismen der Wundinfektionskrankheiten*, 1884; GESSARD, *De la pyocyanie et de son microbe*, 1882; NEISSER, *Deutsch. Med. Wochenschr.*, 1882. BUMM, *Mikroorg. d. Gonorrhoe.*, 1885; BOLLINGER, *On animal poisons*, *Ziemmsen's Encycl.*, vol. iii., 1875; KOCH, *Milzbrandimpfung*, 1882; CHAUVEAU, *Comptes Rendus*, xc. to xcvi.; GAFFKY, *Etiology of enteric fever*, *Syd. Soc. transl.*, 1886; OBERMEIER, *Berlin. Klin. Wochenschr.*, 1873; CARTER, *Lancet*, 1879, and *Spirillum fever as seen in Western India*, 1882; KOCH, *Etiology of cholera*, *Syd. Soc. transl.*, 1886; FRIEDLAENDER, *Fortschr. d. Med.*, ii., 1884; FRAENKEL, *Deutsch. Med. Woch.*, 1886; LOEFFLER (*Diphtheria*), *Syd. Soc. transl.*, 1886; KOCH, *Etiol. of tuberc.*, *Syd. Soc. transl.*, 1886; NEISSER, *Etiol. of leprosy*, *Syd. Soc. transl.*, 1886; LUSTGARTEN, *Med. Jahrbücher*, 1885; LOEFFLER and SCHÜTZ (*Glanders*), *Syd. Soc. transl.*, 1886; EDINGTON (*Scarlet fever*), *Brit. Med. Jour.*, 1887, i. 1262 and ii. 830; FRISCH (*Rhinoscleroma*), *Wien. Med. Wochenschr.*, 1882; DIECKERHOFF und GRAWITZ (*Acne contagiosa*),



Virch. Arch., 1885, cii.; KOCH, Traum. Inf. Disease, Syd. Soc. transl., 1880. *Immunity*—PASTEUR, numerous papers in Comptes Rendus; Report of Commission on Hydrophobia, Brit. Med. Jour., ii., 1887, 12; METSCHNIKOFF, Virch. Arch., ciii., cix., cxiii.; HESS, Virch. Arch., cix., cx.

## B.—PARASITIC FUNGI.

The fungi proper consist of cells devoid of chlorophyll and occurring in the form of threads (*Hyphæ*) and round or oval spores (*Conidia*). The fungi are divisible into two classes, the moulds or filamentous fungi and the yeasts or sprouting fungi. In the filamentous fungi the threads or hyphæ form an interwoven network which constitutes the root part of the fungus and is called the *Mycelium*. From this root part upright stems arise, under favourable circumstances, which bear the fructification. The moulds are called, from the presence of this root part composed of hyphæ, **Hyphomycetæ**. The sprouting fungi consist merely of cells, which multiply by budding, and are not known to produce a mycelium, although some of them are supposed to bear spores. It should be added that this distinction only expresses a mode of growth, and that some of the filamentous fungi, under certain circumstances, grow like yeasts, while the sprouting fungi are believed by some to be only a particular phase of fungi which may have in other phases the characters of the filamentous forms. In Fig.

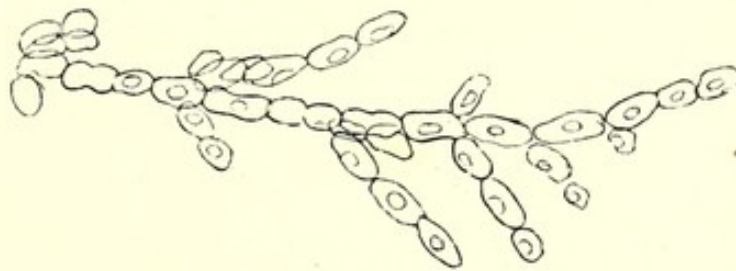


Fig. 109.—Formation of conidia in submerged portion of penicillium.

109, for example, we have a portion of an ordinary filamentous fungus (penicillium) which has been growing submerged in a nutrient material. It shows a mode of growth approaching to that of the sprouting fungi.

## I.—THE SPROUTING FUNGI OR YEASTS.

These consist of oval cells which contain granules and frequently vacuoles. They grow by the formation of buds from the cells, which increase in size and then separate from their parents.

Great importance attaches to the sprouting fungi from the part they play in fermentations, as they present in this respect many analogies with the bacteria. The proper yeast fungus has the faculty, in its growth, of splitting up sugar into alcohol and carbonic acid.

The sprouting fungi are essentially saprophytic, very few of them even occurring as occasional parasites, unless we include the fungus of



thrush as belonging to this class. The proper yeasts are included in the genus *saccharomyces* of which several species are distinguished.

*Saccharomyces cerevisiæ* or *Torula cerevisiæ* is the common yeast plant. *Saccharomyces ellipsoideus* is the fungus concerned in the fermentation of wine, and it is found in all sorts of fermenting juices of fruits. *Saccharomyces mycoderma* is found in the scum which forms on fermenting beer. It used to be supposed that it had to do with the development of acetic acid from alcohol and hence was named the vinegar fungus, but this action is probably produced, not by it, but by a bacterium.

## II.—FILAMENTOUS FUNGI OR MOULDS.

The root part or mycelium of these fungi consists, as we have seen, of hyphæ or threads, from which



Fig. 110.—Aerial growth of penicillium.

may grow up the stems which produce the spores. The common moulds belong, for the most part, to three genera, penicillium, aspergillus, and mucor. The fructification of penicillium is characterized by the formation, on the summit of the stem, of rows of spores (see

Fig. 110), which, from their brush-like appearance, suggest a pencil. The fructification of the aspergillus consists of spherical heads.

The filamentous fungi are mostly pure saprophytes, and some of them, while usually saprophytic, become occasionally parasitic. On the other hand, there are some which are only known as parasites, although they may have also a saprophytic existence. These are confined to the skin and its appendages.

1. **Saprophytes and occasional parasites.**—In the common moulds there are three genera which are frequently represented, namely, *Penicillium*, *Aspergillus*, and *Mucor*, and of these the aspergillus is the only one which is of importance as an occasional parasite.

*Aspergillus glaucus* is one of the common greyish blue moulds. According to De Bary it is not a true aspergillus, but belongs to the genus *Eurotium*. It has been described as occurring as a parasite, but this has probably been a mistake for one of the true aspergilli. It does not grow at a temperature approaching that of the body.

*Aspergillus fumigatus* forms a greenish mould with a granular surface, becoming grey later on. It grows best at a high temperature, and its spores may, after introduction into the body of animals, grow there (see below).



*Aspergillus flavus* is a yellow or yellowish green mould, while *Aspergillus niger* has a dark brown colour. These also grow at the temperature of the body, and may become parasitic.

The *Aspergilli* have been introduced by experiment into the bodies of animals, and the spores of some of them, but especially *A. fumigatus* and *flavus*, are able to live and germinate there. They do not, however, like the bacteria, multiply in the body, but they may by their growth produce destructive effects. The spores carried by the blood settle in certain organs, and may produce lesions not unlike metastatic abscesses. In a paper by Grawitz, for example, there is an illustration showing a condition of the kidney closely resembling that occurring in

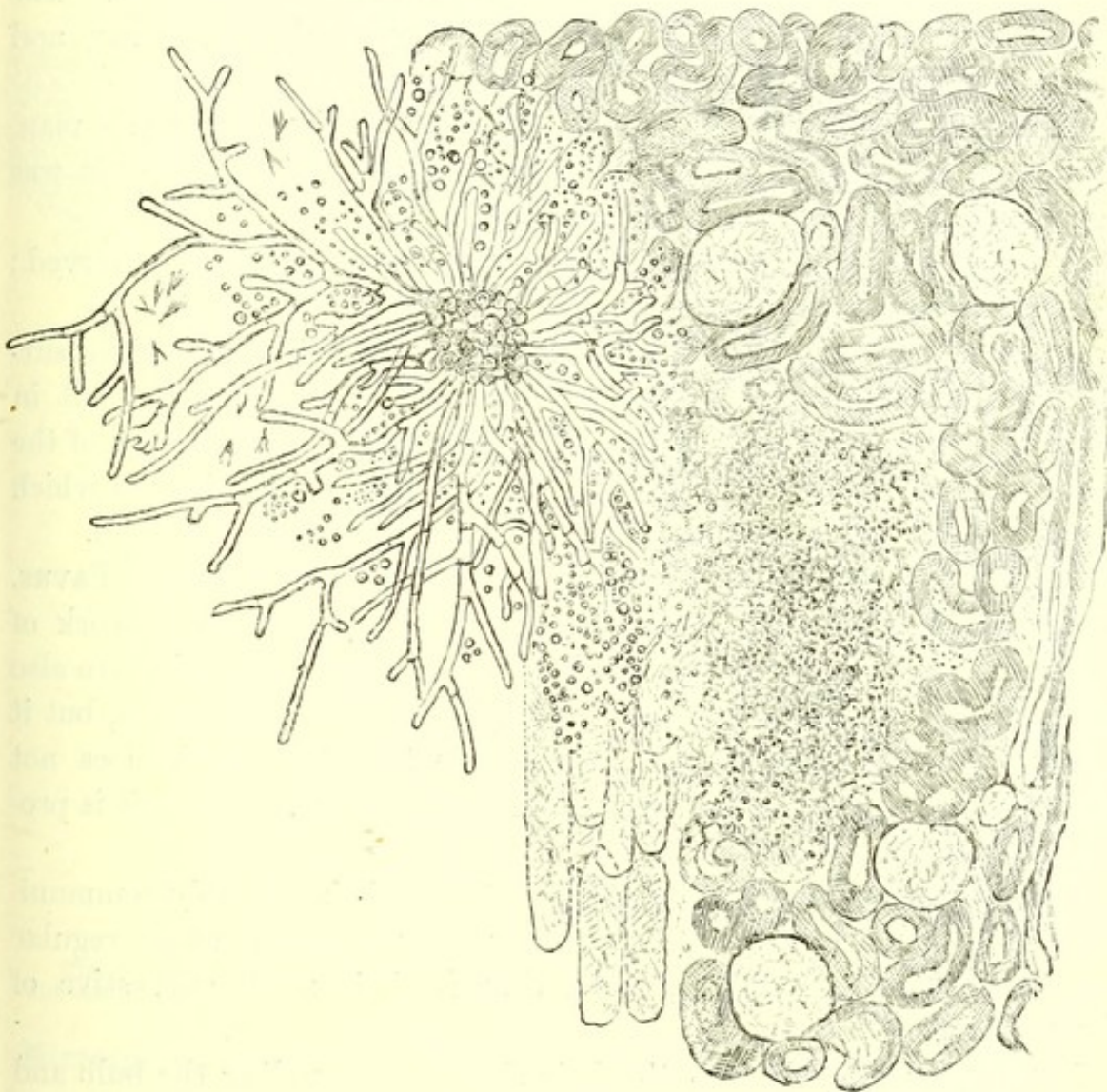


Fig. 111.—Fungus growth in kidney. To the left of the figure the kidney tissue has been cleared up by adding solution of soda so as to bring out the fungus (*aspergillus*). Fat drops and crystals of tyrosin are present, but no spores. (GRAWITZ.)

pyæmia. In each of these lesions a growing fungus is found (see Fig. 111), which is causing necrosis and inflammation of the tissue, but is not producing spores.



Grawitz believed that ordinary mould fungi may, by cultivation, be so altered as to possess pathogenic properties such as those just mentioned, but subsequent observation shows that it is only the aspergilli which have these powers, and that they have them without any special cultivation.

No similar extension of the aspergilli to internal parts has been observed in man.

**Aspergilli** are occasionally found growing in the ear, constituting an **Otomycosis**. They occur by preference on the tympanic membrane and inner third of the external meatus, but when the drum is perforated they may extend to the middle ear. The fungus does not apparently grow unless there be some previous disease causing a breach in the epithelium, and they do not penetrate deeply into the structures. The forms observed have been chiefly *aspergillus fumigatus*, *flavus*, and *niger*, but other kinds have been found.

Leber has also observed a fungus growing on the cornea of a man. It was an *aspergillus*, and he determined by experiment that it was capable of growing on the rabbit's cornea.

A mycosis of the **Nasal mucous membrane** has also been observed; the fungus was *aspergillus fumigatus*.

2. **Pathogenic fungi**.—The true pathogenic fungi occur on the surface of the body, usually attacking the epidermic structures, but in some cases penetrating more deeply. They mostly cause lesions of the hairs and epidermis, and some of them lead to inflammations in which the true skin, and, it may be, the deeper structures are concerned.

**Achorion Schœnleinii** is the fungus of the well-known disease, **Favus**. It consists almost entirely of hyphæ, which form a dense network of threads. Conidia spores of an oval shape and highly refractive are also produced. The fungus has been cultivated on nutrient jellies, but it grows best on blood serum. At ordinary temperatures it does not grow at all, requiring a temperature of about 84° F. Hence it is probably an obligate parasite.

**Favus** is very common in mice and cats, and it is probably communicated from them to man. It is peculiar that the fungus whose regular seat is probably the mouse, has a characteristic smell suggestive of these animals.

The fungus extends into the hair sheath surrounding the bulb and penetrating into the shaft. It also penetrates to the deeper layers of the epidermis and even to the true skin.

**Trichophyton tonsurans** is the fungus of **Ringworm**, and of the corresponding diseases of the body and beard, namely, *tinea circinata* and *tinea sycosis*. It consists of hyphæ and conidia spores, but there is no proper fructification. It has been cultivated on nutrient media, and



grows best on blood serum at a temperature of 84° F. The cultures have, by inoculation, produced the typical lesions of *tinea circinata*.

The fungus grows in the epidermis and penetrates into the hair shaft (see Fig. 112). It also irritates the true skin, producing inflammatory lesions.

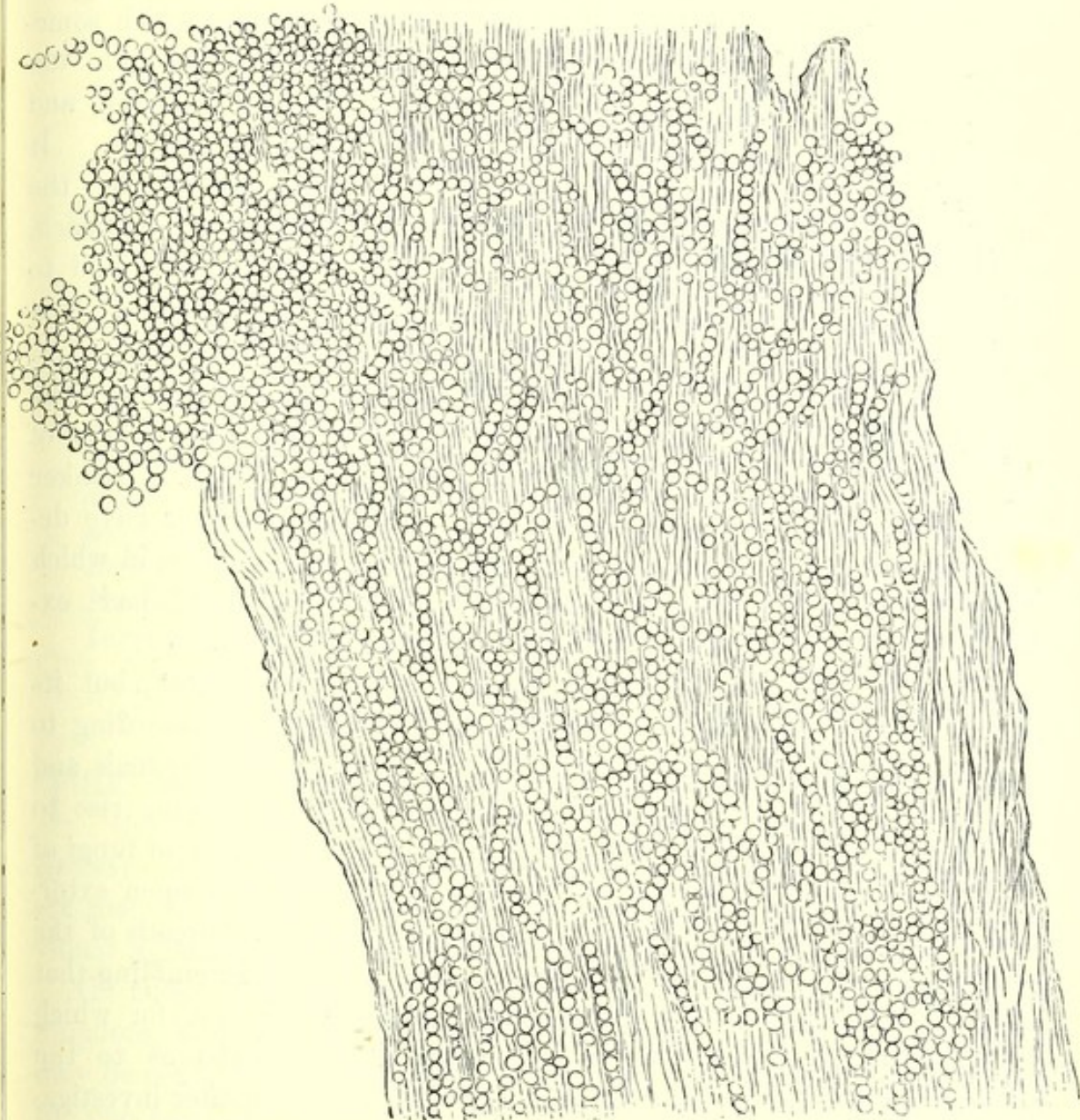


Fig. 112.—Hair from a case of ringworm, showing fungus penetrating and breaking up the shaft.  $\times 350$ .

***Microsporon furfur*** is the fungus of ***Pityriasis versicolor***. It consists of ramified hyphae and spores, which occur in groups almost like the fructification of *penicillium*.

The fungus penetrates very little into the epidermic layers and produces merely a desquamation without inflammation.

***Oidium albicans*** (also called ***Saccharomyces albicans***) is the fungus of **Thrush**. It is by some regarded as belonging to the sprouting fungi,



although it undoubtedly produces long threads or hyphæ. It is in the form of branching threads, and of conidia, which lie in groups. It has been cultivated, and is said to have the power of fermenting sugar.

The fungus grows amongst the epithelium of the mouth, forming, with the epithelium, a soft whitish membrane (see Fig. 113). This is

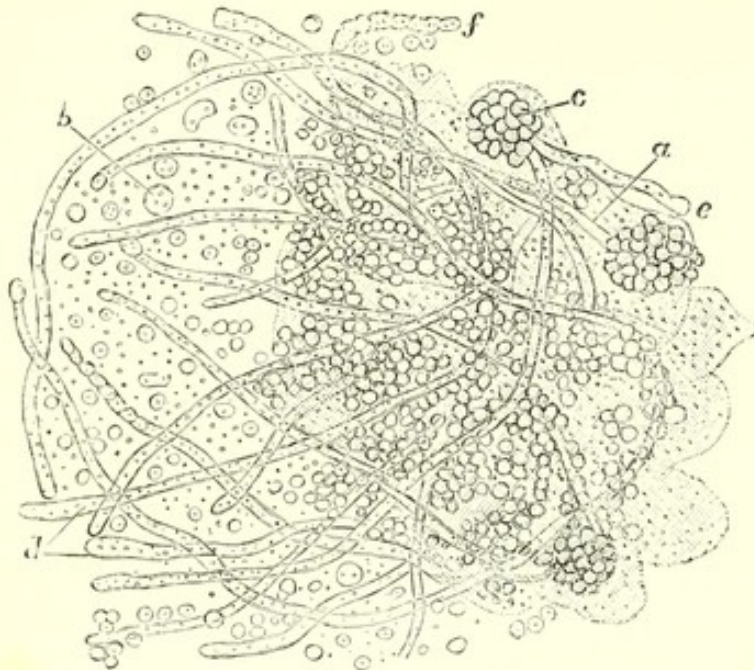


Fig. 113.—*Oidium albicans*, from the mouth in a case of thrush. (KUCHENMEISTER.)

the so-called aphthous condition which sometimes occurs in unhealthy children and even in adults. It may extend to the œsophagus, stomach, and intestine, and to the air passages. It is said to be capable of inoculation on the mucous membrane of the vagina. Zenker and Wagner have described cases in which the fungus had ex-

tended to the blood and caused embolism in the brain.

**Chionyphe Carteri** is a fungus found in the madura foot, but its exact botanical position is not determined. The fungus, according to Carter, penetrates into the substance of the foot, producing canals and cavities among the soft structures and the bones, and giving rise to severe inflammations. In the cavities and canals are masses of fungi of a blackish colour and forming frequent balls. The canals open externally and discharge a foetid pus containing spores and threads of the fungus. There is great thickening and an appearance resembling that of scrofulous disease, and the condition is a chronic one for which amputation must frequently be performed. Doubts exist as to the actual connection of the fungus with the disease, and further investigation is required (see especially Lewis).

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## SECTION XI.

## ANIMAL PARASITES.

- I. ENTOTZOA OR INTERNAL PARASITES, *their general characters and effects.* 1. *Protozoa.* 2. *Trematoda or Flukes, chiefly Distoma hepaticum, sinense, and hæmatobium.* 3. *Cestoda or tape-worms. Taenia solium, structure and development; Cysticercus cellulosæ, its scolex form. Taenia mediocanellata. Taenia echinococcus, forming hydatids; its cysts, brood-capsules, heads, and chitinous membrane. Bothriocephalus latus. Other rare tape-worms.* 4. *Nematoda or Round-worms. Trichina spiralis, its embryonic and adult forms; effects of migrations. Ascaris lumbricoides. Oxyuris vermicularis. Trichocephalus dispar. Dochmius duodenalis. Filaria medinensis. Filaria sanguinis, its periodicity in the blood; relation to chylous urine, and lymph-scrotum.* II. EPITZOA OR EXTERNAL PARASITES. 1. *Arachnidæ, chiefly Acarus scabiei and Pentastomum denticulatum.* 2. *Insecta, chiefly Pediculi and Pulex irritans. Larvæ of insects in wounds, skin, and bowels.*

## I.—ENTOTZOA OR INTERNAL PARASITES.

THE Animal Parasites are divisible into two groups according as they live inside the body (the *Entozoa*), or on the surface (the *Epizoa*). The Entozoa are nearly all included among the worms, and are met with in the alimentary canal and in the tissues. The individual by whom the parasite is accommodated is called its **Host**.

All these parasites belong to the lower classes of the animal kingdom, and they mostly exist in two different active states, which may be compared to the larval and free states of insects. There is on the one hand the sexually mature animal, and on the other hand the embryo. The former has usually a more or less fixed habitation, but the latter generally shows considerable powers of migration, often inside the tissues of living animals. When the mature parasite produces its eggs these nearly always pass away from the locality in which the parent is, and are frequently discharged from the animal in which the parasite lives. The embryo then undergoes a period of migration, which may be partly in the external world, but is chiefly in the living body of a higher animal. This stage of migration is mostly succeeded by a period of rest, in which the embryo becomes



quiescent, and this is usually in the tissues of the animal in which it has been wandering. In this quiescent state it becomes surrounded by a cyst or capsule. The parasite may awaken from this state of rest if it or its host be eaten by another animal. The embryo, having thus been carried into another animal, may undergo development into the sexually mature adult, which in the majority of cases has its seat in the alimentary canal.

It will be seen that the entozoa have two distinct phases of development, in both of which they are parasitic, although the ova and embryos may for a time be free. The embryonic or immature form frequently differs greatly in its size and form, and nearly always in its locality, from the mature, and in many cases it has been a matter of great research to trace the connection. The adult and embryonic periods are rarely passed in the same individual host, and it is not uncommon to find that the adult inhabits one species of animal and the mature worm another.

The effects of parasites vary greatly. All of them live at the expense of the host, and may do harm by consuming its available material. Most adult parasites live in the intestinal canal, and use the juices there for nourishment. But this is generally of slight importance. The embryonic forms usually produce much more considerable disturbances. They do so in part by their migrations. In moving amongst the tissues they may disturb the living structures, and they will leave canals or tracts, which become occupied by new-formed connective tissue. In important organs like the brain this may lead to serious damage. Then, when they come to rest, they may grow to considerable dimensions, forming cysts around themselves, and these by their mere bulk may affect the tissues.

Animal parasites in general are much more common in the lower animals than in man. It is interesting to observe that in nearly all cases each parasite has its own particular species of animals for its host, and this applies both to the mature and embryonic stages. It may be said, indeed, that each species of animal has its own special representative parasites, and when a parasite of one species of animal strays into another then it usually dies very soon. As a general rule, also, the adult and embryonic forms occur in different species of animals, so that the mature worm is attached to one species and the embryo to another. The number of parasites, if we take in the whole animal kingdom, is thus enormous, and their relationships very complicated.

1. **Protozoa.**—These lowest animals are very minute in size, and unless they be present in large numbers can scarcely be detected.

The **Amœba**, which belongs to the class of Rhizopoda, is a naked,



almost structureless contractile body. They have been observed in the intestine and fæces, and were supposed to have caused dysenteric inflammation (see Leuckart).

The **Gregarine**, a form of the Sporozoa, has also little structure, but is covered by a cuticle. It is parasitic so far as is known only in invertebrates.

**Coccidia** also belong to the Sporozoa, and are more interesting, as they cause definite disease in animals and man. They consist of oval bodies with capsules. They are very common in the liver of the rabbit, where they form whitish nodules, which grow to the size of a hazel nut, and may lead by anæmia, etc., to the death of the animal. On section of the nodules a cheesy or purulent mass exudes, in which, besides debris, there are innumerable oval bodies. In a few cases the liver in man has been the seat of this parasite. In a case by Gubler there were in the liver about twenty tumours of cancerous appearance, and most of them the size of a chestnut, but some as big as an egg. The death of the patient was from peritonitis.

**Meischer's or Rainey's tubes** are elongated granular bodies found in the muscular substance of animals. They are supposed to be parasitic protozoa, but their nature is very obscure. Although found in many different animals, sometimes in large numbers, they have not been observed in man.

2. **Trematoda**.—We have here an order of flat-worms of a more or less oval shape, and many of them somewhat in the form of a leaf. They possess, on the ventral surface, one or more sucking discs by which they attach themselves. They have only one opening of the alimentary canal which is generally forked. These worms are commonly called **Flukes** from the resemblance in shape of the commonest of them to the fish of that name. The various forms inhabit the bile ducts, except the *Distoma hæmatobium*, which is found in the veins of the portal system.

***Distoma hepaticum***.—This is the commonest worm of this order. As the name implies, it is met with in the liver, where it inhabits the bile ducts. The liver fluke is generally about an inch in length (Fig. 114) and rather more than half an inch in greatest breadth. The body is very flat, and anteriorly it ends in an elongated process, forming a kind of head. This head bears the mouth, and a short distance behind it comes the sucking disc. Between these lies the opening of the sexual apparatus, both male and female organs existing in each individual. The uterus forms a convoluted tube behind the sexual opening, and the seminal tubules lie still further back. This parasite is very common in certain of the lower animals, especially sheep. It occurs in enormous numbers in the bile ducts, which are dilated by it. As many as 1,000 have been obtained from a single sheep. It produces in sheep the disease called commonly the rot, which



in some years is very fatal. It is said that in 1830-31 between one and two million sheep perished from it. It occurs also in oxen, where it produces more considerable alterations of the ducts. These become greatly dilated, thickened by inflammation, and incrustated with lime. It sometimes happens that masses of inspissated bile and lime salts form in the liver where the flukes are present. It is probable that in these cases many of the parasites have died and become themselves the seat of incrustation. This fluke has also been met with in horses and asses, and in some rare cases in man. In man it has not been observed in large numbers, but it may produce serious obstruction of the bile ducts.



Fig. 114.—*Distoma hepaticum*. (LEUCKART.)

The eggs of this parasite are small oval bodies, which, in water, develop into embryos which swim about by the aid of cilia. Their further development is unknown, and also the form in which they pass into the animal which becomes the host of the adult.

***Distoma sinense*.**—This parasite has been met with in the East, and described by M'Connell in the *Lancet* in 1875, and independently by Macgregor in the *Glasgow Medical Journal* in 1877. Although a much smaller worm it is of a more elongated shape than the *distoma hepaticum*, as will be seen from Fig. 115. It is rather



Fig. 115.—*Distoma sinense*, a, œsophagus and stomach tubes; b, uterus; c, yolk glands; d, ovary; e, testes; f, termination of water vascular system.  $\times 5$ . (After MACGREGOR.)

more than half an inch in length, and about an eighth of an inch in greatest breadth. When seen in the bile in the fresh state, the edges show a beautiful delicate green colour, tinged with yellow, while the centre is of a deep brown. In the accompanying figure the position and appearances of the various organs are indicated. The eggs are very small, and each animal possesses thousands.

This parasite has only hitherto been observed in man, and is found in very large numbers—not less than 500, probably more—in the ducts of the liver and in the gall bladder. In Dr. Macgregor's cases the persons affected with the parasites were Chinese, and they were the victims of a peculiar paralysis of the legs and arms. Dr. M'Connell's case was also in a Chinaman.

***Distoma lanceolatum*.**—This form of fluke is less than three-eighths of an inch in length, and about the fifteenth of an inch in breadth. It is seldom seen in man, and occurs in comparatively small numbers in sheep and cattle, producing little disturbance.

***Distoma hæmatobium*.**—This parasite, also called the *Bilharzia hæmatobia*, has the male and female organs in different individuals. The male is about half an inch in length, and flat, but rolled up at the edges, especially



behind, so as to form a kind of gutter in which the female lodges. The female is about a half longer, but filiform. The eggs are small and furnished with a spine at the end or at the side.

The parasite inhabits the blood-vessels of its host, chiefly the portal vein, the splenic and mesenteric veins, and those of the rectum and bladder. The penetration of the eggs in large numbers into the mucous membrane of the rectum and urinary bladder produces great irritation, and frequently hæmorrhage. Similar irritation may be produced in the pelvis of the kidney and ureters. This parasite is met with almost solely in Egypt and Abyssinia, and it is said that in Egypt about half the natives are victims of it.

**3. Cestoda or Tape-worms.**—These are in the mature state long flat worms, without mouth or alimentary canal. Anteriorly there is a head furnished with some apparatus for attaching itself to the host. Behind the head and neck the worm forms a series of segments called **Proglottides**, each of which develops a bi-sexual apparatus, and is, so far, a complete individual. The adult worm or **Strobilus** is therefore a colony of individuals. The worm inhabits the alimentary canal and apparently occurs only in vertebrate animals. Besides this adult form there is an intermediate immature form, called the **Scolex**, which occurs in the tissues of animals. The scolex has a head like that of the mature worm, and generally possesses a sac or cyst into which it can retire.

There are representatives of two families of this parasite met with in man, namely, *Tænia* and *Bothriocephalus*.

***Tænia solium*.**—This form is of very common occurrence in this country. The strobilus or mature worm occurs in the alimentary canal, and the head is usually situated in the duodenum or upper part of the jejunum, while the rest of the animal extends downwards in the canal, attaining on an average a length of from ten to twelve feet. As already mentioned, this, like other tape-worms, has no alimentary canal, and supports itself by imbibition of nutritious material from the intestine.

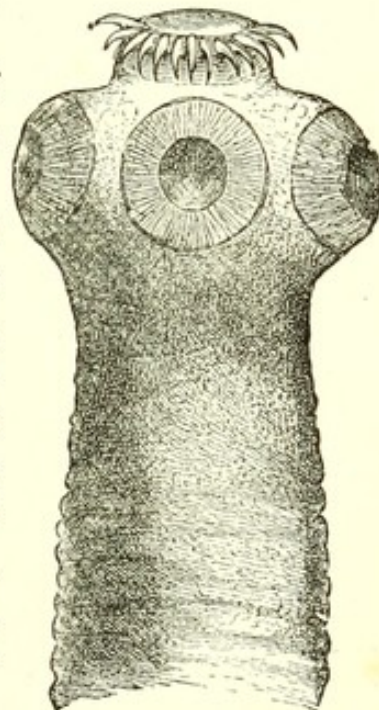


Fig. 116.—Head of *T. solium*.  
× 45. (LEUCKART.)

The head of the worm, which is represented in Fig. 116, is about the size of a pin's head, and of a generally rounded form. In front it is prolonged so as to form a proboscis or rostellum, which is surrounded by a circle of twenty-six hooklets. The wide part of the head has four large sucking



discs. On the head follows a narrow neck, which is so thin that it readily breaks when the worm is handled, rendering it difficult to



Fig. 117.—Two ripe proglottides of *T. solium*, with branches of uterus shown.  $\times 2$ . (LEUCKART.)

obtain the small head. The proper neck is about half an inch in length, and it gradually merges in the anterior part of the body in which fine transverse lines begin to appear as the first indications of the formation of segments. On passing down, the worm increases in breadth, while the segments elongate and become more completely divided. At first the segments or proglottides are homogeneous in appearance, but by and by the sexual apparatus begins to appear. The total number of segments in a worm ten feet in length is about 800. The sexual apparatus begins to appear about the 200th segment from the front, and is mature about the 450th; it consists of the male and female organs which are present in each segment. In the fully matured segment the ova are visible, and when a proglottis is dried on a glass slide they indicate the form of the uterus, which in this tape-worm consists of a central stem and ramifying lateral branches to the number of seven to ten (see Fig. 117).

The male organs consist of a large number of vesicles scattered throughout the segment, as shown in Fig. 118, but more abundant anteriorly, as the female organs

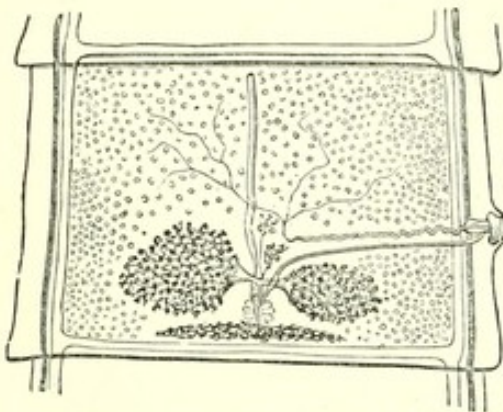


Fig. 118.—Unripe proglottis of *T. solium* showing sexual organs. The small vesicles scattered throughout are the male organs. The other structures shown are seminal tubes, vagina, globular body, yolk body, ovaries, and unbranched uterus.  $\times 10$ . (LEUCKART.)

occupy the space behind. The vesicles are connected with fine seminal tubules which are difficult to make out, and are shown in the figure as fine branching lines. These end in a slightly convoluted tube, the vas deferens, which is generally very distinct, and this passes across the segment to the papilla, a slight projection at the side of the segment into which the male and female sexual organs open. At the papilla the vas deferens ends in a projectile penis which is capable of passing into the extremity of the female organ, the first part of which is called the vagina.

The vagina forms a canal which passes transversely across the segment towards the middle line and tends also backwards, to end in a somewhat globular dilatation, sometimes called the **Globular body**. The connections of this body are difficult to make out, but they may be stated as follows, and understood by the annexed Figs. 118 and 119. In the posterior part of the segment as shown in Fig. 118 are seen on either side the comparatively large ovaries, forming tree-like expansions, consisting of a congeries of closed tubes. The ovaries have ducts which pass into the



globular body. Behind the ovaries and the globular body is the yolk gland which is of a somewhat pyramidal shape and spread out laterally. This also communicates with the globular body in front of it. Besides these communications the globular body, which is thus the central part of the female organs, communicates with the uterus in front. At this period of development the uterus consists of a simple tube extending longitudinally in the middle of the segment (see Fig. 118). It will

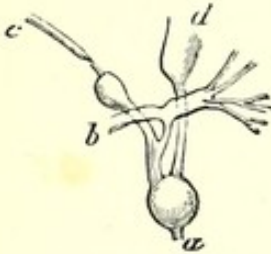


Fig. 119.—The globular body, or Mellis's body and its connections. See text.  $\times 30$ . (LEUCKART.)

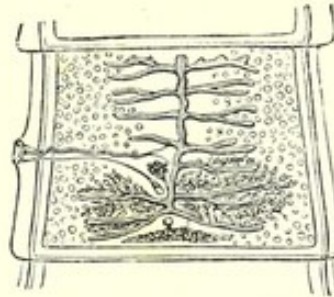


Fig. 120.—Proglottis of *T. solium*, showing branching of uterus.  $\times 5$ .

thus be observed (Fig. 119) that the globular body has communication with four distinct structures, *a* with the yolk-sac, *b* with the ovaries, *c* with the vagina, and *d* with the uterus. The eggs pass from the ovaries first into the globular body, where they receive a covering of yolk, are fertilized, and undergo the beginning of their development. Then they pass into the uterus, which they fill up. As the ova accumulate in the uterus, this begins to throw out lateral branches to the number of seven to ten (see Fig. 120). The lateral branches often show considerable ramifications, in this respect and in their number contrasting with those of the next tape-worm. In the fully mature proglottis only the uterus crowded with ova is visible, the remaining organs having disappeared (see Fig. 117). The prominent ova often make the position and shape of the uterus very distinct, especially if the proglottis be spread out on a glass slide and allowed to dry.

Besides the sexual organs the proglottides possess **muscular fibres**, and a **water-vascular system**. The muscle is non-striated and consists of longitudinal and transverse bundles. The water-vascular or excretory system (shown in Figs. 118 and 120) is in the form of tolerably wide channels, which begin at the head and are continued through the proglottides by two lateral channels right down to the last, where they open outwards. Near the posterior extremity of each proglottis the tubes form transverse communications (see figures). It is possible to inject these tubes from above downwards, but not from below upwards. In addition, the proglottides, as well as the head of the worm, possess numerous round or oval calcareous bodies, which are mainly in the superficial layers of the parenchyma.

As the **proglottides** become mature they sever their connection with the worm and drop off from its lower extremity one by one. They pass down the alimentary canal, and are discharged with the fæces, or else work their way out through the anus by virtue of their contractile power. For a short time after discharge they still show a writhing movement, but they soon come to rest and die. By the decomposition of the proglottis the ova are set free and are ready under suitable circumstances to develop further.



It is mostly in the bodies of swine that the *tænia solium* passes through the next phase of its development, although sometimes it occurs in man.

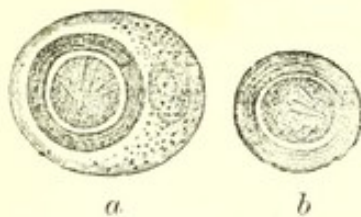


Fig. 121.—Ova of *T. solium*. *a* with yolk, *b* without yolk, as in mature segments. The hard brown shell is indicated. (LEUCKART.)

The Ova (Fig. 121) are surrounded by a dense shell of a brownish colour. Inside the shell the egg develops an embryo which acquires six boring spines. When such ova get into the intestinal canal of the pig, the shell bursts, and the embryo with its spines escapes.

It proceeds to bore its way outwards, and after piercing the alimentary canal, it finds its way to the muscles of the animal where it finds a lodgment.

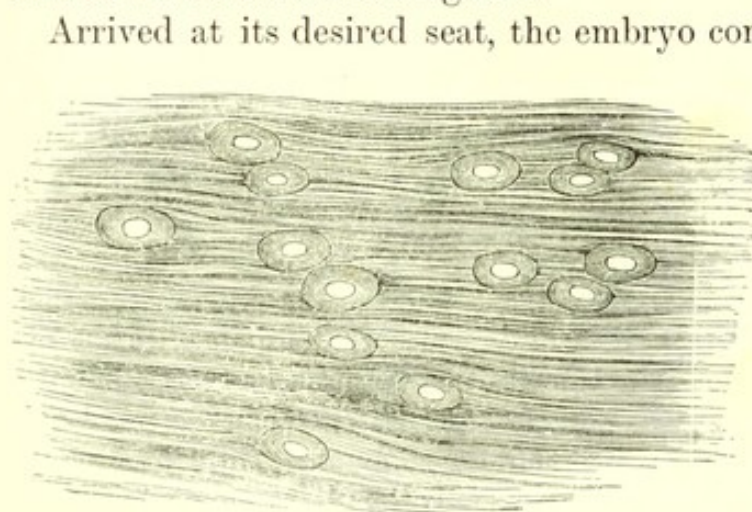


Fig. 122.—Cysticerci of *tænia solium* in muscle. Natural size. (LEUCKART.)

Arrived at its desired seat, the embryo comes to a state of rest, and after a time develops into the scolex, which in this tape-worm takes the form of the so-called **Cysticercus** or **Bladder worm**. The appearance of these cysticerci in the muscular tissue is shown in Fig. 122, which is drawn of the natural size. The complete cysticercus or scolex is com-

posed of a sac, connected with which is a head, which closely resembles the head of the mature worm (Fig. 123).

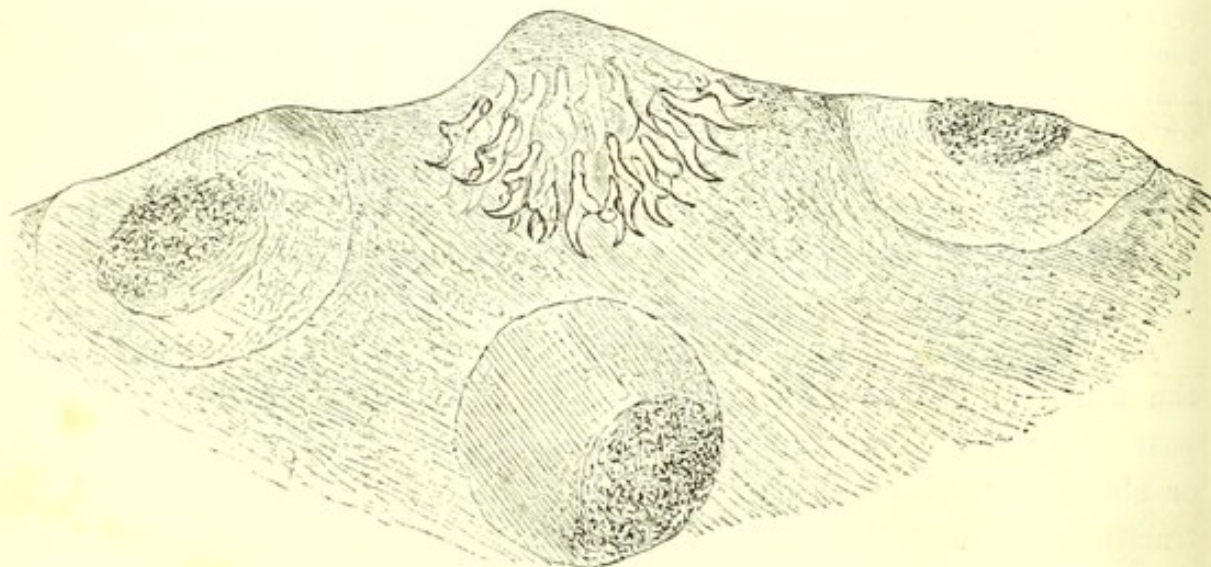


Fig. 123.—Head of *cysticercus cellulose* found in substance of brain.  $\times 90$ .

The observations of Leuckart and others have thrown much light on the develop-



ment of the scolex as they are based on actual experiments in which swine were fed with the ova.

In the first place a vesicle or cyst is formed in the muscle. After a time a slight thickening of the wall of the cyst appears. This grows inwards into the cyst, carrying with it, however, the external wall, so that the projection inwards is hollow with an internal canal continuous with the external surface of the cyst, and so opening externally; this is shown in Fig. 124, where a portion of the wall of the cyst is preserved, and the projection is shown with its internal cavity communicating with the surface of the cyst. This projection enlarges, and by and by the peculiar structure of the head, namely, the four sucking discs and the hooklets, show themselves. But these are formed inside the canal, near its inner extremity, and they are in an inverted position as compared with those of the mature tape-worm. The head with its hooklets is thus at the bottom of the canal, and the four suckers, looking towards each other, follow. After a time the head acquires the power of inverting itself outwards, and thus projecting from the vesicle, or again withdrawing itself within the vesicle as before. This is effected by means of muscular fibres. For the completion of this phase of development a period of from three to four months is required from the time of the ova being taken into the alimentary canal.



Fig. 124. — *Cysticercus* with beginning of development of head. A portion of the original cyst is shown with the projection inwards of a hollow process which communicates externally.  $\times 25$ . (LEUCKART.)

It sometimes happens that this scolex of the *tænia solium* develops in the human subject, and then it is often spoken of as the *Cysticercus cellulosæ*. It occurs chiefly in the brain, in the eyeball, and in muscle. It is to be remarked that in rare cases the scolex assumes in the brain a very peculiar character. The cyst, developing in the membranes on the surface of the brain, presents pouches and swellings which give it somewhat the character of a bunch of grapes, and so has arisen the designation *Cysticercus racemosus*. The cysticercus is usually surrounded by a connective-tissue capsule which is produced from the surrounding tissue and encloses both cyst and head, but not infrequently, especially in the brain and eyeball, it is devoid of this secondary capsule. In that case the vesicle sometimes grows to considerable dimensions, and the head is able to protrude itself and move about in various directions, perhaps in the ventricle of the brain or the eyeball. In these parts the scolex may produce considerable disturbance.

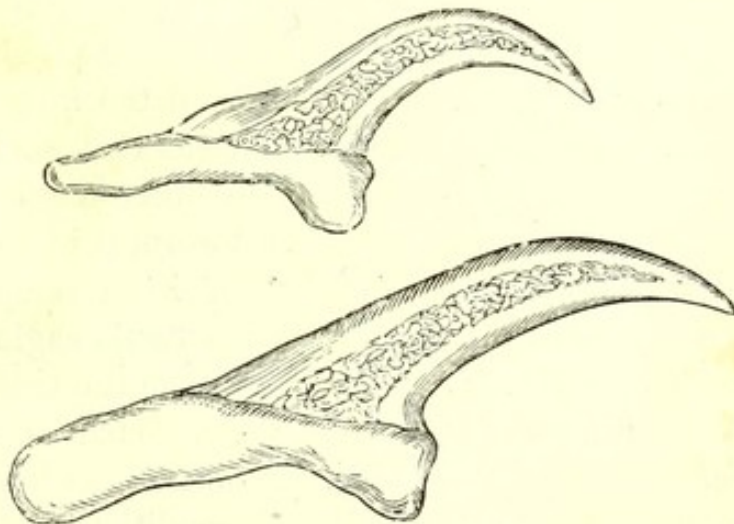


Fig. 125.—Hooks from head of *cysticercus cellulosæ*.  $\times 350$ .





Fig. 126.—Hooks from *tænia echinococcus*.  
× 350.

Although capable of a considerable duration of life, after a time the scolex usually dies, and then it shrinks and becomes, probably, incrustated with lime salts. The hooklets are the most resistant parts, and it is of some consequence to observe the size and general appearance of these. In Fig. 125 are represented the hooklets of this magnified 350 times. They are to be contrasted with those of the *tænia echinococcus* (Fig. 126),

which are magnified to the same extent.

It will be seen that in the *tænia solium* there are two sizes of hooklets, a larger and a smaller, and by comparison with Fig. 123 it will be seen that these are arranged alternately around the rostellum. In this as in other respects, the heads of the scolex and of the mature worm are identical.

We have now to trace the further progress of development where living scolices are taken into the alimentary canal of man. In the first place, the vesicle and everything but the head and neck are lost, and we



Fig. 127.—A single head of *tænia solium* before segmentation has begun. It shows movements of its suckers, etc. × 25. (LEUCKART.)

have a small creature which has considerable power of elongating and moving about its suckers, as shown in Fig. 127. The head now fixes itself to the wall of the alimentary canal, and the body begins to develop from its posterior extremity. It takes eleven or twelve weeks for the worm to assume its full dimensions, and at

the end of that time it begins to shed proglottides. The worm is of tolerably long life, and may inhabit the intestine of its host for many years. It not infrequently happens that several co-exist in the same person; as many as 30 or 40 have been observed.

***Tænia mediocanellata*.**—This worm is in most respects like the *tænia solium* in the various stages of its development and its structure. Leuckart calls this tape-worm the *Tænia saginata*.

The strobilus is a larger worm than the *tænia solium*, measuring from about thirteen feet in the contracted state to about twenty-four feet when extended. Fig. 128 shows the head and neck of this worm in the contracted and relaxed conditions. The difference in length and breadth is very striking. The head has no rostellum or circle of hooks, but it possesses four large sucking discs which are usually



surrounded by zones of pigment. In the greater part of the worm the segments are broader than they are long, attaining a breadth of about half an inch. But as we come to the fully mature proglottides with embryos in the uterus, then they are considerably elongated and at the same time narrower. The number of segments is greater here than in the *tænia solium*, reaching as high a figure as 1,300. There are generally about eight discharged from the posterior extremity daily, and these very often find their way outwards, through the anus, by their own movement.

The sexual organs, except the uterus, are essentially the same as in the *tænia solium*. The uterus, however, presents in the mature proglottis a much larger number of lateral offsets, as many as twenty to thirty, and these mostly branch dichotomously instead of ramifying. (See Figs. 129 and 130.)

The individual segments, like the worm as a whole, possess muscular fibres and are capable of elongating and contracting.

The scolex form of this worm is found chiefly in cattle, and it inhabits mostly the muscles, but is also met with in other organs. The cysticercus measures about the third of an inch, and is of a roundish shape.

This worm is nearly equal in frequency to the *tænia solium* in the human subject. Its cysticercus is not known to occur in

man. The worm may live for many years, at least as long as eleven, and, as some assert, up to twenty or even thirty years.

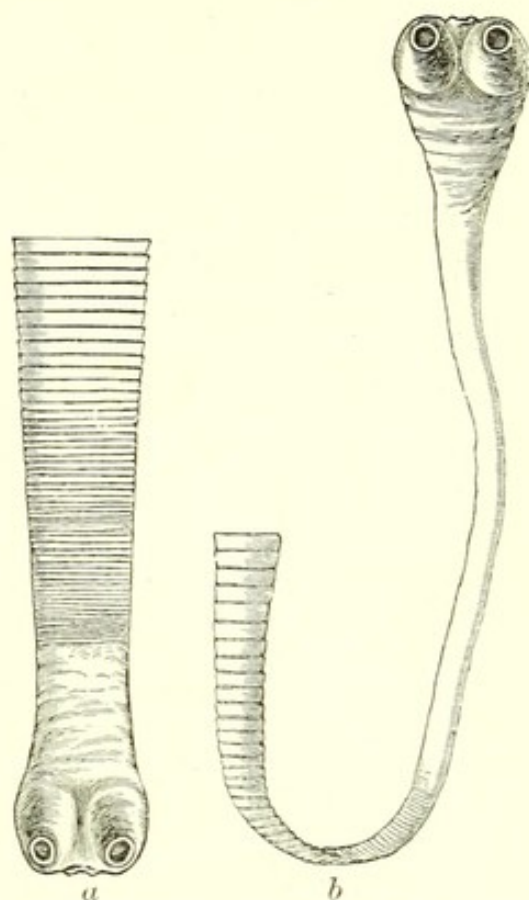


Fig. 128.—Head and neck of *tænia mediocanellata*; *a*, in contracted, and *b*, uncontracted state.  $\times 8$ . (LEUCKHART.)

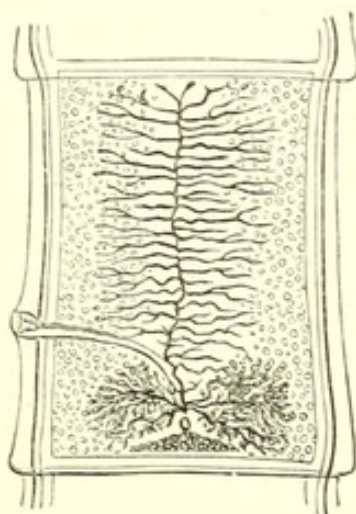


Fig. 129.—Immature proglottis of *T. mediocanellata*.  $\times 5$ . (LEUCKHART.)



Fig. 130.—Mature proglottis of *T. mediocanellata*.  $\times 2$ . (LEUCKHART.)



**Tænia echinococcus.**—In the strobilus form this is a comparatively insignificant worm (Fig. 131). It inhabits the dog, and there are generally several individuals present at the same time. The



Fig. 131.—Adult *tænia echinococcus*.  $\times 12$ . (LEUCKHART.)

total length of the worm is about an eighth of an inch, and it consists only of four segments, including that which carries the head. In the fully developed state the last segment exceeds in length the rest of the worm altogether (see figure). The head is like that of the *tænia solium* in miniature, being very greatly less in size. It has a rostellum with thirty to forty hooklets, and four sucking discs. The last segment develops a large number of eggs, as many as 5,000.

These eggs develop the usual embryos with six spines, and if they find their way into the intestinal canal of man, they pass out into the tissues. Settling in some organ of the body, they show the most extraordinary powers of development, producing the condition commonly called **Hydatids**.

It is proper to say here that hydatids occur in the form of large cysts, often of very complex arrangement, and they should be carefully distinguished from the cysticerci which form small cysts not more than half an inch in size. The hydatids occur in the majority of cases in the liver. Neisser has collected no less than 986 cases of hydatids in man, and he gives the scale of frequency in the different organs as follows:—Liver, 451; lungs and pleura, 84; kidneys, 80; muscle and subcutaneous tissue (including the orbit), 72; brain 68; spinal cord, 13; female organs of generation (including the mamma), 44; male organs, 6; pelvis, 36; organs of circulation, 29; spleen and bones, 28; eye, 3.

When the embryo reaches the liver or other resting place, it soon develops into a cyst which at first is of slow growth. The membrane of the cyst is of considerable thickness, and consists of an external cuticle in several layers and an internal parenchymatous layer containing muscular fibre and a vascular system. Inside the original vesicle arise very frequently secondary vesicles, and inside these even tertiary ones, the successive vesicles being sometimes spoken of as daughter or grand-daughter vesicles. It sometimes happens that the secondary vesicles project outwards, and form a series of external vesicles which may separate from their mother and attain an independent development alongside of her. This latter form is particularly common in the domestic animals, and it is variously designated exogenous hydatids, or *echinococcus scolecipariens* or *granulosus*. There is a third form which has been met with in man, and always in the liver. The parasite develops a congeries of small vesicles, from the size of a grain of wheat to that of a pea. These are embedded in a gelatinous tissue and sometimes possess gelatinous contents. As the whole is surrounded by a firm fibrous capsule, the



tumour is a somewhat solid one, and on section it presents a peculiar alveolar appearance. From this structure it is called the alveolar form, and it should be particularly borne in mind, as this condition has been frequently mistaken for a tumour, especially before Virchow demonstrated its true nature.

Except in the case of the alveolar form, the vesicles, both primary and secondary, enlarge very much and give rise to tumours of very large dimensions, so as sometimes to produce serious disturbance by their mere size. Those of the liver are usually the largest, and they may come to weigh as much as twelve, twenty, or even thirty pounds. The simple vesicles, in which no daughters develop, attain the size of an orange or a fist.

In all forms of hydatids the whole parasite is surrounded by a fibrous capsule, developed by the organ in which it has its seat. As the cysts enlarge, this also increases in size.

We have now to consider the formation of the heads of the worms, which differs in certain respects from that of the other tæniæ.

In the walls of the vesicles, either primary or secondary, are to be seen when they are perfectly fresh, a number of small white points which have their seat in the internal wall. These are called **Brood-capsules**, and it is always in connection with them that the heads or scolices develop. The brood-capsules are little vesicles, in the walls of which the echinococcus head grows. The heads begin (see Fig. 132, in which the development of heads in brood-capsules is shown) as projections outwards of the

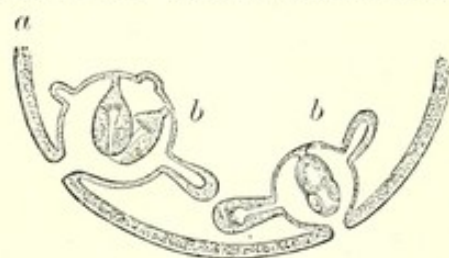


Fig. 132.—Diagrammatical illustration of development of echinococcus heads in brood-capsules. *a*, Wall of cyst. *b b*, Brood-capsules, with heads in various stages of development. (LEUCKART.)

wall of the brood-capsule. The projection is hollow, and communicates with the interior of the brood-capsule. The head develops inside this projection, as in the case of the tænia solium, and very soon acquires the power of inverting itself. When it does so, it projects into the brood-capsule, so that in this respect the brood-capsule is not like the cyst of the cysticercus as the scolex projects outwards from the latter. A single brood-capsule develops several heads, up to twelve, and they may be found either in the extended or inverted position. All the heads are contained in brood-

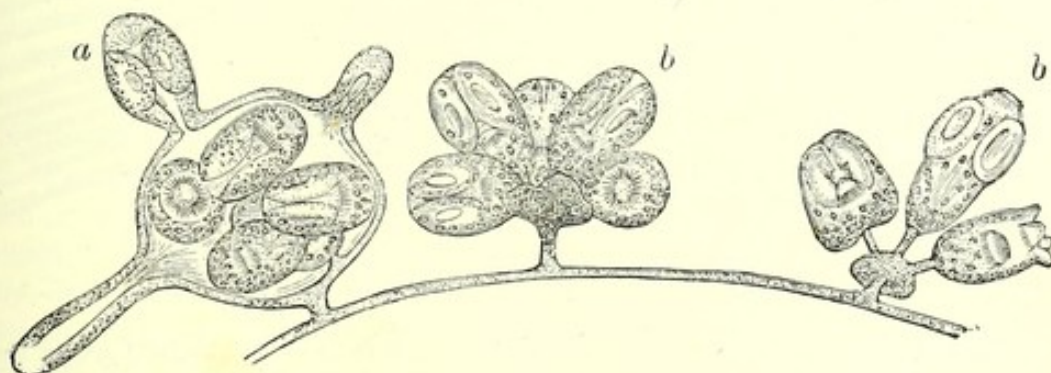


Fig. 133.—Brood-capsules in connection with wall of vesicle. *a*, A capsule in normal unruptured state. *b b*, Ruptured capsules.  $\times 40$ . (LEUCKART.)

capsules, but if after death or during removal the brood-capsule bursts, then an appearance may be produced as if the heads were attached to the wall of the cyst itself. If the capsule bursts, its remains may gather round its stalk and the heads stand up from this as in Fig. 133 *b b*. Heads may also be found lying free if the



capsules have burst. It is to be added that sometimes the vesicles remain barren, neither brood-capsules nor heads developing in them.

**The heads** which are formed in vast numbers inside the complicated system of cysts are exactly like those of the mature worm. They are very minute objects, measuring about  $\frac{1}{70}$ th of an inch in long diameter, and just visible to the naked eye. They possess a proboscis with a ring of hooklets, and four suckers. They are also provided with a water-vascular system, and in their parenchyma abundant calcareous particles are to be found.

The animal may die spontaneously, or be killed by the fluid which fills the vesicles being drawn off. In that case the vesicles shrink, and their contents become converted into a fatty debris, which afterwards may become infiltrated with lime salts. In this way the hydatid mass may be represented by a cyst filled with an atheromatous material. This may dry-in, and at last we may have nothing left but a stony or mortar-like mass, in which careful search may still discover the hooklets (Fig. 126). Besides the distinctive hooklets, or even without them, there are usually in these old cysts bits of the chitinous membrane of the parasite. The wall of the hydatid cyst consists of two layers, an external (called by Huxley the ectocyst) and an internal (the endocyst). The ectocyst consists of a structureless stratified membrane, composed of a chitinous substance, from which circumstance it is often called the cuticle. As this chitinous membrane is very resistant it may be found in the midst of the grumous contents and enable the structure to be recognized. In the case from which Fig. 134 was taken the diagnosis

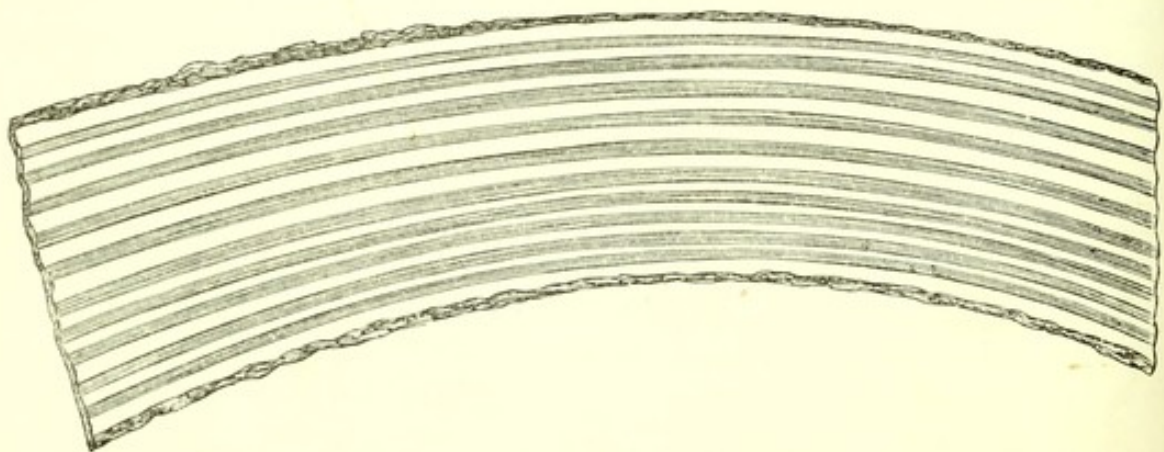


Fig. 134.—Transverse section of the chitinous membrane from an old hydatid cyst of the kidney.  $\times 90$ .

was made chiefly by flakes of this membrane being found, the search for hooklets being in vain.

In regard to the distribution of the echinococcus, it is of pretty frequent occurrence in all known lands, but it is particularly common in Iceland where the men, living in close companionship with the dogs, are much exposed to infection. It is also said to be very common in Australia and neighbouring colonies.



**Bothriocephalus latus.**—This belongs to the family of Bothriocephalidæ, and is the largest tape-worm which occurs in man. It attains a length of from 16 to 26 feet, and possesses from 3,000 to 4,000 segments, which are mostly much broader than long, although the last ones (see Fig. 136) become longer and narrower so as to assume more of a square shape. The breadth at the widest part is about half an inch. The worm is also thick and heavy.

The head (Fig. 135) is oval, and about the twenty-fifth of an inch in breadth.

It is blunt at the extremity, and possesses neither hooklets nor suckers, but fixes itself by means of a slit-like groove on either side of the head (see figure).

The sexual organs, and especially the uterus, occupy the middle part of each segment, where they form a rather prominent knot or rosette (Fig. 137). The uterus is composed of a convoluted tube which gives the rosette-like appearance just mentioned. The sexual organs open in the middle line near the anterior extremity of the proglottis. The eggs are oval in form and are covered by a brown shell.

The scolex form of the worm long eluded observation. It was known that a six-spined embryo formed in the eggs in the usual way, but the habitat of the cysticercus was unknown. Braun has finally demonstrated its existence in the pike and turbot. It was found in

the muscles, sexual organs, liver, spleen, etc., of these fish. Braun proved that it was the scolex of this animal by feeding dogs with it. A tape-worm identical with the bothriocephalus developed. By this discovery all the tape-worms which occur in Europeans have been traced in their mature and cysticercus forms.

This worm is of rather frequent occurrence in Switzerland and north-east Europe, but it is not unknown in this country. The frequency of

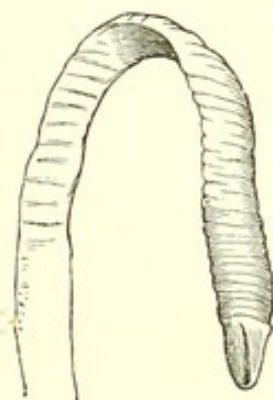


Fig. 135. — Head and portion of body of *Bothriocephalus latus*  $\times 8$ . (LEUCKART.)

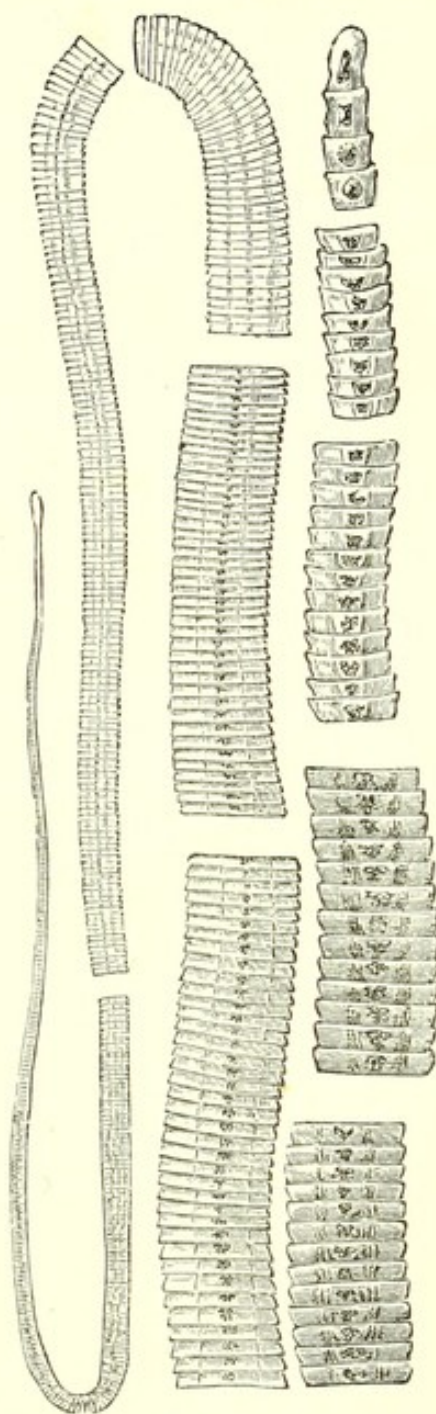


Fig. 136. — The *Bothriocephalus latus*. Natural size. (LEUCKART.)



fresh-water lakes in Switzerland explains its common occurrence there from the use of the fish which form the hosts of the scolex form. Like the other tape-worms the mature worm occurs in the small intestines.

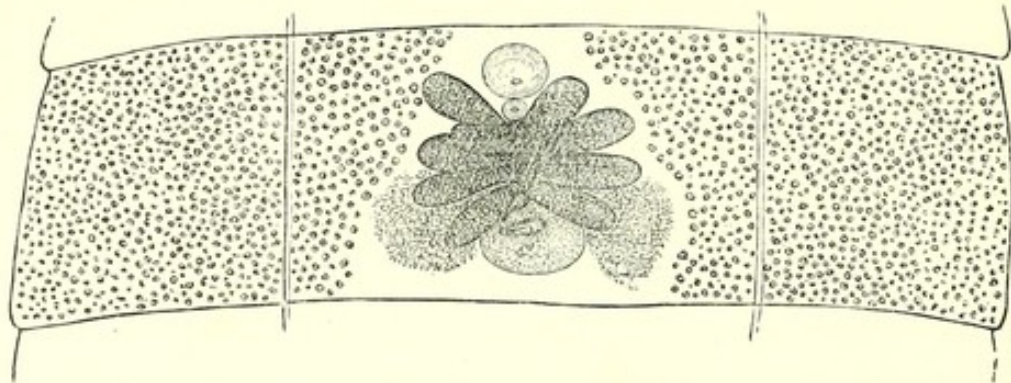


Fig. 137.—Proglottis of *Bothriocephalus latus*, showing female organs.  $\times 12$ . (LEUCKART).

Several other small and infrequent tape-worms have been met with in man.

The *Tænia nana* has only been observed once and in Egypt. It is very small, scarcely an inch in length, and about the fiftieth of an inch in breadth. It possesses a rostellum with hooklets and four sucking discs.

*Tænia flavopuncta*.—It has also been observed but once, in America. It is about a foot in length, and about one twentieth of an inch in breadth. The head of this worm is not known, but it is believed to possess hooklets.

*Tænia Madagascariensis*.—This form has been found in an island on the Madagascar coast. It is about 3 inches long, and the head is not yet known.

*Tænia cucumerina*.—This worm is from 7 to 10 inches long. Its head possesses a rostellum with a quadruple circle of hooklets to the number of about 60. The proglottides reach a breadth of about the twelfth of an inch. It occurs very frequently in dogs and cats, and is often present in large numbers, especially in dogs. It has been found in several cases in man, and it seems to be not infrequent in children. The scolex form has lately been found in the dog-louse (*Trichodectes canis*), and it can be readily understood how in the process of licking itself the dog often swallows its host and becomes itself the host of the strobilus. It may be conveyed to children from the tongue of the dog.

*Tænia marginata*.—This is a tape-worm of large size, and of common occurrence in the dog. Its usual length is about 5 feet, but it may be as long as 8 feet. It is very like the *tænia solium*, the head possessing a rostellum with hooklets of about the same size but more numerous. On the whole, however, the worm is smaller, and so are the proglottides. The scolex form inhabits swine and the ruminants, and as it often develops a large vesicle or set of vesicles, it is liable to be confused with the *tænia echinococcus*. The scolex form is called the *cysticercus tenuicollis*. Neither mature worm nor *cysticercus* is met with in man.

The dog also frequently contains two other *tæniæ*, namely the *T. Serrata* and *T. Cœnurus*. These also resemble the *tænia solium*, and the latter is important, as its scolex form frequently attacks lambs, and, lodging in the brain, is the cause of the very fatal disease "staggers."

**4. Nematoda or Round-Worms.**—The round-worms have elongated bodies and possess a well-developed digestive apparatus, with mouth, œsophagus, stomach, intestines, anus. The sexes are separate. Some of



them bear living embryos, while others produce eggs which become free and afterwards develop embryos.

**Trichina Spiralis.**—This worm is met with in the muscular substance of man, and occurs there in immense numbers, producing the disease **trichinosis**. We shall see afterwards that this is not the mature form of the worm, but it is in this form alone that, for the most part, it is accessible to us, and it will be convenient to begin with its description here.

The affected muscles, as seen with the naked eye, seem, for the most part, to be dusted throughout with fine white particles like sawdust. These are most abundant near the places where the muscular fibres are inserted into the tendons. As a rule the particles are most abundant in the muscles of the trunk, the diaphragm, the intercostal muscles, and those of the abdominal wall, but they may extend to all the voluntary muscles of the body, even the most distant ones of the hands and feet.

On microscopic examination of the fine particles, they are found to consist each of an oval cyst with a tolerably thick wall (see Fig. 138),

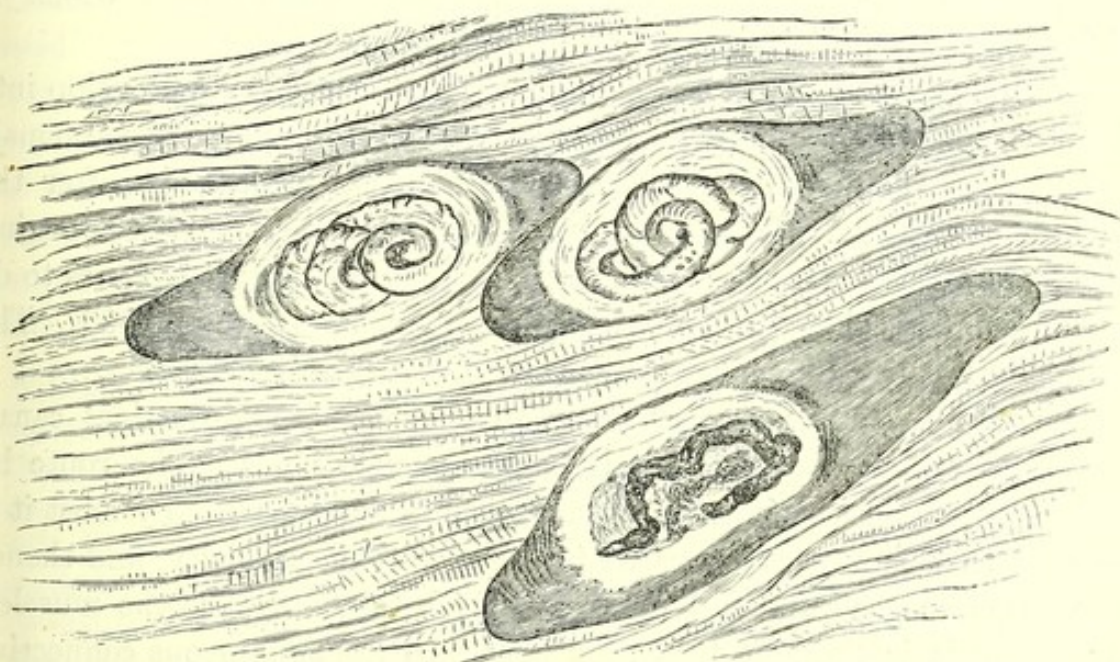


Fig. 138.—*Trichina spiralis* in muscle. The elongated shape of the cysts is due to the fact that these were near the insertion of the muscle into its tendon. In the lowest specimen the worm is dead and calcified.  $\times 90$ .

within which is a small worm coiled up in a spiral manner. The cyst has very often abundant calcareous particles in its wall, especially at the poles, and, if the case is an old one, the impregnation with lime may be so great as to hide the parasite unless the salt be first dissolved out with an acid. When an acid, such as dilute hydrochloric, is used, the lime dissolves with some evolution of gas, and the whole structure becomes very transparent. Sometimes the worm dies in its capsule, and in that case



the wall thickens and the cyst collapses to some extent on the remains of the worm, which itself often becomes infiltrated with lime (see lowest specimen in figure). The worm in the muscle is in the immature state and quiescent, lying rolled up in this manner for years it may be. It may be present in immense numbers, even in millions, in the same person.

The parasite in man is derived from the pig, in whose muscles the embryos occur in the same fashion as in man. If, now, a piece of muscle containing them in their living state—that is to say, not killed by cooking the meat—be eaten, they undergo further development in the intestinal canal. The capsule is dissolved by the gastric juice, and the embryo set free. In the muscle, the embryo, if uncoiled, would measure about the twenty-fifth of an inch in length; but now it grows rapidly, and in the course of two and a half days it reaches the **adult form** when the female is about one eighth of an inch in length; and the male slightly less. The male possesses a testicle consisting of a convoluted tube. The female has an ovary, vagina, and uterus. The adult worm has an intestinal canal from end to end, which is divisible into œsophagus, stomach, and intestine.

The impregnated ova pass into the uterus where they develop into living embryos of minute size. In six or seven days after the female has attained sexual maturity, that is, eight or nine days after the trichinous muscle has been eaten, the birth of living embryos begins. The female gives birth to large numbers, and probably continues to do so for some weeks, thus producing as many as 1000 to 1300. The adults do not live longer than five to eight weeks altogether.

The minute embryos now begin to penetrate the intestinal canal, and they swarm outwards to the voluntary muscles. The route by which they reach the muscles is not absolutely certain. By most it is thought that they pass outwards into the peritoneal cavity, and thence into the connective tissue around, by which they travel to the muscles. By others it is thought that they pass into the submucous connective tissue, thence into the connective tissue of the mesentery, and so onwards. It is probable that they find their way by both these routes, but it is inconceivable that, as some suppose, they get into the blood-vessels, as the vessels available to them are the portal radicles which would take them to the liver.

Swarming outwards from the intestine they reach first the muscles of the trunk, where they are usually most abundant; they then pass to those of the neck and larynx; and, lastly, to those of the limbs. Arrived at the muscles they grow larger, and apparently wander about for a time. They penetrate inside the sarcolemma of the primitive



fibre of the muscle, and destroy the sarcous substance. In about fourteen days they have attained their full size, and begin to settle down. As they pass along inside the sarcolemma they are arrested at the insertion of the fibre into the tendon, hence they are particularly numerous near tendons, and here also the cysts, subsequently formed, are often much elongated (as in the figure). The sarcolemma collapses as the sarcous substance is destroyed, and as the worm coils itself up spirally the sarcolemma forms for it an oval cyst. The worm itself seems also to add to the cyst a layer of its own. It is not uncommon to find two, or even more worms, in one cyst. In the muscles the worms assume a quiescent state, as already mentioned, and may remain so for years (as long as eighteen years has been proved), the cyst being impregnated with lime. They produce considerable destruction by piercing the sarcolemma, and disintegrating the sarcous substance, and there is often to be found a germination of the muscle nuclei around the worm. The death of the host does not cause the death of the trichinæ. They will live in putrid flesh for weeks and remain capable of further development.

During the migration of the embryos considerable irritation is produced. There is in the first week intestinal catarrh (diarrhœa) with fever, and the case may be mistaken for typhoid fever. Later the muscles become stiff and painful, and œdema of the skin, especially of the face, may develop. This œdema of the face, which occurs about the seventh day, is said to be of special diagnostic significance. The symptoms are usually at their height in the fourth or fifth week, and death occasionally ensues.

Besides in man, trichinæ have been found in the muscles of the pig, cat, rat, mouse, marmot, polecat, fox, marten, badger, hedgehog, and racoon. By some it is believed that the rat forms the permanent source of infection, as, when one of these animals dies it is eaten by its neighbours, and so the infection spreads. From their habits, it will be understood how swine sometimes partake of dead rats. The parasite will be communicated to man by eating imperfectly cooked swine's flesh. It is said that a temperature of 50°-55° C. or 120°-130° F., is enough to kill the embryos, but it is quite conceivable that when large pieces of flesh are cooked rapidly, some parts may escape the thorough penetration of the heat.

The search for trichinæ in the muscles of swine before the flesh is sold is compulsory in some countries. For the examination pieces of muscle (preferably from the diaphragm and larynx) are snipped off with scissors and spread out in water on a microscopic slide. Some liquor potassæ may be added to make the preparation more transparent. It is then to be examined with low magnifying powers and afterwards with higher. Several specimens should be prepared from each animal.

**Ascaris lumbricoides.**—The common round-worm is probably the commonest entozoon in the human subject. It occurs very frequently in children, and inhabits chiefly the small intestine. In its colour and



general appearance it resembles the common earthworm. It measures 6 to 16 inches in length, is marked by transverse striæ and tapers to both ends. Like other adult nematodes it possesses an intestinal canal from end to end. The female produces a large number of oval eggs which have a dense shell.

The worm mostly occurs singly or in pairs, but is frequently present in considerable numbers up to one or two hundred. From the intestine it may pass into the stomach and be vomited, or may be discharged per anum. It has been known also to pass up the œsophagus and into the nostrils and sinuses of the head, or by the larynx into the bronchial tubes. Sometimes it penetrates into the bile ducts, which it may obstruct, or through the intestinal wall into the peritoneal cavity.

In the intestine the irritation of the worms produces catarrh, and by reflex action this is supposed to lead to certain nervous symptoms. When present in large numbers, the worms are sometimes rolled up in a ball, and in this condition they may obstruct the intestine. In cases where they have perforated into the peritoneum they have given rise usually to local abscesses pointing chiefly near the umbilicus or groin. More rarely they have led to general peritonitis.

**Ascaris mystax.**—This is a small round worm which occurs in the cat, and is said to be always present in the intestine of that animal.

**Oxyuris vermicularis or Thread-worm.**—This is an exceedingly common parasite. It is white in colour, and the male measures about an eighth of an inch, and the female about three eighths in length. It possesses an alimentary canal from end to end. The eggs are oval, and have a dense shell. The animal inhabits mostly the large intestine. It is stated by Zenker and Heller that the mature female is in the large intestine, the males and young being in the small. The worm often wanders, especially during the night, to the neighbourhood of the anus, where it produces itching. Sometimes it passes over to the vagina, and up into it. It produces catarrh of the bowel, and, as in the case of the ascaris, nervous symptoms are ascribed to it.

**Trichocephalus dispar** (the whip-shaped worm).—It is of frequent occurrence in the cœcum and neighbouring parts of the intestine. It measures  $1\frac{1}{2}$  to 2 inches in length, and has the peculiarity that the anterior portion is much thinner than the posterior, forming a long thread, like the lash of a whip, which is buried in the mucous membrane. The eggs possess a brown shell. The embryos have been traced in water and moist earth.

**Dochmius duodenalis** (*Strongylus Duodenalis*. *Anchylostomum duodenale*).—This worm is not met with in this country, but occurs in



Egypt, Italy, and tropical lands. It has been found frequently among the workers at the St. Gothard Tunnel in Switzerland. It is a third to half an inch in length, and it possesses a mouth armed with four strong teeth. By means of these it fixes itself on the mucous membrane where it sucks the blood. When present in considerable numbers, as it often is among the valvulæ conniventes, it may give rise to considerable loss of blood and serious anæmia.

**Strongylus gigas.**—This is a large worm, reaching a length of over a yard, and a thickness of about three eighths of an inch. It has been met with a few times in the pelvis of the kidney in man, and more frequently in the kidney, bladder, lungs, and liver of dogs.

**Filaria medinensis** (*Dracunculus medinensis*, or *Guinea-worm*).—This parasite is of frequent occurrence in tropical lands, where it is met with in the tissues of the foot and leg chiefly. The female is a long thin worm from 12 to 40 inches in length, and it alone is known as a parasite. The male is much smaller. The worm wanders to some extent in the loose subcutaneous connective tissue, and may give rise to considerable irritation. When mature it presents its extremity at the surface, and a small pustule forms from which the extremity projects. The worm may then be removed gradually by rolling it gently round a quill from day to day as it becomes exposed, care being taken not to break it, in which case the part left in may give rise to severe inflammation.

**Filaria sanguinis hominis.**—This name was originally given to a minute thread-like worm which has been found to be really the embryo of a larger worm. The embryo, as its name implies, has been found in the blood, although it has also been observed in chylous urine and elsewhere. The adult has been seldom found, but is believed to live in lymphatic vessels, and so the name *filaria sanguinis* is not strictly applicable to it. Cobbold has given it the name *filaria Bancrofti*, from the observer who first discovered the adult. As it is inconvenient to give two different names to the adult and embryo forms of the same animal, we shall refer to them under the original designation of *filaria sanguinis*. In some respects the two forms of this worm are comparable to those of the *trichina spiralis*, which also obtains its name from the embryo form.

The adult *filaria* has the sexes distinct, and as yet the female alone has been fully examined, only a portion of a male having been discovered. The female is a long hair-like worm, three or four inches in length, and only  $\frac{1}{100}$  inch in breadth. It has an opaline appearance, and, as described by Manson, it looks like a delicate thread of catgut animated and wriggling as it lies in the tissues.



By the observer just named it was found lying in a lymphatic vessel while he was removing a scrotum which was affected in a way to be mentioned afterwards, and it appears that this is the usual habitat of the animal. The male is probably much smaller than the female, and they are supposed to live together in the same vessel.

The female is believed to live for years in the same position, and gives birth almost continuously to large numbers of embryos. Near the head is the vagina, and behind it the uterus with two horns is found, stuffed with ova, and extending

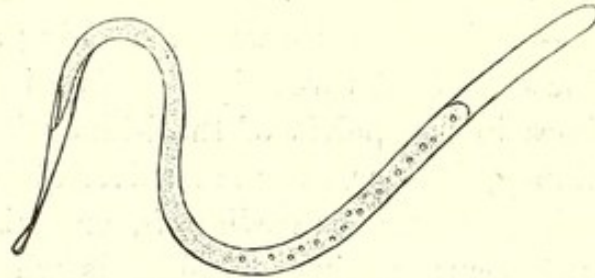


Fig. 139.—*Filaria sanguinis* (embryo) after preservation in weak spirit. The sac is seen at each end of the worm.  $\times 300$ . (LEWIS.)

almost to the tail. Manson states that the embryos can be seen escaping fully formed from the vagina, in exactly the same form as in the blood. But the female sometimes gives birth to ova, as these have been found in the lymph and have been supposed to be the principal factors in producing certain local affections to be afterwards mentioned.

The living embryo (Fig. 139 and 140) is about  $\frac{1}{90}$  inch in length and  $\frac{1}{3600}$  inch in thickness; its breadth therefore nearly corresponds with that of a red blood-

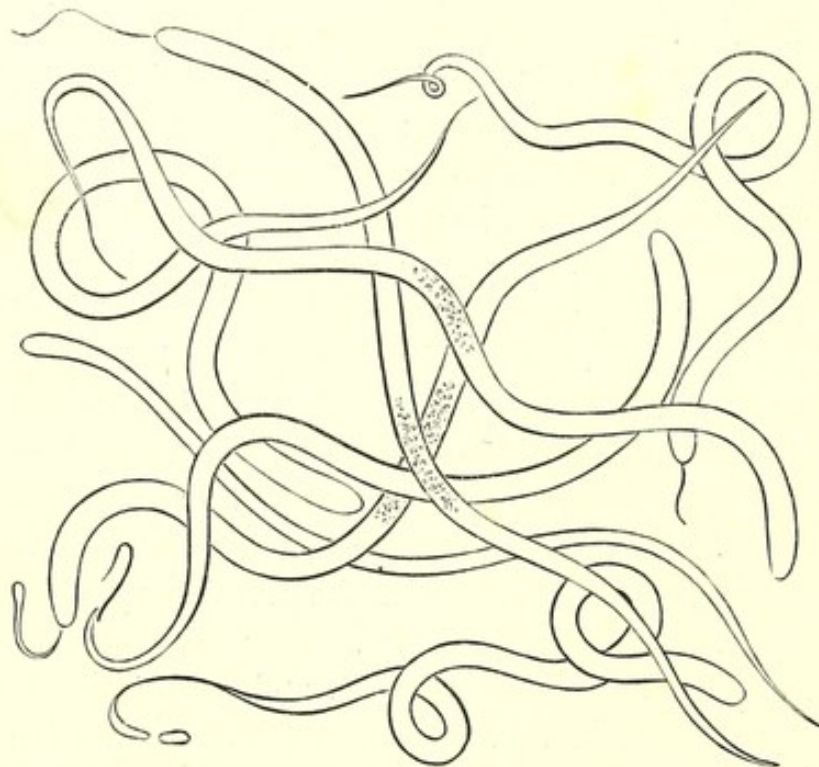


Fig. 140.—*Filaria sanguinis* as they appear in the living state in the blood. From a case of chyluria.  $\times 300$ . (LEWIS.)

corpuscle. It is enclosed in a delicate sac, which is rather longer than itself, so that while moving a portion of the sac extends beyond its extremity. The



movements as seen in the blood and lymph are very active and snake-like in character.

This embryo is present in the blood of about 10 per cent. of the natives where it is endemic, and most of these do not appear to suffer in health from it. A very remarkable circumstance is that, during the day, the parasites are, unless in exceptional cases, absent from the blood, but about six or seven o'clock in the evening they begin to appear, and by twelve o'clock are so numerous that as many as a hundred may be counted in every drop of blood. As morning approaches they diminish in numbers, and by eight or nine o'clock they disappear entirely. This regular rhythm may apparently go on for years. When the periodicity of the bodily functions is interfered with by the person sleeping during the day and doing his work at night, then the parasite is present in the blood in the day time and absent at night.

The periodicity of the *Filaria* is variously accounted for. Manson and others believe that the embryos retire during the day to the internal organs where they lie quiescent. Myers rather believes that they die out daily and a new swarm is produced. This view he supports chiefly on the ground of observations, according to which the embryos, as morning approaches, show signs of languor and loss of condition. He also asserts that the adult female is sufficiently fertile to replenish the stock daily, and that, if this be the case, there must be a daily destruction.

In finding their way from the site of the parent worm in the lymphatics to the blood, it is probable that the embryos travel by way of the lymphatic fluid itself. Being not greater in breadth than a blood-corpuscle, and possessing powers of locomotion, they may be supposed to traverse the lymphatic glands with the lymph, and to reach the blood by the thoracic duct. It is also possible that they may get into the blood by penetrating through the blood-vessels. In the blood they will readily circulate with that fluid, and from their size they are capable of passing through the capillaries and repeating the circuit with the blood-corpuscles. They do not pass through any further stage of development in the blood, the embryos found there being exactly like those which escape from the mother. It is clear also that they do not, either by themselves or their products, act injuriously on the blood or tissues unless by using the nutritious material.

It has been stated above that the female sometimes gives birth to ova instead of living embryos, and it is probably in consequence of this that various forms of disease are found to occur in connection with this parasite. The sac which has been mentioned as enclosing the embryo is really the chorionic envelope of the ovum stretched and rendered elastic. But if ova are born instead of embryos then we have a body  $\frac{1}{80}$  inch to  $\frac{1}{75}$  inch in length. This body is, of course, much thicker than the extended embryo, and when carried by the lymph to the glands it will be unable to traverse these. An infarction of the fine afferent vessels of the gland will thus occur, and as the vessels leading from the part where the mother is situated are plugged one after another, there will by and by be a complete stasis of the lymph in the vessels of these parts. It is probable that dead embryos though extended will similarly plug the lymphatics.



The effect of this complete plugging of all the lymphatic channels seems to be an accumulation of lymph in the vessel below the seat of obstruction. The lymph thus distending the vessels often escapes, apparently by rupture. The locality of the parent will determine the exact nature of the resulting disease. If it be in the lymphatics of the pelvis or lumbar region there may be distension of the lymphatics of the kidneys, ureters, or bladder. In that case **Chylous urine** is found, and the embryos of the parasite have been observed in the urine. If the adult be in the lymphatics of the leg, there will be gradual plugging of the glands at the groin and accumulation of lymph in the legs and scrotum. The accumulation in the scrotum is of very frequent occurrence, giving rise to **Lymph-scrotum**, in which vesicles appear on the surface which occasionally discharge lymph.

It is to be observed that the condition in these various situations is not a proper œdema, which we have already seen to consist in over-filling of the serous spaces, but rather an over-distension of the lymphatic vessels with occasional rupture of them.

It has been asserted that **Elephantiasis** is one of the diseases dependent on this parasite, and is to be placed in the same category as lymph-scrotum and chyluria. But there are serious objections to this view. Elephantiasis has the features, as we have seen, rather of a specific inflammation, than a simple stagnation of lymph, and is more nearly allied to leprosy. It appears, however, that there are cases of lymph-scrotum in which the skin becomes greatly thickened, and these are sometimes called Lymphangiectatic elephantiasis.

The stages by which the embryo attains to the adult form, were, till lately, obscure. The embryo in the human body does not pass beyond the stage which it had reached at birth, and in order to complete the cycle of its development it must pass into the body of another animal. The intermediate host has been found to be a species of mosquito (Lewis, Manson). The particular species of mosquito is only present in districts where filaria occurs, in other districts the prevalent mosquito is incapable of propagating the parasite (Myers). The female of this animal has a proboscis which she inserts into the skin and through which she sucks blood into the stomach. With the blood the embryos are taken in, and they have been found in the stomach even in larger proportional numbers than in the blood of the person from whom they were taken. In the stomach the embryo passes through certain stages of development occupying from four to six days. At this period the mosquito dies, and probably falls into water, the parasite passing from the stomach. The further stages are not known as yet, but by the time the mosquito dies the parasite has already acquired a boring apparatus fitting it to penetrate the tissues. If it reaches the alimentary canal of man, it will then bore through the tissues till it finds its selected site in the lymphatics. It selects this site just as the trichina selects the muscles, or as the echinococcus may select the liver. In its new position it attains to sexual maturity, and being impregnated, gives birth to the embryos as we have seen.

The species of mosquito here concerned is, like other mosquitoes, nocturnal in its habits, so that it attacks affected persons at the time when the parasite is present in the blood. The occurrence of the embryos in the blood at night has been regarded as a provision of nature for the propagation of the species.

In animals, several different species of filaria have been observed. There is the *Filaria sanguinolenta* frequently observed in large numbers in the wall of the œsophagus and aorta in dogs. In certain countries it is said to occur in every third animal. It is here in the sexually mature condition along with eggs in every stage



of development. There is also the *Filaria immitis* s. *hæmatica*, found in the blood of the dog, etc.

## II.—EPIZOA OR EXTERNAL PARASITES.

These do not call for extended treatment here, as they are fully described in works on diseases of the skin. Little more than an enumeration of them will be attempted.

1. **Arachnida.**—Parasites belonging to this class occur both in man and animals. In one the larva inhabits the internal parts, while the adult is external.

**Acarus scabiei, or Sarcoptes hominis.**—This has an oval body just large enough to be visible to the naked eye, the female being one fiftieth of an inch in length, and the male about half that size. The anterior part has a head and four limbs, each of which has a sucker at its extremity. There are also four posterior limbs, all of which in the female have pointed extremities, but in the male two of them have suckers. The female burrows in the epidermis, forming tunnels, in which, as it proceeds, it deposits its eggs; it is usually to be found at the deepest end of the tunnel and the eggs at intervals. The eggs develop, and, as the epidermis desquamates, they come to the surface by degrees, the young being born usually as they reach the surface. The irritation of the animal in the epidermis gives rise to a slight inflammation, causing the formation of a papule. Usually there is great itching, and the scratching leads to further eruptions, especially in pre-disposed persons.

**Acarus folliculorum** (*Demodex folliculorum*).—This is an elongated animal about  $\frac{1}{125}$  of an inch in length, and provided anteriorly with four pairs of short feet. It is found in the sebaceous follicles, especially of the external meatus of the ear and neighbourhood of the nose. It seems to produce no special irritation.

**Pentastoma denticulatum s. tænioides.**—The larval form is very common in the rabbit, and has been not infrequently observed in man. It occurs usually in the liver, but has been observed also in the spleen, lungs, kidneys, and wall of intestine. It is a small animal about a fifth of an inch in length and a fifteenth in breadth. It presents about 90 segments, in each of which are stomata. The mouth possesses four hooks which can be withdrawn into chitinous sheaths. The larva in the liver surrounds itself with a capsule and forms a nodule about the size of a pea. In man they are mostly found dead, and the condition observed is that of a hard nodule surrounded by a fibrous capsule, inside which are the calcified remains of the animal, of which only the hooks may be recognizable.



The larva found in the liver and elsewhere is usually designated the *Pentastoma denticulatum*, while the adult is called the *P. tænioides*. The connection between the two was demonstrated by Leuckart. The adult is found chiefly in the nares of dogs, but also of some other animals, and on one occasion of man (Laudon). The adult is like the larva in form but much larger, the female measuring three inches, and the male about one. The mouth is devoid of hooks. The ova passing from the nostrils of the dog on to the grass are supposed to be taken into the stomachs of hares and rabbits, and to pass thence to the liver.

Many forms of arachnida occur in animals.

**Leptus autumnalis** (Harvest bug).—This is a small red animal, just visible to the naked eye. It is not a necessary parasite, but in some districts it invades the legs and burrows into the skin, thus causing excessive itching. Two other forms of *leptus* are described as occurring in America (Duhring).

2. **Insecta or Insects.**—The parasitic insects occur entirely externally. Some of them are not parasitic at all times.

**Pediculi or Lice.**—The head louse (*P. capitis*) lives among the hairs. It forms a chitinous sheath for its ova, which it cements to the hairs. The young, when they emerge from the egg, are like the adult in form, there being no further metamorphosis. The body louse (*P. vestimentorum*) is like the former but considerably larger. The ova are deposited in the clothing, especially the seams, where also the adults congregate. The crab louse (*P. pubis*) has its popular name from the fact that it has long curved claws with which it attaches itself to the hairs. It occurs in the parts of the body furnished with stiff hairs, chiefly the pubes, but also the axillæ, eyebrows, beard, eyelashes, etc. It is smaller and less elongated than the other two forms.

**Pulex irritans** (Common flea).—This animal is only partly parasitic. Its larvæ, which are about an eighth of an inch in length, occur in quantities in the neighbourhood of mouldering organic matter, in dusty corners of rooms, etc. The mature animal is not necessarily parasitic.

**Cimex lectularius** (Common bug) is still less of a parasite. It lives chiefly about beds, and comes out of retired parts on to the skin to extract blood.

**Pulex penetrans** (Sandflea, chigoe, jigger).—This is common in the West Indies, Central and South America, and southern parts of North America. The female, which resembles an ordinary flea, penetrates the skin, usually of the toes, where it swells up into a sac about the size of a pea, the abdomen being distended with ova. It produces painful inflammation.



The Larvæ of insects or Maggots are occasionally found in the tissues of man. There are a few cases in which such larvæ have, by migrating under the skin, produced considerable inflammation. There are also cases in which, deposited in neglected wounds, or even in the mouth and nostrils of excessively debilitated persons, they have actually produced considerable destruction by feeding on the tissues. In neglected military hospitals wounds are often abundantly tenanted by maggots.

The Larvæ of insects are also sometimes passed by the bowel. The form which has been chiefly observed is the *Anthomyia canicularis*. In some cases enormous numbers of the larvæ of this fly have been passed, stated as quarts in one report. (See Finlayson, in *Glas. Med. Jour.*, xxxi. 225, 1889.)

**Literature.**—The work of Leuckart which is encyclopædic in character is now available to English readers in the translation by Hoyle, vol. i., 1886. See also COBBOLD, Entozoa, 1864 and 1869, Parasites, 1879 and 1882; KÜCHENMEISTER, Syd. Soc., transl. 1857, new edition by Zürn, 1880-82; DAVINE, Traité des Entozoaires, 1877; MÉGNIN, Les parasites, 1880; BRAUN, Die thierischen Parasiten der Menschen, 1883. *Protozoa*—see LEUCKART. *Trematoda*—M'CONNELL (*Distoma sinense*), Lancet, 1885; M'GREGOR (do.), Glas. Med. Jour., 1877. *Cestoda*—fully in LEUCKART und KÜCHENMEISTER; NEISSER, Die Echinococcenkrankheit, 1877; BRAUN, Entwicklungsgesch. des breiten Bandwurms, 1883. *Nematoda*—VIRCHOW, Die Lehre von den Trichinen, 1866; LEWIS, Pathl. significance of nematode hæmatozoa, 1877; BANCROFT, Path. trans., 1878, xxix.; MANSON, *Filaria sang. hom.*, 1883, also Path. trans., 1881, xxxii.; MACKENZIE, Path. trans., 1882, xxxiii. 394; MYERS, Observations on *Filaria sang. hom.*, Shanghai, 1881 and 1886; SCHEUBE, in Volkmann's Sammlung, 1883, No. 232.



## SECTION XII.

## PYREXIA—FEVER.

**NORMAL TEMPERATURE**, *resulting from balance of production and discharge of heat, regulated by a calorific centre in the brain. Limitations of power of regulation when body exposed to excessive cold or excessive heat.* **CAUSATION AND FORMS OF PYREXIA**—*Production from lesions of nervous system. Post-mortem rise of temperature. Pyrexia from contamination of the blood, Fever proper; usually due to action of microbes; phenomena of cold stage, of fastigium, and of crisis.* **THEORIES OF PYREXIA**—*The nervous and the metabolic theories. Arguments in favour of the latter.* **OTHER PHENOMENA OF FEVER**: *cloudy swelling, increased rate of pulse and respirations, fall of blood-pressure and nervous disturbances.* **POST-MORTEM APPEARANCES IN FEVERS.**

**U**NDER the designations pyrexia and fever are included conditions of the body in which the constant phenomenon is an elevation of the temperature of the body. In order to understand this pathological change we must first consider the normal temperature of the body.

**Normal temperature.**—It is a remarkable fact that in man, as in other warm-blooded animals, the temperature of the body is maintained in all climates at a nearly constant level, there being in the healthy person merely slight daily variations. This normal temperature may be stated at about 99° Fahrenheit or 37° Centigrade. This fact implies that the production of heat in the body exactly balances the loss of heat by the body. The production of heat is by a process of combustion, and its amount will be equivalent to a certain quantity of oxygen absorbed and carbonic acid given off, just as any ordinary combustion may be expressed in these terms.

**Normal heat production.**—The seat of heat production is in the living tissues; and their vital processes mostly involve combustion. The chief sources of heat, however, are the active muscles, the secreting glands, and the nervous system, but all the active tissues of the body seem to contribute.

Many facts render it apparent that heat is produced in much greater quantity in **muscular contraction** than in the performance of any other function. We are not only sensible of a great production of heat during muscular exertion, but actual



measurement has shown that during contraction the muscles produce an increased amount of heat.

During severe muscular exertion the body temperature, as measured in the axilla or mouth, has been found elevated as much as from  $0.5^{\circ}$  to  $1^{\circ}$  Centigrade. Also, the amount of heat given off while prisoners were exercising on the treadmill, has been found to be very excessive. Again, the measurement of the temperature of contracting muscles has shown that heat was being produced. The human biceps, even before contraction, has a temperature of about  $1.5^{\circ}$  to  $2^{\circ}$  C. above that of the surrounding connective tissue, this being due to the continual tonic contraction of the muscle; but during contraction the temperature rises  $0.5^{\circ}$  to  $1^{\circ}$  C. (Breschet and Becquerel, Macalister). Bernard also found the blood of the muscular branch of the jugular vein increased in temperature during contraction of the muscles of the jaw. Then, also, there is the great rise of temperature occurring in tetanus, whether artificially produced in animals or occurring as a disease in man. In the former case, when tetanic spasm is produced by electric stimulation, the temperature of the body may rise  $5^{\circ}$  C. In man very high temperatures have been registered in tetanus. Wunderlich has found it as high as  $44.75^{\circ}$  C. ( $112.5^{\circ}$  F.), and after death  $45.4^{\circ}$  C. ( $113.6^{\circ}$  F.).

The muscular tissue of the body seems to be a constant source of heat production. The voluntary muscles are in a continuous state of tonic contraction, and the heart is constantly producing heat by its contractions. It has been calculated by Gréhart that the heat produced by the heart is about equal to a twenty-fourth of that of the whole body.

Next to the muscles the **Secreting glands** seem to contribute most to the heat of the body. Ludwig found that when the submaxillary gland is stimulated the secreted saliva is warmer than the blood of the carotid, and Bernard found the blood leaving the glands warmer than that passing to them. The intestinal glands and the liver are also great sources of heat. Bernard found the blood of the portal vein  $0.1^{\circ}$  to  $0.4^{\circ}$  C. higher than that of the aorta, and the blood of the hepatic vein  $0.2^{\circ}$  to  $0.4^{\circ}$  higher than that of the portal, showing that heat is produced first in the glands of the intestine and then in the liver. The highest temperature in the body is said to be found in the liver.

The central **Nervous system** is also a heat-producer. Observation in animals which had been artificially cooled and in hibernating marmots showed that electric stimulation of the nervous system causes rise in temperature in the brain.

The temperature of the body rises slightly after the ingestion of food. This may be due to the activity of the glands concerned in assimilation, or to the oxidation of the products of digestion in the blood.

The production of heat is approximately measured by the consumption of oxygen and elimination of carbonic acid. In the case of a cold or tepid bath, more heat must be produced in order to compensate for its extraction by the unusually cold medium. Experiment shows that under these circumstances there is an increase in the absorption of oxygen and exhalation of carbonic acid. In man the experiment cannot be long continued, and it cannot be said that the results give more than an approximate indication of quantities. The carbonic acid produced may, for example, be retained to some extent in the body and so the results vitiated.

**Discharge of heat.**—The living body is continually giving off heat in various ways, but chiefly by the skin and lungs. There is a constant radiation and evaporation from the surface of the body, while in the



lungs there is a continual cooling of the respiratory surfaces by the inspired air, and a further loss of heat by the evaporation of moisture from these surfaces. The amount of heat lost will depend on the one hand on the extent of the cutaneous surface, its temperature, the activity of perspiration, and the temperature of the surrounding media, and on the other hand on the depth of the respirations and the temperature of the inspired air.

**Regulation of temperature.**—The fact that in healthy persons the temperature remains close to a constant normal, indicates that there are arrangements in the body whereby the production and discharge of heat are regulated in their amount. These arrangements are under the control of the nervous system, which must contain a **Calorific centre**, located by some observers in or above the pons, and more particularly in the cortex of the cerebrum, about the middle of its lateral or external surface (Wood, Hitzig, Hale White). The regulation of temperature takes place chiefly by modifying the discharge of heat on the one hand, and its production on the other. That is to say, the state of the skin as regards fulness of its vessels, and activity of the sweat-glands is subject to variation according as heat requires to be economized or expended. The respiratory movements are also subject to variation, according to need. Human beings, again, assist in the regulation by clothing themselves according to the requirements of the body. By these means the central nervous system, chiefly through the vaso-motor and respiratory nerves, regulates the loss of heat. The production of heat is also to some extent controlled by the nervous system. When more heat is needed the tonicity of the muscles is increased, they become harder and more braced. Shivering is a kind of exaggerated tonicity whose rationale seems generally an increased production of heat.

**The limits of the power of regulation** are seldom reached in healthy persons. Almost any variation in the amount of heat produced is compensated by alterations in the state of the skin and respiration. Not even the severe muscular exertion of climbing a mountain, producing, as it does, an enormous excess of heat, is able to raise the temperature of the body appreciably. It is interesting to observe, however, that violent contraction of muscles produced by morbid influences will raise the temperature. It is so in convulsions such as those occurring in epilepsy, uræmia, or tetanus.

The power of regulation is not so great against variations in the external temperature. Prolonged exposure to cold, especially when the muscles are relaxed, frequently reduces the temperature. Some very low temperatures have been observed in drunkards who have lain exposed (Macewen, Reineke, Peter). On the other hand a high



temperature in the surrounding media will sometimes raise the body temperature. If the air be dry the body can meet a considerable elevation of temperature, compensation being effected by excessive perspiration, but if the air be moist as well as hot, or if a bath be used, then the temperature will rise.

As the actions of the body involve the production of heat, if the temperature of the surrounding media be even near, without quite reaching that of the body, then a prolonged exposure will raise the latter. When a rabbit is kept in a box at a temperature above  $32^{\circ}$  C. its temperature rises, so that when the air is at  $36^{\circ}$ , the temperature of the animal has risen to  $41^{\circ}$  or  $42^{\circ}$  (Rosenthal). In guinea-pigs Cohnheim observed a similar result. Again, Stapff found during the construction of the St. Gothard tunnel that men working in damp air at about  $30^{\circ}$  C. had their temperature raised to  $40^{\circ}$  C. ( $104^{\circ}$  F.).

The regulation may be rendered inefficient by some interference with the regulating apparatus. Thus, persons in a state of starvation or anæmia are not in a condition to increase the production of heat so readily as others, and hence they are less able to stand cold. Again, interference with the vaso-motor nerves may hinder the compensation effected through them. Thus, the division of the sympathetic nerves in the neck of rabbits, by paralysing the vaso-motor nerves of the ears, produced an active congestion in these parts, which lowered the temperature two degrees C. Animals paralysed by curare or morphia are more readily cooled than normal ones, both because the muscles are relaxed and because the cutaneous vessels are dilated. **Varnishing the bodies** of rabbits (with tar, linseed oil, collodion, etc.), causes a hyperæmia of the skin and thus produces a lowering of temperature which may be fatal. Similarly, extensive burns, by producing an active congestion of the skin, lead to a lowering of temperature. In both cases, if precautions are taken against cooling, the temperature is not reduced. Hence the treatment of burns by wrapping in cotton-wool.

**Causation and forms of pyrexia.**—Among the conditions in which elevation of temperature occurs, there are two distinct classes of cases, which may indeed be used as mutually illustrative of the pathology of fever, but which in the first instance must be kept rigidly apart. There are, on the one hand, cases due to disease or injury of the central nervous system, and, on the other hand, cases in which a poison of some sort is present in the blood. It is to the latter class of cases that the term *Fever* is usually applied, this term including other symptoms besides mere elevation of the temperature.

**Post-mortem rise of temperature** is not infrequently observed, especially when the temperature has been above normal at the time of death. This is explained by the fact that the tissues do not all die simultaneously, and that there may be considerable heat production after the heart has ceased to contract. The cessation of the circulation in the skin will greatly reduce the loss of heat by radiation, and the internal heat production may temporarily counterbalance the loss. A similar explanation is given by Cohnheim of the rise of temperature in the collapse stage of cholera. Here coldness of the surface is associated with rise of internal temperature. The former is ascribed to defective circulation in the cutaneous vessels, due to the thickening of the blood which results from the loss of fluid by the bowels, which is characteristic of cholera.



There is sometimes a rise of temperature **just before death** in diseases of the central nervous system, as first pointed out by Wunderlich. This is sometimes associated with convulsions or spasms, but may occur without these, and is difficult of explanation.

**Pyrexia from lesions of the central nervous system.**—There are many cases on record in which injuries or diseases of the brain have led to elevation of temperature, sometimes to a high degree. These clinical observations have been confirmed by physiological experiment. The clinical cases consist of various lesions of the brain and cord, such as tumours, injuries, hæmorrhages, etc. The experiments were generally such as separate the medulla oblongata from the pons.

Dr. Hale White has made a very admirable collection of cases of pyrexia from disease of the nerve centres. He classifies the cases into twelve groups, which include tumours of the brain and cord, hæmorrhages, especially of the pons, embolism, ill-defined degenerations, insular necrosis, locomotor ataxia, obscure nervous cases (including hysterical pyrexia), mental disease, injuries to the spine and brain. He endeavours to give unity to the cases on the ground that in all of them there is interruption of nerve-fibres passing from a supposed centre in the motor region of the cortex of the brain.

The experiments of Wood of Philadelphia are most important in regard to the production of pyrexia from injury to the brain and cord. A section at the junction of the pons and medulla, if made in such a way as not to injure the vaso-motor centres in the medulla oblongata, causes a rise in the temperature of the animal.

Both Hale White and Wood found that injuries or diseases of the central nervous system sometimes lead to a fall of temperature. Such a result seems always referable to a vaso-motor paralysis of the vessels of the skin, the congestion of these vessels causing an undue cooling of the body.

From these observations an endeavour has been made to establish the existence of a calorific or thermic centre in the brain, that is to say, of a centre which, by its own action, is capable of producing heat. We have seen that the nervous system has arrangements for the regulation of the temperature, but there is not sufficient evidence that there is any single centre having a direct control of the process of heat production. The lesions which lead to elevation of temperature are such as to produce complex derangement of the vaso-motor and muscular functions, and it cannot be said that these have been sufficiently eliminated as to prove the existence of a centre which produces its effects apart from the ordinary processes in the muscles, blood-vessels, etc.

According to the views of Wood, which are partly adopted by Hale White, the heat-producing tissues have a continual tendency to produce too much heat, and the thermal centre is chiefly exercised in controlling or inhibiting the process. As the nervous lesions which cause a rise in temperature are chiefly such as paralyze or divide the nervous connections, it is supposed that they act by removing the inhibition of the thermal centre.



Macalister has endeavoured to separate a thermogenic or heat-producing function in muscle from its contractile power, and to refer pyrexia to a stimulation of the former.

**Pyrexia from contamination of the blood. Fever proper.**—In the great majority of cases pyrexia is produced by the existence in the blood of abnormal matters. These are most commonly the products of the action of microbes, but the microbes themselves need not enter the blood; it is sufficient that the morbid products be present there. Thus putrid matter injected into the blood gives rise to fever, but it does so when all solid particles have been removed, and only the dissolved products used. In the case of putrid wounds or inflammations, we commonly have fever, but there is not usually any actual propagation of bacteria in the blood. In tuberculosis also, fever is usually present, but it seems doubtful whether the tubercular bacillus is at all capable of multiplication in the blood.

But fever may be produced without the agency of microbes. It has been induced by the injection of small quantities of water containing granules of starch or charcoal. These particles having caused obstruction of the pulmonary capillaries, the blood shut off from the circulation undergoes metamorphosis, and its products, being absorbed, cause pyrexia. The products of metamorphosis of the blood, produced in other ways, may lead to fever. Thus the injection of large quantities of pure water, apparently by causing solution of the red corpuscles, leads to elevation of temperature (Billroth and others). Even an extravasation of blood in the tissues and the absorption of its products may lead to pyrexia. Thus Volkmann found that in 14 cases of simple fracture of the femur, fever was present in 11 cases, in 5 it lasted for several days, in one as long as ten.

These facts have surely been altogether overlooked by MacLagan in a very ingenious theory of fever, which makes it depend on the multiplication of micro-organisms in the blood.

Fevers have been divided into three stages, namely, rigor or cold stage, fastigium or acme, and crisis.

During the **Cold stage** there is a marked feeling of cold, and the skin is cold to the touch, and pale or livid in appearance. The feeling of cold is actually due to a reduction in the temperature of the skin, and the shivering which is often pronounced in this stage is a reflex phenomenon, just like ordinary shivering from cold. The coldness of the surface is due to a general spasm of the cutaneous arteries.

While the surface is cold there is a great rise in the internal temperature. This rise may be partly the result of the diminished loss of heat from the surface, but is not entirely so. The rise is too great to



be accounted for in this way. For instance, Liebermeister found that in the cold stage of intermittent fever the temperature in the rectum rose in thirty minutes as much as  $2.31^{\circ}\text{C}$ . ( $4^{\circ}\text{F}$ ). This author also determined that in the cold stage there is a great increase in the process of combustion, as evidenced by the elimination of carbonic acid. The very rapid rise in temperature is therefore due to increased production, with diminished discharge of heat.

**The Fastigium** is characterized by a more or less continued elevation of temperature, which may last for days and weeks and keep near that attained at the end of the cold stage. There is great increase both in the production and discharge of heat. The former is evidenced by an increased absorption of oxygen and discharge of carbonic acid, and the latter has been determined by actual observation. The hot skin of the fever patient is generally a sufficient indication of excessive discharge of heat, but Leyden has demonstrated it by experiment, in which the leg was put into a bath, and the loss of heat measured by the rise in temperature in the water.

As the fever patient eats less than a healthy person, the excessive production of heat takes place, to a large extent, by the combustion of the tissues. Hence, as the fever progresses, there is great **Wasting of the tissues**, both of the adipose tissue and the proper nitrogenous tissues. The consumption of the nitrogenous tissues is expressed by the appearance in the urine of an **excess of urea**, which is the chief ultimate product of the metabolism of the nitrogenous principles in the body.

Under ordinary circumstances the **amount of urea** and other nitrogenous substances in the urine bears a close relation to the diet, being greatly diminished during fasting; hence the excretion of urea in fever can only be appreciated by comparing it with that of a healthy person on the same diet. A young healthy man on ordinary febrile diet will excrete 16 to 18 grammes (245 to 275 grains) of urea, while a similar person in fever will excrete 40 to 45 and even 50 grammes. The excess of urea has been stated by Unruh at 50 per cent., by Liebermeister at 70, and by Senator at 100, that is at double the normal amount.

It is a point of great interest that the increase of urea begins in some cases before the rise of temperature (Sidney Ringer and others). This has been observed chiefly in relapsing fever, and indicates a period in which the fever is latent. There is also usually an excess of urea for the first two days or so of convalescence, the "epicritical" excess. This is, in some cases, due to a retention of urea, whose amount sometimes shows a diminution towards the crisis of the fever, especially in cases characterized by the so-called 'typhoid state.'

Besides the increase of urea, there is an increase of the so-called extractives, which are nitrogenous principles of various kinds. The colouring matter is greatly increased. The soda salts, and more particularly the chloride, are diminished, while the potash ones are much increased. The phosphates and sulphates are increased.

**The State of the skin** during the fastigium is worthy of special attention, especially as it forms an important item in the means of



regulating the temperature of the body in health. The condition varies considerably in different fevers, and even in the same case, from time to time. In a few forms (chiefly acute rheumatism) there is usually profuse perspiration, but as a general rule, the skin is, considering the temperature, remarkably dry. Even apart from this, however, the condition of the skin in regard to blood supply varies greatly, and its temperature also varies. Hence the temperature of the skin does not bear a close relationship to that of internal organs.

**The Crisis or Termination** of fever is often more or less abrupt. It is as if the regulation of the temperature had been re-established in its normal condition, and the temperature rapidly falls to the normal. This is often accompanied by an attack of sweating, this part of the apparatus for regulating the temperature, which we have seen to be disordered, being restored to action. The other secretions are also restored at the crisis, the salivary, gastric, etc. It is remarkable how a temperature which has been for days or weeks persistently above normal, will suddenly and definitely fall, and with this all the symptoms at once improve. Sometimes, however, there is a more gradual fall of temperature, and instead of a crisis we have a **Lysis**.

**Theories of pyrexia.**—The very striking phenomena described above have been somewhat variously explained. In order that the different theories may be understood, let us remember the principal facts in regard to the phenomena. The rise in temperature is due directly to an abnormal combustion in the tissues, but the actual amount of heat-production is not greater than what frequently obtains in health without any rise in the temperature. The mode of heat-production, however, is abnormal, implying a pathological metabolism in the tissues, but there is something abnormal also in the regulation of the temperature, as the body does not dispose of an amount of heat which it is capable of disposing of under normal circumstances.

It is acknowledged in all modern theories of fever that the abnormal heat-production is in the tissues, and is the result of increased tissue-change, and it is agreed that the regulating process by the nervous system is altered, but opinions differ in regard to the exact place which the nervous system takes in the matter. According to one view pyrexia is essentially due to the action of the nervous system, alterations in the heat-centre inducing the increased tissue-change, and at the same time changing the regulatory process. The view opposed to this is that the increased heat-production is due directly to the action of the contaminated blood on the tissues, the abnormal constituents in the former inducing increased chemical change in the latter. The altered regulation is also regarded as related to the state of the blood.



**The Nervous theories of fever.**—The various theories which trace pyrexia to a nervous origin are chiefly based on the facts already indicated, that injuries to the brain, whether produced experimentally in animals or accidentally in man, have been known to cause a rise in temperature, sometimes to a very high degree.

The view of Liebermeister, adopted in this country by Hilton Fagge, is that in fever the high temperature depends on a change in the normal function of heat-regulation, according to which the balance of heat-production and discharge is so arranged as that the temperature is maintained at a higher level. The regulatory apparatus is at work, but it has pitched its normal at a higher level. The object of this change is a curative one. The high temperature has an influence in freeing the blood of the abnormal constituents which we have seen to be present in fever.

The view of Wood, Hale White, D. Macalister, and others is almost the converse of this. According to them the heat-centre is paralysed by the fever-producing agent. This centre when in normal action, as we have seen, is supposed to restrain or inhibit the production of heat, and when paralysed it allows of an over-production.

**The Metabolic theory of pyrexia.**—According to this theory the abnormal production of heat is due to the direct action of the fever-producing agents on the living tissues, while the due regulation of the temperature is interfered with.

To the author there are insuperable difficulties in accepting the purely nervous theory of fever. Pyrexia is produced, as we have seen, by a large number of different agents, each of which when present in the blood produces a rise of temperature. They do so also, up to a certain point, in proportion to the amount of the agent present in the blood. It seems inconsistent that such different agents should act in a similar fashion on the nervous system.

Besides, the production of heat is, apparently, by a process distinct from normal heat-production. In the normal production of heat in muscles, contraction is a constant if not a necessary element, and the production of heat in glands and elsewhere is associated with the performance of their function. In fever, however, the muscles are relaxed and the glands are to a large extent deprived of their function, and the production of heat is due to a destructive combustion of the tissues, and is thus abnormal in its method. It seems more probable that a morbid agent in the blood directly induces this change in the tissues, than that it should be due to a nervous influence.

This also would give us the key to the paralysis of the regulatory apparatus. This apparatus has relations on the one hand with the heat-producing functions, and on the other hand with the heat-discharg-



ing, and it is by the mutual regulation of these that the normal balance is maintained. But if heat-production proceeds from an extraneous cause, and is therefore placed outside the influence of the regulating centre, then the latter may reasonably be expected to be at fault. In addition, the fever-producing agent acts on the nervous centres as well as on other organs and produces a certain paralysis of their functions, just as it paralyses secretion and muscular contraction. In most fevers the cerebral functions are abnormal, although the form of disturbance varies considerably in the different fevers.

It may be a question to what extent the increased metabolism of the tissues is a reaction against the morbid agent in the blood. In this sense the rise in temperature may possibly be related to the elimination of the morbid agent, although the view that the rise in temperature in itself inhibits the morbid poison can scarcely be sustained.

**Other phenomena of fever.**—Most of the remaining phenomena of fever are to be brought into relation either with the rise in temperature or the direct action of the fever-producing agent.

**Parenchymatous degeneration or Cloudy swelling** is a frequent change in the tissues in fever. It affects chiefly the muscles and the secreting glands. Sometimes there is great enlargement of the liver and kidneys from this cause. The weakness of the heart, which is so marked in many fevers and may give rise to dilatation of its cavities, is, partly at least, due to this. The parenchymatous change has been ascribed to the action of the over-heated blood (Liebermeister, Wickham Legg), but this explanation is not sufficient, as it may be absent in cases where the temperature has been high (in cases of acute pneumonia) and is sometimes present without fever (Cohnheim). The action of the altered blood must be taken along with the high temperature.

The **increased rate of the pulse** is generally referred directly to the action of the rise in temperature on the heart. It is known by experiment that when the heart is artificially raised in temperature, by heating the blood which is passing to it or by increasing the temperature of the surrounding air after the heart has been exposed, it beats at an accelerated rate. The **increased frequency of respiration** is also ascribed to the elevation of temperature of the blood. Goldstein, by heating the blood in the carotid so that the temperature in the medulla oblongata was raised, caused great acceleration of the respiration.

**Fall of blood-pressure** is a usual concomitant of fever, although it is not always present. It is present as evidenced by **Dicrotism** of the pulse in typhoid, septic, and puerperal fevers, but absent in the eruptive stages of scarlet fever and small-pox (Recklinghausen). It is to be ascribed to a general relaxation of the arteries due to a paralysis of the



muscular coat similar to that of the voluntary muscles, but also to some extent to the weakness of the heart.

The **nervous disturbances** in fevers vary greatly and are only in part to be ascribed to the high temperature. In relapsing fever and in the rise of temperature which sometimes occurs in simple fractures or under the antiseptic treatment of wounds, there are usually no nervous disturbances. On the other hand, typhus fever is usually accompanied by delirium; typhoid, with dulness of mind, etc.; while in children convulsions occasionally accompany fever. These facts indicate that the nervous symptoms are essentially related to the pyrogenic agent, and differ with the nature of it.

**Post-mortem appearances in fever.**—It will be inferred from what has gone before that there are no constant anatomical changes characteristic of all fevers. The most frequent are the parenchymatous changes already referred to, but even these, as we have seen, are not constant. In the acute specific fevers the blood is usually found after death imperfectly coagulated. As a consequence of this it stains the heart and vessels, and the colouring matter frequently penetrates to surrounding parts, producing frequently deep staining of the skin. The spleen is enlarged in most fevers, and it is frequently very soft, especially in typhus. Sometimes there are wedge-shaped infarctions. The liver is commonly enlarged from parenchymatous degeneration, which is frequently associated with a certain amount of fatty degeneration. The kidneys are also commonly enlarged. The muscles are liable to parenchymatous degeneration, but they are also subject to waxy degeneration, especially in typhoid fever.

Besides these general changes, many of the individual fevers have specially localized lesions, such as the affection of the intestine and mesenteric glands in typhoid fever, of the throat in scarlet fever, of the skin and mucous membranes in small-pox, and of the soft membranes of the brain in epidemic cerebro-spinal meningitis.

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## PART SECOND.—DISEASES OF THE SPECIAL ORGANS AND SYSTEMS.

### SECTION I.

#### DISEASES OF THE ORGANS OF CIRCULATION.

##### A.—THE HEART AND PERICARDIUM.

- I. CONGENITAL MALFORMATIONS, chiefly of the heart and great vessels. *Causation and forms. Cyanosis a common result.* II. COAGULA IN THE HEART; the various forms of thrombi. III. OCCLUSION AND STENOSIS OF THE CORONARY ARTERIES. *Causes. Phenomena in acute and chronic obstruction.* IV. RETROGRADE CHANGES. 1. *Atrophy, including brown atrophy,* 2. *Fatty infiltration,* 3. *Fatty degeneration,* 4. *Calcareous infiltration,* 5. *Other forms,* 6. *Injuries and rupture.* V. HYPERTROPHY. *Causes, including overstrain. Forms of hypertrophy.* VI. INFLAMMATION. 1. *Myocarditis, parenchymatous, purulent, and interstitial,* 2. *Endocarditis, acute, chronic, and ulcerative,* 3. *Pericarditis, acute, chronic, and tubercular.* VII. VALVULAR DISEASE. 1. *Insufficiency and* 2. *Stenosis of mitral,* 3. *Insufficiency and* 4. *Stenosis of aortic valves,* 5. *Valvular disease of right heart.*

##### I.—CONGENITAL MALFORMATIONS OF THE HEART.

MISPLACEMENTS of the heart are of rare occurrence, and the more important of them are merely part of a general malformation of the body. The heart may be transposed, that is to say, placed in a position on the right side of the chest corresponding with that which it normally occupies on the left. With this there is usually transposition of the viscera, but it sometimes occurs alone. Again, the heart may occupy the middle line, as it does in early foetal life. It may be placed outside the thorax altogether (*Ectopia cordis*), but in this case there are



other congenital malformations, and that of the heart only forms a part. (See p. 47.)

**Absence of the pericardium** is a rare congenital malformation, and is mostly associated with misplacement of the heart. It is not to be confused with adhesion of the pericardium and consequent obliteration of the sac which is so frequent in after-life.

**Diverticulum of pericardium** is an unusual malformation. It occurs in the form of a sac with a narrow neck, which communicates with the pericardium. When distended the sac is about the size of a pigeon's egg. (See cases by Bristowe, Path. trans., xx. 101, and by author in Catalogue of Western Infirmary Museum.)

**Malformations of the heart and great vessels.**—These for the most part represent survivals of foetal conditions, and in order to their comprehension it will be necessary here to refer briefly to the state of the heart in early foetal life, and the changes which it subsequently undergoes. The heart at an early period consists of two cavities, an auricle and a ventricle. The simple auricle receives the two venæ cavæ, and the ventricle gives origin to the common arterial trunk. The ventricle, the auricle, and the common arterial trunk subsequently undergo subdivision each into two. This separation in the *ventricle* begins near the apex; the septum gradually rises towards the base, its completion at the base being delayed after the rest of the septum has been formed. Rokitansky distinguishes two parts in the ventricular septum, namely, an anterior (*septum anterius*) which divides the orifices of the aorta and pulmonary artery, and a posterior (*septum posterius*) which comes between the two auriculo-ventricular openings. The **undefended space** (*pars membranacea*) is at the union of the anterior and posterior septa, and will be the last part to close. The common *arterial trunk* begins to show signs of division by a septum about the time that the interventricular septum is approaching the base. A septum passing from both sides of the artery meets and divides the vessel into what are subsequently the pulmonary artery and the aorta. These are so adjusted as to connect with the right and left ventricles respectively. The division of the *auricles* does not begin till the ventricular septum is nearly completed, namely, about the ninth week, and after being fully formed the septum remains partially open during the whole of intra-uterine life.

**Causation of malformations of the heart.**—A large proportion of cases of malformation are related to narrowness or **Stenosis of the pulmonary artery**. This has been variously ascribed to inflammation during foetal life, and defective formation of the parts in the foetus.

By some (Peacock, Meyer) inflammation occurring in early foetal life has been assigned as the cause of the stenosis. We shall see afterwards that inflammation



of the endocardium frequently leads to valvular lesions, which result in obstruction of the orifices. In the adult it mostly occurs in the valves of the left side of the heart, and this is usually ascribed to the fact that the systemic arteries are liable to greater variations in blood-pressure and greater strain than the pulmonary vessels. In the fœtus a much larger proportion of the circulation is dependent on the right ventricle, the abdominal aorta and umbilical arteries being fed by this ventricle. The umbilical arteries are, from their position, exposed to variations in pressure, and this may tell on the pulmonary artery at its origin.

On the other hand, Rokitansky seeks to ascribe the frequency of defect of the pulmonary artery to deficiency in the original formation of the septum dividing the primary common arterial trunk. This is probably the more correct explanation, as there are seldom traces of inflammation visible in the endocardium at birth, and, besides, the lesion is not simply one of the valves, which inflammation produces, but frequently a real narrowing or defect in the artery, as if in the division of the primary arterial stem the greater part had been monopolized by the aorta.

The stenosis of the pulmonary artery is commonly associated with defects in the septa, and these may be ascribed to a mechanical interference with the complete closure of the septa. Let us suppose that the common arterial trunk, instead of dividing in the normal way into pulmonary artery and aorta, does so imperfectly, and so there is a large aorta and a small pulmonary artery, or even an entire absence of the latter. In the case last mentioned the blood from the right ventricle, as well as that from the left, would pass into the aorta, and the constant recurrence of this passage of blood at each systole of the ventricle would **prevent the closure of the septum** at the base, and cause the aorta to take permanent origin from the right ventricle as well as from the left. On a similar principle the obstruction of the pulmonary artery will, by raising the pressure of the blood in the right auricle, interfere with the closure of the foramen ovale.

Instead of stenosis of the pulmonary artery, we may have a similar condition of the aorta. The consequence of this will be defect of the septa and alterations in the circulation, the latter differently located to those already mentioned.

**Forms of malformation.—1. Defects of the septum ventriculorum.—**As already indicated, this usually goes along with defect of the great vessels. When the stenosis, as is mostly the case, is in the pulmonary artery, it is chiefly the anterior part of the septum which is defective; while in the case of aortic stenosis it is the posterior (Rokitansky). The defect is usually in or near the **undefended space** (see Fig. 141), which in the adult is normally composed only of the two layers of the endocardium, and may even present a small aperture. The defect may be so great as that there is virtually no septum, the ventricle being composed of a single cavity, or it may present various degrees of divergence from this extreme.



2. **Defects of the septum auriculorum.**—The auricular septum normally presents up to the time of birth an aperture, the foramen ovale. The least degree of defect of this septum is the permanent or partial

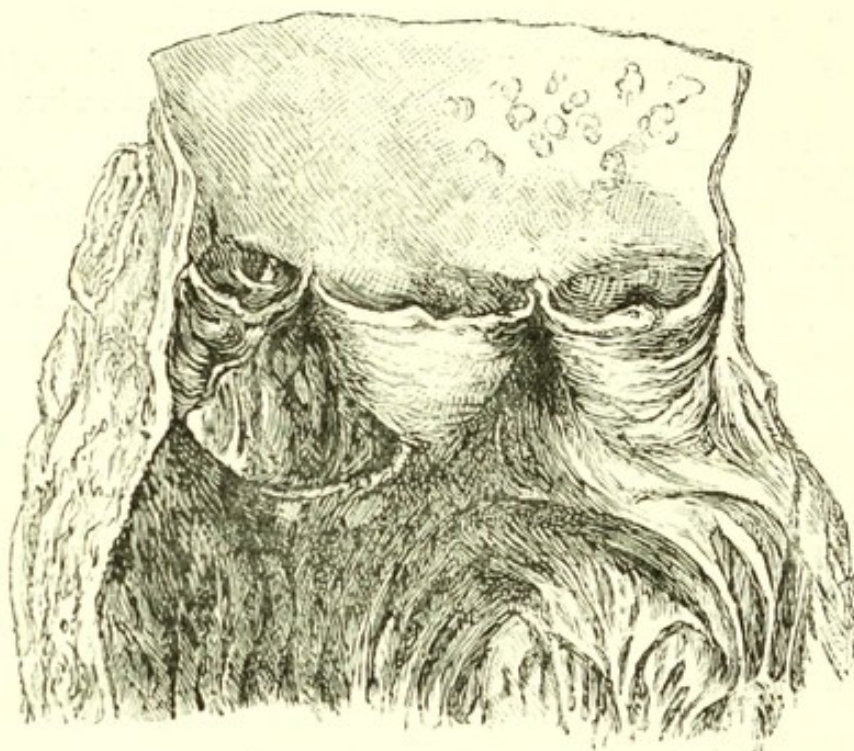


Fig. 141.—Defect of inter-ventricular septum. The gap is situated at the base, in the position of the undefended space in the normal heart. (PEACOCK.)

**Patency of the foramen ovale.** A certain degree of this is frequently present in adult hearts, but the aperture is usually so small or so valved that there is virtually no communication between the contents of the two auricles. The non-closure of the foramen may thus be a mere accident and of no pathological significance, but it may coincide with and be caused by stenosis of the pulmonary artery or aorta, and in that case there will be an actual and persistent communication between the two cavities.

In more extreme cases the septum itself will be defective. It may be entirely wanting or only defective at its lower part. This lesion frequently coincides with a defect of the posterior ventricular septum.

3. **Stenosis of the pulmonary artery.**—This may be to a slight or to an extreme degree. In most cases the ventricular and auricular septa are also defective. (Kussmaul in 192 cases found the ventricular septum complete in only 21.) The most extreme case is that in which the pulmonary artery is entirely wanting, and the aorta receives the blood from both ventricles. In this case the lungs are supplied from the aorta by means of the *Ductus arteriosus Botalli*, which remains patent, the blood passing, however, in the reverse direction to that which obtains in the fœtus. In its minor degrees stenosis of the pulmonary



artery may be associated with a complete ventricular septum, but the foramen ovale is likely to remain open. Stenosis of the pulmonary artery leads to hypertrophy of the right ventricle just as a similar obstruction does in the adult.

4. **Stenosis of the aorta.**—In this case also there is usually defect of the septa. The narrowness of the aorta may be at its origin or further on near the insertion of the ductus arteriosus. In the foetus there is a narrow part between the origin of the subclavian and the point of entrance of the ductus arteriosus called the *isthmus aortæ*, this part being scarcely in use in the foetus. The narrowness may partially persist after birth. Stenosis of the aorta, even amounting to atresia, may have little effect on the circulation during foetal life, the systemic circulation being carried on by the pulmonary artery through the ductus arteriosus. But at birth the great accession of blood from the lungs coming to the left auricle may have difficulty in getting away from the ventricle. In cases of stenosis of the *isthmus aortæ* the circulation will be carried on by the aid of anastomosis between the subclavian artery and the thoracic and abdominal aorta.

5. **Persistence of the ductus arteriosus Botalli.**—This communication between the pulmonary artery and thoracic aorta is patent up to the time of birth. After birth it becomes converted into a solid cord. It may remain patent when from any cause the circulation is abnormally directed through it. This may occur by reason of obstruction of the pulmonary artery or aorta, or from stenosis of the mitral orifice. It has also been observed without any other malformation in cases of obstruction to the pulmonary circulation from insufficient inflation of the lungs (atelectasis).

6. **Transposition of the great vessels.**—This consists in a reversal of the relative positions of pulmonary artery and aorta, so that the former takes origin from the left ventricle and the latter from the right. The blood from the systemic veins will be sent direct into the aorta without first passing through the lungs, and the right ventricle will undergo hypertrophy, and become like the left. In such cases the conditions of life are very unfavourable, and yet in one case life was prolonged for three years. Rokitansky has pointed out that even though the great vessels may be transposed, the position of the ventricular septum may be so altered as to restore the normal connections with right and left ventricles respectively.

7. **Malformations of the aortic and pulmonary valves.**—These may be part of congenital lesions of the main vessels themselves, but are frequently of independent origin. The valves may be in the form of a diaphragm in which there are merely indications of a tripartite forma-



tion (see Fig. 142). In this case the diaphragm is often protruded into the vessel in the form of a funnel. Again, we not infrequently meet with some variety in the size or number of the semilunar folds. There may be only two curtains, usually a large and a normal one, the larger one commonly showing indications of a partial division (see Fig. 143). Then with three curtains there may be two large segments and a small rudimentary one between.



Fig. 142.—Congenital adhesion of the curtains of the pulmonary valve. The valve is viewed from above. (PEACOCK.)

Many of these malformations are probably to be referred to endocarditis occurring in the foetus. In adult life endocarditis

often produces adhesion of the curtains, but there is also very great contortion of

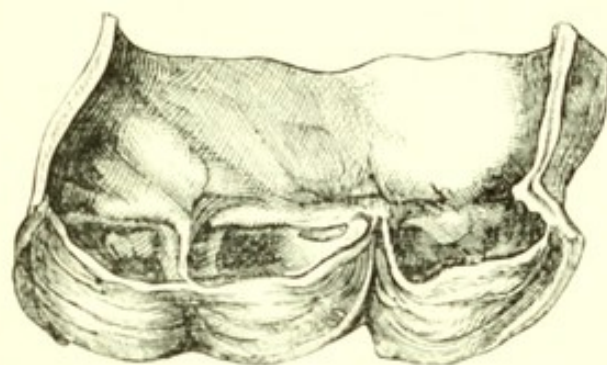


Fig. 143.—Congenital malformation of aortic valve. There are two curtains, but the larger one shows a partial division. (PEACOCK.)

the valves. But, in the foetus, the plastic power and adaptability of the structures is much greater, and the three coalesced valves may form a well-shaped diaphragm, or the two a single larger semilunar fold. In either case there are still indications of the coalescence in the form of thickenings along the line of union.

It is obvious that these malformations will frequently interfere with the functions of the

valves. In the extreme cases of complete union of the three curtains (as in Fig. 142) there will be great obstruction as well as imperfect closure. When there are only two curtains, the middle part, where coalescence has occurred, is thicker, and the curtain may be more rigid at this point, and so its function will be interfered with.

According to Peacock, congenital malformations of the valves are apt to lead to more definite disease from endocarditis in after-life. This may be a recurrence of a foetal endocarditis, but, in addition, the imperfect adaptation of the valves will itself afford a certain irritation and predispose the structures to inflammation. So it happens that valves malformed in this way are peculiarly prone to disease in after-life, even though their function is not at first imperfect.

Besides the variations already noted the curtains of the aortic and pulmonary valves are sometimes **abnormally numerous**, especially those of the latter. There may be four curtains instead of three, and they may present all varieties of size relative to one another.

The semilunar valves sometimes present a condition somewhat **approaching that of the auriculo-ventricular valves** which are of the cuspid form. It is to be noted that during the closure of a semilunar



valve the curtains do not come in contact by their free margins, but that the line of contact is somewhat removed from the margin. During closure, therefore, when the artery is full, a certain portion of the curtain floats free in the blood. This portion between the line of contact and the free margin is frequently the seat of apertures or fenestrations, and that without affecting the function of the valve.

This fenestration may be very extreme and may graduate towards a condition in which (see Fig. 144), instead of a piece of tissue, there are merely a series of tendinous bands passing from the curtain, near the line of contact, to the wall of the vessel at the point of insertion of the curtain.

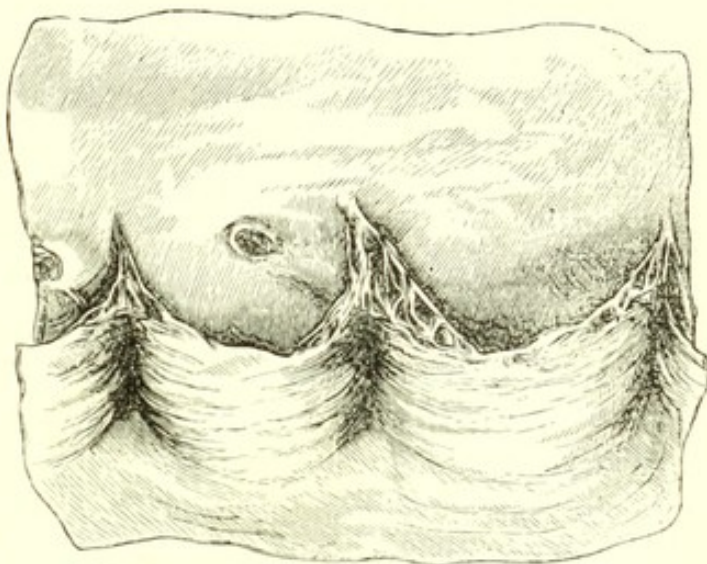


Fig. 144.—Fenestration of the marginal parts of the aortic valve. There is an appearance of chordæ tendineæ, and the bands from two proximal curtains are inserted into a projection from the aorta.

These tendinous bands resemble the chordæ tendineæ of the cuspid valves, and the resemblance may be increased by the neighbouring edges of the curtains being joined, and the tendinous bands from the adjacent borders of two curtains passing together to the wall of the vessel. There may be even on the wall of the vessel an elongated prominence into which the bands are inserted, and resembling a musculus papillaris (see figure). These bands may have a considerable free course from the middle of the curtain to the wall of the vessel. A certain amount of this condition is very frequent in both the aortic and pulmonary valves.

It may be asked, Does such a malformation interfere with the function of the valve? Probably not much, but still to a certain extent. Each form of valve is appropriate to its own place. The cuspid form is adapted to an aperture between two cavities, the semilunar to the orifice or course of a vessel. The latter takes up less room, and by reason of the complete separation of the curtains it falls back completely when the blood rushes past it. But if the valve approaches to the cuspid form, especially if the margins of the curtains are united, and the tendinous bands from two have a common insertion, then there can hardly be that complete falling back which occurs when the function is perfect. This will probably cause a trivial obstruction, and the curtains being unduly exposed to the force of the wave of blood may be specially liable to inflammation.

#### 8. Congenital malformation of the auriculo-ventricular valves.—

These valves are, like the semilunar ones, occasionally the seat of mal-



formations which usually cause narrowing of the orifice. This may be in the form of coalescence of the curtains, produced, in some cases, probably by a foetal endocarditis, so that in place of the valve there may be a more or less immobile diaphragm. The affection is observed chiefly in the right auriculo-ventricular valve. It will produce somewhat similar effects to those induced by stenosis of the pulmonary artery. There will be patency of the foramen ovale, and the blood from the venæ cavæ will, in part or wholly, pass through this foramen into the left auricle, thence into the left ventricle, and then, through a defect in the septum, partly into the right ventricle, so as to supply the pulmonary artery.

**Cyanosis.—Morbus cæruleus.**—Children born with malformations of the heart are usually liable to symptoms connected with the respiration and circulation, which are in many respects comparable with those from valvular disease acquired in after-life, and consist chiefly in attacks of dyspnœa and lividity. These symptoms may occur at birth or soon after it, but they may be postponed even for years, and develop apparently by some extra stress laid on the circulation, or they may not develop at all. Occurring at first intermittently the dyspnœa and lividity very often, to some extent, become permanent. This is particularly true of the lividity, which frequently forms such a marked characteristic of such persons that they are visibly the subjects of Cyanosis or Morbus cæruleus. The lips and finger nails are blue, and there is, perhaps, a deep lividity of hands, feet, and cheeks as well.

Two explanations of this symptom have been offered. The first is that, as, in these cases the septa of the heart are mostly imperfect, the lividity results from the mixing of the currents, venous blood being mixed with the arterial in the systemic vessels. This looks a very likely explanation, but there are two serious objections to it. Cyanosis has been found in cases where the currents did not mix, and it has been absent where they undoubtedly did. Besides this, it is found that the degree of cyanosis is not at all proportionate to the mixing of the currents. The other explanation is that the cyanosis is due to venous engorgement, just as lividity occurring in valvular disease in the adult. In nearly all the cases there is obstruction of the pulmonary artery which we have seen to be at the basis of most malformations, and this has the effect of bringing about a general venous engorgement. During the early periods of life the blood-vessels are more yielding and plastic, and so the permanent congestion tells much more on them than it does in adult life. Cyanosis developed in after-life sometimes approaches in intensity to that from malformation, but it rarely reaches it.



**Duration of life in malformations of the heart.**—If there is merely slight imperfection of the septa, the defect is of little importance, and we have already seen that there is imperfect closure of the foramen ovale in a large proportion of the cases met with in the ordinary course of post-mortem examination. If there is moderate contraction of the pulmonary artery while the heart is otherwise well formed, the right ventricle will probably hypertrophy, and this may almost completely compensate, so that life is scarcely shortened. If the foramen ovale is distinctly patent, this generally implies a greater degree of obstruction of the pulmonary artery, and life is usually abbreviated. Peacock has collected twenty cases of this kind, and only eleven of these lived to the age of 15 and upwards, but some lived as long as 34, 40, and 57. In three cases the ductus arteriosus was also open, and these died at the ages of 10 months, 15 months, and 29 years. If the interventricular septum is imperfect, this implies an obstruction at an earlier period of foetal life, and the duration of life is shorter. Of sixty-four cases, only fourteen survived the age of 15, but still three lived as long as 25, and one to 39 years. Where the pulmonary artery is entirely impervious, the duration of life is still shorter; of twenty-eight such cases only seven lived over a year, and the longest duration was 12 years. Where there is still greater arrest of development, and the heart consists of but one ventricle, with one or two auricles, the period of survival is usually very limited, but it is interesting to find that four persons thus affected have lived to the ages of 11, 16, 23, and 24 years. Transposition of the pulmonary artery and aorta might appear to be a malformation almost incompatible with life, and yet of twenty-one such cases, four lived between 2 and 3 years. When the aorta is obstructed at its isthmus, and the descending aorta is supplied, wholly or partially, by the pulmonary artery, the duration of life is usually very limited. The lungs seem to be deprived of blood, because it passes to the abdominal aorta, and the children die with symptoms of dyspnoea and syncope. If the obstruction, however, be only slight, the person may survive to adult or middle life, even though the ductus arteriosus remains pervious; thus there are cases of survival to 24, 32, and 43 years of age. If the constriction be so slight that the ductus arteriosus closes, it may yet become much more considerable afterwards, or the aorta may even be obliterated at the point indicated. Yet such patients may survive long, as even with obliteration the ages of 45, 50, and 57 have been attained.

**Literature.**—PEACOCK, *Malformations of the heart*, 2nd ed., 1861; ROKITANSKY, *Defecte der Scheidewände des Herzens*, 1875; FÖRSTER, *Missbild. des Menschen*, 1861; Many cases in *Trans. of Path. Soc., London*, by PEACOCK, GREENFIELD, FINLAY, COUPLAND, etc. *Absence of pericardium*—BALY, *Path. trans.*, iii. 60; BRISTOWE, *do.*, vi. 109; WEISSBACH, *Wien. med. Wochenschr.*, 1868.

## II.—COAGULA IN THE HEART.

**Thrombi** are of various kinds and different significance, and it may be said that coagulation of blood within the heart during life is of frequent occurrence. We have already incidentally considered most of these forms of thrombi, and it will be necessary here to do little more than enumerate them. Thrombi are frequently designated vegetations, but it is not advisable to use this word in place of the more accurate one



thrombi. We may distinguish three forms of thrombi: warty, globular, and polypoid.

**Warty thrombi** occur in acute endocarditis, owing to the coagulation of the fibrine on the inflamed and roughened surfaces.

**Globular thrombi** are of common occurrence in dilated and hypertrophied hearts, originating in the retired parts of the cavities, such as the auricular appendages, the apices of the ventricles, and behind the columnæ carneæ. (See Fig. 145). They are usually multiple, and the

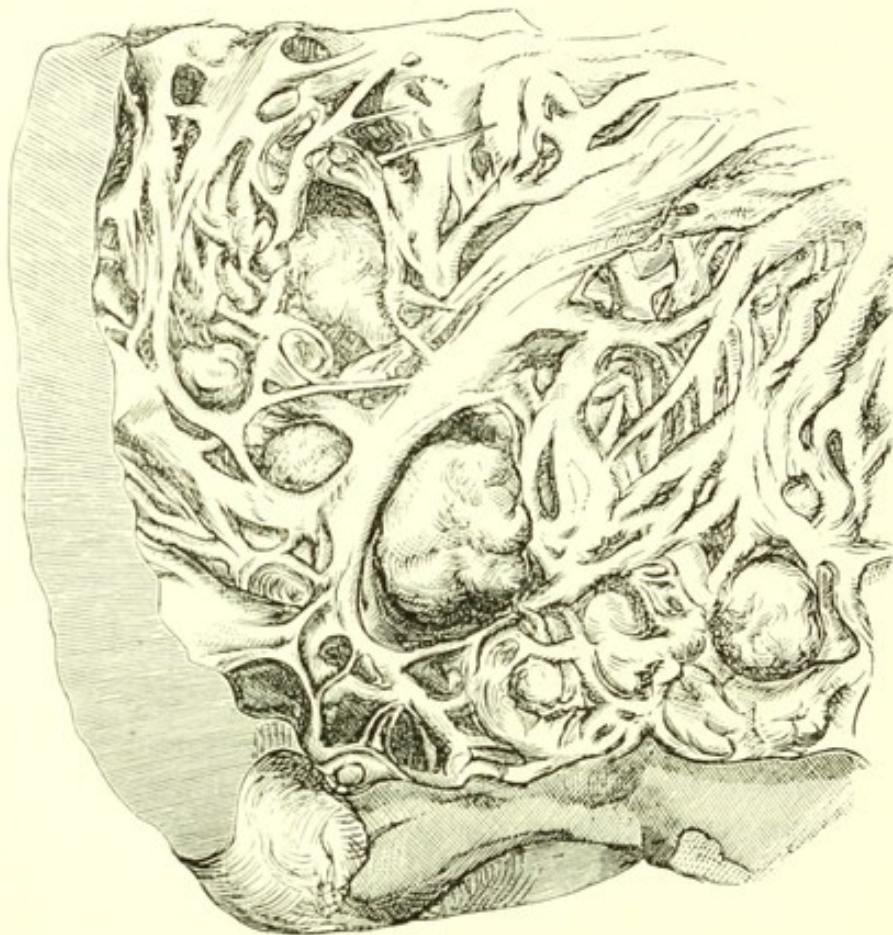


Fig. 145.—Globular thrombi near the apex of the left ventricle. Several of these are seen to project from between the muscoli papillares.

smaller of them may appear as pearly white bodies presenting a rounded projection between the trabeculæ. They may, however, grow out from these positions, and assume considerable dimensions. It is not uncommon to find them distending the auricular appendage, and sometimes filling the greater part of the auricle. The larger ones soften in the centre, forming a whitish or brownish juice, resembling pus in its naked-eye characters, but consisting merely of debris. It is quite common to find the thrombus converted into a sac, composed of a thin rind and a cavity filled with this puriform fluid. Rupture may occur during life and pieces of the thrombus may be torn off, or a thrombus may be detached bodily and carried into the pulmonary



artery or aorta so as to produce embolism. The globular thrombi are most common in the right auricle and ventricle, as these cavities are more liable to dilatation than the left, hence, embolism is more frequent in the lungs than in the systemic system. The formation of thrombi in the right cavities often coincides with thrombosis in the veins, as causes which induce dilatation of these cavities are similar to those which lead to general venous hyperæmia. Hence in a particular case of pulmonary embolism it may be a question whether the source is in the right heart or in the veins.

The **Polypoid thrombi** are much more uncommon than the other two forms. It sometimes happens that a thrombus is formed on a valve or on the internal surface of the heart and from this point grows out by successive deposition to a considerable size. The author has met with a case in which the left ventricle was filled with massive festoons thus formed, and great hypertrophy and dilatation had occurred. In this case also the coagula had undergone a partial impregnation with lime. He has also seen a case in which a thrombus, formed of firm fibrine and attached to the wall of the right auricle, hung free in the auricle and assumed a nearly globular form. It was so placed as to hang down into the tricuspid orifice, which it greatly obstructed, like a ball-valve.

In the first of the cases referred to above, the left ventricle was occupied by a large firm fibrinous mass, which consisted of twelve or thirteen polypoid coagula having a firm attachment to the anterior wall of the ventricle at about its middle. It was estimated that these coagula weighed two ounces. The coagula were obviously old and the basal parts had undergone a kind of fibrous transformation resembling a tendon, while the red part looked like the fleshy part of a muscle. There was no softening, but in the substance of the polypus and partly on its surface, a somewhat massive deposition of lime salts had occurred, forming at one place a sort of stem an inch in length. (See full account in *Glasg. Med. Jour.*, Feb., 1870. The specimen is in the museum of the Royal Infirmary, Glasgow.)

The second case is one in which a stenosis of the tricuspid valve had been diagnosed by Professor Gairdner many years before death. (See clinical account of case in *Gairdner's Clin. Med.*, 1862, p. 602. This unique specimen is in the museum of the Western Infirmary.)

**Literature.**—STEVEN and COATS, *Glasg. Med. Jour.*, Feb., 1870; WICKHAM LEGG (Loose balls of fibrine in left auricle), *Path. trans.*, xxix. 49.

### III.—OCCLUSION AND STENOSIS OF THE CORONARY ARTERIES.

The coronary arteries are frequently affected either at their orifices or in their course by lesions which interfere with the circulation through them. As these arteries are, at the most, possessed of very imperfect anastomosing communications, obstruction implies a very serious dis-



turbance in the function of the heart, amounting, in the case of a sudden obstruction of a large branch, to sudden paralysis of its action and death.

Cohnheim asserts that the coronary arteries are end-arteries. Ligature of one of the larger branches in the dog produced first irregularity and then stoppage of the heart in from thirty seconds to a minute. Wickham Legg and West believe that there are free anastomoses between the branches of the arteries, while Steven asserts that there are communications, but only among the finest arterioles. The author is convinced, from observation of Steven's injections, that the communications are of the very finest vessels, partly capillaries and partly minute arteries.

**Causes of obstruction of the coronary arteries.**—These arteries are peculiarly liable to **Atheroma**. Perhaps the fact that, coming off directly from the aorta, they are exposed to higher pressure and greater variations of pressure than other arteries of their size, may account for this. Atheroma, as explained further on, leads to narrowing of the calibre of the artery, and it often induces **Thrombosis** which may increase the stenosis or even lead to occlusion. This is most common where calcareous infiltration follows atheroma. After a prolonged stenosis there may be a sudden complete occlusion. Again, atheroma sometimes leads to a small aneurysm of the artery. **Atheroma in the aorta** leads not infrequently to **Obstruction of the orifices of the coronary arteries**. The aorta is the most frequent seat of atheroma, and as this condition leads to thickening of the internal coat there will sometimes be a bulging of this coat over the orifices of the coronary arteries. Not infrequently the prominent intima around the orifice coalesces and completely covers the aperture.

**Embolism** is also liable to occur in the coronary arteries. It used to be stated that during the systole of the ventricle the coronary arteries were closed by the curtains of the aortic valve falling against them. It is stated by recent observers, however, that this is not the case, and that the orifices are exposed during the systole, and hence, in cases of acute endocarditis, small portions of thrombi are liable to pass into the coronary arteries. The arteries obstructed will nearly always be small, and the obstruction is often multiple. In ulcerative endocarditis and pyæmia there may be septic embolism and the formation of abscesses.

**Effects of obstruction of the coronary arteries.**—The results vary somewhat according to the suddenness of the obstruction, the size of the vessel, and otherwise.

**Sudden obstruction of a considerable branch**, usually brought about by thrombosis occurring in consequence of atheroma, leads to **Infarction of the heart**. The affected area is usually in the wall of the left ventricle, the artery most frequently obstructed being the descending



branch of the left coronary artery. If death occurs immediately after the obstruction the part will be found of normal consistence, but pale yellow in colour. Very soon it becomes soft and yellowish white or brown in colour, and the part is depressed below the surface. In some cases the part becomes almost fluid. This condition of softening, which has been called by Ziegler **Myomalacia cordis**, may be associated with hæmorrhage, and thus assume the usual features of the hæmorrhagic infarction. Under the microscope the muscular fibre is found more or less broken up, its transverse striæ have disappeared, and the fibres have assumed a hyaline or waxy appearance (waxy degeneration or coagulation-necrosis). There is often, at least in the peripheral part, a great infiltration of round cells, from inflammatory reaction.

The softened portion of the wall may give way before the pressure of the blood, and the result may be an **Acute aneurysm** of the heart or even **Rupture**. The condition under review is probably the most frequent cause of rupture of the organ.

If the patch of softening be small, then through time the muscular tissue is absorbed and the connective tissue is increased in the way to be described further on.

A more **Chronic obstruction** of a considerable branch or a sudden obstruction of a smaller branch, frequently leads to a condition which has been designated **Fibrous transformation**, or (less happily) fibroid degeneration. This condition is really that present in most cases of so-called **Interstitial myocarditis**. The gradual or more sudden deprivation of blood causes atrophy and degeneration of the muscular fibres, and the connective tissue comes to form the chief or entire constituent of the heart-wall, generally reinforced by a certain newformation due to chronic inflammation. The patches of fibrous transformation vary greatly in size according to the artery obstructed. They may be simply small tendinous areas in the midst of the muscle (see Fig. 146), or they may affect extensive tracts. The tendinous or cicatricial appearance is sometimes visible in the muscoli papillares when the ventricles are laid open, but it may only be discovered by slicing the muscular tissue, which is best done by sections parallel to the surface of the heart. Where an extensive area has undergone this transformation the wall of

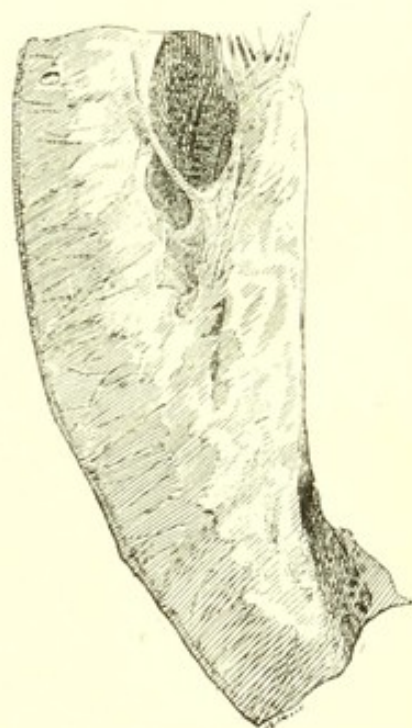


Fig. 146.—Section of left ventricle and a musculus papillaris showing fibrous transformation.



the heart may be bulged outwards, thus forming a **Chronic aneurysm** of the heart. Under the microscope the fibrous patch consists chiefly of wavy connective tissue, with very little appearance of inflammation. There may be no muscular tissue in the midst of the patch, but sometimes atrophied fibres may be visible (see Fig. 147).

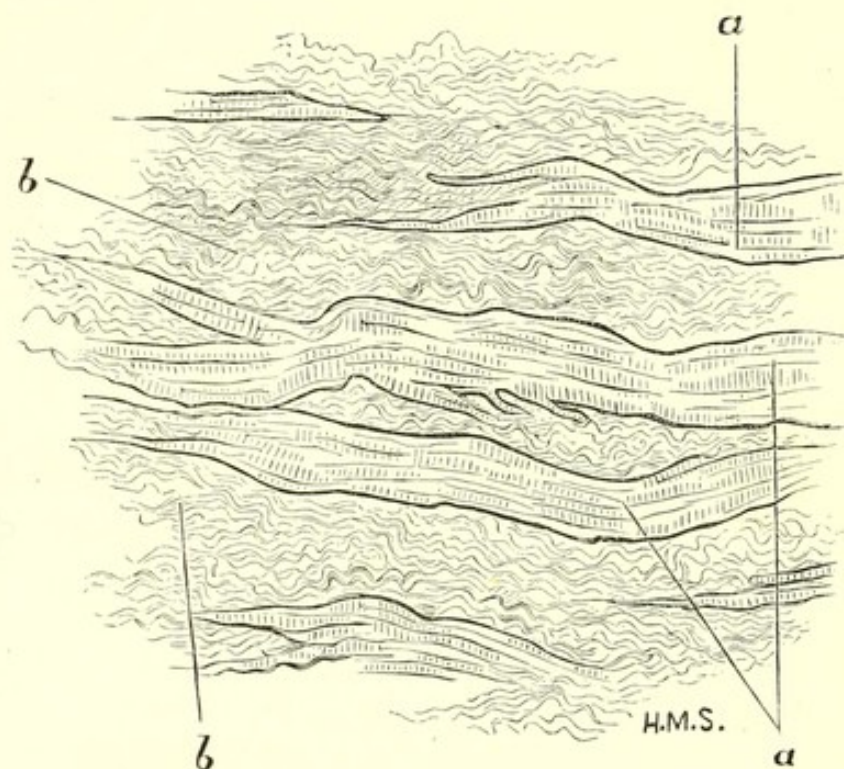


Fig. 147.—From a fibrous patch in heart. *a a*, Muscular fibres; *b b*, connective tissue. (STEVEN.)

**Fatty degeneration** is also a result of obstruction of the coronary arteries. This is most pronounced in cases of obliteration of the orifice of one or both arteries by atheroma of the aorta.

In a case observed by the author where one orifice was completely occluded and the other somewhat narrowed, there was generalized and typical fatty degeneration, such as one sees in pernicious anæmia. There was here a slowly advancing deprivation of blood, which only proved fatal when very extreme. Greenfield has recorded a case in which both arteries were occluded, and in which he suggested that the heart might be partially nourished by imbibition.

It is clear that during life there will often be, in consequence of obstruction of the coronary arteries, serious functional disturbance of the heart. There may be urgent dyspnoea, pain and irregularity of the heart, leading up, it may be, to death more or less suddenly. **Angina pectoris** is a frequent feature in the cases of considerable interference with the circulation.

**Literature.**—COHNHEIM, Virch. Arch., lxxxv. 503; WICKHAM LEGG, On cardiac aneurisms, Bradshaw lect., 1884; WEST, Path. trans., xxxv. 110; QUAIN, On fatty heart, 1851; WEIGERT, Virch. Arch., lxxix., 1880; HUBER, do., lxxxix., 1882; TURNER,



Trans. Med. Congress, London, 1881, i. 427; ROBIN, *Gaz. Hebdomadaire*, 1885, No. 51; PAUL, *do.*, No. 10; ZIEGLER, *Handbook*, and *Deutsch. Arch. f. klin. Med.*, xxv.; HILTON FAGGE, *Medicine*, ii. 32; STEVEN (with full literature), *Lancet*, Dec., 1887.

#### IV.—RETROGRADE CHANGES IN THE HEART.

1. **Atrophy of the heart.—Brown atrophy.**—This condition is one of comparatively frequent occurrence, but is for the most part merely a part of general atrophy, or emaciation of the body. In emaciating diseases where the muscular system as a whole has undergone great reduction in bulk, the heart is found to take part in the same process. Taking the normal weight of the heart as 9 ounces for the female, and 10 or 11 ounces for the male, we may find it reduced to 6 or even 5 ounces. Viewed as a whole the heart is obviously smaller, and it has a darker colour than normal, while the coronary arteries stand out unduly, often as somewhat prominent tubes. This change of colour and the prominence of the arteries are largely due to the loss of the sub-pericardial fat, which normally covers the greater part of the surface and accompanies the coronary arteries, partially embedding them in the adipose tissue.

The muscular substance when incised is found to be deeper in colour and tougher than usual. On account of this the name of **Brown atrophy** is given. With the atrophy of the muscular fibres there is an accumulation of pigment granules around the nuclei. In nearly all old persons there is at the two poles of each nucleus a group of brown pigment granules, and these increase greatly. There is also frequently a distinct demarcation of the cells which constitute the muscular cylinders, an appearance which is not normally present. This has, without sufficient grounds, been interpreted as an indication of disintegration of the muscle. Brown atrophy may be associated with fatty degeneration, the brown pigment being distinguishable by its colour from the fat granules.

2. **Fatty infiltration of the heart.**—The normal heart is well known to present on its surface a certain amount of adipose tissue. This fat is beneath the pericardium, lying between it and the muscular substance of the wall. It is normally most abundant along the course of the coronary arteries, along the inferior border of the right ventricle, at the apex and at the origins of the great vessels. In different individuals the amount of the adipose tissue and the extent to which it covers the muscular substance so as to conceal it from view, vary greatly, but it may be said generally that a considerable part of the surface of the right ventricle and the greater part of that of the left are usually free of fat. Sometimes this fat increases greatly, both in super-



ficial area and in thickness, so that the entire right ventricle may be covered while a portion of the left is still free ; or, the whole heart may be coated with a thick mantle of fat.

The adipose tissue does not always confine itself to the pericardium, but frequently extends into the connective tissue lying between the muscular fasciculi in the proper muscular wall of the heart. The superficial layers of the muscular wall may thus be largely replaced by adipose tissue, which may even appear in isolated patches immediately beneath the endocardium. It is not uncommon to find the proper muscular substance of the right ventricle largely replaced by adipose tissue, only a thin layer of red muscle appearing inside the thick layer of fat, and even this interrupted by areas of adipose tissue. Of course, in this case, there is great loss of the muscular power of the heart, the right ventricle being more affected than the left.

In some cases the increase of the external fat is merely part of a general obesity, in which the fat in all its various localities throughout the body takes part. But sometimes its significance is much more serious, and this applies especially to the cases in which adipose tissue forms between the muscular bundles. The space occupied by the fat must be obtained at the expense of the proper muscular substance, and the question arises whether the atrophy of the muscle is the primary condition or the fatty infiltration. We have to take into consideration the fact that a fatty infiltration of an exactly similar character occurs, as we have already seen, in voluntary muscle, and is there associated with disuse of the muscle. In this case the loss of function is the primary condition and the fatty infiltration is secondary. And so in the case of the heart, we meet with fatty infiltration in cases where there is no general obesity, often in old debilitated persons, or even in those who have been subject to some emaciating disease such as cancer. In that case we may infer that the weakening and atrophy of the muscle has been primary and the infiltration of fat secondary.

**3. Fatty degeneration.**—This condition is of exceedingly frequent occurrence in the heart, especially in its minor degrees. Any disease which causes a serious deterioration of the blood will produce it, and it is seen in its most pronounced form in the various kinds of anæmia, in leukæmia, and in the acute fevers. It is also brought about in a very pronounced form in poisoning by phosphorus and arsenic. In a minor degree it is seen in debilitating diseases, and is often the more immediate cause of death in hypertrophy of the heart.

It is remarkable that, though the determining cause is a deteriorated state of the blood, yet the fatty degeneration frequently shows itself in patches, so that the muscular tissue is seen to be, as it were, flecked with pale spots or streaks. This is best seen on examining the muscular tissue from within (see Fig. 148), as the endocardium generally produces but little obscuration of these markings, and they are most



abundant in the inner layers of the muscular substance. This flecked appearance is not always present, and it would be a mistake to infer the absence of fatty degeneration from its absence. The heart again is generally flabby, friable, and pale in colour, but a very flabby heart may be very little fatty, and fatty degeneration may co-exist with a comparatively firm muscular tissue. Microscopic examination should therefore always be resorted to.

Under the microscope in those cases in which the degeneration is in patches, the transparent muscular tissue is seen with a low power to be interrupted by opaque patches, as in Fig. 149. The general outline of the muscular cylinders is preserved, but they are evidently replaced by some foreign material. Under a high power, as in Fig. 150, the individual fat drops become apparent. These fat granules are frequently seen to be in rows, representing the original muscular fibrillæ, and the contractile substance is obviously lost or converted into oil. In minor degrees the degeneration is, as a rule, more uniformly distributed in the muscular substance, and we can see that the fat granules generally appear first in the neighbour-

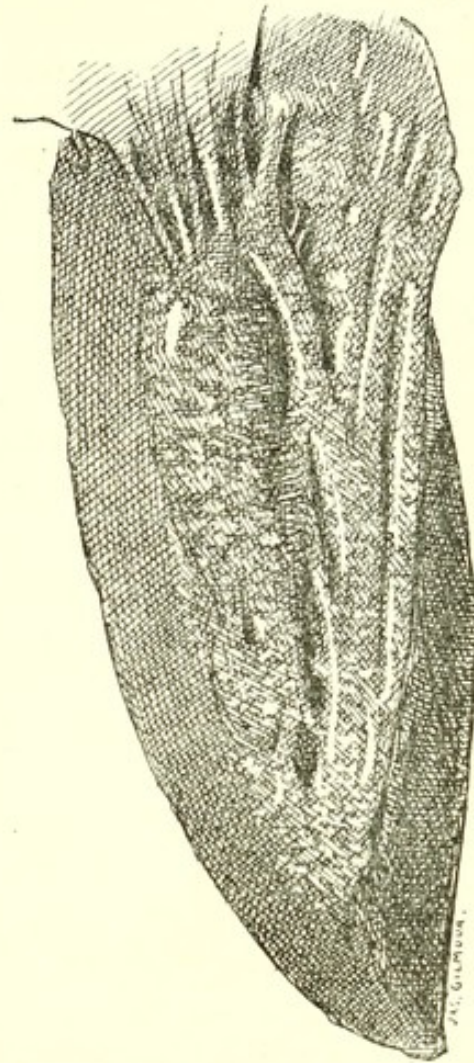


Fig. 148.—Fatty degeneration of the heart. The muscle flecked with light spots.

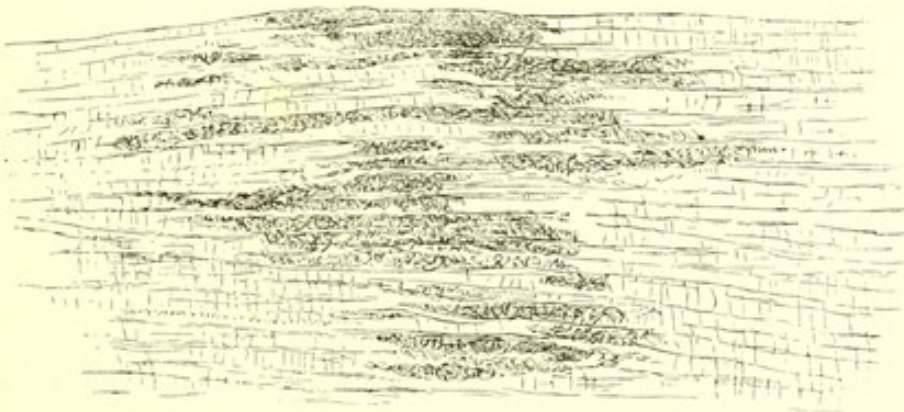


Fig. 149.—Fatty degeneration of the heart. The occurrence of the lesion in patches is indicated by the dark appearance of the fibres.  $\times 90$ .

hood of the nuclei of the muscular fibres forming elongated collections extending from either pole of the nucleus.



It is clear that muscular fibres which have undergone this transformation in its extreme form, are incapable of recovering their original condition. The sarcous substance has undergone transformation. It is to be inferred, therefore, that when such a heart recovers there is an absorption of the fat and

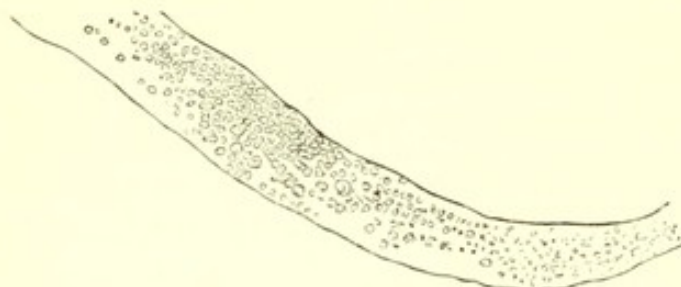


Fig. 150.—Fatty degeneration of the heart. A single fibre with oil drops in it, as seen in the fresh state.  $\times 350$ .

an actual newformation of muscular tissue. This process must be a comparatively frequent one when we consider how common fatty degeneration is in severe anæmias and in acute fevers. The fatty heart is usually a dilated one, even where there is no dilatation to begin with. On recovery the ventricles will resume their former dimensions.

**4. Calcareous infiltration of the muscular substance of the heart.**—In this place we have to consider calcareous impregnation of the muscular substance of the heart, which is of rare occurrence.

It is necessary, in the first place, to refer to certain other forms of calcareous impregnation in order to clear the ground for that more immediately before us. In connection with pericarditis it is not very uncommon to meet with calcareous infiltration of old fibrine or dried-in pus which may remain on the surface. Again, where, in the pericardium, there has been great newformation of dense connective tissue from prolonged pericarditis, lime salts may be deposited in the hard tissue. In this way we may have the heart almost enclosed in a firm shell. But, further, in the muscular substance, old connective tissue may calcify, or an abscess may dry-in and become impregnated with lime salts. In this way we may have stony masses developed in the muscular wall, these being actually in the connective tissue of the wall.

The author has met with two cases of calcareous infiltration of the

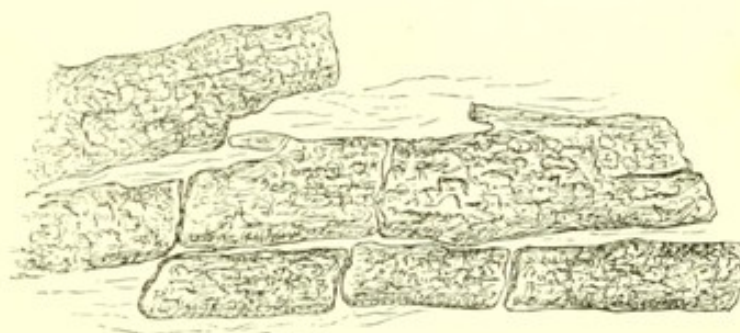


Fig. 151.—Calcareous infiltration of the muscular tissue of the heart. The branching muscular cylinders are shown in a petrified condition with transverse fractures. From a case of pyæmia.  $\times 350$ .

muscular substance of the heart, but they differed somewhat from each other. In one the lime salts were deposited in massive form converting the muscular fibres into solid cylinders, while in the other there was a fine granular deposition, causing the fibres to resemble those in fatty degeneration.



In the one there were pale patches seen with the naked eye in the muscular substance, somewhat like those of fatty degeneration, but larger and situated in the superficial layers just under the pericardium. These patches had, even to the naked eye, a streaked appearance, the streaks following the direction of the muscular fibres and indicating that the condition affected the muscular substance. On cutting into the patches a gritty feeling was experienced, and under the microscope the appearances seen in Fig. 151 were visible. The muscular fibres were converted into solid cylinders which had a markedly crystalline appearance. Many of the cylinders were fractured transversely. On adding hydrochloric acid to these patches, there was an abundant evolution of gas and a solution of the lime salts. After the lime salts were dissolved the muscular fibres were restored so far as their outline was concerned, but their transverse striæ were gone. The case in which this occurred was one of pyæmia, and it is probable that the arteries in connection with these patches had been obstructed, causing a necrosis of the portion of tissue which subsequently became impregnated with lime salts.

In the other case the condition was very different. A certain portion of the muscular substance of the left ventricle was found of a pale colour suggesting fatty degeneration, but the colour was continuous, and it was the external layers that were affected, and that mainly towards the apex. On microscopic examination the muscular fibres were found clouded with fine granules not unlike fat granules (Fig. 152). The granules, however, were dissolved by

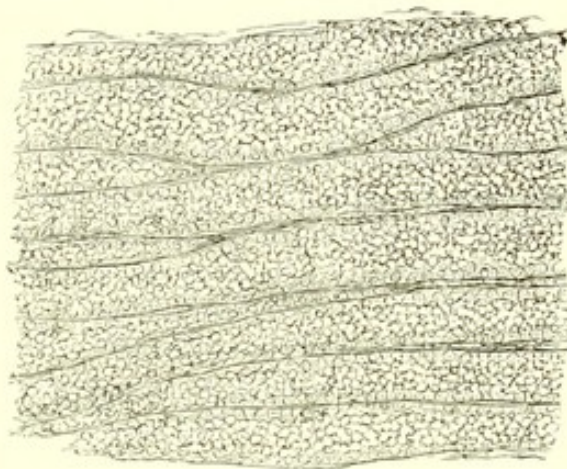


Fig. 152.—Calcareous infiltration of the muscular fibre of the heart. Fine granules occupy the muscular fibres.  $\times 300$ .

hydrochloric acid, but without evolution of gas. Köster has met with a somewhat similar case, and he believes that the salt here is, in part at least, sulphate of lime. In this case the true pathology of the condition was obscure.

**5. Other forms of degeneration.**—**Amyloid degeneration** is rare in the heart, but in extreme cases it may exist in the endocardium and in the intermuscular vessels. **Hyaline degeneration** we have seen to occur in infarction.

**6. Injuries and rupture of the heart.**—**Wounds** of the heart are not by any means necessarily fatal, although, of course, commonly so. A needle or similar small instrument may be passed into the organ without obvious injury. In most cases of penetrating wound of the heart there will be fatal hæmorrhage, but such wounds, especially if they do not divide the muscular fibres transversely, may heal, and be finally closed by connective tissue. There are even cases in which the point of a knife or a rifle ball has lodged in the heart and become surrounded by a connective tissue capsule.

**Spontaneous rupture** of the heart rarely occurs except in consequence of softening from obstruction of a coronary artery, as has been



already described. It sometimes occurs in pyæmia from the formation of an abscess in the wall of the heart. It may also result from the bursting of an aneurysm of the heart, but this also is the more remote result of disease of the coronary arteries. Rupture of the heart is sometimes ascribed to fatty degeneration of the muscular fibre, but there seems no evidence that this alone can cause it. The rupture mostly occurs in the left ventricle and near the apex or towards the aortic orifice. There is usually a large escape of blood into the sac of the pericardium, and the patient dies rapidly.

**Literature.**—*Atrophy and fatty changes*—ALBERS, *Atrophie des Herzens*; in Casper's *Wochenschr.*, 1836, 50; CHURCH (small heart weighing 3 oz. 1 dr.), *Path. trans.* xix., 1868, p. 147; R. QUAIN, *Fatty dis. of heart*, 1851; BARLOW, *Fatty degen.*, 1858; PERL (*Fatty heart in anæmia*), *Virch. Arch.*, lix. 39; also PONFICK, *Berl. med. Wochenschr.*, 1872. *Calcification*—COATS, *Glas. Med. Jour.*, Aug. 1872; also HESCHL, *Zeitschr. f. pract. Heilk.*, 1860, and ROTH, *Correspbl. f. Schweizer Aerzte*, 1884. *Amyloid degeneration*—HESCHL, *Wien. med. Wochenschr.*, 1870; KYBER, *Virch. Arch.*, lxxxi.; WILD, *Amyl. und hyalin degen. des Bindegewebes*, 1885. *Rupture*—see Schrötter, in Ziemssen's *Encycl.*, vol. vi. Various cases in *Path. trans.*, etc.; HADDON (*Rupture in pyæmia*), *Path. trans.*, xxxv. 121; ROBIN, *Gaz. hebdom.*, 1885, No. 51.

#### V.—HYPERTROPHY OF THE HEART.

This term is applied to hypertrophy of the muscular substance of the heart, giving rise according to circumstances to enlargement of particular ventricles or auricles, or of the heart as a whole.

**Causation.**—It will be found that this newformation of muscular tissue is in nearly all cases **Compensatory**, that is, intended to make up for some defect in the heart itself or in the circulation. The needs of the organism assert themselves by the nervous connections of the body on the cardiac ganglia, and the contractions become more forcible. Within the limits of health, and without any increase of the volume of the cardiac muscles, there are great variations possible in the force of the cardiac contractions. But where the heart is for a long period impelled to unusually violent exertion, it becomes hypertrophied.

In many of these conditions there is a mechanical interference with the flow of the blood either in the heart itself or in the arteries, and as a consequence the heart is overloaded with blood, but in some the mechanical cause is not very obvious. As a general rule the cavities are dilated and the dilatation may indeed be the primary condition, the hypertrophy occurring as a secondary consequence. It is usual, therefore, to consider dilatation and hypertrophy together, there being commonly some dilatation along with the hypertrophy.

**Hypertrophy from overstrain.**—**Idiopathic hypertrophy.**—**Patho-**



ogists have frequently observed that hypertrophy of the heart has existed without any mechanical hindrance in the circulation being discoverable. Some of these cases have been traced to functional disorders of the heart, the organ contracting more frequently and violently than it should. But some cases are really due to frequent and violent exercise, which has been so prolonged as to have taxed the contractile power of the heart beyond its normal powers. An acute overstrain may be produced by prolonged muscular exertion, such as in hill-climbing (Allbutt). It has been induced experimentally by Roy and Adami by narrowing the aorta. In that case dilatation of the heart, followed by incompetence of the auriculo-ventricular valve was the result.

When frequently repeated, such overstrain will lead to hypertrophy, which may only partly compensate, and there may be a permanent over-dilatation with incompetence of the valves.

This has been observed under a variety of circumstances. It has been described as occurring amongst the Cornish miners, who after their work was over had to reach the surface of the earth by climbing ladders for an hour (Peacock), or in the case of people in hilly countries who are in the habit of carrying loads uphill (Münzinger). It has been observed also in young soldiers, who as a result of severe drill, have frequently attacks of palpitation. The ability of the heart varies greatly in different individuals; in some the strain of drill reveals a weakness which has come to be recognized in the army as "irritable heart" (Maclean and Myers). During campaigns also, long marches may so overstrain the heart as to lead to dilatation and hypertrophy (Fräntzel).

As further examples of the causes of hypertrophy of the heart may be mentioned, **Adhesion of the pericardium** and **Valvular disease**, in both of which interference with the circulation exists, as will be seen further on.

Hypertrophy is a frequent result of **Obstruction to the circulation in the lungs**. In emphysema, for instance, there is great obliteration of the pulmonary vessels and the right ventricle contracts more vigorously to compensate. A similar result may follow other chronic diseases of the lung, and even extensive pleural adhesions.

Interferences with the systemic circulation, especially **Aneurysms and Rigidity of the arteries**, are frequent causes of hypertrophy of the left ventricle.

**Chronic Bright's disease** leads to hypertrophy of the left ventricle by raising the general systemic blood-pressure. (See further under Diseases of the Kidneys.)

**Forms of cardiac hypertrophy.**—From what has gone before, it will be apparent that hypertrophies of the heart vary greatly in amount and in the distribution of the enlargement. The term **General hypertrophy** is used to express an enlargement of the heart in all its parts, while



**Partial hypertrophy** expresses an enlargement limited to a part of the organ. As it is the ventricles which are specially exposed to the causes of hypertrophy the partial forms are divisible into two, namely, hypertrophy of the right and left ventricles respectively.

In **General hypertrophy** the general shape of the heart is not much altered. The heart is enlarged in all its parts, the ventricles and auricles are increased in capacity, and their walls thickened. The heart is like that of a bullock in size, so that the name *cor bovis* is often applied to it.

When the **right ventricle** is mainly affected, the heart assumes a somewhat quadrilateral form (see Fig. 153). On examining the normal heart as it lies on its posterior surface, after removal from the body, the right ventricle is seen to occupy the greater part of the anterior aspect. The normal position of the septum, as shown in the accompanying figure, is slightly to the right of the left border, and it reaches the apex region slightly to the right of the true apex. In the heart the position of the septum is always indicated on the surface by the coronary artery which, with its padding of fat, occupies a groove corresponding with the anterior border of the septum. In hypertrophy of the right ventricle, as shown by the dotted line in the figure, the apex is unduly obtuse, and it is often difficult to determine what is its exact seat. The septum is nearer the left border than usual, and it reaches the apex region rather to the left than the right of the most projecting point. The right ventricle also monopolizes the anterior aspect of the heart still

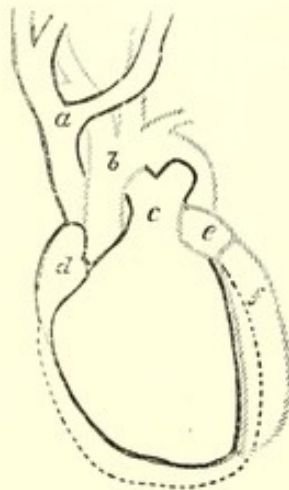


Fig. 153.—Hypertrophy of right ventricle. The alteration in shape is indicated by the dotted line; a, superior vena cava; b, aorta; c, conus arteriosus; d, right auricle; e, left auricle; f, left ventricle. (RINDFLEISCH.)

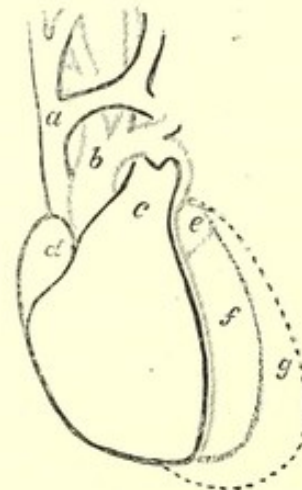


Fig. 154.—Hypertrophy of left ventricle. The alteration in shape indicated by dotted lines; g, the hypertrophied ventricle. (RINDFLEISCH.)

more than in the normal condition. When the heart is laid open, the undue thickness of the right ventricle, as well as the enlargement of its cavity, become apparent.



In **Hypertrophy of the left ventricle** the relations are very different, as shown in Fig. 154. The heart as a whole is more elongated and pointed than normal, and this is often very striking. The apex part especially appears greatly prolonged. When the heart is viewed on its anterior aspect the septum is seen to lie more to the right than is normal, and the true apex is much further to the left of the point at which the septum reaches the apex region. On laying open the heart, the thickening of the wall of the left ventricle is very obvious, and the septum is often greatly thickened. The septum belongs partly to the left and partly to the right ventricle, but as the left ventricle is much thicker than the right the septum belongs more to the left. It will partake, in the hypertrophy of either ventricle, and, in the case we are considering, the thickening is often very striking. The thickened septum frequently bulges into the right ventricle, sometimes diminishing its capacity greatly, and even producing actual obstruction of the conus

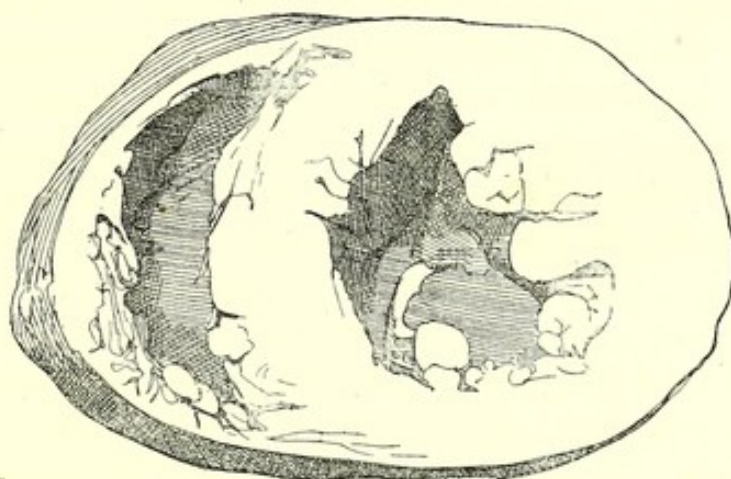


Fig. 155.—Transverse section of ventricles of heart showing hypertrophy of the left, from a case of chronic Bright's disease.

arteriosus. In transverse section the hypertrophy of the left ventricle is sometimes very manifest, the right forming merely a crescentic appendage (see Fig. 155).

The total **Increase in weight** in hypertrophy of the heart is greatest in cases where both ventricles are enlarged; and the weight of the heart in such cases not uncommonly reaches from 27 to 30 ounces. It is least where the right ventricle alone is enlarged, because this ventricle, as a whole, weighs much less than the left, but in pure right ventricular hypertrophy a weight up to 17 ounces is not infrequent. In hypertrophy of the left ventricle, as in Bright's disease, the weight is frequently over 20 ounces.

The hypertrophied heart often presents a peculiarly firm condition of its wall, and this has been ascribed by Sir William Jenner to a **Passive congestion of the heart**. Cases of cardiac hypertrophy are very frequently such as to lead to a general venous engorgement, in which the heart itself, being related to the general venous system, partakes. Now, prolonged venous hyperæmia produces in organs, as we have



seen already, a certain hypertrophy and increased density of the connective tissue. Some part of the hypertrophy in such cases may even be due to increase of the interstitial connective tissue. Hence the result is that the walls of the heart are more rigid than normal, and when the cavities are laid open they do not collapse, but stand out with their outline retained, the walls having a tough leathery character. The muscular substance also is frequently of a very red colour, this being largely due to the excess of blood in the vessels.

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## VI.—INFLAMMATIONS OF THE HEART.

The inflammations of the heart are divisible into three forms, according as the myocardium, the endocardium, or the pericardium is the primary seat. As the myocardium is closely in contact with pericardium and endocardium it will sometimes partake to some extent in their inflammations.

1. **Myocarditis.**—This name designates inflammation of the muscular substance of the heart. Several forms are distinguished.

**Parenchymatous myocarditis** is a general inflammation of the proper muscular fibre of the heart. The term is applied chiefly to cases of aggravated parenchymatous degeneration, occurring in acute infective diseases. It is met with chiefly in septic and pyæmic conditions and in diphtheria.

It must be said that this form of disease is somewhat indefinite, and in any case is secondary in its origin. There is reason to believe, however, that certain of the morbid poisons attack the heart more especially, and act directly on its muscular substance. This is true especially of diphtheria and perhaps also small-pox.

**Purulent myocarditis** is also a secondary affection, occurring in consequence of the transportation of material containing infective micro-organisms. It is frequently embolic, being part of the phenomena of pyæmia or ulcerative endocarditis. The infective matter is distributed by the coronary arteries and gives rise to multiple softenings going on to the formation of **Abscesses**. Such localized suppurations will soften the wall of the heart and may lead to **Aneurysm** or **Rupture**. They do not burst into the cavities of the heart, but are liable to extend to the pericardium, where they give rise to a purulent pericarditis. A suppurative inflammation of the myocardium may also occur by extension from the endocardium in ulcerative endocarditis. The microbes on which this disease depends may propagate into the muscular substance, and so



cause ulceration (*acute ulcer of the heart*) or may even lead to an abscess burrowing in the wall of the heart. This also may lead to aneurysm or rupture of the wall. It must be seldom that such abscesses as those will heal, but there are cases in which calcification either of the abscesses or of portions of the heart's substance which had been softened by embolism has been observed in pyæmia. A case observed by the author and referred to under calcareous infiltration was probably of this kind.

**Interstitial myocarditis** consists in an inflammatory increase of the interstitial connective tissue. There is no doubt that in the great majority of cases the cicatricial or tendinous patches met with in the heart are due to obstruction of the coronary artery, as already described, and in all such cases these arteries should be examined. A more direct local inflammation may be due to other specific causes. Thus in pericarditis or in endocarditis, the inflammation may extend to the muscular substance, causing induration of the superficial layers on the one hand, or of the internal layers on the other. And again in chronic endocarditis affecting the mitral valve we often find, along with thickening of the chordæ tendineæ, a partial conversion of the muscoli papillares into dense fibrous tissue, in fact, an interstitial inflammation with destruction of the muscular tissue.

A localized interstitial myocarditis may be the result of **Syphilis**. There is sometimes a definite gumma, around which a great newformation of connective tissue has occurred, but there may be a local cicatricial condition without any gumma being detected. In these cases there has probably been a gumma at an earlier period. Syphilis may also, perhaps, produce fibrous transformation by causing syphilitic lesions in the coronary artery such as already described.

It is doubtful whether a general interstitial myocarditis occurs. Some have asserted its existence in dilated and hypertrophied hearts, and especially in the hypertrophy of the left ventricle in Bright's disease (see especially Turner). There is, no doubt, in the congested heart hypertrophy of the connective tissue, but the author has failed to find evidence of a true inflammatory condition either here or in the hypertrophy of Bright's disease.

**2. Endocarditis.**—The various forms of endocarditis are somewhat closely related to each other, but it is possible to distinguish three forms, a simple acute, a simple chronic, and an ulcerative or infective form.

(a) **Simple acute endocarditis** (*Endocarditis verrucosa*) occurs as a secondary effect of certain acute febrile diseases.

**Causation.**—Chief amongst the causes is **Acute rheumatism**, but chorea is not infrequently complicated by it, and more rarely scarlet fever, measles, typhoid fever. According to Bamberger 20 per



cent. of the cases of acute rheumatism are complicated with acute endocarditis. Whatever view we take of the origin and nature of acute articular rheumatism, it must be admitted that the blood is in itself of an unusually irritating nature, or else it carries an irritant. The occurrence of acute inflammations in several joints often removed considerably from one another, and the frequent supervention of inflammation in the pericardium and endocardium, are sufficient evidences of this. The irritant, whatever be its nature, seems to act specially on connective tissue membranes, and on such as are exposed to friction of their surfaces. It affects the joints where the synovial membranes lie against each other and in the movements of the joints are moved on one another. It attacks the pericardium where the movements of the heart cause continuous rubbing, and when it attacks the endocardium it affects exactly the localities where the surfaces come in contact. It is as if in addition to the irritant in the blood, the mechanical irritation of friction were necessary to the occurrence of inflammation, and it may be added that in the adult the inflammation is almost limited to the valves of the left side of the heart, where the higher tension of the blood and greater force of the heart make the mechanical force of friction greater than on the right side. We shall see afterwards how this fact bears on the localization of the endocarditis.

According to some authors the acute inflammation is due to the presence of micrococci (Köster, Klebs, Osler). Micrococci have been occasionally detected in the warty vegetations, but they are mostly absent, although possibly in these cases they may have been present at an earlier period. It may be that acute rheumatism is due to some form of micrococcus, but this is not yet determined. Köster asserts that the micrococci reach the valves by the vessels of the endocardium, that is by a process of embolism, while others believe that they are deposited from the blood. If simple endocarditis is really dependent on a micrococcus, it must be a much less virulent one than that which is undoubtedly present in ulcerative endocarditis.

**Characters of lesion.**—The most characteristic effects produced in acute endocarditis are the so-called **Warty vegetations**, which are pale, irregular projections from the surface of the endocardium, generally of small size and somewhat shaggy in appearance. (See Fig. 156.)

The vegetations are composed partly of the swollen, inflamed tissue of the valve and partly of fibrine deposited on the inflamed surface. The inflamed connective tissue produces round cells and is converted into granulation tissue, and the affected parts are thus increased in bulk, and rendered more friable so that irregular projections are produced. The projections are enlarged by deposition of fibrine, which may be regarded as a kind of fibrinous exudation, but is derived from the blood flowing over the surface, and is not really an exudation from the vessels



of the part. It is, perhaps, more correctly a thrombosis, and as the blood is in motion the **White thrombus** is the form produced. The fibrine generally forms the greater part of the bulk of the vegetations.

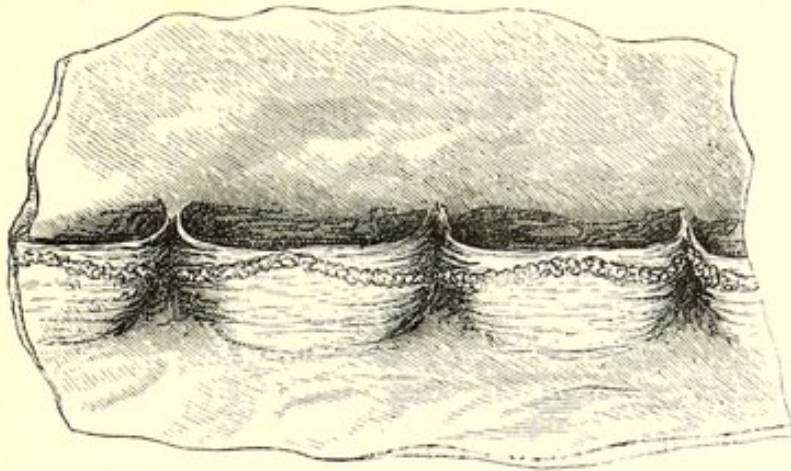


Fig. 156.—The aortic valve in acute endocarditis. The warty vegetations, occupying the lines of contact, are shown.

On their first occurrence the vegetations are limited to the parts of the valves which come against each other in the closure of the valves, and this localization continues more or less throughout.

When after removal of the heart a stream of water is sent into the aorta cut transversely a short distance above the valve, we can look down on the valve closed by the force of the water. It will then be seen that the curtains are not in contact by their margins, but that the line of contact is slightly removed from their edges, and a certain portion of the valve floats free in the water, taking no direct part in the closure of the orifice. The line of contact is nearest the edge of the curtain at the middle of each segment or the corpus Arantii, and forms on either side of this a curved line with the convexity downwards. Between the line of contact and the edge of the curtain the valve is often perforated, and it may even, as we have seen before (p. 365, Fig. 144), be partially resolved into tendinous cords, without interfering with the closure of the valve.

In the mitral valve the line of contact is also removed from the edges of the curtains. In the case of the aortic valve the line of contact is of course on the ventricular side of the curtains, but in the mitral it is on the auricular side, and in order to see the vegetations in acute endocarditis it is usually necessary to examine the orifice by looking in from the auricle. Acute endocarditis of the mitral often escapes notice from this not being done.

In acute endocarditis the warty vegetations frequently demarcate very accurately the lines of contact of the aortic and mitral valves, and the appearances produced in the former case are indicated in Fig. 156.



When the inflammation extends to the valves of the right side, the same principles apply. In the case of the pulmonary valve the vegetations appear along the line of contact on the ventricular aspect of the curtains, and in the tricuspid they are to be seen by looking down through the auricle.

The occurrence of these changes in the tissue renders it **unduly brittle**, and it is not surprising to find that bits of the vegetations are frequently broken off and carried by the arteries to distant parts, to produce embolism there. These broken-off pieces are mostly small, and, beyond the ordinary phenomena of embolism in small arteries and capillaries, they do not by their own nature produce much disturbance, in this respect contrasting with the emboli of ulcerative endocarditis. The softening of the tissue may result in one of two further lesions, either of which may interfere with the function of the valve: these are rupture of the chordæ tendineæ and valvular aneurysm.

**Rupture of the chordæ tendineæ** sometimes occurs in the mitral valve when the inflammation happens to extend to these structures. The result will be that during the systole of the ventricle the valvular curtain will be allowed to some extent to pass upwards towards the auricle, and so allow of regurgitation through the orifice.

**Aneurysm of the valves** is the condition in which a pouch exists, projecting from a valve and with a narrow neck. It occurs in ulcerative as well as in simple acute endocarditis. With the aid of the accompanying diagram, the mode of formation of the aneurysm may be

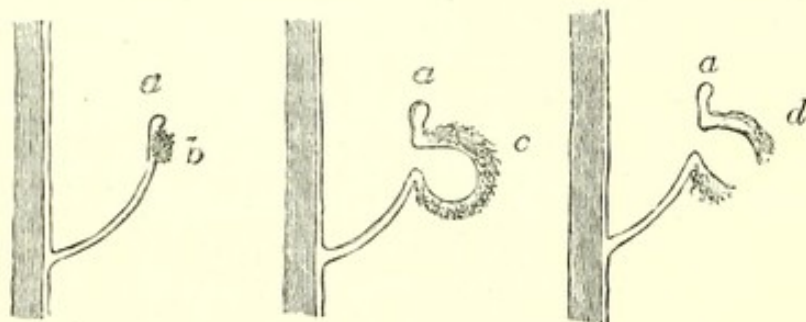


Fig. 157.—Diagram of mode of formation of aneurysm of aortic valve. The curtain (*a*) formed of two layers; at (*b*) its outer layer roughened and softened; at (*c*) the aneurysm, which has burst at (*d*), so as to perforate the valve.

illustrated in the case of the aortic valve, which is its most frequent seat. The semilunar curtains which form the valve are each composed of a double fold of endocardium, as represented at *a*. In acute endocarditis it is the ventricular layer which is principally involved along the line of contact, as indicated at *b*. The aortic layer is usually smooth and unaltered to the naked eye. At the affected part of the ventricular layer the tissue is softened, and during the closure of the valve, the single aortic layer may be unable to support the full pressure of the



blood. In this way it may be pushed towards the ventricle, carrying before it the softened ventricular layer, as at *c*. It will be apparent that, in the case of the aortic valve, the aneurysm will always project into the ventricle. In the case of the mitral valve, on the other hand, the softened layer is on the auricular surface of the valve, and the pressure of the blood during closure of the valve being exercised towards the auricle, the aneurysm consequently projects towards that cavity.

As the aneurysm owes its origin to acute endocarditis, its surface is usually covered with vegetations, which are often very abundant, and may so conceal the aneurysm as to lead to its being overlooked. The aneurysm, again, may rupture, and so produce a perforation of the valve. It is sometimes as if the bottom had been blown out of the aneurysm and a short tube left, surrounded by shaggy vegetations (as at *d* in figure). Even in that case, however, if the neck of the aneurysm be examined, it is often found that the endocardium, as it passes into it, is smooth and unaltered.

(*b*) **Chronic endocarditis** commonly follows on the acute form, and, like it, is related to acute rheumatism, but it may be of more independent origin, not infrequently occurring in the aortic valve along with the similar disease of the aorta, namely, **Atheroma**. (See further on, under Insufficiency of the Aortic Valve.) It is also stated by Roy and Adami that overstrain of the heart, by acting mechanically on the valvular structures, induces oedema and subsequent thickening. In the ordinary rheumatic form it appears as if the irritation were prolonged in a less intense form, and the changes in the valvular structures extend beyond the localities which we have seen to be mainly affected in acute endocarditis.

The chronic form is chiefly characterized by newformation of connective tissue. The granulation tissue of the acute stage develops into connective tissue, and the process extends slowly to the remaining structures of the valves.

In this way arise great thickenings of the valves (Fig. 158), and, as the connective tissue is of

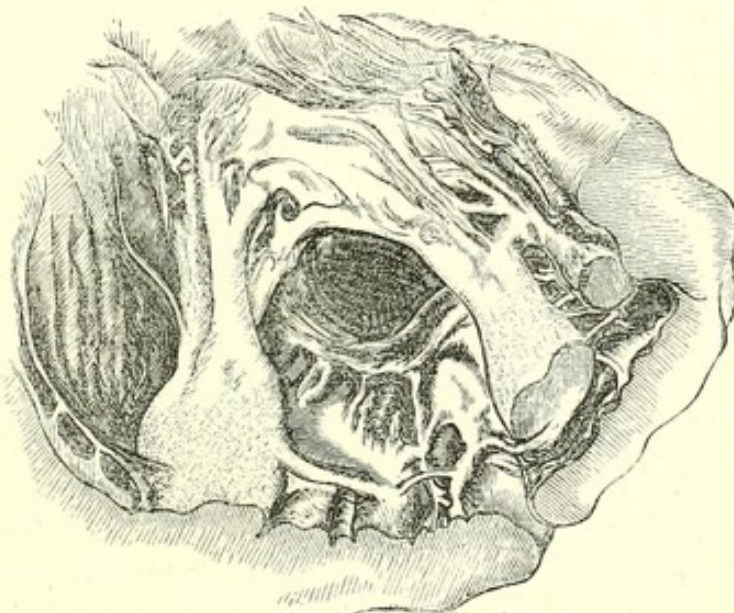


Fig. 158.—Great thickening of the chordae tendineae of the mitral valve, the result of chronic endocarditis.



that dense nature common in chronic inflammations, the thickened valvular structures are often exceedingly **rigid**. The new-formed tissue also **contracts**, and in this way we may have great retractions of the valves leading to serious deformities, as we shall see in studying valvular diseases. Again, it frequently happens that two opposed inflamed surfaces **coalesce**, and we may have still further deformity from this. We have already seen that the inflammation may extend from the endocardium to the muscular substance, leading to cicatricial transformation of it.

In the great majority of cases, the endocarditis is limited to, or has its centre in, the valvular structures, but it sometimes happens that in other parts of the heart an apparently independent endocarditis is set up. This may be along with valvular endocarditis, but separated from it by sound tissue, or it may be without any valvular lesion. We may find an isolated patch of thickening on the surface of the left ventricle, and we have already seen that the disease may penetrate into the muscular substance.

The thickened and rigid connective tissue frequently becomes the seat of **secondary changes**. Fatty degeneration may occur. But this is much less frequent than **Calcareous impregnation**, which may be taken as evidence that the hard, dense, cicatricial connective tissue has, to a great extent, lost its vitality. This condition is of such frequent occurrence that it may be regarded as the normal termination of chronic endocarditis. It sometimes occurs with a very moderated degree of thickening, especially in affections of the aortic valve already referred to as connected with atheroma, and its extent and the date of its occurrence are doubtless determined by individual peculiarities. It may occur in the form of a moderate calcification in the deeper parts of the thickened tissue, or the lime salts may be deposited in a more bulky form so as to give the feeling of considerable stony masses. The occurrence of calcareous infiltration is often of serious import. The valvular structures are rendered still more rigid, and there enters the new element of brittleness. The calcified portion of the valve is exposed very often to mechanical violence in the closure of the valve, and it is common to find that the valve has been broken and a piece of calcareous matter carried off. So far as the valve is concerned, this is not very serious, but as the piece carried off is usually of some size, the resulting **Embolism** is frequently of great consequence. Embolism of the cerebral arteries leading to extensive softening is much more frequent in chronic than in acute endocarditis, and probably the same applies to aneurysms of the larger cerebral arteries, which, as we shall afterwards see, may have their origin in embolism. Embolism of the



spleen and kidneys is also a frequent result. The rough surface left by the breaking-off of the calcareous piece gets coated with fibrine, and here again it is said that micro-organisms may be present. The fibrine may, by getting detached, form a fresh source of embolism.

It is necessary carefully to distinguish from chronic endocarditis the formation of **Opaque patches** on the valves with very little thickening. In this case there is little more than a **fatty degeneration of the endocardium**. Such patches are not uncommon on the mitral valve, and are often erroneously regarded as due to endocarditis.

(c) **Ulcerative endocarditis** (also called *Malignant* and *Diphtheritic endocarditis*) is regarded by some as merely a form of acute endocarditis, but it presents such differences that it seems necessary to place it in a class by itself. The special features in the disease are the activity of the destructive process in the heart, its connection with the existence of micrococci, and the virulence of the metastatic processes when emboli are carried to distant parts.

**Causation.**—The disease is due to the implantation of pathogenic microbes on the endocardium. There seems reason to believe that pyogenic micrococci constitute the form of microbe, but the source of these is sometimes obscure. The disease has been observed to occur in the course of a number of acute febrile affections. It is met with in pyæmia, puerperal fever, acute rheumatism, small-pox, etc., and Osler has pointed out that in a large proportion of cases acute pneumonia has been the primary disease. There are, however, cases in which no definite connection with any other disease can be traced. In the case of pyæmia and puerperal fever the source of the micro-organisms is not far to seek, but in the other cases it is more difficult. In order to the occurrence of ulcerative endocarditis, it seems probable, from the results of experiments on animals, that some previous damage to the endocardium is usually to be inferred. This damage may be the result of a simple endocarditis, and the way being thus opened, the microbes may find entrance.

The micrococci from cases of ulcerative endocarditis have been cultivated on nutrient media in order to determine their characters. They have been found sometimes to possess the usual characters of the micrococci of suppuration. Thus the *Staphylococcus pyogenes aureus* and *albus* and the *Streptococcus pyogenes* have been found (Weichselbaum).

Experiments have been made by injecting cultures of these micro-organisms into the blood of rabbits. It has been found that as a rule their simple presence in the blood is not sufficient to cause ulcerative endocarditis, but that when the valves were at the same time injured then this affection supervened (Orth and Wissokowitsch). On the other hand, it was found by Ribbert that when an emulsion was made of a



culture of these micro-organisms on potatoes the injection of the emulsion was followed by ulcerative endocarditis. Apparently small particles of potato adhered to the endocardium and planted the micro-organism.

**Characters of lesion.**—In its local manifestations this form presents some resemblance to simple acute endocarditis. The disease affects, usually, the valvular structures, and produces an enlargement and roughening of them. But there is not the same localization along the lines of contact, the process generally developing in a defined area, sometimes removed from the valve. Again, the disease, as the name implies, is a much more destructive one, the parts concerned breaking down more readily. In this way perforation or aneurysm of the valve more readily occurs. Sometimes an actual suppuration manifests itself in the valvular structures, but the frequent passage of the blood prevents any considerable accumulation of pus. The ulceration sometimes passes to the muscular wall of the heart, especially when the patch of ulceration is away from the valves. The destructive process spreads rapidly in the myocardium, and a distinct abscess may be the result. In this way also an acute aneurysm of the heart may supervene.



Fig. 159.—Portion of valve in ulcerative endocarditis. *a*, fibrine with colonies of micrococci; the colonies are indicated by the roundish clumps; *b*, endocardium becoming raised by inflammatory infiltration; *c*, elastic layer of endocardium; *e*, round cells infiltrating endocardium, at *d*, passing into superficial layer of fibrine and micrococci.  $\times 22$ .

In their more intimate characters, also, the conditions in ulcerative endocarditis differ from those in the ordinary simple form. As the accompanying figure shows, there is a very marked infiltration of the valvular structures with round cells, almost a suppurative condition. This is immediately overlaid by a fibrinous coagulum, as in the case of simple endocarditis, but mixed with the fibrine there are colonies of micrococci which give quite a striking character to the layer. The



appearances in distant parts are evidence that pieces are frequently carried off from the valves, and, looking to the soft character of the superficial parts we are not surprised at the occurrence of **Multiple embolism**.

Perhaps the most striking feature in this disease is the occurrence of **Metastatic abscesses** in distant parts. These are found in the heart itself, in the spleen, in the kidneys, in the skin, etc. They are everywhere of small size, and usually in large numbers. These abscesses are obviously related to emboli carried off from the endocardial lesion, and lodged in the finer arteries or capillaries. The accompanying figure represents a small artery in the midst of an incipient abscess in the



Fig. 160.—From the kidney in ulcerative endocarditis. An artery is shown, plugged with a dark material containing micrococci. Around these are myriads of leucocytes which are infiltrating the necrosed wall of the vessel and the kidney tissue around.  $\times 90$ .

kidney. It is seen that its calibre is plugged by a material in which are occasional masses of micrococci. At the distal part the wall of the artery is obscure, apparently from necrosis, and the vessel is buried in an enormous aggregation of inflammation cells. The micrococci are



frequent in the capillaries and Malpighian vessels, but not generally with obvious inflammation; apparently they are in that case of recent development, possibly to some extent post-mortem, and their products have not had time to produce inflammation.

We have seen that the micrococci present, locally, an intensely irritating action, and that necrosis of the tissue attacked is a common result. It is probable that they also produce changes in the constitution of the blood. Patients affected with this disease frequently present, like those in pyæmia, a yellow colour of the skin approaching to that of **jaundice**. The probable explanation of this is, that the blood-corpuscles undergo solution, and the colouring matter stains the skin. Besides that, we often have little ecchymoses of the skin, and small **Hæmorrhages** in the pia mater, and even in the brain substance. These have been found associated with the presence of colonies of micrococci in the capillaries, and are to be ascribed to the action of these in weakening the wall, and allowing escape of blood.

3. **Pericarditis**.—The pericardium is comparable in its anatomical and pathological relations to the serous sacs, and, to a certain extent, to the synovial. The pleura, peritoneum, and pericardium are to be regarded, as we previously found in studying œdema and dropsy, as large lymphatic spaces. These sacs are composed of connective tissue, and lined with a single layer of flat endothelium. By means of numerous stomata they are in communication with the lymphatic vessels, and to some extent, with one another. The pericardium is in less direct communication with the pleura and peritoneum than these are with each other, but by circuitous routes there is some communication, especially with the pleura. It is to be remembered also that through each serous cavity there is a certain circulation of serous fluid. This fluid does not accumulate in the sac, because it is carried off as quickly as it is transuded from the vessels, but if the transudation increases greatly there may be an accumulation and consequent dropsy.

(a) **Acute pericarditis**.—In considering the **Causes** of this disease it is of some consequence to note that the inflammation usually affects the whole surface at once. This seems to indicate that an irritant has found admission to the pericardial sac, and by the motions of the heart and the natural currents of the fluid in the sac, has been carried hither and thither throughout it. There are many cases in which the tubercular virus is the irritant; we find tubercles mixed up with the inflammatory products. (See afterwards under Tubercular Pericarditis.) The majority of cases of simple pericarditis are associated with acute rheumatism. There are some cases in which the disease appears to be of spontaneous origin; it is ascribed to cold.



In regard to the **Phenomena** which manifest themselves at the onset of an acute pericarditis, we may presume that the irritant induces the changes in the vessels which have been described in treating of inflammation in general, but opportunities are wanting for observing the consequent redness, as patients survive this early stage. **Exudation** from the vessels soon follows, and **serous fluid** begins to accumulate in the sac. As the inflammation affects the surface of the sac, the endothelial lining is very directly involved. The flat endothelial cells are to a considerable extent shed, being apparently killed by the irritant, but they may also be found showing signs of germination. The exudation consists primarily of exuded liquor sanguinis with contractile cells, but soon **Fibrine** is deposited on the inflamed surface. The detachment of the endothelium seems to be the circumstance which determines the coagulation, on principles already explained. The deposition of fibrine occurs on both visceral and parietal layers of the sac, but it is usually thickest on the visceral surface, where it may present shaggy masses on the surface of the heart. It is whitish in colour, and of soft, almost gelatinous consistence. The fibrinous layers on the opposed surfaces of the pericardium are usually separated by serous fluid which occupies the sac, but an appearance is often presented which suggests the application and withdrawal of the layers while still in a soft plastic condition. This appearance is variously described as the **Honeycomb** or **Pine-apple** condition, and it has been aptly compared to that presented when two pieces of bread, thickly buttered, are stuck loosely together and then separated. This honeycomb appearance, it will be understood, is most markedly present on those parts of the pericardium where the heart in its movement comes most frequently against the parietes.

The exuded fibrine has the usual characters, as seen under the microscope, forming a fine reticulum, in whose meshes are leucocytes, and sometimes red corpuscles.

If the inflammation has been slight and transient, there may be little beyond a small serous and fibrinous exudation, which is gradually absorbed. But as a general rule further changes develop, and these are mainly in the connective tissue of the pericardial sac. This shows evidence of inflammation by the presence of innumerable cells, so that by degrees it is converted into **Granulation tissue**. This inflammatory transformation, according to the intensity of the irritation, penetrates deeply, extending frequently to the interstitial connective tissue in the muscular substance of the heart. A layer of vascular granulations thus forms beneath the fibrinous exudation. The layer of granulations has the general tendencies as well as the structure of granulation tissue elsewhere, it tends to develop into connective tissue as soon as the inflam-



matory irritation becomes sufficiently mild. With the subsidence of the inflammation there will be a reduction of the serous exudation. The fibrine is also disposed of, partly undergoing fatty degeneration and so becoming absorbed, and partly eaten into from beneath by the granulation tissue. The result of this is that the granulations are, as it were, laid bare, and a vascular layer occupies the place of the former fibrinous deposit.

With the absorption of the exudation the two layers of the pericardium come in contact, and a coalescence of the granulating surfaces, more or less complete, occurs. The vessels intercommunicate, and the two layers, so far as they are in contact, become virtually one. As the granulation tissue passes on in its development into connective tissue, the **pericardial sac undergoes** partial or complete **obliteration**, the uniting agent being vascular connective tissue, and so we have the condition of **Adherent pericardium**. The connection will at first be delicate, and may be torn through, but as time goes on it gets firmer, and a condition results in which the two layers are absolutely inseparable. Under certain circumstances the coalescence of the two layers is not complete, and there is only a partial adhesion; in that case the adhesions are sometimes greatly stretched by the movements of the heart so that tags or ligaments may unite the surface of the heart to the perietal layer.

In some intense and prolonged cases of acute pericarditis, **Suppuration** occurs, and the serous exudation in the pericardium gives place to pus. This, however, is a rare occurrence. On the subsidence of the inflammation the pus dries in by the absorption of its fluid, and its debris remains as dead matter. In this way caseous material is produced, which subsequently gets infiltrated with lime salts; this may ultimately become consolidated so as to form calcareous plates in the midst of thick adhesions.

**Septic inflammations**, such as those which occur in pyæmia when an abscess in the substance of the heart extends to the surface and bursts into the pericardium or gives off septic microbes, or in the rarer case of perforation of an ulcer of the stomach or œsophagus into the pericardium, are purulent from the outset.

**Adherent pericardium** frequently leads to **Hypertrophy and Dilatation of the heart**. It does so both directly by interfering with the muscular tissue, and indirectly by requiring the heart to use increased exertion in performing its task.

It is to be remembered that adhesion of the pericardium takes origin in inflammation, and from this circumstance it results that in several ways the action of the heart is interfered with. During the acute stage of the inflammation there is fluid



in the sac of the pericardium, and by the mere mechanical pressure of this fluid the cardiac contractions are interfered with, and if the effusion continue long enough there may be hypertrophy to overcome the obstacle. But again, the inflammation extends a certain distance into the muscular wall of the heart beneath the pericardium. A certain portion of muscle is thus interfered with in its action and more vigorous contraction is required of the rest. There may even be considerable thickening of the pericardium by development of connective tissue, and this extending some distance in the connective tissue between the muscular fibres may seriously compromise them. But further, when adhesion is complete, the heart in contracting must drag in with it the parietal as well as the visceral pericardium. In the normal state the two surfaces of the pericardium slide on one another, and the parietal layer accommodates itself to the movements of the heart. If there is adhesion, however, unless the adhesion be very loose, there can be no such sliding, and there must be some loss of force in dragging the parietal layer inwards. But the parietal layer is normally attached to surrounding parts, and the attachment may be rendered closer by the inflammation. The heart will drag on these parts in contracting, and this will add to its work. It is well known that dragging in of the intercostal spaces is a common sign of adherent pericardium. As these causes vary to a considerable extent, the amount of hypertrophy varies in proportion. As the causes also act nearly uniformly on the heart, the hypertrophy is general, that is, it usually affects all the cavities of the heart. Such a hypertrophy may almost completely compensate, so that a person with adherent pericardium and a very large heart may have no cardiac symptoms.

(b) **Chronic pericarditis.**—As observed above, acute pericarditis often becomes chronic, and, in that case, usually results in adhesion of the pericardium. A more direct chronic inflammation results in the condition designated **White spots**, **Milk spots**, or **Soldier's spots**, which are very common pathological conditions. They occur in about half the cases examined post mortem, and their frequency seems nearly in direct proportion to age. They are in the form of well-defined, whitish, opaque areas on the surface of the heart, of very various size, sometimes very small, at other times so large as almost to cover the anterior surface. Their edges are usually abrupt and well defined, but they may merge gradually in the pericardium. They have often a brilliant white tendinous appearance, but may be more dull. They are most frequently situated on the anterior surface of the right ventricle, and next on that of the left ventricle, especially near the apex. They are also met with on the posterior surface, especially near the base of the heart, and on the intrapericardial portions of the great vessels. They are more uncommon on the parietal layer of the sac.

These spots actually consist in a thickening of the pericardium, presenting merely dense connective tissue covered with endothelium (see Fig. 161).

We have here a circumscribed inflammation affecting by preference certain districts, and the cause must be a local one. It seems to be due



to the irritation resulting from the projection of the heart against its surroundings. The commonest seat is where the anterior surface of the right ventricle comes against the sternum at the place where the edges of the lungs turn aside and expose the pericardium. The sternum is less yielding than most surrounding parts, and so the irritation is greater here. The spot where the left ventricle near the apex strikes against the fifth rib, is the next most frequent site.

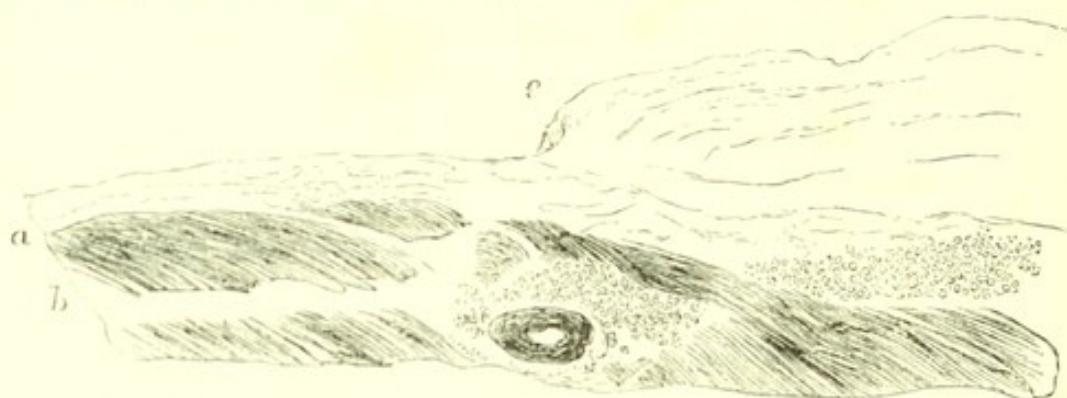


Fig. 161.—Soldier's spot on pericardium. *b*, muscular wall of heart; *a*, pericardium, thickening at *c* so as to form spot.  $\times 25$ .

(*c*) **Tubercular pericarditis** has commonly a sub-acute or chronic course. The infection is sometimes derived from the pleura, the virus reaching the pericardium by the connections between the sacs. It may reach the pericardium, however, according to Weigert, by the lymphatic glands of the mediastinum. These, having become affected as a result of tuberculosis in the parts from which their vessels are derived, and having acquired adhesions to the parietal layer of the pericardium, may impart the disease to the latter.

Occasionally we meet with an acute fibrinous pericarditis in which, when we remove the fibrine, numerous small white fresh tubercles are to be seen on the pericardium. More frequently, however, the two layers of the pericardium are firmly united by connective tissue, and it is impossible to separate them. In the midst of the thickened layers are to be seen yellow caseous masses, sometimes of considerable size. These represent old tubercles, and the whole process is a more or less chronic one, even from the outset.

In the thick adherent pericardium it is usually possible to distinguish two layers of tubercles, one belonging to the visceral, and one to the parietal layer of the pericardium.

Under the microscope there will be found large masses having the indefinite characters of caseous necrosis with occasional fresh tubercles, in the midst of tissue which presents in some parts evidences of recent inflammation and in others merely dense connective tissue.



## VII.—VALVULAR DISEASE OF THE HEART.

In studying endocarditis, we have seen that the valves are frequently altered in their structure; we have now to consider these alterations more specifically, and their effects on the heart and circulation. It is not usual to designate anything as valvular disease unless it interferes with the function of a valve or orifice. The function of a valve is to close an orifice under certain circumstances, and we speak of valvular disease when the alterations are such as either to obstruct the orifice or to interfere with its closure by the valve. Hence valvular lesions may be divided into two kinds, namely, narrowing or stenosis of the orifice, and insufficiency of the valve. In referring to these same lesions, as they affect the current of blood in the heart, we speak of obstruction of an orifice and of regurgitation through the orifice. Of course it would be a very incorrect use of language to speak of obstruction of a valve, or insufficiency of an orifice.

Valvular disease occurs much more frequently in the valves of the left side than in those of the right, and hence we have chiefly to do with the mitral and aortic valves. We have already seen that in the foetus it is the valves of the right side which are most frequently affected, and we have connected this with the fact that these valves are more liable to variations of pressure in the foetus than in the adult.

**1. Insufficiency or Incompetency of the mitral valve.**—This is a condition in which, during the systole of the heart, some portion of the blood passes back into the left auricle instead of the whole being forced into the aorta.

The actual physical conditions are somewhat various, but most of them are related to chronic endocarditis. The commonest is that in which the valvular structures are thickened by the new-formed connective tissue and retracted and shortened from its contraction. This applies to the curtains themselves, but still more to the chordæ tendineæ, which become thickened and shortened, and frequently grow together, so that they hold the curtains rigidly drawn down and do not allow them to go together during the systole of the ventricle (see Fig. 158, p. 387). Again, much more rarely, in acute endocarditis, the valve may be perforated, as in the case of valvular aneurysm, or the chordæ tendineæ torn so as to allow a portion of the valve to flap upwards through the orifice. Lastly, without much alteration of the curtains, there may be a *relative insufficiency* of the valve. That is to say, the cavity of the ventricle sometimes enlarges greatly, and produces enlargement of the orifice, which the valve is no longer able to cover. There are some cases of permanent hypertrophy and dilatation of the left ventricle (as from overstrain) where this occurs, but it may be met with where the dilatation is temporary, as in the flabby fatty heart of typhus fever and anæmia. It is not to be supposed that the so-called anæmic murmurs are usually due to this cause, but there is in some cases an actual mitral regurgitation. When recovery occurs, and the heart resumes its former vigour, the valve will again cover the orifice.



The results which follow constant insufficiency of the mitral valve are frequently very serious and far-reaching. At each ventricular systole blood regurgitates into the left auricle, and the most direct result is over-distension of this auricle occurring at successive intervals. The wall of the auricle is weak, and does not offer much resistance to the distensile force. There is another result which often follows, apparently from the unduly forcible impact of the blood against the endocardium, and the over-stretching of this membrane, namely a thickening of the endocardium. We may find it generally thickened and opaque, or there may be patches of opacity.

But the results do not confine themselves to the auricle—the abnormal blood-pressure is reflected to the **Pulmonary veins** which feed the auricle, and they become distended. The distension is further reflected to the pulmonary capillaries and arteries, and finally to the right ventricle. The right ventricle is over-distended, and, as a consequence, **Hypertrophy of the right ventricle** is a common result of mitral insufficiency. The obstruction to the circulation may extend to the *venæ cavæ*, the right ventricle and auricle being over-dilated. In this way **General venous engorgement** comes about with all the consequences which will be described in the next section in dealing with mitral stenosis.

The amount of blood which the left ventricle sends into the aorta will be diminished in proportion to that which passes back into the auricle. In consequence, the systemic circulation will be partially starved. In some cases the left ventricle undergoes hypertrophy. This is mainly due to the fact that at each diastole it will receive an excess of blood, namely, that which would normally arrive from the auricle along with that which regurgitated from the ventricle. The ventricle having thus to deal with an increased mass of blood will undergo dilatation and hypertrophy.

2. **Obstruction of the mitral orifice. Mitral stenosis.**—This name is applied to the condition in which the mitral orifice is not large enough to allow of the usual

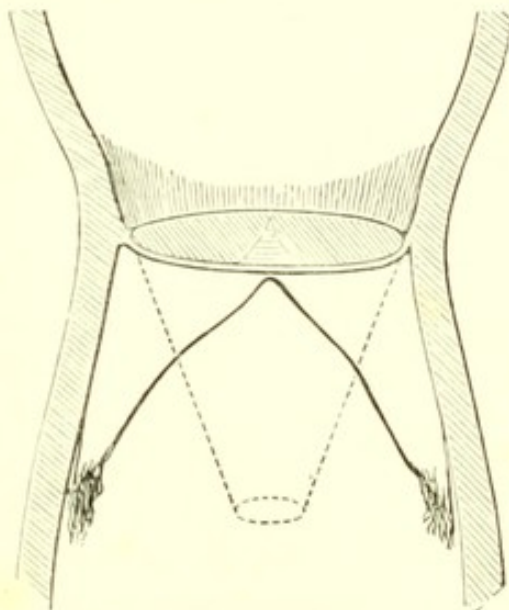


Fig. 162.—Diagram of funnel-shaped deformity of mitral valve. The dotted lines indicate the coalesced curtains forming a funnel projecting into the ventricle with a reduced aperture at the apex.

quantity of blood passing from the auricle to the ventricle. The normal width of the mitral orifice may be roughly estimated with the fingers; in the adult it should allow the index and middle fingers to pass freely through as far as the first joint. The contraction may be very slight or it may be to such an extent that hardly a crow-quill can be admitted into the orifice. In the case of stenosis of the mitral as well as in that of the aortic orifice, the obstruction is usually caused by the curtains of the valves becoming thickened and rigid, but especially by their coalescence. The thickened curtains grow together by their edges, and so the valve is converted into a funnel with its apex turned down into the ventricle—the so-called **Funnel-shaped deformity**. The normal orifice is at the base of the curtains, but when the curtains coalesce, the orifice while becoming contracted is moved downwards, and comes to have its site at the apex of the funnel.

This will be understood from the accompanying diagram (Fig. 162), in which black lines represent the orifice and curtains in their normal condition during the diastole



of the ventricle, the curtains lying back against the wall of the ventricle, and the orifice at their base. The dotted lines represent the coalesced funnel-shaped valve; the contraction of the orifice and its removal downwards being shown. The chordæ tendineæ are commonly thickened and often partly incorporated in the funnel (see Fig. 158, p. 387). On laying open the ventricle this thick, rigid, funnel-shaped deformity is often strikingly prominent. These conditions result from chronic endocarditis, and it will be readily understood that the rigid valve is frequently incompetent, so that this condition is often combined with the one before. There are also not infrequently changes in the aortic valve.

Obstruction is occasionally produced by thrombi growing on the valve, or having their seat in the auricle and projecting into the orifice. This is a rare cause of obstruction, and a still rarer is the presence of tumours growing in such a way as to obstruct the orifice.

It might be supposed that the vegetations occurring in acute endocarditis would obstruct the orifice, but although these rough projections undoubtedly interrupt the even flow of blood, and may produce during life what is technically a murmur of mitral obstruction, yet their actual influence on the function of the orifice must be very slight, and we are not to look for any definite evidences of their influence on the circulation.

We have now to consider the **results** to the circulation of mitral obstruction. The most direct effect will be **dilatation of the left auricle**, as the blood is, to a certain extent, hindered in its passage into the ventricle. As a consequence, the whole pulmonary vessels will be loaded and the right ventricle distended with the accumulated blood. On the principles already laid down there will be increased action and consequent **Hypertrophy of the right ventricle**, and this is commonly more extreme than in mitral insufficiency. The contraction of the orifice interferes with the passage of blood into the **left ventricle**, which, in extreme cases, is, as it were, starved of blood. The increased force of the right ventricle may in great part make up for this deficiency, and sometimes there is also aortic insufficiency, so that the ventricle is fed from the aorta as well. According to these various circumstances will be the state of the left ventricle. It may be actually atrophied and appear as a small appendage to the enlarged right ventricle, or it may be normal in size or even hypertrophied. In any case the hypertrophy of the right ventricle is the predominating condition. The shape of the heart is more quadrilateral, the apex is blunt and formed by the right ventricle. During life instead of the defined apex beat of the left ventricle, there is the more diffused heaving of the right.

As a further consequence, we have a permanent **Passive hyperæmia of the pulmonary circulation**, with consequent brown induration of the lungs. There will also be a tendency to slight hæmorrhages, showing itself in the sputa. Œdema of the lungs readily develops. The dilatation of the right ventricle, when followed by thrombosis, also frequently leads to embolism of the pulmonary artery and the hæmorrhagic infarction. The hyperæmia is reflected to the **systemic venous circulation**, especially if the dilatation of the right ventricle lead to relative insufficiency of the tricuspid valve, and we find evidences of passive hyperæmia of the liver (nutmeg liver), kidneys and other organs. Not infrequently serious œdema of the skin and dropsy of the serous cavities develop. Thrombosis in the veins of the legs often complicates the condition, and this again may be a source of pulmonary embolism.

**3. Insufficiency of the aortic valve.**—This is the condition in which, after the completion of the ventricular systole, a portion of the blood regurgitates into the left ventricle through the imperfectly closed semilunar valve. It is usually brought



about by chronic endocarditis. The individual semilunar folds are thickened and

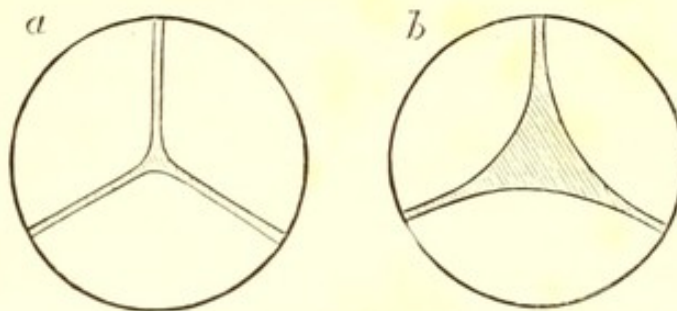


Fig. 163.—Diagram of aortic insufficiency. *a*, the normal valve closed as seen from above; *b*, the valve with curtains shortened and leaving a triangular space.

shortened, the actual length of free margin being reduced. The consequence is that, during the closure of the valve, the edges have not sufficient length to meet perfectly and so a triangular aperture is left. This is illustrated diagrammatically in the accompanying figure. The contraction may reach such an extent as to leave only a nodulation on the

wall of the aorta in place of the curtains. These changes are very commonly accompanied by adhesion of the adjacent folds of the curtains, and this necessarily causes contraction of the orifice; indeed, the curtains as such may disappear, leaving only a diaphragm with a permanent aperture in its middle, the condition being similar to that illustrated in Figure 142, p. 364. An unusual cause of aortic insufficiency is **Perforation of the valve** as a result of acute endocarditis or the bursting of a valvular aneurysm. Of course the perforation of the curtains beyond the line of contact already referred to is not to be mistaken for a pathological perforation. A rare cause of aortic insufficiency is the **Tearing of one of the curtains**. During severe exertion the blood-pressure in the aorta may be so much raised as to rupture a curtain, and such a wound will hardly unite as it will be torn asunder at each closure.

It may be added here that aortic disease is often accompanied by mitral disease, chronic endocarditis, having its origin in rheumatism, attacking both. Chronic endocarditis of the aortic valve again is often connected, as we have seen, with endarteritis or atheroma of the aorta, and in that case it is not so likely to be associated with mitral disease.

It may be interesting here to observe that the origin of the endocarditis has an important bearing on the **Age at which these valvular lesions occur**. Acute rheumatism is a disease of youth and manhood, and most cases of valvular disease take origin in it. Accordingly, diseases of the valves are most common, at least in their inception, between the ages of ten and thirty. But there are some cases of aortic disease which, as we have seen, stand in a different category. Chronic endarteritis or atheroma is a disease mostly of advanced life, and so it is more common in old persons to meet with aortic disease than with mitral.

We have now to consider the **Effects** of this insufficiency of the aortic valve on the circulation. As the semilunar valve does not close completely, the aorta in its recoil after the ventricular systole will force blood back into the ventricle as well as forward into the systemic arteries. This extra mass of blood driven with considerable force into the left ventricle will overfill it and forcibly distend it, while the systemic circulation will be proportionately starved. The natural result is **Dilatation and hypertrophy of the left ventricle**, which may almost completely compensate. In this disease, therefore, the primary and prominent fact is the enlargement of the left ventricle. As the ventricle propels a much larger amount of blood into the aorta, and with abnormal force, there is sometimes a resulting **Dilatation of the arch and great vessels**. In this way there may even be an actual aneurysm of the arch.

It is to be added, that as hypertrophy of the left ventricle is associated with



dilatation of that cavity, there is frequently a consequent widening of the mitral orifice. As a result of this, we may have a relative incompetency of the mitral valve, which is incapable of completely covering the dilated orifice. In this way the consequences already considered of **Mitral insufficiency** may be brought about; but they are usually much less pronounced than in the primary form of this lesion, and of late occurrence. It is to be remembered, also, that mitral disease often co-exists with aortic.

**4. Obstruction of the aortic orifice. Aortic stenosis.**—In this lesion the passage of blood from the left ventricle into the aorta is interfered with. It is, in the great majority of cases, caused by chronic endocarditis. The conditions already described as leading to insufficiency of the valve mostly produce also obstruction of the orifice, by causing rigidity of the curtains, and this is all the more marked when calcareous infiltration ensues. Where the valve is, in the way already mentioned, converted into a rigid diaphragm, then there must be great obstruction of the orifice as well as insufficiency of the valve. These two forms of lesion are, therefore, usually found associated. In acute endocarditis the roughening of the curtains may to some extent obstruct the flow of blood, but the interference is trivial, and will hardly lead to any of the secondary results of aortic stenosis.

The obstruction at the orifice prevents the blood getting away fully during the systole of the ventricle, and there comes to be an overfilling of the ventricle. The ventricle, as in the previous case, is stimulated to increased exertion, and so here also the primary phenomenon is **Hypertrophy of the left ventricle**. This may completely compensate for the obstruction, and persons may go about comparatively well with an obstructed orifice and enlarged left ventricle. But this is not so likely as in the previous case, and any extra need for exertion on the part of the heart, or weakness of its muscle, may lead to incomplete compensation. In such a case the ventricle will get abnormally dilated, and the auricle will not be able to empty itself fully into the dilated ventricle. The **Pulmonary circulation** will become engorged and the **Right ventricle** overloaded, and so we may have all the evil consequences of mitral disease. It will be observed, however, that, as the left ventricle is much more capable of undertaking additional work than the right, it succeeds much more frequently in bringing about a complete compensation. The hypertrophied left ventricle having to dispose of an increased mass of blood, generally does so slowly, and the pulse is consequently slow and regular.

It will have become apparent that in many cases aortic disease is associated with mitral, and that there is frequently combination and complication of the resulting changes in the heart and circulation.

**5. Valvular disease of the right heart.**—We have already seen that, except in the foetus, this form of disease is uncommon as a primary lesion. In cases of acute or chronic endocarditis with well marked lesions on the left side, however, there are very often distinct indications of inflammation in the valves of the right heart.

**Relative insufficiency of the tricuspid valve**, although secondary, is often of considerable consequence on account of the effects to which it leads. We have seen that in mitral disease the right ventricle usually dilates and hypertrophies, and with this change in the ventricle the orifice widens. The valve may thus be unable to cover the enlarged orifice, and become insufficient. In other forms of dilatation and hypertrophy of the right ventricle, as in that due to obstruction to the pulmonary circulation, the same thing may occur. The tricuspid orifice normally admits readily three fingers up to the first joints, and when enlarged it is not uncommon to



meet with cases in which it admits four, five, six, or even seven fingers. If the valve is thus incompetent to close the enlarged orifice the blood will, during the systole of the ventricle, regurgitate into the auricle. The wave will be propagated into the veins of the neck, and there will probably be an aggravation of existing congestion of the systemic veins.

It need only be added that if chronic endocarditis attacks the tricuspid or pulmonary valves, it may produce results similar to those effected in the mitral and aortic. The tissue here, however, is less substantial to begin with, and the inflammation is usually much less intense, and so the changes are rarely of any great consequence.

#### VII.—INFECTIVE TUMOURS, TUMOURS PROPER, AND PARASITES OF THE HEART.

**Miliary tubercles** are sometimes met with, occupying the endocardium, or the pericardium, or even the muscular substance, in cases of acute general tuberculosis. **Syphilitic gummata** are occasionally met with in the muscular substance, and they are accompanied by interstitial inflammation. The gumma has the usual indefinite and varied structure, in the more recent stage softer and more cellular, in the more advanced stage largely composed of dense connective tissue, with probably a caseous centre. There may be a single tumour replacing a portion of the wall, and surrounded by the cicatricial tissue of chronic inflammation, in which the endocardium or the pericardium may be involved. In the latter case there is adhesion of the two layers of the sac. Or the gummata may be multiple, in which case there is a more diffused interstitial inflammation. In some cases there is a localized interstitial inflammation without a gumma, the latter having probably been absorbed. The result is a more or less contracted cicatrix. The **Malignant lymphoma** (Hodgkin's disease) originating in the mediastinal or bronchial lymphatic glands, not infrequently extends to the pericardium and heart. In **Actinomycosis** there are occasionally abscess-like lesions produced by embolism, similar to those in pyæmia.

**Primary tumours** are exceedingly rare in the heart, but a sarcoma of the endocardium has been described, as also primary fibroma, myoma, and lipoma.

**Secondary tumours** are also rare. **Sarcomas** occurring in the neighbourhood may spread to the heart, and especially those of the mediastinum, involving first the parietal and then the visceral pericardium. Sometimes also a **Cancer** of the œsophagus extends to the pericardium. Cancers when they become generalized very often occur in the heart, in the form of round pale tumours.

Of **Parasites**, the echinococcus and the cysticercus cellulosæ have been found in the heart. The cysticercus of the tænia solium is frequent in the heart of swine, and that of the tænia mediocanellata in cattle, but it is very rare in man.



## SECTION I.—CONTINUED.

## B.—BLOOD-VESSELS.

- I. DISEASES OF THE ARTERIES. 1. *Thrombosis and Embolism*, 2. *Obliteration*, 3. *Acute inflammation*, 4. *Chronic endarteritis or Atheroma*; its causes, resulting lesions and effects on the circulation, 5. *Endarteritis obliterans*, 6. *Retrograde changes including fatty degeneration of the intima, and calcareous infiltration with ossification*, 7. *Aneurysms*; causation, weakness of wall and increase of blood-pressure; localization; coats of artery in aneurysms; thrombi; condition of branches; effects on the heart; and on parts around; terminations, by cure, by pressure effects, by rupture. *False and spurious aneurysms*; *cirsoid aneurysm or aneurysm by anastomosis*; *traumatic aneurysm*; *dissecting aneurysm*; *varicose aneurysm*. 8. *Syphilitic and tubercular affections of arteries*.
- II. DISEASES OF THE VEINS. 1. *Thrombosis*, 2. *Inflammations, including septic phlebitis and pyæmia*. 3. *Varix, causation and character of changes*; *Hæmorrhoids*; *varicocele*. 4. *Newformations in veins*.

THE blood-vessels are to be regarded as tubes of which the essential constituent is the intima. According to circumstances the intima becomes clothed with external and middle coats, and so we have arteries and veins. We have already seen that, in nearly all newformations, blood-vessels are produced as well as the proper tissue, and it is first a tube composed of intima which is formed, or a capillary. This primary vessel is capable of enlargement and further complication in the way just indicated, so that a transformation of the primary capillaries into arteries and veins may take place. This formation of vessels and their further development according to the requirements of the tissues is an exceedingly common occurrence, and may be regarded as equivalent to that which occurs in the formation of the tissues during the period of growth of the body as a whole. A process of a similar nature is sometimes seen when the obstruction of an artery causes the current to be in great part diverted into other channels. We know that in this case the anastomosing vessels enlarge, small arteries becoming converted into large ones, and perhaps even capillaries into arteries. The vascular system is thus an exceedingly plastic one, and possesses great powers of newformation and development according to the needs of the tissues.



## I.—DISEASES OF THE ARTERIES

1. **Thrombosis and Embolism.**—These conditions having been somewhat fully discussed in previous pages, it remains here to refer to the more local changes.

**Thrombosis** occurs as a secondary result of disease of the walls of arteries, chiefly in cases of atheroma and aneurysm. In the former it is often the cause of the final closure of the vessel. There is also thrombosis as a result of ligature of arteries. Acute inflammations in arteries also induce thrombosis; this is especially the case in septic inflammations. There may be in this way a condition similar to the more frequent septic thrombosis in veins (thrombophlebitis.)

**Embolism** is very frequent in arteries. If the embolus be a simple one, such as a piece of thrombus, the portion of artery affected will undergo a process of chronic inflammation and the plug will become organized in the manner about to be described. A septic embolus on the other hand, will become the centre of a suppurative inflammation. We have also embolism from the penetration of tumours and parasites.

2. **Obliteration of arteries.**—In several of the affections to be considered in the succeeding paragraphs, partial or complete occlusion and obliteration of arteries occur, and the processes although differing somewhat in detail have many points in common. In all of them there are usually conjoined the two processes of thrombosis and inflammation.

**Obliteration by ligature** affords the simplest illustration. When an artery is ligatured, the internal and middle coats are torn through, as shown in Fig. 164. In consequence of the injury and stagnation of blood thrombosis occurs, and the thrombus, which will contain a considerable excess of leucocytes, will extend to the nearest branch. This is followed by inflammatory changes affecting primarily the internal coat. This tunic becomes cellular and swollen so that it bulges inwards and impinges on the coagulum. The inflamed internal coat becomes vascularized, new-formed vessels penetrating into it from the vasa vasorum. The internal coat thus sends buds or projections inwards which replace the thrombus by vascularized granulation tissue. The new-formed vessels are produced by budding from the vessels of the external and middle coats, and these tunics also take some part in the inflammatory process, but the middle coat is much less active than the external and its special character disappears in the process. The granulation tissue thus formed has the usual tendency to form connective tissue, and the final result is that the portion of artery concerned is resolved into a piece of dense connective tissue which may form part of the cicatrix of a wound.



When Arteries have been wounded a somewhat similar process occurs. If the artery is cut across, the muscular coat by its contraction narrows the calibre of the artery and withdraws it within its sheath. The blood flowing out through the orifice deposits leucocytes, and a clot forms within the sheath and at the orifice of the vessel, by and by completely obstructing it. This coagulum will be a white thrombus. The ensuing processes will be similar to those just described.

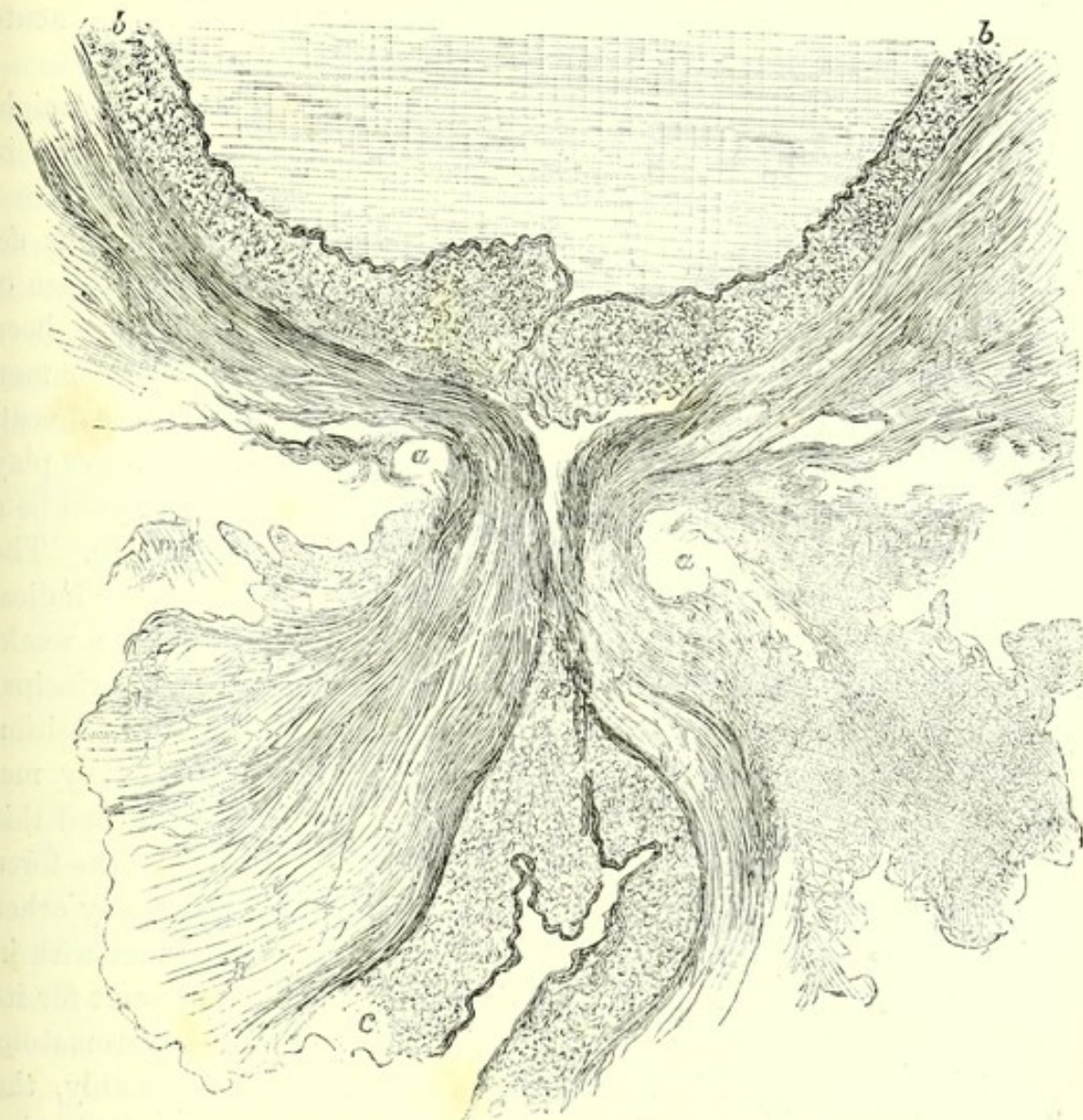


Fig. 164.—Longitudinal section of an artery at seat of ligature. *a, a*, apertures in which the silk ligature was found. The external coat is drawn in at this place, while the middle and internal coats are absent, being absolutely disjoined. These coats are seen above at *b* and below at *c*.  $\times 35$ .

In the case of obliteration from other causes, the process of final organization is frequently interfered with by the diseased condition of the wall of the artery or by the character of the obliterating agent. If the latter be of an irritating nature, as in septic embolism, then there can be no organization, and if the vessel wall be seriously diseased, as is often the case in atheroma, then the whole process of organization may remain absent.



3. **Acute inflammation of arteries. Acute arteritis.**—Arteries often taken part in inflammations in their neighbourhood, but an independent acute inflammation of the walls is very rare. An endocarditis affecting the aortic or pulmonary valve sometimes extends to the neighbouring parts of the aorta or pulmonary artery, producing warty projections on the internal surface of these. More particularly, when in acute endocarditis a considerable thrombus forms on the aortic valve and comes against the wall of the aorta, it may lead to an acute endarteritis there.

**Purulent arteritis** is the result of septic inflammation in the neighbourhood or of septic embolism. The process is similar to that in septic phlebitis.

4. **Chronic endarteritis. Atheroma** (*Arterio-sclerosis, endarteritis deformans, endarteritis nodosa*).—These names are applied to a disease of very frequent occurrence in arteries, the nature of which has been differently regarded at different times. According to the more common view of the process the disease is an inflammation of the internal coat, and this view is so far a correct one; but degenerative processes play such an important part in the course of the disease that it would be a mistake to regard it as merely an ordinary chronic inflammation. The fact that the disease is peculiarly one of advanced life is another indication of its degenerative nature, and it may almost be said that a weakness of the wall of the vessel must be regarded as one of the principal agents in the causation of the disease. While there is this predisposition the actual supervention of the disease is apparently caused by mechanical irritation. Its principal seat is the arch of the aorta, and this is doubtless due to the fact that this part is more exposed to the force of the wave of blood during the systole of the heart than any other portion of the arterial system. The disease is frequently met with in the arteries of the brain, and here it is more difficult to account for its occurrence on the theory of mechanical irritation, but the atheromatous patch is often situated just at a bifurcation, where, presumably, the vessel wall is more exposed to the force of the current. It is also common, as already remarked, in the coronary arteries, where we may presume that the blood-pressure is higher than in other arteries of their size. Again, it is met with in the pulmonary artery in cases of hypertrophy of the right ventricle, the excessive impulse of the blood from the hypertrophied ventricle apparently determining its occurrence. As we shall see afterwards, syphilis produces a disease of arteries in some respects similar to atheroma.

The disease consists in a more or less localized thickening of the internal coat. The thickening is nearly always distinctly limited in



area, so that we speak of **Atheromatous patches** (hence the name *Endarteritis nodosa*). When we examine the aorta in the earlier stages we see elevated areas with tolerably abrupt edges, and usually of a dead white colour as compared with the surrounding intima. These patches are hard, and cut like cartilage. In the arteries of the brain the diseased parts are seen, without opening the vessels, as white opaque patches, and the vessel is more rigid than normal, so that it does not collapse; the calibre also is diminished by the inward projection of the patch. On cutting into the patch, in either the aorta or a cerebral artery, there is often an opaque yellow colour revealed in the deeper parts and this is an indication of fatty degeneration. Very often, too, there is, especially in the aorta, calcareous infiltration of the deeper parts of the patch, but these two conditions will be more fully discussed further on.

In considering more particularly the details of the process, it is instructive to examine microscopic sections, including the edge of the patch and the neighbouring parts of the vessel (see Figs. 165, 166, and

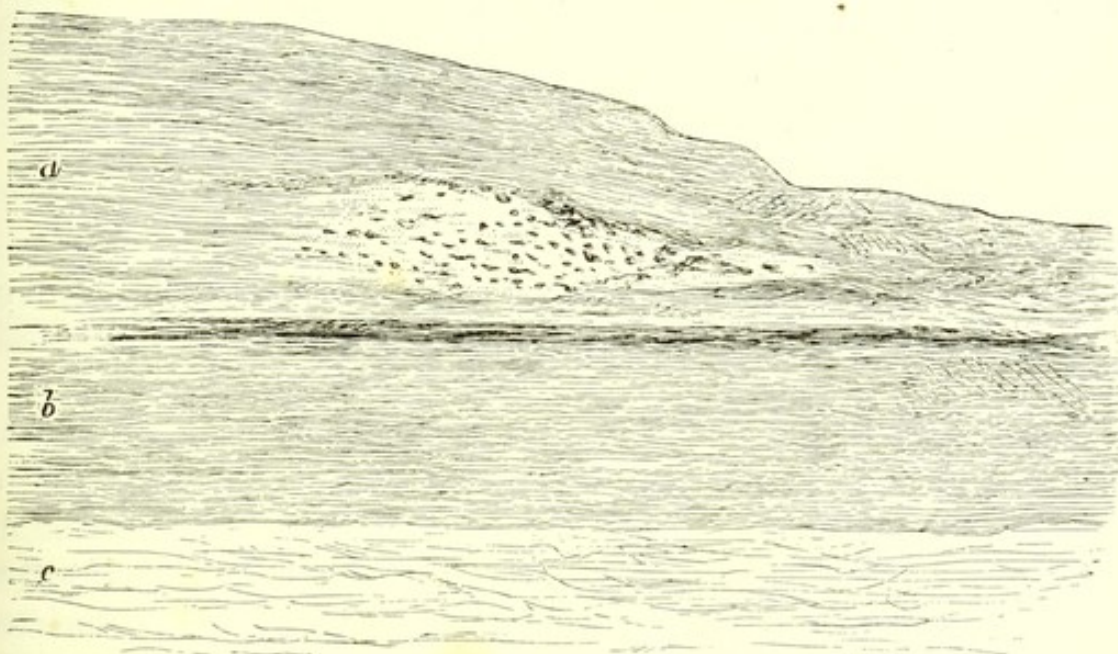


Fig. 165.—Atheroma of aorta. The internal coat (*a*) is seen to be thickened. In its deeper layers there are darker markings indicating the commencement of fatty degeneration. *b*, middle, and *c*, external coat.  $\times 22$ .

168). In well-preserved recent cases it can be seen that the endothelium of the intima is continued over the patch, and in nearly all cases there is little difficulty in observing that the patch is really a thickening of the internal coat, as shown in the figures. The thickened intima is composed of a dense connective tissue, which in the early stages contains many round, oval, and stellate cells. In a fully formed patch the structure is often exceedingly indefinite, and there are few cells that are at all prominent even when staining agents are used. The structure



is indeed half obsolete, and as already indicated, fatty degeneration readily occurs. The fatty degeneration begins usually in the deeper layers of the patch, so that on making a section one generally finds in the portion of the patch bordering on the middle coat an opaque yellow streak (see also Fig. 165). The fatty degeneration at first affects the cells of the intima, but as time goes on the intercellular substance gives way, and the tissue breaks down so that a cavity containing fatty debris with crystals of margarine and of cholesterine is formed. From the character of this fatty debris the name **Atheroma** is derived, and the cavity thus formed is sometimes called an **Atheromatous abscess**. (See Fig. 166.) The tissue superficial to the cavity may ultimately give way, and so expose the cavity, thus forming

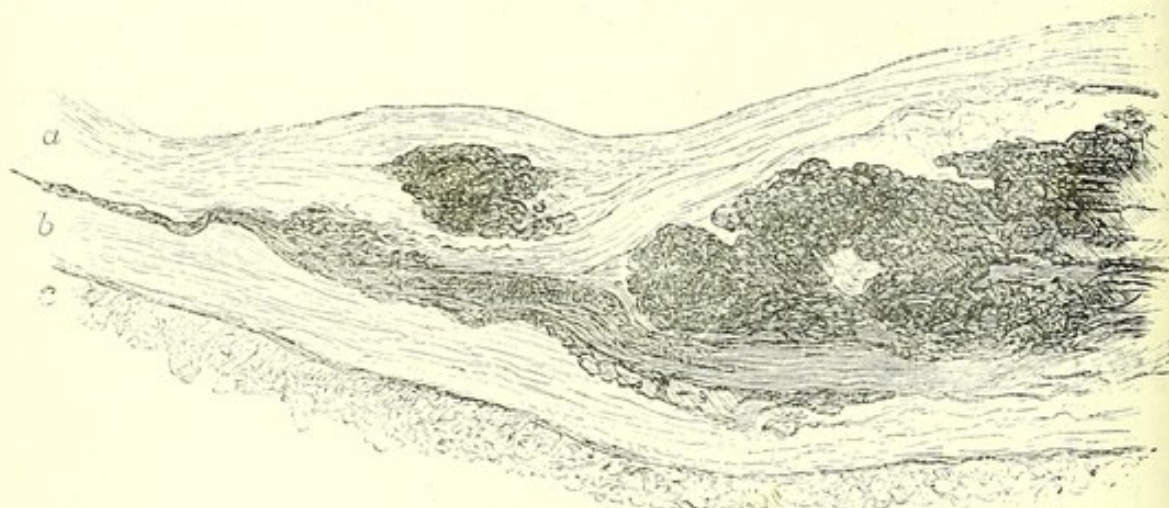


Fig. 166.—Atheroma of femoral artery. The greatly thickened internal coat (*a*) is shown. In the midst of it, towards the right, are dark masses consisting mainly of fatty debris, and representing the atheromatous abscess. At one part the middle coat (*b*) is considerably encroached on from within.  $\times 22$ .

an **Atheromatous ulcer**. The ulcer sometimes presents calcareous masses in its walls, and in any case it may induce the deposition of fibrine on its surface. The fatty material may, on the other hand, be absorbed without forming an actual ulcer, and in that case there will be produced depressions and irregular foldings and puckerings of the internal coat. These depressions and foldings have often a kind of cicatricial appearance.

**Calcareous infiltration** is a frequent result of the disease under consideration. It is generally stated that the calcareous salts are deposited in the fatty degenerated structure. But this is not always the case. The dense tissue of the patch may become infiltrated with lime salts without a preceding fatty degeneration. This is peculiarly the case when the tissue is very hard and dry. We shall see afterwards that some individuals present a very great tendency to the deposition of lime salts in the walls of their arteries, and there are cases of atheroma in which this tendency is very marked, so that with compara-



tively little atheromatous thickening there may be very extensive calcareous deposition. At first the salts are deposited in fine granules, and in the deeper layers of the patch, but as time goes on these aggregate into consistent masses so that we have **Calcareous plates** of various forms and sizes, sometimes attaining to a square inch in area. These plates, having abrupt edges, not infrequently tear through the remaining layers of the intima, and present an edge or angle inside the vessel. They may even become to a considerable extent separated, and hang into the calibre attached only by a strip of intima like a hinge. The rough calcareous edge very commonly induces a deposition of fibrine, and the thrombus thus formed may subsequently get detached, and form an embolus. The determination of the preponderance of the fatty or calcareous change appears to depend on individual peculiarities. Both forms are very frequent in the aorta. The fatty change preponderates greatly in the cerebral arteries.

There are some cases in which the atheroma has a much more distinctly inflammatory course than that described above. The thickened intima is infiltrated with round cells, and there is also inflammatory infiltration of the middle and external coats, the round cells aggregating specially around the blood-vessels, so that the whole wall of the vessel is affected. Such cases are perhaps hardly to be placed in the same category as ordinary atheroma, but as this condition occurs like the other, mainly in the aorta, and the result is a thickening of the internal coat in patches, it is difficult to draw a distinct line of demarcation between them. By some, this condition has been specially distinguished under the designation **Acute aortitis**.

The **Changes in the middle and external coats** in atheroma are important. At first the middle coat is hardly at all affected, but if the process advances far, and especially if the degeneration of the patch causes much deformity of the internal coat, then the middle coat becomes softened and loosened. It also becomes affected with fatty degeneration, the muscular fibre-cells being first involved (see Fig. 167). In advanced cases, especially in the cerebral arteries, we may even find,

along with fat granules, crystals of cholestearine and margarine and pigment, so that through time the middle coat may be considerably

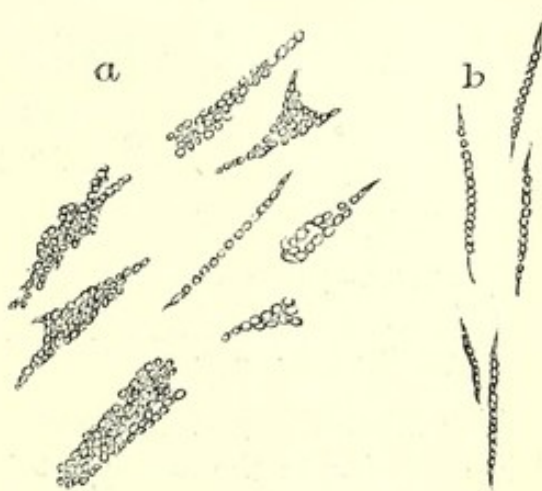


Fig. 167.—Fatty degeneration in atheroma, *a*, from internal coat; *b*, muscle cells from middle coat.  $\times 350$ .



broken up. This process may be present in the middle coat while the hard thickening of the internal coat is still present, and we may find a localized dilatation of the vessel occurring, into which the thickened intima dips. According to Thoma the primary lesion in atheroma is in the middle coat, generally a weakening or atrophy of it. The thickening of the intima is, in his view, compensatory to the lesion in the media. The external coat is much less seriously affected than the middle; if at all involved it presents increased thickening and vascularization.

We have now to consider the **Effects of atheroma on the circulation.** There are three alterations which it produces, each of which may, according to circumstances, have important effects on the circulation. The disease causes *narrowing of the calibre, loss of elasticity and rigidity of the wall, and interference with the muscular contractility of the vessel.*

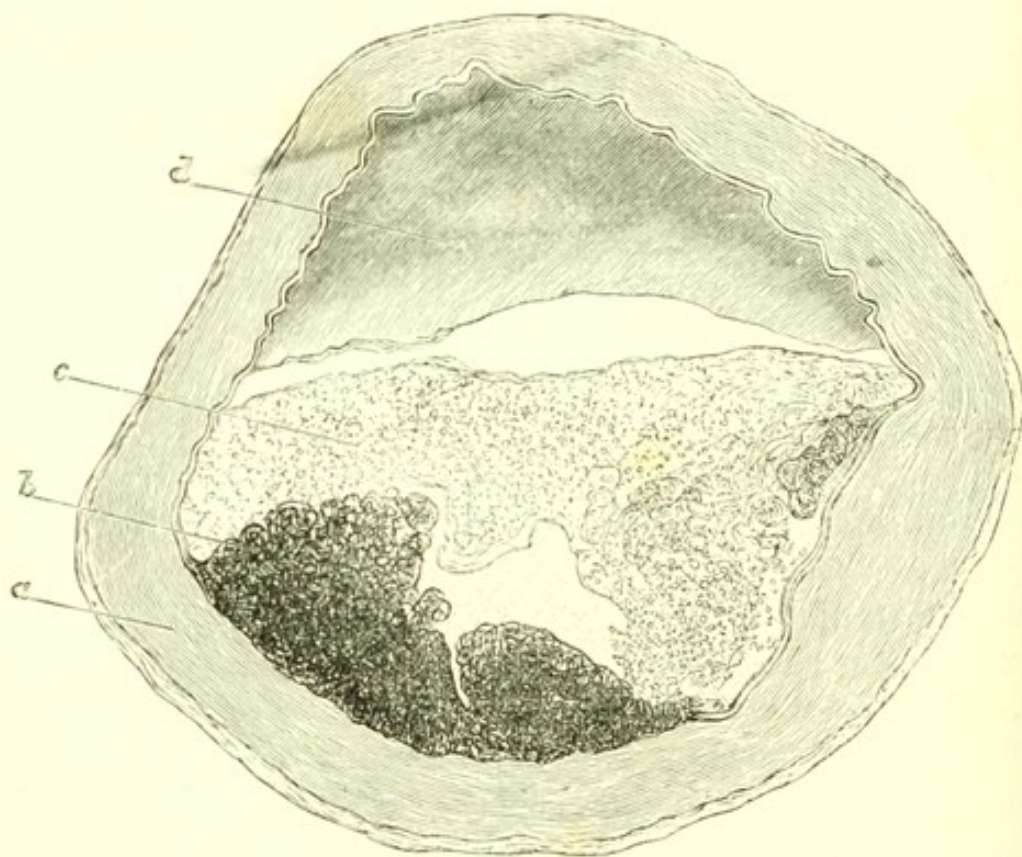


Fig. 168.—Atheroma of cerebral artery. The greatly thickened internal coat is seen. In its substance are dark masses (*b*) in which margaric crystals were found. On the surface is a paler layer (*c*) consisting of partially organized thrombus; (*d*) blood occupying the remaining calibre.  $\times 34$ .

The **Narrowing of the vessel** will be of little consequence in such large arteries as the aorta, but in the case of the cerebral vessels, the cardiac arteries, and those of the legs, the interference with the circulation may be considerable (see Fig. 168). We have seen this illustrated in the case of the coronary arteries of the heart, especially when thrombosis supervenes, where the disturbance of the circulation sometimes produces severe angina, and may lead to sudden death.



The **Interference with the contractility** will also affect mainly the arteries of smaller dimensions, and in the case of the arteries of the brain, the absence of that control of the circulation which is afforded by their varying calibre may lead to serious consequences.

**Rigidity and weakening of the wall** are very important consequences of atheroma, especially in the aorta, where they are probably

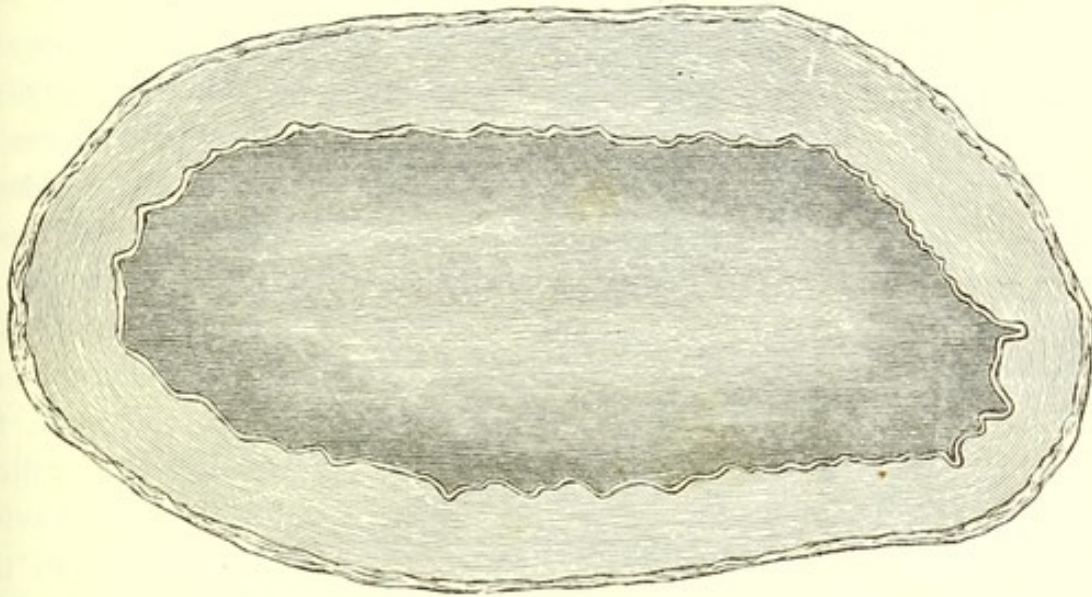


Fig. 169.—Transverse section of normal cerebral artery to contrast with Fig. 176.  $\times 34$ .

the chief factors in the causation of aneurysm. The immediate consequence of rigidity of the aorta is that during the systole of the heart the vessel does not dilate, and at the end of the systole it does not recoil, so that the force of the elastic recoil is lost to the circulation, and in distant parts there is apt to be more or less stagnation. As a result of this we have hypertrophy of the left ventricle, which is often very marked in cases where much calcareous infiltration exists. The hypertrophied ventricle sending the blood forcibly into the rigid aorta produces commonly a diffuse dilatation of the arch. The influence of atheroma in producing aneurysm will fall to be considered subsequently.

The time of life at which atheroma is most frequent is a point of some importance in relation to the causation of aneurysm. According to Rokitsansky, it is commonest between the ages of forty and sixty. It is still pretty frequent down to thirty years of age, but rapidly diminishes in frequency from that age downwards. It is extremely rare under twenty years, and when it does occur it is mostly in connection with congenital anomalies of the great vessels or heart, such as stenosis of a main stem with defect of the septum, etc.

**5. Endarteritis obliterans.**—This is not an independent disease, but is frequently of considerable importance as a part of the phenomena of other conditions. It affects the finer vessels of certain organs, and consists, like atheroma in a thickening of the internal coat (see Fig. 170)



which, in these fine arteries, frequently leads to complete obliteration.

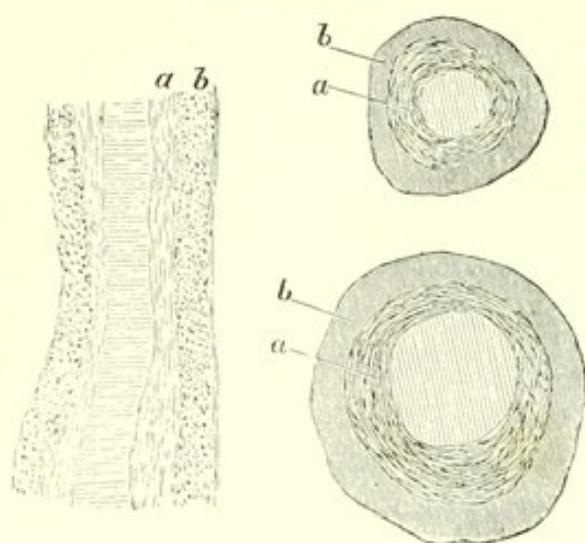


Fig. 170.—Endarteritis obliterans in arteries of kidney, *a*, thickened and fibrous internal coat; *b*, middle coat.

This form of lesion is seen especially in interstitial inflammations of organs, and is particularly frequent in chronic interstitial nephritis, where it will be again referred to.

6. **Retrograde changes in arteries.**—We have already referred to retrograde changes in connection with atheroma. Some more independent forms have still to be considered.

**Fatty degeneration of the intima of arteries.**—This condition is apt to be mistaken for

atheroma. We frequently see, especially in the aorta, yellow markings slightly raised above the internal surface, and having the appearance of superficial figurings. They are to be found in the aorta, pulmonary artery, and other parts of the arterial system. If a portion of such a patch be examined under the microscope by removing a thin layer by a section parallel to the surface, it will be found that the condition consists in a fatty degeneration of the cells of the intima. The flat branched cells are demarcated by the presence in them of abundant fat drops. If the fatty degeneration is much advanced then the intercellular substance becomes also the seat of fat drops, and the cells are no longer demarcated. It sometimes happens that when the degeneration is very advanced the little patch softens and an erosion forms. This erosion is very superficial, and is not to be mistaken for the atheromatous ulcer.

This condition is sometimes met with in the bodies of comparatively healthy persons, but we have already seen that in anæmia and in emaciated persons it is frequent, and it is to be classed in the same category as fatty degeneration of the muscular tissue of the heart.

**Calcareous infiltration of arteries.**—We have already seen that in atheroma there is very frequently a calcareous infiltration of the affected structures, and it has been stated that individual peculiarities appear to play an important part here. In some cases a more independent calcareous deposition occurs, and here individual peculiarities are of still greater consequence. Calcification of the middle coat is the most frequent and important form. This is very frequently associated with atheroma, but it is noteworthy that the atheroma and calcareous



deposition very often affect different arteries or different parts of the same artery. We may have, for instance, atheroma of the aorta and calcification of the middle coat in the femoral and smaller vessels; or there may be atheroma as well as calcification in the femoral, but in their extreme degrees the two are present at separate parts of the artery. For instance, Fig. 166, p. 408 and Fig. 172 are from the same femoral artery, and from parts near each other; but the one shows atheroma with fatty degeneration, while the other exhibits very advanced calcareous infiltration of the middle coat. It may here be remarked that calcification affects by preference the arteries of medium and smaller size. It is often very pronounced in the femoral and brachial, and extends to those of smaller size, but not to the finest arteries.

The deposition occurs primarily into the muscular fibre cells of the

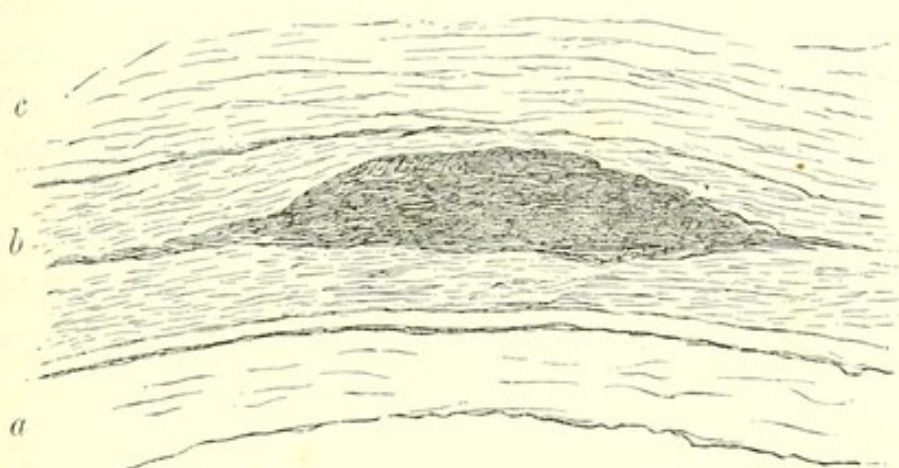


Fig. 171.—Calcareous infiltration of the middle coat (*b*) in an artery, early stage, with atheroma (*a*). At the edges of the affected part, the general outline of the muscle fibre cells can be made out.  $\times 60$ .

middle coat, and at first marks these out by the presence of fine opaque granules (see Fig. 171). The granules flow together till a patch is formed of an opaque appearance, but confined to the middle coat. As the calcareous deposition increases, the patch sometimes assumes a

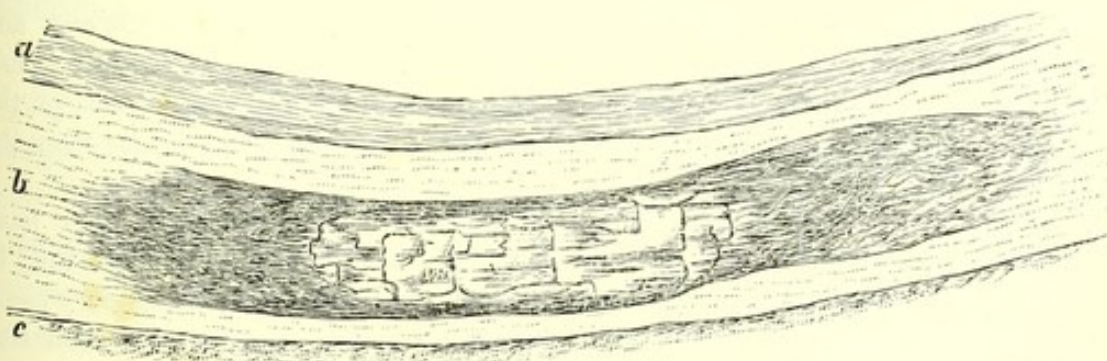


Fig. 172.—Calcareous infiltration of the middle coat of an artery. The lime salts have aggregated together so as to produce a crystalline appearance. This was taken from the same artery as Fig. 166.  $\times 22$ .

crystalline appearance, and the material may become broken and irregular, as in Fig. 172. This condition necessarily gives greatly increased



rigidity to the walls of the arteries, and the feeling of rigidity so often felt in the radial and other smaller arteries is mostly due to this cause, and not to atheroma. The fact, however, that calcification of the middle coat is so often associated with atheroma renders this rigidity to some extent an indication of the existence of atheroma in the larger arteries. To the naked eye the calcified parts frequently manifest themselves by the appearance of a circular opaque striation, visible especially when the artery is laid open and viewed from within. The appearance is better seen if the artery be allowed to dry partially, as then the normal tissue becomes more transparent and the chalky structures more prominent. In that case complete or partial rings are seen, not unlike the irregular cartilaginous rings of the bronchi, but smaller. Another method of rendering the calcification prominent is to tear off the internal coat, which is usually somewhat loosely attached.

The calcified middle coat sometimes undergoes a true **Ossification**, so that spiculae of bone may be mixed with calcareous masses. The ossification is secondary to the calcareous infiltration, and is always small in extent compared with the latter. It is probable that it is really the result of a chronic inflammation around the calcareous masses, the new-formed connective tissue, penetrating amongst these, becoming transformed into true bone by reason of the rigidity of the structures and the abundance of lime salts.

The author has observed a similar development of true bone in a collapsed hydatid cyst in the liver. In this case the new-formed connective tissue had penetrated amongst the degenerated products of the parasite, and in some parts had developed osseous tissue with regular lacunae and canaliculi.

The calcareous infiltration of the middle coat renders the artery peculiarly rigid, and, when extensive, induces hypertrophy of the left ventricle just as rigidity of the aorta does. The rigid vessels are also liable to dilatation, as in the case of the aorta. But besides that, the calcified middle coat is brittle, and affords a much less effective resistance to the distensile wave of blood. An additional strain on the circulation, or some special movement of the body, may break the brittle coat and directly lead to aneurysmal dilatation of the artery. It is not unlikely that peripheral aneurysms are frequently induced in this way, and especially those of the popliteal region, where the artery is peculiarly liable to mechanical injury from the movements of the limb.

A peculiar **Calcification of arteries by metastasis** has been described. We have already seen that when great destruction of bone is occurring the lime salts may be deposited in distant parts, the lungs and mucous membrane of the alimentary canal being the chief seats. Kuttner has described a case in which the incrustation occurred in the arteries. The deposition increased in amount the further the arteries were removed from the heart, and it was especially manifest where the current was permanently impeded, as where a small branch issued from a comparatively large stem. In this case it was the intima that was incrustated, and it was peculiar that the veins were not in the least affected. Apart from the incrustation,



the arterial coats were apparently normal. The source of the lime salts was an acute caries of the vertebral column from the first dorsal to the last lumbar. In relation to the pathology of the case, it is to be added that there was a purulent interstitial nephritis, so that the excretion of lime salts was presumably obstructed. It is remarkable that in this case the metastasis was to the arteries, and that the lungs and mucous membrane of the intestine were entirely free.

**7. Aneurysms.**—An aneurysm is a localized dilatation of an artery. Taking this as the definition, it will follow that in the aneurysm the coats of the artery are stretched, and, to some extent, retained as the covering of it. We shall afterwards see that there are aneurysms in which this is hardly the case, in which the coats rather give way and rupture. Such aneurysms are often designated **Spurious aneurysms**, of which it will be necessary afterwards to describe several varieties. Even of the aneurysms which come under the above definition there are several varieties, but we have to consider here, in the first place, those which arise by a limited dilatation of an artery. The artery may be dilated in its whole circumference, thus forming a spindle-shaped or **Fusiform aneurysm**, or the dilatation may be at a limited part, and we have then a **Sacculated aneurysm**.

**Causation of aneurysms.**—It may be said that permanent undue dilatation of an artery can hardly occur unless the wall of the vessel be weakened. It is perhaps conceivable that the blood-pressure might be so increased as to dilate an artery unduly, but such increased pressure would, in its extreme form, be temporary, and unless there were some weakness of the vessel-wall it would hardly produce a permanent dilatation. In the actual production of the ordinary aneurysm both of these conditions, namely, weakness of the wall and increased blood-pressure are nearly always to be traced.

The more important of the two conditions is **Weakness of the wall of the artery**. By far the most frequent cause of this weakness is **Atheroma**. We have already seen how atheroma with great calcareous deposition may lead to dilatation of the arch of the aorta. But aneurysm is even more frequent in cases where the ordinary fatty change takes place in the patches, causing irregularity in the internal coat. By the formation of the atheromatous abscess and ulcer, or still more frequently by the absorption of the fat and consequent shrinking, depressions and furrows are formed in the internal coat. It is to be remembered also that the atheromatous part is rendered unduly rigid by the thickening of the internal coat, and that any extra distending force will act specially on parts which have been weakened. Another element of equal importance is the weakening of the **middle coat**, which we have seen to be a frequent consequence of atheroma. Fatty



degeneration is of frequent occurrence in the media under the atheromatous patch. In the case of the peripheral arteries calcareous infiltration of the media may occur with or without atheroma, and the calcification may be such as to render the media exceedingly brittle. It is true that, according to old experiments of John Hunter and Home, destruction of the internal or even internal and middle coats of arteries in animals is not sufficient to cause dilatation of these vessels, but with a seriously weakened middle coat any considerable increase of the blood-pressure may be sufficient to cause dilatation.

While atheroma, especially when associated with weakening of the middle coat, is concerned in the causation of aneurysms, there are certain circumstances which indicate that this of itself is not a sufficient cause. One of these is the age at which aneurysms occur. We have seen that atheroma is a disease of advanced life, being most frequent after forty, although still very common between thirty and forty; but aneurysms are most common between the ages of thirty and forty. Another circumstance is the great preponderance of aneurysms in the male. It is true that atheroma is commonest in the male, but not to anything like the same extent as aneurysm. Bizot found in 189 cases of the latter, 171 in males and 18 in females, and Hodgson found in 63 cases 56 in males and 7 in females. It is noteworthy further that in some localities aneurysms are uncommon, while atheroma is by no means so rare. The great frequency of aneurysm in the army as compared with civil life is another circumstance requiring explanation.

All these circumstances are to be explained on the view that, in addition to weakening of the vessel-wall, **Increase of blood-pressure** is needed to produce aneurysms. In severe continued exertion we find the most frequent cause of increased blood-pressure. The engineer who has to manipulate a piece of hot iron while wielding a heavy hammer, or the soldier who has to perform long marches with heavy accoutrements, must put a strain on his heart and larger vessels which ordinary persons are not liable to. It will be apparent that men are much more exposed than women to such excessive stress on their vascular systems. It is again between the ages of twenty and forty that men are mostly exposed in this way. Between the ages of twenty and thirty atheroma is uncommon, but between thirty and forty it is tolerably frequent, and if it exists the liability to excessive stress will render the occurrence of aneurysm more probable. On the same principle we explain the greater frequency of aneurysm in some countries as compared with others. The excessive stress to which workmen in our engineering and shipbuilding establishments are frequently put in this country goes far to explain the frequency of aneurysm here.



Another circumstance of importance is the **Localization of aneurysms** in different arteries. Nearly half the cases of aneurysm occur in the aorta, and the great majority of these in the thoracic portion. We have seen that atheroma is most frequent in this vessel, but, in addition to that, the aorta is most exposed to the **excessive pressure of the blood** when the heart is stimulated to unduly forcible action. Next to the aorta the popliteal artery is most frequently the seat of aneurysm. It has already been pointed out that this vessel is especially liable to injury from sudden flexures of the leg, especially when the middle coat is rendered brittle by calcareous infiltration. But besides that, the vessel is so situated as to be liable to localized increase of blood-pressure. As it issues from the popliteal space the artery passes between the two heads of the gastrocnemius, and is liable to constriction by the contraction of the muscle. On this principle has been explained the frequency of popliteal aneurysm in flunkies whose principal occupation is to exercise their gastrocnemii in standing. But apart from that, when a person makes a severe exertion in a standing posture, the gastrocnemii contract vigorously, thus producing a partial obstruction of the artery and an increase of pressure above the obstructed part, while the general blood-pressure is also increased.

The following table from Crisp gives a statement of the frequency of aneurysms in different situations. It is to be observed, however, that it greatly underestimates the number of aneurysms of the cerebral arteries, and also those of the pulmonary artery in phthisis pulmonalis.

	551 Cases in English Records.	364 from London Museums.
Aorta thoracica (including arch), - -	175	207
„ abdominalis (and main branches), -	59	46
Art. pulmonalis, - - - - -	2	2
„ iliaca com., - - - - -	2	2
„ iliaca int., - - - - -	0	1
„ iliaca ext., - - - - -	9	7
„ glutæa, - - - - -	2	0
„ cruralis, - - - - -	66	12
„ poplitea, - - - - -	137	50
„ tibialis postica, - - - - -	2	2
„ innominata, - - - - -	20	3
„ carotis, - - - - -	25	9
„ cerebrialis, - - - - -	7	1
„ temporalis, - - - - -	1	0
„ ophthalmica, - - - - -	1	0
„ vertebralis, - - - - -	0	1
„ subclavia, - - - - -	23	12
„ axillaris, - - - - -	18	8
„ subscapularis, - - - - -	1	0
„ brachialis, - - - - -	1	0
„ radialis, - - - - -	0	1



Before leaving the subject of the causation of aneurysms it should be mentioned that anything which weakens the wall of an artery will render it liable to dilatation. Thus a blow whose effects extend deeply enough to affect the arterial coats, a piece of fractured bone pushed against an artery, may be regarded as causes. Again it has been pointed out by Ponfick that embolism is a frequent cause of aneurysm in the cerebral vessels, the embolus injuring the walls of the artery.

**The Coats of the artery in aneurysms.**—An aneurysm begins as a localized dilatation or as a little pouching of a limited portion of the artery. In the former case we have the commencement of a fusiform, and in the latter case that of a sacculated aneurysm. Even at this early stage the middle coat is already impaired, and it has probably been affected to begin with. Small pouches, into which one can put the tip of the finger, are often to be seen in an atheromatous aorta. The little pouch enlarges, and while its opening remains small, it enlarges outwards in all directions into a distinct sac. The walls of the sac are sometimes folded back around the aperture so as to apply themselves to the external surface of the artery, and these may become mutually adherent. In that case the aperture has a sharp edge, and it appears as if the wall of the artery were folded over so as to form the wall of the aneurysm.

**The Internal coat** enters variously into the constitution of the aneurysmal wall. In the case of a fusiform aneurysm it is continuous over the internal surface, and probably presents very marked atheromatous changes. In the sacculated form it is usually to be traced some little distance from the edge of the aperture on the wall of the aneurysm, and even in the midst of the internal surface of the sac pieces of internal coat, greatly altered as a rule, may be discovered.

**The Middle coat** appears to be passively affected in the formation of the aneurysm. Its fibres are loosened and stretched as dilatation occurs, and from the first are liable to degeneration. Even in fusiform aneurysms it is often difficult to trace the middle coat far from the beginning of the dilatation.

**The External coat**, on the other hand, may be regarded as forming the chief constituent of the sac. It is composed of connective tissue, and, as the aneurysm grows, inflammatory processes occur and new connective tissue is formed, so that the sac does not necessarily undergo thinning as it enlarges. The connective tissue of the external coat usually makes common cause with that of neighbouring structures, and so the aneurysm acquires adhesions, and the surrounding structures come to form, to a certain extent, constituents of the sac.

**Thrombi in aneurysms.**—Blood-clots are of nearly constant occurrence



in aneurysms, and they may almost be regarded as constituents of the sac, as they doubtless aid to a great extent in preventing rupture. Coagula are most frequent and most important in sacculated aneurysms. We meet with them in the form of firm, dry layers, which present a distinct stratification, generally parallel to the wall of the aneurysm. The coagula often form a kind of sac inside the proper sac, and after removal retain the shape of the aneurysm. The coagula are primarily **White thrombi**. The white blood-corpuscles adhere to the rough internal wall of the aneurysm, and after they have accumulated to some extent coagulation occurs and a thrombus is formed. This process is repeated, and the formation of fibrine is thus in successive layers. Not infrequently the layers become partially separated, and the blood insinuates itself between them. A layer of whole blood is thus formed, and when it coagulates we have a red thrombus mixed with the white. As time goes on the clots become firmer, dryer, and more stringy. The layers next the sac are often of a pearly whiteness, and may be taken for connective tissue. Under the microscope, however, they are seen to be devoid of definite structure, and acetic acid brings out no elongated nuclei as in the case of connective tissue. (See under Hyaline Degeneration, p. 171.) There seems little tendency to the organization of these coagula unless the whole aneurysm becomes filled and its cavity obliterated by them. The continual distension of the cavity seems to interfere with the process of organization; but, if the cavity be obliterated by the formation of clots, then organization proceeds, and by and by the aneurysm is converted into a connective tissue nodule which contracts more and more.

**The condition of branches** given off at the seat of an aneurysm is a matter of great importance. These vessels are frequently obstructed, and there are various ways in which this may come about. The atheromatous process may occur to such an extent around the orifice of a branch as to narrow or even obliterate it. This is most frequent in small arteries like the intercostals, but is not uncommon in larger branches. Again, the coagula may come to overlie the aperture, already narrowed, it may be, by atheroma. Further, it will be apparent that, as an aneurysm enlarges, especially a sacculated one, it will often drag on and contort vessels whose apertures are in or near its walls. The aperture may thus be reduced to a fissure, and the edge may be so placed as to valve the aperture. This is particularly the case in the sacculated aneurysms of the arch of the aorta. Sometimes by the enlargement of an aneurysm the aperture of the branch is to be found at the summit of the aneurysm. In that case the aperture may be obstructed in one of the ways already described, but it not infrequently



remains at least partially free. The coagula may even be tunnelled in order to allow the current to flow into the branches (see Fig. 173). Another mode of closure is by the pressure of the aneurysm itself on the branch beyond the aperture. If a branch be closed in any of these

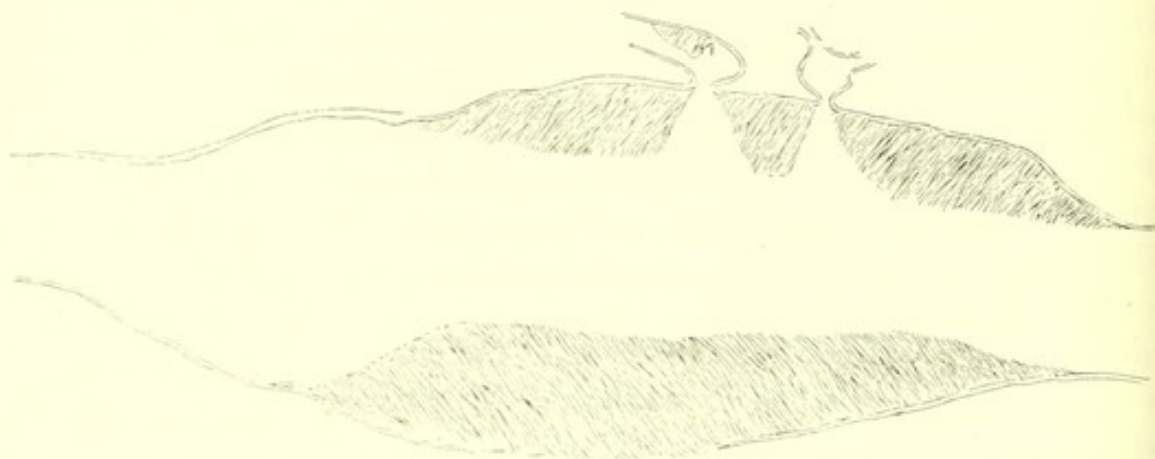


Fig. 173.—Aneurysm of the abdominal aorta with clots tunnelled so as to allow the blood to reach the branches. The general lie of the stratification of the clots is shown. The coeliac axis and superior mesenteric artery are seen to be narrowed at their orifices. The posterior wall of the aneurysm is absent where it impinged against the vertebræ. Half the natural size.

ways it becomes the seat of a thrombus, and in the usual way becomes converted into a solid cord. In regard to the condition of branches it is to be observed that the aneurysm may, as it were, be continued into the branch, the first part of the latter being dilated along with the aneurysm.

**Effects on the heart.**—Aneurysms affect the heart somewhat variously. It is frequently depressed by the mere presence of the aneurysm at its base. Room must be afforded for the increasing tumour, and as the position of the aorta forbids much movement upwards, the heart is pushed somewhat downwards. The extent of displacement will, of course, depend on the position and size of the aneurysm. Besides this, the heart often undergoes enlargement, especially the left ventricle. It is clear that in distending the aneurysm a considerable amount of the force of the left ventricle is wasted, and on principles already considered the ventricle must act more powerfully; the hypertrophy is therefore compensatory.

**Effects on parts around.**—The effects of the aneurysm on other parts will depend on the amount of pressure exercised, and the nature of the structures involved. An aneurysm often **presses on nerves**, causing primarily irritation of them and sometimes ultimately interruption and consequent loss of function. Thus they often produce violent pains, and even symptoms of angina pectoris when the nerves of the heart are pressed on. By irritation of the recurrent laryngeal nerve



they may cause spasm of the laryngeal muscles, or by destroying the nerve lead to their paralysis.

The **various canals within the chest**—the trachea, bronchi, œsophagus, venous stems—are often narrowed or completely obstructed, so that a great variety of symptoms are produced. When an aneurysm meets with a firm structure such as bone, it **erodes** it, as we have already seen. Cartilage resists more than bone; and in the case of an aneurysm coming against the vertebræ, we often find the bodies of the vertebræ deeply eroded, while the intervertebral cartilages stand prominently out between them (see Fig. 174).

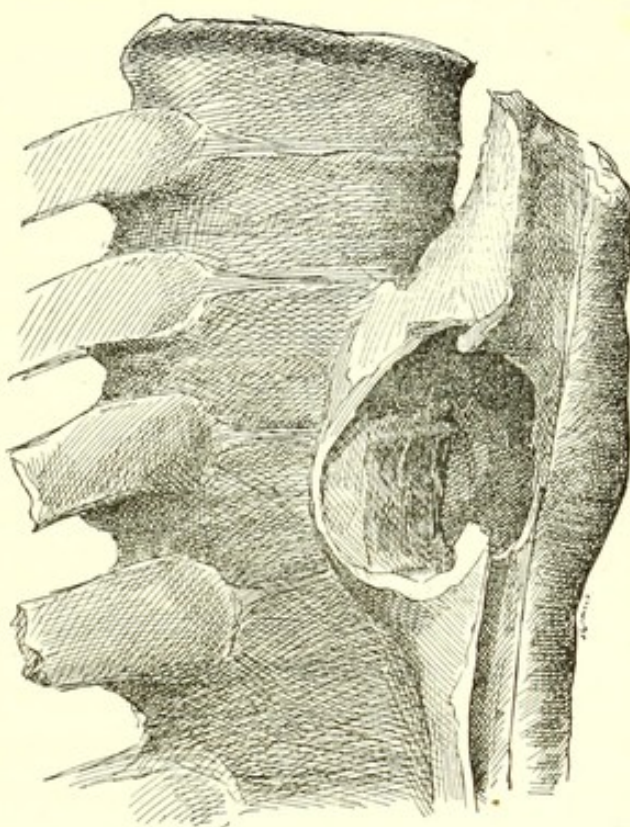


Fig. 174.—Aneurysm of aorta eroding vertebræ. The intervertebral cartilages are seen to remain prominent.

We may find also in an aneurysm advancing to the front of the chest the sternum and osseous ribs much eroded, while the cartilaginous ribs, almost isolated, are very little destroyed. In these cases it will be seen that as the dense structures are exposed in the aneurysm, the proper wall of the latter is, by so much, deficient.

**Terminations of aneurysms.**—Aneurysms have somewhat various terminations, much the commonest being unfortunate. Sometimes, although very rarely, a **spontaneous cure** occurs. The artery is sometimes pressed on by the bulging aneurysm itself, and obliterated. Or the aneurysm having burst, the blood collecting outside, by its pressure, obliterates the aneurysm. It is well known that obliteration of the artery is often effected by **ligaturing** it. If the artery is obliterated, the aneurysm fills up with clot, and by and by organization occurs, and the aneurysm is cured.

But the majority of aneurysms are beyond the reach of surgical interference, and for the most part they continue enlarging till they lead to a **fatal issue**. This may be from the effects of **Pressure on parts around**, or upon the heart, so that death may result from hyperæmia and œdema of the lungs, from general venous engorgement and œdema, from marasmus, and so on.

The aneurysm sometimes ends by **Rupture**, which occurs in various



directions according to the circumstances of the aneurysm. It may here be said that the probability of rupture is by no means in proportion to the size of the aneurysm. A large aneurysm, by the amount of pressure it exercises, will probably, by its irritation, and by causing counter-pressure from displaced organs, produce sufficient support, unless it comes actually to the surface of the body or bulges into a cavity. When rupture occurs it is either into a cavity of the body, as the sac of the pericardium, the pleura, an auricle of the heart, etc.; or into a canal, as the trachea, bronchus, or another blood-vessel, such as the pulmonary artery; or into the substance of an organ, as the lung or brain; or among the muscles or connective tissue of a part. The mode in which the rupture comes about varies somewhat. The aneurysm may rupture into a cavity or on the surface of the body for want of support to its wall, a tear occurring by and by in the sac. Or the aneurysm may undermine a mucous membrane or the skin, and produce a necrosis, rupture occurring on the separation of the slough. Or the aneurysm may induce inflammation of the mucous membrane, and the softened tissue may give way. Again, in the case of an aneurysm meeting an osseous structure a gap may occur in the wall of its proper sac, and the edge of the gap may become detached from the bone, and so occasion a rupture. The rupture is not usually at the very first fatal; there is, to begin with, a slight leakage of blood through irregular apertures which get occasionally closed with blood-clot. But from some accidental increase of blood-pressure or other cause the aperture is enlarged, and a fatal hæmorrhage is the result. Or death comes from exhaustion due to the prolonged drain and the interference with function that the aneurysm otherwise produces.

Aneurysms of the **Ascending aorta** are most liable to burst into the pericardium, but they also rupture into pleura, lung, œsophagus, vena cava superior, pulmonary artery, auricle, ventricle, or externally. Those of the **Arch of the aorta** rupture into the trachea or bronchi most frequently, but also into pericardium and on other neighbouring surfaces. The aneurysms of the **Thoracic aorta** rupture chiefly into the pleuræ, but also into the œsophagus, bronchi, and lungs. Those of the **Abdominal aorta** mostly rupture behind the peritoneum, and enormous quantities of blood may accumulate beneath that membrane, bulging it and the abdominal contents forwards. There may be a subsequent tear of the peritoneum, and blood is, in that case, present in the abdominal cavity. These aneurysms also rupture into the left pleura, inferior vena cava, lungs, colon, pelvis of kidney, and posterior mediastinum.

**False and Spurious aneurysms.**—There are certain lesions to which the name aneurysm is commonly given, but which do not accord with the description given above.

The **Cirroid aneurysm** and the **Aneurysm by anastomosis** are somewhat allied forms. The cirroid aneurysm is sometimes called varix



arterialis, and the name is suggestive of the condition presented. A portion of artery with its branches becomes elongated and widened, and the vessels become convoluted like varicose veins. Sometimes the dilatation extends to the corresponding capillaries and veins. This form of aneurysm occurs most frequently in the arteries of the scalp and face, especially the temporal and occipital, and the enlarged and tortuous vessels are to be felt under the skin. The aneurysm by anastomosis is almost an arterial vascular tumour. There is enlargement and lengthening of a large number of small arteries, with probably newformation of arteries, and the mass of vessels can be felt like pulsating worms under the skin. The enlargement may extend to the capillaries. The affection forms a distinct growth of a bluish red colour, and with a somewhat granular surface. Its most frequent seat is the scalp.

The **Traumatic aneurysm** arises in consequence of an injury to a vessel. An injury has been sustained, and after a longer or shorter period an aneurysm appears. The mode in which such aneurysms arise varies. In some cases an artery is wounded, and the blood makes a cavity for itself, constituting a spurious aneurysm. The cavity remains in communication with the artery, and forms a kind of aneurysmal sac. This is not a common result of wounds of arteries, as these usually close, but it does occur, and most readily when the wound has been an oblique or longitudinal one. It may result from a wound penetrating from the surface, but of such a form as to prevent the escape of the blood from the surface, or it may result from a broken bone tearing the coats.

But there are some cases of traumatic aneurysm which are more slow of formation and in which it is not probable that any distinct tear through the whole coats has occurred. In the fully developed aneurysm it is impossible to trace the exact mode of origin, but it is probable that in many cases a fractured bone projected against a vessel injures or even ruptures the middle coat, and possibly the internal as well. We know that in applying a ligature those two coats give way, and we can conceive a violent force applied to the wall producing the same result. A simple blow may act in a similar way, but it will do so the more readily if the middle coat be already brittle from calcareous infiltration. Many of the aneurysms of the femoral and popliteal arteries are referred to blows and injuries; but these are not all to be regarded as pure traumatic aneurysms, as the arterial coats are frequently so altered by atheroma and calcareous infiltration as to make the injury merely the determining cause.

Traumatic aneurysms, as appears from what has been said as to their



origin, have usually less defined sacs than spontaneous ones. This is especially the case in those which arise directly from wounds.

The **Dissecting aneurysm** is a somewhat interesting form, arising apparently by rupture of the internal coat, and partially of the middle. It sometimes arises from an injury, and is thus a traumatic aneurysm, but it occasionally occurs spontaneously, and there is no form of aneurysm which is so frequently multiple as this. The author has met with a case in which there were four separate dissecting aneurysms on the aorta and its branches. It may be inferred from this fact that in these cases there is often an abnormal brittleness of the internal coat, apparently not due to atheroma or any other definite disease of the coat, but an inherited or acquired brittleness. The internal coat is torn through, and the blood passes among the fibres of the middle coat. It is not that the middle coat is dissected up from the internal coat, but the layers of the middle coat itself are separated, and the blood lies between an external and an internal layer

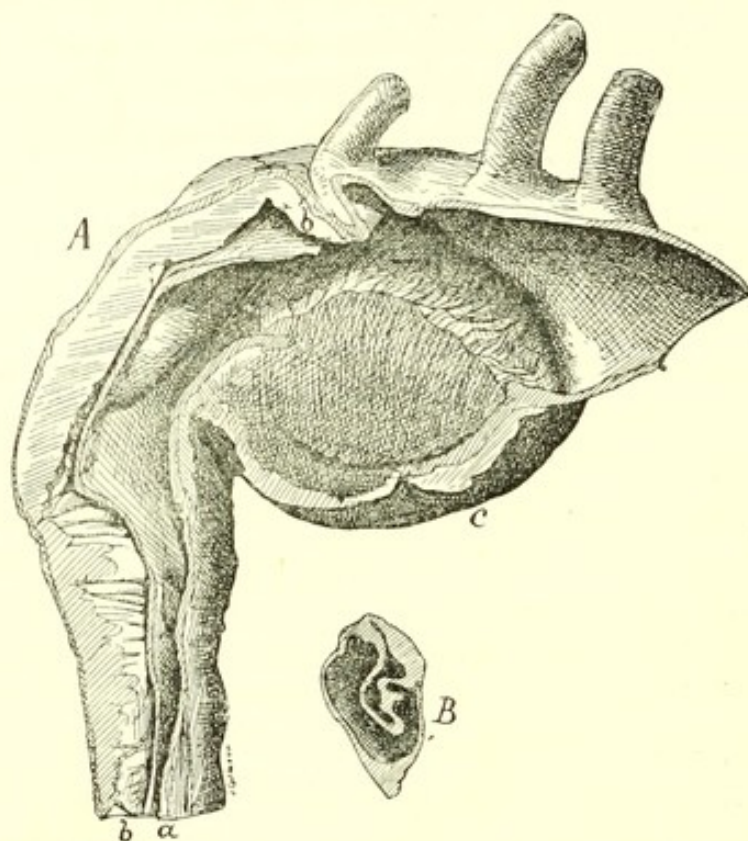


Fig. 175.—Dissecting aneurysm of aorta. In A, a longitudinal section shows the two tubes, the natural calibre *a*, and the aneurysmal one *b b*, the latter largely occupied by clot. The hollow of the arch is largely occupied by clot at *c*, there being here a partial rupture of the coats. B is a transverse section. The proper calibre is greatly reduced, and the separation has extended round about three fourths of the circumference. (From a preparation in the Western Infirmary Museum.)

of the middle coat, as in Fig. 176. The splitting up of the middle coat may be carried a considerable distance, and after passing along in the wall of the vessel the blood may make another tear in the interna coat



and pass back into the vessel. The aneurysm may thus come to have two apertures, and a condition may occur as if the vessel were formed of a double tube, divided longitudinally by a septum composed of the internal and a part of the middle coat (see Fig. 175). The circulation may even be carried on to a great extent through the aneurysm, the blood passing in at the original aperture and out at the secondary one. In that case the proper calibre of the artery may become considerably narrowed. The approximation of such an aneurysm in character to the regular channel is still further increased by the fact that the aneurysm by and by acquires an internal lining resembling the internal coat. It is a homogeneous membrane of about the thickness of the internal coat, so that in a section it may be difficult, even under the microscope, to say whether a particular place is wall of vessel or wall of aneurysm (see Fig. 176). In the case already referred to, one

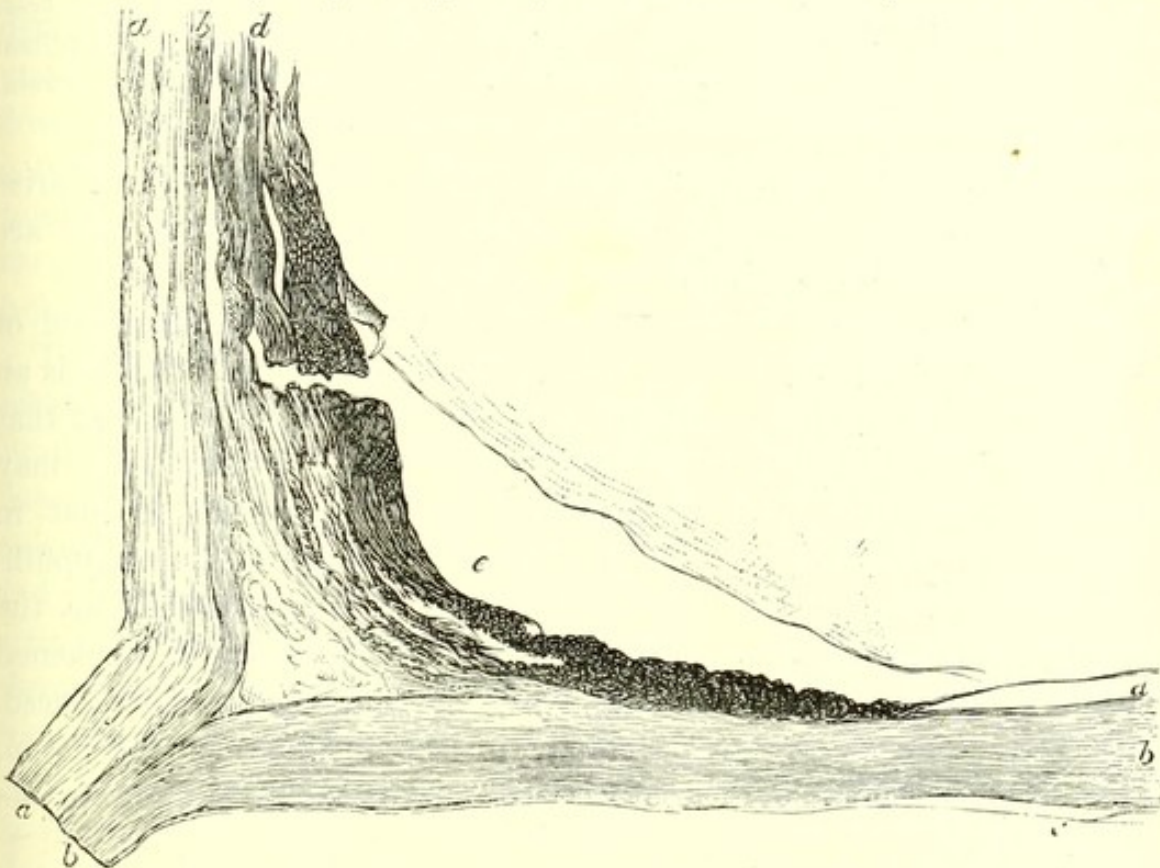


Fig. 176.—Section of portion of a dissecting aneurysm of the aorta. At the left there is the entire wall of the artery, *a* being external, *b* middle, and *c* internal coat. The middle coat is split so as to form the aneurysm. At *d d*, a kind of internal coat is formed to the aneurysm, but this is the seat of fatty degeneration at *e*.  $\times 16$ .

of the aneurysms began in the thoracic aorta, and was continued down into the iliac arteries. There were two apertures, one in the thoracic, and the other far down in the abdominal aorta, and between these the aneurysm had evidently carried on the circulation to a greater extent than the natural channel.

Dissecting aneurysms being confined to the wall of the vessel will appear externally as thickenings or dilatations of the wall, and they



may produce bulging or bagging of the vessel to a remarkable degree. Occasionally, a dissecting aneurysm tears through the remaining layers of the middle coat and the external coat, and so a spurious aneurysm is attached to a dissecting one.

**Varicose aneurysm** is a term applied to the case in which an artery and a vein acquire a permanent abnormal communication. It may arise spontaneously or traumatically. This form used to be not uncommon at the bend of the elbow, when, in performing venesection, the lancet wounded artery and vein together. It occurs spontaneously when an aneurysm ruptures into a vein. In some cases there is an actual aneurysmal sac between artery and vein, and to this form the name **Varicose aneurysm** is more properly applied. But the communication may be direct, and while the artery is little dilated the vein is enlarged, tortuous and pulsating. To this form the name **Aneurysmal varix** is applied. The interference with the venous current produced by the violent injection of the arterial blood, especially if it be suddenly produced, may lead to serious venous engorgement and œdema, which may even have a fatal issue. Spontaneous communications of this kind have been observed between the aorta and venæ cavæ, between the crural and popliteal arteries and the corresponding veins, between the splenic artery and azygos vein, and between the internal carotid and the sinus cavernosus.

8. **Syphilitic and Tubercular affections of arteries.**—These arise usually by extension of the infective process to the arteries and are generally part of a larger and more pronounced lesion.

**Syphilitic affections** of the arteries occur in the neighbourhood of gummata, more especially of the brain and its membranes. There is an extension of the specific inflammation to the walls of the artery so that the coats are infiltrated with round cells. The internal coat may undergo great thickening so that the condition resembles that in obliterating endarteritis (see Fig. 51, p. 196). A more unusual syphilitic lesion is a similar localized inflammation of an artery without the presence of a gumma in the neighbourhood. The artery is thickened and has a cartilaginous consistence, while the calibre is greatly reduced. This condition, which occurs chiefly in the arteries of the brain, resembles atheroma of these vessels, but is more limited in its distribution.

There is considerable doubt as to the existence of a special gummatous endarteritis as described by Heubner, and the influence of syphilis in affecting the arteries has been exaggerated. A lesion of an artery is not to be regarded as syphilitic unless it be either directly connected with a syphilitic lesion, or associated with syphilitic disease elsewhere.

**Tuberculosis** of organs not infrequently extends to their arteries. The wall of the artery is infiltrated with round cells and its tissue opened out and softened. There may result aneurysm or even hæmorrhage although the advancing tuberculosis usually closes the artery by thrombosis.



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## II.—DISEASES OF THE VEINS.

1. **Thrombosis.**—This is of very frequent occurrence in veins. The full description of this process in a previous section applies especially to the veins. As the blood in the veins is normally at a low pressure and flows slowly, it is readily brought to a standstill, especially in the pouches of the valves. Hence, any cause which produces prolonged passive hyperæmia or weakness of the circulation is apt to induce thrombosis. Thrombosis is also a result of inflammation of veins, such inflammation being mostly septic.

Thrombosis induces an **Inflammation of the wall** of the vein, whether the thrombus be specially irritating or not. In the case of a simple thrombus its presence induces a chronic inflammation which extends through the wall of the vein to its adventitia and surrounding connective tissue. There are thus **Adhesions** formed between the vein and its sheath, and these may extend to the sheath of the neighbouring artery, causing considerable matting around. A vein containing an old thrombus is therefore, as a rule, somewhat difficult to dissect clean from the surrounding tissue.

2. **Inflammation of veins. Phlebitis.**—The simplest form of phlebitis is that just referred to in which a thrombus causes a chronic inflammation of the wall of the vein. We may also have an inflammation of the wall from inflammations in the neighbourhood which may or may not induce thrombosis secondarily.

**Septic phlebitis or Thrombo-phlebitis** is a condition formerly of frequent occurrence and much discussed in medical literature. This condition mostly results from the entrance of septic matter into a vein, whose mouth lies open in a wound, but it may occur from the extension of a septic inflammation along the sheath of the vein and thence inwards.

Septic matter introduced into the calibre of a vein induces coagula-



tion of the blood which, from the circumstances of the case, is usually already stagnant. The thrombus thus formed is not a simple one, but contains septic microbes, namely, pyogenic micrococci. There is thus a strong irritant inside the vein, and this induces an acute inflammation of its wall. There is an exudation of leucocytes and of blood-plasma, which accumulate in the interstices of the wall and infiltrate the external and middle coats. The exudation rapidly becomes purulent, and the coats are still further opened out. Viewed from within there may be little collections of pus visible beneath the internal coat like pustules. The pus passes into the calibre of the vein, and as the thrombus softens the vein will often come to be filled with pus. The softening of the thrombus is partly the effect of the pus, but is partly also from septic decomposition. The contents of the vein, by the softening of the thrombus, are allowed to pass into the circulation, although fresh thrombosis may from time to time shut them off. The phenomena of **Pyæmia** result from the carriage of septic matter by the blood and the consequent infection of distant parts. From the veins, the most direct connection is with the lungs where septic embolism is set up. But there is frequently an extension beyond the lungs by the systemic arteries to heart, kidneys, etc.

**Literature.**—HUNTER, Works, edited by Palmer, 1837, iii. 581; HODGSON, Dis. of art. and veins, 1815; ROKITANSKY, Handb. d. path. anat. ii., 1844; VIRCHOW, Ges. Abhandl., 1857; EBELING, Ueber phlebitis, 1880; PANTON, Uterine phlebitis, Glasg. 1840; LEE, Origin of infl. of veins, 1850.

3. **Varix or Phlebectasis.**—Varix is dilatation of veins just as aneurysm is dilatation of arteries.

**Causation.**—We saw that some weakening of the wall is always necessary to the occurrence of aneurysm, but no such condition is required in the production of varicose veins. The walls of veins are already thin, and the blood-pressure within them is low. In the case of the veins of the skin we are familiar with the fact that, as a rule, the blood-pressure is not sufficient to overcome the atmospheric pressure to any considerable extent and the veins are flattened or partially collapsed, a slow and weak current passing through them. The circulation must be considerably excited, or the veins must be obstructed in order to make them stand out as cylinders under the skin. Such being the case it may be conceived that the thin-walled veins, accustomed to a slight blood-pressure, will readily dilate when exposed to an increased blood-pressure. Varix is always due to some cause which is calculated to increase the blood-pressure within the veins, such as obstruction of the veins by thrombi, tumours, the gravid uterus, passive hyperæmia from heart disease, and so on. It should be added, however, that the



tendency to varicose veins is often transmitted by inheritance, so that there may be a certain weakness of the walls allowing of more ready dilatation from comparatively trivial causes.

**Character of the changes.**—In the early stages of varix, as we have frequently an opportunity of seeing it in the legs, there is simply an exaggeration of this natural dilatation above the valves. When we stand erect the column of blood in the veins of the legs is, as it were, supported at each valve, and the downward pressure tells on the valve and the portion of the vessel forming the valvular sinus. Hence this part of the vessel is the first to dilate when the blood-pressure is increased, and the first sign of varix is an exaggeration of the knotted state of the vein. At the outset each valve forms a kind of fixed point, the dilatation occurring at its level, and diminishing as the valve next above is approached. As the dilatation increases and extends up from the valve, the vein increases in length as well as in calibre, and in order to be accommodated it begins to form **Curves or Convolutions**. Thus begin those sinuosities which are so characteristic of varicose veins, and which tend to increase as time goes on. The dilatation of the vein has a tendency to render **the valves incompetent**, and this occurs all the sooner as the region of the valves is, as we have seen, the part where the pressure is most exercised. When the valves become incompetent the pressure tells much more on the walls of the veins, as the column of blood is now arrested at longer intervals. The pressure acts most on the dependent parts of the sinuosities, and will increase the projection of these. In this way we may have wide sinuses with their convexity downwards, and in some cases even diverticula or pouches projecting from the veins. In these exaggerated dilatations the blood stagnates greatly, and it is not uncommon for **Thrombosis** to occur. The vein is obstructed more or less completely by the thrombus, which may organize. On the other hand, the latter often dries in and becomes impregnated with lime salts. In this way varicose veins frequently become the seat of vein-stones or **Phleboliths**.

**Chronic inflammation** is common in the tissues around varicose veins wherever they may be, and so in the skin we often have very persistent eczema with a brown coloration of the skin, which may be referred to hæmorrhage by diapedesis from the hyperæmic vessels. The skin is indurated and thickened, and this along with the swelling of the veins may be so great as to produce an appearance like that in elephantiasis. Ulceration is often induced, and the ulcer is sometimes deep and sluggish and may persist for years.

A varicose vein **may burst**. It may be opened by the ulcerative process, or it may come to the surface and by its increasing dilatation at last



rupture. In the case of the leg the results are sometimes exceedingly serious. As one effect of the dilatation is to render the valves incompetent, the veins of the entire leg may come to be virtually devoid of valves. But the veins in the abdomen are normally devoid of valves, and so it may happen that from the lower leg up to the heart there may be a single column of blood without an arresting valve. If now a vein rupture in the leg, the whole system up to the heart may be, as it were, tapped, and if the person is in the erect posture a fatal hæmorrhage may result. The hæmorrhage will cease if the person lies down, but cases have been known in which an immense amount of blood has been shed into the boots without warning.

**Hæmorrhoids** are varicose veins of the lower rectum. At the lower part of the rectum a network of small veins surrounds the bowel, lying immediately under the mucous membrane. These veins communicate with the inferior mesenteric vein which is a tributary of the portal, and with the internal iliac which is a tributary of the vena cava. Any obstruction in the portal circulation, or abnormal pressure within the abdomen is apt to induce dilatation of this hæmorrhoidal plexus. The dilated veins push the mucous membrane before them and protrude as bluish flat nodules either immediately within or without the anus. They may assume considerable dimensions. The knots are composed originally of a congeries of small convoluted veins (Fig. 177 *B*), which may communicate so as to form a kind of cavernous tissue, but sometimes there are large ones (Fig. 177 *A*).



Fig. 177.—Hæmorrhoids of rectum, in section, natural size. *r* is internal surface of rectum, and *m* the mucous membrane continued over the hæmorrhoid. At *m'* is represented the muscular coat, the dilated veins being in the submucous tissue. At *A*, the veins are few but much dilated, one especially so, just beneath the mucous membrane. At *B*, the dilated veins are more numerous, but smaller. Both occurred in the same case. (VIRCHOW.)

Through time the mucous membrane undergoes alterations. The piles at each movement of the bowels are exposed to mechanical irritation, and so the mucous membrane is almost constantly in a state of catarrh. Then at intervals, when the blood-pressure is unusually great by reason of



an attack of inflammation or otherwise, the veins rupture, and so there is bleeding both in the substance of the mucous membrane and on to the surface. The hæmorrhoids thus get greatly altered. They become condensed from inflammatory newformation of connective tissue. They sometimes become the seat of collections of blood, which may form blood-cysts. Phleboliths may also form in the veins. More severe inflammations sometimes occur, resulting in abscesses, fissures, fistulas, ulcers.

**Varicocele** is a dilatation of the veins of the spermatic cord and the external scrotal veins. It affects in a greatly preponderating proportion the left side, the explanation of this being apparently that the left spermatic vein has a circuitous course, and enters the renal vein at a right angle, while the right opens into the lower vena cava. The varicosity generally begins at the external inguinal ring, and extends downwards as far as the testicle. There is not infrequently atrophy of the testicle, and sometimes hydrocele or hæmatocele.

Varix may occur in other veins, such as those of the neck of the bladder and prostate. It is also met with in the female in the vesical plexus and veins of the vagina, and this may be combined with varix of the broad ligament. Varicosity also occurs in the veins of the dura mater.

4. **Newformations in veins.**—**Syphilis** rarely attacks veins, but a gummatous inflammation has been observed in the portal vein and in the umbilical vein in new-born children. Leprosy may attack the large veins of the extremity.

**Tuberculosis** affects veins in organs which are the seat of this process, and produces results similar to those in arteries. As the walls of veins are thin the process more readily penetrates to their calibre, and the tubercular bacillus may thus find entrance to the blood. Tuberculosis of veins is the most frequent cause of general tuberculosis, as already described.

Primary tumours of veins are excessively rare. A few cases of small **Myomata** have been described, and a larger tumour which was regarded as a myo-sarcoma (Perl). Tumours not infrequently penetrate into veins, on account of the thinness of their walls, but do not produce tumours in them.

**Literature.**—BRODIE, Lectures on Path. and Surgery, 1846; NUNN, Varicose veins and ulcers, 1852; CHAPMAN, On varicose veins, 1856; GAY, Varicose dis. of lower extrem., 1868; A. COOPER, On spermatocele, Guy's Hosp. Rep. vol. iii.; VIRCHOW, Virch. Arch. iii.; COHNHEIM, do., xxxvii.; KÖSTER, (Phlebectasis in intestine) Berl. Klin. Wochenschr., 1879; LESSER, Virch. Arch. ci.; EPSTEIN, do., cviii.; PERL, Virch. Arch. liii.



## SECTION I.—CONTINUED.

C.—DISEASES OF THE LYMPHATIC SYSTEM, SPLEEN,  
AND BONE-MARROW.

- I. THE LYMPHATIC VESSELS.—*Inflammations. Dilatations. Obstruction of the thoracic duct. Cancerous obstruction.* II. THE LYMPHATIC GLANDS.—*Their structure, and general pathology. Pigmentation leading to induration. Degenerations. Acute lymphadenitis, Bubo. Chronic lymphadenitis. Tuberculosis, primary and secondary. Syphilis. Tumours.* III. THE SPLEEN. *Its structure. Malformations and Malpositions. Active congestion and inflammation; the acute splenic tumour. Chronic inflammation; the chronic splenic tumour. Passive hyperæmia. The embolic infarction. Rupture. Amyloid disease; the sago spleen; the lardaceous or waxy spleen. Tuberculosis, syphilis, and tumours.* IV. SOME AFFECTIONS OF THE BONE-MARROW.

## I.—THE LYMPHATIC VESSELS.

WE have already had occasion to observe that the lymphatic system is intimately related to the blood-vascular system, and may be regarded as a part of it. We have also seen that this system has close relations to the connective tissue, the spaces in the latter being lymph spaces lined with endothelium and having direct connections with the lymphatic vessels. Wherever there is increased exudation from the blood-vessels the serous spaces and lymphatic vessels are distended, and the current through them increased. We have seen this to be the case in inflammations and œdemas.

**Inflammation of the lymphatics. Lymphangeitis.**—The lymphatics are liable to inflammations of a more independent kind, and warranting the name lymphangeitis. In order to such inflammation there must be in the vessels some irritant, and the irritant is similar to that which we find in the veins in suppurative phlebitis, namely a septic poison. In connection with wounds which have been exposed to decomposing juices, such as dissecting wounds, we sometimes find the course of the lymphatic vessels marked by red streaks in the skin. These represent inflammation of the lymphatics and surrounding connective tissue. The inflammation not infrequently goes on to suppuration, so that abscesses form at intervals in the course of the vessels or in



the lymphatic glands. The inflammation is due to the presence of pyogenic microbes which propagate inside the vessels and so cause the inflammation to extend.

**Erysipelas** and other phlegmonous inflammations of the skin are inflammations primarily of the lymphatics and of the serous spaces which form the radicles of the lymphatics. Microbes are present in the lymphatics of the skin in such cases (see under Diseases of the Skin) and they may be so numerous as to render the lymphatic vessels as well as the serous spaces peculiarly prominent, when the sections have been stained by aniline dyes. The phlegmonous inflammation of the parotid which occasionally occurs in cases of septic wounds is probably of a similar nature. Epidemic parotitis (mumps) may perhaps be placed in the same category.

We have already seen that **Elephantiasis** frequently begins in recurrent attacks resembling erysipelas, in which the lymphatic vessels are obviously engaged. There are often red streaks passing up the limb, and the lymphatic glands may be enlarged. The irritation here is slighter and more chronic. The inflammation results in great thickening of the connective tissue, but there is often along with it dilatation of the lymphatic vessels, so that, when the part is cut into, an abundant lymphatic fluid exudes which sometimes contains oil.

**Lymphangiectasis.**—The lymphatic vessels are occasionally the seat of *dilatation*. Obstruction of a lymphatic stem may produce a varicose dilatation of the vessels, and even their rupture. We have already seen this to be the case when the lymphatics are obstructed by the ova of the *filaria sanguinis*, resulting in **Lymph-scrutum**, **Chyluria**, etc. In the enlargement of the tongue called **Macroglossia**, which is frequently congenital, there is commonly a great enlargement of the lymphatics, which may form considerable cysts. The dilatation here appears to be due to obstruction of the lymphatics from interstitial inflammation of the connective tissue, although a different origin has been asserted.

There is sometimes a localized dilatation of many lymphatics so as to form a distinct vascular tumour. These tumours are usually cavernous in structure and are classed as **Cavernous lymphangiomas**. Sometimes they develop actual cysts, which are filled with serous fluid, and are designated **Hygromas**.

**Infective and other tumours.**—**Tuberculosis** of the lymphatics arises by extension from neighbouring parts. This is sometimes of importance as the tuberculosis may pass on by the lymphatics to the blood, more particularly if the thoracic duct be affected. (See under General Tuberculosis, p. 207.)

We sometimes find lymphatic vessels dilated in the neighbourhood



of a **Cancer**, the material in the vessels being derived from the tumour. This occurs mostly in the case of cancers which are undergoing softening. It is most frequently seen in connection with mammary cancers, where the lymphatics may have the form of firm cylinders as large as quills, filled with white material. The same thing is seen in the case of secondary cancer of the lung, where there are nodules immediately under the pleura. The subpleural lymphatics around the nodule are found, as it were, injected with white material. So is it sometimes in the capsule of the liver when cancerous nodules are near the surface. These naked-eye appearances indicate how prone cancerous material is to pass into the lymphatics, and, under the microscope, at the margins of a growing cancer the lymph-spaces may sometimes be found filled with the epithelial masses. (See under Cancer, p. 266.)

**Obstruction of the thoracic duct.**—This is not an infrequent accident, arising from various causes, such as pressure of tumours, aneurysms, etc., in its neighbourhood, but more particularly from thrombosis of the jugular and innominate veins. Experiments on animals show that ligation of the duct leads to dilatation and sometimes to rupture of its dilated origin in the abdomen, which forms the cysticerna chyli (Cooper). In man there may be a similar rupture, either in the abdomen or pleura, leading to accumulation of fluid in these cavities. As the fluid in the thoracic duct contains fat, the exudation in the peritoneum or pleura will be chylous (see Chylous Ascites and Hydrothorax). Rupture is, however, only an occasional result of obstruction of the duct, as the anastomosing connections may compensate, so that there may even be comparatively little dilatation.

**Literature.**—VELPEAU, *Arch. gén. de méd.*, 1835, viii.; BRESCHET, *Le système lymphatique*, 1836; CARTER, (Lymphangiectasis and elephantiasis) *Trans. of Med. Soc. of Bombay*, 1861; ISRAEL, *Ueb. Lymphangioma*, 1885; BRADLEY, *Injuries and dis. of lymphatic system*, 3rd ed., 1875; COOPER, *Med. records and researches*, 1798; BÖGEHOLD (Injuries of thor. duct), *Arch. f. klin. Chirurg.*, xxix. 443; COATS, *Museum Catal. of Western Infirmary*, pp. 55, 56 (Obstr. of thor. duct from thrombosis of jug. vein, 2 specimens).

## II.—THE LYMPHATIC GLANDS.

All lymphatic vessels connect themselves directly with lymphatic glands so that no lymph passes into the circulation without first traversing a lymphatic gland. Dissolved substances introduced into the serous spaces of the skin and subcutaneous tissue are rapidly carried up the lymphatics and through the glands into the general circulation; this is matter of experience every day in the subcutaneous injection of medicinal agents. Granular material, however, does not pass through the glands, which form a filter for all solid particles.



The apparatus by which the **Filtration of the lymph** is effected is illustrated in Fig. 178. When the afferent lymphatic vessel coming from the periphery reaches the gland, it first forms a plexus in the capsule. On the other hand the efferent vessel emerges from a plexus of vessels at the hilus, and this plexus is connected with the medullary portion of the gland. In passing from the afferent to the efferent vessel the lymph traverses a series of sinuses *c, c*, in figure. The sinuses are not clear spaces but contain a reticulum or mesh work shown in the figure. These sinuses

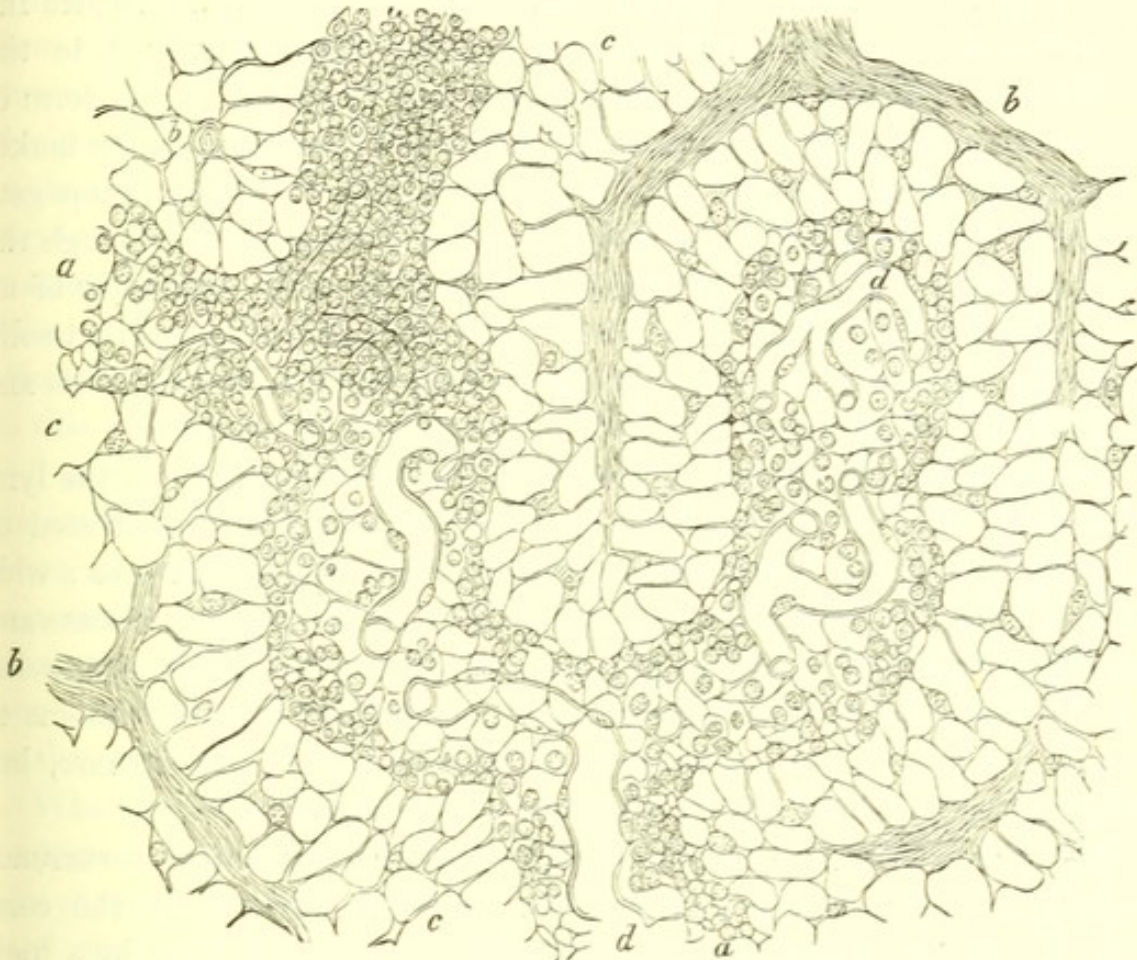


Fig. 178.—Section of normal lymphatic gland; *a, a, a*, follicular or lymphoid cords; *c, c*, lymph sinuses; *b, b*, connective-tissue trabeculae; *d, d*, blood-vessels.  $\times 300$ . (RECKLINGHAUSEN.)

surround the proper glandular tissue which is in the form of rounded follicles or cords composed of densely packed lymph cells (*a, a*, in figure). The lymph first passes from the afferent plexus into the sinuses of the cortex, then into those of the medullary part, where originate the radicles of the efferent vessel. The sinuses being occupied by the fine reticulum will catch solid particles brought to the gland, and such particles will therefore be most readily caught at the cortical portion of the gland.

In the normal gland, according to the researches of Flemming and Baumgarten there is an active newformation proceeding by means of **Karyomitosis**. This occurs at a number of centres in the cortex whose cells are less deeply stained than those elsewhere, but in which the nuclear figures of karyomitosis are visible. There is also nuclear division in other parts, but to a less degree. These appearances probably indicate the newformation of leucocytes.

In the operation of **Tattooing** granular pigment, such as Indian ink or vermilion, is introduced into the skin. Part of it remains perman-



ently in the corium, but part is conveyed up the lymphatics and is caught by the glands, where it remains fixed.

So it is in pathological conditions. The lymphatic glands are very frequently associated in morbid processes which are occurring in parts in which the vessels passing to the glands take origin. Dissolved and solid materials are thus conveyed to the glands, and according to the nature of the material will it affect the gland or not. In **Septic inflammations**, for example, the microbes are often conveyed to the glands and produce inflammation there. As these are in the form of fine particles they are arrested at the gland, and the inflammation makes a pause there only to proceed further, if the organisms propagate onwards. But even in the case of **Simple inflammations**, although the products are for the most part dissolved, still they are frequently of an irritating nature and the glands inflame. We have seen also that solid material from cancers is often conveyed along the lymphatic vessels and is caught at the glands.

In all these cases the material may produce no irritation in the lymphatic vessels along which it is being conveyed. If concentrated or very virulent it will do so, but it often does not, and in that case a wide interval of healthy structure intervenes between the primary disease and the lymphatic gland. Hence it is common to speak of the disease being conveyed to the glands by metastasis. Indeed, most of the lesions to which the glands are liable are secondary to diseases elsewhere, but there are some which may be regarded as idiopathic.

**Degenerative changes** in the glands are not of common occurrence apart from tuberculosis and syphilis. **Amyloid disease** is the commonest form of degeneration, but it does not usually occur in a high degree, and it is only occasionally that the glands are enlarged and rendered firmer by the degeneration. For the most part it is detected only when the glands are examined microscopically with the aid of iodine or methylviolet. The disease appears to attack mainly the reticulum of the gland, and one can sometimes make out the swollen hyaline network, or even detect how the normal reticulum swells out into the degenerated network. In old persons a **Colloid or Hyaline degeneration** is met with. The gland shows a series of alveoli filled with colloid material, or the blood-vessels and trabeculae of the gland become hyaline, the appearance resembling that in amyloid disease but without the reaction.

**Pigmentation of the glands.**—We have seen how pigment artificially introduced into the lymphatics, is arrested at the glands. Similarly carbonaceous pigment is carried from the lungs and deposited in the bronchial glands. Again, in cases of extravasation of blood, or in



inflammations with considerable hæmorrhage, the red corpuscles are conveyed, sometimes in large numbers, to the glands, which may thus be deeply coloured. The dissolved colouring matter in hæmorrhages may also be conveyed and stain the glands.

The pigment is carried, in part at least, by the lymphoid cells, which are to be found in the lymphatic fluid. Arrived at the gland, the pigment is first caught by the stellate cells which form the reticulum in the sinuses of the cortex, and hence in the earlier stages we may have the sinuses marked out by the pigmentation. If it continues to arrive at the gland, it extends into the follicles and finally the whole gland may be deeply and uniformly coloured. When present in such quantity the pigment, although generally non-irritating, produces a **Chronic inflammation**. There is increase of the connective tissue forming the trabeculæ, and a corresponding atrophy of the glandular follicles. The affected gland in addition to the pigmentation is enlarged and hard, and in this state it is probably to a large extent **stuffed up** by the pigment and rendered impervious. When this is the case the lymph probably passes by anastomosing vessels to other glands which in their turn become pigmented. In this way we may explain how pigmentation sometimes extends from gland to gland in the central direction. Thus in anthracosis of the lungs we may have not only the bronchial glands but also those of the mediastinum, and even those of the abdomen and at the porta of the liver pigmented.

What has been said in regard to pigment applies also to other solid materials introduced similarly into the lymphatics. In stone-masons the particles of stone which get into the parenchyma of the lung are in part carried to the glands, where they produce similar changes to those mentioned, but even more readily, as from their mechanically irritating characters they are more apt to produce chronic inflammation.

**Acute inflammation of lymphatic glands, Lymphadenitis.**—As already indicated, this condition is nearly always a result of inflammation in the peripheral parts from which the vessels come to the gland. The enlarged and inflamed gland is called a **Bubo**. The most serious of the acute inflammations are connected with infective processes, as in the case of dissecting wounds, erysipelas, phlegmonous inflammations, diphtheria, scarlet fever, splenic fever, and the plague (Bubo-plague).

The inflamed gland is enlarged, its vessels hyperæmic, and the tissue softened. The enlargement is due to a great increase of lymphoid cells which accumulate largely in the sinuses. The stellate cells forming the reticulum of the sinus also swell, and their nuclei divide. The sources of the added lymphoid cells are several. They arise by division of the existing cells of the gland, as evidenced by the observation of karyo-



mitosis in these cells; but they have their origin in part in the emigration of white corpuscles from the blood-vessels in the gland, and further, the primary inflammation of the periphery may send its contingent. The inflammation may subside and the gland return to its normal condition; the new-formed cells may possibly pass on into the efferent vessel and be disposed of in the circulation.

Not infrequently suppuration occurs; the tissue of the gland breaks down and an abscess is the result. We have thus a **Suppurating bubo**. The inflammation usually extends to the capsule and surrounding tissue which become involved in the suppuration. In this way the abscess advances, and the pus is finally discharged at a surface. It is a common occurrence for such abscesses to discharge on the surface of the body, but they may burst into bronchi, into serous cavities, etc. Some of these severe inflammations have a hæmorrhagic and even a necrotic or gangrenous character.

**Chronic lymphadenitis.**—This term is sometimes used so as to include conditions occurring in leukæmia, Hodgkin's disease, and scrofulous disease of the glands. These, however, belong to special categories, and the name is properly used for a simple chronic inflammation in connection with prolonged irritation of the glands. This is most directly produced by the introduction of foreign substances, where the solid particles are conveyed to the glands (see above). It occurs also in connection with prolonged peripheral irritation, as where an eczema of the leg or the scalp produces an enlargement of the corresponding glands. The inflammation here results in increase of the entire elements of the gland, or else in a preponderating increase of the connective-tissue stroma with atrophy of the proper gland tissue and induration of the gland.

**Tuberculosis of the lymphatic glands (*Scrofula*).**—Tuberculosis occurs in the lymphatic glands as elsewhere, by the action of the tubercular bacilli. In some cases there is an obvious propagation of these micro-organisms from an existing local tuberculosis at the periphery, but there are many cases where the primary seat appears to be in the glands.

**Primary tuberculosis** occurs for the most part in glands which have connections with mucous membranes having absorbent functions. There is usually at the outset catarrh of the mucous membrane, and it may be presumed that here as elsewhere the exuded products of inflammation are conveyed to the glands. There is no reason to believe that these catarrhs are, in the proper sense, tubercular, but it appears that, along with the inflammatory products, tubercular bacilli may be carried from the surface. In susceptible persons these will multiply in the glands and produce a tuberculosis. Such primary tuberculosis mostly occurs in glands whose radicles are connected with the mouth, pharynx,



fauces, or intestine, and they are mostly situated about the jaws or neck, and in the mesentery.

The tuberculosis is characterized by the formation of tubercles, which takes place chiefly in the follicular cords. This is accompanied by inflammatory swelling of the gland, just as in other forms of tuberculosis. There is usually a great newformation of cells which, in the more chronic cases, have mostly the character of epithelioid cells. The amount of swelling and its acuteness vary greatly. There may be a rapid increase of size with redness and other appearances of inflammation, or it may be a slow and indolent process.

**Caseous necrosis** soon presents itself; at first in a number of centres, but as it proceeds involving more and more of the gland, while the areas coalesce. The extension of the tuberculosis goes on while the older parts are caseous, till the caseous condition comes to involve the whole gland. The completely caseous gland is hard, and on section is firm to the touch, presents a homogeneous yellow aspect which has been aptly compared to that of the cut surface of a raw potato. Sometimes at the periphery there is a grey transparent zone representing the still-advancing tuberculosis, but the caseous condition may extend to the capsule.

The **Infective character** of the process is manifested by its extension to neighbouring glands. There is sometimes in the neck a complicated tuberculosis of many glands, some lying deeply among the muscles and fasciæ. It is noteworthy, however, that there is little or no tendency to extension through the capsule unless it be first ruptured or destroyed. This accords with the fact that tuberculosis does not usually penetrate connective-tissue membranes.

The caseous condition being thus brought about, the disease, so far as the individual glands are concerned, may undergo certain further developments. The caseous matter may simply remain as a piece of dead material, surrounded by connective tissue. It often becomes **infiltrated with lime salts**, and converted finally into a putty-like or stony mass. This is most liable to occur in glands which are protected, such as the mesenteric and other abdominal glands. It is quite common to meet with stony and pultaceous masses in the mesenteries of adults, these being the obsolete remains of a tuberculosis in childhood.

In the case of superficial glands, which are exposed to mechanical irritation, there is, on the other hand, frequently a **Softening** associated with inflammation. A process akin to suppuration ensues, in which the matter is partly composed of the debris of the caseous matter and partly of pus. The capsule is perforated and the matter passes towards the surface where it opens as a chronic abscess. In its passage the matter



produces a tuberculosis of the parts with which it comes in contact so that **Tubercular sinuses and ulcers** are frequent results. It is well known that recovery from this condition is only insured by careful removal of all infective material.

**Secondary tuberculosis** is very common in glands connected with organs which are the seat of local tuberculosis. This applies especially to the bronchial glands which are always secondarily involved in phthisis pulmonalis. The process here is often a very chronic one, and the glands sometimes instead of caseating become indurated as in chronic inflammation. Indeed, in fibroid phthisis the two processes of induration and tuberculosis often go together. Secondary tuberculosis also arises in the mesenteric and other abdominal glands from tubercular ulcers of the intestine.

The glands are also commonly affected in **General tuberculosis**. They are the seat of small miliary tubercles and are generally enlarged and hyperæmic.

**Syphilitic disease of lymphatic glands.**—This, as we have seen, is a constant result of the primary syphilitic infection, and constitutes the syphilitic or **Indurated bubo**. The glands slowly enlarge by newformation of round cells, and they remain enlarged for months or years. There is little caseous necrosis in the glands, but the connective tissue trabeculae are thickened. During the secondary stage other glands may swell, such as those of the neck.

**Gummata** sometimes form in the glands in inherited syphilis and in the late tertiary stage of ordinary syphilis. There is enlargement and, it may be, extension to neighbouring glands. The enlarged glands may or may not become caseous. (See account by Birch-Hirschfeld.)

**Tumours of lymphatic glands.**—We have already described several forms of newformation of lymphatic tissue, some of which attain to the dignity of actual tumours, and are designated lymphoma or lymphadenoma. There is the typhoid lymphoma, the leukaemic lymphoma, and the malignant lymphoma or Hodgkin's disease. All of these partake of the character of infective processes, and it is only in the case of the last mentioned that any claim can be made for a place among tumours proper. Under this view it is sometimes called lympho-sarcoma. (See under Malignant Lymphoma, p. 217.)

Of the tumours proper the primary forms are rare but the secondary frequent. The **Myxoma** is sometimes primary, and the **Chondroma** has been observed as a secondary tumour.

In a case by Virchow a chondroma occurred in the axillary glands in connection with a primary tumour of the scapula, and in one by Paget in the glands of the groin in a case of chondroma of the testis.



**Sarcoma** occurs **primarily** as a hard or soft tumour. The harder form is rare and is usually a fibro-sarcoma. The soft form may be a spindle-celled tumour but is more frequently round-celled. It may be an alveolar sarcoma. The commoner round-celled sarcoma resembles in structure the lymphoma and is liable to be mistaken for it, but it does not present the tendency to spread from gland to gland, and in its metastasis does not affect the lymphatic system specially. The tumours usually grow rapidly and form large fungating masses. Secondary growths frequently form in the lungs. These tumours originate chiefly from the retroperitoneal, mediastinal, and bronchial glands. In the alveolar form the newformation occurs chiefly in the adventitia of the vessels.

Sarcoma does not readily occur **secondarily** in the lymphatic glands, but according to Butlin it does so more frequently than is usually stated. Sarcomas of the foot, tonsils, testicles, and probably the kidney are liable to affect the lymphatic glands secondarily. The pigmented and softer forms of sarcoma are more likely to affect the glands.

**Cancer** does not occur as a primary growth in the lymphatic glands but is very frequent as a secondary affection. Cancerous tumours by their irritation may lead to a simple inflammatory enlargement, but, even in those which are very slightly enlarged, there are usually cancerous developments detectible by the microscope. (See further under Cancer, p. 266.)

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### III.—THE SPLEEN.

The exact nature of the function of this organ is somewhat obscure. In order to understand the various changes which it undergoes, it will be necessary to bear in mind certain facts as to its structure. The splenic artery as it enters the organ is accompanied by connective tissue which forms a continuous sheath around its branches. In this sheath



there develop at intervals little masses of lymphoid tissue, forming the Malpighian bodies, which are therefore lymphatic follicles closely related to the arterial branches. The arteries break up into capillaries which

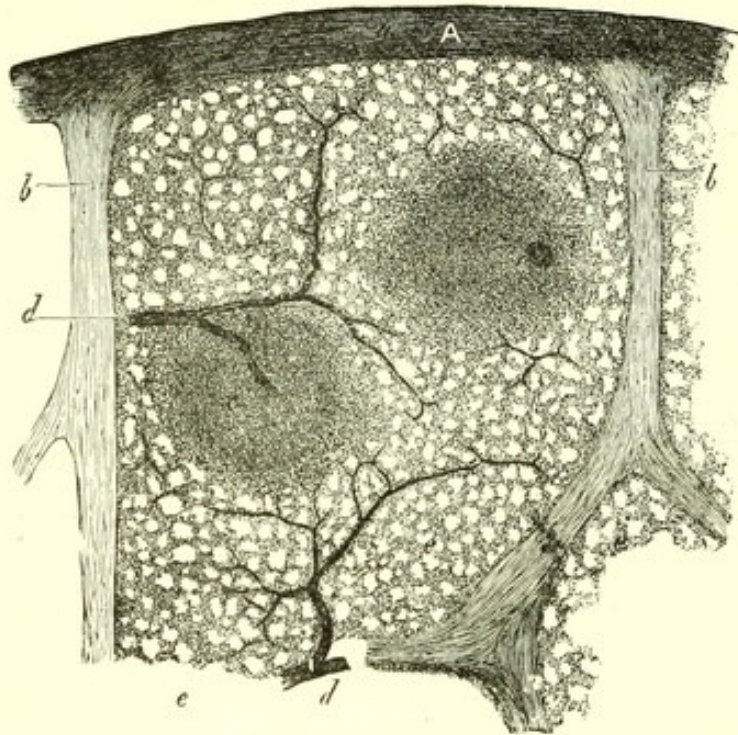


Fig. 179.—Structure of spleen. *A*, capsule; *b*, trabeculae; *c*, *c*, Malpighian bodies; in that to the right an artery, cut transversely, is seen; *d*, *d*, injected arteries, one running into a Malpighian body; *e*, honeycomb-like pulp. (KÖLLIKER.)

are mostly distributed in the Malpighian bodies, where they form a somewhat wide-meshed network. At the periphery of the Malpighian bodies the capillaries open into the splenic pulp, which is, as it were, interposed between the capillaries and the veins. This consists of a honeycomb-like structure, with frequent trabeculae and meshes (see *e*, *e*, Fig. 179). The meshes contain blood, but also large cells and cells containing blood-corpuscles and pigment.

The mesh-work of the pulp develops occasionally wider channels or sinuses which are virtually the radicles of the veins. The pulp has a brownish colour both from its pigment and from the fact that it is full of blood. The Malpighian bodies being lymphoid in structure, and sparsely vascularized, are whitish in colour. It is clear from this structure that the blood passing into the spleen by the arteries will linger long in the organ, and will especially stagnate in the meshes of the pulp. It seems probable that in the pulp the red corpuscles are to some extent destroyed. The existence in the normal spleen of cells containing red corpuscles and pigment seems to indicate this. The spleen also probably gives origin to white corpuscles, but it shares this function with the lymphatic glands and the medulla of bone, as well as probably with the widely diffused connective tissue throughout the body. Some have supposed that the spleen also forms red corpuscles, but this is exceedingly doubtful.

From these remarks it will be understood that the amount of blood in the spleen will be subject to great variations according as the pulp is more or less distended. The pulp forms by far the greatest part of the substance of the spleen, and according as its honeycomb structure is more or less full of blood will the size of the organ vary. Accordingly



there is no organ in the body which varies so much in size, even under physiological circumstances. During digestion there is an active hyperæmia which causes enlargement of the organ. Again, the capsule and trabeculæ of the spleen are abundantly supplied with smooth-muscle fibre-cells, and its size is influenced by the state of contraction of these. It is well known that by electric stimulation the size of the organ may be diminished, the muscular tissue contracting.

**Malformations and Malpositions.**—In addition to the spleen in its usual site, and of its ordinary size, there are very frequently small **Supernumerary spleens** lying in its neighbourhood. These are generally round in shape, and have the dark red or blue colour and soft structure of the ordinary organ. There are sometimes several of these present, and although usually seated in the neighbourhood of the spleen they may be away from it, having even been found embedded in the head of the pancreas. Cases have occurred in which there have been two spleens of nearly equal size. The spleen is also liable to variations in shape and position. These may be congenital, but **Alterations in position** may occur during life, chiefly when the organ is enlarged and increased in weight. Again, there may be an unusual length of the ligaments so as to allow of an undue mobility of the organ. This also may be congenital, or it may be due to a stretching of the ligaments from increased weight.

**Active or Congestive hyperæmia and Inflammation.**—We have already seen that an active hyperæmia of the spleen occurs physiologically during digestion. It is met with as a distinctly pathological condition, and frequently in a very exaggerated form, in a variety of acute general diseases. In this country it is often seen in the highest degree in typhus fever, but it is found also in other **Acute fevers**, in pyæmia, pneumonia, diphtheria, erysipelas, etc. In this group of diseases the blood is in a peculiarly irritating condition, and we know that the various parenchymatous organs, the muscles and the glands, are enlarged as a result of cloudy swelling, which is to be traced to the irritating condition of the blood. In these organs there is doubtless dilatation of the vessels, but in the case of the spleen the hyperæmia produces much more marked results on account of the structure of the organ. The hyperæmic spleen is greatly enlarged, and the capsule stretched. In some cases of typhus and intermittent fever the enlargement has been so great and sudden as to rupture the capsule. The name **Acute splenic tumour** is sometimes given to this hyperæmic enlargement. It will be inferred that in this condition the splenic tissue is exceedingly soft, sometimes almost diffluent, and the colour of the cut surface is a dark red.



But the condition does not continue as a pure hyperæmia. The cells forming the honeycomb structure of the pulp enlarge, the blood-vessels and the meshes of the pulp come to contain more round cells or white blood-corpuscles. These are all indications of inflammation, and a further indication is sometimes afforded by the deposition of fibrine on the capsule of the spleen. The **Malpighian bodies** sometimes undergo enlargement in this stage, but not in all the diseases named. This is most frequently seen in the later stages of **Typhoid fever** and **Small-pox**, and when it occurs the consistence of the organ is firmer than is usual in the acute splenic tumour. In this stage, in which more definite inflammatory processes are superadded to the congestion, the spleen is even larger than in the earlier stage, and it may reach two, three, or even four times the normal size. The organ is unduly soft, and on section may look half-diffuent, although it is firmer than in the cases without enlargement of the Malpighian bodies. The colour of the cut surface is considerably paler than in the earlier period, being more of a greyish or whitish red. On scraping the surface a thick juice is obtained which is not unlike pus mixed with blood. There is not infrequently hæmorrhage in the substance of the spleen so affected, and it may take the form of the wedge-shaped hæmorrhagic infarction.

As a general rule the acute splenic tumour diminishes as the primary disease passes off, and the spleen may be left soft and loose with wrinkled capsule and unduly prominent trabeculæ. Sometimes there results a chronic inflammation with thickening of the trabeculæ, but this hardly occurs unless there have been repeated attacks of hyperæmia as in malarial fevers, and occasionally in typhoid fever.

It is a very rare circumstance for suppuration to occur in the inflamed spleen, but this has been met with in intermittent fever. In such cases the pus appears in numerous little points which represent the Malpighian follicles, or there is a more diffuse suppuration of the spleen. If the abscess bursts a fatal peritonitis results.

Among the cases of suppurative inflammation of the spleen should be mentioned those in which **Ulcerative endocarditis** or **Pyæmia** is the primary disease. In that case minute emboli are carried to the spleen as to other structures, and being of a septic nature they each form the focus of an acute inflammation, which has at first a hæmorrhagic character, and afterwards passes on to suppuration. The spleen as a whole is enlarged by active hyperæmia.

**Chronic inflammation.**—In cases where the inflammatory enlargement is repeated frequently, or there is a more continuous irritation of the spleen, we have a chronic inflammation, causing the **Chronic splenic tumour**. This is very common in malarious districts, and is regularly



met with in persons who have been repeatedly subject to attacks of ague. But in such districts it occurs even in persons who have had no apparent ague, so that almost every post-mortem examination reveals a chronic **Ague cake**. Syphilis in the acute stages produces an acute splenic tumour, which may result in the chronic form, but by no means constantly. Again, the chronic form may occur in congenital syphilis and in rickets.

The enlargement in ague may depend on a **General hypertrophy** of the spleen, so that the consistence of the organ is nearly normal. There is, however, considerable increase of the lymphoid elements and the trabeculæ are thickened. There is frequently also increased pigmentation. In other cases there is a marked newformation of connective tissue in the trabeculæ, the pulp is firmer and the blood spaces reduced in size. This condition warrants the designation **Fibrous induration of the spleen**. The organ as a whole is not so much enlarged as in the other form, but it is much firmer and harder, so that the name ague cake is peculiarly applicable here. In extreme cases there is atrophy of the proper constituents of the spleen; the pulp is much encroached on, and the Malpighian follicles are greatly destroyed. There is usually also much deposition of pigment in the splenic tissue, especially in the pulp, but also to a slight extent in the Malpighian bodies, and in the thickened connective tissue around the vessels (see *Melanæmia*).

In both forms of chronic tumour of the spleen the capsule is generally thickened, a condition being thus brought about which is sometimes called **Perisplenitis**, and is analogous to a similar thickening of the capsule of the liver, *perihepatitis* (see also under *Chronic Peritonitis*). The thickening is usually irregular, and we have tendon-like patches or loose shreddy connective tissue on the surface. Sometimes the capsule acquires adhesions to neighbouring structures, and the spleen may be so buried in adhesions as to make it difficult to dissect it out. It is to be remembered, however, that such adhesions are very frequent apart from any proper disease of the spleen at all, being due to inflammations in its neighbourhood. There are, indeed, occasional patches of dense fibrous or cartilaginous consistency occupying the capsule of the spleen, whose origin it is difficult to ascertain, but which are to be regarded as the results of local peritonitis. In most cases of perisplenitis originating outside the spleen the organ is small and atrophied.

**Passive hyperæmia.**—This condition occurs in the spleen when there is any considerable obstruction of the portal circulation. Hence we meet with it in cirrhosis of the liver, and in cases of heart disease which have gone on to general venous engorgement, the hyperæmia being, as it were, transmitted through the liver. It may, indeed, occur



in consequence of any local or general obstruction whose effects extend to the splenic vein. The organ is enlarged, but not to such an extent as in the congestive form. As in other cases of passive hyperæmia, the connective tissue increases in firmness, and so the whole organ is denser than usual, while the tissue presents a deep red colour. The thickening of the connective tissue affects the trabeculæ and the sheaths of the blood-vessels.

**The embolic infarction.**—This is probably of more frequent occurrence in the spleen than in any other organ, except perhaps the lungs. The explanation of this is probably the large size of the splenic artery. We have already seen that this artery and all its branches are end-arteries, and that obstruction of one of them can hardly fail to cause the infarction. The infarction has sometimes the hæmorrhagic character, but this is frequently absent except at the marginal parts, and the greater part is white or yellow. Even when originally red it soon loses much of its colour. The infarction has more or less the form of a wedge with its base at the capsule. On handling the organ it is to be detected by its density, forming a hard mass in the midst of the soft tissue. As a rule, there are several infarctions in the same spleen, and as the embolus in a larger trunk may break up and be distributed irregularly to various branches, we may have very complicated forms assumed by the infarction. For example, it is not uncommon to find the superficial parts of the spleen involved over a large area, while the infarction does not extend much beneath the surface, as if the emboli had been swept into a number of terminal branches near the surface. **The spleen**, as a whole, **is enlarged**, the obstruction of the arteries leading to a collateral hyperæmia by dilatation of those remaining open. This enlargement may reach considerable dimensions. On the capsular surface of the infarction there is often a deposition of fibrine, and it is interesting to know that during life the existence of this has been sometimes diagnosed by the discovery of friction sound.

The affected area of splenic tissue is in a state of coagulation-necrosis, and presents the usual characters of that condition, chiefly the absence of the nucleus when the tissue is stained. The blood-colouring matter is dissolved out, being partly absorbed and partly deposited in solid granules throughout the infarction or at its marginal parts. Around the infarction a chronic inflammation occurs with the usual newformation of granulation tissue which eats into the infarction. As the infarction becomes absorbed, the connective tissue which develops from the granulation tissue draws together, the final result being a cicatrix in which some cheesy matter may be found representing the original infarction. During these processes the spleen gradually returns to its normal size



and condition, with, it may be, some thickening of the capsule. We frequently meet with one or more cicatrices in the spleen from old embolism. These are always visible at the surface and penetrate inwards, but often they have little depth.

The condition of the spleen in **Leukæmia** has been already described, and it is to be remembered that the spleen may be enormously increased in size, up to a weight of 40 pounds. The condition in malignant lymphoma or **Hodgkin's disease** has also been described.

**Rupture of the spleen.**—This occurs occasionally, as we have seen, in acute enlargements of the organ. But traumatic rupture is much more frequent. This is effected by blows or falls on the abdomen, and also by injuries to the chest by which the lower ribs are forced against the organ. There may be considerable rupture without any external marks of injury. It is important to remember that the enlarged and hard spleen is more liable to rupture than the normal one, both on account of its more brittle character and its size. This is important in a medico-legal aspect, especially in malarious districts. The rupture may lead to fatal hæmorrhage, but bleeding is often associated with other injuries, and only plays a part in the result. On the other hand, the hæmorrhage may be slight, and the wound may heal and leave a cicatrix.

**Degenerative changes.**—Of these by far the most important is **Amyloid disease**. It has already been mentioned that the spleen is more frequently the seat of this disease than any other organ, and that it appears in a majority of cases to be the organ primarily affected. There are two forms of amyloid disease, called respectively the **Sago form** and the **Diffuse form** or waxy spleen. In the former the Malpighian bodies are mainly engaged; in the latter the pulp. We are entirely ignorant of the conditions which induce these differences in the situation of the degeneration, but it may be said that the sago spleen is peculiarly that which occurs in phthisis pulmonalis, while the waxy form occurs in syphilis.

**The Sago spleen** is a moderately enlarged organ. On section we observe on the cut surface, instead of the normal small Malpighian bodies, transparent glancing areas which have been very aptly compared to grains of boiled sago. These are dotted over the surface in great profusion. On applying a solution of iodine the affected areas stand out as brown spots, which become of a deeper colour on adding dilute sulphuric acid. Microscopic sections (see Fig. 180) show transparent hyaline areas of round shape and larger or smaller size, often so large that they are continuous with one another at their peripheries. These areas represent Malpighian bodies, whose normal structure, when the disease is advanced,



is entirely replaced by a nearly homogeneous transparent material. In the early stages, however, the addition of methylviolet or iodine brings out a beautiful network in the Malpighian body, and it is obvious that the reticulum is first attacked. Even in the early stage it can be seen that the lymphoid corpuscles are also to some extent involved, and as the disease progresses they are more and more affected, although to some extent they may be destroyed by the enlargement of the reticulum.

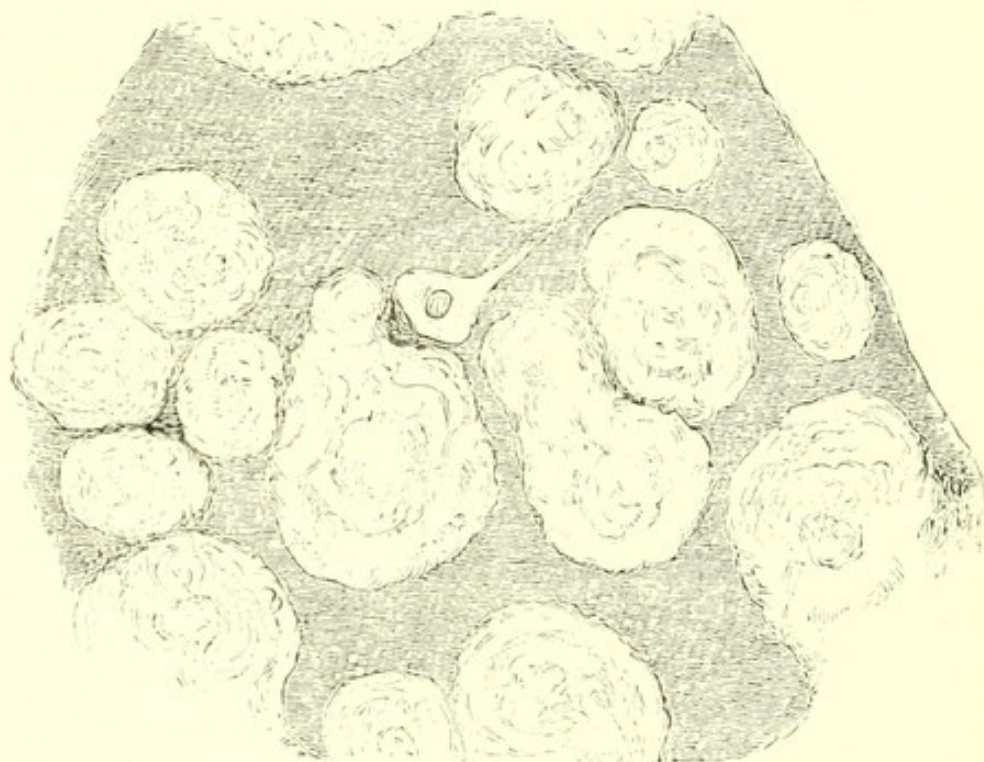


Fig. 180.—Section of a sago spleen. The enlarged and translucent Malpighian bodies are seen. In the middle an artery with amyloid walls.  $\times 20$ .

The arteries whose lymphoid bodies are thus affected are often themselves degenerated, but they may remain unaffected. In advanced stages of the disease the enormous enlargement of the Malpighian bodies causes atrophy of the pulp, and we may have the spleen presenting little beyond large round, sago-like bodies. The pulp may, however, itself take part in the amyloid disease at the periphery of the Malpighian bodies.

**The diffuse amyloid spleen** presents much greater enlargement than the sago form. It is in the highest degree hard and heavy, and the edges rounded. On section the tissue seems homogeneous and inelastic, and of a dark transparent appearance, which has been compared to that of wax or bacon, hence the names Waxy and Lardaceous spleen, which are most appropriately applied to this form. The degeneration affects the greater part of the spleen, but there may be islands of normal pulp visible. The Malpighian bodies are hardly visible, and the cut surface has a smooth uniform appearance. The application of



iodine produces a general deep brown coloration of the tissue. Under the microscope (as shown in Fig. 181) it is seen that the trabeculae of the pulp are involved. On account of the greater density of the amyloid material the honeycomb structure of the pulp is often brought out in a much more striking manner than is possible in a normal spleen. This is more marked if the sections be stained with methyl-violet, the trabeculae assuming a bright colour and becoming very manifest. In the later stages the appearance is more uniform, but indications of the trabecular arrangement may still be visible. By the enlargement of the pulp the Malpighian bodies are greatly atrophied, but they may show traces of amyloid disease. In this form the walls of the arteries and veins are usually amyloid.



Fig. 181.—Diffuse amyloid spleen. The swollen and translucent tissue of the pulp (a) is shown.  $\times 400$ . (After KYBER.)

**Infective and other tumours of the spleen.**—**Tuberculosis** does not occur in the spleen as a primary disease, but in acute general tuberculosis we frequently find tubercles in great abundance. The tubercles are usually seated in the neighbourhood of arteries, and are often difficult to distinguish from the lymphoid Malpighian bodies which have a similar seat. The existence of the giant-cell structure and the tendency to caseous degeneration will assist in distinguishing them. Occasionally we meet with large caseous masses in the spleen, which may reach the size of walnuts. These occur mostly in children who are the subjects of scrofulous disease of the lymphatic glands of the abdomen. The general distribution of the nodules suggests that the infective material has been carried by the blood, and in some cases there may have been a rupture of a softened gland into the splenic artery. **Syphilitic gummata** are very rare in the spleen, and have been met with chiefly in congenital cases. In Hodgkin's disease the spleen is generally the seat of newformations, and there is usually great enlargement of the organ (see p. 217).

**Tumours proper** are equally rare, but cases of Fibroma, Sarcoma, and one case of a pulsating Cavernous angioma have been observed. Sarcomas occur secondarily in the spleen with greater frequency than any primary tumour, and this is especially true of melanotic sarcomas, which



may possibly originate in the spleen. Secondary cancers are very rare and only occur in cases where, from the general diffusion of such tumours, we may infer that the infective material has found its way into the blood.

**Parasites** are very unusual in the spleen. The **Echinococcus** is the most common, forming hydatid cysts, sometimes of large size.

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#### IV.—CERTAIN AFFECTIONS OF THE BONE-MARROW.

The bone-marrow is to be regarded from two aspects, first as a part of the osseous system, and second as an organ presenting some special relations to the blood, and partaking in many of the diseases in which the spleen is involved. In its structure the normal bone-marrow is divisible into two forms, namely, the red marrow, which is composed of round-celled or lymphatic tissue, and the yellow, which is simply adipose tissue.

While the red marrow is present in nearly all the bones of the fœtus, it is greatly limited in its distribution after birth. In the adult it is present in the bones of the head, and most of those of the trunk, being in many cases, however, associated with a considerable amount of adipose tissue. The bones of the extremities contain almost exclusively yellow marrow, but there may be some red marrow in the heads of the femur and humerus.

The pathological changes in the bone-marrow are essentially related to those of the blood. In its physiological relations to the blood it may be primarily involved along with the other blood-forming organs, or it may be secondarily affected when the constitution of the blood is altered. In all of these it closely follows the spleen in its lesions.

In **Leukæmia**, as mentioned in a previous page, there is an increase in the lymphoid tissue of the bone-marrow, a real hypertrophy or hyperplasia of this tissue. In a certain sense this is a restoration of the red marrow which existed in the fœtal state, but as the cellular growth increases the marrow often ceases to be red, and becomes yellow like pus. (See under **Leukæmia**.)

In a case lately observed by the author, the bone-marrow had rather the characters usual in Pernicious anæmia, presenting chiefly large granular cells, many of



them fatty. It was very peculiar that the blood, which to the naked eye was greatly altered, contained large numbers of similar cells, many of them also fatty. These cells were distinguishable from leucocytes by their larger size, and by the possession of large oval or round nuclei. The spleen was also greatly enlarged.

In **Pernicious anæmia**, and to a certain extent in other anæmias, there is what at first sight looks like a conversion of the yellow marrow into red. It is true that the yellow marrow is replaced by a red, almost diffuent structure, but this is not a proper lymphatic tissue. We have not the small lymphoid cells, but rather large granular cells, some containing red corpuscles, and various forms of red corpuscles, some of them nucleated. This change is probably secondary to the anæmia, and is perhaps to be regarded as a retrograde change rather than as a hypertrophy.

In **Acute fevers** the bone-marrow is affected like the spleen. It is hyperæmic, and there are large granular cells visible, especially in the coats of the vessels. There are sometimes hæmorrhages, which may take the form of infarctions. According to Golgi, the bone-marrow in **Hæmorrhagic small-pox** shows everywhere diffuse hæmorrhages.

The more definite septic inflammations, included under the term **Osteomyelitis**, fall to be considered along with the affections of the bones.

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## SECTION II.

## DISEASES OF THE BONES AND JOINTS.

- A. OF THE BONES.—*Introduction as to development and structure.* I. MALFORMATIONS. II. RICKETS, a disease of growing bones, caused by general ill-health. Character of lesions at ossifying cartilage and under periosteum; Deformities due to rickets in long bones, chest, vertebræ, pelvis, head. Recovery. III. RETROGRADE CHANGES (1) Atrophy; (2) Osteomalacia or Mollities ossium. IV. INFLAMMATIONS—*Introductory.* (1) Acute and suppurative inflammations, traumatic and infective, the latter important; (2) Chronic inflammations, implying rarefaction and newformation. V. HYPERTROPHY. VI. NECROSIS—*Causation.* The resulting lesions mainly inflammatory; separation of sequestrum; absorption of dead bone. Phosphorus necrosis. VII. REGENERATION. (1) Healing of fractures; production and structure of callus. (2) Transplantation of bone. VIII. SPECIFIC NEWFORMATIONS. (1) Tuberculosis; causation; lesions; including caries, cold abscess, etc.; healing of lesions. (2) Syphilis; gummata and lesions of congenital syphilis. (3) Actinomycosis. IX. SPINAL CURVATURES—*Introduction.* Forms of curvature, antero-posterior or lateral. X. TUMOURS, chiefly exostoses and sarcomas.
- B. THE JOINTS. I. DISLOCATIONS AND MISPLACEMENTS, congenital forms; talipes; traumatic; spontaneous. II. ANCHYLOSIS. III. INFLAMMATIONS. IV. SYPHILIS AND TUBERCULOSIS. V. LOOSE BODIES.

## A.—OF THE BONES.

**I**NTRODUCTION.—Bone differs materially in structure and function from most of the other tissues of the body, and these peculiarities exercise an important influence on the pathological changes to which it is liable.

During the period of growth bone is in a condition of great activity, the growth of bone being effected by processes which are, in some respects, special and peculiar. The proper bony tissue (*tela ossea*) is composed of a rigid calcified matrix, enclosing the living cells or bone corpuscles in their lacunæ. This tissue once formed is, like the mason work of a house, fixed in form and insusceptible of expansion or plastic alteration in shape. It can only be altered by being taken down and rebuilt. But in the growing bones there is necessarily a very active process of reconstruction going on. The bone as a whole is expand



ing, and this is effected, not merely by apposition of new bony tissue to the old, but it involves a process of destruction and reconstruction. It has been shown by Meyer and Wolffe that bones have a special architectural construction, their trabeculae being arranged so as to meet the various lines of pressure to which they are opposed. In a growing bone the architecture is continually requiring modification as the bone expands, and this implies a constant reconstruction. The extent to which reconstruction is necessary is emphasized when one remembers that the shaft of a young child's femur is no greater in diameter than the medullary cavity of that of an adult. In fact, in the course of growth the whole bone is, in many cases, taken down and reconstructed several times over.

The growth of bone is effected by the so-called process of **Ossification**. In a longitudinal section of a growing long bone, such as that illustrated in Fig. 182, the process of ossification from cartilage may be seen.

At the upper part of the figure (*a*) there is the normal appearance of hyaline cartilage. As the ossifying margin is approached the cells multiply, and at first they form irregular groups, as at *b*. Still nearer the margin the multiplying cells become arranged in rows parallel to the long axis of the bone, the elongated groups being separated by narrower elongated pieces of hyaline matrix. The beginning and end of this zone of multiplying cartilage cells are quite abrupt, and form nearly straight transverse lines. This zone appears to the naked eye as a narrow bluish band. Next occurs an infiltration of the hyaline matrix between the groups with lime salts, the matrix becoming opaque and granular (*c* in figure). This forms a yellow zone as seen with the naked eye. This zone is also suddenly interrupted, and at a definite level the groups of cartilage cells give place

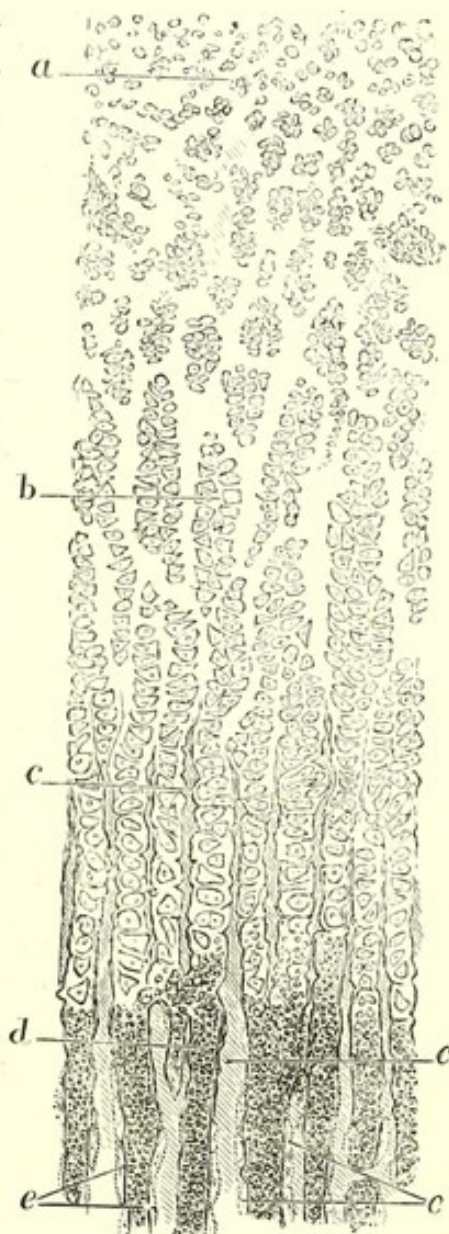


Fig. 182.—Longitudinal section of bone ossifying from cartilage, *a*, commencement of proliferation of cartilage cells; *b*, cells arranged in longitudinal groups; *c*, matrix of cartilage impregnated with lime, this calcified matrix is continued beyond the ossifying border (*c* on right side); *d*, medullary spaces containing round cells replacing cartilage cells; *e*, formation of bone at borders of calcified matrix, the osteoblasts each forming a piece of bone and remaining as the bone orpuscle.  $\times 90$ . (After THIERFELDER.)



to cavities filled with foetal marrow, consisting of round cells and blood-vessels (*d*). It is from these cells that the formation of true bone occurs. The cells contained in the spaces have the power of producing bone, and are hence called **Osteoblasts**. These fix themselves to the walls of the cavities, which, as we have seen, are composed of calcified cartilaginous matrix, and each cell forms around itself a certain portion of bone-tissue, apparently by a process of secretion. The *tela ossea* thus formed encloses the cell which has formed it, and which remains as a bone corpuscle. In such a section as that under review one constantly sees at the borders of the cavities layers of osteoblasts with oval areas of new-formed *tela ossea* around them, *e* in figure. Thus the calcified cartilaginous matrix is encroached upon, and the bone is formed.

Besides this ossification from cartilage, we have what is commonly but not quite correctly called Ossification from membrane. In all such cases we have cells similar to those in the cavities mentioned above, which have the characters of osteoblasts, and form bone in a similar fashion. These osteoblasts form a layer under the periosteum, they exist also in the spongy parts of bones, and at the ossifying margins of flat bones.

The destruction of bone is also effected by active cells which are distinguished by the name of **Osteoclasts**. By several authors the giant-cells or myeloplques which exist in foetal bone-marrow are regarded as having this function. Where absorption is going on they have been found attached to the bone, and lying in gaps apparently formed by them and called Howship's lacunæ. It should be added that this view is not universally accepted. Some believe that the myeloplques are not necessary for absorption, and some hold by the opinion of Virchow, that the Howship's lacunæ arise by the action of the bone-corpuscles, the rounded gap representing a district belonging to a bone-corpuscle which has undergone solution.

When the period of growth is finished, the osteoblasts and the osteoclasts have completed their work, but the bone remains penetrated in every part by soft tissue. We may accept the view of Ranvier, according to which the bone in the adult state is enveloped in, and penetrated by the remaining potential osteoblasts. This tissue constitutes the bone-marrow and the internal layer of the periosteum (called *Periosteal marrow* by Ranvier), while between these two there is a connecting network traversing the Haversian canals.

From these observations it appears that in childhood the bones are in an extraordinary state of activity, a process of enlargement and reconstruction continually going on. At the same time, the function of the bones as substantial props is in full activity. It is not remarkable, there-



fore, that in childhood the bones are peculiarly liable to disease, which may take origin from without or be due to inherent weakness.

In the diseases both of children and adults again, there may be a partial return to the conditions of the period of growth, and osteoblasts and osteoclasts resume their operations, the processes being now pathological.

Another characteristic of pathological processes in bone is the occurrence, formerly alluded to, of **Metaplasia**. Bone, cartilage, and connective tissue are fundamentally the same tissue, and apparently the same cells may be, according to circumstances, osteoblasts, fibroblasts, or chondroblasts. This will be illustrated under Fractures.

#### I.—MALFORMATIONS OF BONE.

The more important malformations of bone are connected with general malformations, and have been considered in an earlier part. We have defects of the skull in anencephalus, and of the vertebræ in spina bifida.

Besides these, we have congenital reduplication of bones, such as of the vertebræ, the ribs, the fingers, and toes. **Supernumerary vertebræ** are met with in any of the regions of the column. **Supernumerary ribs** may be cervical or lumbar. In **Supernumerary fingers and toes** the bones may be absent, or may be represented only by cartilage.

**Malformations of the skull** have been ascribed to premature coalescence of the bones at the base, or of the sutures of the cranium, a condition designated **Synostosis**. The basilar parts of the occipital and sphenoid bones are formed from cartilage. The basilar part of the sphenoid is originally in two parts, presphenoid and postsphenoid, which unite before birth. The sphenoid and occipital bones are separated by cartilage up till the twentieth year, and as this cartilage has the characters of ossifying cartilage, the growth in length of the base of the skull depends on it. A premature synostosis of these two bones will check growth and lead to a **Shortening of the base** of the skull. To this is ascribed the existence of abnormal retraction of the nose.

The bones of the calvarium grow at their margins, or in other words, at the sutures. A premature coalescence of the bones or **Closure of the sutures** will stop this growth. According to the sutures affected the result varies. If the whole sutures close there will be a general smallness of the head, a **Microcephalus**. If there be a premature synostosis of the coronal and lambdoidal sutures, the growth in length is checked, and, as a compensatory growth in breadth occurs, the head is abnormally wide and flat on the vertex, conditions designated by the terms **Brachy-**



**cephalus** and **Platycephalus**. When the sagittal suture is affected, growth in breadth is interfered with and an abnormally long cranium is produced, a **Scaphocephalus**.

There seems no doubt that microcephalus arising in this way may lead to arrest of the growth of the brain, some cases of idiocy being thus explained. On the other hand, smallness of the brain will be associated with smallness of the skull, but this will not be associated with premature synostosis.

## II.—RICKETS OR RACHITIS.

**Causation.**—This disease occurs in children during the earlier periods of growth, and so far as the bones are concerned it is a disease of development, the normal process of ossification being interfered with. Cases have been observed of congenital rickets, but these are rare, and in the majority of cases it begins in the first or second year, its commencement being very rarely delayed beyond the fourth or fifth year. In a large proportion of cases its onset seems to be in the latter half of the first year or the first half of the second.

An endeavour has been made to make out that rickets is merely a manifestation of hereditary syphilis, but this view has not received acceptance in this country. It has been pointed out again and again that rickets is very common in cities where syphilis is uncommon. In the city of Glasgow, for instance, rickets is very common among the poorer classes, but syphilis is very uncommon. It is so also in Aberdeen, Edinburgh, Belfast, and elsewhere. We shall see afterwards that syphilis produces changes in the bones in some respects analogous to rickets, and atrophy of the cranial bones, sometimes designated **Craniotabes**, although formerly regarded as rachitic, is probably in most cases syphilitic; the two diseases, however, are essentially separate.

Rickets occurs amongst children who are injudiciously fed and badly housed, and the disease in the bones is to be regarded as part of a general loss of health. The body is weakened, and, besides the bones, there are other parts that suffer. According to Macewen, it supervenes in a considerable number of cases on an attack of one of the acute fevers, as measles or scarlet fever, such an attack leaving the child in an ill-nourished condition. There are usually symptoms of indigestion and intestinal catarrh, sometimes with fever, and in a certain proportion of cases there is enlargement of the spleen and sometimes also of the liver and lymphatic glands. The bones, being the most actively growing parts at this period of life, are, however, specially affected.

Rickets, or a condition analogous to it, has been produced artificially in growing animals by feeding them with small doses of phosphorus, or by giving them lactic acid, by the mouth or subcutaneously. In both cases the diet must be deficient in lime salts. In the latter case it has been supposed that the lactic acid, being a ready



solvent of lime, prevents its deposition in the bones, and it has been inferred that in the human subject gastric derangement causes the formation of an excess of lactic acid, which is the cause of the disease in the bones. There is, however, the objection to this view, that rickets is not always accompanied by any considerable gastric disorder, and such a view does not explain its occasional occurrence in the fœtus. Besides, the disease is clearly an error in development, and were it due merely to absence of the lime salts, we should expect the bony matrix to be formed without the lime salts. The very opposite frequently occurs; we have often an impregnation of the cartilage with lime salts, a calcification of the cartilage (see Fig. 183, *b, b*), while true ossification lingers. Rickets can, in most cases, be cured by supplying sufficient food and proper lodging, and this points to its origin in the opposite of these.

**Character of lesions.**—The lesions in rickets may be summed up in the statement that the preparatory stages of ossification are exaggerated, while the completion of the process is delayed.

**At the ossifying cartilaginous border** the conditions may be studied by examining Fig. 183, and comparing it with Fig. 182, which represents normal ossification, both figures possessing a similar degree of magnification. The most pronounced difference from the normal is the enormous exaggeration of the zone of multiplication of the cartilage cells (*a, a*). The groups of cells are in much larger numbers and occupy a much greater space; at the same time they have not always the normal arrangement in rows parallel to the long axis of the bone. The cells of the groups also vary considerably in size. In the normal bone the cells in the deepest part of this zone are large, but in rickets, while some are large, as at *b*, there are others even at the same level, or deeper, of small size.

Next, in regard to the zone in which the cartilaginous matrix becomes calcified, this also is represented here in the most irregular fashion. Calcification of the matrix occurs, and calcification of the cartilage cells (as at *b* on left), but it is very irregular. There will be calcification of the cartilage high up, as at *b*, while there are numerous groups lower than that uncalcified. In many cases also, whether calcified or not, the cartilage cells lose to a considerable extent their grouping into rows (as below *b* on left).

Then as to the formation of medullary cavities and development of bone, the figure shows how irregularly this occurs. Blood-vessels appear with considerable frequency in the midst of the cartilage, and towards the deeper parts there are medullary spaces with osteoblasts in them. These may even be seen forming new bone, as at *d*. But the medullary cavities do not occur with any greater regularity than the other stages in the process, and it is not always apparent that they have to do with the formation of the bone. It will be seen in the figure that at a certain somewhat indefinite level, which is lower at the





Fig. 183.—Longitudinal section of ossifying margin of a long bone in rickets. *a*, proliferating cartilage cells, the area of these very greatly extended and the arrangement quite irregular; *b*, *b*, calcification of the cartilaginous matrix at different levels, but not followed by formation of medullary cavities. The formation of medullary cavities (*c*, *e*) and of bone is occurring quite irregularly, the level being higher at the right (*c*) than at the left (*c*). At *d* the osteoblasts are forming bone. In various places, especially at *d*, pieces of cartilage are seen in the midst of bone, and an apparent transition of the one into the other is seen.  $\times 90$ . (THIERFELDER.)



left side than at the right, bony trabeculae appear (*c, c*). But it will also be seen that the bone is directly continuous with cartilage, the difference in the two being indicated chiefly by the difference in the cells. It appears also that in the cartilage, which is thus immediately continuous with the bone, the individual cartilage cells are separated by a matrix just as the bone corpuscles are (see figure). In fact there is here a conversion of cartilage into bone, and there are even appearances suggestive of the gradual transformation of cartilage cells into bone corpuscles. It will be seen in various places, but especially at *d*, that pieces of cartilage survive in the midst of the new-formed bone. The cartilage may be calcified or not.

To the naked eye the changes described above are indicated in a longitudinal section of a rickety long bone. The blue zone, which forms a straight narrow transverse line in the normal bone, is here greatly increased and its boundaries are irregular. The yellow zone of calcification of the cartilage is still more irregular, and yellow pieces crop up in the midst of the blue zone. Blood-vessels appear also at different levels. In the area of the blue zone the cartilage is greatly swollen as the enormously multiplying cells take more space than normal.

**Under the periosteum** the normal thin layer of osteoblasts is increased till it forms a layer of considerable thickness. Bone is formed to a very limited extent and very imperfectly, so that instead of a proper dense bone, such as should be formed on the surface of the shaft, there is a loose irregular spongy bone. But even the trabeculae of this spongy bone are not properly formed. Immediately under the thick subperiosteal layer the new-formed bone does not show in its matrix the homogeneous character of the matrix of bone; it is granular, and the lime is obviously deposited without combining with the matrix in the normal manner. It is a calcification rather than an ossification, and the trabeculae are rather osteoid than osseous. On passing inwards the tissue becomes more strictly osseous however, although hardly acquiring the regular arrangement of the dense bone of the shaft.

Corresponding with the structure described are the naked-eye characters. The subperiosteal layer is seen in the form of a red vascular layer of some thickness. This is so obvious in some cases that, at one time, it was described as if a blood-clot were formed under the periosteum. Beneath this the spongy character of the bone can be seen, while the shaft is considerably thicker than normal and obviously more easily bent.

**Deformities of the bones due to rickets.**—In considering the deformities so often produced by rickets, it is to be remembered that the two chief changes in the bones are, in the first place, enlargement



and softening of the epiphyseal extremities of the bones from the affection of the cartilage, and, in the second place, thickening with loss of resistance of the shaft from the periosteal lesion. These conditions do not always go strictly parallel, one or other being frequently the more prominent in a particular bone. The various bones of the body also present very commonly different degrees of rickets, although the disease, being a general one, usually affects many bones. Even in bones affected to a similar extent, however, the resulting deformities present very great variations; the deformities consist largely of curvatures of the bones from the application of external forces, and the bones are differently placed in relation to such forces.

**Deformities of the long bones.**—The most obvious change at the outset is **swelling of the cartilaginous ends** of the long bones, giving a clubbed or knobbed appearance to the limbs. This is common to all the long bones, and is the condition generally taken in practice as evidence of the existence of the disease. The bones are also arrested in their growth, so that they are **stunted** while at the same time they are unduly thick.

The remaining deformities occur in the majority of cases in the lower limbs, and are the result of the **weight of the body** acting on the bones in their weakened condition. Similar deformities are met with in the bones of the arms in cases where external force is frequently applied to them, as where a child in creeping leans on the arms, or where a nurse frequently lifts a child by one arm (Macewen). The deformities consist mainly of curvatures of the shafts of the bones, along with **shifting of the epiphyses** in some cases.

**Shifting of the epiphyses** is due to the condition of the ossifying cartilage. The extended blue zone at the cartilaginous border, being soft, allows the epiphysis to change its position according to the direction of pressure, and so at the ankle the epiphysis is sometimes displaced inwards, and, as it were, overhangs the joint on its internal aspect. The same may occur at the knee-joint, and the internal condyle may exceed in length the external even after the disease has been cured.

The **curvatures of the bones** are due mainly to the weight of the body acting on the softened shafts, and as the pressure acts mainly on the lower limbs the two principal forms are *genu valgum* and *genu varum*.

**Genu valgum**, or knock-knee, is a condition in which the thigh and leg form an angle at the knee with its apex inwards. This deformity depends usually on several alterations. There is generally a curve of the lower third of the femur, with its convexity inwards, the effect of this being that the internal condyle is lower and the external higher than normal. Along with this there is usually the lengthening of the internal condyle already referred to. As a rule the tibia is not bent, the two conditions named taking the chief part in producing the deformity, but sometimes the shaft of the tibia is at an angle with the epiphysis, as if the latter were to some extent overhanging the former. In addition the femur or tibia sometimes shows an anterior curvature, which of course does not increase the valgum condition.

**Genu varum**, or bow-legs, is the converse condition to *genu valgum*. The shafts of the femur and tibia are curved, with the convexity outwards, but these bones take part in the deformity in very different degrees, the tibial curve being more frequent and usually more pronounced than that of the femur.



The long bones being soft and flexible are not so liable as normal bones to complete fractures, but they are specially liable to **Partial fractures or infractions**. If the bone is suddenly bent it does not break across, but is partially torn as when an attempt is made to break a green stick. In this case the concave surface of the bent bone gives way and the convex surface does not. The marrow is torn by the broken concave portion, the broken edges of which may be projected through the marrow to the opposite internal wall. This kind of fracture has been aptly compared by Virchow to the breaking of a quill. These infractions occur most frequently at the lower part of the tibia, also in the pelvis and ribs, and less frequently in the bones of the arm.

**Deformities of the chest.**—The junction of the cartilaginous and bony ribs is analogous to the ossifying cartilage of a long bone, and undergoes a similar thickening in rickets. These parts of the ribs are therefore knobbed, and there is thus a row of knobs on each side of the chest—the so-called **Rachitic rosary**. The chest is also liable to deformity from the flexibility of the ribs. During inspiration the lateral aspects of the ribs are drawn inwards from being unable to withstand the atmospheric pressure, so that instead of the natural arch with the convexity outwards, these lateral portions are flattened or even rendered concave. As the ribs take thus a straighter course the sternum is pushed forward, and the antero-posterior diameter of the chest increased.

**Deformities of the vertebræ.**—The vertebræ very often escape in the milder forms of rickets, but in many cases they also are composed of abnormally spongy bone, and allow of the occurrence of curvature. Such curvatures are mostly exaggerations of the normal antero-posterior curves, but lateral curvature also occurs. Any considerable curvature will cause narrowing of the chest, and if this be associated with the deformity mentioned above the interference with the circulation and respiration may be serious.

**Deformities of the pelvis.**—These are of great importance in the female in relation to the possible occurrence of pregnancy in after life. The chief deformity is produced by the weight of the body acting through the vertebral column on the sacrum. This bone is pushed forward and the pelvis undergoes a corresponding displacement of its parts, the antero-posterior diameter being diminished. The growth of the bones here also is stunted, and the pelvis therefore remains unduly small.

**The bones of the head** frequently undergo very marked deformities. The bones of the face, like the bones generally, are stunted in their growth so that the face is small. It is also stated that the jaws by the action of the muscles undergo changes in shape by which the lower jaw is shortened and the upper jaw lengthened, so that the teeth of the latter overlap very much those of the former. While these are the conditions in the face the bones of the cranium present striking peculiarities. The flat bones ossify from membrane, and in rickets we have the ossification lingering behind so that at their borders these bones present somewhat broad areas in which there is soft tissue like that under the periosteum. The effect of this is to cause an apparent widening of the sutures and extension of the fontanelles. The closure of the fontanelles is also delayed. The cranium is enlarged in rickets, at least its circumference is greater, and at the same time it is commonly flattened on the summit. The enlargement of the cranium with the stunted condition of the face, causes the well-known overhanging of the brow so often seen.

Another occasional consequence of rickets is the condition designated **Craniotabes**. As we shall see afterwards this is frequently a consequence of syphilitic disease of



the bones, but it occurs also in rickets. The bone in rickets is soft, possessing less power of resistance than normal bone. If the child be lying constantly on one spot, or if the contents of the skull be increased, as in chronic hydrocephalus, then the pressure on the bone may cause it to waste. This occurs most frequently when both these conditions are present, and the bone is as it were between two pressures. In this way may occur thinning and actual perforation of the skull, so that in the midst of the bones there will be holes, where the brain is covered by the soft parts alone. From the nature of the case it will be understood that these apertures are mostly in the occipital or the parietal bone, according as the child lies mostly on its back or side.

**Recovery.**—This takes place in rickets by the removal of the insanitary conditions, or by the termination of the period of growth. The ossification advances in the cartilage and under the periosteum. The spongy bone produced under the periosteum becomes dense, and so the bone may be unduly heavy and thick, while it is stunted. The deformities having occurred in a rigid structure are rendered even more fixed by the completion of the ossification. There is, however, in the course of time an effort on the part of nature to restore the normal architecture of the bones; where they are bent absorption occurs on the convex surfaces and increased newformation on the concave; there may be considerable restoration of the proper shape in the long bones, but usually much less in the pelvis, head, and thorax.

**Literature.**—GLISSON, *De rachitide*, 1650 and 1671; BROMFIELD, *Chir. obs. and cases*, 1757, vol. ii.; STIEBEL, in *Virchow's Handb. der spec. Path. u. Therap.*, Bd. i., 1854; AITKEN, in *Reynold's Med.*, i., 1866; SMITH, in *Internat. Syst. of Surg.*, i., 1882; KASSOWITZ, *Die normale Ossification*, 1881; POMMER, *Osteomalacie und Rachitis*, 1885; NEUMANN, *Ueber fötale Rachitis*, 1881; MACEWEN, *Osteotomy*, 1880.

### III.—RETROGRADE CHANGES.

1. **Atrophy.**—The bones are liable to various forms of atrophy. One of the commonest forms is that described as **Atrophy from pressure**, which, however, is not a true atrophy but an absorption of bone. It is illustrated by the case of **Aneurysms** which in their growth meet with bony structures, such as the vertebræ (see Fig. 174, p. 421), or the sternum and ribs. The bone is eroded, while any related cartilage is preserved. In this way the intervertebral cartilages or the cartilaginous ribs may be rendered unduly prominent by destruction of the bone. A similar erosion of the bone is sometimes produced in the skull by the **Pacchionian bodies**, which may produce deep pouches in the internal table. Tumours also lead by pressure to destruction of bone.

**Senile atrophy** of the bones is a true atrophy. It consists in a general diminution in the organic matrix of the bone so that the



proportion of lime salts increases, but there is at the same time frequently a diminution in thickness or an increased porosity of the bone. In old age the neck of the femur is frequently shortened and forms a wider angle with the shaft. The calvarium in old people is frequently diminished in thickness and very translucent, but this is by no means constant, nor is it limited to old people.

**Atrophy from disuse and from nervous lesions** occurs chiefly in cases of paralysis. A case of pseudo-hypertrophic paralysis, for example, in the museum of the Western Infirmary, shows great atrophy of the bones of the lower limb. In infantile paralysis and locomotor ataxia there is also frequently atrophy of the bones at the joints. After **Fractures** which have not united, the bones may atrophy, sometimes from disuse but at other times from interference with the nutrient artery.

Finally, atrophy occurs in the bones without manifest cause. It may be a simple thinning or increased porosity of the bones, but in other cases it takes the form of softening such as is described in next section. Increased porosity may be associated with enlargement of the bones, as in the disease described by Paget under the name *Osteitis deformans*. (See under Hypertrophy.)

**2. Osteomalacia or Mollities ossium.**—This condition is liable to be confused with rickets on the one hand and simple atrophy on the other. It differs from rickets in being a disease of mature bone, rarely occurring in children. It leads, like rickets, to weakness of the bones, rendering them liable to various curvatures and deformities.

In its causation the disease is often obscure, but pregnancy seems in many cases the determining cause. It may come on during pregnancy and run a rapid course. It has also been observed in cases of insanity, and here the softness of the bones may lead to fractures from comparatively slight violence.

The disease usually begins in the bones of the pelvis, vertebræ, or ribs, but extends to the rest of the skeleton, continuing to progress up till death. In extreme cases the bones may be bent in any direction, and the parts exposed to pressure are curved inwards. In less severe cases there are various curvatures and partial fractures. The pelvis is specially liable to deformity, the acetabula being pushed inwards and the cavity narrowed. The bones as seen after death are soft and light so that they will float in water, and they are easily cut with the knife or scissors.

The lesion consists in a solution of the bone with a corresponding increase in the medullary spaces. It begins, as shown in Fig. 184, in a decalcification of the bony trabeculæ, in the parts next the medullary



spaces and Haversian canals, so that instead of these trabeculæ having in the fresh state a homogeneously opaque appearance, they show at their peripheries a transparent zone in which the bone corpuscles are visible but without their canaliculi. The appearance of these parts is precisely that of bone artificially decalcified by steeping in an acid. The decalcified parts may be rendered prominent by staining with carmine, the unaffected bone remaining uncoloured while the affected parts are stained. The disease advances by the decalcified parts dissolving so that the medullary spaces increase, while the decalcification further encroaches on the bony trabeculæ.

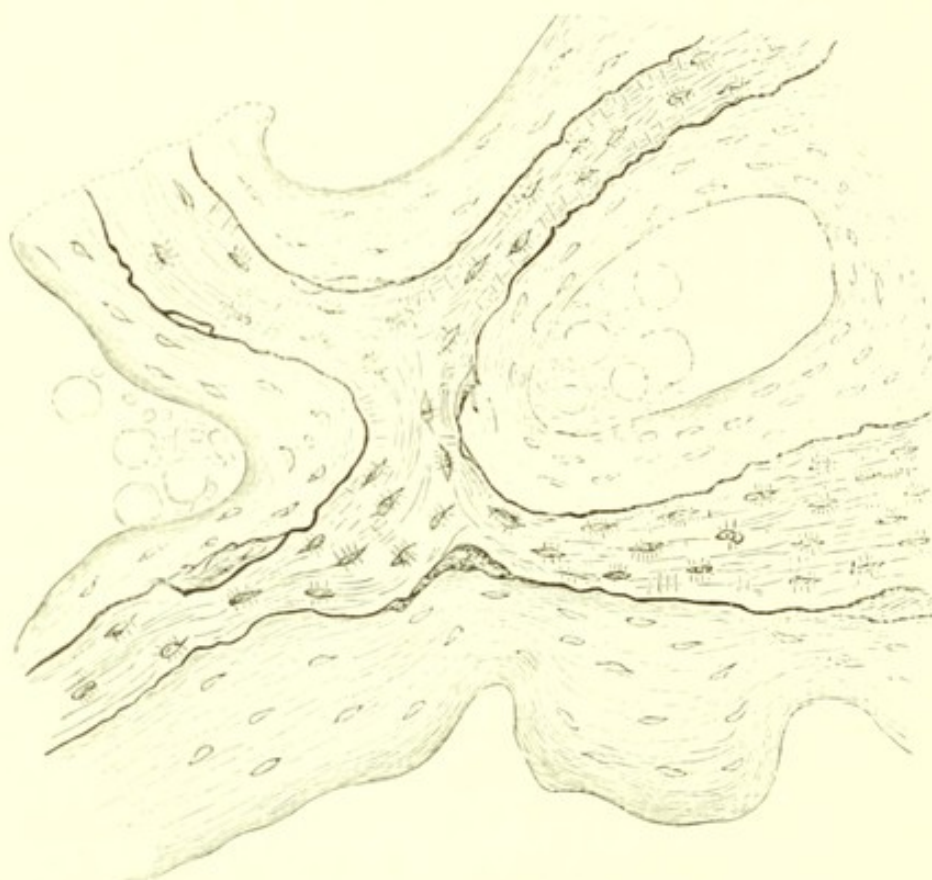


Fig. 184.—A fragment of bone from a case of osteomalacia. The central part shows the usual appearance of bone, while the marginal parts are transparent, being devoid of lime salts, although still showing bone corpuscles.  $\times 90$ .

The bone-marrow is in some cases replaced by round-celled tissue as if the process partly partook of an inflammatory character ; in other cases the adipose tissue of the marrow seems to undergo increase, so as to fill the enlarged spaces. This appears to have been the case in specimens examined by John Hunter, which he thus describes : "The component parts of the bone were totally altered, the structure being very different from other bones, and wholly composed of a new substance resembling a species of fatty tumour, and giving the appearance of a spongy bone deprived of earth and soaked in soft fat." In these cases the medulla forms a bright yellow, pink or deep crimson material, and when



examined microscopically presents free oil in great quantity with crystals of margarine.

**Necrosis** is considered in a special section after inflammation.

**Literature.**—JOHN HUNTER quoted by Paget, Lect. on Surg. Path., 1870, p. 103; STANSKY, Rech. sur l'ostéomalacie, 1851; ROKITANSKY, Handb. d. path. anat.; RIND-FLEISCH, Path. Hist.; COHNHEIM, Path. Anat., i.; POMMER, Unters. über Osteomalacie und Rachitis, 1885; REHN (a case in childhood), Jahrb. f. Kinderheilk., xii.; VINCENT, in Internat. syst. of surg., 1886; SHATTOCK, Path. trans., xxxviii., 1887.

#### IV.—INFLAMMATIONS OF BONE. OSTITIS, PERIOSTITIS, OSTEOMYELITIS.

**Introductory.**—Inflammations of bone are sometimes distinguished according as they affect the periosteum (*Periostitis*), the bone proper (*Ostitis*), or the bone-marrow (*Osteomyelitis*). Such a distinction, however, is not consistent with actual fact. We have already seen that in the growing bone the proper tela ossea is enveloped in an active cellular tissue, which is beneath the periosteum, in the substance of the bone filling its spaces, and in the medulla. In the adult the remains of this tissue is still present and capable of activity. In inflammations it is this tissue, which by some writers is together regarded as the bone-marrow, which is specially affected. The majority of cases of inflammation of bone occur in childhood, when this tissue is still active, and when inflammation occurs in adults the state of activity is restored. While this is true it will be understood that the phenomena of inflammation are frequently most manifest on the surface or in the marrow of bones, because the soft tissue is most abundant there. It is also to be remembered that the surface of bones is most exposed to injury, and that inflammations may thus arise which will chiefly affect the periosteum and superficial layers of the bone.

It will be convenient to divide the inflammations into the acute suppurative, and the chronic, although the former frequently pass into the latter.

**1. Acute and suppurative inflammations of bone.**—It has been pointed out in the earlier pages that suppuration is virtually always due to the action of pyogenic microbes. These may be directly introduced from without or may reach the bone by the blood, and hence two forms may be distinguished.

(a) **Acute traumatic ostitis.**—This occurs chiefly in cases of **Compound fracture** where septic processes have occurred in the wound. It used to be very common before the introduction of antiseptic surgery. The suppurative inflammation extends between the broken ends of the bones, and amongst the surrounding soft tissues, and abscesses fre-



quently form around the seat of injury. There may also be pus in the medullary cavity, disintegrating the bone-marrow. Extension of the septic process to the veins is not uncommon, so that thrombo-phlebitis and pyæmia result.

A suppurative inflammation involving the periosteum and superficial layers of the bone may occur in connection with a suppurating wound or abscess which has penetrated to the bones.

(b) **Acute infective ostitis.**—This name may be applied to a form of disease which has attracted much attention of late years and has been described under various names, such as **Acute periostitis**, **Malignant osteomyelitis**, **Epiphyseal ostitis**, **Infective ostitis**, **Pseudo-rheumatic ostitis**, **Typhus of the limbs**. These names refer to special features presented by different cases. The disease consists in an acute inflammation which nearly always goes on to suppuration, and occurs without any apparent sufficient cause.

**Causation.**—The essential cause here is a pyogenic microbe which in a number of cases has been identified as the *Staphylococcus pyogenes aureus*. In a case observed by the author a plate cultivation showed the presence of great multitudes of this microbe. Experiments on animals, already alluded to, show that when a bone is injured and cultures of this microbe are injected into the animal, the former will settle in the injured bone and produce a suppurative inflammation. In man the bone is always so far predisposed in respect that the disease is one of **Adolescents** in whom the tissues are excessively active, and there is usually also the history of an injury inflicted before the beginning of the illness. The bones most frequently affected are those most exposed to injury. The fact that boys are much more frequently attacked than girls, points in the same direction. It may, perhaps, be said that when, by accident, the microbes in question are present in the blood, perhaps by absorption from the alimentary canal, an injury, otherwise trivial, may determine their settlement in a bone, and so the disease may be induced.

**Character of lesions.**—The inflammatory process presents varying degrees of severity and extent, as well as considerable variety in its seat. The bones most frequently affected are the long bones, more especially the tibia, but it occurs in the pelvic bones, the vertebræ, especially the atlas, and in other situations.

In some cases the periosteum is chiefly affected, and to these the name of **Acute periostitis** is often given. The periosteum is swollen and hyperæmic, and suppuration usually ensues rapidly. The pus accumulates between the periosteum and the bone, and the periosteum is sometimes gangrenous. The bone beneath frequently undergoes necrosis, but this result is by no means constant.



In other cases there is a more diffused inflammation, affecting the bone and medulla as well as the periosteum, or there may even be a special involvement of the medulla. In these cases pus is present in the medullary cavity and in the substance of the bone. According to Ollier the inflammation frequently localizes itself in the neighbourhood of the ossifying borders of the shafts of long bones. This region as it is next the epiphysis (though a portion of the diaphysis) he designates **Juxta-epiphyseal**. The reason of this localization will be apparent from the fact that this is the region of greatest activity.

Along with these local phenomena there is commonly more or less fever, and this may be very great in comparison with the local phenomena. The designation, 'typhus of the limbs,' points to the occasional occurrence of such severe febrile phenomena.

**Necrosis** may occur in the several forms of this disease, but it is by no means uniformly present, its occurrence depending chiefly on local interference with the blood-supply, and perhaps also on the condition of the person affected. (See under Necrosis.)

**Pyæmia** occasionally develops as a result of the extension of the septic process to the veins.

(c) **Post-febrile osteitis**.—An acute osteitis in many respects similar to that just described, sometimes follows the acute fevers, such as acute rheumatism, typhoid fever, scarlet fever, measles, dysentery, etc.

(d) **Abscess in bone**.—Abscesses are met with in the cancellous tissue of the ends of the long bones, where they are probably the result of an infective or post-febrile osteitis. The abscess expands the end of the bone, which may attain large dimensions.

2. **Chronic inflammations of bone. Chronic osteitis**.—In its **Causation**, chronic osteitis is very varied. All sorts of irritants acting on the bones lead to it. Traumatic agents causing wounds and more especially fractures are a frequent cause. The processes concerned in the healing of fractures are essentially those of chronic inflammation. Necrosis, from whatever cause, leads to a chronic osteitis around the dead piece. Again, inflammations of joints, whether tubercular, traumatic, rheumatic, or other, lead to inflammations of the neighbouring parts of the bones, the agent which causes the inflammation in the joint acting also by absorption and in a diluted form on the bone. Syphilis and tuberculosis, although producing specific lesions, also lead to osteitis. Phosphorus produces a peculiar form of osteitis associated with necrosis. (See under Necrosis.)

The chronic inflammation produces activity in the soft structures of the bones. These become cellular so as to be converted into **Granulation tissue**. This will occupy the various situations already referred to



as those occupied by the active tissue of growing bones. It will be present in the medullary spaces, in the Haversian canals, and on the surface of the bone beneath the periosteum. It is an active tissue and produces changes analogous to those of the similar tissue in growing bones. These changes consist in absorption of the existing bone on the one hand and newformation of bone on the other. It is thus possible to distinguish a rarefying or destructive and a formative effect. These two processes commonly co-exist in the same case, but the one or other may predominate.

The **Rarefaction** of bone (sometimes called *rarefying osteitis*) occurs by enlargement chiefly of the smaller spaces and channels, such as the Haversian canals. It is manifested chiefly in the shafts of long bones, where the dense bone may be converted into a loose cancellous tissue. There is usually at the same time newformation of bone, so that the thickness of the shaft is increased. The granulation tissue acting on the bony trabeculae causes absorption of the bone, and it is believed that the giant-cells or myeloplagues are chiefly engaged in the process.

**Newformation** is a more constant and important effect. In all forms



Fig. 185.—Newformation of bone in connection with inflammation. The osteoblasts are getting surrounded by bony matrix and becoming bone-corpuscles.  $\times 85$ .

of inflammation there is liable to be thickening of the bones from newformation on the surface. As this takes place under the periosteum it is generally ascribed to periostitis, and this membrane is generally credited with a special power of bone-formation. It is, however, merely the local circumstances which determine the situation of the newformation, and there is not infrequently an encroachment on the medullary cavity as well as apposition on the surface. Moreover, the thickening is not necessarily all due to apposition. With rarefaction there may be an expansion outwards of the granulation tissue in the bone. The new-



formed bone has a rough tuberculated surface such as that shown in Fig. 186, p. 471. The newformation may follow on the process of rarefaction so that a dense thickened bone may be the result (*condensing ostitis*).

The process of newformation is similar to that in ordinary ossification. The granulation cells act as osteoblasts (see Fig. 185), surround themselves with osseous matrix, and remain buried in this as the bone-corpuscles.

**Literature.**—STANLEY, On diseases of bones, 1849; KLEBS, Beitr. zur path. Anat. der Schusswunden, 1872; ROSENBACH, Mikroorganismen b. d. Wundinfectionskr., 1884; GARRE, Fortschritte der Med., 1885; OLLIER, in Internat. Encycl. of Surg., vi., 1886.

#### V.—HYPERTROPHY OF BONE. HYPEROSTOSIS, PERIOSTOSIS.

The distinction between hypertrophy of bone and chronic inflammation is sometimes a difficult one to draw. In the former there is a newformation, as in the latter, and it may be only the absence of a definite inflammatory irritant which may determine the distinction. In true hypertrophy new bone is formed of a strictly normal character, and the result is a general or local enlargement of the bone. A general enlargement of a bone is called **Hyperostosis**, while a partial enlargement is **Periostosis**.

The **Causation** is in most cases obscure. It may be determined, as already mentioned (p. 180), by a prolonged hyperæmia, which gives rise to increased nutritive activity, but in general we have to do with a gradual, apparently causeless, enlargement.

A **General hyperostosis**, in which the bones generally are enlarged, has been described by Paget under the designation **Osteitis deformans**. As the name implies, Paget regards the condition as the result of inflammation. The disease usually affects in the first place the bones of the lower limb and the cranium. These undergo great enlargement, while at the same time their tissue is opened out as by a rarefying ostitis. The bones are also softened so that various curvatures are produced. The enlarged medullary spaces are filled as in inflammation with a tissue which is abundantly cellular. The name **Osteoporosis** is also given.

A peculiar feature in this disease is the frequent co-existence of tumours, which may be either sarcomatous, cancerous, or lymphatic (as in a case of Goodhart's), but are not seated in the bones.

A hyperostosis affecting single bones has been observed chiefly in the skull and face. These bones may be enormously enlarged, and the face in particular may take on characters which have suggested Virchow's name of *Leontiasis ossea*.



A **Periostosis** is more unusual apart from inflammatory enlargement, but sometimes partial enlargements of the processes of bones have been observed.

**Literature.**—PAGET, *Med. chir. trans.*, vol. lx., 1876; TREVES, *Path. trans.*, xxxii., 1881; SILCOCK, *ibid.*, xxxvi., 1885; ROBINSON, *ibid.*, xxxviii., 1887; GOODHART, *ibid.*, xxxix., 1888; CLUTTON, *ibid.*

## VI.—NECROSIS OF BONE.

Death of bone is of frequent occurrence and it leads to such obvious phenomena that the term Necrosis has been almost monopolized by surgeons for this condition.

**Causation.**—Necrosis in bone is nearly always the result of interruption of the blood-supply. From its frequency it might be supposed that bone readily succumbed to such deprivation, but the reverse of this seems to be the case. Macewen's observations in regard to transplantation seem to show this. (See under Transplantation.) Bone receives its supply of blood partly through small vessels which pass from the periosteum and partly by larger vessels, of which there is generally a special one for each bone, which penetrate the bone and are distributed from within, such vessels being specially named the nutrient vessels.

Necrosis is mostly brought about by the periosteum being raised, sometimes traumatically but generally as a consequence of inflammation, especially when pus accumulates between the membrane and the bone. In this way the periosteal supply is cut off, and the nutrient vessels may also be severed. According to Macewen the supply of the nutrient vessel of a long bone is generally sufficient to keep the bone alive even when the whole periosteum is stripped. This author records a case in which the whole diaphysis of the tibia was denuded of periosteum, but in which the persistence of the nutrient vessels caused all the bone to survive except a small superficial scale.

Necrosis is an occasional result of inflammation in the bone itself, and it occurs in the form of caseous necrosis in tuberculosis. Injuries, by separating a portion of bone, will sometimes produce necrosis, and will do so almost inevitably if septic processes coincide.

**Resulting conditions.**—The dead piece of bone is in itself inert, and undergoes merely passive changes. It retains its form and general appearance, but being macerated by the juices of the body it assumes the characters of macerated bone. It generally appears white and dry, and, as it contains little organic matter, it feels hard to the probe and gives a sound on being struck.

**Inflammation** is a constant concomitant of necrosis. If the latter has not been due to inflammation, then it leads to it secondarily. The dead



piece of bone seems to stimulate the surrounding living tissue, and to lead to a chronic inflammation. By means of a rarefying osteitis the Haversian canals and medullary spaces enlarge and, the bone immediately around the dead piece being replaced by granulation tissue, the necrosed piece becomes a **Sequestrum**.

At the same time newformation commonly occurs, chiefly in the subperiosteal tissue which has probably been separated from the dead bone, but also in the medulla. There is thus produced an irregular layer of new bone, as shown in Fig. 186. This new bone is largely subperiosteal and in many cases it forms an external shell which may exercise the function of support instead of the bone which has died. This external shell frequently confines the dead piece rendering it difficult of access for removal by the surgeon. It is generally provided with apertures called **Cloacæ** through which the pus of the original supuration has passed, and which have remained as canals or gaps in the new-formed shell.

The ultimate disposal of the dead bone occurs in different ways according to circumstances.

**Absorption of dead bone** is of frequent occurrence. In simple fractures it is probable that detached pieces of bone frequently die and become absorbed. Similarly in compound fractures pieces are not infrequently visible in the wound, having the dead white colour and hard feeling of sequestra, but if the wound remains aseptic they are absorbed. Again, at the ends of stumps sequestra often form, and they are sometimes, at least partially absorbed. In all these cases, and even in acute infective osteitis, the dead bone may be absorbed if septic changes be averted or overcome.

The absorption is effected by the **Osteoclasts**. In Figs. 187 and 188 the appearances are shown as observed by the author in a case of limited necrosis at the end of a stump, in which absorption was in progress. Here the Haversian canals and medullary spaces, as seen in Fig. 187, are enlarged and their borders have a worm-eaten appearance.

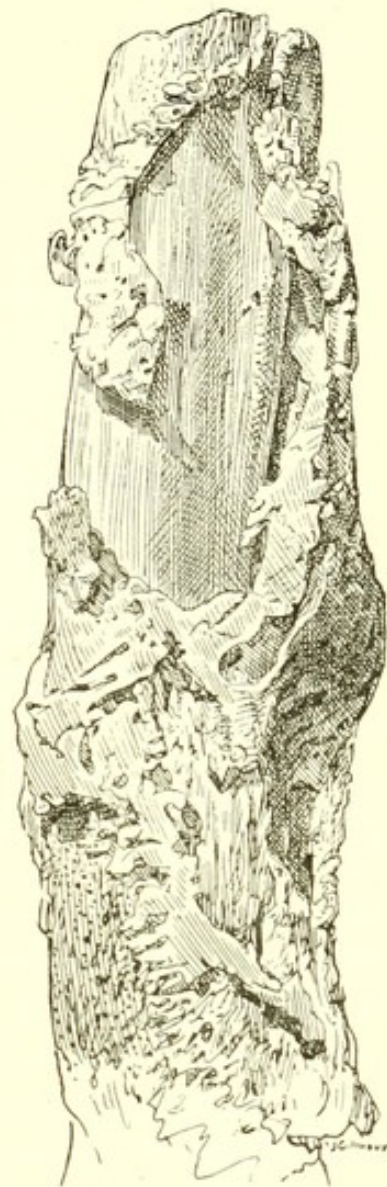


Fig. 186. — Necrosis of femur. The smooth necrosed piece is seen with irregular new-formed bone around.



This latter is shown in Fig. 188 to be due to the existence of Howship's lacunæ in which are giant-cells exercising their function of osteoclasts. Sometimes a regular row of such cells was visible, as if feeding on the

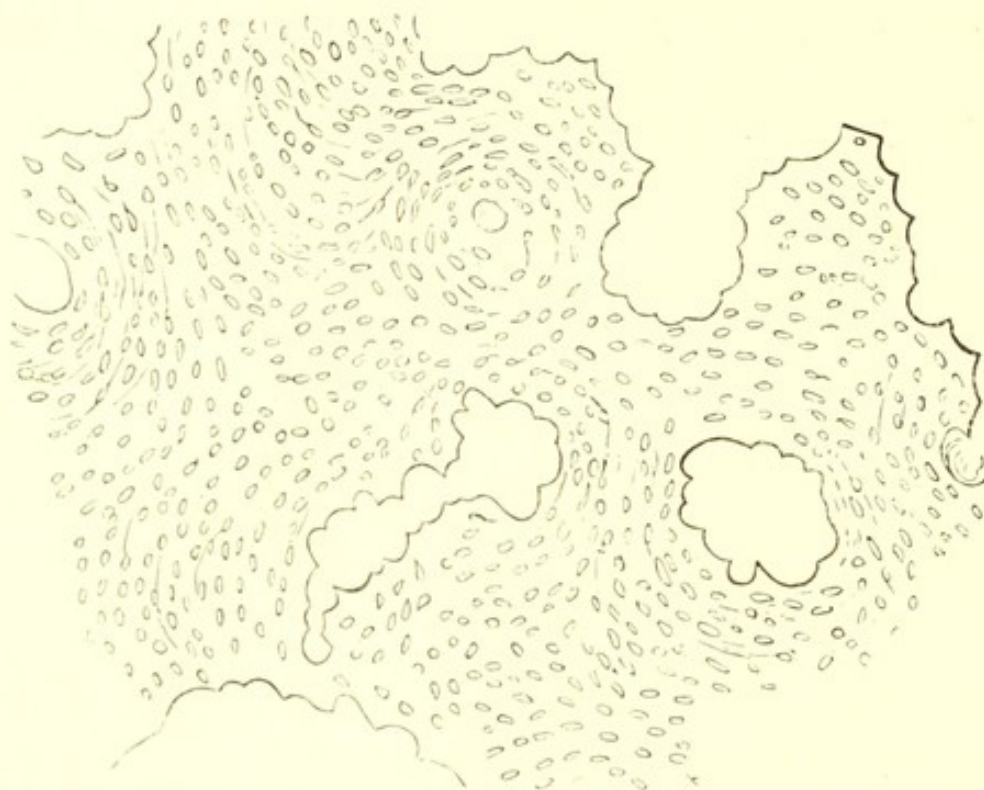


Fig. 187. —Section of a piece of necrosed bone which had been undergoing absorption. The margins of the bone at the medullary spaces and Haversian canals have an eaten-out appearance from the presence of Howship's lacunæ.  $\times 80$ .



Fig. 188. —Howship's lacunæ with giant-cells (osteoclasts) in them, and free giant-cells. From same piece of necrosed bone as preceding figure.  $\times 350$ .

bone, and they were occasionally almost buried in flask-shaped spaces (as at the left in Fig. 188). By this process bone may be entirely absorbed.



A similar process of absorption occurs, not only in pieces of bone which have died in the body, but in pieces of ivory or bone which have been introduced for therapeutic purposes, such as ivory pegs used in cases of ununited fracture, bone drainage tubes, etc.

**Absorption does not occur** where the dead bone, having become septic, is itself a source of serious irritation. Thus a piece of dead bone may lie for years in a suppurating cavity almost unchanged. Cornil and Ranvier, for example, figure a transverse section of a sequestrum of the femur which had remained thirty years bathed in pus. In this time it had undergone no corrosion and had all the characters of macerated bone. The position of the sequestrum inside the shell of new bone often causes its retention, and its removal is only effected by partial destruction of the external shell.

**Phosphorus-necrosis.**—This condition occurs chiefly in persons who are employed in workshops where lucifer matches are manufactured, and who are thus exposed to the vapour of phosphorus. The vapour acts locally, on the jaws chiefly, and it is said that persons with carious teeth are particularly liable to be affected. The phosphorus produces an inflammation mainly of the periosteum, resulting in a great newformation of cancellated bone on the surface of the jaw. This bone may afterwards become condensed by the ossification encroaching on the medullary spaces. After a time the inflammation leads to suppuration, the pus forming between the new-formed bone and the original bone of the jaw. From this results a necrosis of the jaw which may involve the entire bone, and may be accompanied also by necrosis of the new-formed bone. The resulting suppurations are usually fatal in their results, but after removal of the sequestrum healing may occur.

From the observations of Wegner it appears that phosphorus acts as a general stimulant to the structures concerned in the formation of bone. The phosphorus vapour acting directly stimulates so violently as to produce the inflammation and necrosis just described, and this result has been produced also in rabbits exposed to the vapour of phosphorus. When given in small doses internally, phosphorus produces in growing animals a distinct stimulation of the process of formation of bone. In such cases the bone produced at the ossifying border of the cartilage is not a spongy bone, but a dense layer, and there is also an unusually dense bone produced beneath the periosteum. It is noteworthy that in growing animals to whom small doses of phosphorus were given, while insufficient quantities of lime were supplied, there was a great production of osteoid tissue, somewhat like that produced in rickets.

**Literature.**—MACEWEN, *Annals of Surgery*, 1887; WEGNER, *Virchow's Arch.*, lvi., 1872; COATS, *Glasg. Med. Jour.*, vi., 1874; BIBRA und GEIST, *Krankh. d. Arbeiter in der Phosphorfabriken*, 1847; TRÉLAT, *De la nécrose causée par le phosphore*, 1857; THIERSCH, *Arch. d. Heilk.*, 1868; WEGNER, *Virch. Arch.*, lv., 1872; CAMERON, *Glasg. Med. Jour.*, xvi. 112, 1881.



## VII.—REGENERATIVE PROCESSES. HEALING OF FRACTURES, TRANSPLANTATION.

In the various regenerative processes the structures above referred to as having to do with the growth of bone are concerned. These structures become cellular, their existing cells multiplying by karyomitosis, and they thus renew their powers of newformation. The processes concerned are really those of chronic inflammation.

1. **The Healing of fractures.**—In the case of simple fractures or of those which are protected from septic contamination, the process of healing generally begins soon after the infliction of the injury.

The actual injury induces an acute inflammation. There is hyperæmia, serous exudation and exudation of leucocytes, and there is also a certain amount of hæmorrhage. These inflammatory manifestations are present in the torn periosteum, and in the medulla. The acute inflammation, however, usually subsides in a few days; its prolongation, as in septic cases, is inimical to healing. Any considerable accumulation of blood is also inimical to healing, but as a general rule the effused blood is carried off by the inflammatory exudation or transudes into the tissues around and so passes into the lymphatics.

On the subsidence of the acute inflammation the soft structures show active changes. By a process of karyomitosis (as observed by Krafft) the subperiosteal and medullary cells proliferate, so as to form granulation tissue. A similar process is visible in the endothelium of the vessels, by which means new vessels are formed. The tissue of the bone itself partakes in the process, so that it is opened out by a rarefying ostitis. We have thus a vascular granulation tissue formed in these three situations, but its amount varies greatly according to circumstances.

From this tissue the **Callus** is formed. This name is given to the new-formed tissue which in the first instance unites the broken ends of the bones. The callus, as shown in Fig. 189, consists under varying circumstances of **Different forms of connective tissue**. The formative cells produced in the way mentioned above have the characters variously of osteoblasts, fibroblasts, or chondroblasts, and may develop bone (*c, c, c* in figure), cartilage (*b, b, b*), or connective tissue (*d, d, d*). The form of tissue thus produced seems to depend on the condition of the parts as to movement. In parts where there is little movement bone will form, where there is much movement fibrous tissue, while cartilage will take an intermediate position. From this it arises that in man, where the bones are usually kept rigid by splints, the callus is formed chiefly of bone, whereas in animals it is largely formed of cartilage and



fibrous tissue. But in man, where circumstances do not allow of the parts being kept at rest, then we have cartilage and fibrous tissue along



Fig. 189.—Longitudinal section of a fractured rib, showing callus etc.; the section is imperfect at the upper part. *a, a, a, a, a*, the broken extremities of the rib dove-tailing into each other; *b, b, b*, new-formed cartilage constituting part of the callus; *c, c, c*, new-formed bone constituting the external layers of the callus; *d, d, d, d*, new-formed connective tissue constituting the more immediate bond of union between the broken ends,  $\times 11$ .

with bone. This was the case in the specimen from which Fig. 189



was taken. A number of fractures had occurred in the ribs in an insane person, and they were only discovered after death.

Circumstances also determine the **Amount of callus**. When the parts are kept rigid the amount is small, probably because the irritation is slight, but where the parts are moveable there will be much callus, as we find in fractures of the ribs in man and in fractures generally in animals.

It not infrequently happens that a **Piece of bone is detached** and isolated. This, however, does not interfere with the process of healing. If small, the piece of bone will survive, and, having acquired vascular adhesions, will undergo a rarefying osteitis and assist in the production of callus. If large, or if unfavourably situated, it may die, and in that case it undergoes absorption by the granulation tissue (see under Necrosis). In compound fractures it is not uncommon to see a piece of dead bone eaten into and penetrated by granulations.

The callus produced as above is generally described as **Provisional callus**, because much of it is removed before complete restoration. Names are also given according to the position of the callus. **External callus** is formed at the surface from the subperiosteal layer, **Internal callus** is formed from the bone-marrow and endosteum, while **Intermediary callus** is between the ends of the bones and produced, to some extent at least, from the bone itself.

When the bone has become fully united the parts of the callus not necessary for the preservation of the continuity of the bone are disposed of. The intermediary callus becomes, for the most part, the permanent bond of union, while the external and internal disappear except in so far as they are needed to fill up gaps where the bones have not been exactly in apposition. Sometimes the bones are considerably out of position, and a large amount of new bone remains permanently as the bond of union.

From what has gone before it will appear that sometimes the amount of bone produced will be small. If the parts are not kept at rest or if the blood-supply be insufficient, from tearing of vessels or otherwise, there may be an imperfect production of callus. In some cases there is even an atrophy of the broken ends of the bones. In this way may arise **Ununited fractures**, in which the bones are connected with fibrous tissue. In some cases there is even a kind of **False joint** or **Pseudoarthrosis** formed, with smooth articulating surfaces and an approach to a synovial cavity.

2. **Transplantation of bone**.—It has usually been stated that, when pieces of the periosteum are transplanted, they proceed to the formation of bone. This is only true in regard to the deep layer of the



periosteum, or more properly the subperiosteal layer. The periosteum proper, consisting of fibrous tissue, is incapable of producing bone. On the other hand the bone itself, or the tissue in its various cavities, as well as the endosteum, are capable, as we have seen, of forming bone. Hence a transplantation of periosteum, including the subperiosteal layer, or of pieces of bone, may be followed, not only by survival of the pieces, but by newformation of bone from them.

**Transplantation of the periosteum** is not an uncommon pathological process. The periosteum may be raised by the accumulation of pus beneath it, or it may be stripped off by an injury, and in that case, if the ossifying layer be preserved, ossification will occur. This fact is sometimes taken advantage of by surgeons, who will detach the periosteum, and displace it with a view to the ultimate formation of bone.

The **Transplantation of bone itself** has been demonstrated chiefly by Macewen. It is apparent from his observations that bone, contrary to what might be expected, has great power of retaining its vitality when detached from its connections. A piece of bone transplanted will, even when of considerable size, survive long enough to acquire fresh vascular connections, and, in accordance with what occurs in transplanted structures generally, it will often proceed to grow in its new position. Accidental transplantation must often occur in connection with fractures, and the detached piece of bone, as mentioned above, usually survives.

Advantage has been taken of these facts in an ingenious way by Macewen. In the operation of trephining, a circular piece of bone, often of considerable size, is removed from the skull. This piece of bone may be preserved during a subsequent operation on the brain and reimplanted in the gap, at the close of the operation. If septic contamination is prevented, the bone survives and grows into its place, filling up by new-formed tissue the interval made by the saw.

Still more interesting are the results of transplantation in the treatment of a case in which the shaft of the humerus had been lost by necrosis. In this case pieces of bone removed from the femurs of other patients (in the operation of Osteotomy) were broken up into small pieces and inserted amongst the muscles into a sulcus where the shaft of the bone should have been. By three operations the bone was restored to the extent of four and a quarter inches, and there was a subsequent growth in the course of seven years of an inch and three-quarters. This case demonstrates that pieces of the bone removed from one person to another not only survive but grow in their new position. Transplantation from one of the lower animals does not succeed, the foreign bone acting as an irritant.

**Literature.**—MALGAIGNE, *Traité des fract. et luxat.*, 1855; GURLT, *Handb. d. Lehre v. d. Knochenbrüchen*, 1862; KRAFFT, in Ziegler and Nauwerk's *Beiträge*, 1884; OLLIER, *Traité de la régénération des os*, 1867; MACEWEN, *Phil. trans. of Roy. Soc.*, 1881, and *Annals of Surg.*, 1887.



# VIII.—SPECIFIC NEWFORMATIONS. TUBERCULOSIS, SYPHILIS, ACTINOMYCOSIS.

1. **Tuberculosis of bone.**—This is an exceedingly frequent affection, and one attended usually by very serious results. The lesions to be here described are not universally recognized as tubercular in character, although, as observation widens, the dissentients from this view diminish. The term **Scrofulous**, here as in other structures, is equivalent to tubercular. **Caries**, is another term which, as will be explained below, is nearly equivalent to certain forms of tuberculosis.

**Causation.**—The tubercular bacillus in most cases reaches the bones by the blood. In a large proportion of cases both joints and bones are affected, and it is not determined to what extent the one or the other is the primary seat. It is demonstrable that primary disease of the bones frequently extends to the joints, and it may afterwards attack other bones entering into the articulation. On the other hand when primarily affecting the joint it may extend to the bones concerned in the joint.

The tuberculosis begins in the spongy parts of the bones. It is very common in bones which are largely composed of spongy tissue, as the

vertebræ and the bones of the hands and feet, while in the long bones it affects chiefly the articular extremities. In the case of the vertebræ, it mostly affects simultaneously the proximate surfaces of two bodies (see Fig. 190), as if it took origin in the intervertebral cartilage. In this respect the author's observations correspond with those of Wilks and Moxon.

While spongy bone evidently forms a favourable nidus for the tubercular bacillus, there are two further predisposing elements frequently traceable. Tu-

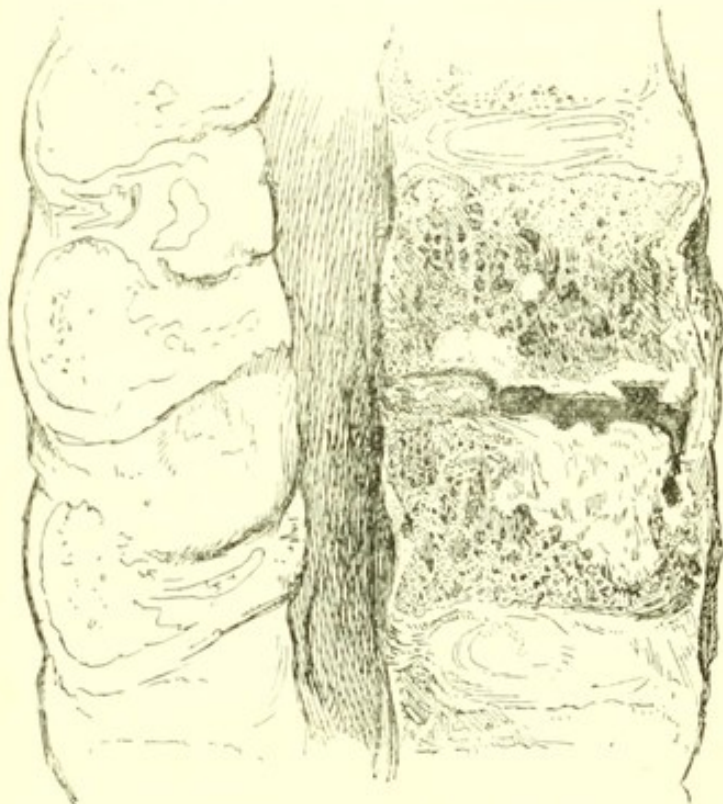


Fig. 190.—Tuberculosis of vertebræ. The intervertebral cartilage is destroyed and the disease, as shown by the white colour (due to caseous necrosis), is invading the bodies. In the upper there is a small isolated patch.

berculosis is most frequent in young persons, the actively growing bone



apparently predisposing to it. This applies especially to the bones of the extremities, and to a less degree to the vertebræ. Injury is the other predisponent. This is a fact of clinical observation which has been confirmed by the experiments of Schüller, who found that after injecting tubercular bacilli into the blood, he could induce tuberculosis of the joints by inflicting injuries such as otherwise would be readily recovered from.

**Characters of the lesion.**—The tuberculosis affects primarily the soft tissue or medulla contained in the meshes of the cancellated tissue, and secondarily the bony trabeculæ. These spaces become occupied by round-celled or granulation tissue, in which tubercles with their typical giant-cells are visible. The granulation tissue eats into the bony trabeculæ, so that the latter may be entirely destroyed in the affected area. More frequently necrosis occurs in the tubercular newformation before the bone is entirely destroyed. The form of necrosis is caseation, and in a case of any standing the presence of the disease is evidenced by the presence of yellow caseous matter occupying a certain portion of the cancellated tissue. This is shown in Fig. 190. In this area there are the thinned remains of the bony trabeculæ which are also necrosed. As the disease is an advancing one, the more recent affection will extend beyond the caseous area.

The bone so affected has lost much of its mechanical power of resistance, so that in the case of the vertebræ the bodies will collapse and lead to acute curvature, as in Fig. 191, while in other cases the bones will be eroded and worn down by the friction at the joints. In some cases the necrosis leads to the formation of a **Cavity** in the bone. This may be from softening of the caseous tissue, but in some cases the necrosed piece is separated as a kind of **Sequestrum** which may be found lying in the cavity. In either case the walls of the cavity will be lined with tuberculous granulation tissue. In the case of small bones, such as those of the foot or hand, which are thus excavated, there may be a complete collapse of the bone, so that it will be more or less destroyed.

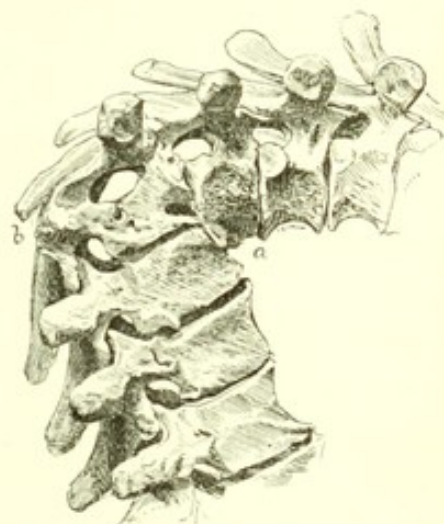


Fig. 191. — Acute curvature from tuberculosis. The bodies of two vertebræ have been destroyed, while the ones above and below have coalesced at *a*. The spinous processes are also ankylosed.

It is to some of the conditions indicated above that the term **Caries** is sometimes applied. The characteristics of caries are, rarefaction of the bone, which here is produced by the encroachment of the tuberculous tissue; undue softness of the bone, so that it is eroded and generally exposed at an articular surface or in the wall



of a cavity; the presence of caseous matter, along with prominent and flabby granulation-tissue, whence the name fungous caries. (See further under Diseases of the Joints.)

The tuberculous bone is a centre of irritation, and there usually follows a suppurative process which is frequently slow in developing. The resulting **Cold abscesses** are most typically seen in connection with tuberculosis of the vertebræ. The matter, consisting of caseous debris with pus-corpuscles and serous fluid, may travel considerable distances, forming, according to the place where it comes to the surface, the **Lumbar, Psoas, or other abscess**. The whole track of the abscess is liable to be infected by the tuberculosis so that there may be an extensive tubercular surface.

**Healing of the tuberculosis** occurs not infrequently. It may take place before any considerable extension of the process, the tissues

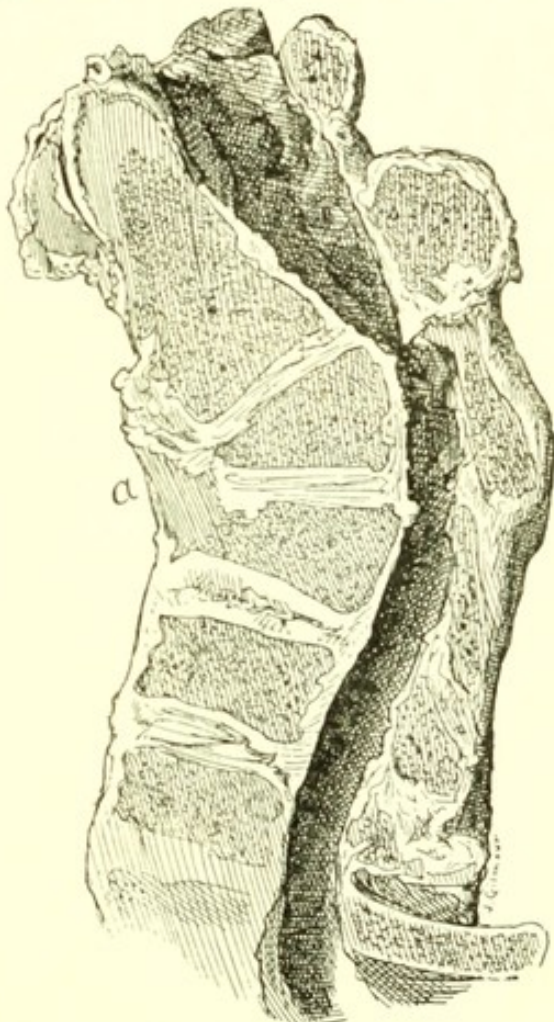


Fig. 192.—Healed tuberculosis. The bodies collapsed anteriorly. Dense bone produced at *a* to form a support. Permanent curvature and narrowing of spinal canal.

overcoming the tubercular infection, and ultimately absorbing the necrosed products. But it also occurs after the process of softening and formation of abscesses. In that case when the tuberculous matter has been cleared out, healthy granulation tissue will be produced. This may produce new bone, so as to some extent to regenerate and replace that which has been lost. Where there has been much erosion or collapse, the newformation will do little towards restoration, and will rather tend to fix the bones by **Anchylolysis** in the position which they may have assumed. In Fig. 192 for example, the anterior parts of the bodies of the third and fourth cervical vertebræ have been destroyed and a piece of dense bone (at *a*) has been formed to act as a support. This has confirmed the acute curvature which is shown by the direction of the spinal canal.

In Fig. 191 also, there has been a complete collapse of two bodies, whose spinous processes have coalesced (at *b*) while the bodies of the vertebræ above and below have also coalesced (at *a*).



While healing may thus occur, the tubercular virus may still linger about the parts, and on the occurrence of a favourable opportunity it may renew its advance. In the case from which Fig. 191 was taken, for example, the history showed what appeared to be a complete recovery (with curvature) three years before death, but a more recent onset ending in tubercular pleurisy led up to the fatal issue.

The tubercular process is accompanied by the ordinary phenomena of chronic inflammation, so that in the neighbourhood there is usually newformation of bone, chiefly subperiosteal, and the surface of the bone may be rough with irregular projections.

**2. Syphilitic affections of bone.**—The **Lesions** of tertiary syphilis have usually their seat of origin in the periosteum, although the subjacent bone may be simultaneously involved. The condition may be briefly described by stating that gummata are produced, while inflammation occurs in the neighbourhood.

Gummata of the usual structure are produced in the internal layers of the periosteum, and there is thickening of the periosteum around. This membrane being tightly stretched over the tumour, the swelling is hard to the feeling, but often with a certain elastic resistance. The gumma undergoes caseous necrosis in its central parts, but advances peripherally, and as it advances against the bone it causes erosion of it. The advance is in the first instance along the vessels, and as erosion occurs around them a worm-eaten appearance is produced in the bone. Beneath the gumma, which has most frequently its seat on the bones of the skull, especially the frontal bone, there may be thus considerable loss of substance, so that even the entire thickness of the skull may be penetrated. The caseous material often undergoes softening, and suppuration ensues; there may be very obstinate and even progressive ulcers produced. Without ulceration the gumma may undergo resolution, and a cicatricial depression result, or after ulceration healing may occur.

Along with these processes immediately related to the gumma, there are, in the neighbourhood, conditions referable to inflammation. The bone around is condensed by newformation filling up the medullary spaces, and there is sometimes a thickening of the bone by subperiosteal inflammation. This is much less common in the skull than in the long bones, especially those of the legs, where it sometimes leads to a very striking hypertrophy of the bone, whose surface is exceedingly rough from the loss of substance in some parts, and the irregular newformation in others. The bone around the syphilitic defect or ulcer is dense bone, and in this respect contrasts very markedly with that around the ulceration in tuberculosis.



**Congenital syphilis** leads to important changes in the bones, which, according to Wegner, occur in a large proportion of syphilitic fetuses. The lesions affect the structures concerned in the processes of ossification, and in this respect resemble those of rickets, but they partake more of the inflammatory character, thus agreeing with the lesions of hereditary syphilis generally (see p 197).

In the **Cranial bones** the periosteum is thickened by inflammation, and this leads to softening of the bone beneath. **Craniotabes** is a frequent result, the bone being lost in parts exposed to pressure when the child is lying, and the brain being so far covered only by the soft parts.

In the **Long bones** the ossifying border is the part affected. The cartilage cells undergo excessive proliferation, as in rickets, and there is an irregular calcification of the matrix. Beneath this the proper ossifying zone frequently shows a newformation of granulation tissue or pus, so as to produce a soft layer almost interrupting the continuity of the bone. There may be a partial necrosis of the calcified cartilage. The whole process occurs very irregularly, the ossifying part of the bone being swollen and occupied by calcified and proliferating cartilage and inflammatory tissue, while true ossification is delayed and occurs irregularly. A not infrequent consequence is a partial fracture of the bone or a **Separation of the epiphysis**.

The disease affects many bones, but may present different degrees. According to Wegner the usual seats, in order of frequency, are, lower end of femur, lower ends of bones of leg and forearm, and upper end of tibia.

3. **Actinomycosis in bone**.—This affection occurs in the bones of the face, and sometimes, by extension, the vertebræ, sternum, etc. The lesions somewhat resemble those of tuberculosis, the new-formed granulation tissue opening up the bone and leading to caries. The granulations, however, are more exuberant and there is no caseous necrosis, but rather a tendency to suppuration.

**Literature.**—*Tuberculosis*—NELATON, L'affection tuberculeuse des os, 1857; VOLK-MANN, Arch. f. Klin. Chir., iv.; MENZEL, Die Häufigkeit der Caries in den verschied. Knochen, *ibid.* xii; FRIEDLÄNDER, in Volkmann's Sammlung, No. 64; SCHÜLLER, Ueber die Entstehung d. tub. Gelenkleiden, 1880; VINCENT, in Internat. Encycl. of Surg., vi., 1886 (with French literature). *Syphilis*—VIRCHOW, Krank. Geschwülste, ii., 1865; CANTON, Path. trans., xiii., 1862; RICORD, Traité des malad. vénér., 1851; LANG, Vorles. üb. Syph., 1885; WEGNER, Virch. Arch., l., 1870; STILLING, *ibid.* lxxxviii., 1882; MÜLLER, *ibid.* xcii., 1883; PARROT, Arch. de Phys., iv., 1872; CORNIL et RANVIER, Man. d'hist. path., i. 444, 1881.



## IX.—SPINAL CURVATURES.

**Introductory.**—The spinal column is composed of vertebræ whose bodies are separated by elastic fibro-cartilages. The vertebræ articulate with each other at four other points, two on the upper and two on the lower surface of the arch. They are also bound together by firm ligaments in front of and behind their bodies. The effect of these various connections is that, even when the spine is separated from all its attachments except the ligaments, it retains its general form, and its natural conformation may be studied after its removal from the body.

When seen in profile the spine presents the well-known antero-posterior curves, the convexity being forwards in the cervical and lumbar regions, and backwards in the dorsal. These curves are capable of considerable variation in the movements of the body. The whole of the curves may be obliterated and converted into a general convexity backwards by stooping forwards, as when, with the arms extended and the legs straight, an attempt is made to touch the toes with the tips of the fingers. By arching the body backwards the dorsal curve may be partially obliterated, and a general convexity forwards produced. It appears, therefore, that the spine is capable of considerable antero-posterior movement. These antero-posterior movements imply a considerable degree of compressibility of the intervertebral cartilages. The combined cartilages occupy about a fifth of the entire length of the spinal column, and their compressibility may be inferred from the fact that during the retention of the erect posture the entire length of the column gradually diminishes, so that the stature is usually half an inch to three quarters less at night than in the morning. This is believed to be due chiefly to the compression of the cartilages, which recover at night when the recumbent posture is assumed. The antero-posterior movement of the spine is freest in the cervical and lumbar regions, and most limited in the dorsal.

The spine is capable to a much more limited extent of lateral movement. The articulating processes, being situated on either side of the arches, prevent any considerable lateral deviation, as they become locked against each other when that occurs. If the surfaces of these processes were horizontal, facing one another above and below, then they might allow of freer lateral movement, but the more they assume the perpendicular position, and the more they face inwards and outwards, the greater is the impediment to lateral movement. It will be found that on passing from above downwards the articulating surfaces assume more and more of a perpendicular position. In the cervical region they are oblique, and face slightly inwards and outwards; in the dorsal they are more perpendicular and face nearly forwards and backwards, while in the lumbar region they are nearly perpendicular, and face each other nearly inwards and outwards. In this way it occurs that while lateral movement is limited in all regions it is almost impossible in the lumbar region. For a similar reason, twisting of the spine on its axis is possible to a very limited extent.

The question of the existence of a natural lateral curvature has been matter of dispute. It is generally stated that there is a slight lateral deviation to the right in the upper dorsal region, and this is usually ascribed to the more frequent and



forcible exertion made with the right arm; but the existence of this curve has been seriously questioned (Adams). The late Dr. Foulis in 110 post-mortem examinations found lateral deviation in no less than 58 cases. He did not observe it specially in the upper dorsal region or towards the right, and concluded that it was due to the positions habitually assumed by the persons at their various trades. We may perhaps conclude that normally there is no lateral curvature, but that a very slight permanent deviation is often assumed when a frequently repeated position of the body predisposes to it.

**Forms of spinal curvature.**—The function of the spine is to support the structures attached to it, and to hold the head erect, the latter function being in man the more prominent one. Any single curvature of the spine will have the tendency to remove the head from the erect position, and tilt it backwards or forwards or to one side, and in order to preserve the erect position there is required a curvature in the opposite direction. The natural antero-posterior curves are in this sense mutually compensatory, the lumbar restores the position lost by the sacral curve, and the cervical that of the dorsal. When abnormal curvatures occur there is a tendency to a similar compensation, so that these curvatures may be divided into **Primary** and **Secondary**. It will not be necessary to consider in detail the secondary curvatures; their amount and direction may be inferred from those of the primary ones. It may be stated, however, that there are, frequently, several secondary or compensating curves, the spine presenting several sinuosities in order to reach the stable position for the head.

1. **Antero-posterior curvature.**—There are two quite distinct forms of antero-posterior curvature, the curve in the one form being rounded, and mainly an exaggeration of the natural curvature, and in the other sharp or angular.

(a) **Angular curvature or Pott's disease of the spine.**—This depends on disease or injury to the bodies of the vertebræ. For the most part it is a **Local tuberculosis** of one or more of the bodies with the caseous necrosis and caries described above. The softened bodies of the vertebræ give way under the superincumbent weight, and the spine is bent at a sharp angle, the spinous processes becoming unduly prominent behind. The affection of the bodies may be unsymmetrical, and if collapse be more at one side than the other the angular curvature will not be exactly antero-posterior.

This form of curvature is more rarely due to traumatic causes, as crushing of the bodies by heavy weights falling on the head or back, or by a fall from a height. This condition is shown in Fig. 193 where a vertebra is seen crushed and with part of it displaced against the cord. This may lead immediately to an angular curvature, or may be the starting-point of a necrosis and suppurative inflammation.



It is worthy of special notice that angular curvature is much more obvious in the dorsal region than elsewhere, as it here increases the natural posterior curvature. In the cervical and lumbar regions, where the natural convexity is forwards, there may be a rounded posterior curvature produced, or, even with extensive disease, there may be very little posterior curvature visible externally.

The spinal cord is not necessarily injured by angular curvature, but in many cases it is crushed and interrupted. It may be so even during the process of healing, the sclerosis or condensation of the vertebræ sometimes causing further shrinking and increase of the curvature.

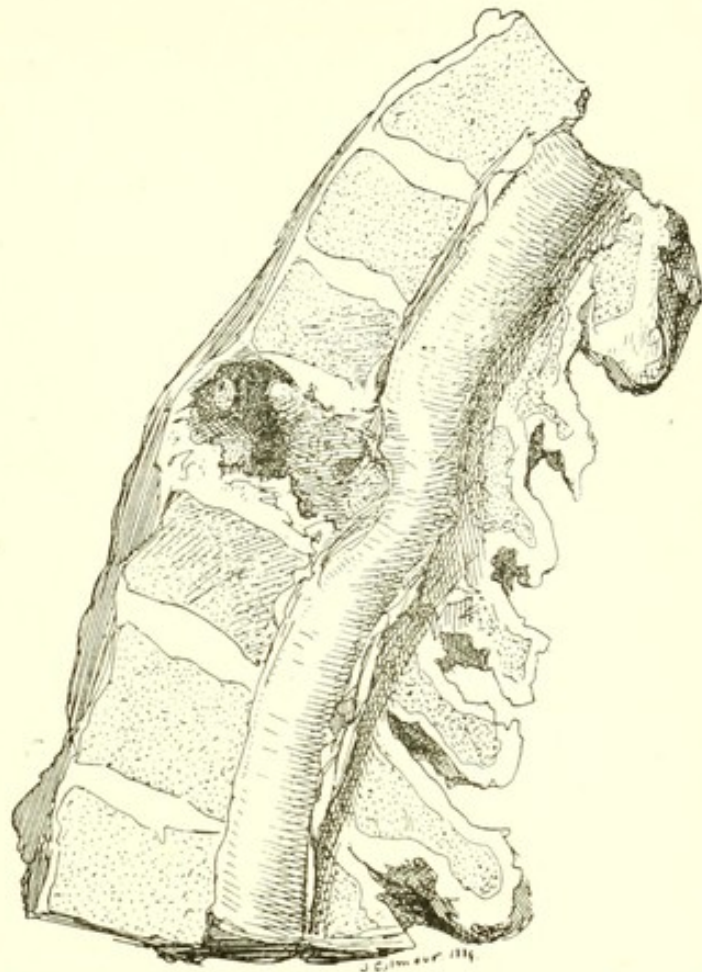


Fig. 193.—Vertebra crushed and part of it displaced backwards so as to injure the cord. There was acute curvature and paralysis. (The curvature is not shown in figure, as the parts have been replaced so as to display the lesion.)

(b) **Kyphosis or Posterior curvature.**—In angular curvature the convexity is backwards, and the name kyphosis is sometimes applied. The term is more commonly employed to indicate a more gradual rounded curvature. This is mainly an exaggeration of the normal curvature with the convexity backwards in the dorsal region, and is due chiefly to muscular weakness or a habit of stooping. It is most frequent in children and old people, leading in its most exaggerated form to hump-back, and in a lesser degree to round shoulders.

(c) **Lordosis or Anterior curvature.**—In this condition the convexity of the curve is forwards, and the tendency of it is to throw the head backwards. It is most common in the lumbar region, and in the majority of cases is due to rickets. Rickets when it affects the pelvis causes the sacrum to assume a more horizontal position than normal, and in order to retain the erect position the lumbar anterior curve is exaggerated. It may also be produced by congenital dislocation of the hip-joint (stated by Adams to be of considerable frequency), and ankylosis of the hip. There may be also, but rarely, a direct lordosis



in the lumbar region from rickets, the natural curvature being increased by reason of the softness of the bones. In the dorsal and cervical regions lordosis is very uncommon.

2. **Lateral curvature, Rotatory curvature, Scoliosis.**—In the introduction to this subject we have seen that the mechanism of the spine allows of exceedingly limited lateral deviation. But it often happens that from habitual faulty positions at work or otherwise there is a frequently repeated tendency to lateral deviation. In that case, as a direct displacement is not possible, there may be, especially in persons constitutionally weak, a deviation with partial **Rotation** of the vertebræ. The faulty position may be merely the result of a bad habit, of standing on one leg for instance, so as to cause obliquity of the pelvis; or it may be from sitting in a constrained position at study or manual labour, so that either the pelvis is oblique or the arms are used particularly in one direction so as to displace the centre of gravity to one side; or there may be a hysterical contraction of the muscles of the scapula lasting for a long period and altering the centre of gravity; or there may be an obliquity of the pelvis from one leg being shorter than the other, as

sometimes in rickets. There is also softening of the bones in rickets, and very severe curvature may occur when this is associated with obliquity of the pelvis.

The mechanism of this rotation will be understood from the accompanying diagram (Fig. 194). The dark outline indicates the natural position of the vertebra, the dark straight lines indicating the natural axis and the direction of the transverse processes. When rotation occurs the body moves round so as to present to one side in the direction of the lines *aa* and *aa*, while the spinous process deviates very little. In the figure the centre of rotation is just behind the tip of the spine,

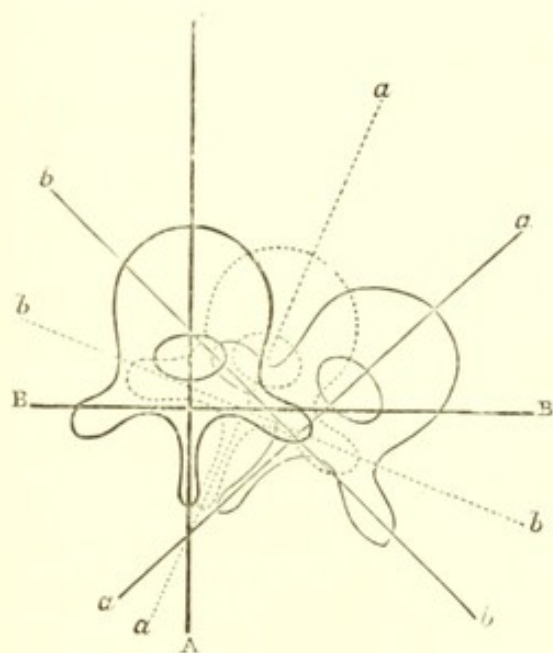


Fig. 194.—Diagram showing rotation of vertebra in lateral curvature. The various lines show the positions of the body and transverse processes in the different degrees of rotation. The axis of rotation is behind the tip of the spinous process, where the three lines meet. (ADAMS.)

and there is a slight deviation of it; but the centre of rotation may be at the tip of the spine, and with very marked rotation of the vertebræ there may be no deviation of the spines.

Lateral rotatory curvature is met with chiefly in the dorsal and lumbar regions, being, as a rule, much more extreme in the former, pro-



bably from the greater mobility there. There is usually curvature in both these regions, the two curves being in opposite directions, and one of them in a certain sense secondary. It is not possible, however, to distinguish rigidly between primary and secondary curves, as the two form nearly *pari passu*; a slight deviation to one side will result in a similar curve to the opposite side, and they will increase together.

We have seen above in connection with Fig. 194 that the plane of the transverse processes is altered by the rotation of the vertebræ. An examination of that figure will show that the transverse process on the side of the convexity is considerably displaced backwards, and in actual cases it may be felt projecting under the skin. If the curve be in the dorsal region the rib will be similarly displaced, and its angle especially will project. As shown in Fig. 195, the rib in that case commonly

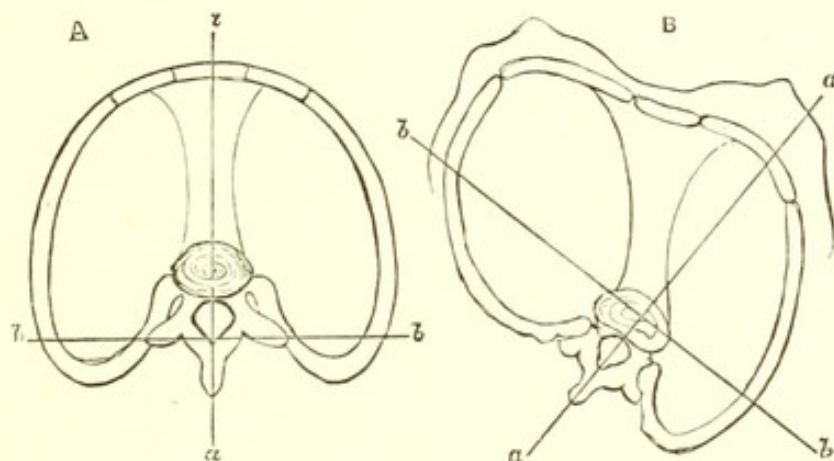


Fig. 195.—Diagrammatic section of normal chest and that with rotatory curvature. In the latter it is seen that the body of the vertebra is rotated into the right side of the chest, the rib on this side turning at an unduly sharp angle. The prominence of the transverse process on this side and of the breast on the opposite side is shown. (ADAMS.)

makes a sharp curve at its angle, so that it is greatly flattened laterally. The figure also shows that, in such a case, the capacity of the corresponding side of the chest is greatly diminished. The body is rotated into that side of the chest and the ribs are flattened towards the vertebræ, these conditions sometimes attaining to such a degree that the bodies of the vertebræ approach the internal surface of the ribs. There is consequently great reduction in the capacity of this side of the chest, the lung being correspondingly compressed and curtailed in its movements. It will be seen also that posteriorly there is prominence of the angles of the ribs on the side corresponding with the convexity of the curve, while anteriorly the breast on the opposite side is prominent.

The term **Spondylolisthesis** has been recently introduced to designate a rare condition in which, as a result of injury or inflammation, the fifth lumbar vertebra is displaced forwards so as to overhang the base of the



sacrum. The vertebra carries with it the whole spinal column above it and projects into the pelvis and narrows it.

**Literature.**—POTT, Remarks on that kind of Palsy which is often found to accompany curvature of the spine, 1779 ; Further remarks, etc., 1782 ; WARD, *Prac. obs. on distortions of spine, etc.*, 1822 ; SHAW, in Holmes' *Syst. of Surg.*, 1883 ; ADAMS, *Lectures on curvature of spine*, 2nd ed., 1882 ; LITTLE, in Holmes' *Syst. of Surg.*, 1883 ; NEUGEBAUER, *Zur Entwicklung d. spondylolisthesis des Beckens*, 1882 ; SWEDLIN, *Arch. f. Gynaek.*, xxii., 1883 ; KRUKENBERG, *ibid.*, xxv., 1884.

#### X.—TUMOURS OF BONE.

The tumours of bone spring from the active tissue which enters into the composition of bone, and this has been shown to be, on the one hand, the medulla, forming not only the marrow of the long bones, but occupying also the spaces of the spongy bone and the various canals of the dense bone ; and, on the other hand, the subperiosteal layer, which really constitutes the same tissue as the medulla. The tumours of bone may be distinguished according as they arise from the medulla, in which case they may be called **Central** or **Myelogenous**, or from the subperiosteal layer when they are designated **Peripheral** or **Subperiosteal**. Originating in these tissues, they frequently present ossifying characters.

The primary tumours of bone belong to the connective tissue series, but from the near connection of some of the bones (especially the jaws) with epithelial structures, we may have tumours composed of epithelium, such as cysts and cancers growing as primary tumours. Of the typical tumours the commonest are the Osteoma, Chondroma, and Fibroma, while the Myxoma, Lipoma, and Angioma constitute rare forms. Amongst the atypical, Sarcoma in various forms is somewhat common as a primary tumour, while Carcinoma is frequent as a secondary growth. Cysts are frequent in the bones of the jaws.

**Osteoma or Osseous tumour.**—The osseous tumours in connection with bone have been distinguished as exostoses when they grow on the surface, and enostoses when they are central. The former are much the commoner. According to structure they are designated ivory exostoses when they are composed of dense bone, and spongy exostoses when of cancellated bone. In some of the exostoses the bone is formed from membrane, while in others it is from cartilage, so that a layer of cartilage covers the surface of the tumour so long as it is growing. This form is sometimes called the **Cartilaginous exostosis**.

The remarks made in regard to the exostoses at p. 224 need not be repeated here. It may be added that **Multiple exostoses** are of occasional occurrence. In the case from which Fig. 196 is taken (Virchow)



the patient, a boy ten years of age, had suffered, during the course of three years, from repeated attacks of rheumatism affecting the joints and muscles. The result was the formation of sixty-five exostoses on various bones of the body. Exostoses also occur not infrequently at the insertion of tendons, growing into the latter and sometimes even separate from the bones. These are connected in their origin with the contraction of powerful muscles, occurring chiefly where such muscles are inserted, and sometimes induced by specially violent exercise of the muscles.

**Fibroma.**—These tumours are mostly peripheral, but they have been observed also centrally in the lower jaw, vertebræ, and ends of the long bones. The peripheral ones mostly occur in the bones of the face and cranium. They sometimes grow into the nares, forming nasopharyngeal polypi. In structure they consist of an intricate meshwork of fibrous tissue. They do not form limited tumours, but grow out from the periosteum over a considerable area, and are firmly adherent to the bone.

**Chondroma.**—This occurs both as a central and as a peripheral tumour. The characters of the cartilage vary somewhat in different cases.

The central chondromata originate in the medulla, especially in the small bones of the hands and feet, and they are often multiple. They may grow till they distend and even rupture the bony shell (see p. 233, Fig. 65).

The peripheral chondromata occur on the long bones, the bones of the trunk and those of the head. They are generally nodulated.

The name **Osteoid chondroma** has been given by Virchow to a form of tumour which merits a more special description. The tumour is composed essentially of tissue such as we find in the deep layer of the periosteum of a growing bone, or in callus, its main constituents being osteoblasts. The cells are smaller than cartilage cells and mostly round; they do not possess a capsule. There is a dense intercellular substance which has a somewhat fibrous appearance. In the midst of the tissue bits of true cartilage may be found. Being formed of tissue analogous to that which is preliminary to ossification it frequently undergoes calcification and even conversion into true bone. Such tumours form, mostly, under the periosteum of the long bones and their seat of election is the lower end of the femur or the upper end of the tibia, originating probably in the layer of osteoblasts there, and they may grow to large dimensions. They thus form club-shaped expansions of the long bones (Fig. 197). On section the tissue is found to be dense, and it becomes osseous or calcareous on passing



Fig. 196.



Fig. 197.

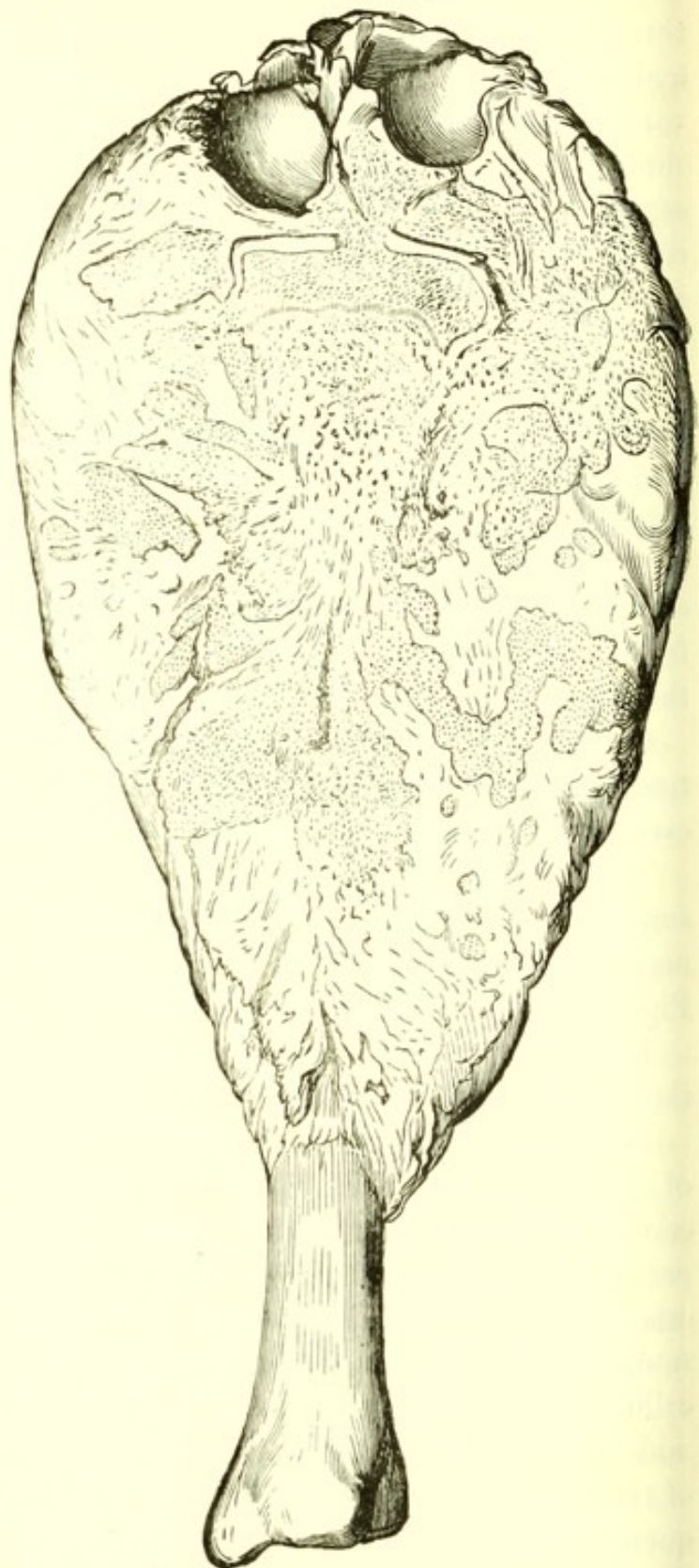


Fig. 196.—Multiple exostoses of femur, tibia, and fibula. These are grouped towards the ends of the diaphyses. At its lower end the fibula is flattened by the exostoses growing out from the tibia. From a boy 10 years of age. One third the natural size. (VIRCHOW.)

Fig. 197.—Osteoid chondroma of the tibia, divided obliquely. At the upper extremity the tumour has surrounded the cartilages of the knee, and new-formed cartilages have been produced. Internally the bone is condensed by the newformation of bony masses. From a boy 13 years of age. (VIRCHOW.)



deeply, where it is incorporated with the bone. The medullary cavity of the bone is often filled with new-formed bone. This form of tumour sometimes presents a tendency to become sarcomatous, and even without that it may recur after removal.

**Myxoma.**—This form of tumour is very rare, but cases have been observed of both central and peripheral origin. A pure myxoma is rare, the tissue being usually mixed with cartilage or other tissue. The growing tumour, whether central or peripheral, causes atrophy of the bone. The commonest seat is the jaw-bones.

**Lipoma** is still more rare. Cases have been described in the upper jaw and tibia.

**Angioma.**—Tumours composed of blood-vessels are very rare in the bones, but some cases are recorded in which cavities filled with blood have communicated with vessels. These cavities have been chiefly at the ends of long bones. The cavities are variously regarded. Some consider them true vascular tumours, others believe they are produced by rupture of arteries, and are **False aneurysms**, while others assert that they originate from sarcomas whose vessels have enlarged. The name **Hæmatoma** is also applied in the view that they originate in hæmorrhage.

Besides these tumours, others may be so highly vascular as to be pulsatile during life. This applies especially to the soft sarcomas.

**Cysts and Cystoma.**—Cysts occasionally arise in bone by softening of the tissue of other tumours or of the bone-marrow itself. Fig. 198 represents a case in which the head of the fibula was distended by a unilocular cyst which contained serous fluid\* and was lined by a vascular membrane.

**Cysts of the jaws** merit special attention. These originate for the most part near the alveolar processes, and are probably related to the teeth. Some of them are single, **Unilocular cysts**, and have their origin in teeth which

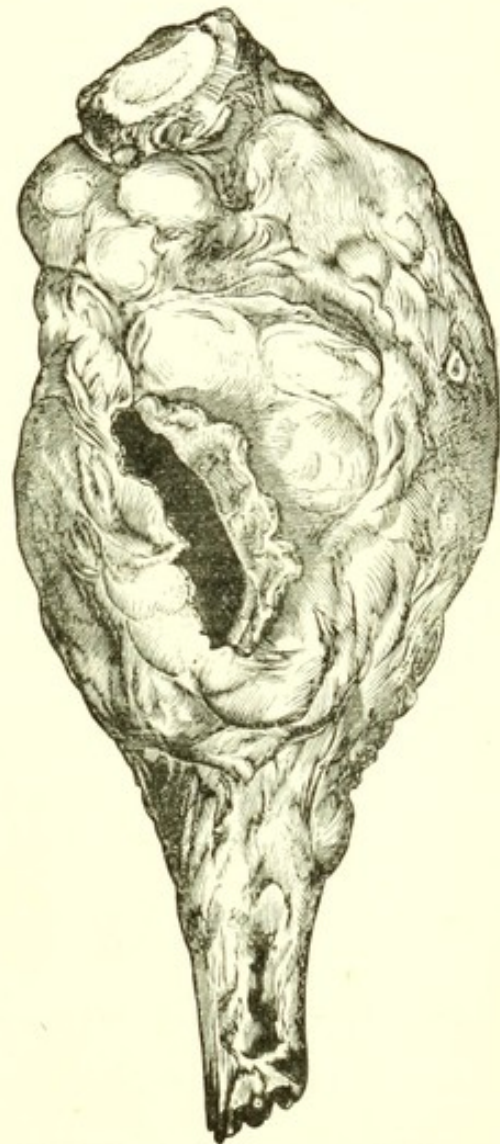


Fig. 198.—Cyst of upper end of fibula. The upper part of the bone is expanded into a cyst which contained a reddish serous fluid.



have not undergone the usual eruption. A tooth or pieces of hard substance (dentine) are usually present in the wall of the cyst. Others are **Multilocular** and develop out of a gland-like tissue, the cells of which undergo colloid degeneration, and in this way form cysts. In its original structure the tumour might be called an adenoma, and a considerable portion of it is sometimes composed of solid glandular or epithelial tissue. According to Eve the tumour arises by penetration downwards of the epithelium of the gums, and he relates this to the normal penetration of this epithelium in the foetus in order to form the enamel organ. These multilocular cysts are mostly innocent, but sometimes the epithelium is less regularly arranged, and they approach in structure and tendencies to the cancers. Both forms of cysts originate in the interior of the jaws and distend them; those in the upper jaw have a special tendency to pass into the antrum which they may fill out.

**Sarcomas.**—These form the most important group of tumours of bone. They are divisible into central and subperiosteal forms.

**The Central or Myelogenous sarcoma** occurs in the lower jaw and in the cancellated tissue of the long bones, especially the femur, tibia, and humerus. They present a somewhat varied structure. Many of them are giant-celled or myeloid sarcomas. Some are composed of large round cells, which may resemble epithelial cells; some are spindle-celled. There are also cases in which large cells exist in nests, forming an alveolar sarcoma, which resembles cancer.

The central sarcomas, originating in the bone-marrow or in the substance of the bone, destroy in their growth the bone-tissue. In this way they frequently work their way to the surface, destroying the continuity of the bone. **Spontaneous fracture** is the result.

In some cases the sarcomatous tissue almost entirely replaces the proper bone, the condition suggesting a newformation in the entire medulla. This is strikingly the case in a preparation in the Museum of the Western Infirmary. Here the entire humerus is replaced by a bulky tumour consisting of round and spindle-shaped cells, no trace of the bone persisting except the condyles. The arm was amputated, but a precisely similar tumour developed in the other humerus, which was also amputated.

**The Subperiosteal sarcomas** are chiefly round-celled or spindle-celled, but in the case of the jaws they may be giant-celled. The tumour may enclose a considerable portion of the bone, especially in the case of the long bones, and may present a somewhat radiating arrangement from the shaft. The tissue is very liable to undergo ossification. In some the ossification is so striking a feature that the whole tumour is represented by a mass of dense bone, on the surface of



which the sarcomatous tissue is present as a comparatively thin layer. To this form the term **Osteoid sarcoma** may be applied. The cells on the surface of this tumour are variously shaped, and the bone seems to arise directly from these cells, which act as osteoblasts.

The sarcomas of bone show very varying degrees of **Malignancy**. They grow locally at the expense of the bone, and not infrequently encroach on neighbouring structures, sometimes penetrating into the joints. Many of them show merely this local malignancy, but others extend by metastasis to internal parts. Even the highly ossifying forms may lead to secondary tumours in the lungs, the metastatic tumours showing a similar ossifying tendency.

**Carcinoma.**—A primary cancer can scarcely originate in bone, where there are no epithelial elements. In the jaws, however, we meet with what are practically primary cancers. The near connection of the jaw-bones with the gums and with the teeth, whose enamel organ is an epithelial structure, explains the origin of such tumours. Cysts of the antrum have been already referred to as frequently originating in adenoid tissue, and this tissue may take on a cancerous character. In both jaws, also, cancer, originating at the alveolar surface, may grow into and involve the bone.

**Secondary cancer** is very frequent in the bones. Cancer, when it becomes generalized, is very liable to become implanted in the cancellated tissue, especially of the vertebræ, ribs, and ends of the long bones. It there grows as a central tumour, distending and destroying the bone, and sometimes leading to spontaneous fractures. In most cases the tumours are multiple.

**Secondary thyroid adenoma** is a rare form of tumour in bone. A few cases have been observed in which, with enlargement of the thyroid, secondary tumours composed of glandular tissue have developed, chiefly in the bones of the skull and vertebræ. In a case observed by the author the tumours occurred in the diploë of the skull, and in growing destroyed the bone. One of the tumours in the occipital region caused a gap in the skull an inch and three quarters in diameter. This was occupied by the tumour which projected inwards and outwards. It formed during life a bulky **Pulsatile tumour**.

**Parasites** are very rare in bone.\* The **Cysticercus cellulosæ** has been observed, but the **Echinococcus** more frequently. The hydatid cyst has mostly been met with in the bones of the pelvis or in the tibia. After growing and distending the bone it may burst through it and develop further in the soft parts around.

**Literature.**—WEBER, *Die Exostosen und Enchondrome*, 1856; VIRCHOW, *Geschwülste* ii.; LÜCKE, in Pitha and Billroth's *Handb.*, 1869; CORNIL et RANVIER, *Man.*



d'hist. path., i., 1881; EVE, Cystic tumours of jaws, Brit. Med. Jour., 1883, i.; HEATH, Injuries and Dis. of Jaws, 3rd ed., 1884; BUTLIN, Oper. surg. of malig. disease, 1887; JONES, Dis. of bones, 1887; COATS, Path. trans., xxxviii., 1887.

## B.—THE JOINTS.

### I.—DISLOCATIONS AND MISPLACEMENTS.

1. **Congenital dislocations.**—Children are sometimes born with the joints in faulty positions, some of these being really traumatic and others of more obscure origin. The traumatic cases arise for the most part during parturition, either from the natural forces engaged in delivery, or by dragging on the part of attendants. The **Hip-joint** is not infrequently dislocated in this way, and usually the displacement is on both sides at once. The head of the bone is usually displaced upwards so as to rest on the dorsum of the ilium above and behind the acetabulum. Through time the head makes for itself a new joint, and the acetabulum fills up. Congenital dislocations of other joints are exceedingly rare.

2. **Talipes or Club-foot and Club-hand.**—These names are applied to distortions of the feet, or more rarely of the hands; the bones assume certain abnormal positions in which they are retained by the contraction of muscles.

**Causation.**—A certain proportion are **Congenital**, the child being born with one, or more commonly both feet, turned rigidly in, so as to form a Talipes varus. The muscles keep the foot in this position. According to Eschricht and Berg, the foot in early foetal life is inverted, and at birth normally retains a degree of inversion, the soles being turned in. An exaggeration of this, or a retention of the early foetal condition, constitutes a congenital club-foot.

The **Non-congenital** forms are, in most cases, due to infantile paralysis, a disease in which some muscles are paralysed, while others are not. In some cases the deformity is due to the rigid contraction of the unparalysed muscles, a condition to which the name **Paralytic contracture** is often given. In many cases, however, the faulty position is largely due to the fact that the muscles are not able to keep the foot in its proper position, so that it tends to fall into the attitude it would assume if there were no muscles. These positions are also variously modified by the use made of the foot in walking, the parts being brought against the ground in the position most suitable to stability in the maimed condition of the limb. In a similar way club-foot may occur in pseudo-hypertrophic paralysis.

Besides paralysis, it seems that spasm may lead to talipes, although



the explanation of the local spasm is often very obscure. It is undoubted that children sometimes acquire a talipes after a convulsive attack.

**Forms of Talipes.**—These do not call for detailed description here. There are four principal forms, some of which may be combined. In **Talipes varus** the toes are inverted, and the inner margin of the foot is raised upwards; the *tibiales anticus* and *posticus* are the muscles which are chiefly contracted. In **Talipes valgus**, which is one of the rarer forms, the foot is turned outwards, the outer border is raised, and there is usually also some elevation of the heel; the *peronei*, *extensor longus digitorum* and *gastrocnemius* are the muscles chiefly engaged. In **Talipes equinus** the heel is raised and the foot extended, so that the person walks on the distal extremities of the metatarsal bones; the *gastrocnemius* is the muscle contracted. In **Talipes calcaneus** the heel is depressed and the foot flexed at the ankle, this deformity being the reverse of the preceding one; the contracted muscles are chiefly the *tibialis posticus*, the *peronei* and the *extensors*.

By long retention of the fixed position the **Bones** become variously **modified in shape**, undergoing atrophy where exposed to prolonged pressure. Where the cartilage is no longer used in the movements of the joints, it also atrophies. The bones frequently acquire adhesions in their new positions, and the ligaments, by elongating or shortening, accommodate themselves to the new position. New ligamentous attachments are sometimes formed, and the bones may become joined together by ankylosis.

**3. Traumatic dislocations.**—We have here to do with cases in which the bone is pushed out of its place by some external force acting on it.

The bones are kept in their places mainly by the ligaments of the joints, but no inconsiderable aid in this regard is given by the muscles which act on the bones. As a general rule when external pressure is exercised on a bone at a joint, the muscles are so braced as to enable the bone to retain its place in spite of the external force. But if a force is exercised on a bone unexpectedly or when the muscles are generally relaxed as by alcoholic stupor, then it may be displaced, although the force under ordinary circumstances would be insufficient to produce this effect.

When the bone is dislocated, the same contraction of the muscles which normally aids in preventing dislocation, generally offers serious obstacles to the return of the bone to its normal place.

A bone may even be displaced by the action of the muscles themselves, where a certain group acts very vigorously while their antagonists are relaxed. There are indeed persons who can produce **Dislocation voluntarily** of almost all the more moveable joints, and that by mere muscular effort. In order to this we must suppose a



certain laxness of the ligaments, but there is also a power acquired by education of strongly contracting certain muscles, while others which usually contract along with them are relaxed. We know that for the most part the muscles in their contractions are co-ordinated, and most people are unable to contract individual muscles apart from their co-ordinates, but there are exceptional persons who possess this power, some in a limited degree and others very remarkably. Many persons, for instance, cannot shut one eye without shutting the other, and most persons when they shut one require to make an active effort at opening the other, in order to prevent it shutting too. But there are persons who can close the eyelids of one eye as easily as they can close the fingers of one hand.

In traumatic dislocation there is usually tearing of the ligaments to some extent, and in the case of some joints much laceration is necessary before dislocation can occur.

If restoration does not occur soon then the bone acquires adhesions in its abnormal situation, the adhesions being the result of chronic inflammation set up by the irritation of the bone. The displaced bone generally comes to press with its head against a neighbouring part of the bone with which it formerly articulated, and the adhesions attach it to the periosteum in its new position. Friction by degrees wears down to some extent the opposed piece of bone, and as new bone is produced around by the irritation there may be a kind of hollow joint formed. By the wearing of the bone the cancellated tissue would be exposed, were it not that dense bone is produced on the surface so as to cover in the spongy tissue. A smooth hollow surface may thus be produced, and a tolerably perfect joint, although cartilage is not formed to cover it, but only a layer of smooth polished bone. The head of the displaced bone also loses its cartilage, and may even get worn away considerably. If the bone remains permanently displaced the old hollow of the joint becomes filled up, bone growing when the friction of the opposing bone is no longer exercised.

**4. Spontaneous dislocations.**—This name is applied to dislocations which occur without any considerable violence, and they usually imply a previously diseased state of the joints. The disease is generally inflammation or tuberculosis, which soften the ligaments and alter the form of the joints.

**Literature.**—ADAMS, *On Club-foot*, 1866; ESCHRICHT, *Deutsch. Klinik*, 1851; MICHAUD, *Arch. d. phys.*, iii., 1870; HOLL, *Langenbeck's Arch.*, xxv., 1880.

## II.—ANCHYLOSIS.

By this name is meant fixation of a joint by union of the opposing bones by means of firm adhesions. The expression "false ankylosis" is sometimes used to designate the condition in which the joint is fixed, not



by adhesion between the bones, but by rigidity and shortening of the surrounding soft parts.

For the most part ankylosis is the result of inflammations of joints, where the cartilage has been destroyed and healing has subsequently occurred. In the process of healing the inflammatory tissue on the opposing surfaces develops into connective tissue, and as the two surfaces have to a considerable extent coalesced, fibrous tissue unites them permanently. In this fibrous bond of union there are often bony plates, and occasionally the union is effected by bone itself. In the latter case the term **Synostosis** is applied. This, however, is a very unusual occurrence, as even slight movement of the joint is sufficient to prevent the formation of bone. The fixation of the joint, however, is often so firm as to resemble an actual coalescence of the bones.

The term **Spondylitis deformans** is given to a condition in which the vertebræ are ankylosed together. There is a synostosis of the arches and articular processes, while the heads of the ribs are ankylosed to the spine. The condition is a gradually advancing one, and the back becomes rigid.

### III.—INFLAMMATION OF JOINTS—ARTHRITIS.

In most cases of arthritis the inflammation affects, more or less, all the structures which enter into the construction of the joint. The irritant is usually present in the joint itself, and is distributed over it, with the synovial fluid, by the movements of the joint. We may expect, therefore, that the synovial membrane and the cartilage, as they cover the surface, will in most cases be primarily affected. The cartilage, being non-vascular, is less liable to inflammatory changes than the synovial membrane, which in most cases is primarily and chiefly engaged, especially in acute inflammations. But in chronic cases the irritation extends beyond these structures to the ends of the bones, to the neighbouring periosteum, and even to the ligaments.

1. **Simple acute arthritis.**—This condition is produced most directly by the opening of joints and the occurrence of septic decomposition in their fluids. But it also occurs sometimes by exposure to cold, from injuries, and from the extension of phlegmonous inflammations from surrounding parts.

In its anatomical details, the inflammation resembles closely that of the pleura and pericardium. The synovial membrane and cartilages are covered with fibrinous exudation, and the cavity contains serous fluid in which flakes of fibrine are visible. This fluid is sometimes very considerable and distends the joint. In this stage the disease may resolve and the joint return to its normal condition. On the other hand, especially



if the joint has been laid open and exposed to decomposition, the inflammation may go on to suppuration. The synovial membrane becomes swollen and dull, being infiltrated with inflammatory cells, and gradually converted into granulation tissue. If the inflammation is very acute, we may have suppuration by the mere exudation of leucocytes without much change in the synovial membrane. When the disease has gone on to the formation of granulation tissue it is now in a subacute condition which is apt to be prolonged. The inflammation also extends beyond the synovial membrane. In the cartilage the cells undergo active proliferation, and the matrix breaks down; thus softening occurs, and ulcers form. There is inflammation of the bone, a rarefying osteitis. The ligaments also take part in the inflammation, they are softened and, with the synovial membrane, take part in the formation of granulation tissue. The periosteum is inflamed and new bone is formed, so that irregular projections occur near the ends of the bones. With all this there is, of course, usually an abundant purulent discharge from the joint, which may weaken the patient and prove fatal, perhaps with amyloid disease.

If the inflammation subsides, the various masses of granulation tissue develop into connective tissue, and, by the contraction of this, great rigidity of the joint may be produced. The granulation tissue lining the joint does also to a great extent coalesce, and the result is a partial or complete obliteration of the joint. The bones thus become finally united by fibrous or osseous adhesions, and a permanent ankylosis is the result.

2. **Pyæmic arthritis.**—In this disease septic microbes are deposited in the joint and spread over the surface by the synovial fluid. The result is an acute inflammation with fibrinous exudation, but generally going rapidly on to suppuration. It is remarkable that, when the joint is full of pus, there is sometimes very little structural change in the synovial membrane, the whole condition being almost confined to the blood-vessels from which an excessive exudation has occurred. The inflammation usually affects several joints simultaneously.

It is to be added that occasionally in **Dysentery, Diphtheria, Scarlet fever, Erysipelas**, etc., a similar acute arthritis occurs. In these diseases there is a breach of a cutaneous or mucous surface and septic or other microbes may find entrance. The affection of the joints is thus similar in its origin to Ulcerative endocarditis, which sometimes complicates these affections.

3. **Gonorrhœal arthritis.**—An acute arthritis sometimes develops in gonorrhœa, but the connection between the two diseases is disputed by some. The inflammation is usually slight, like that in Acute rheumatic



arthritis, or it may, in exceptional cases, assume a suppurative character like that in pyæmia.

4. **Acute rheumatic arthritis.**—Like pyæmic arthritis this is due to an irritant which is present primarily in the blood and affects the structures of the joints like other connective tissue structures. The result is an acute inflammation accompanied by serous and sometimes by fibrinous exudation into the joint. The synovial membrane is injected and swollen, but in most cases the inflammation passes off without leaving any permanent change.

In exceptional cases the inflammation lingers in one or more joints, just as it does in the heart when the valvular structures are affected by the same rheumatic poison. In that case the chronic inflammation produces thickening and rigidity of the ligaments, sometimes with fibrous union of the bones.

5. **Gouty arthritis.**—In this disease uric acid, in the form of urate of sodium, is deposited in the tissues of the joints. It is first deposited in the cartilage, and, according to Charcot, always, to begin with, at the middle of the articulating surface, that being the point furthest removed from the blood-vessels. The urate is sometimes in the form of stellate crystals (Fig. 199), the cartilage cells forming the middle points of the bunches of crystals. But the salt is also deposited in the matrix and often in the form of irregular needles. It is also frequently deposited in the synovial membrane, and in the bones, ligaments, and soft parts around the joints. Sometimes the salts are deposited even under the skin, forming visible projections (the so-called tophi or chalkstones).

According to Moore the deposition of urates is always preceded by degenerative changes in the cartilage. This author also associates gout very intimately with chronic interstitial nephritis, pulmonary emphysema, and chronic inflammation of the aortic valves.

The deposition seems to lead to an acute irritation of the structures of the joint, chiefly an intense hyperæmia, often with serous exudation. It never goes on to suppuration, however. Complete recovery usually takes place, but sometimes there remains a chronic inflammation, leading to stiffness and deformity of the joint.



Fig. 199.—Cartilage of joint in gout, with crystals of urate of sodium. The salt is in stellate crystals which are nearly continuous at upper part of figure, which corresponds to surface of joint.  $\times 200$ . (CORNIL and RANVIER.)



6. **Chronic rheumatic arthritis.**—As a rule this disease affects many joints. As the changes are virtually the same as in **Arthritis deformans**, and the two conditions run into one another, we may describe them together. In arthritis deformans the disease is partial, being confined perhaps to a single joint, and the lesions attain a much greater development than in general articular rheumatism. It is also much more a disease of old people, and is met with chiefly among the poor. In both these diseases the nature of the irritant is entirely obscure.

The inflammatory phenomena appear first in the synovial membrane and the cartilages. The synovial fringes enlarge by a slow process of inflammation, and the villous projections increase in number and become more prominent. Not uncommonly pieces of cartilage develop in the fringes, originating in the cartilage cells which exist normally there, and these pieces of cartilage, being usually pedunculated, act very much like free bodies in the joint and may become loose. This cartilage also sometimes undergoes ossification in whole or in part. At the same time there is an increase of the synovial fluid, but without any formation of fibrine or pus.

The cartilage cells undergo proliferation and the matrix presents a peculiar fibrillation, so that the cartilage assumes a soft velvety condition, and readily undergoes destruction from the friction of the opposing surfaces. It is stated by Rindfleisch that the fibrillæ of the matrix undergo mucous degeneration, and that mucus may be found in the synovial fluid.

The bone beneath the cartilage also undergoes considerable alteration, especially when the cartilage gets worn away. To prevent the cancellated tissue from being opened, there is formed, as the cartilage is destroyed, a layer of dense bone, which is smooth and polished, and takes the place of the cartilage. This forms a much less efficient covering, and the ends of the bones work roughly against each other. This layer of bone may also be worn down, and so considerable atrophy of the head of the bone may result, a layer of dense smooth bone being always reproduced.

At the borders of the articular surface there is often considerable newformation of bone, the result of the inflammation where the parts are not exposed to friction. It appears from the observations of Ranvier that the bone here develops largely from cartilage. The synovial membrane, at the edge of the joint, covers in some of the marginal portions of the cartilage, and being thus protected, the cells may proliferate so as to lead to newformation of cartilage, which may ossify. The periosteum may also produce new bone, and even the ligaments attached to the periosteum may be the seat of bony processes.



With all this there is considerable thickening of the ligaments by inflammatory newformation of connective tissue, and often fibrous union between opposing parts of the joints. Indeed, if the joints are kept at rest, there may be a complete union of the parts around the joints opposite each other, leading to ankylosis.

Without ankylosis there is stiffness of the joints, whose movements are greatly curtailed. There is frequently great deformity from the thickening of the tissues and newformation of bone.

An extreme degree of arthritis deformans is sometimes seen in the hip joint, where it gets the special name of **Morbus coxæ senilis**. Here the wearing down of the head of the bone is sometimes very extreme, so that ultimately the articulating surface may lie between the trochanters. As newformation of bone occurs simultaneously at the borders of the articular surface, a kind of artificial head is produced, and the appearance is presented as if the neck were atrophied and the head displaced. In like manner an apparent widening of the acetabulum may occur. The original articular surface is worn away, but by the formation of new bone under the periosteum around, a wall is formed, giving the appearance of the borders of a widened acetabulum.

**Charcot's disease.**—This name is applied to conditions of the joints arising in consequence of diseases of the spinal cord, especially locomotor ataxia. The lesions consist in an atrophy of the articular ends of the bones including the cartilages. There is, in consequence of the exposure of the bone, a wearing down of the bones, without the newformation of bone such as appears in chronic rheumatic arthritis. (See under Locomotor Ataxia.)

#### IV.—SYPHILIS AND TUBERCULOSIS OF THE JOINTS.

**Syphilis** does not frequently attack the joints, but rheumatic attacks in syphilitic persons may have some relation to the specific virus. According to Lancereaux, there may be, in the secondary stage, an inflammation like that of acute or subacute rheumatic arthritis, and, in the tertiary stage, manifestations like those of chronic arthritis.

**Tubercular arthritis.**—This disease is also called *strumous synovitis*, *gelatinous degeneration* of the joint, *tumor albus*, *fungous caries*, etc. The tubercular virus is the essential factor in the **Causation** of the disease. In many cases it reaches the joint after having attacked the bone, but the proportion of cases in which it does so as compared with those in which it comes directly is matter of dispute. Tuberculosis of bone will extend to the joints much more readily in the case of some bones than others. As the head and neck of the femur are exposed in the hip-



joint in immediate contact with the synovial membrane, extension very readily occurs here.

The disease is mostly one of children, a fact which favours the view that it generally takes origin in the bones. It is chiefly weakly children who are attacked.

The disease begins usually in the **Synovial membrane**, and consists in a formation of tubercles and a chronic inflammation with great production of granulation tissue. The soft, pulpy granulation tissue gives the synovial membrane a gelatinous appearance, from which one of the names of the disease is taken. There is also a slow enlargement of the joint from the increased bulk of the synovial membrane.

The **Ends of the bones** constituting the joint are affected simultaneously or soon after, and here the characters presented are those of tuberculosis of bone. The medullary spaces become filled with granulation tissue and enlarged by destruction of the bony lamellæ. In this granulation tissue there are also tubercles. There is thus, as it were, a pad of granulations under the articular cartilage.

The cartilage also at the sides is partly encroached on and overlapped by the altered synovial membrane which advances over it. In this way it is partly enclosed between two layers of granulation tissue, and it gradually becomes eaten into. The granulations, chiefly those of the medulla, extend into the cartilage, and their encroachment is assisted by proliferation of the cartilage corpuscles, which enlarge and cause softening of the matrix around.

By the absorption of the cartilage the whole joint may be converted into a cavity lined with granulation tissue, and the ligaments also are frequently transformed in a similar way.

Generally suppuration results, and the joint becomes filled with pus. Abscesses also not uncommonly form around the joints. Through time the pus generally finds its way outwards, and is discharged by an aperture in the skin. A fistulous canal is the result, forming a communication between the cavity of the joint and the surface, and this canal is also lined with exuberant granulations, which pout out at the opening in the skin. Among the granulations here, as elsewhere, tubercles are found.

The rubbing of the two ends of the bones, which are now covered by granulations, leads to an ulcerative destruction of those soft structures. The inflammation extends more deeply in the bone as the superficial parts are ulcerated, and so we have progressive caries. For some distance beneath the surface the medullary spaces are filled with granulations and the bony trabeculae thinned. It will be understood from this how the disease is apt to recur unless the whole carious por-



tion be removed, for tubercles are present in the granulation tissue filling the medullary spaces, and unless they be removed a fresh extension may occur.

In an early period of the disease, before suppuration has occurred, there may be recovery; but after the occurrence of suppuration, there is seldom a spontaneous restoration, which at best is a slow process. If recovery takes place the granulating surfaces unite more or less, and the joint being partly or completely obliterated, a fibrous union may come about, leading, it may be, to ankylosis.

The author has met with a case in which an early and pure tuberculosis of the synovial membrane presented peculiar characters. There was very great thickening, so that the synovial membrane was converted into a bulky soft grey tissue which overlapped the cartilages, and was so prominent that when the joint was opened for the purpose of excision, the idea of a tumour was suggested. In this case there were large numbers of the most typical tubercles, many consisting almost entirely of giant-cells and epithelioid cells. It is from one of these that Fig. 53, p. 201, is taken.

**Literature.**—LANCEREAUX, *Traité de la syph.*, 1874; KÖNIG, *Tuberculose der Knochen und Gelenke*, 1884; CROFT, *Path. trans.*, xxxii., 1881.

#### V.—LOOSE BODIES IN JOINTS.

These occur most frequently in the knee joints, but also in the hip, shoulder, maxillary, and other joints. They consist generally of more or less rounded pieces of tissue, and we may have fibrous tissue, bone, cartilage, and adipose tissue entering into their composition. They are nourished by the juices of the joint and may even grow in their abnormal position.

The loose bodies have various origins, but usually arise by separation of pieces of tissue which may either be parts of the normal cartilage or bone, broken off by violence, or else parts of abnormally prominent structures which have in the movements of the joints been torn off. Thus the synovial fringes may enlarge by excessive growth of adipose tissue or even of cartilage in them, or by inflammatory newformation. Again, in chronic rheumatic arthritis, the prominent bony excrescences are liable to be broken off.



## SECTION III.

## DISEASES OF THE NERVOUS SYSTEM.

- INTRODUCTION. *The plan of the nervous system.* A.—THE PERIPHERAL NERVES. *Anatomical introduction.* 1. *Effects of injury and division of nerve stems.* 2. *Hyperæmia and acute inflammation; Neuritis.* 3. *Infective newformations and tumours.*
- B.—THE SPINAL CORD AND MEDULLA. *Anatomical introduction.* I. SECONDARY DEGENERATIONS. *Grey degeneration or sclerosis; descending and ascending; degeneration after amputations.* II. INFLAMMATIONS. A. TRANSVERSE MYELITIS, 1. *Acute; acute softening,* 2 *Chronic; chronic compression.* B. SYSTEMATIC MYELITIS. *General causation.* 1. *Sclerosis of posterior columns, lesions, and symptoms; Hereditary ataxia.* 2. *Lateral sclerosis.* 3. *Acute ascending paralysis.* 4. *Acute anterior poliomyelitis.* 5. *Chronic anterior poliomyelitis.* 6. *Progressive muscular atrophy.* 7. *Bulbar paralysis.* 8. *Pseudohypertrophic paralysis.* III. TUMOURS.

## INTRODUCTION.—PLAN OF THE NERVOUS SYSTEM.

THE simplest idea of a nervous system is that of a central ganglion cell with an afferent or centripetal fibre and an efferent or centrifugal one. An approach to this simplest form of nervous system is afforded us in the case of the heart. We have here in the substance of the organ certain groups of ganglion cells, which possess on the one hand centripetal fibres coming chiefly from the endocardium, and on the other hand centrifugal fibres passing to the muscular fibres of the heart. It is to be presumed that impressions conveyed from the endocardium induce the development of impulses which are conveyed by the centrifugal fibres to the muscle and bring about its contraction.

But these intrinsic ganglia of the heart, although forming with their connections a complete nervous system, are not entirely isolated and independent. They are under the command of higher centres which control their action and through them affect the contractions of the heart. From these higher centres fibres reach the heart by two paths, by the vagus and by the sympathetic, and by means of these fibres the action of the intrinsic ganglia is restrained or stimulated.

Taking a general survey of the nervous system, we find that, among



the innumerable centres, there are grades or orders to be recognized, the lower or simpler being under the control of the higher and more complex. Leaving aside the peripheral centres and the sympathetic system, we may fitly illustrate this in the case of the cerebro-spinal axis.

In the **Spinal cord** there are, chiefly in the anterior cornua, groups of ganglion cells which form distinct individual centres. Many of these appear to be of the simplest kind, representing, as it were, single muscles or limited groups of muscles. The stimulation of such simple centres would produce no properly co-ordinated movements, but simply the contraction of a muscle or muscles. But in the cord itself there are centres of a higher order than this, representing, not single muscles or very limited groups, but more considerable groups of associated muscles, so that movements of some complexity are brought about by their stimulation. The centres of lower order are under the control of the higher, and it is to be presumed that the higher, in bringing about movements, do not act directly on the muscles, but stimulate in the first instance the lower centres, which then act directly on the muscles. Even the higher centres in the cord are, as compared with those in the brain, of a very low order, and are only capable of effecting such simple actions as the extension of the toes, the drawing up of the leg, etc.

The **Medulla oblongata** may be regarded as simply an extension upwards of the spinal cord. Its centres are scarcely of a higher order than those of the cord, and the movements which may be effected by it alone are of the simplest character. In it are massed the great centres which have the control of the respiratory movements, and the contraction and dilatation of the blood-vessels. The muscles of the tongue, mouth, pharynx, etc., are represented here as are those of the legs and trunk in the spinal cord.

Passing to the centres next in order above the cord and medulla oblongata we reach the so-called **Middle brain**, including the centres in the pons varolii, the corpora quadrigemina, and, as perhaps of a still higher order, the cerebellum. Many animals can go through very elaborate movements when deprived of all parts above this middle brain. A pigeon can fly, a frog can leap, and a rabbit can run. There is, however, a want of spontaneity in the movements, which present many of the characters of complex reflex or automatic actions. A rabbit will remain quiet till its foot is pinched, and will then set about running. The movements effected by means of the middle brain require the action of the same muscles as those in which the spinal cord alone is concerned, but the combinations are more complex and the grouping of the muscles more intricate. In effecting these more complex move-



ments the higher centres act in the first place on the lower, and, through them, on the muscles, the lower centres in the cord being thus a necessary link in the chain.

In man the middle brain appears to be much less independent than in the lower animals. In many animals, as we have seen, a stimulus coming from the periphery may induce such complex acts as flying, leaping, running, but it is not so in man. If the centres for such acts are situated in the middle brain in man, they are so dependent on the higher centres that when their connection with these is severed they are only able to act very imperfectly. A certain degree of independence is shown in man by the fact that when a person, completely paralysed on one side by the connection being divided between the middle and upper brain, yawns, the paralysed arm will often move in an exaggerated fashion entirely independently of the will. Yawning is an exaggerated inspiration, and in order to elevate the chest the arm is stretched upwards and backwards so as to bring the pectoralis into action on the chest wall. When we have command of ourselves we can control these movements, but when the middle brain is disconnected the paralysed arm may act in an exaggerated fashion.

The **Basal ganglia** of the cerebrum form a series of centres of a very high order. When such animals as the dog and cat are deprived of all centres higher than the corpus striatum they are capable of running about, these movements being, of course, automatic. But in man, and also in monkeys, although the general movements of the body may be regarded as gathered together in these ganglia, they are not sufficient for the more complex acts of locomotion, etc. The movements of the body, although represented in a complex form in these ganglia, are represented higher up in a still more complex form, and at the same time the lower centres are less independent of these higher ones.

In the **Convolutions** of the cerebral hemispheres we have the highest order of centres, and in man the **Motor area** may be taken to form the seat of all the centres which are concerned with the more complex voluntary acts. In the motor convolutions we have the movements of the body as it were written larger, occupying much more space than in the corpus striatum, and more individualized.

In regard to **Sensation**, we are not to look for a succession of centres such as we have in the case of motion. There are peripheral organs of a highly specialized character, which are engaged in the transmission of the various special kinds of sensation. Between these and the highest centres there are virtually no others interposed, the intervening structures being only concerned in conduction, perhaps with arrangements for fortifying the impressions as they are conducted through



greatly elongated paths. Besides the apparatus engaged in sensation, there are afferent fibres which are related to reflex actions, and probably the same fibres to some extent subserve both functions.

In studying the specific diseases of the nervous system it will be necessary to carry these physiological considerations along with us, and in the case of each disease it will be proper to take into account the effect which it will have on the physiological action.

Lesions occurring in nervous structures produce various effects. They may **irritate** the centres either directly or by means of their communicating fibres. If a **Motor centre** be irritated there will be muscular movements, spasm, convulsion. If a **Sensory centre** be irritated there will be subjective sensations as of sight, smell, touch, hearing, taste. If a **Mental centre** be irritated there will be subjective mental phenomena, that is, mental phenomena which are beyond the control of the individual, peculiar thoughts, illusions, etc. On the other hand, lesions may **destroy** centres, in which case we shall have paralysis of motion, or loss of sensation (anæsthesia), or mental degeneracy.

Lesions which are large and palpable are often called **Coarse lesions**, as where a tumour or a clot destroys or irritates, or does both. Coarse lesions are thus distinguished from those finer changes which are matter for microscopic observation. In some cases, indeed, the existence of actual physical changes is matter of inference, the anatomical demonstration of them being not yet furnished.

**Literature.**—An excellent exposition of the construction of the nervous system is given in HERBERT SPENCER'S *Principles of Psychology*, vol. i.; also in many papers by HUGHLINGS JACKSON, whose influence in advancing the pathology of the nervous system has been very great. A systematic study of the physiology of the nervous system in FERRIER'S excellent work, *The functions of the brain*, 2nd ed., 1886. See also, for *Diseases of the nervous system*, ROSS, *A treatise on dis. of nervous system*, 2nd ed., 1883 (which contains frequent references); and GOWERS, *A manual of diseases of the nervous system*, 1886 and 1888.

## A.—THE PERIPHERAL NERVES.

**Anatomical introduction.**—A nerve stem, whether met with embedded in the tissues of an organ or lying free, is composed of one or more bundles of nerve fibres united together by connective tissue. The accompanying figure (Fig. 200) shows the general arrangement of this connective tissue in a stem composed of a single bundle of nerve fibres. There is an external layer of connective tissue, the perineurium (*a*), binding the whole bundle together. But inside the bundle there is connective tissue binding the individual nerve fibres together and forming the endoneurium (*b*), the nuclei of which are prominently seen in the figure. In a nerve stem made up of several bundles these also are bound together by connective tissue, the epineurium or neurilemma.



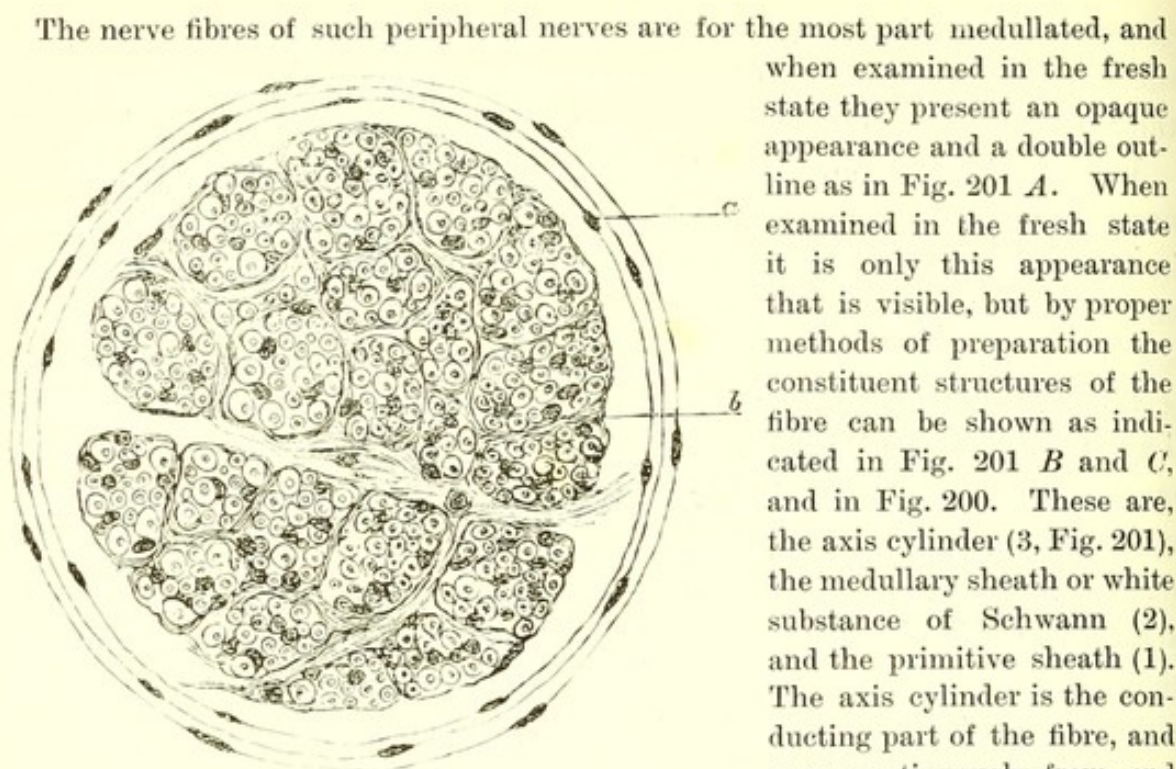


Fig. 200.—Transverse section of a nerve consisting of a single bundle; from a specimen stained and mounted by Clarke's method. *a*, perineurium; *b*, endoneurium. Inside the perineurium is the lymphatic space between it and the nerve bundle. The nerve-fibres are represented by rings with a central dot—the axis cylinder.  $\times 120$ . (KLEIN.)

double contour. This substance is semi-fluid, and when the nerve fibre is broken up, either during life or after death, it is apt to flow out, and so we may have free

when examined in the fresh state they present an opaque appearance and a double outline as in Fig. 201 *A*. When examined in the fresh state it is only this appearance that is visible, but by proper methods of preparation the constituent structures of the fibre can be shown as indicated in Fig. 201 *B* and *C*, and in Fig. 200. These are, the axis cylinder (3, Fig. 201), the medullary sheath or white substance of Schwann (2), and the primitive sheath (1). The axis cylinder is the conducting part of the fibre, and runs continuously from end to end. The medullary sheath is composed of a fatty substance (myeline) and is prone to undergo a kind of coagulation which gives rise to the

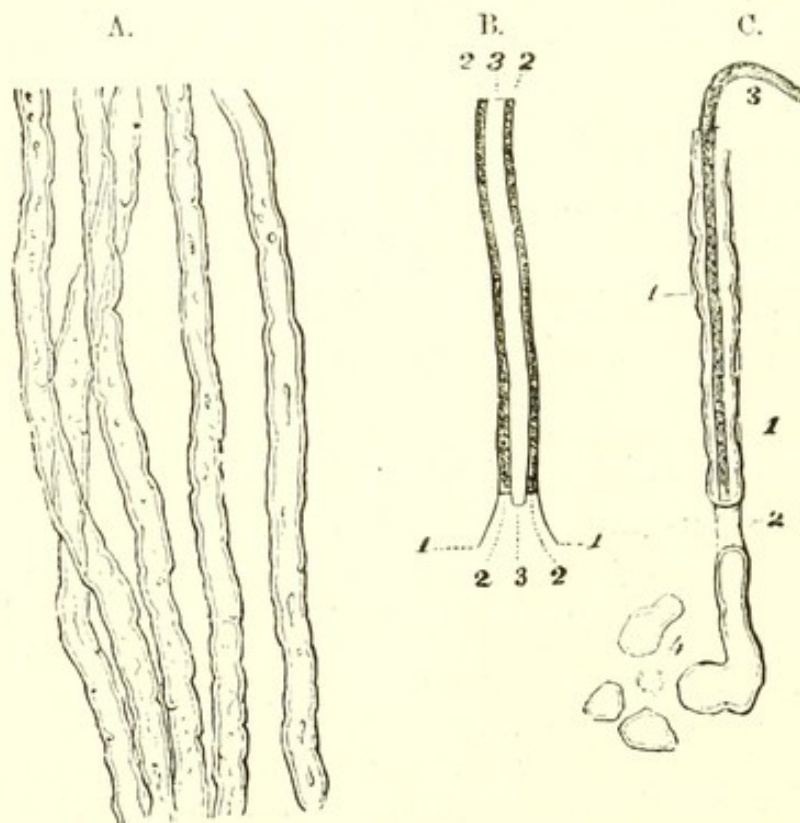


Fig. 201.—Medullated nerve fibres. *A*, the natural appearance; *B* and *C*, diagrams showing the constituent structures as in text. (QUAIN.)



drops of myeline which have a strongly refracting outline (4, Fig. 201). The primitive sheath is a transparent membranous tube which covers the fibre and keeps the medullary sheath together. When transverse sections of a nerve which has been hardened and stained are examined these various constituents appear as in Fig. 200. The axis cylinder is a coloured point in the middle of each fibre. The medullary sheath around this is transparent and colourless. The primitive sheath forms a coloured ring around the fibre.

If a medullated nerve fibre be examined after preparation with osmic acid, it will be seen that, as Ranvier has shown, the medullary sheath is not continuous, but is interrupted at intervals, the axis cylinder and primitive sheath being alone present. These *nodes* divide the nerve fibre into sections therefore, and each section receives a further individuality from the fact that about its middle an oval nucleus is present inside the primitive sheath, between it and the medullary sheath.

Non-medullated or pale nerves have no medullary sheath, and consist essentially of axis cylinders each covered with a primitive sheath in which nuclei occur at intervals. As the white appearance of ordinary nerves depends on the medullary sheath, non-medullated nerves are grey in colour. Most nerves at their peripheral terminations lose the medullary sheath and become pale, but some are so throughout, chiefly the olfactory nerve and the sympathetic.

**1. Injury and division of nerve stems.**—When a nerve is divided there occurs motor and sensory paralysis in the peripheral parts dependent on it. But it frequently happens that conduction is after a time restored, the divided ends having re-united, and this sometimes so rapidly that immediate union has been supposed to occur. Most frequently, however, the restoration of function is tardy, and processes occur in the nerves which have been carefully studied by experimentation on animals. The changes which ensue on the division of a nerve occur mainly in the peripheral portion of it, and they consist in the first place in a **Degeneration** by which the structure is largely destroyed, and in the second place in a **Regeneration** by which it is restored.

The **Causation** of this peculiar degenerative process is connected with the nutritive arrangements of the nerves. The nutrition of nerve fibres in general is under the command of certain ganglionic trophic centres, and when the fibres are cut off from these centres they suffer in their nutrition. For the sensory or afferent fibres the trophic centres are in the ganglia of the posterior or sensory roots. For the motor or efferent nerves the centres are in the anterior cornua of the spinal cord. Most nerve stems consist of both afferent and efferent fibres, but the division of the stem will cut the fibres off from their trophic centres, whether they be afferent or efferent. It is interesting to observe that when the posterior nerve root is divided on the proximal side of the ganglion, its fibres degenerate in the central direction, or towards the spinal cord.

The changes about to be described have been studied chiefly by ex-



perimentation on animals, but similar conditions have been observed in man after division of nerves. In several diseases of nerves also, such as inflammation, leprosy, syphilis, there may be interruptions of the nerve fibres leading to lesions similar to those occurring in consequence of division, but perhaps mixed up with other phenomena.

The **Degenerative process** occurs in the whole peripheral distribution of the divided nerve, and almost simultaneously. The most obvious change is in the medullary sheath. It coagulates, breaks up into drops, and through time disappears by absorption. This disintegration of the medullary sheath occurs gradually, and the granular fat into which it breaks up is partly taken up by the nuclei of the nerve fibre, but partly also finds its way out of the primitive sheath and is found in the surrounding connective tissue and the walls of the capillaries. There is some difference of opinion as to the part taken by the axis cylinder in the process. Erb asserts that it persists after the medullary sheath has been destroyed, but Ranvier states that it is broken up, its interruption corresponding with the abolition of electric conductivity in the nerve. Whether the axis cylinder is destroyed or not, the nerve fibre undergoes a great transformation by the loss of its medullary sheath, and it becomes converted into a pale fibre, interrupted at intervals by the nuclei or by some persisting clumps of myeline.

It is asserted by Ranvier that the nuclei, which we have seen to exist inside the primitive sheath between every two nodes, take an active part in this process. They enlarge and divide, and, by impinging on the medullary sheath, help to break it up. It is by them also, according to this author, that the axis cylinder is interrupted. The enlargement and division of the nuclei is somewhat similar to that which occurs in muscle in certain lesions of the spinal cord to be considered afterwards, and it is regarded as inflammatory in its nature.

At the place of division of the nerve, as there is a wound, there are signs of inflammation. Leucocytes collect between and around the cut ends of the nerve, and even penetrate into the primitive sheath for some distance. These leucocytes, which are most abundant soon after the section, attack the medullary sheath, and assist in breaking it up; the myeline is taken up by the leucocytes so as to give them the appearance of compound granular corpuscles. In the central end of the divided nerve, however, the destruction of the medullary sheath is limited, as the invasion of leucocytes generally stops short at the first node.

After a time the inflammation subsides largely, and the wound, including skin and soft parts, is united by a cicatrix formed in the usual way. The divided ends of the nerves are united by a pale cicatricial band, which does not as yet contain any proper nervous elements, and does not effect a restoration of the conductivity.



Conduction is restored by a process of **Regeneration**. According to the researches of Ranvier, this occurs entirely by the axis cylinders of the central end budding out and extending first into the cicatrix and then into the peripheral end. The axis cylinder enlarges at its extremity and becomes divided longitudinally into several fine fibres, which grow out into the cicatrix. Arrived at the cut end of the peripheral portion they penetrate into it and very frequently pass into a primitive sheath. In this way a number of new-formed axis cylinders may be found inside an old nerve-tube, and there may be alongside of them some pieces of persisting myeline. These new axis cylinders after a time acquire medullary sheaths, and the regeneration of the nerve is completed. According to Remak and others, the new fibres are not formed entirely by budding from the central end, but arise also from the remaining axis cylinders of the peripheral end.

Along with the nerves, the **Muscles** suffer, their nutrition being under the same trophic centres as the motor nerve fibres. The muscular cylinders diminish in diameter, the transverse striation becomes less distinct and the fibres become granular. If regeneration does not occur the muscular fibres lose their transverse striation entirely, become greatly narrower, and may even present hyaline degeneration (see under Coagulation-Necrosis, p. 149). The atrophy of the muscular fibres is often accompanied by an interstitial inflammation resulting in a newformation of connective tissue between the muscular fibres, a kind of cirrhosis, leading, in cases where the conduction of the nerve is not restored, to considerable shortening of the muscle and deformity. If regeneration of the nerve occurs then the muscle is restored, but there is frequently some prolonged or permanent damage. The muscular fibres remain partly of smaller diameter and there may be some permanent interstitial overgrowth of connective tissue.

There are also **Trophic changes** frequently manifested in the skin and other structures, such as will fall to be described later on as **Trophoneuroses**. They consist of atrophy of the skin, œdema, and occasionally the peculiar condition described as "glossy skin." There may also be swellings of the joints.

2. **Hyperæmia and acute inflammation of nerves.**—These conditions are exceedingly common as secondary processes in the neighbourhood of wounds and inflammations, but are rarely primary. Under such circumstances the vessels of the nerve may be highly engorged with blood and there may even be capillary hæmorrhages. Leucocytes may be exuded, and may be found in the connective tissue between the nerve fibres. It is very rare, however, to find a proper suppurative inflammation of a nerve. A stem may be bathed in pus and almost isolated by the suppuration around, and yet there may be almost no pus in the connective tissue of the nerve itself. This seems to indicate that the perineurium forms a barrier between the lymph spaces of the nerve and those of surrounding parts. If a nerve be exposed and isolated, and the wound filled with water in which vermilion is suspended, the leucocytes



which accumulate take up the vermilion and carry it in various directions, but not into the nerve, showing that there is no open path into it. But if a nerve in a suppurating wound be itself wounded so as to lay open its internal structure, the suppuration will readily extend into it.

**Chronic neuritis** is, like acute, for the most part secondary. It may remain after an acute attack, or it may take origin in a prolonged inflammation of some peripheral organ. It may have its starting point in an inflammation of a joint, in dysenteric inflammation of the intestine, in inflammation around the kidney, and so on.

**Inflammation of the pelvic organs**, the bladder, uterus, etc., perhaps most frequently gives rise to neuritis by extension to the nerves. It is very important to observe that inflammations originating thus may extend up the nerves, and we may have an **Ascending neuritis** which may even extend to the spinal cord and its membranes. It is to be presumed that the irritant gradually finds its way along the lymph spaces of the nerve, which we have seen to be somewhat independent of those around. Many cases which have been regarded as reflex paralysis have been due, according to Leyden, to this extension of chronic inflammation to the cord. There may also be an extension downwards to the muscles and a consequent **Interstitial myositis**.

The inflammation manifests itself as an interstitial inflammation affecting the connective tissue of the nerve; the perineurium and endoneurium present an excess of round cells and become gradually thickened. As a consequence of this the nerve fibres atrophy, the medullary sheath being first destroyed. The nerve as a whole may be thickened and indurated, while there is actually an atrophy of the proper conducting fibres. These will be recognized as the general characteristics of chronic interstitial inflammations.

**Multiple or Peripheral neuritis** is a name given to a group of cases whose pathology has only recently been elucidated. The disease consists in a more or less acute inflammation of a number of nerve stems in the limbs. In some cases it seems to be due to a definite morbid poison. Thus it sometimes ensues on diphtheria or leprosy, and it occurs as an endemic disease in Japan where it is known as **Kak-ke**, and in India, where it is called **Beri-beri**. Multiple inflammations of the nerves form part at least of the phenomena of these diseases. The commonest cause, in this country at all events, is alcoholism, and here the term **Alcoholic neuritis** is almost equivalent to multiple neuritis. There are also changes in the nerves in locomotor ataxia which partake somewhat of the characters of multiple neuritis.

The inflammation is acute or subacute and accompanied at first by exudation of leucocytes in the connective tissue, as well as by changes



in the proper nerve fibres. These changes may be variously interpreted as evidences of parenchymatous neuritis, or as secondary degenerations. The changes in the nerve fibres are very similar to those which follow the division of a nerve, the medullary sheath breaking up into granules and becoming absorbed. The interruption of the nerve fibres by the inflammation doubtless plays an important part in this result, though perhaps a direct affection of the nerve fibres may be present in some cases. As the case becomes chronic there is an interstitial induration such as that noted in chronic neuritis.

In most cases it is only the nerves of the limbs which are affected, and it is interesting to note that however severe the affection the anterior roots escape.

The muscles present changes similar to those which occur in consequence of division of nerves, and there may be great wasting in consequence. There are also, occasionally, trophic changes in the skin similar to those already referred to.

**3. Infective newformations and tumours.**—**Syphilitic gummata** are very infrequent in the peripheral nerves, but they do occur, especially on the cranial nerves inside the skull, where they are probably propagated from the membranes of the brain. **Leprosy**, as we have already seen, sometimes manifests itself in the form of tumours of granulation tissue in the nerves.

**Sarcoma** is very rare in nerves. **Cancers** do not occur as primary tumours, but nerves are often involved in the extension of such tumours. It often happens that a cancer or sarcoma grows around a nerve, which passes through its midst without becoming the seat of the tumour tissue.

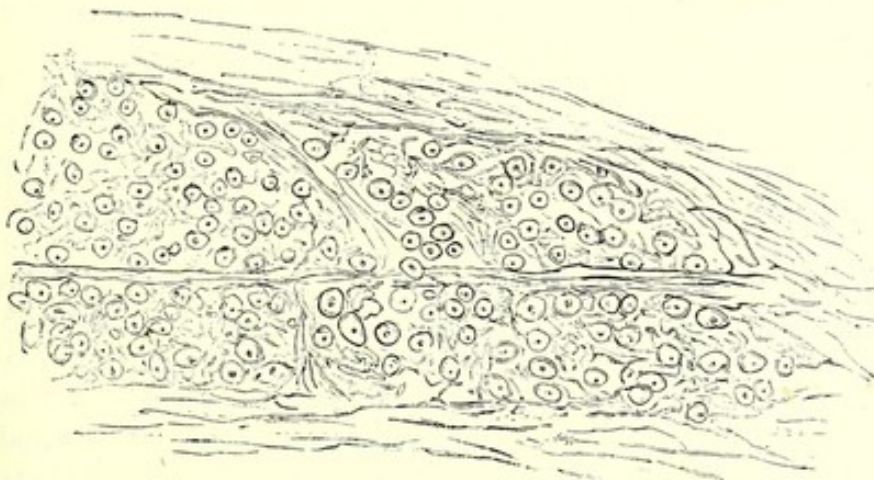


Fig. 202.—Transverse section of a neuroma, showing medullated nerve fibres.  $\times 80$ .

This is again to be associated with the apparent independence of the lymph spaces in nerves. Sometimes, however, a cancer breaks into a



nerve and grows in the lymph spaces between the perineurium and the bundle of fibres. In such cases the nerve fibres undergo degeneration.

The term **Neuroma** is applied to almost all forms of tumours in the course of nerves, and as the majority of these do not consist of nervous tissue, they are to be regarded as false neuromata. In the true neuroma there is newformation of nerve fibres which form a considerable portion of the tissue (see Fig. 202). The false neuromata are mostly fibrous tumours of nerves, and there are two forms which deserve special mention.

**Plexiform neuroma** consists of a series of thickened cords composed of connective tissue in the midst of which the nerve fibres are contained. The connective tissue may, to a considerable extent, take the characters of mucous tissue.

**Multiple neuromata** are really fibrous-tissue tumours, and as such are described at p. 228. The painful subcutaneous tumour is also a fibrous neuroma (see p. 241).

**Literature.**—RANVIER, *Leçons sur l'histologie du système nerveux*, 1878; CORNIL et RANVIER, *Histol. path.*, 2nd ed., 1881, i. 661; WOLBERG, *Deutsch. Zeitschr. f. Chir.*, xviii. and xix., 1883 (with literature); PITRES et VAILLARD, *Arch. d. phys.*, v., 1885. *Multiple neuritis*—see full account by BUZZARD, *Harveian Lectures*, in *Lancet*, 1885, vol. ii., also separate publication; DRUMMOND, *Peripheral paralysis*, 1888; FINLAY, *Trans. Med. Chir. Soc.*, 1887. *Tumours*—see literature under Neuroma, p. 242.

## B.—THE SPINAL CORD AND MEDULLA OBLONGATA.

**Anatomical Introduction.**—The cord is made up of grey substance, forming the ganglionic centres and consisting of ganglion cells in the midst of a fine network, and of white substance consisting of medullated nerve fibres having essentially the structure of those in the peripheral nerves.

The grey substance of the cord is arranged in the well-known form of an anterior and a posterior cornu on either side. The ganglionic centres have their seat chiefly in the anterior cornua, and form tolerably definite groups whose arrangement may be followed by reference to Fig. 203, which shows sections of the lumbar and cervical enlargements. There are the lateral groups divided into antero-lateral and postero-lateral groups (*al* and *pl*), the anterior group (*a*), and the internal group (*i*). Towards the centre of the horn is the central group (*c*). The median group (*m*) is much larger in the cervical than in other parts of the cord, and so causes the horn to be extended laterally (see Fig. 203). Lastly, there is a group of cells, generally called Clarke's vesicular column, situated near the internal border of the posterior horns close to the posterior commissure. The group is present only in the lower part of the cervical enlargement, in the dorsal region and in the upper part of the lumbar enlargement (see *vc* Fig. 204).

The white substance, consisting of nerve fibres, forms connections in the first place with the various orders of centres in the cord itself and in the second place with the higher centres above the cord, in the brain. We may thus distinguish two sets of



nerve fibres, one forming connections within the cord and medulla and the other forming communications between the cord and the cerebellum and cerebrum. These two sets of fibres will be divisible again into afferent and efferent.

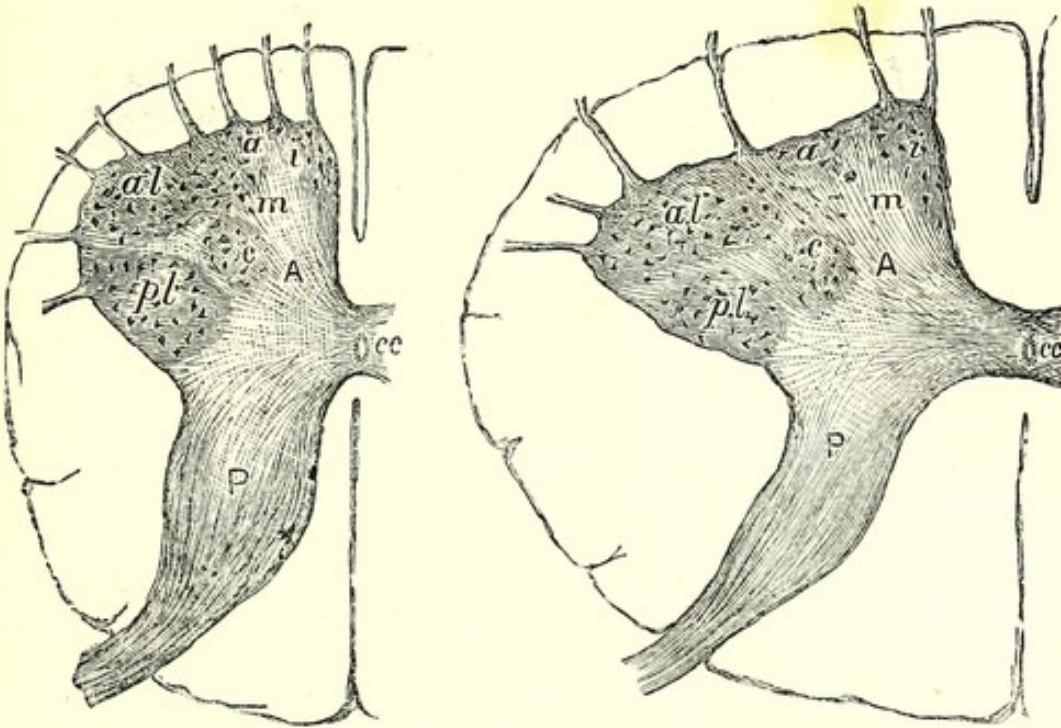


Fig. 203.—Sections of spinal cord from middle of lumbar and cervical enlargements, showing groups of ganglion cells; *al* and *pl*, antero-lateral and postero-lateral, *a*, anterior, *i*, internal, *c*, central, *m*, median, groups. (Ross.)

It is a fact of very peculiar interest, that the two sets of fibres distinguished above seem to be developed not only separately but at different periods, and so the aid of embryology has been sought to enable us to distinguish between them. The fibres which connect the different parts of the cord and medulla with each other may be regarded as the **Primary or Fundamental** fibres, and it is found that they are the first formed, while those forming higher connections are of subsequent development, and may be named **Secondary or Accessory** fibres. As nerve fibres are first developed without the medullary sheath, and recently formed tracts are therefore much paler than those which have acquired it, we have in this a means of distinguishing the fundamental from the secondary. What follows will be understood by reference to Fig. 205, which represents a transverse section of the cervical cord in the foetus of nine months.

The first developed fibres immediately surround the grey matter, and are called the **Anterior and Posterior root-zones**, also the columns of Burdach. As they form communications between one part of the grey

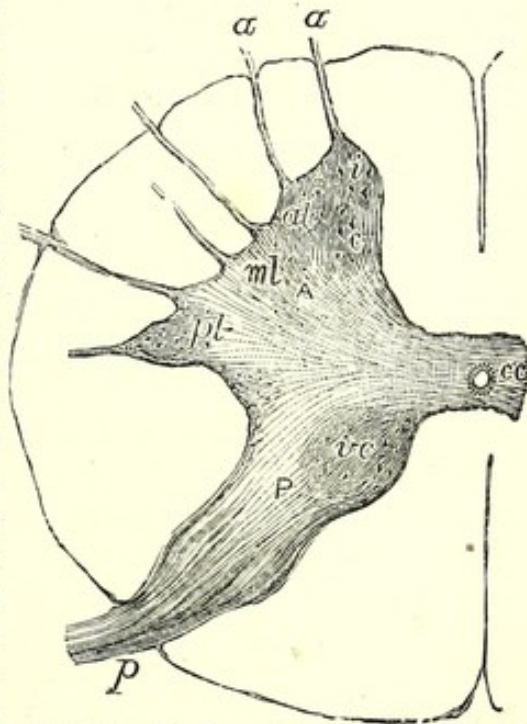


Fig. 204.—Section of dorsal region of cord. Letters same as in previous figure, with the addition that the antero-lateral and postero-lateral groups are separated by a medio-lateral area (*ml*), and Clarke's column (*cc*) is shown. (Ross.)



matter and another, the fibres are comparatively short and their number is generally in proportion to the amount of grey matter, or at least of ganglion cells in the horns. These fibres therefore do not diminish from above downwards as do the others.

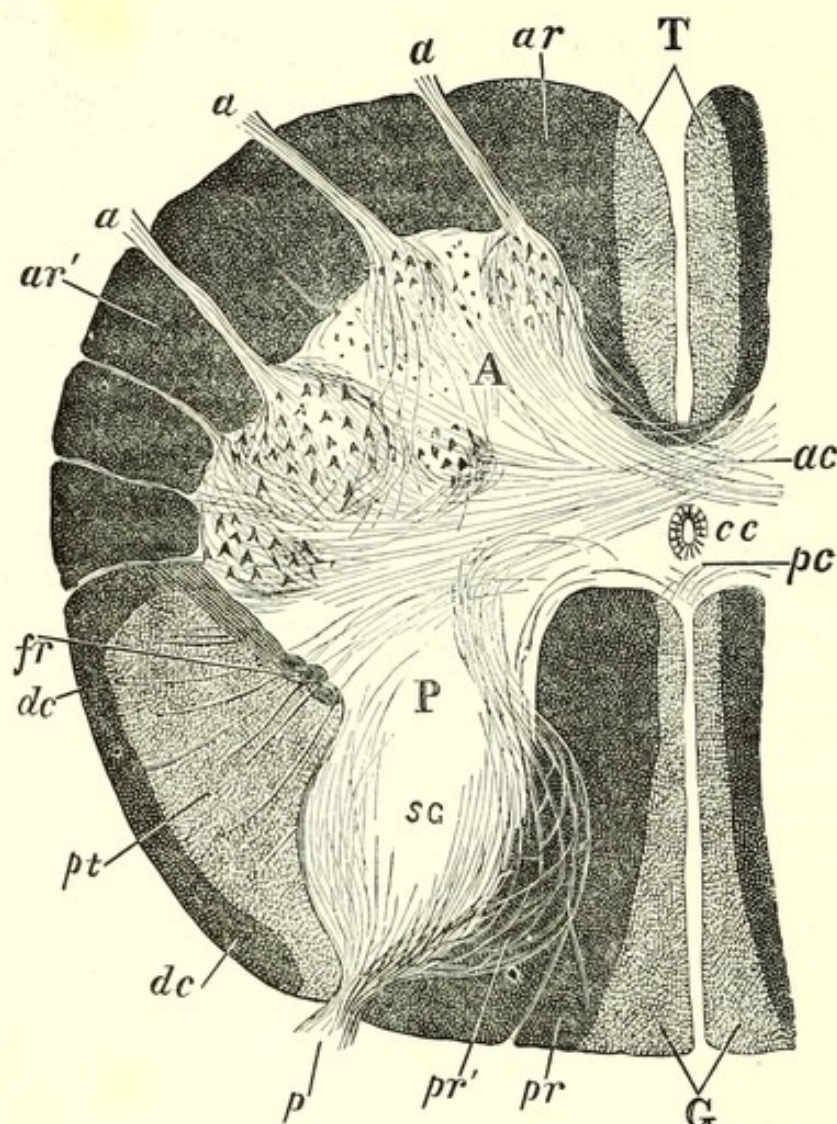


Fig. 205.—Cervical enlargement of cord in a foetus of nine months. A and P, anterior and posterior cornua; G, columns of Goll; T, columns of Türk; ar and pr, anterior and posterior root-zones; pt, pyramidal tract; dc, direct cerebellar tract. (Ross.)

Of the fibres forming communications between the brain and cord, the best known are those which convey the motor impulses from the brain to the cord. These form the **Pyramidal tract**. We shall afterwards trace them from the cortex of the brain downwards, but at present we take them up at the medulla oblongata. Here they form the anterior pyramids and most of the fibres decussate so that in the cord they occupy the opposite side to that which they have in the brain. Some of them, however, do not decussate, but remain in the anterior parts of the cord, forming a small band on either side of the anterior longitudinal fissure, the **Column of Türk** (*T* in figure). The great mass of the fibres, having decussated, pass to the lateral column of the cord, where they occupy a definite position in its posterior parts (*pt* in figure). The fibres in both these positions diminish in number from above downwards, as they pass into the grey substance of the cord at successive levels in order to connect with the centres in the anterior cornua.



The secondary centripetal fibres, or those which form sensory connections between the cord and brain, are represented by a tract in the posterior columns lying next the posterior longitudinal fissure, and occupying a position somewhat similar to that of the columns of Türk anteriorly. These are called the **Columns of Goll** (*G* in figure) and they also diminish from above downwards.

Besides this there is a tract which forms communications between the cerebellum and the cord, but which is not of late development. This is the so-called **Direct cerebellar tract** (*dc* in figure), which lies in the lateral column outside the pyramidal tract, and as if flattened against the surface. The function of this tract is not known, but it is composed of centripetal fibres and diminishes from above downwards even more quickly than the pyramidal tract, so that by its disappearance the latter may come to the surface.

According to Gowers there is a centripetal tract anterior to the direct cerebellar and pyramidal tracts, and lying for the most part close to the surface of the cord, where it extends almost to the anterior median fissure. This is called the **Antero-lateral ascending tract**. It is sometimes compared with the cerebellar tract.

In the **Medulla oblongata** the grey and white substances of the cord may be traced upwards, but they undergo considerable dislocation. From the examination of Fig. 206 it will be seen that, as the central canal passes backwards and finally opens out in the fourth ventricle, the grey matter departing from its arrangement into cornua but still aggregated in the neighbourhood of the central canal and ventricle, forms various masses in the posterior part of the medulla.

These masses have special importance as being the nuclei of origin of certain nerves, and will be afterwards more particularly referred to, in connection with bulbar paralysis. The white substance gradually comes to occupy the middle and anterior parts of the medulla, and the olivary body is intercalated in its midst. The pyramidal tract is easily recognized here, forming the anterior pyramids (*p*), which decussate at the lower part of the medulla (*A*). The motor fibres having assumed a position in front remain anterior to the sensory in the rest of their course upwards in the brain.

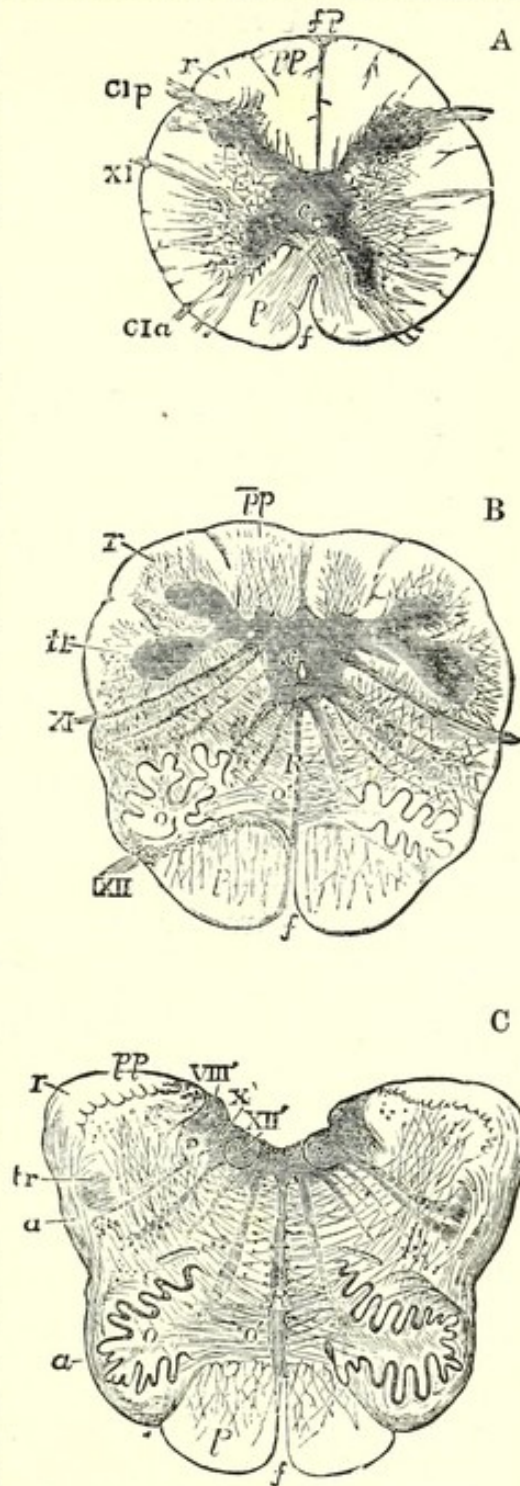


Fig. 206.—Medulla oblongata at various levels, *A*, at decussation of pyramids (*p*), the general shape of the cornua still retained. *B*, higher up, the grey matter passing backwards, and pyramids becoming more isolated. *C*, in fourth ventricle, the nuclei in the floor of which are shown. (QUAIN.)



**Literature.**—For a very full exposition of the structure and development of the cord see Ross's *Nervous system*, vol. i. An important contribution is FLECHSIG, *Die Leitungsbahnen im Gehirn und Rückenmark des Menschen*, 1876. For diseases of spinal cord, Ross and GOWERS are very complete, also ERB's very excellent work in Ziemssen's *Encyclopædia*, CHARCOT, *Lectures on dis. of nervous syst.*, Syd. soc. transl., 2nd series, 1877 and 1881. BRAMWELL, *Dis. of spinal cord*, 1882; HAMMOND, *A treatise on dis. of nervous system*, 6th ed., 1876; LEYDEN, *Klinik der Rückenmarkskrankheiten*, 1875.

#### I.—SECONDARY DEGENERATIONS IN THE CORD.

**Grey degeneration. Sclerosis.**—We have already seen that when a nerve stem in an animal or in man is divided, the peripheral portion of the nerve degenerates. The most prominent changes are in the medullary sheath which coagulates, then breaks up, and is finally absorbed. The degeneration occurs from the point of section towards the periphery, and we have seen that the explanation of this seems to be that the nerve fibres are cut off from their trophic centres in the cord.

In the central nervous system a somewhat similar secondary degeneration occurs when the fibres are interrupted, either by division, as in experiments on animals, or by coarse lesions in man. The degeneration is of the same character as in peripheral nerves, and results in the destruction of the medullary sheath. It is known that the opaque dead white colour of the white or fibrous nerve substance is due to the medullary sheath, which is a highly refracting fatty substance. If this medullary sheath be lost, then the white nerve substance becomes grey, and so we speak of **Grey degeneration** in all cases where the medullary sheath is lost, whether from the cause we are considering or not. A tract of white nervous tissue affected with this degeneration will be like a tract of grey substance.

In preparing the tissue of the nervous system for microscopic examination, some form of the method introduced by Lockhart Clarke is generally used. In this method staining by carmine or other dye is employed, and as the medullary sheath does not take on the staining at all, and as it is the most bulky constituent of the white substance, the normal white substance as a whole is less deeply stained than the grey substance. When secondary degeneration has occurred, the effect of staining is to make the altered white substance appear like the grey substance. In figures representing grey degeneration therefore the affected parts will be shaded like the grey substance.

When a degenerated area is examined more particularly under the microscope the most marked change is the great reduction in the number of nerve fibres, as shown in Fig. 207. The connective substance is greatly increased, and the nerve fibres appear only at intervals, a few surviving in the midst of the general destruction. There is some-



times also considerable shrinking of the degenerated area, but this is much more manifest in the medulla oblongata and parts above it than in the cord.

In addition to simple degeneration there is sometimes a newformation of connective tissue, which some regard as inflammatory in character. The inflammatory character is little marked in secondary

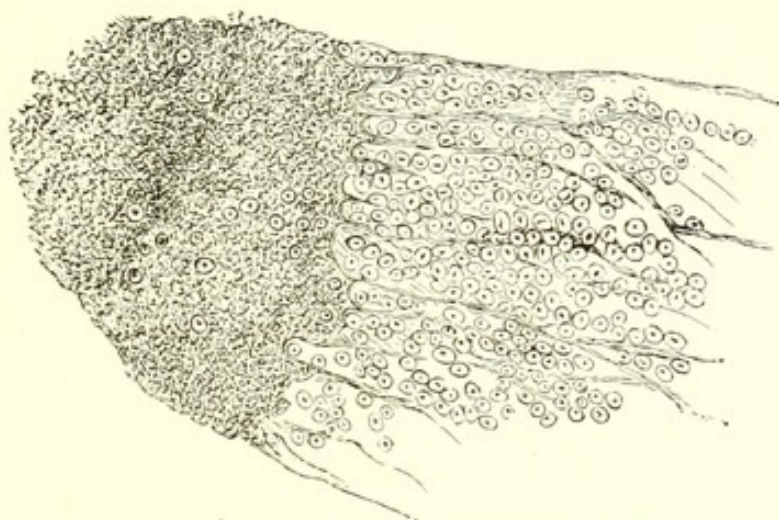


Fig. 207.—Sclerosis or grey degeneration of cord. To the right is normal white substance. To the left the degenerated tissue shows a granular basis-substance in which a few nerve fibres are still visible. From a case of descending degeneration.  $\times 80$ .

degeneration, but there are some forms of grey degeneration in which the inflammatory newformation is perhaps the primary factor, and the degeneration of the nerve fibres the result of it. Whether there be new-formed connective tissue or not, and whether the process be inflammatory or not, the absence of the soft medullary sheath causes a hardening of the tissue, and so the term **Sclerosis** when applied to the white substance is nearly equivalent to grey degeneration.

It has been said above that the degeneration occurs in the fibres which are cut off from their trophic centres, and it seems that in the cord the trophic centres are at the lower termination of the centripetal fibres and the upper termination of the centrifugal. Hence the degeneration follows the direction in which the nerve conducts, passing upwards from the seat of lesion in the case of centripetal fibres and downwards in the case of centrifugal. So we speak of ascending and descending sclerosis.

The degeneration takes some time to develop. According to experiments in dogs it begins fourteen days after the infliction of an injury, but it is many weeks before the appearances are fully established.

**Descending grey degeneration** (*Descending sclerosis*).—As we have just seen, this lesion affects centrifugal or motor fibres, including the fibres of the anterior root-zone and the pyramidal tract. The former



are short fibres, and when they are interrupted there is a descending degeneration extending only a short distance downwards. But the fibres of the pyramidal tract are continuous from the brain downwards to the extremity of the cord, and wherever interrupted they show degeneration in all parts situated below the lesion.

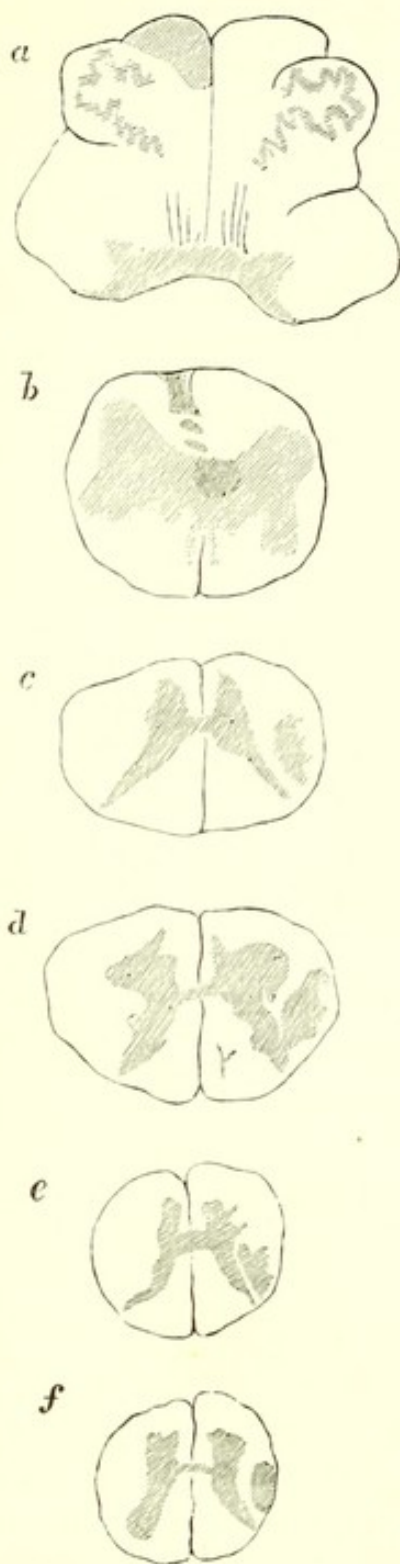


Fig. 208.—Descending degeneration in medulla oblongata and cord. *a*, Medulla at fourth ventricle, *b*, at decussation, *c*, *d*, *e*, *f*, cord in upper and lower cervical, dorsal, and lumbar regions.

The degeneration of the pyramidal tract is most frequently brought about by a **Lesion of the brain**, and in that case it exists in the parts of the tract above as well as in those below the medulla oblongata. We have here to consider it in the medulla and cord. In the medulla oblongata it occupies the anterior pyramid (see Fig. 208 *a*), where it frequently produces great shrinking. At the lower part of the medulla (*b*) the degenerated fibres decussate and the degeneration takes up its position at first at the side of the central canal (see figure), afterwards, in the cord, passing into the lateral columns in the regular position of the pyramidal tract (*c*, *d*, *e*, *f*). In some cases it entirely decussates, but the column of Türck in the anterior white column is also affected in most cases. In its whole course the lesion diminishes from above downwards, but is traceable down to the lumbar region.

The pyramidal tract may be interrupted in any part of its course, and it may be so **in the cord**. If the entire cord be divided there is, immediately beneath the point of section, a degeneration of the anterior root-zones which is continued but a short distance. There is also a degeneration of the pyramidal tract on both sides continued downwards the whole length of the cord. The interruption either in the cord or brain may be incomplete, and in that case the

degenerated fibres will be fewer in number and the area less distinctive in appearance.

**Effect on function of descending sclerosis.**—We have already seen



that the pyramidal fibres end in the cord at successive levels, passing into the ganglionic centres. When the fibres are degenerated the ganglion cells will not be affected, except that they are cut off from the higher centres and left more to themselves. They are still connected with the muscles, which retain their contractile power. Voluntary motion is lost, but certain involuntary muscular phenomena may be even exaggerated.

**Late rigidity**, occurring in hemiplegia, or in paraplegia from injury to the cord, comes on a considerable time after the onset of the paralysis, and may be regarded as coinciding in time with the full development of the sclerosis. There is here a more or less permanent contraction of some muscles with absence of contraction in others, producing sometimes fixation of the members of the body in special positions so as to have the appearance of deformity. This fixed condition, due to the muscular spasm, is often called **Contracture**. It implies a continuous impulse to the contracted muscles originating in the cord or elsewhere. Charcot suggests that the inflammatory process which he supposes to be involved in the sclerosis may irritate the fibres passing to the anterior cornua and so result in the stimulation of the muscular centres there. But this view can hardly be accepted; there may be almost no signs of inflammation in the affected part and yet marked rigidity. Besides, it is difficult to understand how irritation of degenerated and virtually lost fibres should cause stimulation of the ganglion cells. A more probable explanation is suggested by Hitzig. Taking as an example the case of the arm in hemiplegia, it appears that the contraction occurs most readily in those muscles which are in a position to contract most easily. The hemiplegic generally lies in bed with his forearm across his chest, and even when walking about he supports it across his chest, and it is the biceps which becomes rigid. Then it has been shown that when the fingers are released from the action of the muscles and left to assume the position to which the bones and ligaments best accommodate themselves, they assume a semi-flexed position, such as we see in the dead body. It is obvious that the flexor muscles will most easily contract under these circumstances, and it is they which get rigid in hemiplegics. Take along with this the fact which Volkmann points out, that muscles can actively contract but cannot actively relax, and it is seen that any slight impulse is apt to be cumulative when there is no action of antagonistic muscles. The ganglion cells of the anterior cornua are still in connection with the muscles, and although cut off from the upper brain are still exposed to irregular and, as it were, accidental stimulation. There are reflex stimuli, and there are stimuli from above conveyed in a roundabout way through the still open communications in the cord. A



multitude of slight stimulations will reach the ganglion cells and feeble impulses will be conveyed to the muscles. Those which from their position are stretched will not contract, but those which are so placed as to contract easily will do so, at first feebly but with cumulative force. This view is supported by the fact that, in the early stages of late rigidity, there is often considerable relaxation of the muscles after prolonged rest, as in sleep, so that a limb which was rigid at night is found in the morning soft and moveable. The ganglion cells in sleep are protected from external stimulation and they cease acting.

The exaggerated **Tendon reflex** may be regarded as due to the isolation of the muscular centres in the cord. When the control of higher centres is removed, lower centres usually act more readily. The skin reflex is frequently decreased, but the muscular centres seem to be more powerfully acted on by stimuli coming from the muscles.

The so-called tendon-reflex has been shown not to depend on the tendon but on the muscle, and doubt has been thrown even on its reflex character (see Gowers vol. i., p. 13). It is certainly so far reflex as that it is abolished by anything which interrupts the reflex circle in any of its parts.

It may be added that these two symptoms are prominent in the spastic paralysis of Erb, where there is, as we shall see farther on, a spontaneous sclerosis of the lateral columns.

**Ascending grey degeneration** (*Ascending sclerosis*).—This condition occurs as a result of any cause which interrupts the ascending or sensory fibres of the cord. It may be the consequence of pressure by tumours, hæmorrhages, abscesses, fractured or displaced vertebræ, inflammatory products, etc. It occurs in parts above the seat of interruption, and the degenerated fibres are centripetal.

The degeneration affects the centripetal fibres, and these we have already seen to be of two kinds. There are the short fibres communicating between different segments of the cord, and forming the posterior root-zones, and there are the longer fibres communicating between the cord and brain, and forming the columns of Goll, the direct cerebellar, and partly the antero-lateral tracts. Immediately above the lesion, all of these are affected so that the degeneration has considerable lateral extension involving the whole of the posterior columns and the direct cerebellar tracts. But the lesion soon limits itself to the columns of Goll, the direct cerebellar, and partly the antero-lateral tracts (see Fig. 209), and in these parts it can be traced up to the restiform body on the one hand, and the cerebellum on the other. According to Schiefferdecker, the degeneration in these situations diminishes from below upwards, the diminution occurring at intervals as if, at definite levels, fibres passed into the cord.



Some cases have been recorded in which a tumour pressing on the cauda equina, or a severe traumatic lesion of the sciatic nerves, had led to ascending degeneration of the cord. In the lumbar and lower dorsal regions the whole posterior white columns were degenerated, but on passing upwards the degeneration became limited to the columns of Goll and the cerebellar fibres as above. We may infer from this that the fibres of the posterior column are, to a large extent, continuous with the posterior nerve-roots, and that they have their trophic centres in the ganglia of these roots outside the cord.

It will be evident that a lesion which interrupts the cord will lead to an ascending degeneration above its seat and a descending degeneration below it, as in figure 198. The degeneration will diminish in both cases as we pass from the lesion, quickly at first, but afterwards more gradually.

**Degeneration in the cord after amputations.**—The removal of a limb abolishes the function of the nervous structures concerned in the movements and other actions of the limb, and so these structures undergo atrophy from disuse. The posterior roots of the nerves are often slightly atrophied, but the white columns of the cord are not affected. The principal change is in the anterior cornua, where some of the ganglion cells completely disappear, and others are shorn of their processes and atrophied. This occurs at a part of the cord corresponding with the amputated limb, and on the same side of the body. If the amputation has been made comparatively late in life, there is usually little change in the cord; the earlier the age of the person at the time of the amputation the more likely are these changes to occur.

In a case recorded by Edinger, in which intra-uterine amputation of the forearm had occurred, and the person lived to the age of 52, there

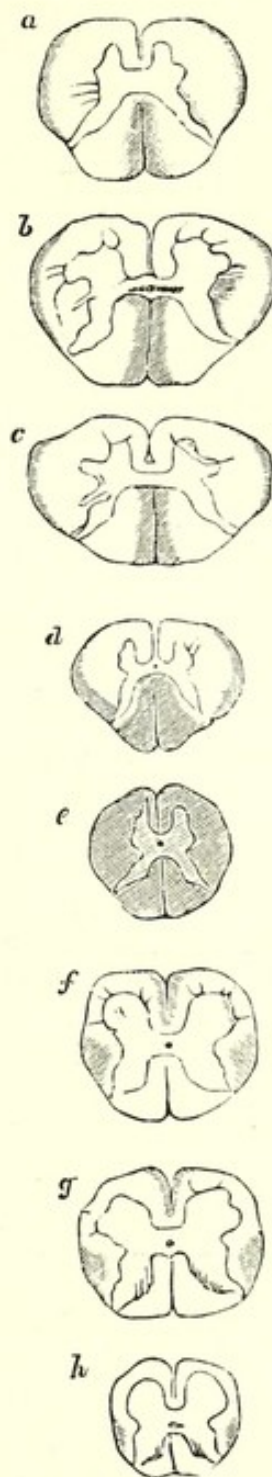


Fig. 209.—Transverse myelitis (*e*) with secondary ascending and descending degeneration. At *d*, the ascending degeneration affects the posterior root-zones, the columns of Goll, and the direct cerebellar tracts, but above that it is confined to the two latter. At *f* to *h*, the columns of Türk and the pyramidal tracts are affected. (ERB.)



was considerable atrophy of the nerves and of the corresponding half of the cord, especially of the grey matter. There was also some atrophy in the motor region of the convolutions in the brain.

**Literature.**—Besides general works already noted, CHARCOT, Lectures on localization of cerebral and spinal diseases, Syd. Soc. transl., 1883; TÜRCK, Sitzungsber. d. Akad. d. Wiss. Wien, vols. vi., 1851, and xi., 1853; LEYDEN, Deutsch. Klinik, 1863; GOWERS, Diagnosis of dis. of spinal cord, 2nd ed., 1881; SCHIEFFERDECKER, Virch. Arch., lxvii., 1876; DRESCHFELD, (Cord after amputation) Jour. of Anat. and Phys., 1879, xiv. 424; GENZMER, Virch. Arch., 1876, lxvi. 265; EDINGER, Virch. Arch., 1882, lxxxix. 46.

## II.—INFLAMMATIONS OF THE SPINAL CORD.

In its widest acceptance, inflammation of the spinal cord, or myelitis, includes a large number of widely different conditions, each of which will be considered separately. The cases may for convenience be divided into two groups. In one of these the affection has a limited longitudinal extension, but involves the cord rather in its thickness. For this class of cases it is customary to use the expression **Transverse myelitis** in order to distinguish them from those in which a considerable length of cord is attacked, while the inflammation has a limited transverse extension. The former are usually due to a definite cause which acts on the cord at a particular level. In the second group the inflammation follows a particular physiological system in the cord, as for instance a tract of white substance, and extends in it for some distance from above downwards. Such inflammations may involve the white substance or the grey substance, and they are often distinguished by the names **Poliomyelitis** ( $\pi\omicron\lambda\iota\acute{o}\varsigma$  = grey) and **Leucomyelitis** ( $\lambda\epsilon\upsilon\kappa\acute{o}\varsigma$  = white). These inflammations have for the most part no assignable cause, and are sometimes called spontaneous. In some of them the process shows, perhaps, more of a degenerative than an inflammatory character. Closely allied to these **Systematic inflammations** there is a form of spontaneous myelitis in which the distribution is somewhat irregular, but still has a certain tendency to localize itself. We shall consider the two classes indicated above under separate headings.

### A.—TRANSVERSE MYELITIS.

1. **Acute transverse myelitis** (or simply *Acute myelitis*).—Acute inflammation of the spinal cord is nearly always secondary to some other condition. It may be produced by fracture, dislocation, or acute curvature of the vertebræ, with direct injury to the cord. It may be caused by extension of inflammation, as when an abscess penetrates into the



spinal canal, producing inflammation first of the meninges. A tumour may produce it by compression, but as the compression is usually gradual there is more frequently a chronic myelitis, and the same applies to most cases of curvature of the spine. It also occurs in consequence of exposure to cold, especially when the body has been overheated. It may be a complication or sequel of an acute specific fever, as typhus, and it is said to be not uncommon in syphilitic patients. The inflammation of the cord is often associated with inflammation of the membranes, and the condition is then designated **Meningomyelitis**.

The myelitis is limited in longitudinal extent, and as the usual result is softening of the cord, the expression **Acute softening of the cord** is almost equivalent to myelitis. The softening is usually most manifest in the grey substance and may even be apparently confined to it, but it really involves the white substance as well, and usually the whole thickness of the cord.

The softened nervous tissue presents, in different cases, considerable variations in colour, so that red, yellow, grey, white, and even green softening have been described. If much blood has escaped from the vessels, there will be red softening merging into yellow.

The characteristic morbid changes are the breaking-up of the nervous tissue and fatty degeneration of the other structures. In the white substance drops of myeline escaped from the medullary sheath are found, and the axis cylinders are swollen. The fatty degeneration is manifested by the presence of abundant compound granular corpuscles, which, to some extent, are probably leucocytes or neuroglia cells, which have picked up the disintegrated myeline; but there may be abundant leucocytes apart from these cells. There is also fatty degeneration of the walls of the blood-vessels, the fat here having a similar source.

The myelitis rarely goes on to suppuration, but usually passes into a chronic stage. The fat, both of the medullary sheath and of the compound granular corpuscles, is absorbed, and a condition of grey softening remains. As the inflammation becomes more chronic there is new-formation of connective tissue of a cicatricial character. In the midst of the connective tissue there are often to be found large cells with radiating processes, sometimes called "spider cells" or "Deiter's cells." These are very obscurely visible in the normal cord where they form part of the neuroglia, but here they are much enlarged. In this way the cord at the part affected may be replaced by a cicatrix, or by a cyst, and its conduction interrupted. In some cases the interruption is not complete, and there may even be a partial regeneration of the conduct-



ing fibres, and a partial recovery from the consequent paralysis. As a consequence of the interruption in the white substance, there is ascending and descending grey degeneration of the usual distribution. There may be also an extension of the inflammation upwards in the pyramidal tract for a short distance from the seat of the lesion.

Gowers has described an appearance as if a newformation of nerve fibres were taking place, and certainly the fact that there may be a partial return of power after motor paralysis which has lasted twelve months, would seem to indicate that there may be some such newformation.

**2. Chronic transverse myelitis.**—We have just seen that this condition may follow on an acute myelitis, but the inflammation may be chronic from the first, when it is produced by an irritant which acts gradually. It is most frequently the result of **Chronic compression of the cord**, as in the case of curvature, or pressure by a tumour. It may also be propagated from the membranes, a spinal meningitis passing into a myelitis. We have already seen that it may arise by extension from a peripheral nerve, the inflammation travelling upwards to the cord. It occurs also from exposure to cold, from strains, etc., as in the case of acute myelitis.

The changes produced are those common to chronic inflammations. There is newformation of a dense connective tissue, developing in the usual way from granulation tissue. The new-formed tissue causes compression of the proper nervous elements and their atrophy. In the white substance the nerve fibres disappear, and in the grey substance the ganglion cells. With the growth of this dense tissue there is an induration of the cord, and the term **Sclerosis** is applicable. In the white substance this is manifested by the usual appearances of grey degeneration. There is also a degree of shrinking of the parts, and this is often especially manifest in the grey substance.

As the nervous structures are partially or completely destroyed there is a greater or less interruption of the cord, and this results in the usual secondary degenerations above and below the seat of the lesion.

**Literature.**—POTT, Palsy from curvature of spine, 1776 and 1782; BRODIE, Injuries of the spine, 1837; JACCOUD, Des Paraplegies, 1864; FROMMANN, Normal und path. Anat. des Rückenmarks, 1867. For more complete literature see Ross, ii. 89.

## B.—SYSTEMATIC MYELITIS AND DEGENERATION.

As already indicated, in the diseases included here the affection follows certain physiological systems, and we have, in the first place, to inquire whether any explanation of this can be suggested.

**General causation.**—In almost all the examples of inflammation



studied in other parts of this work, the extension of the process is along definite paths. The irritant has been brought to an organ by the blood-vessels, or by some path of transit, such as a mucous canal, and it has extended, it may be, by the lymphatics or along a surface. In the lesions here to be considered, however, the problem is much more difficult. The localization of the disease bears no relation to the blood-vessels or lymphatic vessels, and if the irritant is brought by the blood there must be local peculiarities leading to differences in the susceptibility of different structures.

In studying the various structures which constitute the cord we saw that in the development of the cord a certain difference is apparent. In the white substance there are certain **Fundamental structures** constituting the root-zones, and certain **Accessory tracts** which are separately developed, and mostly of later formation. In the grey substance also there are differences in the periods at which the groups of ganglion cells appear, so much so that some of them which appear later than the rest have been named **Accessory nuclei**. These are chiefly the median and medio-lateral groups. These various tracts of white substance and groups of ganglion cells have apparently different powers of resistance, and the more recently developed or accessory structures are least resistant. It is not unlikely indeed that in some cases these may be, from their origin, so unstable that without any obvious exciting cause they may tend to degenerate. In this way the boundary line between actual inflammations and simple degenerations is reached. There are some of the diseases here to be considered which have undoubtedly the characters of inflammation, and even of acute inflammation. But there are others in which the characters are rather those of a chronic degeneration or atrophy of the structures concerned, with very little of an inflammatory character.

There are thus various degrees of susceptibility in the different structures of the cord and medulla oblongata, and when an irritant exists in the blood, developed perhaps in connection with exposure to cold or otherwise, there will be, according to circumstances, various manifestations.

1. **Sclerosis of the posterior columns** (*Locomotor ataxia*, *Tubes dorsalis*).—**Causation**.—The disease has the characters more of a degeneration than an inflammation, and hence is not frequently traceable to any direct injury to the cord. An apparent exception to this is the fact that syphilis is a frequent cause (according to Gowers, in about two thirds of the cases). But there is no definite syphilitic newformation, and it seems rather as if an attack of syphilis left the cord more liable to degenerative changes.



**Characters of the changes.**—If the cord is examined in well advanced cases of this disease, the **Posterior white columns** are found grey and shrunk, and the posterior roots are also atrophied. There is often meningitis, the soft membranes being thickened and adherent to the cord beneath as well as to the dura mater on the surface. In early stages of the disease the changes in the cord may be invisible to the naked eye, but can be detected with the microscope. In advanced stages there may be similar changes in the optic nerve, the oculo-motor, the hypoglossal, etc.

The minute changes consist in atrophy of the nerve fibres and increase of the connective tissue, similar to that described as occurring in secondary degenerations. As to the degree in which actual inflammatory processes exist opinions differ. The result is a sclerosis of the affected regions, and the tissue may have many amyloid bodies in it. These are round or oval bodies three or four times the size of blood corpuscles, and presenting a peculiar bright glancing appearance (Fig. 210). They frequently present a concentric striation like grains of

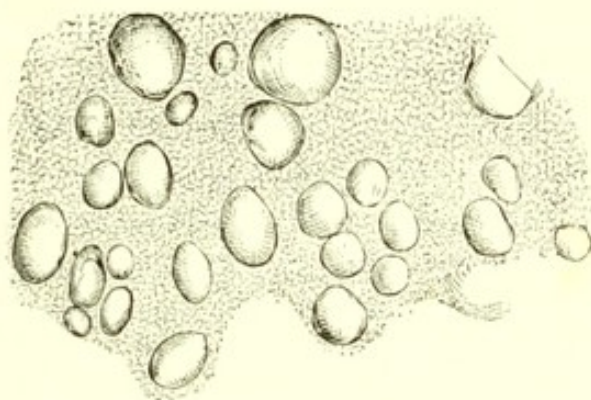


Fig. 210.—Amyloid bodies from cord.  $\times 300$ .

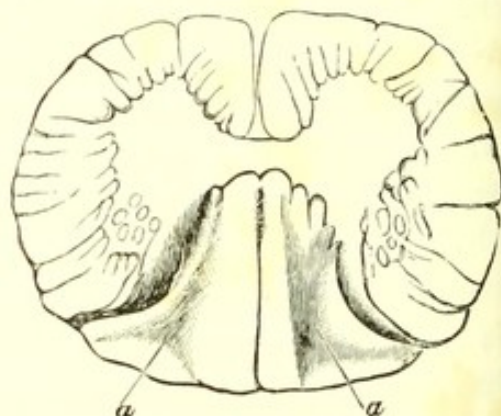


Fig. 211.—Posterior sclerosis in an early stage, confined to the external part of the posterior columns. (PIERRET quoted by ERB.)

potato starch, and with iodine they take on a deep brown coloration. These bodies are sometimes present in such numbers as to form the most prominent feature under the microscope.

The posterior columns are the parts affected, but the disease does not invade these columns uniformly. Its special seat is the outer parts of the posterior columns, namely, the **Posterior root-zones**, or columns of Burdach, and it is stated that these alone are affected in the earlier stages (see Fig. 211). The **Columns of Goll** are secondarily invaded, as the result of an ascending sclerosis. That this is so appears from the actual examination of the cord at various levels. The disease begins, and is usually most advanced, in the lumbar region, but in the lowest part of that region the lateral parts of the posterior columns may be alone involved. On passing upwards the columns of Goll become also affected,



so that in the upper half of the lumbar swelling the degeneration may be co-extensive with the posterior columns. This continues in the dorsal region, but in the cervical the sclerosis begins to diminish externally, and shades off into the columns of Goll, in which it may be continued up into the restiform body. The actual primary lesion is therefore that in the external parts of the posterior columns, and the affection in the columns of Goll is really an ascending secondary degeneration, these columns being, as already stated, the principal seat of ascending sclerosis.

Besides extending to the columns of Goll, the disease commonly affects also the **Posterior roots**, which are often much atrophied. It may also extend to the **Posterior grey cornua**, in which case anæsthesia occurs, or to the **Lateral columns**, in which case paralysis ensues; or it may even pass through to the **Anterior cornua**, when muscular atrophy occurs in addition to paralysis. There is sometimes an extension to the **Medulla oblongata**, as evidenced by inco-ordination of the muscles of the eyeball and of those of speech.

In a large proportion of cases the **Peripheral nerves** are affected similarly to the posterior columns of the cord. There is atrophy of the nerve fibres with increase of the connective tissue. It is the sensory nerves which are affected, and chiefly the fine filaments in the skin and joints, but probably the sensory fibres of the muscles also. The lesion diminishes on passing from the finer filaments to the larger stems. There are frequently similar changes in the **Optic nerve** and in the **Ascending root of the fifth**.

**Relation of lesion to function.**—The lesion in the cord and nerves affects the sensory tracts, and yet the most prominent symptom is inco-ordination of motion. Violent pains are indeed commonly present at an early stage, and there is sometimes anæsthesia in later periods. The inco-ordination is not a motor paralysis, but due to interference with the centripetal nerve fibres, especially those which connect the muscles with the cord. The ganglion cells in the anterior cornua of the cord are, as we have seen, the immediate centres for the muscles, being centres of a low order dominated by higher ones. These centres receive centripetal fibres and give off centrifugal ones, thus forming a complete nervous system of a low order. The centripetal fibres come to a large extent from the muscles, and keep the centres informed, so to speak, of the position and state of contraction of the muscles. Any break in the circle will set the whole apparatus at a disadvantage and interfere with the completeness of the action. Here there is a break in the centripetal system, probably more particularly in the fibres coming from the muscles, as evidenced by the very early loss of tendon reflex (according



to Erb, whenever the ataxia is developed). In consequence the contractions of the muscles will be to some extent at random and without that accurate adjustment which exists normally. Probably also the normal tonicity of the muscles is, as Lockhart Clarke suggested, due to continuous stimulation of the ganglia in the cord by stimuli conveyed from the periphery. The interruption of the centripetal fibres will interfere with this stimulation, and the contraction of the muscles will start at a disadvantage. The absence of the knowledge of the state of the muscles which is implied in a break in the reflex circle may be partly compensated by using the eyes to direct the movements, the motor cells in the cord being, as it were, informed from above of the state of the muscles.

The occasional supervention of anæsthesia in locomotor ataxia is to be held as indicating an extension of the disease to the grey matter of the cord. The occurrence of true paralysis indicates that the neighbouring motor tracts in the lateral columns have become involved.

**Trophic lesions.**—In addition to the conditions already referred to, Charcot has called special attention to certain trophic lesions which occasionally occur in locomotor ataxia. These are of three kinds, namely, certain cutaneous eruptions, muscular atrophy, and affections of the joints.

The **Cutaneous eruptions** occur in the earlier periods, usually coinciding with the attacks of lightning pains, and they are in the form of local eruptions of herpes, lichen, pemphigus. As these occur in the earlier periods, and are coincident with the pains which are evidences of irritation of the centripetal fibres, we may infer that the trophic fibres of the skin are situated in the posterior roots, and that it is because these are involved in the process that the cutaneous eruptions occur.

**Muscular atrophy** is not very common in locomotor ataxia, and is of later occurrence. The muscles may waste as a direct result of paralysis—a slow atrophy from disuse. In some cases, however, there are localized atrophies of special muscles similar to those in progressive muscular atrophy and in bulbar paralysis. In these cases the disease has extended to the anterior cornua in which are situated the trophic centres of the muscles, and there is destruction of the ganglion cells, just as in progressive muscular atrophy.

The **Affections of the joints** consist in comparatively acute swellings, with exudation, followed by atrophy of the ends of the bones. Dislocations of the joints may occur in consequence. These trophic disturbances of the joints seem, like those of the muscles, to occur in cases where the disease extends to the anterior cornua, the trophic centres for



the whole organs of locomotion having their seat there. The propagation of the disease to the anterior cornua, leading to muscular atrophy or trophic affections of the joints, seems to be not uncommon, and may occur at a comparatively early period. This can hardly be regarded as remarkable when we consider that there are, for purposes of reflex action, direct connections between the posterior root-zones and the anterior cornua.

General paralysis of the insane is often complicated with ataxia, and the spinal cord presents sclerosis of the posterior columns similar to that in ordinary ataxia.

**Friedrich's hereditary ataxia.**—A considerable number of cases have been described in which motor inco-ordination has appeared at an early age in several members of the same family. There is, in these cases, probably a congenital faulty development of the cord, and the ataxia is the consequence rather of a degenerative than of an inflammatory process.

The lesion is not confined to the posterior columns but affects also the lateral columns, involving usually the pyramidal and the direct cerebellar tracts, and sometimes extending forward at the periphery of the cord. There seems little tendency to extension to the grey matter, and hence proper paralysis and muscular atrophy as well as anæsthesia are uncommon. There is seldom any evidence of irritation in the sensory fibres such as we have in ordinary ataxia, evidenced by the lightning pains.

The disease frequently extends upwards to the medulla oblongata, as evidenced by the occurrence of disturbance of speech and nystagmus.

**2. Spontaneous lateral sclerosis (*Erb's Spastic paralysis*).**—In this disease we have an independent disease of the pyramidal tract, having a similar anatomical distribution to that in descending grey degeneration. It usually begins in the lumbar cord, and attacks simultaneously both pyramidal tracts. These cases are of somewhat frequent occurrence, but seldom come to post-mortem examination unless there be an extension to other systems in the cord, especially the anterior cornua. Hence few cases have been observed in which pure lateral sclerosis has been proved by anatomical observation. In a case recorded by Dreschfeld, however, sclerosis was found in the lateral columns and very little in the anterior cornua.

The symptoms already referred to as characteristic of descending sclerosis are here very pronounced, namely, spasm of the muscles, chiefly of the lower extremity, with exaggerated tendon reflex. There is also, of course, paralysis.

The sclerosis appears to extend not infrequently from the pyramidal tract to the anterior cornua of the grey substance. In that case there is muscular atrophy in addition to the other phenomena. To conditions of this kind, Charcot has given the name **Amyotrophic lateral sclerosis**.



The name *Ataxic paraplegia* has been given to cases in which the lesions and symptoms of sclerosis of both the posterior and lateral columns have been present.

3. **Acute ascending paralysis** (*Landry's Paralysis*).—In this disease there is a rapidly extending paralysis, commencing in the legs, extending to the trunk and arms, and usually causing death in a few days, although not always fatal. Although the symptoms point to a serious affection of the cord involving the motor tracts, no lesion has been yet discovered. The acuteness of the symptoms and the occurrence of enlargement of the spleen and lymphatic glands have suggested the existence of a morbid poison (Gowers), and the disease has been looked upon as perhaps analogous to diphtheritic paralysis.

4. **Poliomyelitis anterior acuta** (*Infantile Paralysis, Acute Atrophic Spinal Paralysis*).—This is a disease mostly of infancy and early childhood, but not unknown in the adult.

The disease is an acute inflammation of the grey substance of the anterior cornua of the cord. In an observation by Drummond in which a child died after a few hours' illness, there was redness of the anterior cornua, with minute extravasations, swelling of the ganglion cells, etc. In later cases, but still comparatively early, the anterior cornua are seen under the microscope to be altered, not continuously, but in patches. They contain numerous round cells and compound granular corpuscles. The ganglion cells in certain of the groups have disappeared or have shrunk considerably. At the periphery of the affected patches round cells are aggregated, and there is already some shrinking of the patches. The condition is most manifest in the lumbar and cervical enlargements. The anterior roots are also somewhat atrophied, and show evidences of degenerative changes.

Many cases have been examined years after the onset of the disease, at intervals varying from seventeen to sixty-one years, and the changes have been very obvious even to the naked eye. These consist of sclerosis with shrinking, mainly of the anterior grey cornua, but also of the anterior and antero-lateral columns of white substance (see Fig. 212). These changes are very manifest when the cord has been hardened and fine transverse sections made. The shrinking of the horn affects certain of the groups of cells specially, and the shrunken part consists largely of connective tissue in which there are visible no ganglion cells or only deformed and pigmented ones. There is connective tissue with abundant nuclei and enormous numbers of amyloid bodies. The lesion in the anterior cornua is by no means homogeneous or symmetrical. One cornu may be atrophied and the other normal, and on examining sections at different levels there is great variety in the longitudinal distribution. The anterior roots are also atrophied at the parts corre-



sponding to the most affected portions of the cord, and they may appear to the naked eye small and grey.

The disease usually begins acutely and the paralysis generally assumes its full development almost at once. This extensive paralysis seems due in part to pressure by the distended vessels and inflammatory exudation, and in some cases by extravasated blood. In this period some ganglion cells may be destroyed, but those not absolutely destroyed may recover as

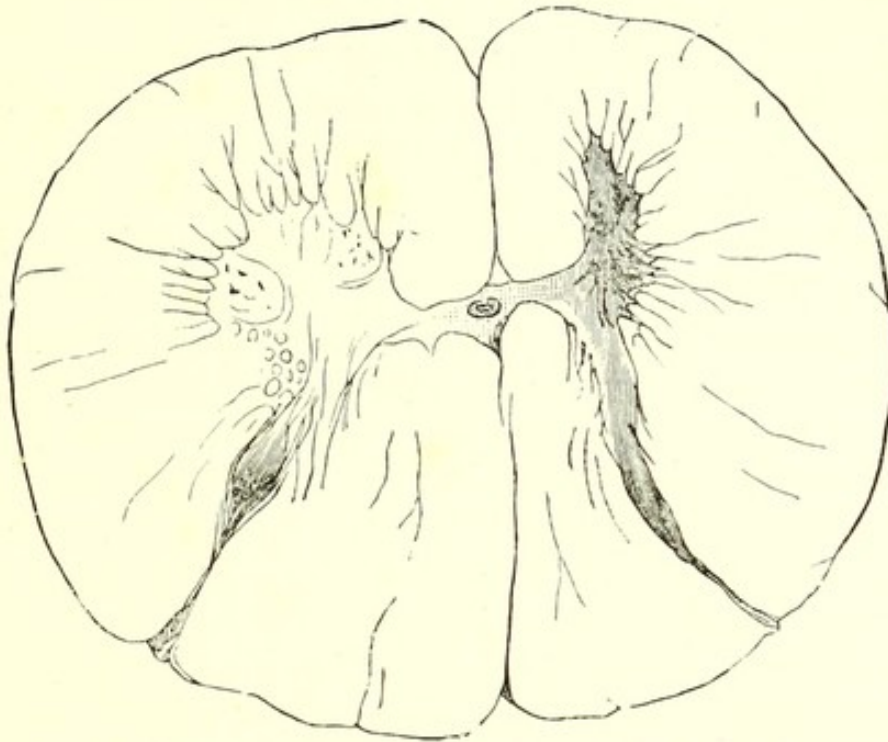


Fig. 212.—Anterior poliomyelitis. The right anterior cornu in the cervical region is shrunken, and there is atrophy of all the white columns on that side. From a woman, aged 50, who was the subject of infantile paralysis of the right arm. (CHARCOT.)

the acute inflammation passes off. In many parts the inflammation is slight and subsides completely, in others it is severe and goes on to sclerosis, and there is permanent destruction of the ganglion cells. Hence it is that a paralysis which has been at first almost universal may be recovered from almost completely. The improvement goes on till all the cells which are capable of it have recovered, and there is a residue which have been permanently lost and cannot be restored. These permanently lost cells represent single muscles and groups of muscles, and there is a corresponding localized paralysis.

**Trophic changes in muscles, etc.**—The centres which have the direct control of the contractions of the individual muscles are probably the centres which command the nutrition of the muscles and their nerves; or the trophic centres, if separate, must at least lie alongside the muscular centres. For the permanently paralysed muscles soon undergo a marked and rapid atrophy and the fibres of the anterior roots also de-



generate. The muscular atrophy is not simply from disuse, for it is much greater and more rapid than in cases of paralysis where the anterior cornua are not affected. The muscles rapidly get soft and emaciated, and may entirely disappear. Besides the loss of substance there may be proliferation of the nuclei of the interstitial connective tissue, and an increase in this tissue; sometimes there is infiltration of fat in the interstitial tissue, so that the muscle may appear less atrophied than it really is. The defective development of the bones and joints met with may be partly from disuse, but also probably to some extent from destruction of trophic centres.

At the onset there are usually signs of acute inflammation (fever, sometimes convulsions), but these symptoms may be very slight, and the first thing noticed may be the paralysis. Whether accompanied by fever or not, the paralysis develops rapidly, and generally attains its full extent almost at once. It may affect both arms and both legs, or the legs alone, or the arms alone, or the leg and arm on one side, or one leg or one arm alone. After some weeks, the paralysis begins to improve, and may go on improving for some time. Complete recovery rarely results, and generally the paralysis affects certain muscles permanently. The whole muscles of a limb may remain paralysed, or the paralysis may be limited to a group or two of muscles. In the permanently paralysed muscles a rapid atrophy occurs which becomes very extreme. The bones of the paralysed part do not grow normally, and the articulations are imperfectly adjusted. Certain deformities ensue, the commonest and most prominent of which is club-foot. Curvature of the spine is also a comparatively frequent result. All through the disease the patient may maintain good general health, and after recovery from the initial fever he may present nothing abnormal but the paralysis and atrophy. Through time he learns to use his remaining muscles to the best advantage, and may pass through a long life maimed by the infantile attack.

Although most common in children a good many cases of this disease have now been recorded in **Adults**. Here also it is ushered in by acute symptoms such as fever, pain in back and extremities, vomiting, headache. The paralysis develops mostly in a few hours, but it may be as late as a day or two. After a time recovery begins, and is more frequently complete than in the case of children. It is frequently incomplete, and then we have paralytic deformities, which, however, as the bones are fully formed are not so striking as in the case of children. Here, also, there is rapid atrophy of the muscles, which is not so liable to be concealed by fatty infiltration as in the case of children.

5. **Poliomyelitis anterior chronica**.—Some cases have been recorded, chiefly in adults, in which, with little or no fever or disturbance of the general health, a motor paralysis has occurred, and in the course of a few days or weeks has extended to the entire lower limbs and then rapidly to the upper. It rarely takes the opposite course. The muscles become slack and soft and lose their reflex irritability, and they rapidly atrophy. There have been few post-mortem examinations as yet, but



they seem to show a chronic inflammation or sclerosis of the anterior cornua with loss of the ganglion cells. The disease is frequently recovered from, but recovery is slow and complete restitution may take years. On the other hand, the disease may extend upwards and produce death. In this disease, also, it will be observed, centres for contraction and trophic centres seem to be simultaneously affected. Many cases of **Multiple neuritis** have been mistakenly diagnosed as belonging to this class.

**6. Progressive muscular atrophy.**—This disease, although its name points to an affection of the muscles, is yet considered here, because there is every reason to believe that an affection of the spinal cord is the primary lesion, the muscular condition being secondary.

In its clinical aspects the main feature is a gradually progressive atrophy and consequent paralysis of the muscles. It very commonly begins in the muscles of the hand, but progresses from one muscle to another till the death of the patient. In its later stages it frequently becomes associated with a corresponding form of disease of the medulla oblongata, namely, bulbar paralysis, to be considered next.

In the muscles the change consists in what may be regarded as a chronic inflammation of them. The muscular nuclei increase in number, and as the contractile substance diminishes, the sarcolemma may come to be filled with cells, the result of this proliferation of the muscular nuclei. At the same time the interstitial connective tissue shows active changes, increase of nuclei, and newformation of connective tissue. In the muscular substance various forms of degeneration have been observed, chiefly fatty and waxy, or simple atrophy. In any case the muscular fibres are lost by degrees and the connective tissue increased but not sufficiently to make up the bulk of the lost muscular substance. Sometimes a fatty infiltration of the connective tissue occurs, so that adipose tissue comes to occupy the place of the muscle to a large extent. This change, if it occurs, is only local, and pure atrophy may exist side by side with atrophy with formation of adipose tissue. This formation of adipose tissue is mostly a late, and by no means a characteristic, occurrence in this disease.

In the Spinal cord the essential changes are in the anterior cornua, but in most cases the pyramidal tracts are also involved. The lesion in the anterior cornua consists in a gradual atrophy of the large ganglion cells, affecting the parts corresponding with the atrophied muscles. These cells may have entirely disappeared or there may remain only small angular bodies representing them. There is at the same time an increase of the connective tissue so that there is little change in the size or shape of the cornu.



According to Ross, the central grey column (the grey substance on either side of the central canal) is most affected, being traversed by large canals and fibrillated.

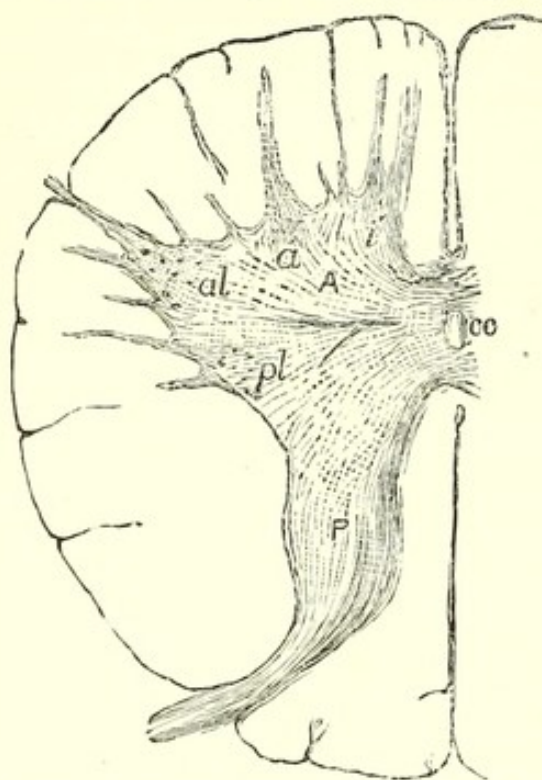


Fig. 213.—Progressive muscular atrophy—section of cord in cervical region, from an advanced case. The central portion of grey substance is fibrillated; the median group of cells has disappeared, and the other groups are atrophied. (Ross.)

In the annexed figure (Fig. 213) it is seen that the cells of the median group have also entirely disappeared, while the other groups of the anterior cornua are limited in size, the peripheral cells of the groups having disappeared, leaving only the more central ones. It very commonly happens that, short of absolute destruction of the ganglion cells, they are greatly atrophied and pigmented (*pigmentary atrophy*).

**The Pyramidal tracts**, both those in the lateral and those in the anterior columns are affected probably in all cases (Gowers), and the lesion has been traced up through medulla oblongata, pons, peduncles, internal capsule and corona radiata to the motor convolutions of the brain where the large ganglion cells have been observed to be atrophied.

**The Anterior nerve-roots** are also atrophied. The nerve fibres may

have almost entirely disappeared, and the nerves may be almost replaced by connective tissue. The degeneration exists throughout the peripheral distribution of these nerves, and the terminal fibres in the atrophied muscles are greatly altered.

Leyden has recorded a case of progressive muscular atrophy in which there was the peculiar condition of the cord designated *Syringomyelia*. (See further on.)

**Relation of lesion to function.**—In this disease we have a degeneration or chronic inflammation affecting the motor system, chiefly in its lower parts. The disease centres in the ganglion cells of the anterior cornua, but it affects the nerve fibres arising from these, which are really prolongations of their processes, and in most cases also the pyramidal tracts above them. The degeneration of the pyramidal tracts is not to be regarded as in any sense secondary, but as part of the prolonged lesion of the motor system. Indeed the order in which the different parts are involved varies somewhat, and we may have the pyramidal tracts, at least in some parts of their course, affected in advance of the anterior cornua, although this is not common. We have already seen that spasm of the muscles is the characteristic result of degeneration of the pyramidal tracts, but this can only occur



when the anterior cornua are unaffected. Hence in this disease rigidity of the muscles is not common, although in some cases it does occur locally, especially in the legs. Charcot has endeavoured to group this class of cases separately on the supposition that in them the lesion has begun in the pyramidal tracts, but it is doubtful whether this is a sufficient basis of distinction.

It will appear from the above that, as in most cases the anterior cornua are affected before or simultaneously with the pyramidal tracts, there will usually be no evidence during life of the affection of the latter.

**7. Bulbar paralysis, or Glosso-labio-laryngeal paralysis.**—This condition is called bulbar paralysis from the fact that the part affected is the medulla oblongata, which is frequently designated the bulb. There is progressive atrophy and paralysis of muscles supplied from the medulla oblongata. The muscles are mainly those of the tongue, lips, arches of palate, pharynx and larynx, and in consequence there is progressive interference with articulation, chewing, swallowing, and even with the production of the voice. The disease may pass on later to affect the more vital functions of the medulla oblongata, those concerned in respiration, etc.

We have already seen in connection with the normal structure of the medulla oblongata that, on passing from the cord, the grey substance is dislocated backwards, and that in the posterior region of the medulla a set of grey nuclei appear, which, as the spinal canal opens up in the fourth ventricle, present themselves in the floor of that ventricle. These nuclei are mainly motor, corresponding with the anterior cornua, and they form the immediate centres for certain cerebral nerves (see Fig. 214). In the lower half of the fourth ventricle we find the hypoglossal nucleus (xii') occupying the part next the middle line. Outside it there is the spinal accessory nucleus which begins in the cord, and does not extend far up in the floor of the ventricle; it is not shown in the figure, but gives place to the nucleus of the pneumogastric or vagus (x'). Outside the vagus appears the glosso-pharyngeal, which partially divides the vagus nucleus into two. As we pass upwards the vagus nucleus gets smaller and the hypoglossal and glosso-pharyngeal approximate to each other. Above that again come in the nucleus of the sixth (the abducens) in the middle line, and outside that

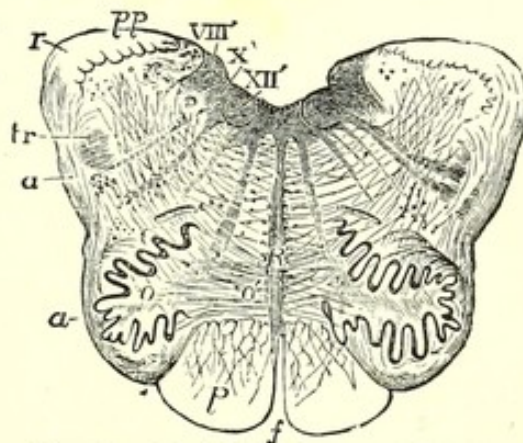


Fig. 214.—Medulla oblongata at the level of the fourth ventricle, showing the position of the grey nuclei. (QUAIN.)



the motor nucleus of the fifth and that of the facial. Outside these again are sensory nuclei, those of the acoustic and of the fifth.

In bulbar paralysis there is atrophy of these grey nuclei. It is seldom that opportunity is afforded of examining the medulla in recent stages, but according to Benedikt, who had such an opportunity, there are definite signs of inflammation, hyperæmia, thickening of the walls of the vessels, and numerous round cells. In later periods increase of the connective tissue, with round cells and amyloid bodies, have been observed. These changes centre in the nucleus of the hypoglossal, and this nucleus is usually most seriously damaged, but they pass soon to the accessory and vagus, while the glosso-pharyngeal sometimes, but not usually, escapes. The nucleus of the facial is often attacked, and sometimes that of the motor branch of the fifth. The disease seldom extends to the abducens, and never attacks the sensory nuclei of the acoustic and fifth nerves.

There is in many cases degeneration of the nerve fibres in the anterior pyramids similar to that in the pyramidal tracts in progressive muscular atrophy. There is even reason to believe that in some cases the lesion may be essentially in the pyramidal fibres which connect with the nuclei in the medulla oblongata, in which case the disease would be analogous with lateral sclerosis of the cord rather than with progressive muscular atrophy.

The corresponding nerves usually show considerable atrophy, especially the hypoglossal, and next to it the accessory, vagus, and glosso-pharyngeal. In them may be found a fatty degeneration involving destruction of the medullary sheath and subsequent overgrowth of connective tissue. In this way the nerve-root may come to be almost nothing but a connective-tissue strand.

In the affected muscles the change is exactly parallel to that in progressive muscular atrophy with which sooner or later this disease is so frequently associated. There is increase of the muscular nuclei and connective tissue, with destruction of the proper contractile substance, and consequent atrophy of the muscle as a whole. Thus the tongue, palatine arches, lips, pharynx, and larynx may have their muscles intensely atrophied. Sometimes also the muscles of the neck, especially the trapezius (supplied by the spinal accessory), are affected. In some cases the atrophy is obscured, as in progressive muscular atrophy, by the interstitial tissue becoming adipose.

**8. Pseudo-hypertrophic paralysis.**—The resemblance of this disease to progressive muscular atrophy strongly suggests that it is primarily due to a lesion in the spinal cord. But repeated examinations of the cord have shown no constant lesion there, and in most cases the anterior



cornua have been found intact. The muscular nerves are also apparently unaffected, and the general opinion is now that it is a primary disease of the muscular system. It occurs mostly in childhood, and is frequently met with in several members of the same family. It may be regarded as having its origin in a faulty development of the muscular system. Considering the close relationship in function and even in nutrition between the nervous system and the muscles, as shown by the affections already described, it is perhaps justifiable to consider this disease along with those of the spinal cord.

In many forms of disease in which the muscles are disused or paralysed there is an overgrowth of the interstitial connective tissue sometimes with excessive development of fat. We have noted this in some of the affections considered above, and it occurs also when the muscles are disused by reason of fixation of joints, etc. (See under Fatty Infiltration, p. 159.)

In the present disease there is an apparently spontaneous atrophy of the muscular substance with a newformation of connective and adipose tissue between the muscular fibres. The newformation, at least in some situations, usually more than compensates in bulk for the atrophy,

and there is an apparent hypertrophy of the muscle as a whole. By the time the patient dies there is generally a very extraordinary and widely distributed affection of the muscular system. Many of the muscles on being cut into look like pieces of fat, in which to the naked eye no trace of muscular tissue is visible. Under the microscope occasional narrowed muscular fibres

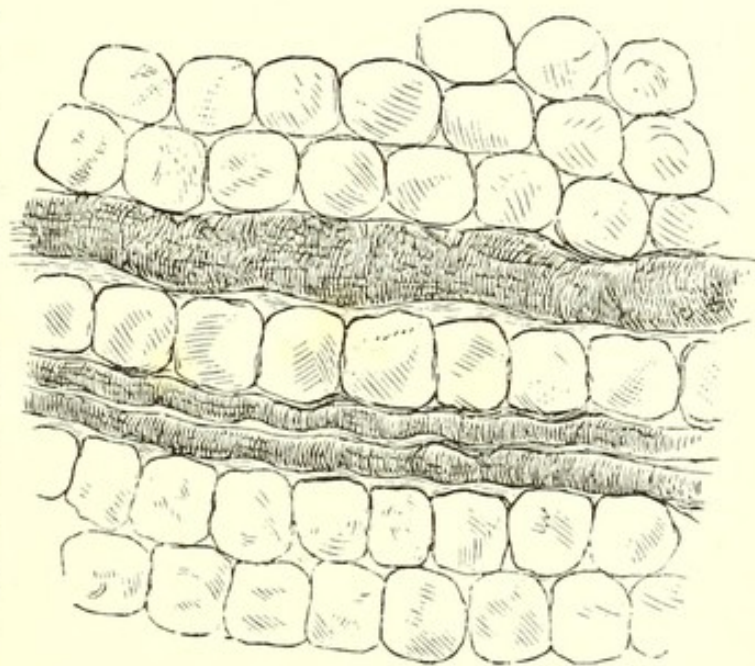


Fig. 215. —Muscle in pseudo-hypertrophic paralysis. There is chiefly adipose tissue with a few atrophied muscular fibres.  $\times 180$ .

may be visible in the midst of the adipose tissue (see Fig. 215), usually accompanied by strands of connective tissue. The muscular fibres, although varying in size and in general greatly narrowed, for the most part preserve their transverse striation. In some situations the fat may have, to a large extent, disappeared, so that fibrous tissue replaces the muscle.



An **idiopathic muscular atrophy** has been described in which the wasting of the muscles has not been associated with increase of connective tissue or fat. This condition shows many analogies with that described above, occurring like it in several members of the same family, but not generally beginning in childhood.

### III.—TUMOURS OF THE CORD AND MEDULLA.

Tumours of the cord and medulla proper are very rare, but it is not uncommon to find tumours of the meninges, and even of the bones, affecting the cord by pressure.

**Glioma** of the cord is less common than that of the brain. It has usually a very small transverse extent, but may occupy a considerable length of the cord, which is correspondingly swollen. The boundaries of the tumour are frequently rather ill-defined. **Myxoma** and **Sarcoma** have also been observed, usually glio-sarcoma, but in one case a proper spindle-celled sarcoma. A case has been recorded by Ganguillet of a so-called **Cylindroma**, in which there were branching bodies whose ramifications presented gelatinous mantles.

The **Solitary scrofulous tubercle** is much less common in the cord than in the brain. It forms a hard cheesy mass, usually with a softened zone of tissue around it. The larger have usually their seat in the lumbar portion of the cord.

**Syphilis** we have seen to be apparently a frequent cause of locomotor ataxia. It seldom gives rise to proper gummata. When it does so the tumour, here as in the brain, takes origin in the meninges, and is hence at first superficial, although it may subsequently extend into the substance of the cord. The membranes over it are thickened and usually adherent.

**Cysts** do not occur in the cord as independent formations, but a cystic condition will be described further on as **Hydromyelus** and **Syringomyelia**.

All forms of tumour, whether of the cord or meninges, are liable to interrupt the conduction of the cord, and so to produce paralysis. There will be also the usual ascending and descending secondary degenerations.

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## SECTION III.—CONTINUED

## C.—THE ENCEPHALON

- C. THE ENCEPHALON.—ANATOMICAL INTRODUCTION, *arrangement of fibres and centres. Functions of convolutions. Arteries of brain.* I. MALFORMATIONS. 1. *Congenital smallness, microcephalus*; 2. *hypertrophy*; 3. *heterotopia*. II. INJURIES; *Laceration of brain*. III. LESIONS AFFECTING THE CIRCULATION. 1. *Hyperæmia, anæmia, and œdema*. 2. *Occlusion of arteries; by embolism or thrombosis; effects, including Softening of the brain substance; occlusion in chorea*; 3. *Thrombosis of the cerebral sinuses*. IV. CEREBRAL HÆMORRHAGE. 1. *From large arteries, chiefly due to aneurysm, sometimes to atheroma; influence of increase of blood-pressure*. 2. *From nutrient arteries, may be from miliary aneurysms or atheroma*. 3. *From capillaries, chiefly complicating other forms. Appearances in hæmorrhage, and results; the apoplectic cicatrix and cyst*. V. INFLAMMATION. 1. *Acute localized encephalitis, chiefly abscess of the brain*. 2. *Chronic localized encephalitis*. 3. *Diffuse encephalitis, (a) in fevers, (b) in dementia paralytica, (c) in hydrophobia, (d) in tetanus*. VI. ATROPHY AND DEGENERATIONS. *Senile atrophy and other forms; secondary degenerations*. VII. TUBERCULOSIS AND SYPHILIS. 1. *The tubercular tumour*. 2. *Syphilis of brain and membranes; disease of arteries may lead to softening*. VIII. TUMOURS AND PARASITES, *chiefly glioma and sarcoma*.

**Anatomical Introduction.**—In the encephalon the course of the fibres is even more important than in the spinal cord and medulla, as they are liable to be interrupted in a more isolated fashion. It is of great consequence that the pathologist should have an acquaintance with the general relations of parts so as to identify the position of lesions. In this section nothing more is attempted than to indicate these general relations, the more intricate particulars being left to the special works on anatomy. In the actual work of post-mortem examination, it is important to note the exact locality of lesions on the spot. To assist in this, tracings may be made of the figures which accompany this section, or of similar ones, the position of the lesions being entered in shading.

In the medulla oblongata, as we have seen, motor fibres, which have been in the lateral columns of the cord, come forward and form the anterior pyramids, in which they decussate. The sensory fibres are now behind, and they remain posterior to the motor in all the succeeding parts. In the *Pons varolii* (Fig. 216) the motor fibres are in front, forming bundles, as shown in lower half of figure, but overlaid by the transverse fibres from the cerebellum. Besides these fibres, which are mainly in the anterior half of the pons, there are grey nuclei (shown in figure), which continue up the series which lie in the medulla oblongata in the floor of the fourth ventricle. In



In the pons the principal nuclei are those of the sixth and facial, the motor and sensory roots of the fifth, and one of the nuclei of the auditory nerve. It is to be remembered also that the fibres of these nerves in part at least traverse the pons, and are liable to be involved by a lesion, even when the nuclei are not reached.

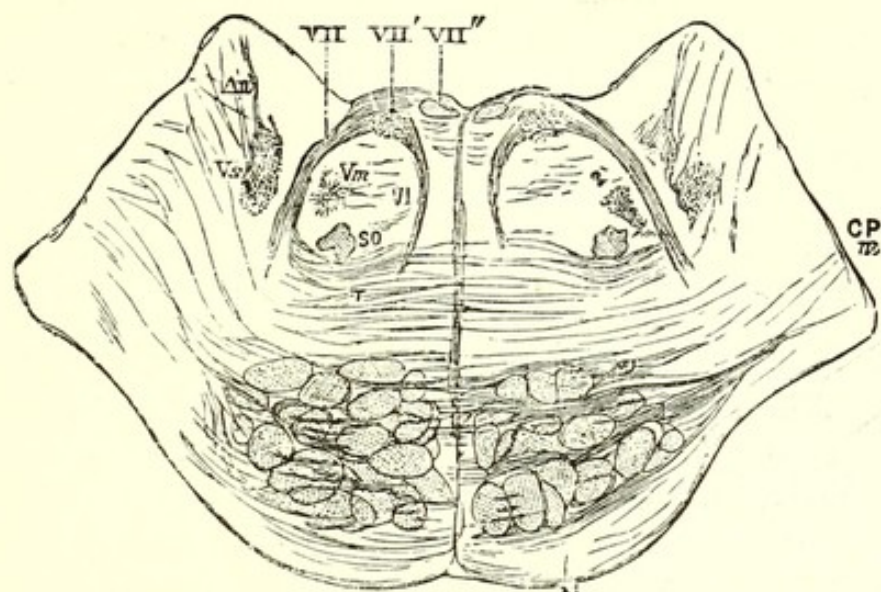


Fig. 216.—Pons varolii. In lower half the transverse section of nerve bundles continued upwards from the cord is seen, the pyramidal tract being in front (or lower in figure). The Roman numerals and letters indicate the nuclei of cerebral nerves. (QUAIN.)

In the *Crura cerebri*, or cerebral peduncles, the motor fibres are in front and internal, forming the greater part of the crura. Sensory fibres occupy about the external fourth of the crura. Behind the crura is the locus niger, behind which again is the tegmentum, containing a rather complex mass of fibres and grey matter.

Above the crura the fibres are continued upwards, the motor still anterior and the sensory posterior in the mass of white substance called the **Internal capsule**. It is here necessary to be somewhat more minute in the description of the relation of the structures, as these relations are of great importance. When one of the lateral ventricles of the brain is opened, certain masses of grey nervous tissue are seen in its floor. In front there is a long brown prominence, rounded anteriorly, and tailed behind. This is the **Nucleus caudatus**, which is often designated the corpus striatum, although really only one piece of it. Behind the nucleus caudatus is the **Optic thalamus** which is more bulky and rounded. It is to be remembered that in opening the lateral ventricle almost no nerve fibres need to be cut except the commissural ones of the corpus callosum. The great masses of nerve fibres passing upwards are as it were pushed outwards by the lateral ventricle, and we have to cut into its floor in order to reach them.

From the accompanying figure, which represents a horizontal section of the brain just below the floor of the lateral ventricle, the relations of parts may be gathered (Fig. 217). Beneath and outside the nucleus caudatus (*NC*), and continuous above and externally with the great central white substance of the hemispheres, the corona radiata, there is a mass of white substance. This is the **Internal capsule** in which may be distinguished an anterior division *IK'*, a posterior division *IK*, and a middle part, the knee *K*. To the outside of and beneath the internal capsule lies a mass of grey substance, the **Nucleus lenticularis** (*LN*), which on section has a triangular shape with the base turned outwards, and is seen to be divided into three pieces (shown on right side). It is generally regarded as a part



of the corpus striatum and motor in function. The nucleus lenticularis extends a considerable distance from before backwards, and in its posterior parts the internal capsule (*IK'*) lies between it and the optic thalamus, which has now largely taken the place of the nucleus caudatus. Outside the nucleus lenticularis there is a narrow band of white substance (*EK*), the **External capsule**, the capsules being named from their relation to the nucleus lenticularis. Outside the external capsule again and close to the convolutions (here the Island of Reil) there is a narrow band of grey sub-

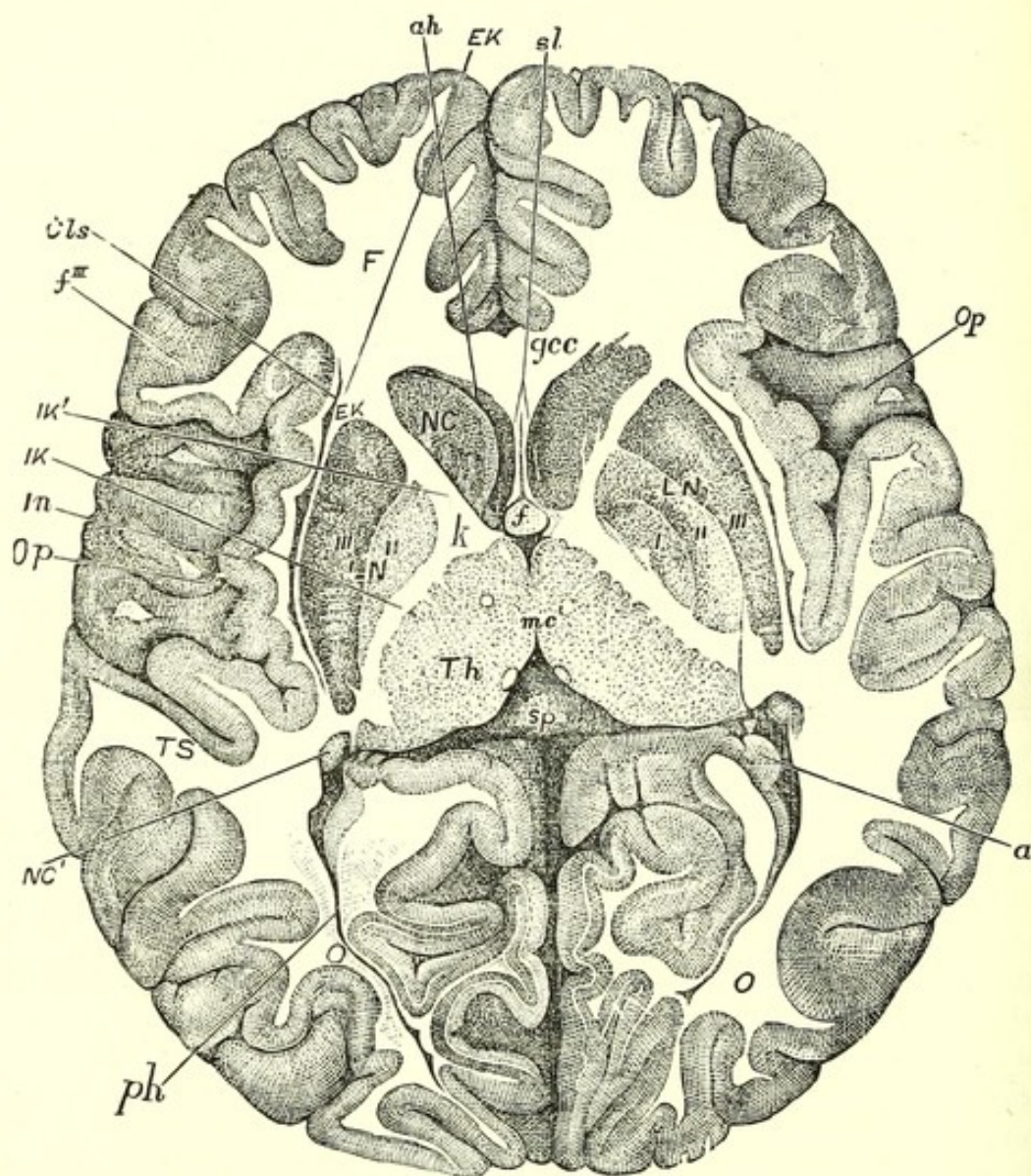


Fig. 217.—Horizontal section of brain of a child nine months old, the right side at a somewhat lower level than the left. *F*, frontal; *TS*, temporo-sphenoidal; and *O*, occipital lobes; *Op*, operculum; *In*, island of Reil; *Cls*, claustrum; *f'''*, third frontal convolution; *Th*, thalamus; *NC*, nucleus caudatus; *NC'*, tail of same; *LN*, nucleus lenticularis; *I*, *II*, *III*, its first, second, and third divisions; *IK*, internal capsule, posterior division; *IK'*, anterior division, and *k*, knee; *ah* and *ph*, anterior and posterior horns of left lateral ventricle; *gce*, knee of corpus callosum; *sp*, splenium; *mc*, middle commissure; *f*, fornix; *sl*, septum lucidum; *a*, cornu Ammonis. (Ross from FLECHSIG.)

stance (*Cls*), which is fancifully compared to a tape-worm, and is called the **Nucleus tæniæformis** or **Clastrum**. The anterior parts of the internal capsule contain motor fibres, and the posterior sensory; it is generally said that the anterior two thirds are motor, and the posterior third sensory.

It is to be remembered that the three nuclei we have referred to, the nucleus caudatus, nucleus lenticularis, and thalamus opticus, receive fibres from and give







Of the convolutions the easiest to determine are those bounding the fissure of Rolando, one of which belongs to the frontal and the other to the parietal lobe; these are called the **Ascending frontal** and **Ascending parietal convolutions** respectively. In front of the ascending frontal, the frontal lobe presents three layers of convolutions (*a*, *a'*, and *a''*) which lie transversely and pass by their posterior extremities into the ascending frontal. These **Transverse frontal convolutions** are distinguished as the first, second, third, or superior, middle, and inferior. It is not to be supposed that these are single simple convolutions, they are rather layers or strata of convolutions. The inferior (*a''*) is a very important one, and it can generally be easily recognized as it curves round the short ascending branch of the fissure of Sylvius (*s*). After curving round this branch it becomes continuous with the lower end of the ascending frontal. At this point the two convolutions form a somewhat triangular piece, and as this lies over and partially covers the island of Reil, it is often called the **Operculum** (in the position of *x A* in Fig. 218). In addition to these we have still in the frontal lobe the **Supraorbital convolutions** (*a\**) which have no arrangement that needs to be detailed.

In the parietal lobe the **Ascending parietal convolution** (*B*, *B*) is already known. Another easily recognized one is that which lies immediately above the fissure of Sylvius and is called the **Supramarginal convolution**. The posterior portion of this convolution curves round the upper end of the fissure of Sylvius, and as it turns thus round an angle it is often called the *angular gyrus* (*b''*, *b'''*), and is important, as Ferrier has supposed it to be the seat of the sense of sight. The rest of the parietal lobe is divided into an upper and a lower portion by a longitudinally placed fissure which is often not very distinct, the intra-parietal fissure, which arises close to the fissure of Sylvius behind the fissure of Rolando to which it at first lies parallel, and then passes backwards. The lobe is thus divided into the **Superior** and **Inferior parietal lobules**, the former being continuous in front with the ascending parietal convolution.

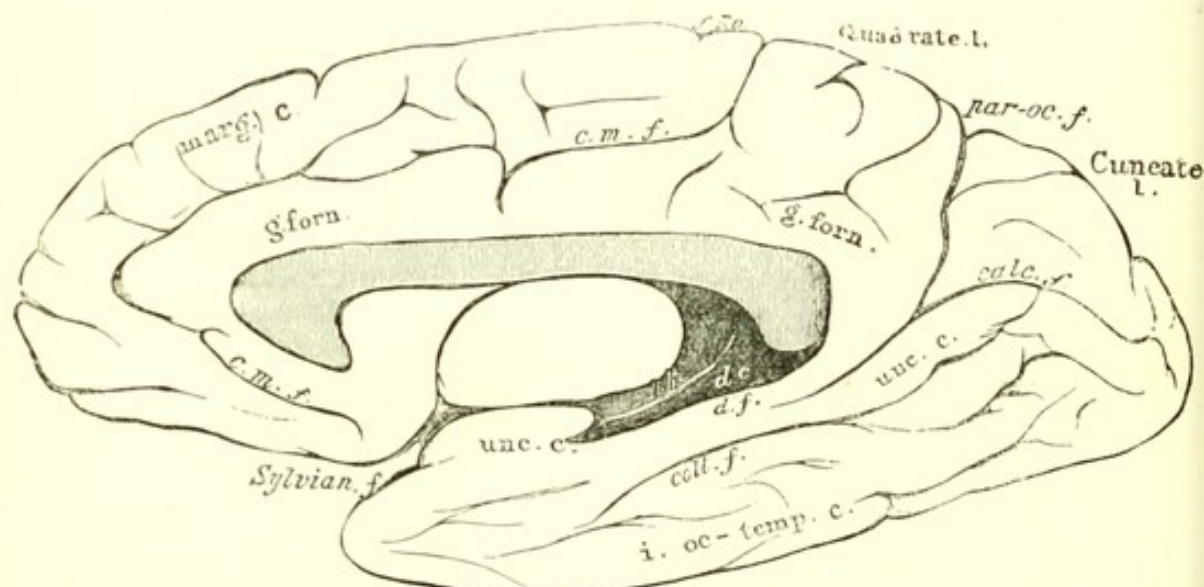


Fig. 219.—Internal aspect of right cerebral hemisphere. Explanation in text. (QUAIN.)

In the **Occipital lobe**, looking at the lateral aspect, **Three transverse layers** of convolutions can be distinguished. These are named as in the frontal lobe, first, second and third, or superior, middle and inferior (*d*, *d'*, *d''*). On the under or basal surface there are two further layers which are continuous in front with those of the temporo-sphenoidal lobe and are named in common with them **inferior occipito-temporal convolutions**.



The **temporo-sphenoidal lobe** presents on its lateral surface again **Three transverse convolutions**, superior, middle, and inferior (*c*, *c'*, *c''*); the superior, bounding the fissure of Sylvius and also called **Infra-marginal**, is continuous with the angular gyrus. The remaining two sets on the basal surface have been already mentioned as forming, with those of the occipital lobe, the inferior occipito-temporal convolutions.

On examining the **Internal aspect** of the cerebral hemisphere (Fig. 219) certain convolutions are to be distinguished. The superior frontal convolution and the ascending frontal and parietal convolutions are here partly visible. More distinctly on the internal surface we distinguish in front the **Marginal convolution** (*marg. c.*) which is continuous with the superior frontal and lies along the superior longitudinal fissure. Immediately behind this convolution we come to the superior parietal lobule which, on its mesial surface, is called the **Præcuneus** or **Quadrangle lobe** (see figure). The parieto-occipital fissure is here very marked, and it is joined at an angle by the calcarine fissure (*calc. f.*) in such a way as to demarcate a triangular surface, the **Cuneus** (*cuneate l.*). On this aspect also appear the inferior occipito-temporal convolutions (*i. oc.-temp. c.*). Within this external ring of convolutions we have now a deeper layer. Immediately bordering on the corpus callosum and following the fornix we trace from before backwards the **Gyrus fornicatus**. Having skirted the corpus callosum from before backwards it turns round at its posterior extremity, and passes downwards into the **Gyrus hippocampi**. This gyrus is also continuous with the gyrus cuneus and the median occipito-temporal. The gyrus hippocampi passes forward towards the anterior extremity of the temporo-sphenoidal lobe where it terminates in a hook-like curve, the **Gyrus uncinatus** (*unc. c.*).

In the cornu Ammonis of the lateral ventricle there lies a convolution whose surface is turned inwards and forms an elongated rounded projection in the cornu; this is the **Gyrus dentatus** (*d. c.*).

**Functions of the convolutions.**—The localization of function in the cortex of the brain is of so much practical importance that it is well, when opportunity offers, to familiarize the mind with the topography of the convolutions in relation to function.

**The cerebellum.**—There are only a few points which require notice here as to the general arrangements of the parts in the cerebellum. Like the cerebrum it is divisible into two lateral hemispheres, the right and left lobes. These are united by a central piece, which is most marked on the under surface, called the vermiform process. The cerebellum is divided by many fissures which run horizontally and leave narrow convolutions called the **Folia**. One of these fissures, deeper than the rest, and called the great horizontal fissure, divides the cerebellum into an upper and a lower portion. In its internal structure it presents white matter which runs outwards from the peduncles diverging towards the folia and forming a tree-like expansion, the **Arbor vitæ**. In the midst of the white substance in each hemisphere there is a small grey nucleus, not unlike the olivary body, called the **Corpus dentatum**.

**Arteries of the brain.**—With a view to the identification of the numerous lesions of the arteries in the brain, it will be proper here to refer briefly to the distribution of these vessels. The **Circle of Willis** gives off at the base three main arteries to the brain, the posterior, middle, and anterior cerebral. The **Posterior cerebral artery**, besides giving certain central branches to be afterwards referred to, is distributed on the surface of the brain, supplying the greater part of the occipital and temporo-sphenoidal lobes with the exception of the upper temporo-sphenoidal convolution. The **Middle cerebral artery**, or the artery of the fissure of Sylvius, is of great importance as being much more frequently the seat of lesion than the others. Besides its central branches, afterwards considered, it supplies the middle district of the brain



all round the fissure of Sylvius, including the parietal lobe, the posterior parts of the frontal lobe, and the superior convolution of the temporo-sphenoidal lobe. The **Anterior cerebral artery** is distributed to the anterior parts of the frontal lobe.

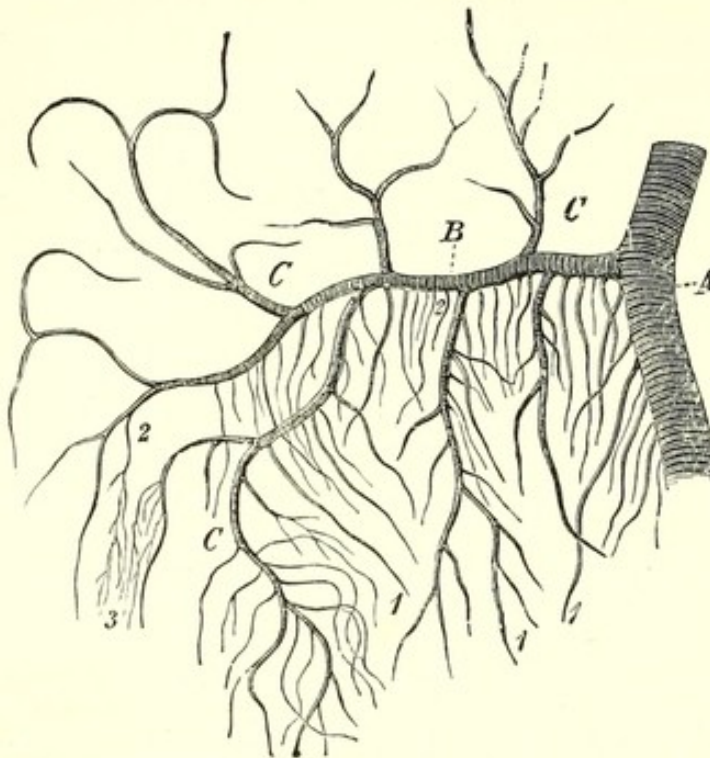


Fig. 220.—Arteries of the brain showing cortical distribution. *A*, a tertiary branch of main artery. *B*, primary twig. *C, C, C*, secondary twigs. The nutrient arteries are seen to pass off from all these branches as fine hair-like offsets. (Ross after DURET.)

These arteries, with the exception of their central branches, run in the sulci of the convolutions, and divide into successive orders of branches which lie in the pia mater. The larger branches anastomose sparsely, while the finer twigs are end-arteries (Duret). All the arteries hitherto considered run on the surface of the brain in the soft membranes. The actual nutrient arteries are branches of these, and penetrate from them into the substance of the brain. We may thus distinguish the larger arter-

ies of the surface and the smaller or nutrient arteries.

The **Nutrient arteries** again are divisible into two groups, which may be designated the cortical and central systems.

The **Nutrient arteries of the cortex** pass off not merely from the finer twigs of the larger arteries, but also from the larger branches (see Fig. 220). They are fine thread-like vessels which pass down perpendicularly into the brain substance. Some of them are short and supply the grey substance of the convolutions, others are longer and reach the white substance, their territory here extending to the boundaries of that of the central arteries. In stripping the membranes from the surface of the brain these nutrient arteries are seen like fine hairs emerging from the brain substance. If a portion of the soft membranes be now floated in water these fine vessels will be seen like bristles passing from all the branches. All the nutrient arteries are end-arteries.

The **Central nutrient arteries** have a somewhat different arrangement from those of the cortex. They are long vessels of larger calibre than the cortical ones, and they pass off from the main arteries very near their origin. The most important are those which come off from the middle cerebral just outside its origin and pass through the anterior perforated space to the basal ganglia. Of these, three sets of branches have been distinguished: (1) the *lenticular branches*, short twigs lying internally which pass to the internal parts of the lenticular nucleus; (2) the *lenticulo-striate* branch, a longer and larger vessel which lies outside the first and supplies the outer parts of the lenticular nucleus and adjoining parts of the internal capsule and caudate nucleus; this branch is most frequently the seat of hæmorrhage; (3) the *lenticulo-optic* branch is posterior to the former and supplies the posterior part of the lenticular nucleus and the anterior part of the optic thalamus. The remain-



ing central branches are supplied by the anterior and posterior cerebral arteries. The former sends branches which supply the anterior part of the caudate nucleus, and the latter sends two sets, an internal and an external, to the optic thalamus. All these central arteries are end-arteries.

The arteries of the pons and medulla oblongata are like the cortical arteries in their small size and like the central ones in respect that they come off from large stems, and pass directly into the substance of the part.

# I.—MALFORMATIONS OF THE BRAIN.

The more considerable of these have been considered in the section on general malformations. The principal forms are *anencephalus* accompanied by *acrania* or *cranioschisis*, *encephalocele* and *cyclopia*. The relation of these to dropsy of the ventricles of the brain (hydrocephalus) has also been referred to. It has been pointed out that hydrocephalus occurring at an early period leads to these serious malformations, which are incompatible with life. But hydrocephalus sometimes occurs at a later period, and the child is born with a *congenital hydrocephalus*, which is itself sometimes regarded as a malformation. It will be taken up, however, along with the other forms of hydrocephalus.

1. **Congenital smallness or Aplasia of the brain.**—This may affect the brain as a whole or parts of it.

**Micrencephalus** signifies a general smallness of the brain, which is usually associated with **Microcephalus** or smallness of the head, the latter term, being frequently used as synonymous with the former. The defect usually concerns the cerebrum chiefly, although other parts may also be involved. The brain may be defective in structure as well as in size, the convolutions being small, irregular or defective. The affection may be due in some cases to a simple defect in formative power in the cerebral matter, but it results sometimes from more definite pathological conditions in the foetus. Thus there are cases in which it co-exists with hydrocephalus. In these the size of the head may not be diminished, but that of the brain may be seriously so. Ahlfeld, again, ascribes many cases of micrencephalus where there is no existing hydrocephalus to a dropsy at an earlier period. Premature union (synostosis) of the cranial sutures and synchondrosis of the basal bones have been assigned as a cause of microcephalus (see p. 455). The cranium cannot expand and the brain remains small. Again, thickenings of the pia mater sometimes co-exist, and this indicates an early inflammation as the probable cause of the defect in the brain. Any considerable degree of micrencephalus is accompanied by idiocy.

The average weight of the entire encephalon in the adult is about 50½ ounces in the male and 44½ in the female, the weight ranging from about 43 to 60 ounces in the male, and 40 to 50 in the female, although there are exceptional cases both below



and above that range. The average weight of the cerebellum is  $5\frac{1}{4}$  ounces in the male and  $4\frac{3}{4}$  ounces in the female, that of the pons and medulla oblongata is about 1 ounce in the male, and the same in the female. (See further in Quain's Anatomy.)

**Parts of the brain** are sometimes congenitally small. The convolutions may be imperfectly formed, or there may be asymmetry of the hemispheres. The cerebellum is also not infrequently imperfect. It may be as small as a walnut, or present various degrees of defect.

In a case reported by Fraser (Glasg. Med. Jour., xiii., 1880) there was a congenital ataxia which was found after death to be due to smallness of the cerebellum ( $2\frac{1}{2}$  ounces). Similar symptoms existed in a sister, who was presumably affected with the same defect.

**2. Hypertrophy of the brain.**—This is a condition rarely observed, but occasionally, without any hydrocephalus, a child is born with an unusually large encephalon. There are also cases in which in later life, a hypertrophy of the brain takes place, sometimes acute, sometimes chronic.

**3. Heterotopia of the brain substance.**—Certain cases of *Hernia cerebri*, in which there is no hydrocephalus, are regarded as belonging to this class. Sometimes also masses of grey substance are met with in abnormal situations, as in the midst of the corona radiata. Some of these cases may really be tumours of the brain, but they are congenital, and due to errors in development.

**Porencephalus** is a term used for defects of the brain in the form of gaps in the superficial parts penetrating more or less deeply into the cerebral substance, sometimes as deeply as the ventricles. The gaps are occupied by the œdematous membranes. The lesion is ascribed to inflammation occurring in the fœtus. There may be associated with it a congenital paralysis and atrophy of parts corresponding with the lesion in the brain.

In cases of **Idiocy** there is defect of the brain, but it may be in various forms. There may be micrencephalus or congenital hydrocephalus, or partial defects. There may be little beyond some traces of inflammation in the membranes. Similarly **Cretinism**, which is endemic in some localities and is usually associated with goitre, is related to various lesions of the brain. In this condition, however, the defect in the brain seems to depend on the condition of the bones. The skeleton as a whole is stunted and deformed, and the bones of the skull show premature synostosis and synchondrosis, so that the elongation of the basal parts and expansion of the vault are variously hindered, the brain being correspondingly defective.

**Literature.**—For greater defects see under Malformations. *Micrencephalus*—VIRCHOW, *Gesam. Abhandl.*, 1856, p. 891; HITZIG in Ziemssen's *Encyclopædia* (with literature); AEBY, *Mikrocephalie and Atavismus*, 1878. *Defect of cerebellum*—FRASER, *Glas. Med. Jour.* xiii., 1880; GOULD, *Path. trans.*, xxxiii. 6; CLAPTON, *Path. trans.*, xxxii. 20. *Hypertrophy*—HITZIG, *loc. cit.*; VIRCHOW, *Ges. Abhandl.*, 1856, *Virch. Arch.*, xxxiii.; LEES, *Dubl. Med. Jour.* xxii., 1842; TUKE, *Jour. of Anat. and Phys.*, xii., 1873; LANDOUSY, *Gaz. méd. de Paris*, No. xxvi., 1874. *Heterotopia*—HITZIG, l. c.; VIRCHOW, l. c., and *Virch. Arch.*, xxxviii., 1886; MESCHÉDE,



Virch. Arch., lvi., 1872; SIMON, Virch. Arch. lviii., 1883. *Porencephalus*—HESCHL, Prag. Vierteljahrschr., 1868, and Jahrb. f. Kindereilk, xv., 1880; DE LA CROIX, Virch. Arch. xcvi.; BINSWANGER, do., cii.

## II.—TRAUMATIC LESIONS OF THE BRAIN.

**Laceration of the brain.**—The brain may be injured directly or indirectly. In injuries involving fracture of the skull the brain is often simultaneously wounded, or the bone may be carried downwards and impinge upon the brain. On the other hand, the brain is often lacerated without a corresponding external wound or fracture of the skull, by so-called *contre coup*. In falling from a height, if the head come suddenly against the ground the solid skull will be simultaneously arrested, while the soft brain substance may expend its force against the internal surface of the cranium, and so be greatly lacerated. The point of injury to the brain will be on the opposite part of the head to that which struck the ground, and there may be a wound or fracture at the latter.

The lacerated brain substance is torn and softened and there is more or less hæmorrhage. The hæmorrhage may be important if any considerable vessel in the meninges be injured, but is not usually great from the brain substance itself. The injured parts are afterwards affected by inflammation, which in the case of compound fracture with a septic wound may be acute, leading to abscess, but in most cases is chronic. Such chronic inflammations, affecting brain and meninges, may persist for long periods, even enlarging their area and producing extensive destruction and shrinking of the cerebral substance. In the affected part the membranes will be adherent and thickened, and the brain substance indurated by new-formed connective tissue. Important nervous and mental phenomena may follow such lesions even when the injury to the head has not been great.

## III.—LESIONS AFFECTING THE CIRCULATION IN THE BRAIN.

These conditions are somewhat variously associated with each other and with lesions of other kinds. They mostly imply local or general alterations in the volume of the blood and lymphatic fluid in the brain, and such alterations have somewhat complex relations, chiefly arising from the fact that the skull, in the adult at least, is a closed cavity with rigid walls, whose contents as a whole are scarcely capable of variation, although the fluid constituents are variously interchangeable.

1. **Hyperæmia, Anæmia and Œdema.**—Hyperæmia in the brain, as elsewhere, is divisible into active and passive. **Active hyperæmia** occurs generally or locally in consequence of hypertrophy or over-action



of the heart, especially when the arteries are atheromatous and unable to control the circulation. It also occurs in inflammations of the brain or meninges, and in various conditions of excitement of the brain, as the delirium of fevers, the early stage of general paralysis, the typhoid stage of cholera. In these conditions it may be regarded as an inflammatory phenomenon.

**Passive hyperæmia** results most directly from thrombosis of the cerebral sinuses, but is also an occasional consequence of pressure on the jugular veins, or of disease of the heart or lungs leading to general venous engorgement.

The appearances visible after death are frequently very insignificant, especially in active hyperæmia, whose existence is usually matter of inference rather than of observation. The overfilling of the vessels is most visible in the meninges, but there may be also a deepened colour of the convolutions, and the venous stems in them and in the white substance may be visibly dilated.

**Anæmia or Ischæmia** results mostly from obstruction of arteries, (see below). It may also be a consequence of general anæmia, in which case it affects the brain as a whole. It may result from local pressure on the brain produced by tumours, extravasations of blood, or inflammatory exudations. Local anæmia in all these cases is liable to result in softening of the brain substance.

**Œdema of the brain** is also usually but a part of some other lesion. Œdema of the brain substance is a rare and in many cases a hypothetical condition. The more definite œdemas affect the membranes and cavities, and will be considered further on. There is, however, an occasional local œdema of the brain substance in the neighbourhood of hæmorrhages, tumours, and veins obstructed by thrombus, and it is also believed to occur at the outset of local inflammations.

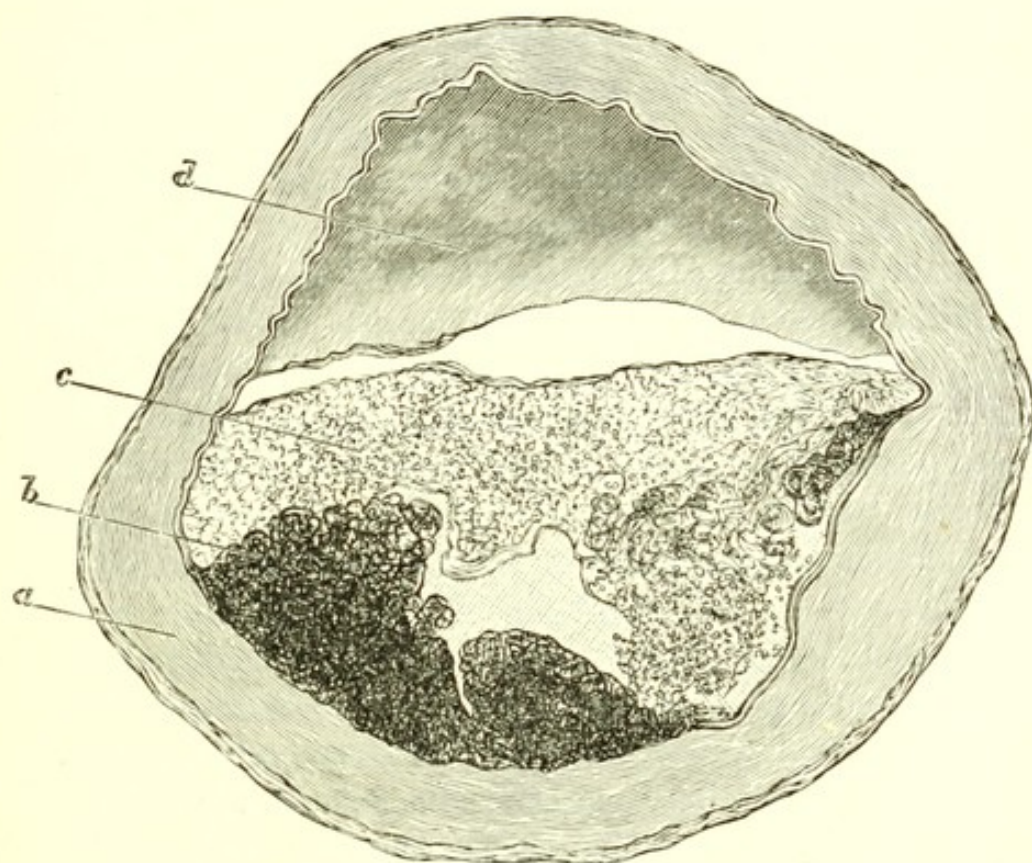
**2. Occlusion of arteries.**—This is a frequent and serious lesion, occurring mostly as a result of embolism or thrombosis.

**Embolism** occurs in the great majority of cases in connection with acute endocarditis or valvular disease of the heart from chronic endocarditis. In the former case the embolus is a piece of fibrine broken off from the inflamed aortic or mitral valve; in the latter case it is mostly a piece of calcareous matter from the indurated and calcified curtain. Besides, we may have globular thrombi in the left auricle or ventricle becoming detached or softening, and so giving rise to plugging. So also may thrombi in the aorta become detached and carried off. Much more rarely do emboli come from the lungs, but sometimes in gangrene of the lungs there is a thrombosis of the veins, and from bits of fibrine carried off we may have septic embolism of the brain. A cancer of



the lungs after penetrating into the pulmonary vein may produce embolism.

It is matter of general observation that in cerebral embolism the **Middle cerebral artery** (or artery of the fissure of Sylvius) is the vessel plugged in the great majority of cases. It is also stated that the left middle cerebral is more frequently the seat of embolism than the right, but this has been doubted by competent observers. The frequency with which the middle cerebral is affected admits of easy explanation. As the vertebral artery arises from the subclavian nearly at a right angle, it is not common for an embolus to pass into it. But the innominate and the carotid, being nearly in the direct line of the current from the aortic orifice, readily receive any fragment. Then



\* Fig. 221.—Atheroma of a small cerebral artery the eleventh of an inch in diameter, with thrombosis. *b*, atheromatous internal coat ; *c*, thrombus on its surface partially organized. Between these two is an irregular clear space, which represents fresh blood which had been insinuated between patch and thrombus. *d*, remaining calibre filled with blood  $\times 34$ .

again, the middle cerebral is the direct continuation of the internal carotid and an embolus will more readily sweep into it than diverge forward or backward. It is important here to bear in mind that the middle cerebral supplies the greater part of the basal ganglia, including nearly the whole of the corpus striatum and internal capsule and a part of the thalamus opticus. It also supplies the greater part of the motor convolutions.



**Thrombosis** of the cerebral arteries arises in consequence of some alteration of the walls of the arteries. It is predisposed to by weakness of the heart, but mere sluggishness of the circulation so induced will hardly cause thrombosis in a healthy artery, although it may do so in the venous sinuses of the dura mater, where the circulation is naturally so much slower. Thrombosis is mostly induced either by **Atheroma** or by **Syphilitic disease** of the arteries. In both cases the internal coat is thickened, and the surface is rough, while the calibre of the artery is already considerably encroached on, as Fig. 221 shows. Atheroma occurs mostly in old persons, and it is usually most pronounced in the larger arteries of the base, where there may be numerous yellow patches. But not infrequently it extends to the finer ramifications, and it is in these that it most readily leads to occlusion by thrombosis. The artery from which the figure is taken, for instance, had an external diameter of the eleventh of an inch. Syphilitic disease of the arteries, on the other hand, occurs most frequently in younger people in connection with gummata, and these may occur at any part of the surface of the brain.

**Effects of occlusion of arteries.**—Here, as in other parts, the effects of occlusion depend chiefly on whether the arteries concerned have sufficient anastomosing communications or not. The arteries of the circle of Willis anastomose freely, and occlusion of one of them leads only to a very temporary derangement of the circulation. The larger branches anastomose sparsely, and occlusion of them has much more serious effects. The nutrient arteries do not anastomose at all, and occlusion of them has very evil results.

The most direct result of occlusion of arteries is **Anæmia** of the part supplied. As the occlusion in the case of embolism is sudden, there is often a very abrupt interference with the cerebral functions. In the case of a large artery, such as the main stem of the middle cerebral, there may be a very extensive anæmia, leading to a fatal issue before the anastomosing circulation can be established.

**Softening of the brain substance** (*Ramollissement*) is a further and more important consequence. This is really a necrosis with fatty degeneration of the nervous tissue, and it only occurs when the conditions are such that the circulation is brought absolutely to a standstill. This is the case when any of the nutrient arteries are obstructed, as these are end-arteries. The nutrient arteries which come off from the first part of the middle cerebral are the most exposed to obstruction from embolism, and hence softening of the central parts of the brain, especially in the region of the corpus striatum, is of somewhat common occurrence. It is important to notice that, for reasons to be afterwards



considered, it is these arteries also which most frequently give rise to hæmorrhage.

But softening not infrequently occurs although the vessels occluded are not end-arteries. This is especially true where thrombosis is the cause of the occlusion. As we have already seen, thrombosis is common in connection with atheroma. Now this is a disease of old people, in whom the circulation is weak. Under these circumstances occlusion of a small peripheral artery may lead to softening, before the force of the blood has brought about an anastomosing circulation, all the more because many of the neighbouring arteries are also partially obstructed by atheroma. Even in the case of embolism there may be considerable cortical softening if the embolus has broken up and plugged several vessels at the same time, so as to interfere with the establishment of the anastomosing circulation.

The **Changes in the brain substance** in softening have already been incidentally referred to. It will be a local lesion limited to the piece of brain substance to which the affected artery is distributed.

Softenings have often been distinguished according to the colour presented by the affected brain substance, so that **white, yellow, and red** softenings have been described. For the most part the colour depends on the blood mixed with the nervous tissue, and the amount and condition of the former can hardly be regarded as a chief characteristic of the softening; the colour is therefore not of primary importance.

We have seen already that softening of the brain substance is really due to a **Necrosis**. The result of the death of the nervous structures is their disintegration. The nerve fibres very rapidly break up; the myeline of the medullary sheath coagulates and escapes from the primitive sheath, and afterwards breaks up into fine fat granules. The ganglion cells are more resistant, but they also become granular and gradually disappear. The cells of the neuroglia and the nuclei of the walls of the vessels undergo fatty degeneration. These structures become filled with finely divided fat, so that the neuroglia cells are converted into characteristic **Compound granular corpuscles** and the nuclei of the vessels are converted into aggregates of fat granules at intervals along the vessels (Fig. 222). So far as the neuroglia cells are concerned it is probable that in some cases they pick up fat arising from the disintegrated myeline. It is not so clear that the nuclei of the blood-vessels do this, and there is here, probably always, a proper fatty degeneration. Besides the neuroglia cells there may be present in the part amoeboid cells, or these may pass into it after the occurrence of the necrosis, and these also become occupied by granular fat. In



this way there are, frequently, large numbers of compound granular corpuscles which are very conspicuous when a piece of the softened tissue is examined in the fresh state under the microscope. A ready means of distinguishing true pathological softening from a mere post-mortem change is afforded by the presence of these cells. In true softenings they are present in large numbers and afford a highly characteristic appearance.

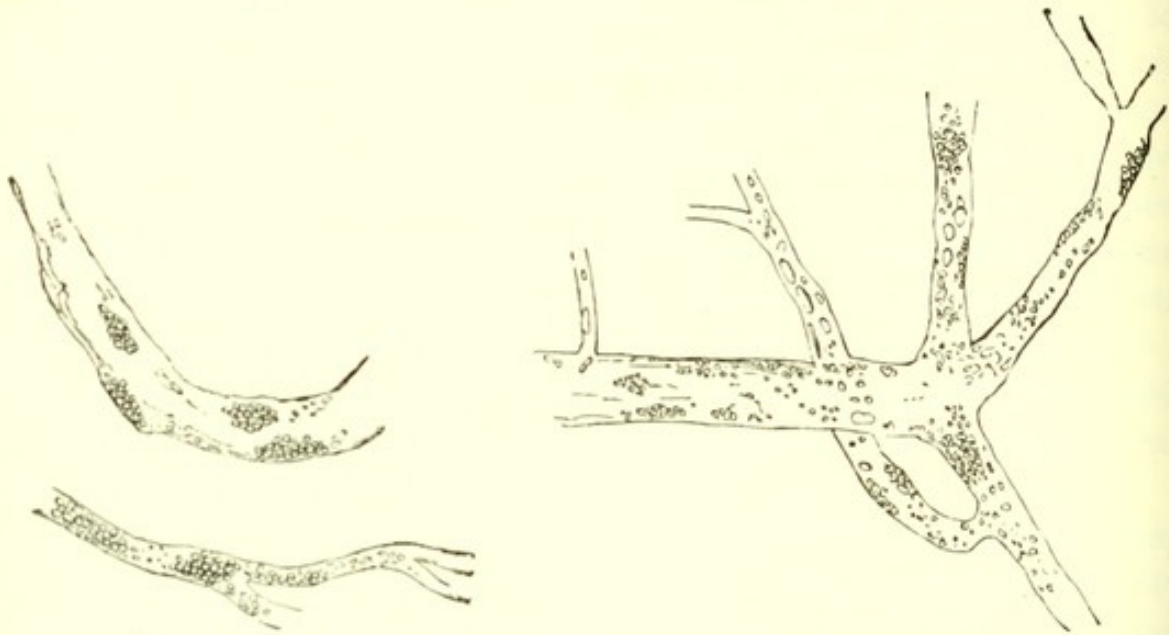


Fig. 222.—Fatty degeneration of the vessels in cerebral softening. (PAGET.)

In some cases, as already mentioned, the softened brain substance is largely mixed with blood. At first the blood gives a red colour to the softening. But as time goes on the colouring matter is dissolved out of the red corpuscles and diffused throughout the softened structures, as well as to some extent in the brain substance around. In that case the colour becomes less intense and merges into yellow, or the colour may be reddish yellow from the outset. It will be seen from this that red and yellow softening run together, and that the latter is often merely a later stage of the former. At the time when the softening occurs the vessels are often filled with blood, which soon coagulates. The fibrine disintegrates as the colouring matter is dissolved out from the corpuscles. Sometimes the dissolved colouring matter is deposited in the granular or even in the crystalline form inside or outside the vessels. If there is very little blood in the vessels of the softened part then the colour is white or grey. This is mostly seen when death has occurred very soon after the occlusion of an artery.

The ultimate disposal of the necrosed piece of brain substance occurs on principles already considered. The softened brain substance has ultimately the characters of a fatty emulsion, which is gradually absorbed. A chronic inflammation occurs around, just as in the case of a



hæmorrhage, and in a similar fashion connective tissue is produced, taking the form of a cyst or a cicatrix, according to circumstances. The resulting cyst or cicatrix is not always to be distinguished from that of a cerebral hæmorrhage. Usually the latter presents more distinct pigmentation, but the former may also present pigment granules and crystals.

**Other forms of softening** are of less consequence than those from occlusion of arteries. Softenings are very frequent around extravasations of blood. They are partly due to mechanical destruction, but also to anæmia from the pressure of the blood. Softenings around tumours are similarly due to anæmia. There is occasionally a softening from localized inflammation, more especially such septic forms as lead on to abscess.

**Occlusion of arteries in chorea.**—The frequent association of this disease with acute endocarditis has suggested the view that it may have its origin in embolism of the arteries and capillaries of the brain, multiple embolism with softening having been found in some cases, but not in all. Embolism, however, may be a coincidence related to the endocarditis, and not necessarily connected with the pathology of chorea. This, indeed, is evident from the fact that acute endocarditis is not infrequently absent in cases of chorea. Other changes in the vessels have been observed, such as dilatation of the arteries and veins throughout the substance of the brain and spinal cord, exudations or small hæmorrhages, sometimes with blood crystals, around the vessels. These lesions, however, are by no means constant, and are to be regarded as consequences of irritation of the brain, or as coincident phenomena.

It seems probable that in chorea there is an irritant present in the blood, and it may be the same irritant as that in acute rheumatism. It attacks the nervous system, producing hyperæmia and exudation from the vessels. While the whole central nervous system is more or less affected there are certain parts specially involved, these parts being what have been called the accessory portions of the nervous system. It may be supposed that the fundamental and simpler parts of the brain and cord are more stable than the accessory parts, and that when attacked by an irritant the latter will give way first. The anatomical distribution of the lesions suggests that this is so, and the symptoms of chorea indicate disorder in "movements which are acquired, and which are probably only learned by a long apprenticeship" (Hughlings Jackson).

**3. Thrombosis of the cerebral sinuses.**—These venous channels are somewhat frequently the seat of thrombosis. The coagulation may have its starting point in an inflammation propagated from a neighbour-



ing structure. The most frequent origin of such inflammations is caries of the temporal bone from disease in the ear, but it may follow on injury to the head, inflammations of the skin of the face and scalp, especially erysipelas, and of the bones. In all these cases the thrombosis usually has its starting point towards the base of the skull, and especially in the lateral and petrosal sinuses.

There is, however, another class of cases in which the thrombosis has a more obscure origin, and seems sometimes even spontaneous. The blood simply coagulates in the sinus, and it is usually in the longitudinal sinus that the coagulation begins. In most of these cases the person is in a state of debility, and the thrombus may be regarded as **Marasmic** in its origin. But there are cases in which there is no obvious weakness of the heart, and the coagulation has no apparent cause. The localization of the thrombosis suggests stagnation of the blood as its cause. The sinuses are rigid tubes incapable of narrowing when the circulation is slow, and they are intersected by bands of connective tissue. The longitudinal sinus also is so situated that, at its middle part at least, the blood passing from the cerebral veins flows upwards to it against the force of gravitation (see p. 67).

The **Effects** of thrombosis of the sinuses will vary according to the cause. If due to the propagation of inflammation from neighbouring structures, then a suppurative phlebitis may result, with **Meningitis**, and even **Abscess of the brain**. There are, however, cases of abscess of the brain without meningitis, in which the inflammation seems to have extended along the veins either in their interior or in their sheath.

In the other form the results are usually much less serious. The veins which open into the sinus are greatly engorged, and the thrombosis may extend into them. It is here chiefly the longitudinal sinus that is concerned, and the veins which open into it are those of the cerebral hemispheres. These may stand out as prominent worm-like cords filled with dark coagula. In most cases the blood finds its way by other routes, and there may even, after a time, be a re-establishment of the circulation in the sinus. Sometimes, however, the obstruction in the veins is such as to lead to **Hæmorrhage**. This is usually in the form of numerous capillary hæmorrhages, but sometimes there is along with this a large effusion of blood in the substance of the brain and under the soft membranes. The seat of such hæmorrhages is usually the superficial and upper parts of the hemispheres, and they are frequently multiple.



## IV.—CEREBRAL HÆMORRHAGE.

By this term is meant bleeding in the substance of the brain. The blood may be large or small in quantity but it pushes aside the brain substance, tearing for itself a cavity where it coagulates. With the doubtful exception of certain diseases of the blood, where the hæmorrhage may possibly be by diapedesis, there is always rupture of one or more vessels.

In a previous page reference has been made to the two orders of cerebral arteries, and the difference in the circumstances of hæmorrhages from these are such that we must consider them separately here.

1. **Hæmorrhage from the larger cerebral arteries.**—We have seen that these vessels run in the soft membranes, and it might be supposed that their rupture would give rise to meningeal rather than cerebral hæmorrhage, and so it is usually stated. But this is not the case. The blood nearly always finds its way into the brain substance, where it is found usually in much larger quantity than in the meninges, so that the case has much more the characters of cerebral than of meningeal hæmorrhage. There is usually some blood in the meninges occupying the sulci between the convolutions, and it may even be considerable, extending to the base and perhaps covering the optic commissure; but usually the amount is small, and, on account of the large cerebral hæmorrhage, is apt to be overlooked. It is very rare indeed for the blood to escape to the surface and appear in the cavity of the dura mater. Any appearance of blood in the meninges, however, should at once direct attention to the larger cerebral arteries as the probable source of the hæmorrhage.

The explanation of the extension of the blood into the brain substance suggests itself when we consider the circumstances of the parts. When rupture of such an artery occurs the blood tears its way around; it passes into the loose connective tissue, and by and by reaches the surface of the brain. Here the tissue, being soft, tears readily, and the blood rapidly passes inwards. On the other hand the connective tissue on the surface is tough and the blood will tear it with difficulty. The blood may work its way from space to space in the connective tissue, but this takes time and probably needs considerable pressure. There is another circumstance which probably has to do with the blood so constantly finding its way into the brain substance. As we shall see afterwards, a large number of the cases of hæmorrhage in this situation are from rupture of aneurysms. Now an aneurysm will probably project more readily towards the surface of the brain where the substance



is soft, than in other directions, and when such an aneurysm ruptures it may do so directly into the brain.

The **Cause** of the rupture of these larger arteries is to be sought for in disease of their walls and increase of the blood-pressure.

The commonest cause is **Aneurysm** of these arteries, which is of remarkably frequent occurrence. The great majority of the cases of cerebral hæmorrhage occurring before the age of fifty years is due to the rupture of aneurysms of these larger arteries. The aneurysms are mostly thin-walled and therefore prone to rupture. Their most frequent seat is on the middle cerebral artery or one of its branches in the fissure of Sylvius, but they may occur on any of the arteries of the brain and are not infrequently multiple (see Fig. 223).

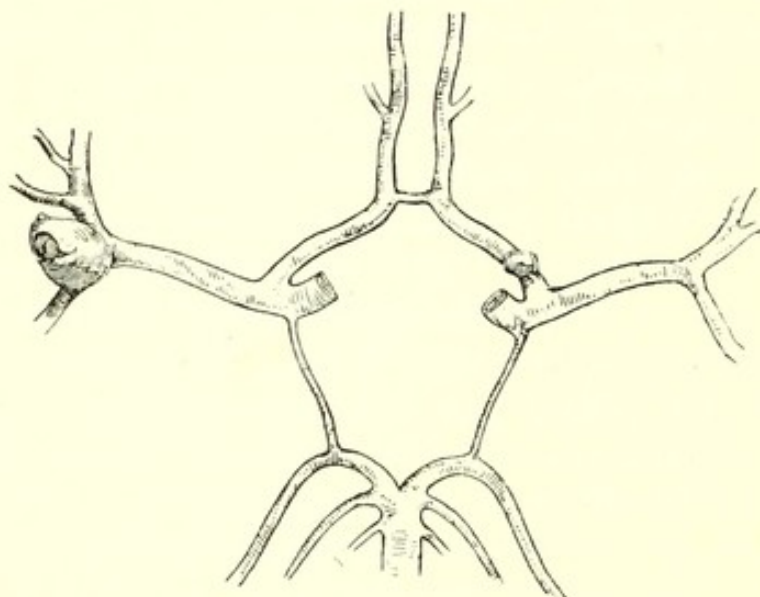


Fig. 223.—Aneurysms of larger cerebral arteries. A large one at the bifurcation of the left middle cerebral. There is an aperture at its summit, from which fatal hæmorrhage occurred. A small one near the origin of right anterior cerebral.

The frequent occurrence of these aneurysms and their serious import suggests an inquiry into the causes of their formation. In the first place the arteries are thin-walled and are placed in a loose tissue, so that they very readily undergo dilatation. Any local injury to the wall may be the starting point of the dilatation. This is often produced by **Embolism**. In a large proportion of cases the aneurysm is associated with valvular disease of the heart, and an embolus imperfectly obstructing an artery, especially if it be a cretaceous piece broken off from a valve, may readily injure the wall so as to allow of dilatation. Ponfick has found that in a considerable proportion of cases of acute endocarditis there is embolism, with either fully formed or incipient aneurysms. When they have an origin such as this the aneurysms will be specially thin-walled and partake of the characters of false aneurysms. This mode of origin also goes far to explain the greater predominance of these aneurysms in the middle cerebral artery, which, as we have seen, is especially liable to embolism. Another occasional cause of cerebral aneurysm is **Atheroma**. This disease injures the vessel-wall and produces obstruction, and it may lead to aneurysm on the one hand by



weakening the wall, and on the other by locally increasing the blood-pressure on the peripheral side of the obstruction. Syphilitic disease is also assigned as a cause of aneurysm.

The subject of **Atheroma** as a direct cause of cerebral hæmorrhage is a somewhat difficult one. This condition is undoubtedly in many cases associated with cerebral hæmorrhage, but it is difficult to believe that it can be the sole cause. Atheroma produces, as Fig. 221 shows, a thickening of the arterial wall. At the same time, however, it weakens the wall, and if the coats get separated and the blood penetrates behind the atheromatous patch, there may be serious risk of rupture. Besides this, atheroma will, as mentioned above, produce, by obstructing the vessel, an increase of blood-pressure on the proximal side of the patch and also in neighbouring arteries. If there be extensive atheroma the mere rigidity of the arteries may produce an increase of blood-pressure in the smaller arteries. But all these conditions seem scarcely capable by themselves of leading to hæmorrhage, and it is in cases where atheroma is associated with a general increase of blood-pressure that this occurs, especially in chronic Bright's disease.

On the other hand, **Increase of blood-pressure** alone can hardly lead to hæmorrhage. In chronic Bright's disease there is frequently a constant increase in the arterial tension, and hypertrophy of the left ventricle of the heart. This condition will conduce to hæmorrhage in both orders of arteries in the brain, especially if this disease be associated with atheroma, or with fatty degeneration of the smaller arteries of the brain, which some assert to be common in Bright's disease.

2. **Hæmorrhage from the nutrient arteries.**—As these vessels run in the substance of the brain the hæmorrhage will be always cerebral and will rarely extend to the meninges. It might be supposed that as the nutrient vessels are small the hæmorrhage from them would be small, and in many cases it is so; but when bleeding has once begun, the blood tearing the brain substance ruptures other vessels, and there is often a considerable effusion of blood.

Hæmorrhage of any consequence rarely occurs from the cortical nutrient vessels, but is very common from the central ones. The circumstances of these latter go far to explain this. They arise from large stems, mainly from the middle cerebral immediately after its origin from the internal carotid. It is clear that the blood here will be at a pressure not much less than that of the aorta, and any variations of pressure will tell readily. On the other hand, the cortical vessels mostly arise from fine vessels in which the blood-pressure has been reduced by successive division and sub-division.

As to the **Causes** of hæmorrhage in these arteries, **Aneurysm** again



plays the most important part. As the arteries are small so are the aneurysms, but they are numerous in the same person. Such aneurysms have been called by Bouchard and Charcot **Miliary aneurysms**. They occur in every region of the brain, but are most readily detected on the surface of the convolutions, where, on stripping off the pia mater from the convolutions, they may be seen as small red or brown spots. When examined under the microscope they have all the characters of ordinary aneurysms. Most of them are sacculated (Fig. 224), but some are fusiform (Fig. 225). It is stated that the

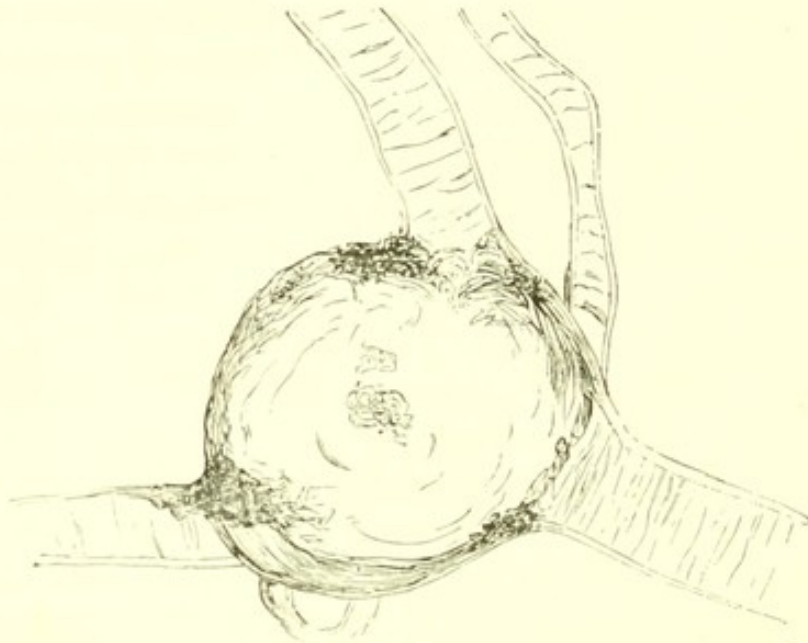


Fig. 224.—Sacculated miliary aneurysm of a nutrient artery of the brain. The aneurysm is about the twentieth of an inch in diameter.  $\times 27$ .

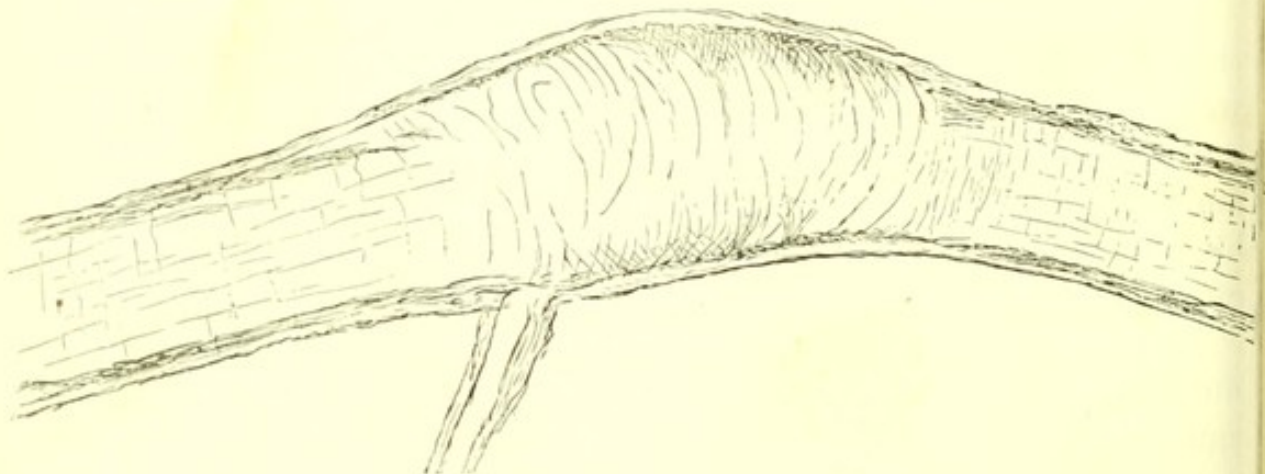


Fig. 225.—Fusiform miliary aneurysm.  $\times 27$ .

cause of their formation is a sclerosis of the walls of the arteries, involving first a formation of round cells in the external coat with subsequent fibrous transformation. There results an atrophy of the middle coat which seems to be the most direct cause of the dilatation. This diffused sclerosis of the nutrient arteries is mostly met with in old



people ; in persons above fifty cerebral hæmorrhage is, in the larger proportion of cases, due to rupture of miliary aneurysms.

In a case observed by the author there was frequently in the arteries a fatty degeneration affecting chiefly the internal coat. This was present in patches in a large number of vessels of small size. In connection with it there was sometimes a partial dilatation, an aneurysm obviously forming, and also fully formed aneurysms. Older and more recent hæmorrhages were connected with the aneurysms, and there was one very large fatal hæmorrhage.

Although the aneurysms are present in all regions of the brain, rupture seldom occurs except in those of the central arteries. The explanation of this has already been indicated, and it has been mentioned that the lenticulo-striate branch is pre-eminently that from which hæmorrhage occurs.

**Atheroma**, with increased blood-pressure, is occasionally a cause of hæmorrhage from the nutrient arteries as from the larger ones. Atheroma is not common in small arteries, but as the central arteries are much larger than the cortical ones, atheroma is commoner in them, and hæmorrhage from this cause is therefore, for a double reason, more frequent in these.

**3. Hæmorrhage from the capillaries.**—A certain amount of capillary hæmorrhage generally accompanies all larger bleedings. The explanation of this seems to be that the pressure of the blood produces such obstruction of the vessels around, that frequent leakage occurs from the capillaries. In thrombosis of the sinuses and veins there is also capillary hæmorrhage (see above). Again, embolism may cause capillary hæmorrhage, and, as we have seen, the blood is often mixed with the softened brain tissue. Septic embolism as in ulcerative endocarditis and pyæmia leads to capillary hæmorrhage. Lastly, we may have leakage from the capillaries in scurvy, purpura, leukæmia, and other morbid states of the blood.

In capillary hæmorrhages the collections of blood are generally small in size, forming a congeries of red puncta. But if very frequent and closely set they may run together and form a considerable effusion.

**Appearances of the brain in hæmorrhage.**—The appearances presented when a person dies soon after the occurrence of hæmorrhage are sufficiently characteristic. The effused blood increases the contents of the skull, and in order to its accommodation there must be some displacement and crushing of the remaining brain substance. If the hæmorrhage be at all extensive we find on opening the skull that the corresponding hemisphere is bulged outwards and perhaps projects beyond the middle line. The convolutions are more or less flattened,



and there is a certain dryness and glazing of the surface which indicate that all available fluid has been absorbed to make room for the addition made to the contents of the skull. These are all indications of increased pressure within the skull, and during life this increase of pressure causes symptoms referrible to the brain as a whole or to parts removed from the seat of hæmorrhage. It sometimes happens that the appearance of blood in the membranes suggests the existence of hæmorrhage before the brain is laid open, and this will be especially the case when rupture of an aneurysm of a larger artery or thrombosis in the sinuses has been the cause. On cutting into the brain substance the appearances will vary to some extent according to the cause and extent of the hæmorrhage. If there are numerous small hæmorrhages closely set there will be much softening of the brain, and the brain substance will be mixed with blood. If the hæmorrhage be large the blood will be more pure. In any case the blood produces softening in the parts around, which may be stained with the blood colouring-matter. As already mentioned, there are usually red spots from capillary hæmorrhage around the clot. The clot itself is mixed with the debris of brain substance, and the internal wall of the cavity in which it lies has an irregular character.

If the patient die almost immediately the clot is exactly like an ordinary gelatinous coagulum. But if he survive a day or two it has already drawn together somewhat and become firmer and has more of a brown colour. This is sometimes peculiarly manifest at the peripheral parts of the clot, so that a kind of capsule may be formed of condensed fibrine.

In washing away the clot from a cavity made by a hæmorrhage one often isolates many small arteries with round knots at their extremities. These are arteries which have been torn across by the accumulating blood. The torn arteries have withdrawn within their sheaths, and these latter have become distended with little plugs of blood which have, in the manner already described (p. 83), contributed to the stilling of the hæmorrhage. These little swellings may be readily mistaken for miliary aneurysms.

**Disposal of the clot. The Apoplectic cicatrix and cyst.**—The further organization of the coagulum is a slow process, and proceeds very much in the same way as organization of clots inside or outside of vessels. An inflammatory process is set up in the neighbourhood, and this results in the production of connective tissue which by and by encapsules the clot. The clot in the meanwhile softens; its pigment is dissolved out, and frequently deposited in the crystalline (see Fig. 226) or granular form in the softened material or in the capsule, often giving the parts a rosy or rusty colour. The blood being frequently mixed with brain



substance and both these constituents being degenerated, various appearances are presented; sometimes the capsule contains material of an atheromatous aspect.

Through time the contents are absorbed and there may result, as in the case of softening of the brain, a **Cicatrix**. In many cases, however, the absorbed matter is replaced by clear fluid and a kind of cyst is

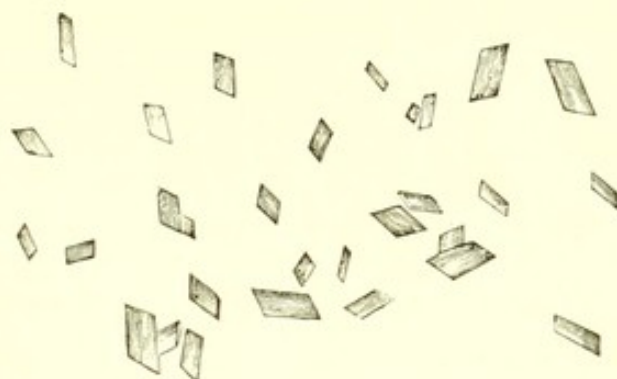


Fig. 226.—Crystals of hæmatoidin from a cerebral hæmorrhage. The crystals form oblique rhombic pillars of a reddish brown colour.  $\times 350$ .

the result, the so-called **Apoplectic cyst**. The cyst is not merely a sac containing fluid, but it is generally intersected by connective-tissue trabeculæ so as to be filled with a network. It is indeed more an œdematous cicatrix than a cyst. It has already been noticed that similar cysts and cicatrices occur as a result of softening of the brain. These latter, however, do not commonly show blood crystals in their walls, whereas the true apoplectic cyst may present them even at a late date.

The apoplectic cyst may be compared in its origin with a cicatrix; it arises by the formation of connective tissue and fills the place of tissue lost, and it is only because, being situated in the midst of the brain substance, it does not readily contract that we have a cyst rather than a cicatrix. If the hæmorrhage has been near the surface of a ventricle or of the brain itself we may have a cicatrix; or a cyst, by thickening of the trabeculæ and gradual drawing together of the parts, may be converted into a cicatrix. In the case of cicatrices occurring thus on the surface of the brain the soft membranes are depressed and puckered and firmly adherent to them. The cicatrices like the cysts often present some remains of blood-colouring matter.

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## V.—INFLAMMATIONS OF THE BRAIN.

These embrace a very extensive and varied set of conditions, some of which have little in common. Thus there are localized inflammations of an acute kind, going on sometimes to the formation of abscesses; chronic localized inflammations; inflammations diffused throughout the brain; inflammations extending to the brain substance from the membranes, etc.

1. **Acute localized encephalitis.**—We have already seen that, around and in the midst of hæmorrhagic foci and cerebral softenings, there are inflammatory manifestations, evidenced by the presence of leucocytes, containing fat granules. The inflammatory process does not go on to suppuration, but results in the formation of connective tissue, constituting a cyst or a cicatrix.

When the brain is injured—lacerated, for example, by a blow on the head or directly wounded by a protruding piece of fractured bone—then there may be a more intense inflammation, which sometimes goes on to suppuration, especially when the brain communicates with an external wound.

**Abscess of the brain.**—This is of somewhat frequent occurrence. It is met with most frequently in the cerebral and cerebellar hemispheres, rarely in the central parts. The abscess is usually single, but there may be several.

**Causation of abscess.**—The abscess is always septic, produced by the action of microbes conveyed from an infected centre near or remote. The centre is in the majority of cases in the neighbourhood of the brain (in 70 per cent., Gowers), being in the ear in nearly half the cases. Next to disease of the ear comes injury of the head, while the nose, orbit, and other parts of the head form rare seats of the primary disease. When the original focus is at a distance the abscess may have various seats, a certain proportion of cases being properly pyæmic. There are also a few cases in which no centre of suppuration is discoverable apart from the abscess.

**Suppuration in the middle ear** is the chief cause of abscess in the brain. There is a septic inflammation in the tympanic cavity and mastoid cells, the tympanic membrane being absent or perforated. The discharges are usually very putrid. This disease is common in children, chiefly as a sequela of scarlet fever, but abscess of the brain is most frequent in adults. This has been ascribed to the fact that at puberty the mastoid cells expand and both these and the tympanic cavity come nearer to the interior of the skull (Adams). Sometimes, indeed, the bone is defective over these parts, which are therefore



covered in by dura mater. The suppuration in the middle ear affects the mucous membrane, which is here closely connected with the bone and acts as a periosteum. Hence the bone is liable to be affected, becoming carious or even undergoing necrosis. Disease of the bone, however, is not necessary to the formation of abscess in the brain.

The tympanic cavity and mastoid cells both send blood into the sinuses of the dura mater by means of veins which pass through the bone; those of the tympanum passing into the petrosal sinus and those of the mastoid cells into the lateral sinus. On the other hand veins pass into these sinuses from the brain, the cerebral hemispheres sending branches into the superior petrosal and the cerebellum into the lateral sinus. The extension of the infective process is mainly by these veins. There is sometimes a **Septic thrombo-phlebitis** set up, which may affect the sinuses to a considerable extent, and even produce a **Pyæmia** in the ordinary way without leading to abscess of the brain. But the septic phlebitis may extend from the sinus to the cerebral or cerebellar veins and so lead to abscess. The extension on the other hand may be by the sheaths of these veins and not by their interior, or it may be a more direct extension from the bone to the dura mater and on to the brain. From the situation of the tympanum and mastoid cells and from the relations of the veins it follows that, when the extension is from the tympanum, the abscess is usually in the cerebrum, while in the case of the mastoid cells it is in the cerebellum, although this rule is not without exceptions. The abscess in ear disease is always an extremely fœtid one, as if the extension were not merely of the ordinary pyogenic bacteria (whose products have not a putrid odour) but more of a gangrenous character.

**Injury to the head** is a somewhat frequent cause, chiefly in cases of compound fracture with laceration of the brain. It is said that laceration by *contre coup* (see p. 551) may be a cause without external wound. **Disease of the nose and orbit** are rare causes, and it will be only when the bones are diseased, as in some cases of syphilis, that extension will occur.

When the septic focus is at a distance its most frequent seat is in the lungs, in cases of suppurating cavities, etc. In two cases of thrush abscesses in the brain have been found which contained growths of the *oidium albicans*.

**Formation and character of the abscess.**—The abscess begins presumably with a small softening which goes on to suppuration. The fully formed abscess contains a thick pus, usually greenish in colour, and in the case of ear disease exhaling a pungent putrid odour. In many cases the abscess is really a chronic one and the pus-corpuscles are dis-



integrated. It is bounded usually by a distinct wall formed of granulation tissue, sometimes partly developed into connective tissue. This frequently forms a definite separable membrane, which separates the abscess from the brain tissue, the latter being often softened in the immediate vicinity. The membrane takes some time to form, being rarely distinct till the third week and not fully formed for two months or even longer. When once formed and encapsuled the abscess may remain long stationary, but it usually enlarges gradually, and may finally burst into the lateral ventricles or externally.

The abscess causes enlargement of the part affected, and the convolutions over it are flattened and softened. Abscess is four times as frequent in the cerebrum as in the cerebellum, and the temporo-sphenoidal lobe is most frequently affected from its proximity to the ear.

2. **Chronic localized encephalitis.**—The processes of chronic inflammation are similar to those in the spinal cord, and the result here, as there, is **Sclerosis**. In the case of the brain there is hardly anything of the systematic sclerosis so frequent in the cord. There is, however, one form of disease which affects both brain and spinal cord, and which presents some analogies with the systematic sclerosis of the cord.

**Disseminated or Insular sclerosis, Sclerosis in patches.**—This condition is characterized by the occurrence of patches of chronic inflammation or degeneration scattered over the cord, or the brain, or both. These patches have all the characters of sclerosis; in the white substance, where they chiefly occur, they are grey, and in both white and grey substance they produce induration and shrinking. Under the microscope the sclerosed white substance shows loss of the medullary sheaths of the fibres, while the axis cylinders, except in advanced stages, are largely retained. In the grey substance, the ganglion cells are shrunk, sometimes with pigmentation. At the peripheral parts of the patch there may be evidences of more active inflammation, in the form of abundant round cells along with compound granular corpuscles.

The distribution of the patches is somewhat irregular. They are found of various sizes in the cord, medulla oblongata, pons, peduncles, corona radiata, convolutions, cerebellum. In such different situations the sclerosis leads to very various functional disturbances. The great frequency of motor tremors, however, suggests some tendency to localization. Erb has pointed out that when tremors exist, patches are specially present on the peduncular parts (pons, medulla, peduncles). If patches are seated there, the motor impulses will be imperfectly conducted but not absolutely interrupted, as the axis cylinders are preserved.

3. **Diffused encephalitis.**—This term may be applied to conditions



in which there is a general irritation of the brain, presumably by an irritant circulating in the blood. It may be held to include a considerable number of varied conditions. As the irritant, being in the blood, will attack all parts equally, we may expect to find evidences of irritation in the spinal cord as well as in the brain.

(a) In **Acute febrile diseases** it may be presumed that the brain is irritated by the morbid condition of the blood. According to Popoff, there are visible evidences of this in the case of typhoid fever in the presence of abundant leucocytes. These are stated to be particularly abundant around the vessels, and evidently spreading out from these. The leucocytes were frequently found in the spaces around the ganglion cells (pericellular lymph spaces), and even, in some cases, in the substance of the ganglion cells.

Middleton has pointed out that in delirium tremens, tubercular meningitis, uræmia, fracture of the skull with injury to the brain, erysipelas, etc., the brain substance is over-run with leucocytes, the appearances being very similar to those to be described as occurring in hydrophobia. In all cases where irritation of the nervous centres is evidenced during life by **Delirium** or otherwise, these signs of irritation may be looked for in the brain after death. In some cases of erysipelas, diphtheria, and septic inflammations, micrococci have been detected forming centres of irritation with minute softenings.

(b) **General paralysis of the insane** (*Dementia paralytica*).—The condition of excitement with which many cases of insanity begin, suggests irritation of the cerebral centres; but it is very difficult to determine the exact relation of the conditions here, especially as the cortex of the brain is very intricate in its structure and difficult to disentangle.

In general paralysis the most usual anatomical appearances are those of a chronic inflammation of the brain and its membranes (*Meningo-encephalitis*). But these appearances are the result of a prolonged disease, and the acute symptoms with which the disease often begins, and which may occur at intervals throughout, would indicate a stage of acute inflammation. Meyer asserts that death often occurs in this acute stage, and according to him there are, in the swollen and hyperæmic brain, multiple foci in which collections of leucocytes and capillary hæmorrhages are visible.

In the more **Chronic periods** of the disease the appearances are those rather of degenerative than of active processes, but the degenerative are presumably secondary to the inflammatory, the disease having very much the characters of an interstitial cerebritis and meningitis with loss of the proper nervous tissue as a consequence, a cirrhosis of the brain.



The appearances are prominently those of **Atrophy of the brain**. On removing the calvarium the dura mater is commonly found wrinkled and the soft membranes œdematous, evidently from shrinking of the brain substance. The œdema of the membranes is most visible in the sulci, which gape and are filled up with œdematous connective tissue. The fluid is chiefly in the membranes over the parietal and occipital lobes, perhaps from gravitation. There is also distension of the ventricles, frequently very great, so that the brain substance lying between the fluid in the ventricles and the superficial œdema of the membranes is greatly shrunk. The surface of the ventricles is beset with little prominent granulations which are often very marked. The pia mater is adherent to the surface of the convolutions, so that on attempting to remove it bits of brain substance come away. These adhesions may be taken as indications of the inflammatory nature of the disease, and their locality affords some evidence of the localization of the lesions. Crichton Browne states that they occur mainly in the anterior three fourths of the brain, affecting the frontal lobe chiefly in its anterior and posterior thirds, and the parietal in all its convolutions.

The brain as a whole is greatly reduced in weight. Taking the normal weight of the brain, including membranes, etc., as fifty ounces for the adult male, and forty-five for the female, the weight in general paralysis often falls to thirty-five ounces. The loss of weight does not affect the brain uniformly; it is mainly the cerebral hemispheres that are affected, the basal ganglia and peduncular parts being much less so, and the cerebellum not at all.

**The membranes** show irregular thickenings. The **Pia mater** presents milky opacities or more obvious patches of thickening. The **Dura mater** is so frequently affected that at one time it was supposed to be the primary seat of disease. It presents patches of thickening, opacities, and even flat bony developments, and it is commonly adherent to the calvarium. Not infrequently there are extravasations of blood of smaller or larger size accompanied by appearances to be described afterwards under **Pachymeningitis chronica hæmorrhagica**. The **Calvarium** itself sometimes presents thickening, which is usually diffuse, and may be chiefly due to the shrinking of the brain; sometimes there are also local prominences or actual exostoses. Sometimes the diploë is converted into dense bone, so that the calvarium is much heavier than normal—so-called sclerosis of the bone.

These appearances are due to inflammation and shrinking of the brain substance, the dropsy of the ventricles and œdema of the membranes occurring in order to fill up the space formerly occupied by the



brain, the cranium being a cavity of fixed size and the dropsy being *ex vacuo*.

The appearances under the microscope are chiefly those of diffused sclerosis and atrophy of the brain substance. We have already seen that there is sclerosis of the posterior columns of the cord, and this may even be an early condition. In the brain there are often traces of the earlier acute condition in the form of little clumps of pigment around the vessels, the remains of former extravasations. But the appearances are more those of degeneration. The ganglion cells are shrunken and frequently pigmented; the blood-vessels show sometimes fatty degeneration, sometimes a homogeneous glancing appearance (colloid); and they are frequently obstructed by glancing masses which are partly calcareous and partly colloid. These appearances are not to be regarded as in any way peculiar to this disease, they are simply evidences of the profound atrophy of the brain, and may occur to a limited extent in old persons where there has been only a senile atrophy.

(c) **Hydrophobia.**—In this disease there are indications of irritation

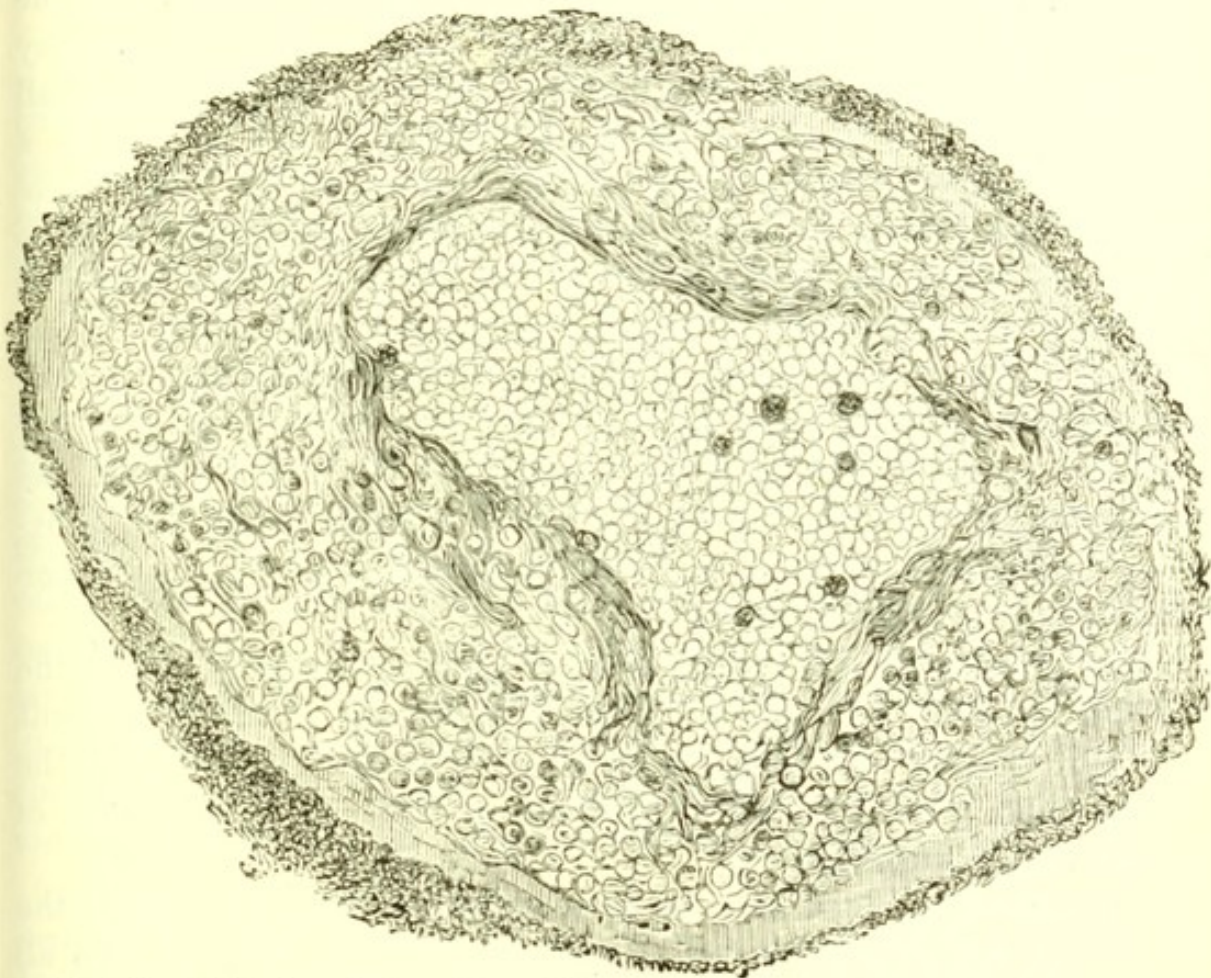


Fig. 227.—A blood-vessel from the medulla oblongata in a case of hydrophobia. Large numbers of round cells are seen in its sheath.  $\times 350$

of certain nervous centres, and of a greatly increased reflex irritability. The centres irritated are chiefly those of the medulla oblongata and



spinal cord, although delirium, indicating irritation of the cerebral cortex, is also occasionally present. The symptoms point mainly to the medulla oblongata; there are various spasms of the muscles of deglutition and respiration which occur to some extent spontaneously, but are also evoked by slight sensory stimulation. The mere sight of water gives rise to the idea of swallowing it, and brings on a violent spasm of the muscles of deglutition. A breath of cold air on the surface of the body causes a violent respiratory spasm or gasp. The centres in the cord are also irritated, as shown by the tendency to spasm of the muscles generally.

In this disease an irritating virus is present in the blood and induces these conditions of the nervous system. Pasteur's researches have proved that the virus is present in the cord and medulla (see p. 311). After death most manifest signs of irritation are usually visible on microscopic examination. They are to be found most characteristically in the medulla oblongata and next to that in the spinal cord, but are not absent in the other parts of the nervous system. The most prominent condition is an accumulation of leucocytes around the vessels in the substance of the cord and medulla oblongata. There may be just a few leucocytes in the sheath, but from this there are all gradations up to a condition in which the vessel is clothed with a mantle consisting of many layers of leucocytes (Fig. 227). The leucocytes are also present elsewhere, and sometimes in such quantities as to induce some observers to speak of the collections as miliary abscesses. The leucocytes find their way into the pericellular spaces and are found keeping company with the ganglion cells in these situations. In the medulla oblongata the main nutrient vessels are towards the posterior part, and as the motor nuclei are in this region, it may be that the localization of the irritation here is partly determined by their proximity. In addition to these conditions minute hæmorrhages have been observed in the medulla and cord.

Signs of irritation are present in other parts of the body besides the nervous system. The salivary glands have been found to present abundant leucocytes between the glandular elements (Fig. 228). In the kidneys also there are signs of irritation in the form of dilatation of vessels and hæmorrhage.

It is clear then that here an intense irritant is circulating in the blood, and the intensity of it may be judged from the fact that all these very marked appearances occur within two or three days of the onset of the nervous symptoms. These structural changes vary greatly in degree in different cases, being sometimes very slight. This would indicate that the virus attacks the nerve structures directly, and that



the inflammatory appearances are concomitants and not necessary parts of the morbid process. They are, however, evidences of the presence of a strong irritant.

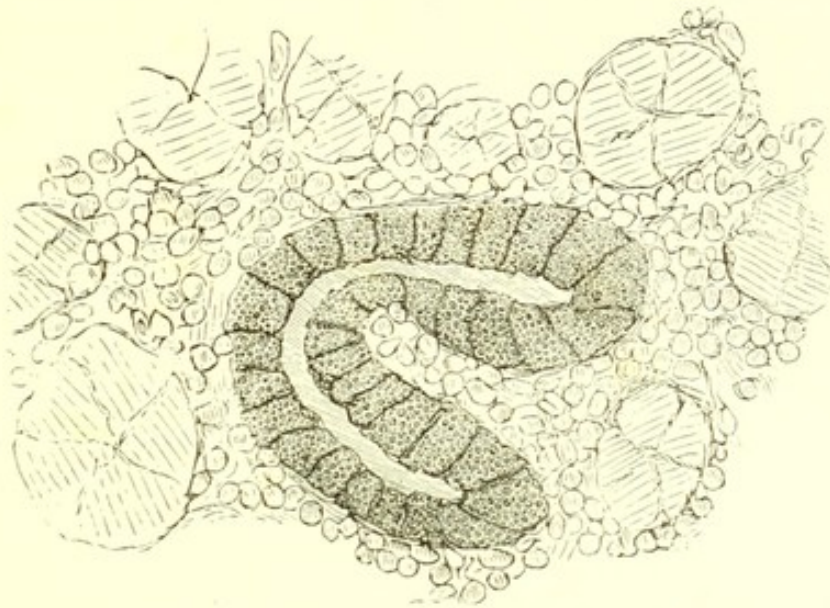


Fig. 228.—From the salivary gland in a case of hydrophobia. In the middle is portion of a duct; abundant round cells surround it as well as the glandular structures, shown in outline  $\times 350$ .

(*d*) **Tetanus.**—In this disease we have another example of an intense irritation of the spinal cord and medulla oblongata. There is here violent spasm of the muscles, but those under the control of the medulla oblongata are not so specially involved as in hydrophobia, although the person frequently dies from spasm of the muscles of respiration.

Here again there are signs of irritation discoverable in the spinal cord and medulla oblongata, although they are very different from those in hydrophobia. In tetanus the most prominent and constant appearance suggests the exudation of fluid from the vessels. The fluid sometimes collects around the vessels and, as in hardened specimens the albumen is coagulated, a granular appearance is produced. In other cases it looks as if the fluid produced a disintegration of the neighbouring nervous tissue, the appearance being that of the “granular disintegration” of Lockhart Clarke. The localization of the disintegration around the vessels suggests its origin in an exudation from them. There is also usually an excess of leucocytes around the vessels and in the grey substance of the medulla oblongata generally.

The cause of the irritation in tetanus is not at all clear. It seems likely, however, that here also there is an irritant in the blood, just as there is in hydrophobia. In both of these diseases the temperature is elevated, and, especially in the case of tetanus, reaches sometimes a most startling height ( $110^{\circ}$  F.). It is difficult to account for this extreme rise of temperature by the muscular spasm, though the possibility of this explanation is not to be absolutely denied. It seems more probable, however, that an irritant acting to some extent generally on the tissues is the



cause of the elevation of temperature. This is the more likely because tetanus has been known to occur in epidemics, and because if a case survives eight days there is considerable probability of recovery, as if the poison ran its course in that period. Some authors have adduced evidence in favour of the view that the poison of tetanus is in the soil.

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#### VI.—ATROPHY AND DEGENERATIONS OF THE BRAIN.

Under the congenital malformations of the brain we have already described several conditions which might be included in the designation congenital atrophy or aplasia. We have also seen that in *Dementia paralytica* there is an atrophy of the brain as a whole.

**Senile atrophy** has somewhat similar characters to that last mentioned. Occasionally the brains of old people undergo a general shrinking, and the space is made up partly by dropsy of the ventricles and œdema of the membranes and partly by thickening of the cranial bones. The atrophy is sometimes partial.

**Degenerative changes.**—It is not necessary to give here a detailed account of these, as they have mostly been incidentally referred to.

The **White substance** of the brain undergoes a process of atrophy under various circumstances—in softening of the brain, in sclerosis, *etc.* It also presents a condition which Lockhart Clarke has designated **Granular disintegration**. This occurs in the neighbourhood of blood-vessels, and is probably due to exudation from them. The white substance degenerates into an indefinite granular material.

The **Ganglion cells** are frequently the seat of atrophy and degeneration. They may undergo a **Simple atrophy**, shrinking and losing their processes. But very commonly the atrophy is accompanied by pigmentation, and so a **Pigmentary degeneration** is the result. Vir-



chow was the first to describe a **Calcification** of the ganglion cells. This condition seems to be of frequent occurrence when these cells are suddenly deprived of vitality. It was found by Virchow originally in cases of commotio cerebri, but has since been seen in softening of the brain, in acute poliomyelitis anterior, etc. The ganglion cells, having died and having undergone coagulation-necrosis, become infiltrated with lime, like other dead structures. A **Hyaline or Colloid** degeneration of the ganglion cells has also been described, especially in cases of insanity.

**Secondary degenerations in the brain.**—These are similar to those already described as occurring in the spinal cord, and they concern chiefly the pyramidal tract. This tract degenerates when it is cut off from the motor convolutions at whatever level. This severance will be due to destructive lesions, usually hæmorrhage or softening, the most frequent seat of such lesions being the region of the corpus striatum. In cases of extensive destruction of the motor convolutions the secondary degeneration has been traced downwards through the pyramidal tract to end only in the spinal cord. A tract of grey degeneration is found in the parts of the corona radiata corresponding with the lesion. It extends to the internal capsule, occupying the anterior two thirds of its hinder limb (IK Fig. 217), thence it passes to the crus cerebri, occupying the middle two fifths of the crusta, extending from the surface below nearly to the substantia nigra above. In the pons the tract is anterior (Fig. 229), and is covered over by transverse fibres,

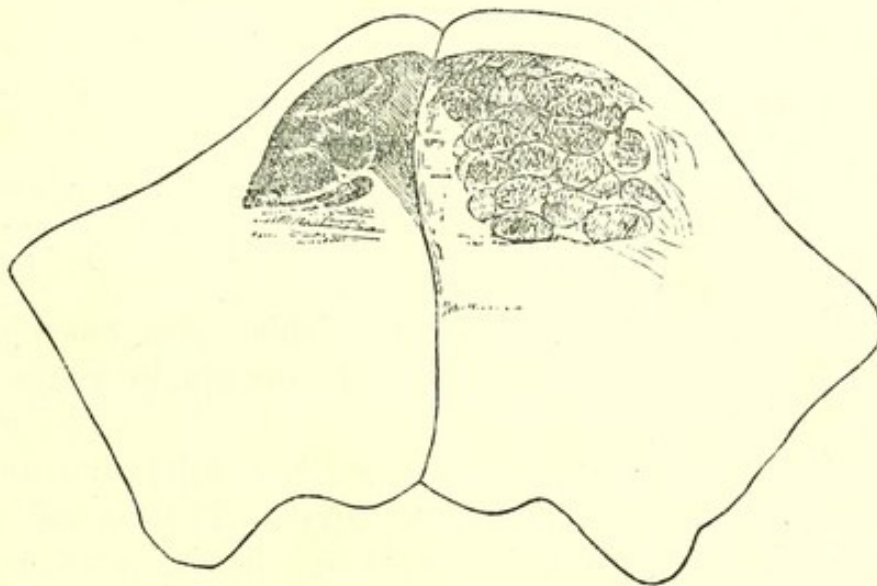


Fig. 229.—Pons in descending sclerosis. The pyramidal tract on the left side is seen to be shrunken and in a state of grey degeneration.

which also pass in and separate the bundles (Figs. 216 and 229). The course in the medulla oblongata and cord has already been traced.

In some cases of cerebral disease there is a degeneration on both



sides of the cord, the motor centres of one cerebral hemisphere being represented apparently on both sides of the body. The degeneration, however, is much more marked in the pyramidal tract belonging to the lesion.

Secondary degeneration also occurs in the fibres passing from the parts anterior to the motor area, namely, those in front of the ascending frontal convolution. These fibres pass through the anterior limb of the internal capsule and the inner part of the crusta of the peduncles, to end in the grey matter of the pons where their degeneration ceases. These fibres are probably prolonged to the cerebellum, forming communications between the frontal convolutions and the cerebellum (fronto-cerebellar fibres).

If of long standing there may be considerable shrinking in the parts affected by the secondary degeneration (see Fig. 229).

#### VII.—TUBERCULOSIS AND SYPHILIS OF THE BRAIN.

Both of these conditions present themselves in the form of tumours, but more especially tuberculosis. In its clinical aspects, indeed, tuberculosis of the brain substance has all the characters of a tumour, and it constitutes about half the cases of tumour of the brain.

##### 1. Tubercular tumour of the brain (*Scrofulous tubercle of the brain*).—



Fig. 230.—Tubercular growth in cerebellum. *a*, Main mass of tumour; *b*, part involving corpora quadrigemina; *f*, part projecting into fourth ventricle; *c*, third ventricle; *d*, pons.

As already indicated, this is a very common lesion, being the most frequent form of tumour of the brain. It is to be carefully distinguished from tubercular meningitis, with which it has scarcely anything in common, either in anatomical characters or clinical features.

The tubercular tumour occurs most frequently in young persons, and is often multiple. We have seen that tubercles are minute round bodies, but here in the brain substance we have solid tumours of a size increasing up to that of a hen's egg and larger. These massive tumours are composed of myriads of tubercles, along with the products of their degeneration. The greater part of the tumour is made up of a firm yellow caseous mass, which resembles very closely in appearance a scrofulous gland (see Fig. 230). Sometimes the cheesy mass is directly continuous with the brain substance, but usually there is a transparent grey zone outside it, and this gradually merges in the brain substance



around. This grey zone, when it exists, indicates that the tumour has been growing up till death, and is often distinctly composed of tubercles of characteristic form. Sometimes there are no rounded tubercles in it, and it forms simply a cellular zone. The grey tissue gradually merges in the brain substance, and both it and the brain substance show inflammatory conditions characterized by the presence of multitudes of round cells. The solitary tubercles are met with in all parts of the brain and spinal cord, but are most frequent in the cerebellum, and next to that in the cerebrum.

The tumours in the cerebrum or cerebellum are almost always near the surface, and they sometimes lead to a local or general tubercular meningitis. In the cerebellum they may be so large as to replace almost the entire substance of a lobe (see Fig. 230).

2. **Syphilis of the brain and its membranes.**—Syphilis attacks the brain chiefly in conjunction with the membranes. It is very rare to meet with a syphilitic tumour in the brain substance without some connection with the surface, and probably the lesion, in almost every case, begins in the membranes. The lesions are those of the tertiary stage, and

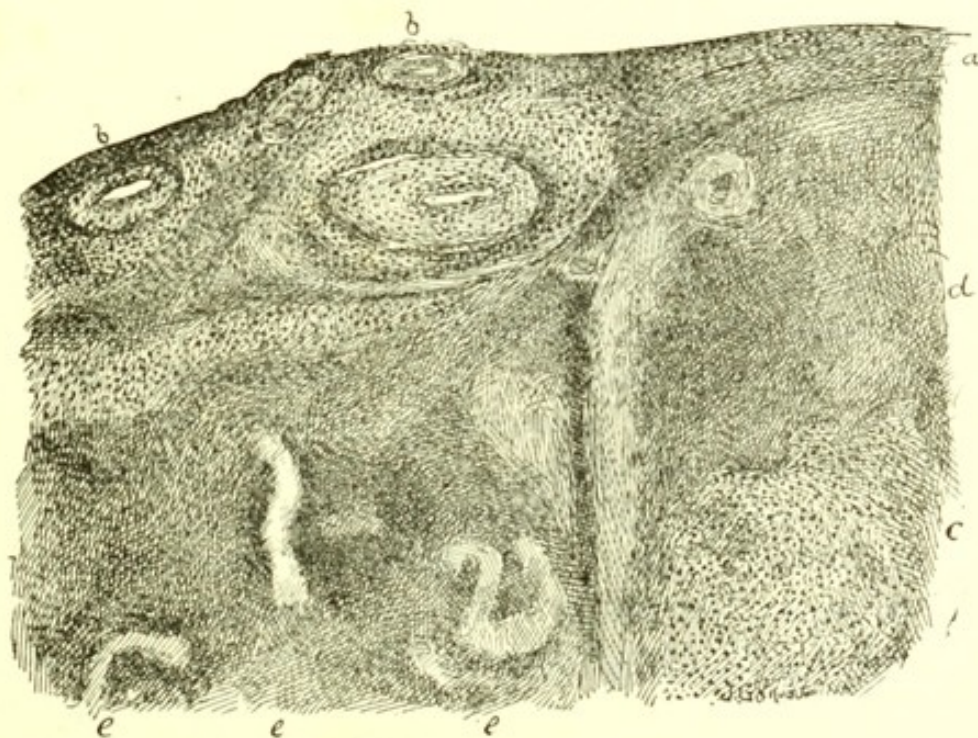


Fig. 231.—Gumma of membranes affecting brain. *a*, Thickened membranes adherent to brain; *b*, *b*, arteries with syphilitic endarteritis; *c*, *c*, round-celled tissue in brain substance; *d*, caseous structures, involving occluded capillaries *e*, *e*, *e*.

have the characters already described. They are, the gumma associated with more or less inflammation around, and the affection of the arteries.

The **Gumma** occurs as a grey or yellow mass, or is variously composed of both grey and yellow material. The grey structure consists of more or less fresh granulation tissue, while the yellow is due to caseous necrosis, and hence is older. The caseous part consists of granular



structureless matter, in which may be recognized some traces of the original tissue, along with fat and blood-pigment. The mass presents these two constituents in varying abundance (see Fig. 231), and the outline of the tumour is indefinite, shading off on the one hand into inflamed and thickened membranes, and on the other into the brain substance infiltrated with round cells.

Sometimes the gumma obviously grows from the **Soft membranes**, the pia-arachnoid. The membranes are thickened and matted, and adherent to the brain beneath. From their under surface rounded masses bulge against the brain substance, adhering to it and causing destruction. There is usually a zone of softening at the advancing margin of the tumour, and there may be considerable areas of softening deeply in the brain substance due to disease of the arteries in connection with the gumma (see below). In other cases the gumma is seated chiefly in the **brain substance**, but it is superficial, and the soft membranes are

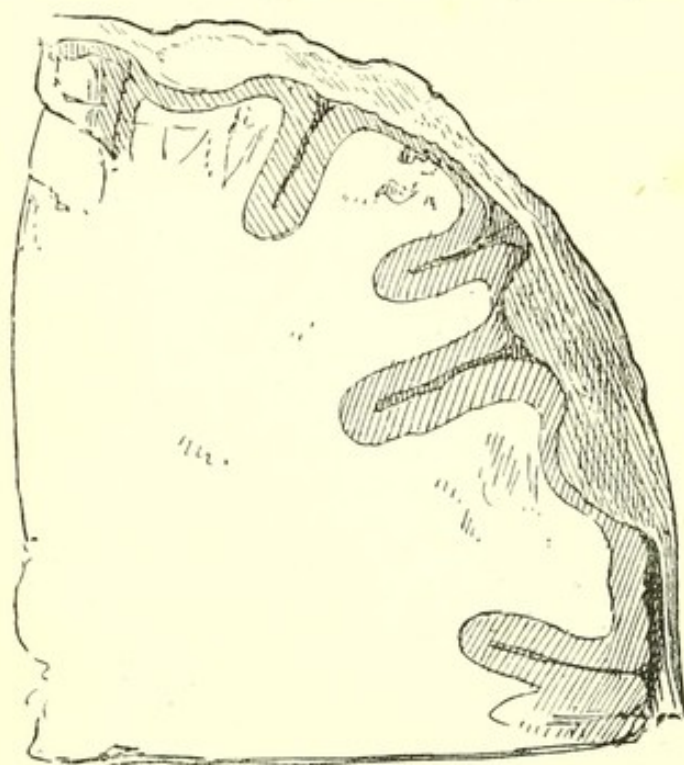


Fig. 232.—Gummata of dura mater impinging on and causing atrophy of the convolutions. (From case in Western Infirmary Museum.)

involved in the process. On the other hand, the **Dura mater** may be the primary seat of the tumour, and here we have tumours of considerable size (see Fig. 232), as large as a pigeon's or even a hen's egg. The dura mater is thickened and adherent to the soft membranes, and the tumour is usually to a large extent caseous. These tumours have a special tendency to form in the duplicatures of the dura mater, such as the falx cerebri. The projecting tumours impinge on the

brain substance like those of the soft membranes, but they also, when seated over the surface, project against the bone, producing erosion which may be considerable (*dry caries*).

The gumma may occur at any part of the convexity of the brain, and is also frequent at the base, affecting especially the optic chiasma, pons, and crura. By the latter it may extend to the optic thalamus, but otherwise scarcely attacks the basal ganglia. It not infrequently involves the cerebral nerves as they issue from the brain, especially the optic, and those emerging from the peduncles and pons (third, fourth, and fifth).



It has been already stated that inflammation and thickening of the membranes occur around the gummata, but sometimes a condition exists which may be designated a **Gummatous meningitis**. The gummatous tissue is formed more diffusely in the form of a grey gelatinous layer, covering a considerable surface at the base or on the convexity. In the former case there may be some resemblance to tubercular meningitis. In the latter, the whole membranes are thickened and matted together by gummatous tissue, and the brain substance beneath is softened. If, under treatment or otherwise, the proper gummatous tissue disappears, then the thickened membranes may present the appearance of an old simple inflammation.

**Syphilitic disease of the blood-vessels**, although not confined to those of the brain, is most frequent and most characteristic in them (see p. 196). The lesion is, in many cases, simply a part of the phenomena in connection with a gumma and presents itself in a number of small arteries, as an endarteritis obliterans (see Fig. 231, *b, b*). But it also occurs as an isolated affection of the larger cerebral vessels, chiefly those which contribute to the circle of Willis. The appearances are like those of atheroma, but the affection is more localized, occurring mostly in a few patches and not, as in atheroma, extending to a large number of branches. The patches are greyish white and opaque, and the wall is firm, so that the vessel preserves its cylindrical shape. The newformation has a dense character, and as it consists in thickened internal coat, the calibre is diminished or even obliterated if thrombosis ensue.

The lesion is a gummatous newformation in the wall of the artery. It has its seat, according to Heubner, especially between the elastic lamina and the endothelial layer. The newformation consists of granulation tissue and rudimentary connective tissue in the form of spindle and stellate cells. The lesion is apt to extend some distance along vessels, sometimes involving communicating branches.

An important result of this affection of the arteries is local **Softening of the brain**. This may occur either in relation to a proper gumma, or independently, from the affection of an artery alone. It is due to the narrowing of the calibre and occlusion, and has the same nature as softening from embolism, etc. The softening may be seated at some distance from the gumma or arterial lesion, especially where a long artery is distributed without intervening branches. According to Heubner the lesion affects most frequently the carotid and its branches, especially the middle cerebral, so that we may have softenings in the basal ganglia as in embolism.

The **Functional effect** of these various lesions depends largely on their **Localization**. The gummata and resulting inflammations of the convexity, occurring as they do on the surface, will, at first, irritate the surface of



the convolutions. If a motor part be affected then there will be muscular spasm, often progressive and ending perhaps in general convulsions of a quasi-epileptic character. Any sort of local spasm may be produced according to the particular part irritated. If the lesion be over a sensory part there will be subjective sensory impressions. We have already seen that the brain substance commonly gets involved in the gumma, and that outside it there is also softening of the brain. This destruction of brain substance involves loss of function, namely, paralysis or loss of sensation; but there will still be irritation of the marginal parts, and though paralyses and anæsthesiæ may develop, the signs of irritation generally remain prominent. At the base the gumma is very apt to involve motor tracts. The motor fibres of the pons and peduncles are anterior, and are therefore more exposed to the inroads of the tumour. But these paralyses are apt to be complicated by the lesion extending to the cerebral nerves which issue here, mainly the optic, third, fourth, and fifth, and so we have very complex conditions brought about, such as crossed paralysis, etc. Again, the syphilitic disease of the arteries occurs most frequently in the carotid and middle cerebral, and the resulting softening is usually most manifest in the region of the corpus striatum and motor convolutions. In this way a hemiplegia may be produced which imitates that from embolism. It will be seen that a great variety of symptoms may be produced from syphilitic lesions of the brain and its membranes.

*Literature.*—HEUBNER, *Die luetische Erkrankung der Hirnarterien*, 1874; GREENFIELD, GOWERS, HUTCHINSON, GOODHART, and others in *Path. trans.*, xxviii., 1877, pp. 249-360; GREENFIELD, *Path. trans.*, xxix., 1878.

#### VIII.—TUMOURS AND PARASITES OF THE BRAIN.

Tumours are of frequent occurrence in the brain, and produce secondary consequences of great importance. These secondary results are mostly related to the fact that the brain is enclosed in a rigid case, and that either by their own bulk or otherwise the tumours are liable to increase the contents of the cranial cavity. Hence there is pressure exercised on the brain. In addition to this there are various destructive effects produced. The secondary effects are, from a practical point of view, the most important consequences of the presence of tumours, and as these effects are, on the whole, similar, whatever the nature of the tumour and whether it be primary or secondary, intra-cranial tumours form a consistent group of lesions. In this view tubercular and syphilitic tumours are on a similar level with other forms.

**Pressure effects** are of great importance. Any tumour in its growth will press on the parts in its immediate neighbourhood, and the usual result is **Softening** of the brain substance around, the proper tissue-



elements being destroyed. When the growth is slow there may be little pressure, and the nerve structures may partly persist around the tumour or even be contained in its peripheral parts, if the growth is an infiltrating one like the glioma. The pressure is also exerted at a distance. The tumour itself by its bulk added to that of the cranial contents increases the intracranial pressure. The action of the pressure will depend somewhat on locality. The dense membranous septa, the falx and the tentorium, but especially the latter, limit its action. A tumour of the cerebellum being confined by the tentorium will often cause compression of the parts beneath the latter, and the pons is not infrequently damaged by such tumours (Gowers). Again, tumours often lead to **Hydrocephalus**, or accumulation of fluid in the ventricles, and this raises the general intracranial pressure. This occurs chiefly when they press on the transverse fissure so as to obstruct veins or lymphatics there (see further on under Hydrocephalus). There may be considerable thinning of the bones from pressure either with or without hydrocephalus, and sometimes even a gap in the bones. In children the sutures may be widened by the intracranial pressure. The relation of increased pressure to affections of the optic nerve is referred to further on.

**Inflammation** is a frequent result around a tumour. The irritation of the tumour frequently leads to prominent symptoms during life, and its presence causes structural changes. The brain substance in some cases shows sclerosis, although softening is more frequent. The membranes on the other hand are usually thickened by chronic inflammation, over the seat of the tumour and even at a distance. In syphilis the inflammation of the membranes is very great, and in tuberculosis there is sometimes an associated tubercular meningitis.

**Glioma.**—This form of tumour is peculiar to the nervous system and is of frequent occurrence there. As its tissue somewhat resembles brain substance in consistence, and as the tumour is an infiltrating one, its boundaries are frequently very indefinite and it may even appear as a simple local enlargement of the brain. Some cases described as instances of local hypertrophy are of this latter kind.

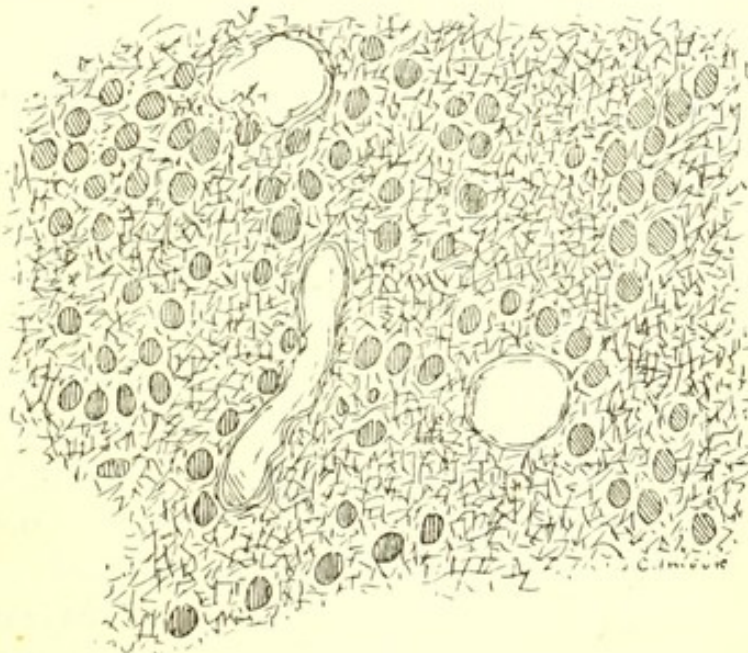


Fig. 233.—Glioma of brain.



On section the tumour generally shows some difference in colour from the brain substance into which it merges at its boundaries. It is generally redder than the white substance, and sometimes deeply red from excessive development of vessels. There is not infrequently hæmorrhage in the substance of the tumour, and this may be so considerable that the whole softened mass has the appearance of an extravasation of blood, and may be mistaken for this.

In its structure the glioma consists of cells usually round and of small size. These are enclosed in a granular, homogeneous, or finely fibrillated inter-cellular substance (see Fig. 233), which is usually very soft and delicate. Sometimes the inter-cellular substance has a distinctly fibrous character, and the tumour is then very firm. Sometimes the intercellular substance has a mucous character, in which case the tumour may be designated a **Myxo-glioma**. On the other hand the tissue may become very cellular and the tumour assume the characters of the sarcoma.

The glioma usually grows slowly. When it reaches the surface there are sometimes defined growths formed, which may be pedunculated. There may be small separate growths formed on the surface (Gowers).

Ziegler describes under the name Neuro-glioma tumours composed not only of neuroglia, but also of ganglion cells and nerve fibres. Such tumours are due to errors of development, and are congenital. They are better regarded as cases of Heterotopia of the brain substance (see under Malformations of the brain).

The **Psammoma** is not a frequent tumour of the brain substance, being much more common in the choroid plexus and dura mater. But

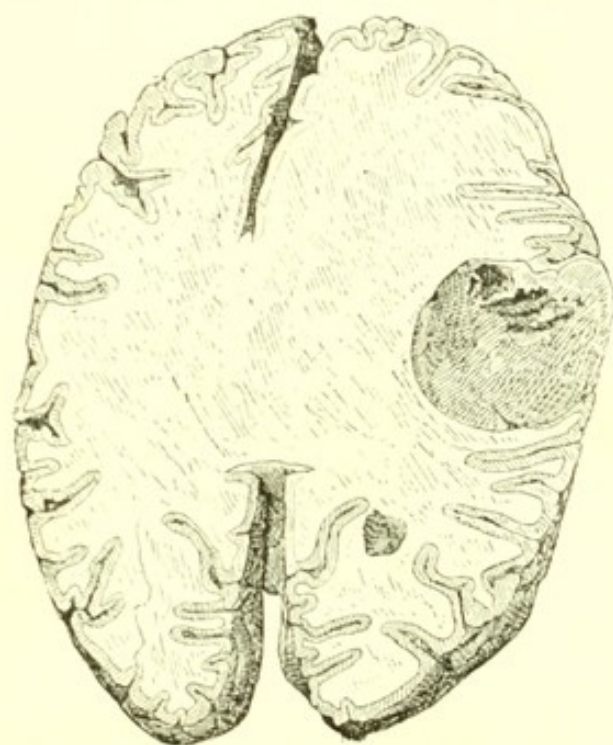


Fig. 234.—Spindle-celled sarcoma of brain. It forms a defined tumour. The affected hemisphere is considerably enlarged by its presence.

it does occur in the substance of the brain, as in the case from which Fig. 79, p. 245, is taken. Here a hard calcareous tumour, which was with difficulty incised, measured  $\frac{3}{8}$  by  $\frac{1}{4}$  of an inch, and to that extent completely replaced the brain substance. It was seated at the surface of the convolutions. As shown in the figure it was composed of the characteristic sand granules held together by a very small quantity of connective tissue. These bodies also occur in other tumours, chiefly the Sarcomas.

**Sarcoma.**—The sarcoma of the brain substance is a circumscribed tumour, in this respect contrast-



ing with the glioma (see Fig. 234). It occurs in the midst of the brain substance, and usually in a more or less rounded form, being free to expand in all directions on account of the softness of the tissue around. The round-celled sarcoma is the commonest form, but spindle-celled tumours and those with variously shaped cells are not infrequent. The sarcomas vary in consistence, the softer ones being usually round-celled. They may soften in the central parts and form cysts. Most demarcated tumours of the brain, excluding tubercular masses, are sarcomas.

**Myxoma** is not a frequent tumour of the brain. The **Cylindroma** is also of occasional occurrence (see p. 259).

Of the other tumours, Osteomas are occasionally seen. Lipomas are very rare. Angioma is also rare.

**Cancer.**—Primary cancer occurs in the substance of the brain, but is perhaps usually connected with one of the ventricles, probably deriving its epithelial elements from these. In the case figured (see Fig. 235) the tumour had its seat chiefly in the fourth ventricle and

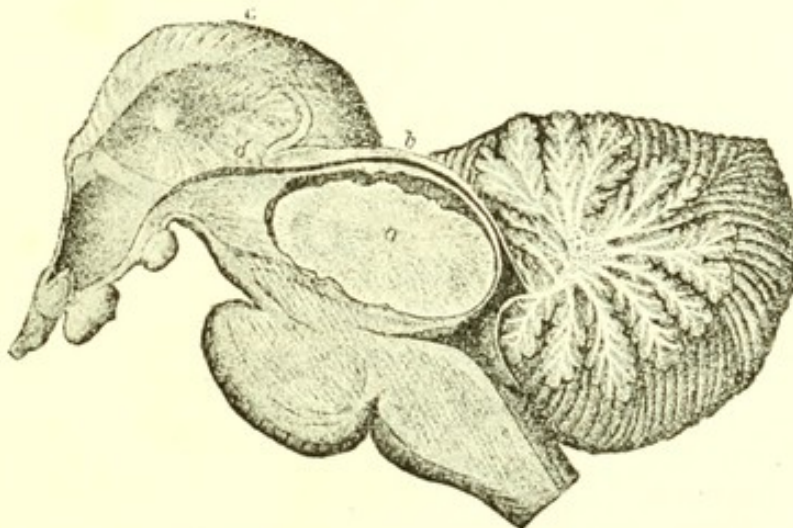


Fig. 235.—Primary cancer of the brain. *a* Tumour. *b*, Corpora quadrigemina stretched over it. The aqueduct of Sylvius is visible between tumour and corpora quadrigemina; the tumour bulges into fourth ventricle. *c*, Thalamus opticus. *d*, Third ventricle

aqueduct of Sylvius, bulging into these. The tumour consisted of a vascular-stroma and cylindrical cells which impinged on the brain substance.

The **Cholesteatoma** or **Pearl tumour** is a rare form of tumour which occurs chiefly in the membranes but is also met with in the substance of the brain. It consists of a cyst containing epidermic cells which have a glancing character and are arranged in rounded masses. The tumour has a soapy appearance. It probably originates from the epithelium of the spinal canal or ventricles, and is therefore an epithelioma. Ziegler mentions that small hairs may be present in the tumour.

**Secondary cancer** is not common in the brain, but it may form single or multiple tumours. In a case recorded by the author in which the primary tumour was presumably in the lung, the secondary tumours



occurred chiefly in the form of **Cysts**, of which there were twenty-four discovered in the brain (see Fig. 236). In one of the tumours which was not yet entirely cystic the cancerous structure was apparent (see Fig. 237).

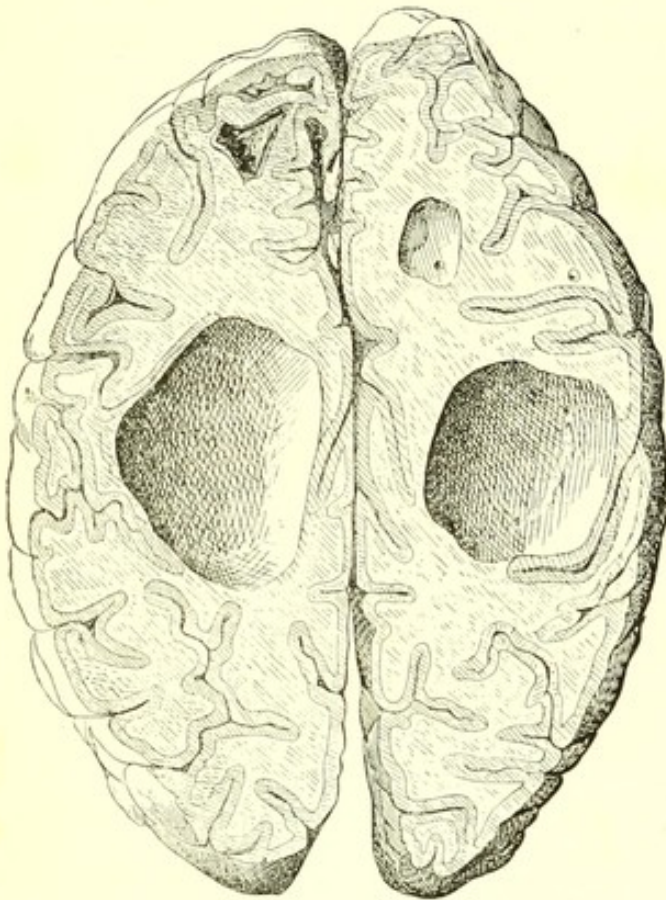


Fig. 236.—Secondary cancer in brain. Cysts developed.

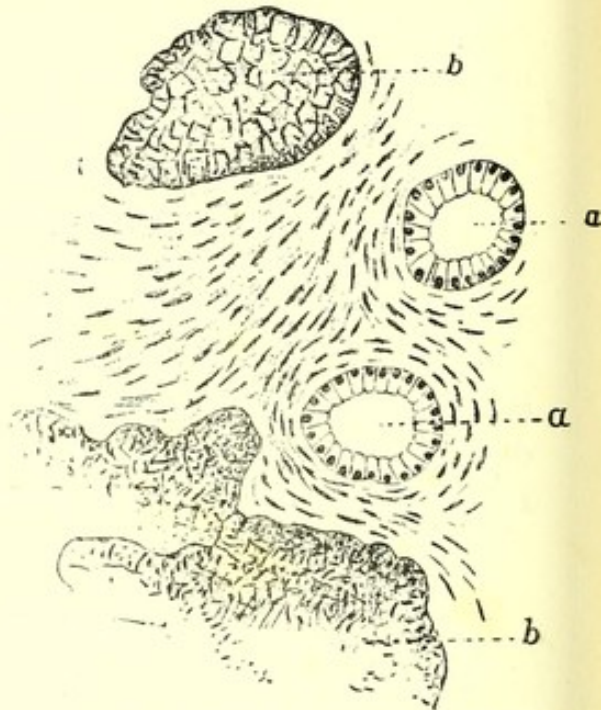


Fig. 237.—Secondary cancer in brain developing cysts. *a a*, duct-like canals, *b b*, larger spaces, the beginnings of cysts.

**Parasites.**—The **Cysticercus** of the *tænia solium* has occasionally its seat in the brain, mostly in the cortical portions, but it may be in any situation. It forms a small round structure enclosed in a connective tissue capsule. Inside this the proper stratified membrane of the parasite, with the head (see Fig. 123, p. 328) and other structures, are to be found. Sometimes the parasite is dead and calcified. As a rule it produces little disturbance, and is often found by accident after death. If situated in the peduncular portions it may produce serious symptoms, but even here its slow growth and small size generally allow of accommodation. Sometimes several cysticerci, as many as 10 to 20, have been found.

The **Echinococcus** more rarely develops in the brain, forming its usual cysts, which are generally in the substance of the cerebral hemispheres. They also occur in other parts of the brain, and between the dura mater and the bone.

**Literature.**—GOWERS, l. c., ii.; HALE WHITE, *Guy's Hosp. Rep.*, xliii., 1886; VIRCHOW, (*Glioma*) *Geschwülste*, ii. 123; Various authors and tabulations of 54 tumours, *Path. trans.*, 1886, xxxvii., p. 6-89; COATS, (*Cancer*) *Path. trans.*, 1888, xxxix., p. 5, (*Multiple cysts*) *do.*, p. 326; EPPINGER, (*Cholesteatoma*) *Prag. Vierteljahrschr.*, 1875; ROKITANSKY, (*do.*) *Handbuch*, ii.; MULLER, *Virch. Arch.* viii.



## SECTION III.—CONTINUED.

D.—THE MEMBRANES AND CAVITIES OF THE BRAIN  
AND SPINAL CORD.

INTRODUCTION. I. ACCUMULATION OF CEREBRO-SPINAL FLUID. 1. *Œdema of membranes.* 2. *Hydrocephalus, acquired and congenital.* 3. *Dropsies of central canal and meninges of cord.* (a) *Hydrorhachis interna, hydromyelia, syringomyelia.* (b) *Spina bifida, in various forms; the tumour of spina bifida; condition of the cord; spina bifida occulta.* II. MENINGEAL HÆMORRHAGE. *Hæmatoma of dura mater; 2. Hæmorrhages in soft membranes, especially the infantile form.* III. INFLAMMATIONS. MENINGITIS. 1. *Pachymeningitis; 2. Leptomeningitis, including (a) simple, acute, and chronic, (b) epidemic cerebro-spinal, and (c) tubercular meningitis.* IV. TUMOURS OF THE MENINGES. V. AFFECTIONS OF THE PINEAL AND PITUITARY BODIES.

INTRODUCTION.—The dura mater is a dense connective-tissue membrane which has much more intimate connections with the bones than with the brain and spinal cord, forming, in fact, a periosteum to the bone. On the internal surface there is no separate arachnoid membrane, but the dura mater is less vascular here than in its external layers. The space between the dura mater and the surface of the brain is sometimes described as a serous cavity analogous to the pleura or peritoneum, but in its pathological relations, at least, it is very different, and it is better designated as the **Subdural space**. The internal surface of the dura mater and the external surface of the arachnoid are defective in vessels, and are hence little liable to inflammations. This cavity forms, in fact, a kind of barrier to the propagation of inflammations either from without or within. Hence inflammations of the external surface of the dura mater seldom extend to its internal surface, and it is very rare to meet with an inflammation affecting the whole subdural space. On the other hand the arachnoid and pia mater really form one membrane, which is partly in two layers. The deep layer or pia is closely connected with the surface of the brain or spinal cord, following the various irregularities of the surface. This deep layer is connected with the superficial one, the arachnoid, more or less intimately. In the sulci between the convolutions there is loose connective tissue with many interstices, constituting the **Subarachnoid space**. But on the summits of the convolutions the two are intimately united. The free surface of the arachnoid is, like that of the dura mater, defective in vessels, and inflammations of the deeper layer scarcely ever extend to this surface. We have thus to distinguish the dura mater connected with the bone, the subdural space, the pia-arachnoid forming the proper meninges of the brain, and the subarachnoid space.



The **Cerebro-spinal fluid** is contained partly in the ventricles of the brain and central canal of the spinal cord on the one hand, and partly in the subarachnoid space on the other. Between these two there are important communications. The pia mater is prolonged into the lateral ventricles through the transverse fissure, as the velum interpositum and choroid plexus, thus forming a somewhat free communication. There is also an open communication between the lower part of the fourth ventricle and the subarachnoid space, by the foramen of Majendie. The subarachnoid space and the ventricles of the brain with the central canal of the spinal cord thus form a continuous system of lymph spaces, and the cerebro-spinal fluid circulates in them. In addition to this, the subarachnoid space communicates with lymph spaces around the vessels and ganglion cells, the **Perivascular** and **Periganglionic** spaces. It is important to observe that the subarachnoid space does not communicate with the subdural space.

#### I.—ACCUMULATION OF THE CEREBRO-SPINAL FLUID IN THE MEMBRANES AND CAVITIES.

It has been already pointed out that the subarachnoid space, with the ventricles and central canal of the spinal cord, forms a single system of lymph spaces which intercommunicate. The lymph in these spaces may accumulate, and so lead to various forms of œdema and dropsy of the brain. In this connection it is important to remember that the skull is a closed cavity, and any increase in the fluid in these spaces implies a decrease in the quantity of blood in the vessels or of the brain substance itself.

1. **Œdema of the membranes.**—There may be **General œdema** of the membranes and spaces, perhaps including the perivascular spaces in the brain substance. This sometimes takes place in Bright's disease, and may occur along with an œdema of other parts, or develop in a more isolated manner. The pressure of the fluid in this case was asserted by Traube to be the cause of the symptoms usually called uræmic, and in some cases it may contribute to the nervous disturbances of that condition.

Then there is an **Œdema ex vacuo**. When the brain shrinks, as we have seen in **General paralysis**, or when it undergoes a more simple atrophy as in **Senile atrophy**, there is a serious loss of substance. The loss of substance thus produced may be compensated by thickening of the cranium or by augmentation of the cerebro-spinal fluid. Thickening of the cranium takes place only to a limited extent, and the space is chiefly filled up by fluid. The ventricles dilate greatly (*hydrocephalus*), and the subarachnoid space is highly œdematous. It is to be particularly observed that there is seldom any excess of fluid in the subdural space, but that the soft membranes, especially between the atrophied convolutions, are highly œdematous.

2. **Hydrocephalus.**—This term expresses a massive accumulation of



fluid inside the skull. The fluid is, in the great majority of cases, in the ventricles, more especially the lateral ventricles, but in some cases it is in the subdural space. Hence the terms **Internal** and **External hydrocephalus**. The latter condition scarcely occurs unless there be a congenital defect in the formation of the brain, it may be even a perforation or absence of the corpus callosum, so that the ventricles communicate with the subdural space.

External hydrocephalus is sometimes used as equivalent to œdema of the membranes, but this is an incorrect use of the term, which really expresses a considerable collection in a cavity.

(a) **Acquired hydrocephalus**.—Apart from the form *ex vacuo*, hydrocephalus as met with in the adult is mostly related to a definite disturbance of the circulation in the brain. The ventricles are supplied with fluid chiefly through the choroid plexus, and any obstruction to the veins here is liable, by causing a passive hyperæmia, to lead to an excessive exudation. It seems also as if the choroid plexus, with its villi, acted as a drain to the ventricles, the fluid passing into its lymph spaces, and so either into the veins or outwards to the subarachnoid space. If the veins are obstructed they can no longer aid in absorption, but on the contrary their obstruction causes, as we have just seen, excessive transudation from the capillaries. Again, if the lymphatic vessels and spaces are filled up, this will interfere with the absorption, so that dropsy may arise in this way. We have therefore two classes of cases, one in which the veins and another in which the lymph spaces are obstructed.

The veins of the choroid plexus gather themselves into the great **Veins of Galen**, which pass outwards between the corpus callosum and corpora quadrigemina to open into the straight sinuses. In cases of obstruction of these veins there may be a rapid accumulation of fluid in the ventricles, and death may even result from the sudden increase of intracranial pressure. These veins may be obstructed by pressure from without, especially by tumours of the cerebellum, or by thrombi occupying their calibre. In most cases the thrombus originates in the sinuses, and grows into the veins of Galen, but Newman has recorded a case in which these veins alone seemed to be the seat of thrombosis.

In **Tubercular meningitis** hydrocephalus is nearly constant, but it is rather difficult of explanation. Tubercles are generally found on the vessels of the choroid plexus, and the exudation in the ventricles is sometimes inflammatory in character. This is rarely the case, however, as the exudation, consisting of a clear pellucid fluid with little albumen, and of a low specific gravity (about 1010), presents the characters rather



of a transudation than an inflammatory exudation. The explanation is probably to be found in part in the exudation outside the ventricles. This is generally abundant where the velum interpositum passes into the ventricles in front of the cerebellum, and here, besides filling up the lymph spaces, it is liable to cause pressure on the veins of Galen as they pass through the comparatively narrow isthmus. This is the more likely to produce hyperæmia and exudation from the fact that the arteries reach the plexus by a different route, and are not so liable to be pressed on.

(b) **Chronic hydrocephalus. Congenital hydrocephalus.**—It has been pointed out in a previous section that various malformations affecting brain and cranium are due to dropsy of the cerebro-spinal canal during early foetal life. A similar dropsy may occur at later periods of intra-uterine life, or even after birth. The condition may be designated chronic hydrocephalus, which, it will be understood, is often congenital, and may by the enlargement of the head produce serious difficulty in delivery. No proper explanation of the dropsy has been offered. It is said that chronic hydrocephalus often goes along with rickets, and it has been suggested that as a rickety skull is more yielding than a normal one, too little pressure is exercised on the brain, and the accumulation of fluid is allowed. This is not, however, a sufficient explanation, especially of the severe congenital cases. The probability is that the disease depends on a congenital defect in the apparatus for the secretion and absorption of the cerebro-spinal fluid.

The dropsy may be general, affecting all the ventricles, but the fourth is often but little dilated, the fluid being mainly in the lateral and third ventricles. Sometimes the lateral ventricles are alone distended, the foramen of Munro being closed. Sometimes the ventricle of the septum lucidum is obviously distended as it lies between the two lateral ventricles. As the accumulation increases the convolutions are flattened out and they may be completely unfolded, the brain forming a smooth globe over the greatly distended ventricles. The distension may be so great as to cause thinning of the covering brain substance to an extraordinary degree, leaving little between the pia mater and the fluid. The surface of the ventricles (ependyma) is usually somewhat thickened, giving a kind of leathery membranous lining to the cavity, sometimes with granular elevations, especially on the surface of the corpus striatum and fourth ventricle. There may even be rounded prominences consisting of grey brain substance (Virchow). The fluid is clear and pellucid, and has a low specific gravity (1001 to 1009). In some very rare cases the corpus callosum and its pia-arachnoid have given way, the fluid coming thus to the surface and filling the cavity of the dura mater (Hydro-



cephalus externus). In that case the hemispheres are folded aside, and the central parts of the brain are exposed, the brain being as a whole pressed down towards the base.

With this great dilatation of the ventricles the head is greatly enlarged, and as the bones are more yielding in some parts than others, an alteration in shape occurs. The fontanelles and sutures are widened and their closure greatly delayed. The frontal bone is pushed forward so that the forehead rises perpendicularly or overhangs the eyebrows; the parietals bulge laterally, and the occipital bones are pushed backwards. The head in this way becomes greatly increased in circumference, while it is usually much flattened at the vertex. The bones of the face, even though they are of normal size, look dwarfed beside the enlarged cranium, and the face has a pinched look. The eyeballs are rendered prominent by the pressure on the roof of the orbit, and enlarged veins are generally seen beneath the thin skin of the head.

Although there is this great thinning of the brain substance, it is remarkable how the functions may be retained. A child in this condition may remain very intelligent, and when recovery occurs may pass through life with no permanent defect in the functions of the brain. The yielding of the skull seems to prevent any such excess of pressure as to seriously damage the brain, which has a remarkable power of accommodating itself to alterations in position of its parts. If recovery takes place, the skull to some extent collapses, the fontanelles and sutures close, often with the formation of additional centres of ossification, forming Wormian bones in the sutures. But the fontanelles are late in closing, and the cranium retains somewhat of the hydrocephalic shape during life.

A **Partial dropsy** of the ventricles is of occasional occurrence, either as a congenital or acquired condition. One lateral ventricle may be distended, or even one horn. There may also be a dropsy of the third ventricle alone, or of the ventricle of the septum lucidum. These partial dropsies will cause displacements according to their situation.

**3. Dropsies of the central canal and Meninges of the spinal cord (Hydrorhachis, Spina bifida, etc.).**—The affections to be here considered are almost all congenital, and they lead for the most part to malformations of the cord.

The central canal of the cord presents normally considerable variations. In its typical form it is a narrow canal lined with cylindrical epithelium. The tissue immediately around it is much more cellular than the rest of the cord, this being due to the fact that it consists almost alone of neuroglia, without nervous elements, in this respect comparing with the ependyma of the cerebral ventricles. From this



typical condition the variations consist in different conditions of the canal itself and of the surrounding tissue. The canal may be widened either generally or in the transverse or median direction. On the other hand the canal may be obliterated and its position only indicated by the cellular area of neuroglia. These are all within normal limits.

The pathological conditions of the central canal consist mainly of dilatations, and these again present certain relations with the important condition, spina bifida.

(a) **Hydrorhachis interna, Hydromyelia, Syringomyelia, Cysts of the cord.**—The term Hydrorhachis corresponds with hydrocephalus, and expresses a dropsy of the cord, either of the central canal (H. interna) or of the meninges (H. externa). Dropsy of the central canal is also designated Hydromyelia and Syringomyelia. The former term is applied to cases where the dilatation is obviously congenital, while the latter is used where it is met with in the adult. Syringomyelia, is however, probably always congenital in its origin, so that it may be said that it originates in a hydromyelia. These terms are applied to all cases of cavities or cysts in the cord filled with serous fluid whether they are demonstrably due to dilatation of the central canal or not.

It will be remembered that the central canal is formed by the arching backwards of the medullary plates (see p. 43), so that the closure is posterior. The central canal will, therefore, at first lie posteriorly, and it is by a gradual coalescence of the posterior portions in the middle line that the canal takes its usual position. Dropsy occurring in foetal

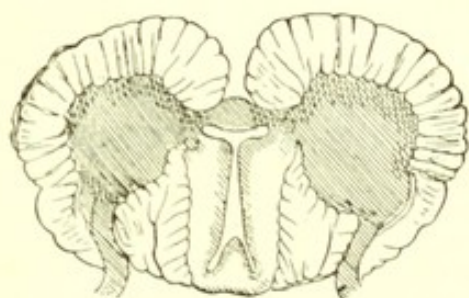


Fig. 238.—Hydromyelia. Enlargement of central canal backwards.

life is liable to cause a permanent enlargement backwards as shown in Fig. 238. The dilatation may vary greatly at different levels, so as to lead sometimes to a cystic appearance in the cord.

In some cases cavities exist in the cord without obvious connection with the central canal, but as these are always in the posterior parts of the cord, they probably take origin in portions of the original canal, which in the process of coalescence have become isolated. These not infrequently extend into the posterior cornua, and may be double (Fig. 239) or single (Fig. 240).

An interesting peculiarity of these various conditions is that the serous cavities produced are surrounded, like the normal central canal, with neuroglia, which in some cases is in considerable excess. The excess may be so great as to amount almost to a tumour (glioma). The importance of this is emphasized by the occasional co-existence of syringomyelia with tumour of the cord, and even with multiple



tumours (see Gowers). Gliomata of the cord mostly originate behind the central canal, even when there is no dilatation of the latter.

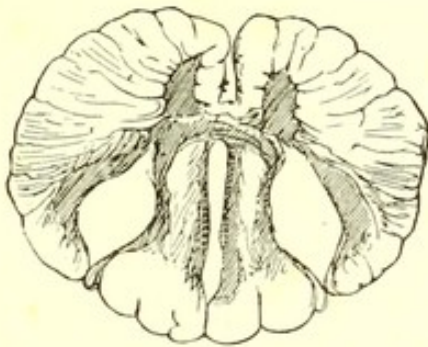


Fig. 239.—Syringomyelia. Cavities in each posterior cornu and in the middle line.

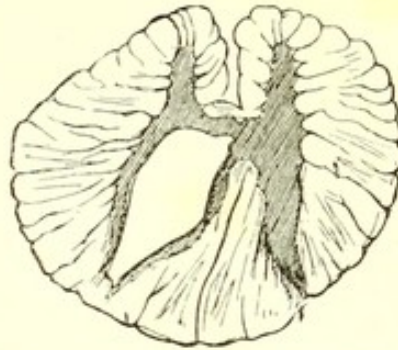


Fig. 240.—Syringomyelia. Cavity in one posterior cornu.

Hydromyelia and syringomyelia do not usually interfere with the function of the cord, but if the dilatation be excessive, or if there is a formation of tumour, there may be considerable damage.

(b) **Spina bifida.**—In the section on General Malformations, this condition has already been referred to. **Rhachischisis**, or spina bifida without tumour, has been sufficiently described (see p. 45), but the form associated with tumour requires fuller consideration.

The term spina bifida is by some limited to the tumescent form, which is therefore distinguished from rhachischisis, but this distinction cannot be fully carried out.

The **Tumour of spina bifida** forms a rounded swelling, which is usually situated in the lumbo-sacral region, but may be in any part of the column. There is usually a defect in the arches of the vertebræ, and the swelling protrudes posteriorly; but there are a few cases in which the bodies of the vertebræ are defective, and the swelling protrudes into the thorax, abdomen, or pelvis. There is sometimes no defect, or a very slight one, of the arches of the vertebræ, the protrusion passing between two adjacent arches.

The protrusion (Fig. 241) is covered with skin over the greater part of its surface, but in the central part there is often an area of membrane which contains none of the special structures of the skin, such as hairs and sebaceous glands. Inside the skin there are the membranes of the cord; but, according to Recklinghausen, the dura mater is frequently defective to a similar extent with the skin, so that the sac is formed of pia-arachnoid, which is usually considerably thickened. There is generally a dimpling or umbilication at the summit of the tumour (see *g'* in figure), and this sometimes amounts to a small canal which may directly communicate with the central canal of the cord. The umbilicus corresponds with an adhesion of the cord to the sac.



The contents of the sac are a pellucid fluid of a specific gravity of about 1007. It is really the cerebro-spinal fluid, with which the contents of the sac are continuous. Indeed, there is often associated with the spina bifida a hydrocephalus.

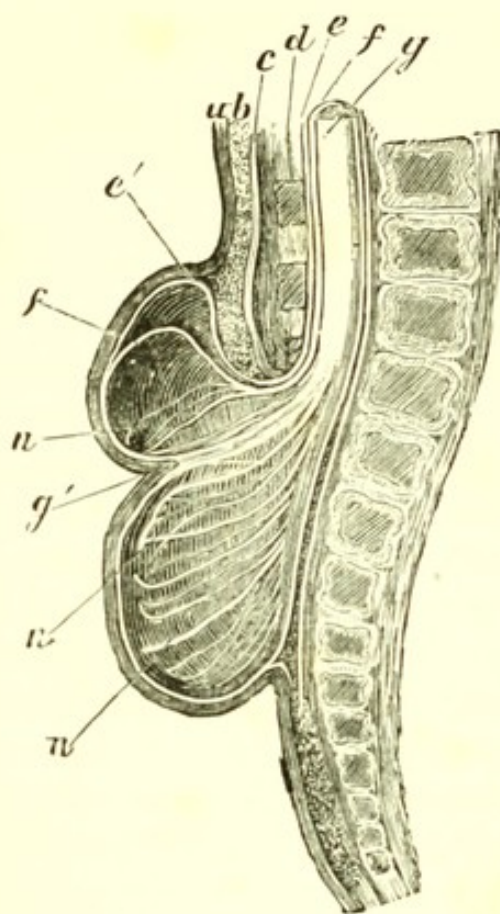


Fig. 241.—Section of a spina bifida of the lumbar region. *a, b*, cutis and subcutaneous tissue; *c*, fascia; *d*, spinous processes; *e*, dura mater, which passes into sac and becomes attached to the skin at *e'*; *f*, arachnoid, which passes into sac forming its internal lining; *g*, spinal cord, which also enters the sac and becomes attached to the skin *g'*, where it had a small opening; the attachment forms a dimple in the sac; *n, n*, spinal nerves which pass from the cord round to the anterior aspect of the sac so as to reach their normal places of issue from the spinal canal. (VIRCHOW.)

Spina bifida implies a dropsy of the structures of the cord, and different names are applied according to the share which the various structures take in the protrusion, these names being constructed on similar principles to those in use for the brain and its membranes. We have (1) *Meningocele*, or protrusion of the membranes only; (2) *Meningo-myelocele*, or protrusion of membranes and cord together; and (3) *Syringo-myelocele*, in which the protrusion obviously consists of a dilated and closed central canal. This form is also called *Myelo-cystocele* (Recklinghausen). Of these three forms the second is much the commonest, although the last-mentioned form is probably more frequent than has been usually supposed (Cleland).

The spinal cord in all forms of spina bifida is prolonged downwards as far as the seat of the tumour. As the latter is usually in the lumbosacral region, this implies that the cord occupies the entire length of the canal as it does in the earlier periods of foetal life. As the cord begins to retract within the canal in the fourth month the affection must be earlier in its origin than that period. It is of interest that even in cases of meningocele where the membranes alone are protruded this elongation of the cord occurs.

With the exception of elongation, the cord in a small number of cases is not involved. The fluid may be entirely behind it as in meningocele, and it may be little altered. In that case the nerves also will be in front of the protrusion.

In *Syringo-myelocele* the central canal is distended backwards, and although the internal lining of the sac is really the expanded central



canal, yet the sac is as in meningocele, behind the nerves and the greater part of the cord.

In the majority of cases, however, comprising the **Meningo-myeloceles**, the spinal cord is actually involved in the sac (see Fig. 241), passing into it and attached to its wall, thus presenting various malformations and dislocations. The nerve-roots also which come off from the part of the cord involved take origin within the sac, and traverse it in order to reach their foramina. The posterior roots are necessarily more involved than the anterior. Sometimes the nerves take origin from the posterior wall of the sac, and the latter may present externally a double row of slight depressions from this origin.

The term *Spina bifida occulta* has been applied by Recklinghausen to an interesting case in which, without a tumour externally, there was elongation of the cord and defect of the vertebral arches. He supposes that there was here an early meningocele, which had subsequently shrunk. An important peculiarity in the case is the existence within the spinal canal and in contact with the cord of adipose and fibrous tissue and striated muscle, in such quantity as to form a distinct tumour (myo-fibro-lipoma). The muscle was in bundles like those of the multifidus spinæ, and he supposes that it may have arisen by a dislocation or transposition inwards of parts of these muscles by the defectively formed vertebral arches. Another peculiarity was the existence of an **excessive growth of hair** over the concealed spina bifida. It is not improbable that such lesions may be not infrequent.

The origin of spina bifida is to be referred to fœtal life. Looking to the series of forms of lesion from completely open spina bifida with anencephalus to meningocele, it seems necessary to connect the affection with dropsy of the cerebro-spinal canal. A further confirmation of this is the frequent co-existence of hydrocephalus. It can scarcely be supposed that a mere dropsy of the membranes, and still less a simple defect of the arches of the vertebræ, would fix the cord in the fœtal position, and we are induced to regard the lesion as primarily one of the cord itself. There may be a very early dropsy interfering with the closure of the canal, and so leading to rhachischisis. But after the closure there may be a dropsy leading to a local distension which may by and by rupture. If it does not rupture then there may be a spina bifida having the form of a syringo-myelocele. If it does rupture then the meningocele or the meningo-myelocele will result. The aperture may be ultimately a very narrow one or may become obliterated, and the central canal may form and even be of normal character. As a general rule, however, the fluid in the sac communicates freely with the cerebro-spinal fluid, and sudden withdrawal of the former sometimes leads to serious cerebral symptoms from the reduction of the cranial contents.

**Literature.**—*Hydrocephalus*—LEUBUSCHER, Path. d. Hirnkrank., 1854; HUGUENIN, (with full literature) in Ziemssen's Handb., xi. 1; WILKS, Guy's Hosp. Rep. 1860. *Hydrorhachis and Spina bifida*—LEYDEN, Virch. Arch., lxxviii.; LANGANS, do., lxxxv.; KRAUSS, do., c.; SCHULTZE, do., cii.; FÖRSTER, Missbildungen, 1865; VIRCHOW Geschwülste, i. and iii.; RANKE, Jahrb. d. Kinderheilk, xii.; W. KOCH, Mittheilungen, 1881; MORTON, On spina bifida, 2nd ed., 1887; CLELAND, in Morton, p. 32, also Jour. of Anat. and Phys., xvii. 257; Rep. to Clin. Soc. of London, 1885, xviii. 339; RECKLINGHAUSEN, Virch. Arch., cv. 243 and 374 (with full literature).



## II.—MENINGEAL HÆMORRHAGE.

1. **Hæmatoma of the dura mater.**—This is a condition concerning which considerable differences of opinion exist. It presents itself mostly in the form of a somewhat massive blood-clot covering the internal surface of the dura mater and compressing the brain substance. When the clot is more particularly examined it is seen to be not exactly free on the surface of the dura mater, but covered with a delicate membrane, which is continued beyond the clot on the surface of the dura mater as a thin soft layer. This membrane generally has a brownish colour, evidently from the blood colouring-matter, and it presents in its substance, as well as between the membrane and the dura mater, numerous smaller hæmorrhages. This condition is of somewhat frequent occurrence, particularly among the insane.

There are two views as to its true nature, and it is quite possible that there may be actually two diseases. According to Prescott Hewett, Huguenin, and others, a hæmorrhage into the cavity of the dura mater is the primary condition. It is undoubted that a hæmorrhage may lead to a condition resembling that described. In a case of aneurysm of one of the larger cerebral vessels, where bleeding had occurred into the subdural space some time before the fatal cerebral hæmorrhage, the author found a layer of soft tissue covering the dura mater and having much of the characters described above. In this and similar cases the coagulum on the surface of the dura mater becomes organized in the usual way, and vessels pass from the dura mater into the rudimentary tissue thus produced. These thin-walled vessels are specially apt to bleed for reasons to be presently referred to, and so there is hæmorrhage in the new-formed tissue and under it.

But many cases have a more spontaneous origin, and agree with the description which Virchow has given of **Hæmorrhagic pachymeningitis**. This disease begins in an inflammation of the dura mater, characterized by hyperæmia. The inflammation being chronic, the result is the formation of a soft membrane on the internal surface of the dura mater, owing to an inflammatory transformation of its internal layer. In its structure this membrane somewhat resembles mucous tissue, containing stellate and spindle-shaped cells in a matrix which gives a precipitate with acetic acid. In it there are large thin-walled blood-vessels in large numbers. The false membrane is easily lifted from the dura mater with forceps, and as this is being done numerous red threads are seen to stretch from it to the dura mater; these are the blood-vessels.

An explanation of the large size and tendency to rupture of these vessels has been suggested by Rindfleisch. To begin with, there is hyperæmia of the dura mater with



relaxation of the arteries. The normal capillaries being in a dense tissue will not dilate much, but the blood in them will be at a high pressure. The new-formed vessels, however, are delicate and lie in a soft tissue, and they communicate with the capillaries of the dura mater in which the blood-pressure is excessive. They are therefore very liable to dilatation, and although they have the structure of capillaries they are mostly three or four times as wide as ordinary ones. These vessels often rupture, so that there is frequent hæmorrhage into the soft membrane. But sometimes a more considerable hæmorrhage occurs, and the blood accumulating dissects up the membrane from the dura mater, rupturing fresh vessels as it advances. In this way a large flat clot as thick as the hand may be formed, the proper hæmatoma. It will be observed that this clot is still covered with the membrane, and it is quite unusual to find the blood escaping into the cavity of the dura mater.

If a fatal hæmorrhage does not occur, the new-formed membrane undergoes organization in the way of other inflammatory structures. It becomes more cellular and finally develops into connective tissue which coalesces with that of the dura mater. The disease, however, is apt to recur, and a fresh soft layer is formed which goes through the same stages, so that there may be several layers in different stages of transformation on the surface of the dura mater, the innermost layer having the characters described above.

This condition occurs chiefly over the convexity of the brain, and is stated to be mainly in the domain of the middle meningeal artery. An acute suppurative inflammation very rarely develops in connection with the hæmatoma.

2. **Hæmorrhages in the soft membranes.**—Most of these are secondary to some other lesion and they are usually of minor consequence. We have seen that aneurysms although situated in the meninges give rise, when they rupture, to cerebral hæmorrhage more than to meningeal. There are again numerous small hæmorrhages occasionally in anthrax, hæmophilia, scurvy, and ulcerative endocarditis. There are also hæmorrhages from injuries to the skull, especially when they involve lacerations of the brain. But there is one form which is primary and of considerable importance.

**Infantile meningeal hæmorrhage.**—In severe and prolonged labours, where the head is much compressed and there is obstruction of the vessels, hæmorrhage sometimes takes place on the surface of the brain. It is important because it may lead to permanent injury to the brain. The hæmorrhage is over the convexity or at the base, and in the former case it occupies chiefly the central region towards the middle line. The brain beneath is sometimes much injured, being lacerated and infiltrated with blood. When at the base, the hæmorrhage is chiefly in the posterior fossa, the blood lying on the pons, medulla, and cerebellum, and generally arising from a laceration of the cerebellum.



As the blood is absorbed there may remain a permanent atrophy of the parts which had been injured. There may be thus a depression over the central convolutions which are dwarfed and indurated, and the usual secondary degenerations may ensue.

During life, there may be little that is noteworthy in the child immediately after birth, but as the powers develop it may show motor weakness and spasms, along with more or less mental defect. There is often considerable improvement as life goes on.

**Literature.**—*Hæmatoma*—PRESCOTT HEWETT, Med. chir. trans., 1845; VIRCHOW, Würzb. Verhandl., 1856, vii.; HUGUENIN, Ziemssen's Handb., xi.; RINDFLEISCH, Path. histol. (Syd. Soc. transl.), 1873, ii. 302. *Infantile hæmorrhage*—LITTLE, Obstet. trans., 1862; M'NUTT, Amer. Jour. of Med. Science, 1885; GOWERS, Dis. of nerv. syst., 1888, ii. 380.

### III.—MENINGITIS—INFLAMMATIONS OF THE MEMBRANES OF THE BRAIN AND CORD.

1. **Pachymeningitis. Inflammation of the dura mater.**—Excluding the condition already described as connected with hæmatoma, the inflammations of the dura mater are, for the most part, secondary to affections of the bones or sinuses. In compound fractures of the skull, suppuration may extend to the dura mater, and in some cases to the soft membranes. Caries of the bones may have a similar result. Thrombophlebitis of the sinuses, especially in cases of disease of the ear, also gives rise to suppurative inflammation, sometimes associated with gangrene of the dura mater. (See under Abscess of the Brain.)

2. **Leptomeningitis. Inflammation of the pia-arachnoid.**—As the arachnoid and pia mater are closely connected and form virtually one membrane, they are always associated in their inflammations. The term **Meningitis** is commonly used without qualification to express inflammation of the soft membranes, which is much more frequent than that of the dura mater. The nutrient vessels of the brain pass in from the vessels running in the subarachnoid space, and their sheaths are direct continuations from the loose tissue of the pia mater. Hence in all forms of inflammation there is liable to be an extension inwards for a certain distance along the vessels. This is important as serious irritation of the cortical substance of the brain is apt to result.

(a) **Simple acute meningitis.**—This name is applied to non-specific inflammations of the pia-arachnoid. Some irritant obtains access to the membranes, and as it is carried by the circulating cerebro-spinal fluid the inflammation is usually of a spreading character. The cause is usually traumatic and the irritant is **Septic** in nature; but it may arise from disease of the bones, especially in ear disease, and in that case the



meningitis may be associated with abscess of the brain. There are cases in which the meningitis has an embolic origin, septic matter being transported in pyæmia or ulcerative endocarditis. It has also been found associated with acute specific fevers, typhoid, scarlet fever, small-pox, and with other acute diseases, especially pneumonia.

In the milder cases there may be simply a serous exudation in the subarachnoid space, constituting an inflammatory œdema. But in most instances the case goes on till pus, or pus with fibrine, is exuded. The first appearances are visible in the neighbourhood of the veins, in the sulci between the convolutions. Very often there is a white or yellow band visible on either side of the vein, and this consists of accumulated leucocytes, it may be with fibrine. As the exudation increases the veins become buried in it, and the whole subarachnoid space becomes filled. The spaces being filled out the pia-arachnoid forms a bulky solid layer which may be separated from the surface of the brain, and form a mould of the convolutions on its under surface. The appearances under the microscope are similar to those shown in Fig. 242. The inflammation extends, as already noted, in some measure along the perivascular spaces into the superficial part of the brain substance.

The inflammation usually begins in a particular locality, generally some part of the convexity, and here the arachnoid and the dura mater are sometimes united by fibrinous exudation. It spreads from this, generally confining itself to the convexity, but sometimes it extends to the base, or to the spinal cord, or even into the ventricles. It is to be noted that, in accordance with what is stated in the Introduction to this section (see p. 585), the exudation is in the subarachnoid space not on the free surface of the arachnoid.

**Simple chronic meningitis** of the brain and spinal cord is usually secondary. There are thickenings and adhesions of these membranes in different forms of insanity, in diseases of the bones, in the various sclerosis of the cord, especially posterior sclerosis. It is important to remember that a chronic inflammation of the membranes of the cord may be propagated from the peripheral nerves along their sheaths.

(b) **Epidemic cerebro-spinal meningitis.**—As the name implies, this is a disease which, like the acute fevers and hydrophobia, depends on a virus introduced into the body. The morbid poison shows a special affinity for the meninges of the brain and spinal cord, although other organs are also in a minor degree affected.

The soft membranes present evidences of acute inflammation, at first in the form of serous exudation with few leucocytes and red corpuscles. This condition is only seen in cases which have died very early after the onset of the disease, for the exudation soon takes on a purulent charac-



ter as in simple meningitis. The exudation is here also in the sub-arachnoid space, and penetrates, along the sheath of the nutrient arteries, into the nervous tissue beneath (see Fig. 242). There is very seldom

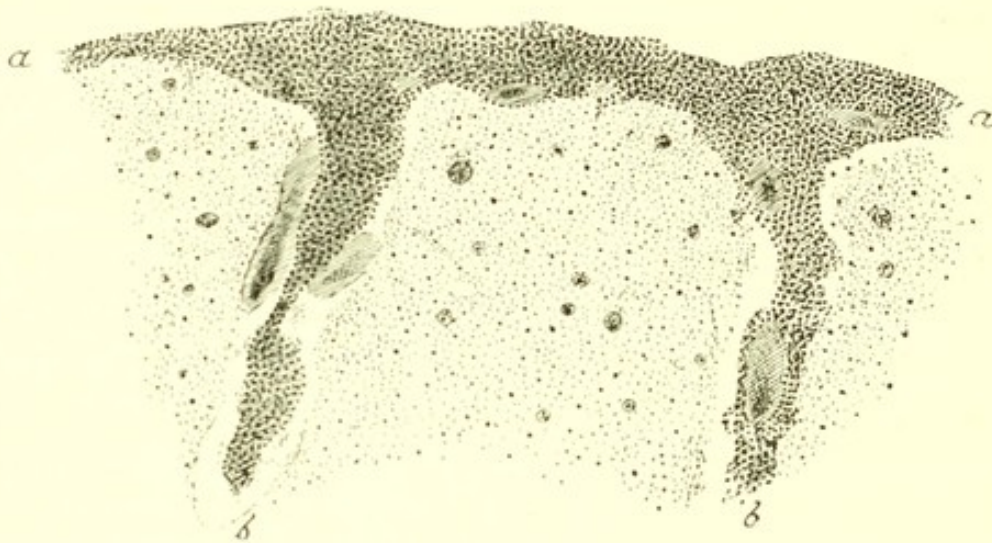


Fig. 242.—From a case of cerebro-spinal meningitis. *a a*, the soft membranes infiltrated with pus, also passing into the sulci, *b b*. Dilated veins are seen in the midst. Vessels also prominent in the brain substance.

any exudation on the surface of the arachnoid, and the dura mater hardly ever takes part in the disease. The exudation is most marked in the sulci between the convolutions of the convexity, in the fissure of Sylvius, on the surface of the pons, and upper surface of the cerebellum. It sometimes extends into the lateral ventricles. In the spinal cord it is most abundant in the lumbar region, and is almost confined to the posterior surface where it often surrounds the posterior roots.

The disease is fatal in over 50 per cent. of the cases, but when recovery occurs it is usually complete. Sometimes, however, there is permanent damage to the nervous structures, as evidenced by deafness, etc.

There can be little doubt that this disease is due to a micro-organism and probably a micrococcus, but the exact form has not been determined. Pneumonia sometimes occurs in conjunction with the meningitis, as if the same morbid poison produced both. Otherwise the organs of the body show appearances similar to those in other acute fevers. The spleen is enlarged but not very greatly so. The follicles of the intestine are swollen. The liver and kidneys are enlarged and show parenchymatous infiltration. There are sometimes abscesses in the muscles and pus in the joints. The skin has a peculiar tendency to show petechiæ during life, and becomes deeply stained by the blood colouring-matter after death.

(c) **Tubercular meningitis** (*Basal meningitis*, *Hydrocephalus acutus*).—This disease, as the name implies, depends on the presence in the meninges of the tubercular virus. The tubercular bacillus is readily detected in the affected structures, lying diffused in the membranes but



more especially in the walls of the arteries. In the majority of cases the virus is carried to the membranes by the blood. The disease is, in many cases, part of a general tuberculosis, and is associated with the usual lesions in other organs, but the meningitis leads to such pronounced symptoms that it often monopolizes the attention of the clinical observer. The primary seat of the infection is frequently in the bronchial, cervical, or mesenteric glands, the tuberculosis extending to the veins, and so to the blood. There may, however, be an extension by the blood without the occurrence of a general tuberculosis. The primary seat may be in the lungs, and the extension to the membranes may be the only secondary extension.

When the extension is not by the blood, there are evidences of a local infection. There may be solitary tubercles in the brain substance, although these do not usually give rise to tuberculosis of the membranes. There may also be an extension from the bones of the skull, as in a case observed by the author, in which a tuberculosis of the basilar bone and the first cervical vertebra had extended to the pituitary body and onwards to the soft membranes.

The meninges are affected by tuberculosis more frequently in children than in adults; it thus seems as if the soft membranes in children were more adapted to the growth of the virus. But tubercular meningitis is more frequent in the adult than is usually supposed, and many obscure head cases are found post mortem to be cases of general tuberculosis with, it may be, a very limited meningitis.

The virus reaching the meninges by the blood, lodges in the lymph spaces of the membranes, and produces inflammation and tubercles in the walls of the finer arteries of the pia mater, and in the subarachnoid space. These conditions are peculiarly localized in the basal parts of the brain, although extending sometimes to the lateral aspects and to the spinal cord. It may be that this is related to the fact that the arteries are distributed from the base, or, more probably, that the cerebro-spinal fluid stagnates here more than elsewhere, and allows of the growth of the virus.

The **Appearance of the brain post mortem** is somewhat as follows. On removing the dura mater, the cerebral hemispheres are generally seen to be fuller than usual, and the surface of the arachnoid is somewhat dry and glazed. This is due to the pressure from the ventricles, which are distended with fluid, usually clear serum, but sometimes slightly turbid with pus. The amount of fluid is sometimes very great, and this prominent feature gave rise to the name **Acute hydrocephalus** applied to this disease (see p. 587). In the neighbourhood of the lateral ventricles, and especially posteriorly, the brain substance is soft and almost diffuent (white softening). On exposing the base of the



brain, the appearances of inflammation in the membranes are to be looked for. These are often somewhat obscure, and in appearance trivial. In the slighter cases they consist merely of a turbidity or opacity over the pons and optic chiasma. The subarachnoid space is occupied by a serous exudation with some pus, giving often a greenish colour. But usually the exudation is more abundant and covers the basal structures, extending to the surface of the cerebellum, and up the fissures of Sylvius, where it often reaches the lateral aspects of the hemispheres. The concentration of the exudation at the base is of great importance, and the covering-in of the optic chiasma is often the most direct and sometimes the only prominent sign of the existence of the disease.

The affection usually extends to the **Meninges of the spinal cord**, although here it may be even more insignificant in appearance than in the brain. There may be little more than an undue redness with a granular appearance on the surface of the cord.

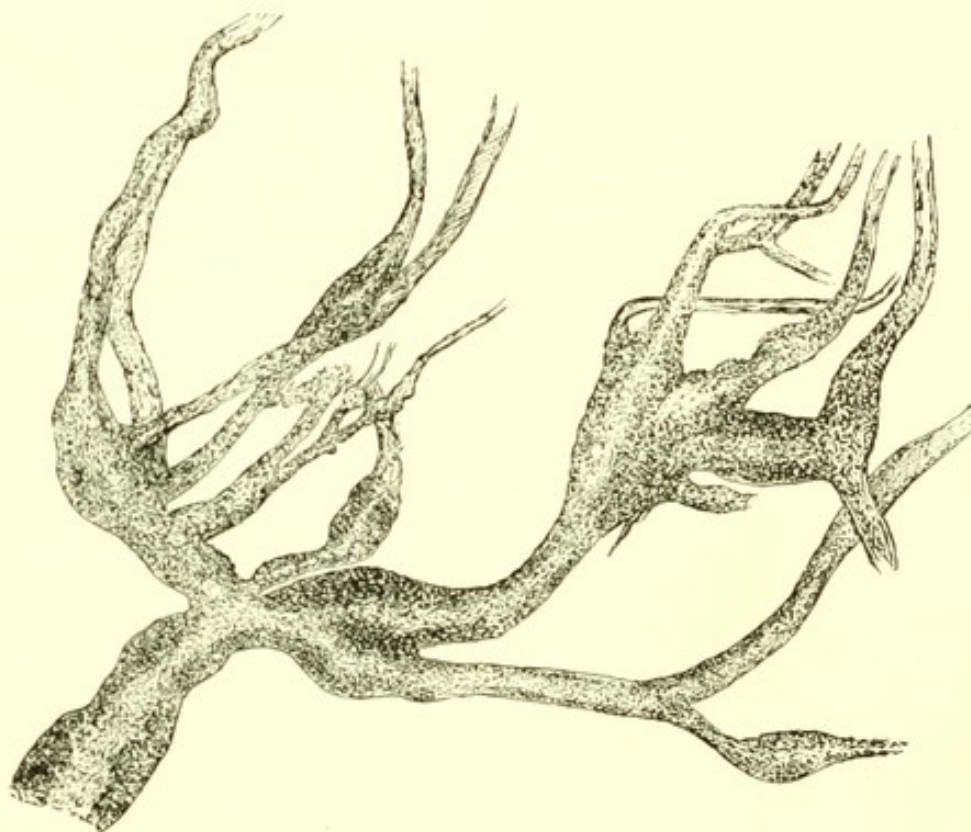


Fig. 243.—Arteries of the pia mater in a case of tubercular meningitis. The frequent spindle-shaped swellings indicate the position of the tubercles.  $\times 16$ .

The evidences of inflammation are much more prominent to the naked eye than the tubercles. These are often distinctly visible only on microscopic examination, as they are so much buried in the exudation and attached to such small arteries. On opening up the fissure of Sylvius, however, where the exudation generally glues the opposed surfaces



of the brain together, it is usually possible to see the little white tubercles, often no larger than pins' heads. If a piece of the soft membrane be removed, however, even from a part where the exudation is very abundant, and examined with a low power under the microscope, the tubercles are readily seen as spindle-shaped prominences on the small arteries (see Fig. 243).

There is not in this form of tuberculosis a distinct formation of typical tubercles with giant-cells and epithelioid cells, the nodules described as tubercles being composed of aggregations of round cells. The walls of the arteries are often specially affected, so that they present localized prominences (see Fig. 243) as if the virus had acted at certain spots. The swellings are mainly due to accumulation of cells in the adventitia (Fig. 244), but the intima is very commonly also the seat of an inflammatory infiltration as shown in the figure. Besides this there are



Fig. 244.—Tubercular meningitis. *a*, an artery with infiltration chiefly of external coat, but also of internal, which shows villous projections. The connective tissue around is also infiltrated, and so are the walls of the nutrient arteries passing into the brain substance in either side, *b b*.  $\times 30$ .

tubercles in the walls of veins and in the tissue apart from the vessels, and there is a general infiltration of the membranes with multitudes of leucocytes (see Fig. 244). The inflammatory infiltration extends along the nutrient vessels into the substance of the brain and spinal cord, as well as into the cerebral nerves. The superficial parts of the brain substance in particular are hyperæmic, and not infrequently the seat of small hæmorrhages.

The **White softening** of the parts around the ventricles has given rise to some discussion as to its cause. It may extend somewhat deeply into the brain substance, involving fornix, septum lucidum, corpus callosum, and even the thalamus opticus



and corpus striatum, and it is sometimes so extreme as to reduce the tissue to the consistence of thick cream. It is clear from the comparative absence of symptoms that this extreme softening does not exist during life. Probably the fluid in the ventricles macerates and loosens out the brain tissue without interrupting the functions, and after death a more pronounced softening occurs. The condition is not an inflammatory one, and it occurs mainly where the fluid gravitates, namely, in the brain substance around the posterior parts of the ventricles.

**Literature.**—NEISSER, *Die Entzünd. der serös. Häute des Geh. u. Rückenm.*, 1845; HUGUENIN, in *Ziemssen's Handb.*, xi. 1. *Epidemic cerebro-spinal meningitis*—MEISSNER, *Schmidt's Jahrb.*, 129 and 136; KLEBS, *Virch. Arch.*, xxxiv.; WILICH, (*Pneumonia and mening.*) *D. med. Wochenschr.*, 1875, No. 23; FICKET, (do.) *Annal. de la soc. méd. chir. de Liège*, 1880; GOWERS, *Dis. of nerv. syst.*, 1888, ii. 317 (literature fully); FREW, *Glasg. Med. Jour.*, 1884, xxii. p. 21; EBERTH, *D. Arch. f. klin. Med.*, xiii.; NAUWERK, (do.), xix.; CORNIL et BABES, *Les bactéries*, 1886, p. 446.

**4. Syphilitic meningitis.**—This has been described in a previous page along with Syphilitic lesions of the brain (p. 577).

#### IV.—TUMOURS AND PARASITES OF THE MENINGES.

Tumours of the membranes of the brain are of importance especially when they press on the brain inside or on the nerves as they issue from the skull. They are of considerable variety.

**Fibromas** have been found arising from the dura mater both of the brain and of the spinal cord. In the latter case they are liable to press on the cord and may even interrupt it. In a case observed by the author a hard fibroma the size of a marble produced serious mischief by pressing on the cervical cord.

The **Chondroma** has occasionally been observed, especially in the dura mater of the cord. In a dog observed by the author, paraplegia was produced by such a tumour in the dorsal region.

The **Lipoma** is a very rare form of tumour in the meninges; that already mentioned on the surface of the corpus callosum (p. 229) probably originated in the pia mater.

The **Psammoma** occurs not infrequently in the dura mater, where it forms a rounded growth on its internal surface (see Fig. 245); sometimes it is multiple. The tumour is usually small, but may be as large as a walnut, and its surface is rough and tuberculated. The psammoma also occurs as a small soft tumour in the choroid plexus and in the pineal gland. In the latter case there may be a tumour of considerable size, whose structure is like that of the normal gland, and so the condition is sometimes designated **Hyperplasia of the pineal gland**. There may also be a **Papilloma**

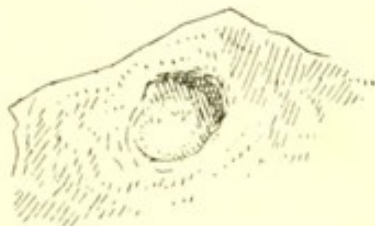


Fig. 245.—A small psammoma of the dura mater, natural size.



arising from the choroid plexus either in the lateral ventricles or the fourth ventricle.

The **Osteoma** hardly occurs in the membranes as a distinct tumour, but it is common to find flat pieces of bone in the dura mater, especially in the falx and tentorium, and even in the arachnoid.

The **Pacchionian bodies** sometimes give origin to tumours which may grow to considerable dimensions. These bodies consist of papillæ which spring from the arachnoid and project in various directions; some of them present towards the skull, where, after thinning the dura mater, they cause pits; others project into the longitudinal sinus.

Cleland has described two tumours of a papillary character which grew from the dura mater and pressed on the brain substance. Although one of these was situated in the falx, which does not normally present pacchionian bodies, yet the structure of the tumours conformed to that of these bodies. In that on the falx there were calcareous masses like those which characterize the psammoma. The tumours measured respectively two inches and one and a half in diameter.

**Sarcomas** are exceedingly important tumours, especially those of the **Dura mater**. Most of these tumours are soft in consistence and contain chiefly round cells. Sometimes there is a distinct alveolar structure, and the cells have a distinct resemblance to epithelium, so that the tumours have been often described as **Cancers**. There are even tumours remarkably like epitheliomas sometimes observed in the dura mater; they are generally regarded rather as endotheliomas. The sarcomas generally present great malignancy. On the one hand, they extend and involve neighbouring parts of the dura mater as well as the soft membranes and the brain; and, on the other hand, they press outwards on the skull, and may, after destroying the bone, work through to the external surface. These tumours have their seat mostly at the base of the skull and often lead to serious nervous lesions. They may involve directly such structures as the pons or medulla oblongata, the sarcomatous tissue growing into and replacing the proper structure; they also grow into the cranial nerves. By involving the bone and enlarging it, they may impinge on the foramina by which the nerves take exit. As they are seated usually at the base, they generally, when they work through the skull, present at the pharynx or nares.

Of the remaining tumours, the **Syphilitic** have been already considered (p. 577). Tubercular growths hardly ever occur except by propagation from disease of the vertebræ. **Dermoid cysts** of the dura mater have been observed in a few cases.

The **Echinococcus** sometimes develops in the neighbourhood of the spinal canal, in the muscles or in the vertebræ. By enlarging it may extend into the canal, pushing the dura mater before it and compressing



the cord. It has even been found to have its seat inside the dura mater. A few cases of *Cysticercus* in the subarachnoid space have been described.

**Literature.**—BEIGEL, (Cysts of choroid plexus) Path. trans., 1869, xx.; KELLY, (Papilloma of 4th vent.) do., 1873, xxiv.; CLELAND, Glasg. Med. Jour., xi., 1861; DAVINE, Traité des entozoaires, 1878; MADER, (Cysticercus in 3rd vent.) Berichte des Rudolfstiftes in Wien, 1872.

#### V.—AFFECTIONS OF THE PINEAL AND PITUITARY BODIES.

The pineal gland is occasionally the seat of tumours. There are simple enlargements from hyperplasia, sometimes with increase of the brain sand constituting a psammoma. There are also cysts replacing the gland.

Bulky tumours of a **complex structure** have been observed originating in the pineal gland, sometimes attaining to the size of a small apple. Weigert has described a teratoma which measured an inch and three



Fig. 246.—From complex tumour of pineal gland. *a*, cartilage, *b*, gland tissue developing cysts; *c*, sarcomatous tissue.

eighths in diameter, and contained hairs, hair-follicles, sebaceous glands, cartilage, fat, smooth muscle, cylindrical epithelium, and perhaps nerves. Falkson and the author have recorded cases to which the name chondro-adenosarcoma might be given, in which the bulk of the tumour was sarcomatous, but associated with cartilage, glandular structures, and cysts (see Fig. 246), and Turner has described one somewhat similar in structure but without cartilage.



From the position of the pineal gland these tumours are specially liable to distend the third ventricle, and they will frequently press on the optic thalami, peduncles and corpora quadrigemina (see Fig. 247).

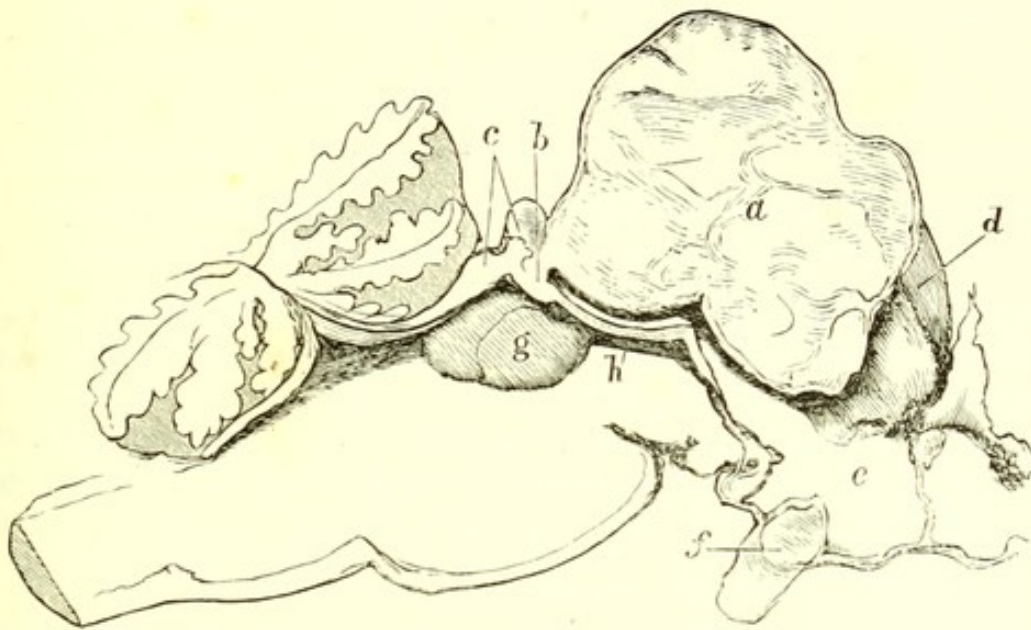


Fig. 247.—Tumour of pineal gland. *a*, tumour; *b*, pedicle attached in situation of pineal gland; *c*, corpora quadrigemina; *d*, thalamus opticus; *e*, third ventricle; *f*, optic nerve; *g*, portion of tumour in aqueduct of Sylvius and fourth ventricle.

At an early period of development the pineal gland contains saccules and follicles lined with cylindrical epithelium, and this structure is retained in adult life in some animals. According to recent observations the pineal gland is an aborted or rudimentary median eye, the **Pineal eye**, which in the reptilia still shows sufficient structure to be recognizable. The eye is of the invertebrate type having the rods and cones in front of the expansion of the nerve instead of behind it as in the vertebrata. These observations may perhaps explain the occurrence of tumours with epithelium, cartilage, and other structures in the pineal gland.

The **pituitary body** may take part in various processes along with the membranes of the brain. It may be the seat of inflammation, or tuberculosis, or syphilis in common with these membranes. Or there may be an extension of tuberculosis from the base of the skull to it. It is also liable to fatty and colloid degeneration, and the latter may cause considerable enlargement of the gland. **Tumours** also occur in the pituitary body. Weigert describes one the size of a hen's egg which consisted of the normal tissue of the gland but highly oedematous. He designates it a struma or goitre of the gland. Sarcoma has also been observed.

**Literature.**—*Pineal gland*—WEIGERT, Virch. Arch., 1875, lxxv.; FALKSON, *ibid.*, 1879, lxxv.; TURNER, Path. Trans., 1885, xxxv.; COATS, *ibid.*, 1887, xxxviii.; MIHALKOVICS, Centralb. f. d. med. Wiss., 1874; BALDWIN SPENCER, (Pineal eye) Quart. Jour. of microscop. Sc., Oct. 1886.



## SECTION IV.

## DISEASES OF THE ORGANS OF RESPIRATION.

- A.—THE NASAL PASSAGES. 1. *Congenital malformations*; 2. *Hæmorrhage*; 3. *Acute catarrh*; 4. *Chronic catarrh*; 5. *Syphilis and tuberculosis*; 6. *Tumours*; 7. *Foreign bodies*.
- B.—LARYNX AND TRACHEA. I. MALFORMATIONS. II. INFLAMMATIONS. 1. *Croup and diphtheria, lesions in fauces and larynx; question of independent croup*. 2. *Acute catarrh, sometimes leading to œdema glottidis*; 3. *Chronic catarrh*; 4. *Subglottic inflammation*; 5. *Inflammation of cartilages*. III. SYPHILIS AND TUBERCULOSIS. IV. TUMOURS AND FOREIGN BODIES.
- C.—THE BRONCHIAL TUBES. I. INFLAMMATIONS, chiefly 1. *Catarrh, constituting ordinary Bronchitis*; 2. *Fibrinous bronchitis*. II. STENOSIS AND DILATATION. III. TUMOURS AND FOREIGN BODIES. *Importance of the lesions resulting from foreign bodies*.

## A.—THE NASAL PASSAGES.

1. **Congenital malformations.**—Such deformities as absence of the nose and its cavities, stenosis, and so on, are usually parts of a general malformation, and have been already referred to under General Malformations. In **Clefts of the lip and palate**, the nasal cavities are in communication to a greater or less extent with the mouth. This communication renders the mucous membrane of the nares liable to inflammation. Normally, the nasal passages are protected against any irritation except that of the air passing through them, and the mucous membrane is correspondingly sensitive. We know how the accidental passage of a piece of solid food into the nares causes great irritation, and we shall afterwards see that the existence of a foreign body in the nares is a frequent cause of prolonged catarrh. In cases of cleft palate the food and secretions of the mouth readily pass into the nares and, although a certain tolerance may be established, persons so affected are peculiarly liable to catarrh of the nares.

2. **Hæmorrhage. Epistaxis.**—Hæmorrhage from the nares is of frequent occurrence. There are persons who have a special proclivity to it, in whom it occurs at intervals without apparent cause. It also occurs in consequence of tumours, especially polypi in the nares, the mucous



membrane over the tumour being liable to bleed. It is an occasional concomitant of fevers (typhoid fever) especially at their onset. The blood usually comes from the anterior part of the nares, and especially from the mucous membrane over the cartilaginous septum.

**3. Acute catarrh. Acute rhinitis. Coryza.**—Acute inflammation of the nares occurs most frequently as a catarrh, constituting the ordinary cold in the head, but there may be a more intense and specific inflammation in diphtheria and scarlet fever.

In the case of the specific fevers, the special irritant concerned in their production may attack the nares and produce acute inflammation there. This is nearly always the case in measles, but sometimes also in small-pox, diphtheria, and scarlet fever. For ordinary catarrhs Cold is usually assigned as the cause. The mucous membranes of the air passages are indeed much more liable to inflammations than those of the alimentary canal, and an exposure to cold air which would not produce any catarrh of the mouth might possibly do so in the nares. Moreover, the fact that the nares are the nearest part of the air passages to the air, and are therefore most exposed to the action of cold, is an indication that such exposure has probably something to do with it.

Mere inhalation of cold air is not enough to produce catarrh, as every one who is liable to cold in the head will admit. A person is able at one time to brave any amount of exposure without the risk of catarrh, while at another time a slight draught, or no perceptible exposure at all, will bring it on. Such circumstances as these, taken along with the fact that the catarrh has a definite course, usually of a week's duration, has induced some to suppose that the disease is really due to a specific morbid poison. In this view it is necessary to suppose that the microbes are usually present in the air, and that they take, as it were, advantage of the mucous membrane, when, at any time, it is in a specially predisposed state. This predisposition may be induced by cold applied either directly to the mucous membrane or to some other part of the body, but other predisposing causes may exist. For instance, when a person is overheated the arteries of the skin and of the respiratory mucous membranes are relaxed, and there is an active hyperæmia. At such a time a catarrh is readily induced apparently because the mucous membrane is less able to resist the action of irritants than usual. There is apparent confirmation of such a view as this in the fact that nasal catarrh undoubtedly occurs as the result of the action of specific irritants. In measles the catarrh of the nares and conjunctivæ is referrible to the specific virus of that disease. In hay asthma the irritation of the pollen of grasses is believed to be the cause of the catarrh. Again, it is commonly stated that nasal catarrh is communicable from person to person, and it is hardly conceivable that this could occur unless the disease had, as a part of its cause, some specific materies morbi.

The catarrh begins with an inflammatory hyperæmia of the mucous membrane. This may, of itself, lead to swelling so great as partially to obstruct the passages, giving the feeling of a "stuffed nose." The inflammation soon passes on to exudation. This finds its way



for the most part to the surface, but in its passage it increases the swelling of the mucous membrane. The exudation is the usual one of inflammation, namely the blood-plasma with leucocytes. At first this is mixed with mucus, but as it increases in amount it rapidly assumes a serous character, and we know that a great abundance of serous fluid may be discharged from the nostrils. As time goes on the leucocytes increase in the exudation, and it may assume a semi-purulent character. Sometimes red corpuscles are present, giving the discharge a greenish-yellow colour.

It is to be remembered that there are various cavities in direct communication with the nares, of which the principal are the **Frontal sinuses** and the **Antrum of Highmore**, and that these frequently take part in the acute catarrh. It will be observed that in all stages of the catarrh there is more or less swelling of the mucous membrane, and that this leads to the obstruction of the passages which is such a marked symptom. This swelling is temporary in the acute disease, but in chronic catarrh it is apt to give way to a more permanent thickening.

Acute inflammation sometimes extends to the nares in **Diphtheria**. The anatomical changes will be considered in next section.

4. **Chronic nasal catarrh.**—This may supervene upon acute catarrh, or it may be of more independent origin. Sometimes the presence of foreign bodies induces it. The prolonged existence of catarrh leads to various changes in the mucous membrane. There may be **Atrophy** with induration, a process similar to cirrhosis. In this case there is frequently an excessive purulent secretion (**Ozæna**), which usually has a very foetid smell. In other cases there is **Hypertrophy** of the mucous membrane, which may even amount to a distinct polypus (see below). The finer ramifications of the olfactory nerves are injured by these tissue changes and smell is lost or impaired, in many cases permanently.

5. **Syphilitic and Tubercular lesions of the nares.**—**Syphilis** not infrequently affects the nares in the tertiary stage. There is the formation of gummatous tissue with inflammation, beginning either in the mucous membrane or in the deeper parts. As the lesion is superficial there is usually ulceration, and this may involve the soft parts and the bones very extensively. There may arise in this way serious lesions, such as perforation of the septum, communication with the mouth, falling in of the nasal bones, etc. When the bones are affected the discharge is liable to be very putrid (*Syphilitic ozæna*). If healing occurs the cicatricial contraction may produce still further deformities.

**Tuberculosis** of the nares is not often observed, perhaps because the nares are not usually examined fully post mortem. The disease occurs



chiefly in connection with tuberculosis of the lungs. There are tubercles in the mucous membrane and ulceration.

6. **Tumours.**—The most frequent tumours of the nares are the **Mucous polypi**. Like other mucous polypi they occur very commonly as a result of chronic catarrh, but they appear occasionally without any such cause. They are usually in the form of rounded projections having a comparatively narrow base or neck, the growth becoming more pedunculated as it enlarges. They often produce much obstruction of the passages. In structure some of them present simply the constituents of the inflamed mucous membrane, connective tissue with rather wide serous spaces, and covered with cylindrical ciliated epithelium. (If the epithelium be examined in fluid immediately after removal, the ciliary motion will be seen.) The connective tissue is usually so infiltrated with serous fluid as to give an œdematous appearance to the polypus, and sometimes a more definite cystic formation occurs. In some cases there are contained in the tumour mucous glands, but it is doubtful if there is a true newformation of gland tissue. Cysts sometimes develop from dilatation of mucous glands contained in the tumour. The polypi are not infrequently multiple.

Polypi are sometimes found of a different character from those mentioned above. There may be true **Papillomata**, or there may be **Myxomata** or **Fibromata** taking the form of polypi. These latter may take origin in the periosteum. Malignant tumours also sometimes assume the polypoid form.

Of the malignant tumours **Sarcomas**, originating mostly in the periosteum or perichondrium, may produce serious obstruction and deformities of the nasal structures. The sarcomas often dislocate the nasal bones, and involve the neighbouring structures in their substance. In this way they sometimes penetrate into the antrum, or involve the hard palate and alveoli. **Cancers** rarely occur in the nares as primary tumours, but may involve them by extension from neighbouring parts.

7. **Foreign bodies.**—Different kinds of foreign bodies are not infrequently found in the nares. They may be introduced into the nostrils accidentally or designedly by children, and consist of various articles. These may become coated with lime salts so as to form nasal calculi or **Rhinoliths**, which may also form without a true foreign body, the lime being deposited in inspissated secretions. Occasionally, maggots find their way into the nares, and sometimes fungi (*Aspergillus fumigatus*) form.

Most foreign bodies irritate the mucous membrane, producing chronic catarrh, frequently accompanied by very putrid discharges (*Ozæna*).



**Literature.**—COHEN, Dis. of throat and nasal passages, 1879; MORELL MACKENZIE, Manual of dis. of throat and nose, ii., 1884; MICHEL, Krank. d. Nasenhöhle, 1876; ZUCKERKANDL, Norm. u. path. Anat. der Nasenhöhle, 1882; BRESGEN, Chron. Nasen- und Rachen-katarrh, 1883.

## B.—THE LARYNX AND TRACHEA.

The lesions of the larynx and trachea are frequently associated with those of the bronchi, more especially the inflammations. Their separation here is consequently somewhat artificial.

**I. Malformations.**—Entire **Absence** of the larynx and trachea occurs only in acephalic monsters which are incapable of living. There are, further, cases of **Communication between the trachea and œsophagus**; generally in these cases the pharynx ends in a cul-de-sac and the œsophagus opens into the trachea. Then we meet with cases of imperfect closure of the original branchial arches leading to the **Congenital fistula of the neck**, already considered (see p. 47). Again, **Individual cartilages**, as the epiglottis, or one or more of the rings of the trachea, may be **absent**, or there may be one or more rings supernumerary. The trachea may divide into three main bronchi instead of two, and in that case two stems pass to the right lung and one to the left. Occasionally the larynx is **congenitally narrow**, or it may fail to undergo the usual changes at puberty, especially in cases of castration before puberty or of non-descent of the testes. Lastly, the trachea has been observed to the left of the œsophagus or even behind it.

**II. Inflammations of the larynx and trachea.**—1. **Croup and Diphtheria.**—Much has been written on the question whether these two diseases are identical or not. Without in the meantime entering on that subject, it may be said that in using the terms here it is intended to refer to the conditions which are designated clinically as croup and diphtheria, and that the distinction sometimes made between croupous and diphtheritic inflammation is not made use of. (See ante, p. 133.)

In **Diphtheria** we have a disease which is undoubtedly infectious, its communicability from person to person being doubtless associated with the fact that it depends on the existence of specific bacteria, whose characters, however, have not been fully established. (See ante, p. 304.) The specific microbe, either in itself or by its products, is an irritant to mucous membranes, and produces violent inflammations.

The inflammations differ somewhat in the fauces and nares on the one hand and the larynx and trachea on the other. In both cases a catarrh is the first sign of inflammation. The mucous membrane is hyperæmic, and there is increased mucous secretion. In the case of the **Fauces and**



**Nares** this is succeeded by the formation of a false membrane, which does not lie free on the surface but involves the superficial layers of the mucous membrane as well. The irritant in fact has caused a partial necrosis of the mucous membrane, which by and by sloughs and leaves an ulcer. The necrosis here is in the form of coagulation-necrosis, the dead tissue becoming converted into something like a fibrinous clot, and enclosing true fibrine in its meshes. The false membrane is therefore composed of fibrine and necrosed and coagulated mucous membrane. When the slough separates or is removed there is apt to be a second formation of false membrane.

In the **Larynx and Trachea** the process is a somewhat different one. The disease usually begins in the fauces, whence it occasionally extends in the nares, and more frequently into the pharynx and larynx. In the larynx, after the catarrhal stage, there is the production of a false membrane, but this does not adhere to the mucous membrane, nor is there usually any necrosis. The false membrane is a whitish layer which is loosely connected with the mucous membrane and is readily raised by the accumulation of mucus under it. There are undoubted inflammatory changes in the mucous membrane in the form of infiltration with round cells, but the only actual loss of substance is the shedding of the epithelium of the surface. The epithelium always undergoes necrosis before the formation of the false membrane.

Differences of opinion exist as to the nature of this false membrane in the larynx and trachea. By Wagner it has been asserted that it arises from the epithelium, being the product of what may be regarded as a coagulation-necrosis of it. It is not to be denied that the epithelium may undergo this process, but it does not seem possible that this can be the source of the whole membrane. We know that the membrane is often removed and reproduced, and that it may be produced in such quantity as to fill up the larynx completely. The thin layer of epithelium is quite inadequate to such a result, and we must believe that fibrine is actually deposited on the inflamed surface. The removal of the epithelium leaves the surface incapable of preventing the disintegration of the leucocytes. The false membrane is, therefore, to be regarded as an inflammatory exudation, and in its histological characters it agrees with this. It consists of a fibrinous network, often very coarse in its texture, and with inflammatory cells in it.

**Croup** is a name generally given to membranous sore throats in which the exudation is mainly or entirely in the larynx and trachea. The name was first applied before the eminently contagious diphtheria was distinguished, and it was commonly understood to designate a peculiarly violent inflammation of the air passages. There seems to be no doubt that, now-a-days at least, the great majority of cases of croup are really cases of diphtheria in which the disease is mainly or entirely in the larynx and trachea. The name is also given to throat affections in



general, in which respiration is accompanied by a crowing sound, this being indeed the original meaning of the term Croup, which is a Scotch word signifying the sound made in such cases.

We have seen that in the larynx diphtheria produces an inflammation in which, after the shedding of the epithelium, a fibrinous exudation occurs. It is asserted by some that the diphtheritic poison is the only agent capable of producing this form of inflammation. Looking at the matter from a purely pathological point of view apart from clinical experience, it certainly seems possible that other irritants may produce similar results. Croup has been produced artificially in rabbits by the injection of ammonia into the trachea (Weigert). In these cases the irritant first kills the epithelium, and then fibrine is deposited. If croup occurs in man apart from diphtheria, the irritant must be strong enough to destroy the surface epithelium. Apart from the action of microbes, such an irritant must be of rare occurrence, but the possibility of its existence is not to be denied. We shall see afterwards that in rare cases we meet with a bronchial croup where there can be no question of diphtheria, and so we may have laryngeal and tracheal croup of a simple inflammatory kind. In such cases there will be no signs of general disease, but all the symptoms will be referrible to the local inflammation and obstruction of the larynx. It should be added that laryngeal croup has been met with in small-pox, measles, pyæmia, etc., and in that case it is to be ascribed to the action of specific morbid poisons, as in diphtheria.

**2. Acute catarrh of the larynx and trachea.**—We have seen that **Acute catarrh** forms the first stage in croup; it is the result of the action of the specific poison. Similarly we have acute catarrh in measles and small-pox, there being here a specific eruption similar to that on the skin, along with acute catarrh. As we have already seen, this may, in exceptional cases, go on to the formation of a fibrinous exudation as in diphtheria. In typhoid fever we may also have acute catarrh, which, according to Eppinger, is of similar significance to the affection of the intestine, being due to the specific agent. Catarrh occurs also as an independent affection, just as nasal catarrh does, and in this case, although usually slight, it may assume a very severe character. Lastly, a catarrh may be set up by the inhalation of irritating chemical fumes.

There is, as in other inflammations, hyperæmia and exudation. The mucous membrane is red as seen during life, but on post-mortem examination the redness has usually disappeared entirely, the vessels emptying apparently by the shrinking of the tissue. The exudation is originally mucous in character, and is not generally very abundant. After a time, as in the case of nasal catarrh, it usually assumes a more purulent character. The swelling of the mucous membrane is not usually great, and there is not commonly any serious obstruction. On the other hand, in children a slight catarrh may bring on a sudden suffocative attack due not so much to the swelling as to spasm of the



muscles. As an unusual complication of acute laryngitis may be mentioned œdema glottidis, the condition next to be described.

**Œdema glottidis.**—This name is applied to a comparatively sudden œdematous swelling, causing often a serious or even fatal obstruction of the larynx. The œdema is in most cases an inflammatory exudation, but it occurs in Bright's disease as part of a general œdema. It may be part of a simple inflammation of the larynx, or may be connected with diphtheria, or the pustular inflammation of small-pox. Or the inflammation may be propagated from the pharynx and fauces, or from the inflamed perichondrium. The condition is not an œdema of the mucous membrane itself; that would produce a very moderate swelling; but it is an inflammation and œdema extending to the submucous tissue. In most parts of the larynx there is little or no submucous tissue, the mucous membrane being bound down to the perichondrium. There are some parts, however, where the tissue is looser, chiefly the base of the epiglottis, and, to a less extent, the whole epiglottis, the ventricular bands, and, most of all, the ary-epiglottic folds. The epiglottis is swollen, especially at its base; the ary-epiglottic folds are usually much tumefied, appearing as rounded tumours projecting backwards from the base of the epiglottis. These rounded swellings form indeed the most prominent appearances. The ligaments passing from the epiglottis to the tongue are also sometimes swollen. Examined from above, the tumefied ary-epiglottic folds conceal the parts beneath, but on laying open the larynx after death it is found that the ventricular bands (false cords) are tumefied, although the true cords are usually very little affected. The œdema may affect the submucous tissue in the trachea for some distance below the glottis. If the swollen parts be cut into a fluid exudes, which is usually sero-purulent and sometimes almost purulent. Occasionally there is blood in it.

3. **Chronic catarrh.**—This is a common result of repeated attacks of acute catarrh, but may occur spontaneously. It is chiefly characterized, like other chronic inflammations, by newformation of tissue; the mucous membrane is thickened, and its surface is irregular. The increase is mainly of connective tissue which, having the usual characters of that resulting from inflammation, gives rigidity to the parts. The moveable structures of the larynx are thus rendered more or less stiff, and hoarseness is the result. Not infrequently flat superficial ulcers or erosions form, and these have their seats most commonly at the posterior commissure. The racemose glands of the larynx may undergo special enlargement so as to appear as rounded prominences. They sometimes ulcerate, and so give rise to small crater-shaped ulcers, which are chiefly to be seen on the epiglottis and ary-epiglottic liga-



ments. The thickening and contraction of the connective tissue are sometimes so great as to produce very great **Stenosis of the larynx**, so that tracheotomy is needed to permit of respiration. Sometimes mucous polypi form on the surface, and add to the irregularity.

4. **Subglottic inflammation.**—This disease, which is not of very frequent occurrence, is an inflammation of the mucous membrane beneath the glottis. It may be acute at its onset, but it generally passes into a chronic stage. It has been observed as a sequel to erysipelas and typhus fever, and may take origin apparently in inflammation of the perichondrium. In acute cases there may be considerable oedematous swelling. In the chronic form there is thickening of the mucous membrane as in ordinary chronic laryngitis. The inflammation is often just beneath the cords, and so may produce fixation of them, but it may occur further down, and is not infrequently in patches interrupted by normal mucous membrane.

5. **Inflammation of the perichondrium. Perichondritis.**—This disease is rarely a primary one, being induced chiefly by syphilitic and tubercular inflammations, especially when there is deep ulceration extending down to the perichondrium. It occurs occasionally as a sequel of typhoid and also probably of typhus fever. It has usually a somewhat chronic course, but may be acute, and in either case it ends in the formation of pus under the perichondrium. The pus, accumulating under the perichondrium, cuts off the cartilage from its source of nutrition, and, just as in periostitis, this is usually followed by necrosis of the cartilage. The destruction of the cartilage may be a slow process, and there may be a kind of caries followed by necrosis. This disease is generally confined to one cartilage at the outset, the cricoid being most frequently attacked, but it may extend to others. When suppuration has occurred the inflammation spreads to structures around, and we may have burrowing of the pus under the mucous membrane for some distance, or even outside the larynx among the structures of the neck.

The necrosed cartilage is, by degrees, separated from the living. It is usually discharged into the larynx, but the pus sometimes forms an external opening through which the cartilage may pass. There is usually great deformity of the larynx, which may be partly due to the primary disease and partly to the collapse resulting from loss of the cartilage. When the cartilage is discharged it is generally found calcified or ossified, and it may be a question how far the calcification preceded the inflammation. Dietrich has suggested that, in the case of the cricoid, ossification may sometimes be the primary condition, and that the inflammation may be induced by the pressure of the hardened cartilage against the vertebral column.



**III.—Syphilis and Tuberculosis of larynx and trachea.—1. Syphilis.**  
—In the secondary period of syphilis the larynx is frequently the seat of catarrhs which are to be classified along with the various inflammations of that stage. There may be in this period slight erosions of the surface, but no proper ulceration.

In tertiary syphilis the larynx is occasionally attacked, and we have here, as in other situations, irregular infiltrations of granulation tissue, gummatous newformations, ulcerations, etc.

The lesions may be at first comparatively superficial, consisting of thickenings and elevations of the mucous membrane so as to form irregular papillary projections (*Condylomata*), which sometimes imitate in appearance epithelioma of the larynx.

Much more characteristic, however, are deep infiltrations and ulcerations. The mucous membrane and submucous tissue are infiltrated and thickened, ulcers develop, first as a general rule in the epiglottis, but they are prone to extend deeply and widely so as to destroy large portions of the epiglottis or the whole of it. We have already seen that the ulceration may lead to perichondritis and necrosis of the cartilage with still wider results. With all this there is great newformation of connective tissue with corresponding deformity, and, if the ulcers heal, the contraction of the cicatricial tissue leads to great deformity and not infrequently to such obstruction of the glottis as to require tracheotomy.

**Congenital syphilis** sometimes manifests itself in the larynx. It may be in the form of a superficial catarrh, sometimes accompanied by oedema glottidis, or there may be a superficial infiltration of the mucous membrane. It is rare to have a deep infiltration and ulceration such as is common in ordinary syphilis.

**2. Tuberculosis. Laryngeal and Tracheal phthisis.**—Tuberculosis of the larynx is usually secondary to pulmonary phthisis, the mucous membrane being infected by the sputum from the lung; it occurs in about 30 per cent. of the cases of tuberculosis of the lung. But it is occasionally primary. Even when the laryngeal tuberculosis is secondary to that of the lung, it may seriously aggravate the latter by the infective material from the ulcerated surfaces being carried into the lung by insufflation.

The author has recorded a case of primary tuberculosis of the larynx. From observation in several cases he is induced to believe that, not infrequently, a slight pulmonary tuberculosis produces an infection of the larynx, and that the latter may advance while the former retrogrades. The laryngeal phthisis may thus slowly go on to ulceration while the pulmonary affection heals. Subsequently the lung may become infected from the larynx. In cases of ulcerated laryngeal tuberculosis, there



is often a very rapid and extensive tuberculosis of the lung, which presents the features of an almost simultaneous infection.

In cases of phthisis pulmonalis the mucous membrane of the larynx is frequently pale from anæmia, but this may be nothing more than a manifestation of a general anæmia secondary to the wasting disease of the lungs. The first result of the actual tubercular disease is inflammatory thickening of the mucous membrane. There is at first chiefly an exudation of serous fluid and inflammatory cells, so that it is mainly an oedematous thickening. It is most marked in the epiglottis and ary-epiglottic folds, these latter often showing themselves as rounded prominences. In this stage microscopic examination shows the presence of tubercles with their characteristic structure, along with inflammatory cells. The tubercles are in the mucous membrane and the submucous tissue, the epithelium being as yet intact.

To the thickening succeeds ulceration, the ulcers being at first small and superficial. These ulcers result from the caseation and softening of superficial tubercles. By coalescence larger ulcers form out of the smaller ones, and there is a continual tendency to spreading. As a rule there are many ulcers, and between them is thickened mucous membrane, which at the borders of the ulcers sometimes presents irregular projections like papillary outgrowths. The ulcers are at first superficial, but as the disease progresses considerable destruction of tissue may result. The vocal cords are not infrequently destroyed, and so there is loss of voice, but the voice may be lost from the rigidity of the structures caused by thickening from chronic inflammation. Again, perichondritis not infrequently follows, with suppuration, and this causes still further inflammatory manifestations.

Ulceration not uncommonly exists in the trachea and bronchi as well as in the larynx. There are many ulcers, and it is not uncommon to find the cartilaginous rings of the trachea extensively exposed. With these ulcerations of the trachea there will be swelling of the remaining mucous membrane and sometimes a perichondritis with necrosis.

The lymphatic glands in the neck are affected secondarily to the larynx; they are the seat of scrofulous processes such as we have already described, and their enlargement may, in some cases, aid in the exact diagnosis of the disease in the larynx.

**Leprosy** produces in the larynx thickenings and ulcerations similar to those of the skin.

**Glanders** also attacks the larynx, producing the lesions already described.

**IV. Tumours of larynx and trachea.**—The most frequent form of



tumour of the larynx is the **Papilloma**. This tumour is often preceded by catarrhal conditions, and is particularly common in persons who, from the nature of their profession, use the voice frequently. But it often occurs without either of these predisposing conditions. The tumours most frequently grow from the vocal cords, where the epithelium is flat. They consist, like other papillomas, of a basis of connective tissue forming numerous conical projections covered with epithelium (see Fig. 248). The epithelium may be thick, and the connective tissue

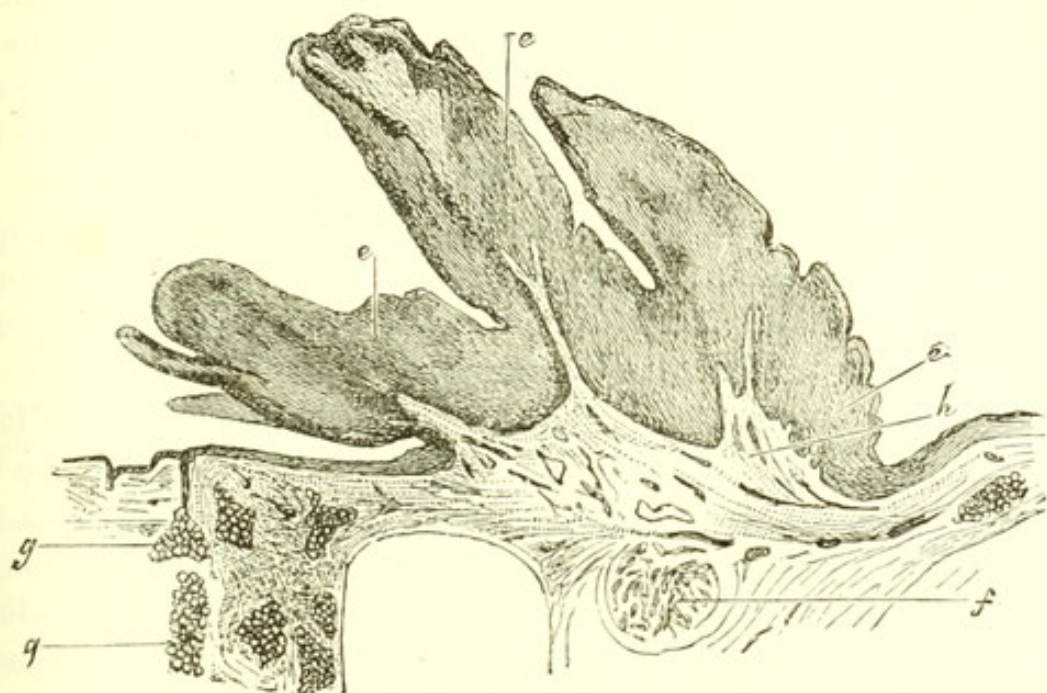


Fig. 248.—Papilloma of larynx. *e, e*, greatly thickened epithelium; *h*, connective tissue; *g*, mucous glands; *f*, a gland at base of tumour atrophied by its pressure.  $\times 20$ . (CORNIL and RANVIER.)

dense, so that the tumour is like a hard wart, or the epithelium may be thin and the connective tissue soft, sometimes richly infiltrated with round cells, and so the growth is soft like a soft wart or condyloma. The growth may occupy a small surface of the cord, being partially pedunculated, but it often has a considerable base, forming a shaggy, irregular outgrowth. Sometimes, after extirpation, these tumours are said to take on a sarcomatous character, or even to develop a cancerous growth.

A kind of false papilloma occurs, as we have already seen, in some cases of tubercular or syphilitic ulceration of the larynx. Sometimes, also, the surface of an epithelioma has a papillary character.

Next to the papillomas the **Fibromas** are the commonest tumours in the larynx. Morell Mackenzie has found in one hundred cases of non-malignant tumours, sixty-seven papillomas and sixteen fibromas. They are tumours of slow growth, mostly seated on the cords or at the base of the epiglottis. They consist of firm or soft connective tissue, those of firm consistence being the commoner. They are usually more or less



pedunculated, and their surface is generally smooth, although it may be irregular or even furnished with papillæ. They are usually small tumours from the size of a split pea to that of an acorn.

**Mucous polypi** occasionally occur, but are much less frequent than in the nares. They very commonly undergo transformation into cysts, and their most frequent seats are the epiglottis and the ventricles of Morgagni.

Other forms of simple tumours are uncommon, but cases of **Lipoma**, **Myxoma**, and **Angioma** have been met with. **Cartilaginous tumours** formed by outgrowth from the normal cartilages have been found. They are usually multiple and sessile. They may project considerably into the larynx, and, being covered with mucous membrane, may be mistaken for one of the commoner tumours mentioned above.

**Sarcomas** of the larynx are of occasional occurrence. They are usually of the spindle-celled form, but may be round celled. They may grow to a considerable size, and are, of course, prone to recur unless the whole larynx be removed along with the tumour.

**Cancer** of the larynx develops in the form of flat-celled **Epithelioma**. It is not infrequently a primary tumour, growing usually from the ventricular bands, but also originating in other parts. An epithelioma may also extend from a neighbouring part, particularly from the tongue. There is first a limited infiltration which extends in area, and gradually advances, involving the parts indiscriminately. Very commonly there is an abundant papillary growth on the surface so that there is a resemblance to the cauliflower cancer or to the papilloma. The central parts of the growth undergo ulceration while the disease is extending at the periphery. In this way great destruction of tissue may result, and the parts present great deformity.

Tumours are of very rare occurrence in the trachea.

**Foreign bodies and parasites** are very uncommon in the larynx. They give rise to violent expulsive efforts by coughing. Various articles may pass into the larynx, especially in children, such as buttons, peas, pieces of food, etc. Round worms have been known to lodge there. If a foreign body lodges in one of the pouches of the larynx it will excite inflammation.

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### C.—THE BRONCHIAL TUBES.

The diseases of the bronchi stand in close relation on the one hand to those of the larynx and trachea, and on the other to those of the lungs, and it is impossible to draw an absolute line of distinction on either side. The affections of the larger and middle bronchi are essentially like those of the larynx and trachea, and this is consistent with the fact that in structure they are similar. As we pass down the bronchial tree, however, the structure changes considerably. The cartilaginous plates become irregular and smaller, and finally disappear; the elastic tissue becomes more completely incorporated with the mucous membrane, so as to form a single layer which becomes thinner as the tube diminishes in calibre; the bronchial glands altogether disappear in the finer tubes. The bronchial tube thus becomes more simple in structure, and approximates to that of the lung alveoli. So it happens that in their diseases the finer bronchi are more allied to the lungs, and are often involved with them. We shall afterwards see that in acute catarrhal inflammation of the lung the disease often begins in a capillary bronchitis, and is sometimes described under that name.

**I. Inflammation of the bronchi, Bronchitis.**—1. **Bronchial catarrh, Ordinary bronchitis.**—This is an inflammation affecting the larger and middle-sized tubes, but not extending to any considerable degree into the finer ones. As to the conditions giving rise to it, much that has been said under nasal catarrh is again applicable here. It is to be added, however, that in many persons there is a special proclivity to recurrent attacks of bronchitis. This may be due to an inherited weakness, but is perhaps more frequently occasioned by an acute bronchitis which has left the bronchi considerably altered in structure, has rendered them, in fact, the least resistant part of the body. In such persons bronchitis may be set up by cold, by disorder of the stomach, or by some other trivial cause. Valvular disease of the heart frequently gives rise to bronchitis by causing a passive hyperæmia of the mucous membrane.

In the slighter forms of bronchitis the larger bronchi and the trachea are mainly affected. In fact, it frequently happens that along with a slight laryngitis there is a tracheitis and a bronchitis of the larger stems, without the name bronchitis being given. In the more definite cases of bronchial catarrh it is the middle-sized tubes that are chiefly involved. As already noted, capillary bronchitis associates itself with inflammation of the parenchyma of the lung, and will be considered in that section.



In order to appreciate the changes which occur in bronchitis it is necessary to refer to some points in the structure of the tubes. The mucous membrane is covered with epithelium whose superficial layer is cylindrical and ciliated. Beneath the epithelium there is a basement membrane which is separate from the proper mucosa under it. The mucosa is composed of connective and elastic tissue and is covered externally by a muscular layer, chiefly of circular fibres, but with some longitudinal ones. Outside this we have the sub-mucosa or adventitia, which is continuous with, and really forms a part of, the general interstitial connective tissue of the lung. In this loose adventitia the cartilages are embedded, and there are abundant serous and lymphatic spaces, which are in communication with those throughout the lung, the perivascular and others.

According to the careful description of Hamilton, acute bronchitis is accompanied first by great congestion, chiefly of the vessels in the proper mucosa. Edema of the basement membrane follows, and this structure becomes greatly thickened. The superficial layer of columnar epithelium is shed, but vigorous germination of the deeper layers occurs, abundant embryonic cells being produced. At the same time there is great infiltration of the mucosa, muscular layer, and adventitia, with round cells of inflammation. This infiltration may be very great, and as the adventitia communicates with the general interstitial tissue of the lungs, there may be a wide distribution of these cells, which pass on by the lymphatic channels to the lymphatic glands.

If the disease becomes chronic the inflammatory infiltration of the layers beneath the basement membrane continues, and there is the usual newformation of connective tissue producing an induration of the tissue. As in other productive inflammations the new-formed tissue often causes atrophy of the proper structures. In many cases the muscular coat is to some extent atrophied, although sometimes it is hypertrophied. The mucous glands are often considerably destroyed, and even the cartilages may be partially atrophied. These alterations in the walls of the tubes no doubt weaken them, and as they occur especially in long-continued cases of bronchitis, we can understand how in such cases the bronchial tubes sometimes undergo dilatation. Dilatation of bronchial tubes is not, however, a very prominent feature of ordinary bronchitis.

The **Exudation** in bronchitis is important as it forms the **Sputum**, and gives its characters to the latter. In the normal condition the mucous glands secrete enough mucus to keep the membrane moist, the secretion consisting of a glairy fluid containing a few leucocytes which are here called mucous corpuscles. The fluid owes its glairy or sticky character to the fact that it contains mucin, a chemical substance allied to albumen



and secreted by the mucous glands. This normal secretion catches and holds the minute particles forming the dust which we inhale with the air, and the vibratile cilia of the epithelium, acting towards the outlets, carry mucus and dust outwards, to be swallowed or expectorated. In bronchitis there is at first very little increase of the secretion, and it is chiefly concentrated, tough and scanty. As the disease progresses it becomes more abundant but less tough and less transparent, and this is brought about by the increase of the inflammatory exudation consisting of serous fluid and inflammatory cells. The degree of toughness depends on the proportion of mucin, and the degree of opacity on the quantity of cells.

The **Sputum coctum**, or ripe sputum, met with at the acme of the disease, is yellowish white or greenish and opaque. The sputum as seen in a vessel appears at first sight like pus, but it is much more tenacious. Under the microscope it also resembles pus, the field being crowded with multitudes of leucocytes. But the tenacity of the fluid is often shown by the manner in which the plastic leucocytes are altered in shape, being drawn out into oval or more elongated forms according as the tough mucus is drawn out. On adding acetic acid the usual development of nuclei occurs in the cells, and the intermediate fluid becomes markedly opaque from the precipitation of the mucin, which can be seen now in fine granules. With acetic acid the sputum assumes to the naked eye a whitish, opaque, and almost membranous character.

In chronic bronchitis the sputum is often very abundant, and is usually sero-mucous with comparatively few leucocytes, and hence transparent. There is a comparatively rich serous exudation with very little increase in the secretion of mucus. The sputum is not so tough as in more acute cases, and is often very frothy. In some very chronic cases there is an abundant cellular exudation—the sputum is almost like pus and has not the toughness of the sputum coctum. Large quantities of pus may thus be expectorated. In these cases it may be supposed that leucocytes are present in the walls and outside the bronchi as well as in the expectoration, and it is in them that we may look especially for dilatation of the tubes.

2. **Fibrinous bronchitis.**—This name, as well as that of **Plastic bronchitis** or **Bronchial croup**, is given to a condition of very rare occurrence and of rather obscure pathology, but yet of great interest. We have seen that in laryngeal and tracheal croup the fibrinous exudation sometimes extends down into the bronchial tubes, and that casts of these are formed occasionally. Taking the other end of the bronchial tree we find that in acute pneumonia the fibrinous exudation which forms in the lung alveoli commonly extends some distance into the finer bronchi, and so we find casts in them. But there are cases in which fibrinous casts form in the bronchial tubes independently, without any disease of the trachea on the one hand or of the



lung proper on the other. These cases are somewhat chronic in character, and the expectoration of casts occurs at intervals during months or even years. The casts are of a whitish grey colour and represent the bronchi of, it may be, a single lobe, with their ramifications, as we can see very beautifully by floating them out in water. Sometimes the fine ends of the casts are swollen out as if they had come from the alveoli. The casts sometimes present on section a stratified arrangement as if the fibrine had been deposited in layers. They show under the microscope fibrine with leucocytes.

The exact pathology and the source of this exudation are somewhat obscure. In some cases where death has occurred shortly after the expectoration of the casts, or where they have been found in situ, there has been little perceptible alteration of the mucous membrane. This has led some to suppose that the fibrine comes from the lung alveoli, and in many cases the lung tissue is considerably altered; there may be phthisis, or pneumonia, or collapse. But it is not apparent to what extent these may be secondary to the bronchitis. A possible indication of the pathology may be afforded by the fact that in a considerable proportion of the cases there has been hæmorrhage from the lungs.

**II. Stenosis and Dilatation of bronchi.**—1. **Narrowing or Obstruction of the bronchi** occurs under a considerable variety of different circumstances. It may be the result of inflammation in the bronchial wall itself. There is some swelling of the bronchial mucous membrane in all acute inflammations, and in the case of the finer bronchi, this along with the inflammatory exudation will cause serious obstruction. Hence, in capillary bronchitis (broncho-pneumonia) respiration is often seriously interfered with.

In **Asthma** there is an obstruction of the finer bronchi throughout the lungs. It is brought on by nervous agencies, but the exact process in the bronchi themselves is not quite determined. The nervous origin suggests spasm of the muscular coat of the bronchi, and the suddenness of onset and recovery are consistent with this. On the other hand, the long duration of the paroxysms in some cases throws doubt on the view that spasm alone is the cause. An acute hyperæmia, due to vaso-motor paralysis and accompanied by œdema, has been suggested as a more probable explanation. From whatever cause there is sudden stenosis of the finer bronchi, probably spasmodic at least to begin with. Leyden observed, in the sputum of cases of asthma, crystals similar to those found in the blood in leukæmia (Charcot's crystals, see under Leukæmia). He supposes that these crystals may set up a reflex spasm of the bronchial muscles, but this view has not been accepted.

Narrowing of the bronchi may be due to **Pressure or Encroachment** from without. **Tumours** not infrequently extend from the root of the lung along the connective tissue around the bronchi, and growing there they may compress and narrow a bronchus. In not a few cases the tumour infiltrates the bronchial wall, and may even grow through it,



so as to present itself in the calibre of the tube, thus narrowing it considerably. These conditions occur chiefly in the case of malignant lymphatic tumours of the mediastinum. Tumours by their mere pressure, or aneurysms impinging on a bronchus, may narrow the calibre.

**Foreign bodies** in the bronchi will produce obstruction, and a large one plugging a main bronchus may lead to serious respiratory trouble. Foreign bodies in the bronchi, however, may give rise to affections of a different kind to be considered in next page.

**2. Dilatation of bronchi. Bronchiectasis.**—By bronchiectasis is meant dilatation of bronchial tubes, a condition of somewhat frequent occurrence, but depending on a considerable variety of circumstances. The normal bronchial tube offers resistance to the dilating force of the air, chiefly by virtue of its elastic or dense walls, and the support which it receives from the distended pulmonary tissue around. If the walls be weakened, or the support of the pulmonary parenchyma withdrawn, or if the distensile force of the air or other contents be considerably increased, then we may have bronchiectasis. It will be seen that one or other of these predisposing conditions may occur under very various circumstances. A prolonged catarrh may cause atrophy of the elastic tissue of the bronchial wall, and so reduce the resistance. On the other hand, obliteration of the finer bronchi and lung alveoli in a district will cause the pressure of the air in inspiration to act especially on the bronchial tubes, the dilatation being here similar in its origin to that in complementary emphysema. Lastly, we may have the bronchi dilated from accumulation of secretion.

The various forms of bronchiectasis may be divided into two, namely, cylindrical and sacculated dilatations. The **Cylindrical form** occurs mainly where the causal condition has been more or less of a general one, such as a prolonged bronchitis. The bronchi, especially the middle-sized ones, are unduly wide, and their walls are thinned. The dilatation may be quite regular, but very often there are little bulgings or pouches. In the **Sacculated form** there has usually been a more local agent at work. The most typical sacculated dilatations are found in fibroid phthisis, under which heading they will be treated of more fully. They may be produced also by any cause which induces shrinking of the lung tissue. The wall of the bronchus is usually thinned, but it still presents a distinct membrane lining the sac. When cavities in the lung have a distinct lining membrane, they are generally of this origin, and they are especially so when the membrane is surrounded by lung tissue which approaches to the normal condition.

As the secretion stagnates in the dilated bronchi, whether cylindrical or sacculated, and may decompose, there may be great irritation and an abundant discharge of very putrid pus.



**III. Tumours of the bronchi and Foreign bodies.**—Primary tumours originating in the bronchi are very uncommon. **Cancer** not infrequently occurs, originating in the mucous glands, although the resulting tumour has rather the characters of cancer of the lung. (See under Cancer of the Lung.)

The bronchi are not infrequently affected secondarily, on the one hand, by tumours seated at the root of the lung, and, on the other, by those in the lung parenchyma. The bronchi also take part very markedly in tuberculosis of the lungs.

**Foreign bodies** in the bronchi give rise to lesions which often affect the lung as a whole and may lead to appearances and symptoms very like those of phthisis pulmonalis. When a foreign body of some size passes down the trachea and lodges in one of the larger bronchi, it becomes a centre of irritation. It irritates the bronchial wall, making for itself a cavity whose internal wall is ulcerated and discharges pus. The discharge stagnates in the cavity and decomposes, and as it is in direct communication with the bronchial tree, it is liable to be insufflated into the finer bronchi and even into the lung alveoli. The whole bronchial mucous membrane is converted into an inflamed and discharging surface, and the discharge is of a highly putrid character. Thus arise some cases of so-called **Fœtid bronchitis**. The inflammation of the bronchial wall renders it less resistant and bronchiectasis will often result, the putrid fluid stagnating in the dilated bronchi. There are thus cavities formed which may ulcerate and imitate those of phthisis. In the lung tissue itself there may be developed an acute or a chronic inflammation, the former having the characters of an acute phthisis and the latter more those of a chronic fibroid phthisis. The difference in the result will depend to some extent on the character of the foreign body. If it be a decomposable substance, such as a piece of meat or of bone with meat attached, it is more liable to lead to acute symptoms, while if the body is in itself inert the results may be more chronic.

These remarks are chiefly based on cases observed by the author, who is impressed by the frequency of serious disease of the lungs, resembling phthisis in many cases, brought about by foreign bodies in the bronchial tubes. A fuller account of two such cases is given by the author in his Lectures on Phthisis.

**Literature.**—BIERMER, Krankh. d. Bronchien u. d. Lungenparench., Virch. Handb., v., 1854; and on Bronchial asthma, in German Clin. Lect. (Syd. Soc.), 1876; GAIRDNER, On bronchitis, 1850; GREENHOW, On bronchitis, 2nd ed., 1878; PEACOCK, (Fibrinous bronchitis) Path. trans., v. 41, 1853; SALTER, *ibid.*, xi. 36, 1860; RIEGEL, Ziemssen's Handb., iv.; SOCOLEFF, Virch. Arch., lxi., 1877; HAMILTON, Path. of bronchitis, etc., 1883; CARSWELL, (Bronchiectasis) Illustrations, 1833-38; JÜRGENSEN, in Ziemssen's Handb. v.; LEYDEN, (Charcot's crystals in asthma) Virch. Arch., liv., 1872; OPITZ, Fremde Körper in Luftwegen, 1858; COATS, in Gairdner and Coats, Lect. to pract., 1888.



## SECTION IV.—CONTINUED.

## D.—THE LUNGS.

- D.—OF THE LUNGS. *Introduction, as to structure.* I. MALFORMATIONS. II. ATELECTASIS AND COLLAPSE. III. HYPERTROPHY. IV. PULMONARY EMPHYSEMA, 1. *Interlobular*; 2. *Vesicular*; causation, either substantial or complementary; anatomical changes, involving atrophy; effects of emphysema. V. DISORDERS OF CIRCULATION. 1. *Active hyperæmia*; 2. *Passive hyperæmia*, chiefly from cardiac lesions and hypostatic; leads to œdema. 3. *Embolism*; 4. *Hæmorrhage*; the *Infarction*, etc. VI. INFLAMMATIONS. *General observations.* 1. *Acute lobar pneumonia*; causation; epidemic prevalence. *Stages of engorgement, Red and Grey Hepatization, and Resolution*; occasional *Purulent infiltration*, etc. *Condition of pleura.* 2. *Acute broncho-pneumonia or capillary bronchitis.* 3. *Diphtheritic pneumonia*, 4. *Embolic pneumonia*, 5. *Chronic pneumonia*, characterized by *induration.* VII. GANGRENE. VIII. PHTHISIS PULMONALIS. 1. *Definition and historical resumé*, 2. *Causation*, 3. *Anatomical changes in (a) Caseous form, and (b) Fibroid form.* 4. *Extension of the tuberculosis*, 5. *Healing process.* 6. *Hæmorrhage*, (a) *early and (b) late.* 7. *Condition of pleura—pleurisy and pneumothorax.* 8. *General effects.* IX. DISEASES FROM INHALATION OF DUST. X. GENERAL TUBERCULOSIS, SYPHILIS, etc. X. TUMOURS and PARASITES.
- E. OF THE PLEURA. 1. *Affections of the circulation*, 2. *Acute pleurisy*, 3. *Chronic pleurisy*, *Pleural adhesions*, 4. *Tuberculosis*, 5. *Pneumothorax*, 6. *Tumours.*

**I**NTRODUCTION.—In examining the lungs after death we seldom meet with them in a perfectly normal condition. They may be abnormally adherent to the wall of the thorax, or unduly pigmented, or there may be cicatrices in them, or œdema, and so on. The explanation of this is perhaps that the lungs are peculiarly exposed to deleterious influences in two directions. The air passing into them is apt to carry irritating materials with it, and the blood circulating so richly through their tissue is liable to variations in its constitution and degree of pressure. We have already seen, for instance, that organic disease of the heart has serious effects on the pulmonary circulation, but apart from that, simple weakness of the heart may, as we shall afterwards see, have important effects on the lung. It must not be forgotten also that there is no organ of the body whose tissue is so intimately related to its blood-vessels as the lungs. These organs are little more than a congeries of blood-vessels with a sufficient supporting stroma. Any deleterious



substance circulating in the blood, therefore, is very prone to affect the lungs, especially if there be any special weakness in this direction. We have abundant illustration of this in the frequency of lung complications in the acute fevers.

There are one or two points in the anatomical relations of the lungs which should be kept in mind. They are supplied with two different sets of blood-vessels, those of the **Pulmonary artery** on the one hand, and of the **Bronchial artery** on the other. We should remember the distribution of these, and not confuse effects due to obstruction of the one with those due to obstruction of the other. Then we speak of diseases affecting the **Respiratory surface** on the one hand, and the **Supporting structures** on the other; that is to say, there are some diseases which affect the surfaces of the finer bronchi and of the alveoli, while others involve the walls of the alveoli and of the bronchi and their supporting connective tissue. It is true that these two are generally involved together, but in different cases the one or the other is primarily concerned, and usually retains the lead. It is obvious that to a certain extent the determining cause of the disease will have something to do with this. An agent which acts by being carried into the lungs with the air will mostly affect the surface of the alveoli and bronchi in the first place, whereas an agent arriving by the blood will be more apt to attack the walls. We shall see, however, that to this there are important exceptions, because, on the one hand, the capillary vessels have such close relations with the surface of the alveoli, and, on the other hand, substances arriving from without very readily penetrate into the substance of the lung. A more important consideration arising from the anatomical relations has reference to the distribution of a lesion in larger or smaller districts of the lung. When a disease is caused by something arriving by the blood then we would expect it to be distributed over a large extent of lung tissue or over the whole; the disease will occupy whole **Lobes**. If it arrives by the bronchial tubes, we would expect it to extend to the proper lung tissue more irregularly, here and there an extension corresponding with a particular minute bronchus; the disease at first involves **Lobules**. In this way we may distinguish roughly diseases which are lobar as due to alterations in the blood, and those that are lobular as related to the bronchial stems.

The **Lymphatics of the lung** are important as they often convey different kinds of solid particles and are the means of dissemination of these. The lymphatics are present wherever connective tissue exists. They exist in the walls of the alveoli and in the interlobular tissue, in the subpleural tissue, and in the connective tissue around the bronchi and vessels, where they form the peribronchial and perivascular lym-



phatics. The internal surface of the alveoli is closely related to the lymphatics, so that solid particles readily pass from the alveoli into the lymphatics and may be carried thence throughout the lymphatic system of the lung, and on to the bronchial lymphatic glands. (See under Inhalation of Foreign Substances).

### I.—MALFORMATIONS OF THE LUNG.

These are not frequent, and are of minor importance. Apart from absence or exceeding smallness of one or both lungs, which occurs as part of general malformations, there are cases where single lobes have been wanting, and their place taken by cicatricial tissue. These have probably their origin in obliteration of a bronchus in early foetal life. Congenital non-inflation of the lung parenchyma may be associated with a complementary dilatation of the bronchial tubes and with a compensatory hypertrophy of the inflated lung. (See under Hypertrophy of the Lung.) Again, the lungs may be normal in form, but very small in size. In such cases the whole body, and especially the circulatory system, will remain ill-developed.

It is not uncommon to meet with abnormal lobulation of the lungs, the regular lobes being divided by the formation of deep fissures. Rokitansky has described a case in which an accessory lobe existed between the base of the left lung and the diaphragm, and quite separate from the lung. It had no bronchus, however.

### II.—ATELECTASIS AND COLLAPSE OF THE LUNG.

These names designate conditions in which the lung alveoli and finer bronchi contain no air, but are in a condition similar to that of the foetal lung before inflation, the internal surfaces of the alveoli being applied to each other. It may be a survival of the foetal state, or it may be subsequently produced by the alveoli being, in some way, emptied of their air.

**Atelectasis.**—In its strict sense this term is applied to an imperfect expansion of the lung at birth. It is frequently found in new-born children, being, indeed, a survival of the foetal state. The lungs have to a greater or less extent remained uninflated. The non-inflation may be due to some obstruction in the bronchi, by meconium or mucus, but in most cases it is merely due to the weakness of the inspiratory efforts. The new-born child usually cries lustily, and in the deep inspiratory gasps between the cries the lungs are fully inflated. But if the child be weak or has not cried freely, certain parts of the lungs are apt to remain devoid of air. The atelectasis of the new-born is most fre-



quent in the lower lobe, and in the posterior parts of this lobe. It may be only in small areas in the midst of the inflated lung tissue, or the greater part of a lobe, or the whole lobe may be affected. In any case the non-inflated part usually shows by its shape that it is the district supplied by one or more bronchi.

The atelectasis shows itself by the smaller volume of the part. If it is in the midst of inflated lung it is depressed below the surface. Like the foetal lung, it is redder than the normal, firmer to the touch, and non-crepitant when handled. It is important to distinguish this condition from condensation of the lung, for which it is liable to be mistaken. In both the lung is devoid of air, but in the case of condensation it is so because the air spaces are filled up with solid material, usually inflammatory exudation. The atelectatic portion is capable of inflation by blowing air into the lung by the bronchi, at least it is so at the outset, while the condensed part is not inflatable.

There is no doubt that a lung which was partly atelectatic immediately after birth may subsequently become perfectly inflated. On the other hand, there is reason to believe that, if the atelectasis persist long, the lung becomes incapable of inflation. If the child survive, the applied walls of the lung alveoli adhere and an actual obliteration of the latter occurs. The part gradually atrophies, and it has been supposed that cicatrices sometimes seen in the adult lung and without any obvious cause may have this origin.

**Bronchiectasis** occurs in consequence of the atelectasis, the bronchi dilating to fill up the space left by the non-inflation, so that the bronchiectasis is complementary. In a case observed by the author, and referred to further on under Hypertrophy of the Lung, the greater part of one lung had failed to inflate, and the bronchial tubes were dilated into considerable sacs.

The question arises here whether, after inflation of the lung, portions or the whole may again collapse. We shall see immediately that collapse occurs in the adult, and there is no reason to suppose that it does not occur in the new-born infant. There are undoubted cases of children who have lived over twenty-four hours and have cried, in whose bodies the lungs have been found with only an island here and there of inflated lung.

**Collapse of the lung or Apneumotosis.**—This is an emptying of the lung of air at any time after its expansion. In some cases it is due to direct **Compression of the lung**; the air is simply squeezed out of it. Usually this arises from the presence of fluid or air in the pleural cavity. In that case the air vesicles may be only partly emptied and readily recover. But if the exudation is great and remains long, then the lung may be pressed upwards and backwards and come to form merely a red



fleshy layer flattened against the chest wall. This condition is often called **Carnification**, and in it the tissue appears darkly pigmented, the absence of blood and the packing together of the lung tissue exaggerating the existing carbonaceous pigmentation. The cause mentioned above is by far the most frequent in producing collapse by pressure, but there are others. Curvature of the spine sometimes causes such a narrowing of a part of the chest that the lung is squeezed or collapsed. Aneurysms or tumours may also compress the lung, but they more frequently cause collapse by obstructing the bronchi. Even distension of the abdomen, by pressing the diaphragm upwards and limiting the chest space, may cause a partial collapse. A great distension of the pericardium may have a similar effect.

A very important cause of collapse is **Obstruction of bronchi**. To a limited extent collapse is exceedingly frequent in bronchitis, much more frequent than is usually supposed. The collapse very often appears in the form of small wedge-shaped depressions at the edges of the lungs, and may be almost concealed by neighbouring emphysematous lung. But sometimes, especially in children, the collapse may be much more extensive.

One mode in which this collapse occurs has been described by Gairdner. If a pellet of mucus obstructs a bronchus it may act to a certain extent like a ball valve; it is pushed out into the larger tube during expiration, and being drawn back against the bifurcation in inspiration stops the tube. In this way the escape of air during expiration is allowed, but the entrance of air during inspiration prevented. The respiratory movements will thus act, to a certain extent, like an air pump, and the portion of lung tissue concerned will be gradually emptied of air.

Another way in which collapse probably occurs when a bronchus is obstructed has been demonstrated by Lichtheim. It is to be remembered that the lung tissue is elastic, and that left to themselves the alveoli collapse and their walls apply themselves together. If a bronchus be obstructed, therefore, and communication with the external air withdrawn, the absorption of oxygen will soon reduce the bulk of the contained air, and cause a reduction of its elastic pressure. The elasticity of the lung tissue, again, will cause pressure to be exercised on the air, and absorption of its gases will be thus promoted. It has been proved that such absorption actually occurs somewhat rapidly, first of the oxygen, then of the carbonic acid, the nitrogen being slowest of absorption. The lung of course collapses as the air is absorbed.

It has just been stated that collapse from bronchial obstruction is most common in the bronchitis of children, but it is necessary to observe that in children, in whom the form called capillary bronchitis is common, it is often accompanied by a condition which is apt to be mistaken for collapse, namely, lobular condensation. The catarrhal process in the bronchi readily passes in children to the lung alveoli, and the catarrhal products fill these up, causing condensation of a portion of the tissue which has a wedge-shaped configuration similar to that of the collapsed portion. Of course, these two conditions may co-exist in the same lung, or we may even have a combination of them, the collapsed lung becoming the seat of catarrh, and so passing into the condition of condensation.



**Literature.**—HELLER, D. Arch. f. klin. Med., xxxvi.; COATS, Trans. of Clin. Soc. of Lond., 1884; GAIRDNER, On bronchitis, 1850; LICHTHEIM, Arch. f. exper. Path., 1879.

### III.—HYPERTROPHY OF THE LUNG.

This occurs as a **Compensatory** process perhaps more frequently than is usually supposed. There is evidence to show that in persons who go to reside in high altitudes, the chest increases in size, the greater requirements of the attenuated air apparently inducing increased respiratory movement and by degrees a permanent enlargement of the lung. It is also probable that in cases of phthisis which recover with loss of a certain portion of lung substance, the loss may be partly made good by a true hypertrophy of the lung, although there is also observed in such cases an emphysema by which the space is filled up without a proper hypertrophy. Hypertrophy in cases of phthisis will be promoted by residence at a high altitude.

An unequivocal compensatory hypertrophy of the lung occurs in **consequence of atelectasis**. There may be in this case a great enlargement of the expanded lung so as partly to fill the place of the non-expanded part. In such cases there is not probably any numerical increase of lung alveoli, but these are enlarged, their walls expanded, and the capillaries elongated or multiplied.

A case occurred to the author which distinctly manifested the characters mentioned above. The left lung was of very small dimensions, especially the upper lobe, which appeared merely as a membranous structure in which dilated bronchi could be felt. There was no pigment in this lobe, and not a trace of lung parenchyma. The lower lobe contained the ordinary carbonaceous pigment in limited amount, and was greatly reduced in size.

The right lung was of very unusual volume, extending across the middle line, so that its edge reached beyond the left nipple. The enlargement seemed to be due to an addition of lung tissue which threw forward the anterior parts. The supra-clavicular part of the lung was normal and its anterior border indicated the position of the normal anterior margin. Below and beyond this, however, the lung extended into the other half of the chest. This part of the lung had not the appearances of emphysema, but was bulky and consisted of sound vascular tissue. The right ventricle of the heart was greatly enlarged, and the pulmonary artery showed a remarkable thickening of its wall, there being scarcely any difference between it and the aorta.

The absence of pulmonary tissue in the upper lobe of the left lung indicated that the collapse had been of long standing, and the entire absence of pigment showed that if not congenital it had occurred in very early life. The enlargement of the right lung was thus compensatory, dating from a period when the organ was in a state of growth. The compensation seems to have been somewhat effective, as the person lived to the age of 46, and it was only during the later months of his life that he suffered from serious dyspnoea, followed by signs of venous engorgement. The remarkable thickening of the pulmonary artery without any appearance of atheroma, also seems to point to a compensatory hypertrophy of the right ventricle during the



period of growth, the vessel in its growth accommodating itself to the increased blood-pressure. It may be added that the left pulmonary artery had only about a third of the calibre of the right, but the main bronchus was of equal size on the two sides. A somewhat similar case is described by Recklinghausen.

**Literature.**—COATS, *Lect. to pract.*, 1888; SCHUCHARDT und RECKLINGHAUSEN, *Virch. Arch.*, ci., 1885, p. 71.

#### IV.—PULMONARY EMPHYSEMA.

This name includes two distinct lesions, in one of which air escapes into the connective tissue of the lung and distends the connective tissue spaces, while in the other the alveoli are over-distended and various other changes result, but the air does not escape from its natural chambers. It will be seen that the first form is comparable with surgical or cutaneous emphysema, while the second is essentially different.

**1. Interlobular or Interstitial emphysema.**—This form, which is very rare, occurs when the air vesicles are ruptured and the air escapes into the interstitial tissue. The air vesicles may be actually torn open by a broken rib coming against the lung, or by the lung being directly wounded. On the other hand, the air vesicles may rupture from acute over-distension. Thus it may be the result of very violent expiratory efforts, generally with, but sometimes without, obstruction of the air passages. It has been met with in whooping-cough, in diphtheria, and in violent coughing from the inhalation of irritating material. The violent efforts with closed glottis cause such compression of the air in the alveoli that at some place the vesicles rupture.

As the parts are seen after death the air appears in the form of minute rows of bead-like bubbles, visible through the pleura. These rows of beads demarcate the lobules. Occasionally there are larger bullæ, which have been known to rupture externally, and so lead to pneumothorax.

The air sometimes travels along the connective tissue for some distance, just as in the case of **Subcutaneous or Surgical emphysema**. It may pass to the root of the lung, and from there up along the trachea and out to the subcutaneous tissue of the neck, and so lead to a surgical emphysema. This has in some cases induced a mistake in diagnosis, as Virchow has pointed out. Interlobular emphysema sometimes occurs in diphtheria, and may lead to subcutaneous emphysema in the way just mentioned. But if tracheotomy has been performed, it may be thought that the emphysema has taken origin in the wound.

**2. Vesicular emphysema.**—In this condition the air vesicles are over-distended, and, by partial atrophy of their walls, to some extent coalesced, but without any actual tearing of them.

**Causation of emphysema.**—There have been differences of opinion as to the exact manner in which the over-distension is brought about. On the one hand, Gairdner asserted that it was produced during inspiration,



while Jenner held that it was due to the expiratory force. Probably each of these has its influence, and the cases may be divided according as they are due to repeated and severe expiratory efforts as in coughing (*Substantial emphysema*), or to inspiratory distension, acting especially on one part in consequence of another part being uninflated (*Complementary or Vicarious emphysema*). In the former case we have a more generalized and in the latter a more localized emphysema.

**Substantial or Substantive emphysema** is due to over-distension during expiration. When the glottis is closed, and **Expiration** violently performed, then the air in the vesicles will be at an increased pressure. The expiratory effort is produced by the muscles causing the moveable walls of the chest to be pressed against the lung. The contraction of the abdominal muscles presses the diaphragm upwards, while the ribs are depressed. The lung is thus compressed, but at the same time it is supported by the structures which compress it, and the same force that increases the pressure helps the lung tissue to resist it by increasing the support. When the glottis is closed the whole lung may be regarded as one cavity, and the pressure will be universally diffused. It may be expected, therefore, that if any part of the lung be insufficiently supported the distensile force will tell especially there. The question therefore arises, Are there any parts where the lung is not fully supported by the chest walls? If a deep breath be taken, the glottis closed, and the act of expiration vigorously performed for a few seconds, we find in our own feelings indications that the lungs are over-distended, mainly at the anterior parts and the parts above the clavicles. We can easily understand how this should be. The anterior part of the chest, by reason of the flexibility of the costal cartilages, is more moveable than the rest of it, and the anterior edges of the lungs, as we can see in an animal whose chest is laid open while artificial respiration is carried on, have very free play. Then, above the clavicle the lung is obviously less supported than where the chest has bony walls. We shall see afterwards that these, with one or two other parts, are those in which emphysema, when due to frequent expiratory efforts, occurs most typically.

It will be obvious that if the lung tissue has lost in elasticity permanent dilatation of the air vesicles will occur more easily, and will readily extend to parts more fully supported than those mentioned. In the form to be afterwards referred to under the name of Senile Emphysema there is a loss of elasticity from the atrophy of old age. But there seems to be in some cases a general loss of elasticity, and it appears as if such a condition were hereditary to some extent. It is true that frequent over-distension will cause atrophy of the elastic tissue of itself, but it is quite apparent that in many individuals there is, to begin with, less elastic tissue, or it is less resistant, and so we have a predisposition to emphysema, sometimes inherited.



A similar loss of elasticity may be induced by disease, as in a case by Hertz in which a cornet player developed a marked emphysema after an attack of pneumonia.

Emphysema is liable to occur when frequent and violent expiratory efforts are made with closed glottis. Coughing implies such efforts, and it is chiefly in diseases where coughing is a prominent feature that we are to look for emphysema from this cause. It is met with pre-eminently in bronchitis. It also occurs in whooping-cough, occasionally in croup, and even in the violent expiratory efforts of parturition. As chronic bronchitis is specially a disease of more advanced life, we may expect that a preliminary atrophy of the lung tissue, implying a loss of elastic tissue, plays an important part in the production of emphysema in a large number of cases.

**Complementary or Vicarious emphysema** arises in consequence of over-distension during inspiration. If a part of the lung does not distend fully in inspiration, there must either be a falling in of the chest to a corresponding extent, or else an over-distension of another part of the lung in order to fill up the space. To what extent one or other or both of these will occur is determined by circumstances, chiefly by the situation of the insufficiently distended part and the time occupied in the occurrence of the lesion.

If a portion of the lung is collapsed, the neighbouring part often undergoes emphysematous distension. It occurs thus in bronchitis, and the wedges of collapse, already referred to, are often fringed with emphysema. We see it also around cicatrices in phthisis, or along with bronchiectasis in fibroid phthisis. Emphysema occurs sometimes to a remarkable extent in connection with general adhesions of the lung, and the emphysema is often very marked at the anterior parts. In this case the adhesions prevent the forward movement and expansion of the lung during inspiration, and there is thus an over-distension in this direction. The opposite lung may project so as to partly fill the space, and its edge may be also emphysematous.

**Anatomical changes in emphysema.**—We have to do with air spaces of irregular shape and separated by partial partitions, and as the distensile force acts from within, its tendency is to distend equally and so render the spaces globular. The ultimate bronchial tubes terminate in the elongated passages called from their shape infundibula. From these passages open, by wide apertures, the air vesicles, which therefore cluster around the infundibula and are mere pouches out from them. The distensile force will act first on the infundibulum with its system of air vesicles, causing distension. The tendency will be to render the outline more globular and the cavity simpler. The partitions separating the alveoli atrophy and the infundibulum expands into a simpler cavity



(*a a* in Fig. 249). Thus while more space is occupied there is less respiratory surface. The infundibulum continues to expand and meets

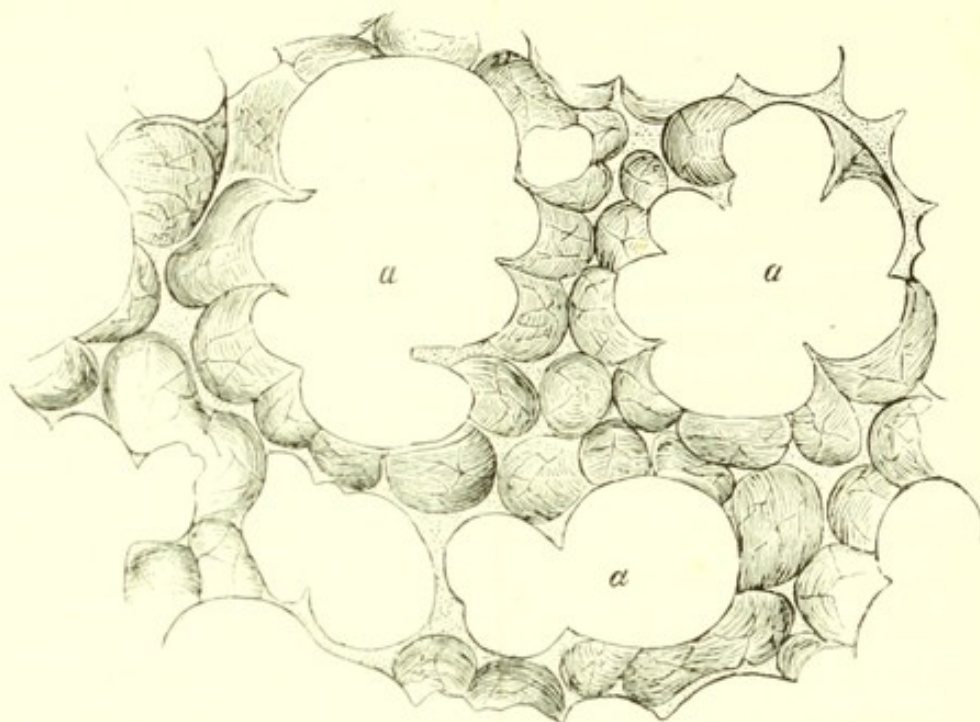


Fig. 249.—Section of the lung in emphysema, early stage. The infundibula, *a, a, a*, are dilated.  $\times 100$ . (RINDFLEISCH.)

with other infundibula undergoing a similar process. By mutual pressure the adjoining tissue atrophies, and the infundibula communicate by such rounded apertures as are shown in Fig. 250. The larger cavity

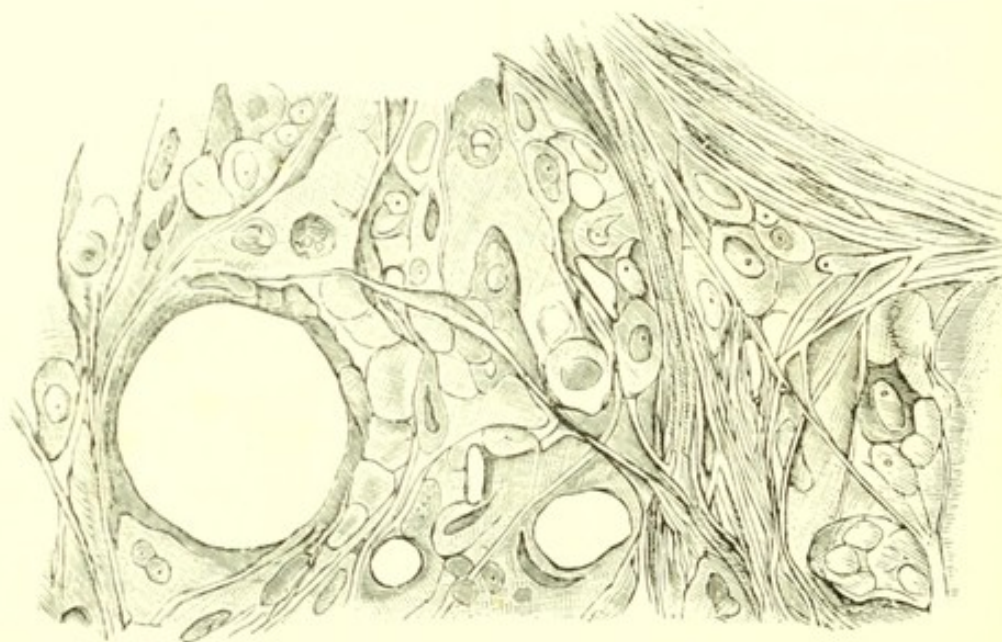


Fig. 250.—Emphysema of lung, portion of the wall of an alveolus. There are several rounded apertures. In some parts, and especially at right hand corner, an aperture seems to be forming by the separation of epithelium which has come to form the entire thickness of the alveolar wall.  $\times 600$ . (THIERFELDER.)

tends to become simpler by atrophy of all partitions and septa, and so the process goes on by coalescence of neighbouring spaces and atrophy of



the partitions. The enlarged air spaces thus produced are of various sizes, up to considerable vesicles or bullæ.

With this atrophy there is great **loss of tissue and of blood-vessels**. Before the actual destruction of the blood-vessels the pressure seems to cause obstruction of them so that they are not able to be injected and appear as white cords. It is stated that this obstruction is effected by the formation of white thrombi. The epithelium for the most part persists in the distended air spaces, but it is frequently in a state of fatty degeneration. Of most importance is the atrophy of the elastic tissue, because on this account the lung tissue will be prevented from collapsing as it normally does.

**The lung as a whole** presents alterations consistent with those in its finer structure. In general emphysema, when the chest is opened, the lungs are seen to be **bulky and pale**; they do not retract within the chest, and they give to the hand a soft downy feeling. They scarcely crepitate when handled and when pressed on the surface a pit is left. These characters are due to loss of elasticity, which renders the lung tissue less capable of retracting. The localization of the lesion is indicated by increase of bulk and change of colour, the affected parts having a light grey or pinkish colour mottled with spots and lines of pigment. Thus the **Anterior margins** are unduly voluminous, and the edges rounded. The enlargement is often somewhat irregular so that pieces are partially isolated by being specially enlarged. This applies more particularly to the **Ear-shaped piece** which projects from the anterior border of the left lung corresponding with the lower part of the heart. **The Base** also, on account of the insufficient support of the diaphragm, frequently shows marked evidences of emphysema, in the form of irregular rounded prominences, and the diaphragmatic surface of the lung may be convex instead of concave. This is especially the case in the left lung where the diaphragm is not supported by a solid organ as it is on the right side by the liver. Again the enlargement is very frequent at **the Apices**. It has been pointed out by Jenner that the right lung often presents a special bulging posteriorly at a place corresponding to the space behind the trachea.

The localization will be more various where the emphysema is complementary, and in that case will depend greatly on the locality of the cause. Thus we see it **around cicatrices** or contracted pieces of lung, but in nearly all cases there is a tendency to affect especially the anterior parts.

In all these places, with increase of bulk in general there is, even to the naked eye, a visible enlargement of the air spaces. Normally, the individual air vesicles are scarcely visible, but in emphysema the



abnormal air spaces are plainly seen through the pleura, often giving a frothy appearance from the size of the vesicles and the delicacy of the partitions. Beyond this we may have all degrees of visible enlargement of the air spaces up to considerable bladders as large as an egg. On cutting into them there is often a creaking, crisp sound, and if the vesicles are large they collapse markedly.

**The effects of emphysema.**—The lungs are permanently increased in bulk, and the **Capacity of the chest is increased.** The chest is kept more or less in the condition of deep inspiration, the diaphragm depressed and flattened, the shoulders elevated, the sternum pushed forwards and outwards, and the chest more or less barrel-shaped. The heart is depressed along with the diaphragm and placed more horizontally, while it is overlaid by the edges of the lungs. The liver is pushed downwards by the flattened diaphragm.

Emphysema has a serious **Effect on the circulation.** We have seen that there is obstruction and destruction of the blood-vessels in the lung, and this, reacting on the pulmonary artery, causes hypertrophy of the right ventricle, which may for a time compensate for the obstruction. But as time goes on the compensation is liable to become incomplete, and the general results of dilatation of the right ventricle and general venous hyperæmia follow. The prolonged strain will try the muscle of the heart and death commonly ensues from heart-failure, the muscular substance being not uncommonly found fatty.

**Senile emphysema.**—This is primarily an atrophy of the lung tissue corresponding with the general atrophy of old age. When the chest is opened the lung in this form retracts and falls backwards against the posterior wall of the chest. Hence the name applied by Jenner of “small-lung emphysema.” The normal weight of the lungs is about 44½ ounces in the male and 35 ounces in the female, but in senile emphysema they may be only two thirds of this weight. The air vesicles undergo changes similar in appearance to those in ordinary vesicular emphysema; their walls atrophy and the infundibula coalesce so as to form larger cavities, but this is by a simple process of atrophy. On examining the lung the increased size of the air spaces may be invisible at first, but if the lung be distended, or if after laying it open by incision it be floated in water, the large vesicles become visible. As old people are specially subject to bronchitis, senile atrophy may strongly predispose to the more ordinary form of emphysema.

**Literature.**—*Interlobular*—VIRCHOW, Arch. f. wissensch. Heilk. iii.; SACHSE, Virch. Arch. li., 1871; GÜTERBOCK, *ibid.* lii., 1871; *Vesicular*—LÉNNÉC, *Traité d'auscult. méd.* i., 348; MENDELSSOHN, *Der Mech. der Resp. u. Circul.*, 1845; GAIRDNER, *Edinb. Monthly Jour.*, xii. and xiii., 1851; JENNER, *Med. chir. trans.* xi. 1857;



RINDFLEISCH, Path. Hist.; THIERFELDER, Atlas d. path. Hist., 1872; HERZ, in Ziemssen's Handb., v. ii; EPPINGER, Prag. Vierteljahrschr. f. pract. Heilk., vol. cxxxii.; DURAND FARDEL, Malad. des vieillards, 1854.

#### V.—DISORDERS OF THE CIRCULATION IN THE LUNGS.

It is to be remembered that the pulmonary arteries are thin-walled, and that their muscular coat is comparatively inactive. Any increase of blood-pressure, therefore, will tell with full force on the pulmonary circulation. The capillaries are abundant and closely set, and are capable of great distension. Their dilatation will cause them to project into the air vesicles, and it is not improbable that they sometimes interfere seriously with the inflation of the air vesicles.

1. **Active hyperæmia.**—This is not frequently met with except as an accompaniment of inflammation or other affection of the lung. It may be produced by the inhalation of irritating vapours. A **Collateral hyperæmia** occurs when, in consequence of disease which has rapidly developed, a considerable portion of the pulmonary circulation is obstructed. In cases of pneumonia where the exudation in the lung alveoli exercises pressure on the vessels, the unaffected parts of the lung are commonly hyperæmic. Embolism of the pulmonary artery will also produce a collateral hyperæmia.

A more equivocal collateral hyperæmia is sometimes produced by the application of cold to the surface or the imbibition of large quantities of cold liquids. The effect of this is to cause great contraction of the arteries in the skin on the one hand, or the stomach and neighbouring parts on the other. The anæmia thus produced causes, when extreme, a rise of blood-pressure in the pulmonary circulation, an acutely developed hyperæmia, which, however, partakes of the nature of passive more than of active hyperæmia. There is in some such cases acute œdema, which may be accompanied by hæmorrhage. (See case by Hertz.)

2. **Passive hyperæmia and Œdema of the lungs.**—These two conditions are often associated in the lungs as well as elsewhere. As already pointed out, passive hyperæmia readily leads to œdema, and it may be said that though œdema does not constantly follow hyperæmia, it is very liable to ensue on that condition. Passive hyperæmia in the lungs is nearly always connected with functional disturbance in the heart, and there are two principal forms of heart lesion which are apt to give rise to it. There are, on the one hand cardiac lesions, chiefly valvular, which interfere mechanically with the pulmonary circulation, and on the other hand, weakness of the heart, which, as we have seen, is a cause of passive hyperæmia in general.



**Passive hyperæmia from cardiac lesions** occurs under all the different forms of heart disease in which the return of blood to the left auricle and ventricle is hindered by valvular incompetence or otherwise (see under Valvular Disease of the Heart). In mitral disease especially there is commonly a direct obstruction to the return of blood from the lungs. This is, in many cases, partly compensated by hypertrophy of the right ventricle, but even when it is so the pulmonary circulation will be at an increased pressure, and a more or less permanent passive hyperæmia will exist. This expresses itself during life in the readiness with which dyspnœa develops itself, and also in the frequent coincidence of bronchitis.

The appearances presented after death are expressed in the designation **Brown induration**, usually applied to this condition. The whole lung is more consistent than normal, and does not retract so fully when the chest is laid open. The colour is brownish, but the depth of colour varies considerably in different cases and in different parts of the lung.

In this, as in other cases of prolonged hyperæmia, the connective tissue is thickened and increased in density; hence the induration, which Rokitansky thought to be due to inflammatory hypertrophy of the connective tissue. At the same time the capillaries are greatly dilated and tortuous, and hæmorrhage by diapedesis is liable to occur as in passive hyperæmia generally. The blood escapes partly into the lung alveoli, giving rise to **Hæmoptysis**, and partly into the surrounding connective tissue, producing the **Brown pigmentation**. The pigment is in the form of brown granules in the connective tissue corpuscles.

The condition of the vessels is well shown by separating a piece of lung tissue by ligature, then placing it in nitric acid and afterwards in alcohol. When sections are made the varicose capillaries will be displayed as a brownish-red network, and the larger vessels will also be seen.

Even extreme and prolonged hyperæmia may exist without the development of œdema, the occurrence of which will usually imply the supervention of weakness of the heart.

**Passive hyperæmia from weakness of the heart** is commonly accompanied by œdema of the lungs. In many cases it is developed **shortly before death**, and is an expression of the fact that failure of the heart is the immediate cause of death. In a large number of debilitating diseases we find just before death the chest full of râles and the breathing much obstructed. In these cases passive hyperæmia and œdema are found after death. On the other hand, these phenomena may develop in the course of diseases which specially involve the heart, more particularly **the acute fevers**, in which passive hyperæmia and œdema are frequently of serious import. Existing cardiac lesions, as



already mentioned, will predispose to œdema if the heart happens to become exhausted or weakened.

Under these various circumstances gravitation plays an important part in the development of the hyperæmia and œdema, and these phenomena are usually most distinct in the parts which during life have been the most dependent. Hence the term **Hypostatic engorgement** is often applied. As a general rule, the parts affected are the posterior and basal portions of the lungs. They present after death a dark blue colour, and the tissue is bulkier and more solid than normal. The physical condition is expressed in the term **Splenization**, the tissue resembling that of the spleen. If the lung be incised and squeezed, a frothy fluid mixed with blood issues from the cut surface. Examined microscopically it is seen that the capillaries are distended and the lung alveoli occupied by serous fluid.

Hypostasis occurs as a post-mortem phenomenon, and this must be borne in mind in the diagnosis of hypostatic engorgement. Even in cases of sudden death from accident the posterior parts of the lung are often of a livid colour from the blood before coagulation having gravitated to the parts of the lung which, in the recumbent position of the body, were dependent.

When passive hyperæmia and œdema have persisted for a time they often pass into a condition closely allied to **Inflammation**. Fibrine and catarrhal cells are present in the lung alveoli, and the lung tissue becomes still more solid, approaching to the condition of hepatization. There is not, however, the complete solidification of proper pneumonia, and an excess of fluid, sometimes of a thickish grumous character, exudes from the cut surface.

**Œdema of the lungs** sometimes occurs **without hyperæmia**, or at least with very little evidence of the latter visible post mortem. This admits of somewhat ready explanation in cases of Bright's disease, where it is related to the general tendency to œdema. A simple œdema may also occur in consequence of multiple fat embolism in the lung. In other cases there is no other apparent cause than failure of the heart, and the absence of hyperæmia may be due to a general anæmia or to some local conditions such as pleural effusion.

In cases of simple œdema the lungs are pale and look bulky. On incision clear frothy fluid exudes, sometimes in large quantity.

Cohnheim has endeavoured to elucidate the pathology of œdema of the lungs by experiment in animals. An extreme passive hyperæmia may be produced by paralyzing the left ventricle, while the right ventricle retains its full powers. Cohnheim supposed that a weakening of the left ventricle might explain the hyperæmia and œdema occurring immediately before death. This explanation does not seem to the author to be a likely one. Great hyperæmia exists in cardiac lesions, produced in a somewhat similar fashion to that in these experiments, namely, by obstruction on



the left side of the heart, without any œdema, unless there is a failure in the cardiac contractions. It seems to be rather the extreme stagnation of blood, which such failure implies, that determines the œdema.

**Literature.**—ROKITANSKY, *Lehrb.* iii.; ZENKER, *Beitr. z. norm. und pathol. Anat. d. Lungen*, 1862; COHNHEIM, *Allg. Path.*, i. 501, 1882.

3. **Embolism of the pulmonary artery.**—The pulmonary artery is probably more liable to embolism than any other vessel. The embolus may be derived from thrombi in the right side of the heart or in the veins; it may consist of fat, forming an oil embolism, or it may have origin in tumours or parasites which have penetrated the veins.

In a considerable proportion of cases the hæmorrhagic infarction results, but by no means in all (see p. 76). In the case of **Fat embolism** there may be many small hæmorrhages with œdema, or even a more considerable area of hæmorrhage. (See pp. 78, 79, and Fig. 8.)

There are many cases of pulmonary embolism without any development of the infarction and, it may be, without any pronounced appearances at all. If the embolus be large enough to obstruct the main artery of the lung, then there can be no hæmorrhage as the whole lung is cut off from its blood supply through the pulmonary artery. When smaller branches are occluded, the infarction may be absent. Cases of sudden death after parturition occur from embolic obstruction of the lungs without any infarction. The suddenness of death in some of these cases seems to indicate that it is not due entirely to the obstruction in the lungs and consequent non-aëration of the blood. It seems rather due in great measure to the fact that, as the blood is prevented from reaching the left ventricle in sufficient amount, the brain and medulla oblongata are deprived of their proper nourishment. In such cases the right ventricle will be found dilated, while the lungs are pale and probably over-inflated by the violent but ineffectual respiratory efforts. These sudden deaths after confinement used to be explained as cases of shock before Virchow pointed out their true significance.

4. **Pulmonary hæmorrhage.**—Hæmorrhage occurs in the lungs under a considerable variety of different circumstances and presents many different appearances.

The **Hæmorrhagic infarction** is the form most frequently met with after death. It results from obstruction of the pulmonary artery, and this is in the great majority of cases due to embolism, but in a few may be the result of thrombosis. In cases of extreme passive hyperæmia, there may be such stagnation of the blood in the pulmonary artery in some parts as to induce coagulation, but this is very rare.

The pulmonary artery in itself and its branches is an end-artery, but various local circumstances frequently interfere with the formation of the infarction. The capillaries of the lung are wide, and may act in the same way as anastomosing vessels so as to keep up the circulation and prevent engorgement. In addition, the bronchial artery nourishes the lung tissue and even forms communications with the



pulmonary artery. The infarction is least likely to occur where the affected piece of lung tissue is completely surrounded by lung tissue whose capillaries communicate. The existence of the pleura at one or several surfaces will interfere with such communication, and hence the infarction is more common at edges, where two or three surfaces are covered with pleura, than in the substance of the lung or at its lateral and posterior aspects.

The hæmorrhagic infarction presents itself as a limited piece of condensed lung tissue, which may be often recognized by its dark colour seen through the pleura, but is more easily discovered by handling the lung, when the solid mass is readily detected. It is more or less wedge-shaped, the blunt end being at the surface. It is usually peripheral and most frequently at an edge of the lung. It presents great varieties in size up to nearly half the lung, but the most common size is from that of a hazel nut to that of a hen's egg. When recent, the infarction looks on section like a recent blood clot, and has almost a similar

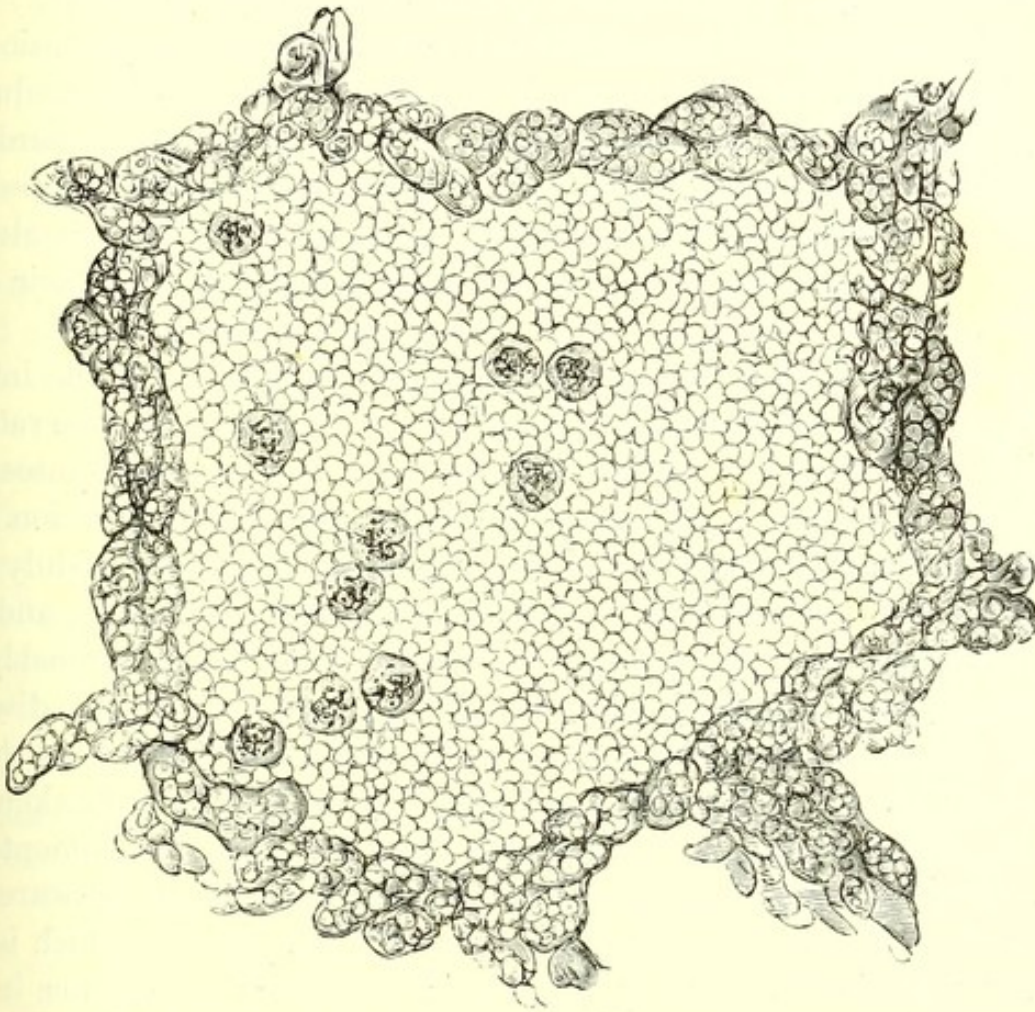


Fig. 251.—From a hæmorrhagic infarction of the lung. The alveolus is filled with red blood-corpuscles, with one or two large catarrhal cells. In the wall of the alveolus the capillaries are greatly distended with blood.  $\times 350$ .

smooth surface, from which circumstances the term **Pulmonary apoplexy** is used to be given to the condition. As time goes on the colour merges



into brown, and the appearance may come to resemble that of red hepatization. As blood fills the air spaces the piece of lung is more bulky than that surrounding it. The infarction, therefore, is often seen as a rounded bulging beneath the pleura, or when the lung is divided it stands at a higher level than the surrounding tissue. The pleura over the infarction is usually covered with a layer of fibrinous exudation of a yellow colour, and the pleural cavity contains fluid, often in considerable quantity.

Search should always be made for the obstruction of the artery. The artery may be filled with coagulum in the infarction itself, but the plugging generally extends beyond its apex, the actual embolism being usually some little distance on the proximal side of the infarction. An endeavour should also be made to find the source of the embolus.

Under the microscope, as shown in Fig. 251, we have simply an enormous aggregation of red corpuscles in the lung alveoli and finer bronchial tubes, with distension of the pulmonary capillaries. The absolute filling out of the alveoli with blood to the entire exclusion of air indicates that there has been a leakage from the capillaries gradually filling up the alveoli and expelling the air. There is usually hardly a trace of fibrine to be seen, merely red corpuscles which have escaped by diapedesis, and some catarrhal cells. Such cells are nearly always present, the blood irritating the epithelium and inducing their production.

If the patient live for some time after the occurrence of the infarction certain changes occur in it, but there are few actual observations bearing on this point. It is very probable that in some cases the infarction clears away; the blood is discharged by expectoration, and the circulation is re-established, but the portion of lung is unduly pigmented. In other cases the portion of lung gradually shrinks, and the ultimate result is a pigmented induration. Indurations, probably of this origin, are not infrequently met with in cases of valvular disease. In still other cases the portion of lung tissue dies and sloughs, so that a cavity forms. This condition has probably been often mistaken for phthisis, especially as it has probably been preceded by hæmoptysis. Sometimes the slough decomposes, and we have all the features of gangrene of the lungs. It may seem strange that necrosis, which is the regular result in other organs, is not of more constant occurrence in the lungs. It is to be remembered, however, that the lung tissue in the midst of the infarction is still nourished by the bronchial artery, and that, while the capillaries and actual walls of the alveoli may die, the interlobular connective tissue may survive, and may even use the necrosed tissue as pabulum.



Besides the hæmorrhagic infarction there are various forms of pulmonary hæmorrhage. We have already referred to the hæmorrhage which is a concomitant of passive hyperæmia. There are also hæmorrhages in scurvy, hæmophilia, hæmorrhagic small-pox, etc. Hæmorrhage is very frequent in phthisis pulmonalis, as will be subsequently described. It is asserted also that there may be rupture of a branch of the pulmonary artery from fatty degeneration of its walls, as we may have hæmorrhage in the brain from atheroma, but this is excessively rare.

**Hæmorrhage** is not uncommon from **Tearing or Rupture of the lung**, by a wound or a broken rib. The blood in this case will partly escape into the pleura, but it will also collect in the cavity torn in the lung and infiltrate neighbouring parts of the parenchyma. According to Rokitsansky blood thus effused may be encapsuled and subsequently infiltrated with lime salts like a foreign body.

When the hæmorrhage is from the bronchial mucous membrane or the lung alveoli, then it appears in the sputum. To some extent, however, the blood remains in the air passages, and if it be in considerable quantity it may even be carried by **Insufflation** into the lung alveoli. The blood, in this case, is mixed with air, and there is no such condensation as that which occurs in the hæmorrhagic infarction. In the lung alveoli the blood acting as a foreign body irritates the tissue and a catarrhal inflammation may result. In such cases large catarrhal cells occupy the alveoli abundantly wherever the blood has penetrated, and these cells may be deeply stained with the colouring matter of the blood.

It is stated that blood insufflated in large quantities may even cause necrosis of the lung tissue in which it lies, and so give rise to a condition like that of the infarction. This may, perhaps, occur when the lung is already seriously damaged, as in phthisis pulmonalis, but hardly in an otherwise healthy lung. It is also possible that blood may be drawn into cavities and form fibrinous plugs there.

**Literature.**—ROKITANSKY, *Lehrb.*, iii.; ZENKER, *Beitr. zur norm. u. path. Anat. d. Lungen*, 1862; HERTZ, in *Ziemssen's Handb.* v.; COHNHEIM, *Allg. Path.*, 1882, i. 501; GERHARDT, *Volkmann's lectures* (Syd. Soc. transl.), 1876, 2nd ser., p. 261.

## VI.—INFLAMMATIONS OF THE LUNG.

The inflammations of the lung vary considerably according to cause, distribution, and the structures specially affected. The irritant which leads to the inflammation may reach the lungs by the blood-vessels or by the air passages. In the latter case it will probably happen that the agent will affect certain bronchi and their connected alveoli so as to



produce a **Lobular** distribution. In the former case there may be, on the one hand, an **Embolic** distribution, or else a general diffusion over a wide tract of lung, such as a whole lobe or more, so as to give a **Lobar** distribution.

The character of the inflammation will depend on the nature of the irritant as well as on its distribution. We have acute inflammations, with excessive exudation, sometimes going on to suppuration; and we have chronic inflammations with newformation of connective tissue. We have also inflammations mainly affecting the lining membranes of bronchi and alveoli, and others involving the deeper structures. In nearly all inflammations the epithelium of the alveoli is more or less involved. In acute inflammations it is soon shed, while in the chronic forms it proliferates and yields large cells (catarrhal cells) which may accumulate in the alveoli.

The chief forms of inflammation may be considered under four headings, namely, acute lobar pneumonia, acute broncho-pneumonia, embolic pneumonia, and chronic or interstitial pneumonia. In phthisis pulmonalis many of the lesions are inflammatory, but, as we have here a true tuberculosis, the inflammatory manifestations will be considered along with the tubercular.

1. **Acute lobar pneumonia** (*Croupous pneumonia*).—This is essentially a disease of the lung alveoli and the most prominent feature is an exudation of fibrine, from which the name croupous pneumonia is derived.

The **Causation** of acute pneumonia has not been fully elucidated. The fact that a definite chemical irritant such as carbolic acid, when administered in considerable quantity, has more than once produced a typical pneumonia, seems to indicate that a poison in the blood is the cause. Many facts indicate that in ordinary cases of pneumonia some form of morbid poison, probably a microbe, exists in the blood and produces, on the one hand the acute fever and, on the other the local inflammation. For example, the fever often precedes the local lesion and hence can scarcely be regarded as a consequence of it. The fever has a definite course and usually ceases abruptly, while there is no such sudden change in the local inflammation. Then pneumonia is sometimes associated with other local inflammations, as with acute meningitis, and even with other forms of fever, such as typhoid, in which case it is to be presumed that the same irritant existing in the blood has produced the several effects. There is also the fact that pneumonia sometimes occurs in an epidemic form.

The question of the form of microbe involved in pneumonia has already been considered (see p. 303). Looking to the various circumstances under which pneumonia is met with, and its occasional occur-



rence along with other forms of inflammations and fevers, it seems probable that no single microbe is always the morbid agent.

Many epidemics of pneumonia have been observed in various quarters of the globe (see Hirsch). It is an interesting fact that occasionally epidemic outbreaks of fever of peculiar and unusual characters have been associated with acute pneumonia. Amongst these may be mentioned an epidemic which occurred in an industrial school in Glasgow, which in some of the earlier cases caused death within twenty-four hours of the onset, and which, in many of the cases which lived for a longer period or survived, was accompanied by a typical pneumonia. This epidemic could not be referred to any of the recognized forms of fever (see Russell).

The disease is an acute inflammation, and as the lung alveoli possess merely a single layer of pavement epithelium which is soon desquamated the inflammation resembles that of serous rather than of mucous membranes. As in the former we have here a fibrinous exudation, and though this occurs primarily and mainly in the alveoli, the fibrine, as we shall see, generally extends to the finer bronchi, forming casts of them.

Pneumonia is divisible into several stages, which, however, to some extent merge into each other.

In the first stage, that of **Engorgement**, there is an active inflammatory hyperæmia; the lung capillaries are highly injected, and there is an exudation of serous fluid into the air vesicles. To the naked eye the affected portion of lung is of a dark red colour, to the touch it is inelastic, and the finger applied to the surface leaves a pit behind. On section a reddish serum flows out, and the tissue does not crepitate under the knife so much as in the natural state. This state, from the resemblance of the cut surface of the diseased lung to the spleen, has been called **Splenization**. So far as the merely anatomical condition is concerned the lung is very much in the same state as in passive hyperæmia and œdema. In pneumonia, however, the splenization is not localized in the dependent parts, but it affects a definite region of lung, generally the lower lobe as a whole, along with, perhaps, a portion of the upper. Like other inflammatory exudations, the serous fluid contains leucocytes and also red blood-corpuscles, sometimes in large numbers. As the alveoli are filled with serous fluid, the air, bubbling in and out among the fluid during respiration, produces the fine crepitation which is the characteristic auscultatory sign of this stage.

In the second stage, that of **Red hepatization**, we have fibrine deposited in the alveoli. In consequence a coagulum (Fig. 252) comes to occupy the lumen of the vesicles and infundibula, instead of the mixture of serous fluid and air which is present in the first stage. The coagulum, like the fluid, contains abundant leucocytes and red corpuscles,



the former often so abundantly as almost to conceal the fibrine. The fibrine may be detected as a coarse network with interlacing fibres (Fig. 253). The capillaries of the lung are in much the same state of over-distension as in the first stage, and the lung parenchyma is likewise little altered.



Fig. 252.—Cast of small bronchus infundibula and air vesicles in pneumonia.  $\times 40$ . (CORNIL and RANVIER.)

The red corpuscles are present in the alveoli in very varying proportion. They are never entirely absent, but in general form the minority of the total cells present; in some cases they are equal to the white ones, in some more abundant. In very rare cases they are so abundant that the exudation has more the character of an ordinary clot than of a fibrinous exudation. In these latter cases, which may be described as **Hæmorrhagic pneumonia**, the lung itself has a deep red colour. These are mostly very severe cases, and imply a previous state of debility in the patient, very commonly referrible to alcoholic excess. In accordance with this exudation of red blood-corpuscles we have, in this and in the preceding stage, the rusty tinge of the sputum which is characteristic of pneumonia.

The appearance of the lung in this stage is somewhat different from that in the first. It retains its red colour, both from the continuance of the congestion of the capillaries and from the red corpuscles in the exudation, but it is now much firmer and heavier—it does not crepitate under the knife or finger, and it sinks in water—no air being any longer contained in the vesicles. On section from a sound part into a hepatized part, it is observed that the latter remains on a level while the sound part sinks away, so that the diseased part appears enlarged. Even on external examination the affected part of the lung looks bulky. On more close examination, the cut surface has not the homogeneous velvety character of the lung in splenization, but a coarse granular appearance, and this will be more readily seen on tearing the tissue and examining the torn surface with a lens. The granulations thus brought out are undoubtedly the plugs of fibrine with corpuscles which fill the air vesicles. On stroking the cut surface with the blade of the knife casts of the vesicles and infundibula may be obtained (as in Fig. 252). The finer bronchial tubes when laid open are generally found to contain casts of soft fibrine, as if the exudation had overflowed from the alveoli into them. The appearance of a section of such a lung has been compared, from its solid character, granular surface, and colour, to that of the liver, hence the name hepatization applied to this stage.

The solidified lung is a much better conductor of sounds than that



filled with air ; hence, during life, we hear the sounds of the trachea and bronchi much more distinctly than usual ; it is as if one put the stethoscope over the trachea itself.

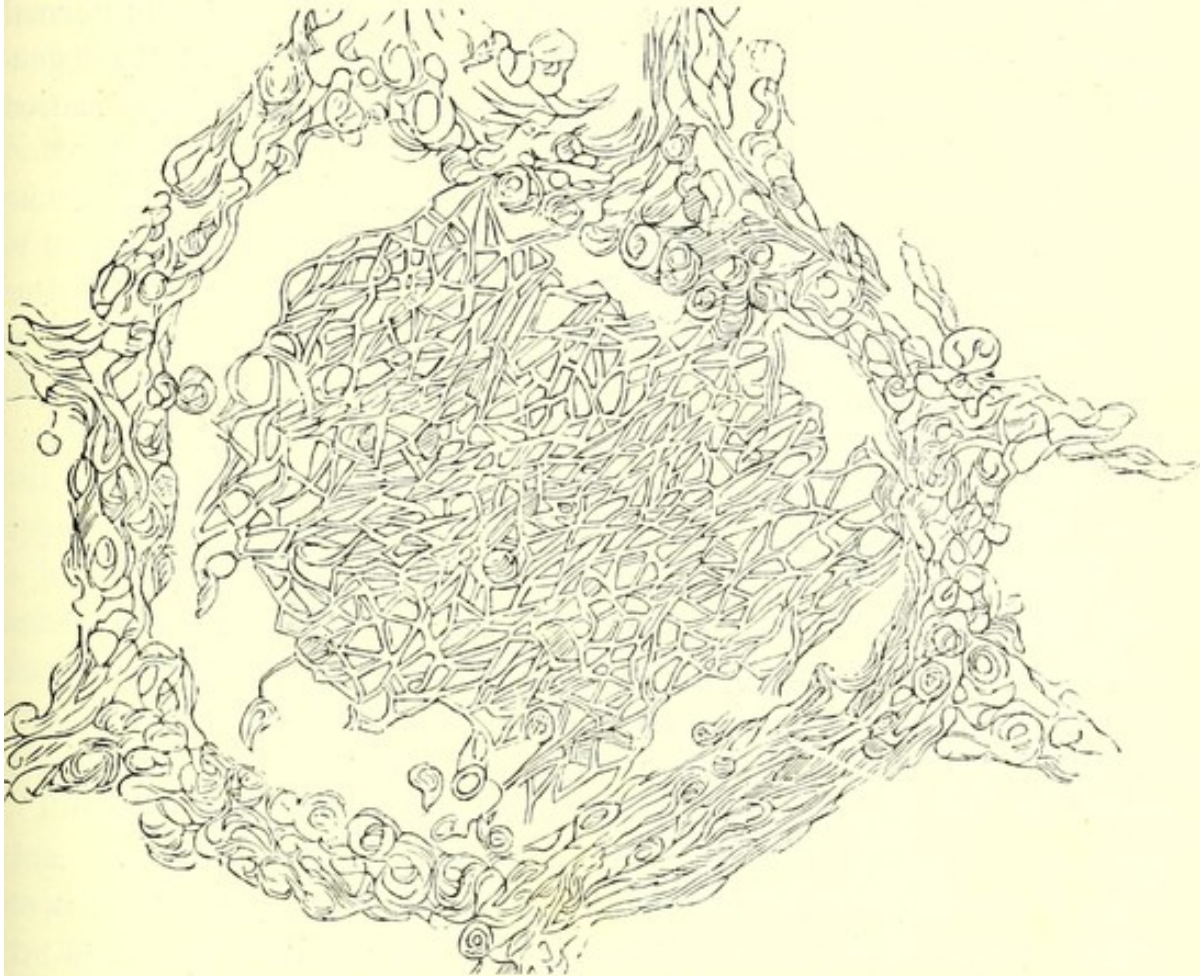


Fig. 253.—Alveolus filled with fibrine and a few leucocytes, from a case of pneumonia in stage of red hepatization.  $\times 350$ .

The third stage, that of **Grey hepatization**, develops naturally out of the second (Fig. 254). In the earlier periods red corpuscles are exuded largely, but except in cases of a hæmorrhagic kind the white very much preponderate in all periods, but especially in the later ones. The leucocytes swarming into the alveoli distend them more and more. The additional material in the vesicles also causes pressure to be exercised on the capillaries, which are thereby emptied. In this way we have, instead of the previous hyperæmia, an anæmia of the tissue. In accordance with the much less abundance of red blood, and the presence of an additional number of colourless cells, the colour of the tissue is changed. It retains the firm character, and the granular appearance of the previous stage, but the colour is grey. The pigment of the lung intermixed with the white colour of the multitudinous cells gives the appearance which has been aptly described as marbled.

It is not to be supposed, however, that during life the vessels are empty. It is always possible to inject the vessels after death, and during life no doubt the force



of the heart is sufficient to keep up the circulation. We are, therefore, scarcely warranted in saying that the grey colour of this stage is anything but a post-mortem appearance.

In the last stage, that of **Resolution**, the lung returns to its normal condition. The cells and fibrine in the air vesicles undergo fatty degeneration, and the plugs soften. Even in the stage of grey hepatization

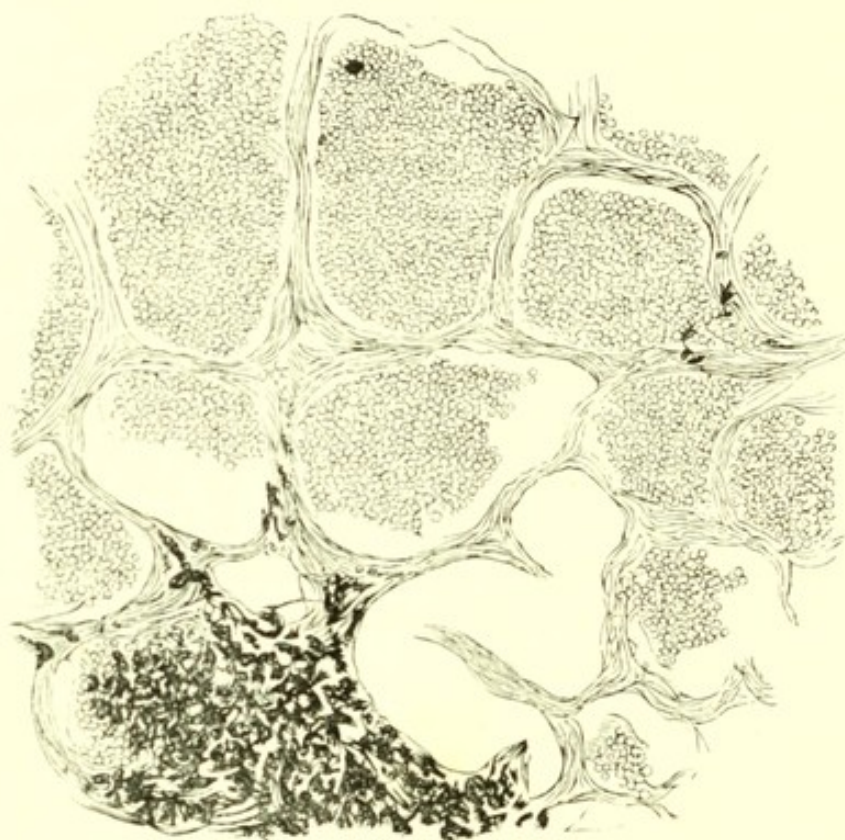


Fig. 254.—Section of lung in grey hepatization. The alveoli filled with plugs in which are multitudes of round cells. In the stroma the usual carbonaceous pigmentation of the lung is seen at lower part of figure.  $\times 90$ .

the leucocytes and fibrine have begun to undergo fatty degeneration, and this process progresses in the present stage. The fatty degeneration and disintegration of both cells and fibrine results in the conversion of the exudation into an emulsion which fills the alveoli, and, having a yellow or greyish-brown turbid appearance, resembles pus, in its naked-eye appearances.

The lung is still solid, still sinks in water, but its firmness is gone, its surface is pale, yellowish, or greyish red, it has lost the granular appearance, and a greyish, dirty fluid oozes out, which makes the surface so slippery that a small portion is with difficulty lifted or held in the fingers. The tissue is also extremely soft, and readily tears under manipulation. In removing such a lung from the body, unless care is exercised, pressure of the fingers may rupture the tissue, and as the pus or emulsion flows into the cavity it may give rise to the appearance of abscesses in the midst of the lung.



The softened exudation is now in a condition to be disposed of, and this is done partly by expectoration, but chiefly by absorption. The exudation must have been thoroughly softened before it will make its way through the narrow neck of the infundibulum into the bronchus, and the fatty emulsion is for the most part absorbed. It is remarkable how rapidly an extensive exudation in the lung may be disposed of. Sometimes, in the course of four or five days from the crisis of a pneumonia, the physical signs will indicate an approach to complete disposal of the exudation and return of air to the alveoli, and this may be almost entirely by absorption, almost no expectoration occurring in the interval. In some cases, however, expectoration materially aids in the disposal of the exudation.

After the infundibula and vesicles are emptied the blood returns in full force into the capillaries. Instead of the anæmia we have indeed a hyperæmia, for the tissue has been weakened by the inflammation, and is less able to resist the blood-pressure than formerly. It should be remembered in practice that the lungs of a pneumonic patient take some time to recover from the effects of the inflammation, and great exertion during convalescence should be warned against, till the tissue has recovered its tone.

**Purulent infiltration** is an occasional termination instead of resolution. In some cases, instead of the inflammation ceasing, it goes on in its acute form, and leucocytes continue to be exuded. In this case, when the fibrine breaks down, its place is taken by pus, and a true purulent infiltration occurs. The condition somewhat resembles that present while resolution is in progress, the fatty emulsion in that case resembling pus in its naked-eye characters.

The purulent infiltration in some cases results in **Abscess**, of which there may be several present. There may be rupture of a considerable vessel in the wall of such an abscess and profuse hæmorrhage. The abscess may subsequently burst into a bronchus or the pleura, in the latter case producing empyema, and perhaps pneumothorax.

Another unfortunate result which sometimes occurs is **Gangrene** of the lung. This is mostly met with in the hæmorrhagic form and in drunkards, but it may occur where there is a bronchiectatic cavity with decomposing contents.

There are some cases of pneumonia which seem to end in **Phthisis pulmonalis**, either in the indurative or in the caseous form. This is a very rare outcome, and it will probably occur only when a tuberculosis has established itself before the onset of the pneumonia. It may be said also that an acute tuberculosis may resemble in its course, and be mistaken for a pneumonia.



Another rare result is **Chronic pneumonia**, whose description follows below. It is more common than the termination in phthisis, and is probably not infrequently mistaken for the latter.

The **Pleura** always takes part more or less in the inflammation of the lung. The pleural surface of the inflamed portion of lung is coated with a white fibrinous exudation, which is sometimes of considerable thickness. There is rarely any considerable serous exudation in the pleura, probably because the lung distended with the solid exudation fills the cavity, and by its pressure prevents the accumulation of fluid. Sometimes the pleural exudation takes on a purulent character, and an empyema may remain after the resolution of the pneumonia. In some cases the inflammation extends to the pericardium, on the surface of which there may be a slight fibrinous exudation.

When resolution occurs the pleurisy will become chronic and the fibrinous exudation will be gradually absorbed. The result will usually be coalescence of the two pleural surfaces, adhesion of the pleura.

Pneumonia is an acute febrile disease, and so produces secondary changes in the organs of the body, generally comparable with those in acute specific fevers. There is commonly, but not always, enlargement of the spleen. The liver is usually enlarged, and shows parenchymatous infiltration.

In some cases of pneumonia the connective tissue of the mediastinum and sub-pleural tissue are the seats of inflammatory œdema, which may extend to the loose tissue between œsophagus and trachea, up to the retropharyngeal tissue, the soft palate, the tonsils, and even to the nares. Sometimes this inflammation assumes a phlegmonous character. This probably occurs by propagation of the specific microbe in the loose tissues (Weichselbaum).

**2. Acute broncho-pneumonia** (*Catarrhal pneumonia, Capillary bronchitis*).—This disease occurs most frequently in children, and is in them, as in adults, associated with catarrh of the finer bronchi. The bronchi are first affected, and so it may be said that the pneumonia springs out of a **Capillary bronchitis**, the tubes affected being those of the finest calibre. In a large proportion of cases the bronchitis originates in measles, or it may occur in diphtheria, or small-pox, or whooping-cough. In adults it may follow typhoid or other infectious fever, or it may be the result of the inhalation of irritating gases, or of the presence of decomposing material or foreign bodies.

It is to be noted that an ordinary bronchial catarrh, such as was described in the section on bronchitis, seldom goes on to a catarrhal pneumonia, but that for the most part the latter is due to the existence of some special irritant such as the virus of measles or decomposing juices in the bronchial tubes.

We have already seen that the presence of blood in the alveoli, acting as a foreign body, may cause catarrh. It is also interesting that a catarrhal pneumonia may be



produced in animals by **Division of the pneumogastric nerves**. The pneumonia here seems due to stagnation of the secretion which is no longer expelled, and to the irritation which this produces. In new-born children there is sometimes a broncho-pneumonia which may be due to insufflation of amnionic fluid or of impure secretions from the maternal parts.

The author has several times had the opportunity of observing how the **Insufflation of putrid juices**, it may be in cases of gangrene of the lung or of the presence of foreign bodies in the larger bronchi, has led to broncho-pneumonia, with a condensation considerably resembling that in phthisis.

As the disease begins in the bronchial tubes and is propagated to the lung tissue, it follows in its distribution the arrangement of the bronchial tubes; that is to say it occurs in a lobular form, hence the name **Lobular pneumonia** which is frequently used. Although the disease is thus primarily lobular, it is clear that it will often occur in several neighbouring lobules, and so a considerable tract of lung may be involved.

The inflammation manifests itself by the production of an exudation mingled with cells, so that the fine bronchi and the connected alveoli become filled. There is thus brought about a lobular condensation. The cells produced are partly leucocytes and partly catarrhal cells, the derivatives of the epithelium. The proportion of each of these will depend somewhat on the acuteness and nature of the irritant, but in acute cases each minute bronchus will often contain a small drop of pus. The affected pieces of lung are firm and reddish brown or grey, and on the cut surface stand above the general level. Even when a considerable district is generally affected the lobular appearance is usually visible.

As the disease begins in a catarrh of the finer bronchi, there is often **Collapse** of the corresponding portion of lung even before any actual inflammatory processes occur in it. The obstruction of the tubes with tough secretion sufficiently accounts for this collapse. It is a lobular collapse, and, as seen from the surface, areas of larger or smaller size are depressed and of a bluish red colour. These are mostly to be found in the first instance at the posterior and inferior parts of the lung. In other parts, and especially in the upper lobes, we may have emphysema.

In some cases the inflammation does not confine itself to the bronchi and alveoli, but may extend to the connective tissue of the lung generally. In such cases the exudation in the bronchi is composed of leucocytes, and cells of a similar character infiltrate the bronchial wall and neighbouring structures, the bronchi evidently forming centres of irritation.

In many cases **Acute pleurisy** accompanies the process in the lung (Fagge).

The inflammatory products in the air vesicles may after a time



undergo fatty degeneration, break down, and be discharged, some part of them being probably absorbed by the lymphatics. In some cases, however, the result is not in every part of the lung so fortunate. For one thing, the collapsed portions may remain uninflated, and as considerable tracts of lung may be affected, marked shrinking of the lung and deformity may result. Or, again, the catarrh may become chronic, and, being associated with an interstitial inflammation, result in a permanent induration, the alveoli being encroached on by the growing connective tissue, as in the case of chronic pneumonia. In this way, the person, though recovering, emerges from the disease with a damaged lung; according to the extent of the permanent damage will be the shrinking of the lung and possible displacement of organs. Again, phthisis pulmonalis may develop out of an acute catarrhal pneumonia, but this is fortunately an unusual result.

It has already been mentioned that the disease begins in the bronchi, and it has been asserted by Buhl that, throughout, the inflammation remains confined to the bronchi, the inflammatory products found in the alveoli being simply insufflated from the bronchi. This view, however, can hardly be maintained, as evidence of acute changes can actually be observed in the epithelium of the alveoli.

**3. Diphtheritic pneumonia.**—In many cases of diphtheria the exudation extends down the bronchi even to their finest ramifications, and sometimes also to the lung alveoli. The bronchi contain casts which do not generally obstruct the calibre entirely. These casts consist of fibrine and leucocytes, the latter in great abundance. Sometimes the finer bronchi generally are filled with leucocytes. There is also exudation of leucocytes into the air vesicles, but not usually fibrine.

**4. Embolic pneumonia.**—(*Pyæmia or Metastatic abscesses*).—In cases of septic thrombosis of veins (thrombo-phlebitis) in connection with wounds or abscesses, a septic embolism is liable to occur, in which the lungs are most directly involved. Pieces of the thrombi containing pyogenic microbes (generally *staphylococcus pyogenes aureus*) are carried to the lung and are caught in small arteries or capillaries.

Each such embolus becomes a centre of acute inflammation going on usually to suppuration and gangrene. If a considerable branch be obstructed there may be at first the regular hæmorrhagic infarction. But soon there is such an abundant exudation of leucocytes that the red colour is obscured, and the result is an area of grey hepatization. A similar grey hepatization will develop if there has been no hæmorrhage. The patch will be wedge-shaped or round according to the vessel affected. The grey hepatization gives place to a purulent infiltration, usually with gangrene of the lung tissue and the regular formation of an abscess.



Such metastatic abscesses are very various in size, form, and number. If the embolism be capillary there may be multitudes of minute abscesses, but very often there are only a few and these may be of some size.

If an abscess be near the surface it gives rise to an acute pleurisy which is apt to be suppurative in character. This may occur without the actual bursting of an abscess into the pleura, the microbes propagating through the inflamed wall into the pleural cavity.

5. **Chronic pneumonia** (*Interstitial pneumonia, Simple cirrhosis of the lung*).—We include here those conditions in which the lung tissue is the seat of a simple chronic inflammation, without anything of a tubercular or other specific nature. The simplest cases are those in which an acute pneumonia, instead of resolving, passes into a chronic condition. In old persons, again, ordinary pneumonia is apt to be chronic, and it may therefore be said that senile pneumonia is included under the present head. To a certain extent the same is true of pneumonia in drunkards and in debilitated persons, especially when it affects the apex of the lung. In addition to that we have a very important group in which chronic inflammation is set up by the inhalation of irritating solid particles, as among potters, stone-hewers, etc. (See further on.)

The chronic inflammation here, as in other organs, is chiefly characterized by a newformation of connective tissue, so that **Induration** of the lung is the result. In this view of it the terms chronic interstitial pneumonia, cirrhosis, sclerosis, are sometimes used as being virtually of the same meaning as chronic pneumonia. But here it is necessary to distinguish very carefully. We shall see afterwards that there is a form of phthisis pulmonalis in which a great newformation of connective tissue occurs, but the inflammation in that case is due to the tubercular virus. In the disease at present under consideration there is no such irritant present, and the inflammation is simple. The proper tubercular fibroid phthisis is a much commoner condition than the simple chronic induration.

The naked-eye appearances presented by the lung in cases of acute pneumonia which have had a prolonged course and have become chronic are not unlike those of the lung in the stage of grey hepatization. The disease is generally confined to one lung, and may affect only a portion of it. The lung is bulky and dense, and feels solid to the touch. When cut into, the solid lung has usually a grey colour, although sometimes with a tint of red, but it has a smoother cut surface than that in hepatization, and the tissue is much tougher. To this condition the name **Iron-grey induration** may be aptly applied.

Under the microscope the conditions are such as are indicated in Fig.



255. The walls of the alveoli are greatly thickened by fibrous tissue, which largely encroaches on the alveoli, the epithelium of which is preserved, and sometimes occupies their interior. When it is considered that the lung as a whole is not reduced in bulk, and that the alveoli are in great part empty, then the overgrowth of connective tissue will be understood to be very great. Along with the interstitial newforma-



Fig. 255.—Chronic pneumonia: The connective tissue is greatly increased, and the alveoli (a, a, a), are represented by contracted spaces lined with well-formed epithelium. The epithelium here is much more distinct than in the normal alveoli.  $\times 350$ .

tion there is commonly thickening and adhesion of the pleura. In the condition hitherto described, there is no considerable obliteration of the air vesicles except in so far as they are encroached on from without.

In some cases there is a very striking appearance as if the fibrinous plugs in the alveoli in acute pneumonia were being eaten into and replaced by connective tissue. There are obvious masses of connective tissue inside the alveoli, sometimes distinctly pedunculated. These have arisen in a manner similar to that in which a thrombus becomes organized, the new-formed tissue moulding itself on the fibrinous plug. This appearance establishes the fact that chronic pneumonia sometimes develops out of acute, a view which has been questioned by some.

If the disease progresses, the new-formed connective tissue takes on a cicatricial character and by its contraction destroys and contorts the proper lung tissue. Just as in the case of cirrhosis of the liver, there



is here an atrophy of the normal structure and a tendency in the organ to shrink. This leads to dilatation of the bronchi on principles already enunciated, so that **Bronchiectasis** is a prominent feature in advanced cases of this kind. The bronchial secretion may stagnate in the dilated bronchi and decompose; the irritation of the decomposing juices sometimes causes ulceration, and ragged cavities may thus form, so that the condition may come to resemble phthisis pulmonalis.

**Literature.**—LÆNNÉC, *Traité d'auscult. méd.*, 1819; ROKITANSKY, *Lehrb. d. path. Anat.*, iii., 1861; STOKES, *Dis. of chest*, 1837; FRIEDLÄNDER, *Ueber Lungenentz.* 1873; JÜRGENSEN, in *Ziemssen's Handb.*, 2nd ed., 1882, and *Volkmann's Lectures* (Syd. Soc. transl.), 1876; GAIRDNER, *Clin. Med.*, 1862; STURGES, *Nat. hist. and relations of Pneum.*, 1876; *Collective Invest. Record*, vol. ii., 1884; HIRSCH, *Geograph. und histor. Path.*, 1886, iii. 125; RUSSELL, *Peculiar outbreak of feb. disease*, 1888; LEYDEN, (*Abscess and gangrene*) *Volkmann's Samml.*, Nos. cxiv. and cxv., 1877.

## VII.—GANGRENE OF THE LUNGS

In this condition necrosis of a definite piece of lung tissue occurs. The necrosis is always accompanied or followed by decomposition, and the irritating character of the decomposing material plays an important part in the processes concerned. The gangrene may itself arise by the action of decomposing material. If a foreign substance, such as a piece of solid food, gets into a bronchus it may induce a bronchitis with putrescence of the secretion, and the irritation of the putrid juices may induce gangrene of the lung. Similarly, putrid juices inspired from ulcers and wounds of the mouth and air-passages, or perforation of abscesses or ulcers into the trachea or bronchi may set it up. Again, the juices in cavities, especially in those arising by dilatation of bronchi, may stagnate and decompose, and lead to gangrene. Wounds and contusions may cause necrosis directly. Sometimes the lung tissue dies in severe cases of typhoid fever or other zymotic diseases. We have also seen that gangrene may occasionally follow the hæmorrhagic infarction or acute pneumonia, and that it is a constant feature of the metastatic abscess. Lastly, there are some cases in which the cause of the gangrene is obscure, but these cases as well as those with a more definite cause, are somewhat common in debilitated persons and those given to alcoholic excess.

It is customary to divide gangrene of the lung into a circumscribed and a diffuse form. In both the lung tissue dies and decomposes, ultimately becoming separated, if the patient survive, as a shreddy slough, which occupies the cavity formed by the loss of tissue. In the diffuse form there are gangrenous patches throughout the lung, or a considerable portion of it, and there is little probability of the



effects becoming limited by reactive inflammation in the neighbourhood. The diffuse form not infrequently develops from the circumscribed, the decomposing juices from the slough causing still further necrosis.

The various changes which occur around a gangrenous piece of lung, and in more distant parts of the organ, are related to the irritating character of the slough. These changes are mainly inflammatory. The immediately neighbouring lung tissue is acutely inflamed, and there is thus a zone of condensation around having the usual features of acute pneumonia, often with a specially hæmorrhagic character. In this inflammatory zone the gangrene may advance. On the other hand the slough may be detached by the inflammatory process, and through time, a more chronic inflammation having occurred, the slough may be separated from the lung tissue by a layer of granulation tissue which produces pus abundantly into the interior of the cavity. If the slough be small enough, the cavity may, after the discharge of the slough, ultimately contract and form a cicatrix, but in the case of larger sloughs a suppurating cavity may long remain.

The effect on the bronchial mucous membrane is of importance. The decomposing juices from the slough and from the inflamed lung tissue find their way into the bronchial tubes, where they set up an acute inflammation of a highly suppurative character. A rich secretion of putrid pus is the result. This secretion carried to the bronchi in other parts of the lung may set up gangrene in numerous small isolated patches, and in this way multiple small abscesses may occur. If the gangrene be near the surface an acute pleurisy is the result, with fibrinous exudation. Sometimes the cavity opens into the pleura, and we have a suppurative pleurisy, perhaps with pneumothorax.

An occasional complication of gangrene is **Hæmorrhage**. As the slough separates the more resistant tissues retain their connection longest. The bronchi and larger vessels sometimes remain as rigid trabeculæ in the midst of the soft slough. The arteries remain longest in connection, but they are usually filled with thrombi and obliterated. Occasionally, however, the gangrene advances around an artery which is still pervious, and in that case hæmorrhage of a serious or even fatal character may result.

Sometimes the gangrene leads to a definite septicæmia, or metastatic inflammations result, having their seats especially in the brain. In these cases the decomposing material gets into the pulmonary veins, having first caused thrombosis of them.

A peculiar feature in gangrene of the lung is the very abundant and highly **Putrid sputum**. The decomposing juices from the slough set up, wherever they are carried, acute suppurative inflammations, and the abundant inflammatory products



also undergo decomposition. The bronchial tubes being weakened by the severe inflammation often undergo dilatation, and the material stagnates in them all the more, and decomposes. So it happens that in the cavity itself and in the dilated bronchi there are usually large quantities of putrid secretion. This is expectorated at intervals, and sometimes so abundantly that it pours out of nose and mouth. The sputum is extraordinarily foetid, and, if allowed to stand, deposits triple phosphates, crystals of margarine, etc. It also contains abundant pus-corpuscles, many of them broken down by decomposition, pieces of lung tissue, and bacteria isolated and in colonies. Sometimes the sputum contains also spirilla.

**Literature.**—LÆNNEC, *Traité d'auscult*; LEYDEN, *Volkmann's Sammlung*, No. 26, 1871; HERTZ, in *Ziemssen's Handb.*, v. 514, 1877; HANOT, *Progrès méd.*, 1876, No. 14.

### VIII.—PHTHISIS PULMONALIS. PULMONARY TUBERCULOSIS.

1. **Definition.**—The term phthisis pulmonalis was originally used to designate a wasting of the body associated with disease of the lung. In its modern use it is applied to cases in which the lungs are affected by a progressive lesion, the ordinary and regular result of which is destruction of the lung tissue and the formation of cavities. The idea of wasting is thus transferred to the lungs, and associated with the anatomical character of the lesion. It has always been recognized that tuberculosis plays a considerable part in the pathology of phthisis pulmonalis, but it is only of late years that a more complete demonstration has been furnished of the fact that virtually all cases conforming to the above definition are really cases of tuberculosis of the lung. There may be a few cases of actinomycosis, and possibly some of syphilis, which produce lesions somewhat similar in character, but they are so few that phthisis pulmonalis may now be regarded as synonymous with local tuberculosis of the lung.

It may be well here to refer briefly to the various phases through which the views as to the pathology of phthisis have gone since the time of Lænnec, especially as many of the terms in common use in connection with the disease are related to some of these views.

Lænnec believed that there was a particular tubercular matter which was liable to be deposited in the lungs or elsewhere. It might be deposited in isolated places, forming miliary tubercles, or infiltrated into a considerable portion of lung, forming infiltrated tubercle. In both cases the deposit usually began as a **Grey transparent** structure, which, however, was prone to change into a yellow or whitish material which was drier and harder. This yellow material was called **Yellow or Crude tubercle**, whether occurring in the isolated or in the infiltrated form. All cases were regarded as tubercular in which there were either isolated nodules or extensive infiltrations, whether these were grey or yellow.

By and by it came to be seen, however, that many of the conditions in phthisis are simply inflammatory. The minute histological characters of what we now call the tubercle were discriminated, and the essentially inflammatory processes were



sought to be separated from the tubercular. It was shown that the existence of caseous material is no evidence of tuberculosis, since the ordinary products of inflammation and other newformations, such as tumours, may undergo this change, which, in its essence, is really a necrosis with degeneration of the structures concerned. In phthisis, then, the process is largely an inflammatory one, with the special tendency in the products of inflammation to undergo a caseous metamorphosis. In this way arose Virchow's designation **Caseous pneumonia**—an inflammation with a caseous tendency in its products, just as scrofulous disease of the glands is an adenitis with a similar tendency.

When the lungs in phthisis were more particularly examined, however, it was found that the condition is not such a purely inflammatory one as Virchow's position would indicate. In all stages of the disease **Tubercles** are to be found alongside the inflammatory products. The tubercles undergo changes similar to these, and it is often difficult to discriminate between the two, especially when caseous metamorphosis has occurred. But, in nearly all cases where the disease is advancing, proper tubercles are to be found along with the inflammatory conditions.

The more modern position brings us back more nearly to that of Lænnec. Again we regard phthisis as a tubercular disease, but not merely in the general sense of Lænnec. We are to observe carefully the inflammatory processes and distinguish their effects on the lung tissue. Our position differs also from Lænnec's in respect that he regarded a particular state of the constitution as the essential cause of the tuberculosis. It is not to be denied that the lungs must be in a state of susceptibility before they can be affected by the **tubercular Virus**, but the same may be said concerning any form of tuberculosis, and indeed concerning ordinary inflammatory processes. We know that different persons, or the same person at different times, are very variously susceptible to catarrhs, and to inflammations of all sorts.

We are to regard phthisis pulmonalis as a **Local tuberculosis** in which inflammatory processes and the actual formation of tubercles play their parts, and both lead on to necrosis and ulceration.

2. **Causation.**—In what has been said above it has been implied that the causation of phthisis pulmonalis is connected with the tubercular bacillus. All that has been said in regard to the causation of tuberculosis at page 198 applies here.

There is, in this as in other forms of tuberculosis, not only the action of the specific microbe to be considered, but also the susceptibility of the individual, which may be inherited, but is often acquired. It is acquired principally in the case of persons so placed as to have the general health reduced, and in whom especially the respiratory functions do not get justice. Persons living in close dwellings, especially when, at their work in factories and otherwise, they are in the habit of breathing vitiated air, in which, it may be, finely divided dust is abundantly suspended, frequently acquire a tendency to phthisis although not originally predisposed.

In relation to the resulting lesions, the **Path of entrance** of the irritant to the lungs is a matter of importance. In the study of the lesions met with we shall find that they all start at the finer bronchi. A



catarrh of the finest bronchial tubes, usually occurring in a number of these simultaneously, is the starting point of a variety of lesions, which, however, for a considerable time remain related to the bronchi in their distribution. This is an indication that the agent finds access to the lungs by the inspired air.

As a general rule the bacilli probably find entrance in small numbers and by accident, but there are cases of a more considerable **Insufflation** of tubercular matter. Thus in a case observed by the author a tubercular lymphatic gland adherent to a bronchus burst into the bronchus with the result of an acute tuberculosis in a limited district of the lung. There may also be a somewhat rapid extension from insufflation in tuberculosis of the larynx.

The **Localization at the apex of the lungs** of the earliest lesions is probably related to the fact that the apices of the lungs are the least expansile portions. The first rib even in women is very little raised in inspiration, and in persons with weak respiratory movements the air is apt to stagnate at the apex. This view receives some confirmation from the fact that phthisis so frequently improves when the patients go to reside in high altitudes where the rarefied air requires more vigorous respiratory efforts. In such persons the size of the chest as a whole generally undergoes an increase. Remembering that the tubercular bacillus is of slow growth, we may presume that it is more likely to obtain a footing when it is left undisturbed in parts where the air is more or less stagnant.

3. **Anatomical changes in phthisis.**—In studying the changes in the lung it will be necessary to give descriptions of the various processes separately, and to a certain extent these processes are separable, but at the same time it will be understood that many of them go together and by their simultaneous occurrence frequently mask each other. It may be said in general that the disease, beginning in the finer bronchi, tends to spread in one of two directions, or in both of them at once, namely, along the bronchus to the lung alveoli, or else from the bronchus to the surrounding connective tissue and on into the general connective tissue of the lungs. In both cases we find tubercles developed in all stages of the process, and in both there is inflammation, but of different kinds according to the structures involved. In the one case the inflammation produces newformation of connective tissue and consequent induration. In the other case there is a catarrh of the alveoli, although the alveolar wall may be at the same time involved. It is exceedingly difficult to say what determines the one or the other mode of extension, but it is probable that it depends on individual peculiarities. Some persons are peculiarly liable to catarrh



of the alveoli, and some are not. It is important to observe that the tubercles which occur in both cases partake, to a considerable extent, in the peculiarities of the inflammations. In the case of extension to the connective tissue the tubercles tend to undergo fibrous transformation. In the other case they are liable rather to caseous necrosis.

According to the preponderance of one or other of these groups of lesions we may distinguish two principal forms of phthisis, which may be called the caseous and the fibroid forms respectively.

(a) **The Caseous form.**—When the lungs are examined in an ordinary case of this kind after death they are found adherent to the chest-wall and they are usually seen to be the seats of several cavities, chiefly in the upper lobes. The cavities are of various sizes and very irregular in outline. There are very often one or more large ones divided by partial septa, evidently formed by the coalescence of several smaller ones. The cavities may contain a curdy pus, or they may be comparatively empty, but their internal surface is usually coated more or less with a yellow curdy matter. The tissue immediately around is usually condensed and pigmented.

It is necessary to look away from the cavities in order to observe indications of the **Initial lesion**. One can nearly always distinguish in the midst of the crepitant and comparatively normal tissue isolated

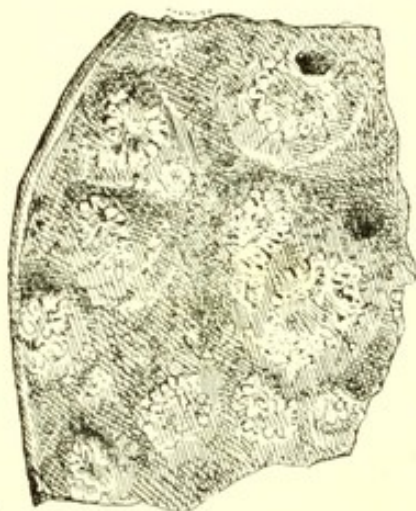


Fig. 256.—Lobular condensations in the caseous form of phthisis. Rounded areas are seen which, being solid, stand out from the general level. From piece of lung in the fresh state.

areas of condensation, such as those illustrated in Fig. 256. They can be felt as solid masses in the soft tissue, and on section they stand out above the general level of the cut surface. They are usually more or less rounded in area and present on section indications of a lobular arrangement, there being a central stem and bodies grouped round it like grapes on a bunch, or more correctly like the carpels of a berry. The central parts are generally whitish or yellow and opaque, and this appearance may involve the whole area, but the peripheral parts have usually a grey translucent character. The coalescence of areas having these

characters gives rise to considerable condensations, in which no such lobular arrangement may be visible, although at the margins there are generally indications of it.

On microscopic examination of the more recent of the initial lesions, appearances will be found which may be illustrated by Fig. 257. The lesion begins and centres in a small bronchus. The bronchus is plugged



with what is at first merely an inflammatory exudation (*b*) consisting of desquamated epithelium and round cells. The outline of the tube is preserved (*a*) but its wall is considerably infiltrated with round cells (*c*).

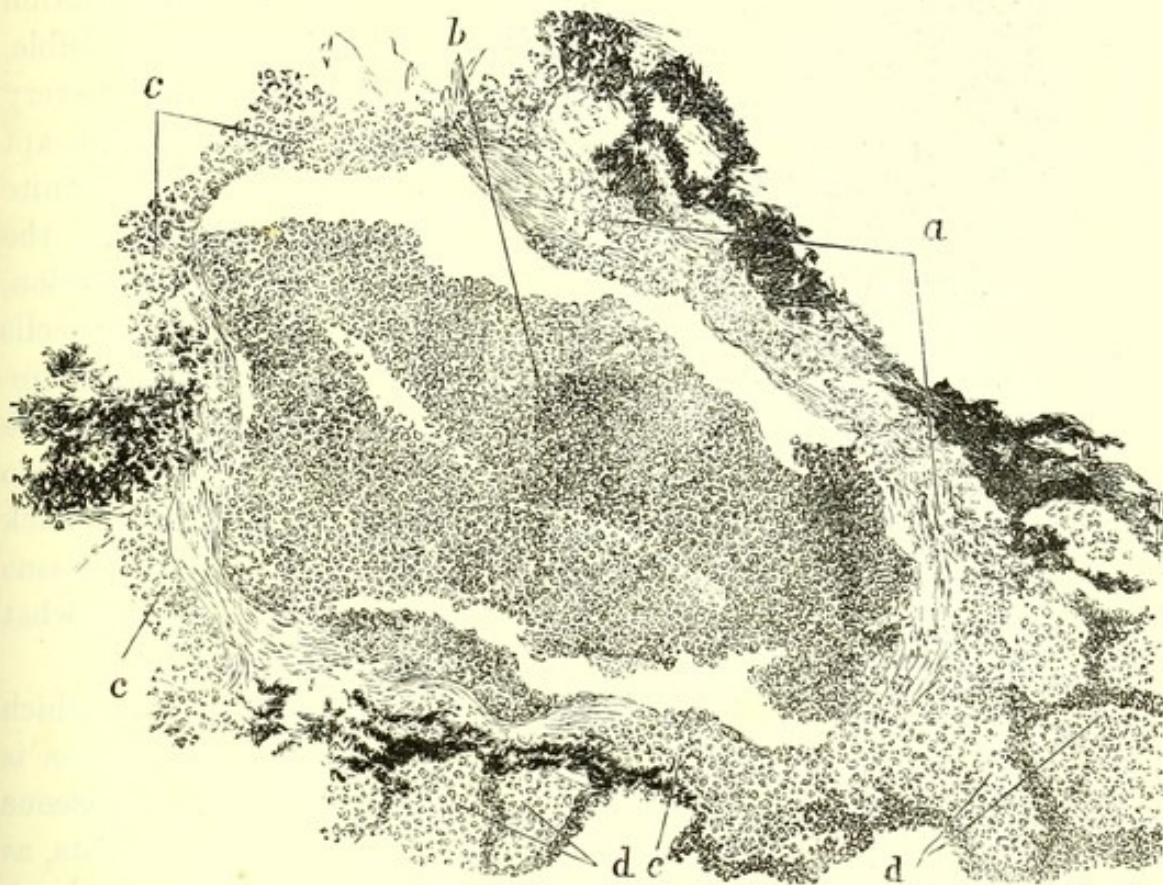


Fig. 257. —Caseous phthisis; recent centre. *a*, wall of bronchus with pigment externally; *b*, plug in bronchus; *c*, round cells infiltrating wall of bronchus; *d*, alveoli filled with blood and catarrhal cells.  $\times 60$ .

The plugged bronchi form the central stems and branching twigs of the areas under consideration, but there is an extension to the lung alveoli and here also we have the effects of inflammation. The inflammation is generally of the parenchymatous or catarrhal character, and the alveoli are occupied by large catarrhal cells (Fig. 258) which are the derivatives of the alveolar epithelium. Sometimes the epithelium itself is seen enlarged and it may be desquamating. Blood is very often present along with the catarrhal cells, sometimes in such abundance as to fill the alveoli. In the specimen of which Fig. 257 is a drawing, for example, the alveoli contained much blood (see further on under Hæmorrhage).

The exudation in the alveoli is sometimes more like that of an acute inflammation, consisting to some extent of round cells, and there are cases in which even fibrine is present.

Besides these inflammatory conditions we have **Tubercles** present in the affected parts. The bronchial wall, as we have seen, is infiltrated with leucocytes which, as they accumulate, obscure the structure of the



wall. They also extend to the connective tissue outside the bronchial

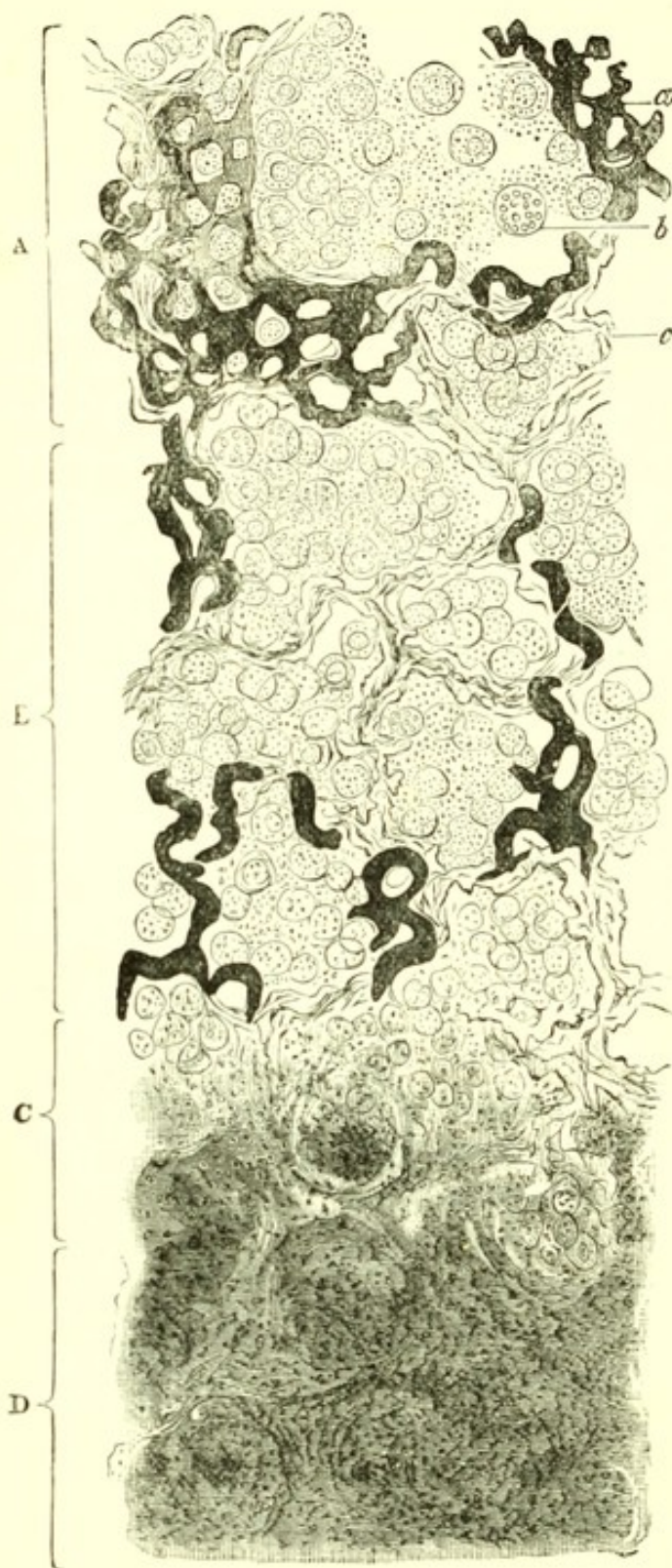


Fig. 258.—Lung in caseous form of phthisis; the capillaries injected (shown black). At *A* there are catarrhal cells (*b*), in alveoli, and the capillaries are well filled. At *B*, the capillaries are less injected. At *C*, caseous necrosis has begun; and at *D*, it is complete, the structure being here quite obscured in a general granular opacity.  $\times 350$ . (HAMILTON.)

tissue outside the bronchial wall, and to the alveolar wall. In the midst of these evidences of inflammation definite tubercles are visible, sometimes typical in every respect. But they are apt to be somewhat indefinite from the existence of the inflammatory infiltration, so that only the giant-cells may be definitely distinguishable. The giant-cells not infrequently take into their substance the black pigment of the lung tissue and so may be somewhat strikingly manifest.

Another feature which distinguishes the process is the occurrence of **Caseous necrosis**. This consists, as we have seen, of the death of the structures accompanied by the production of finely granular fat. As this change reduces everything which it affects to a homogeneous granular condition, it greatly obscures the structure and renders the identification of the individual elements very difficult. It occurs in all the structures already described as affected by the inflammation, the plug which fills the bronchus, the bronchial wall, the contents of the alveolus,

the alveolar wall, and the tubercles. If the process is not very advanced then it may be seen that the caseous necrosis begins in



the bronchus, as in Fig. 259, the outline of the tube being still visible. In this illustration the change has extended to the alveoli immediately around the bronchus and their outlines are still obscurely visible in the

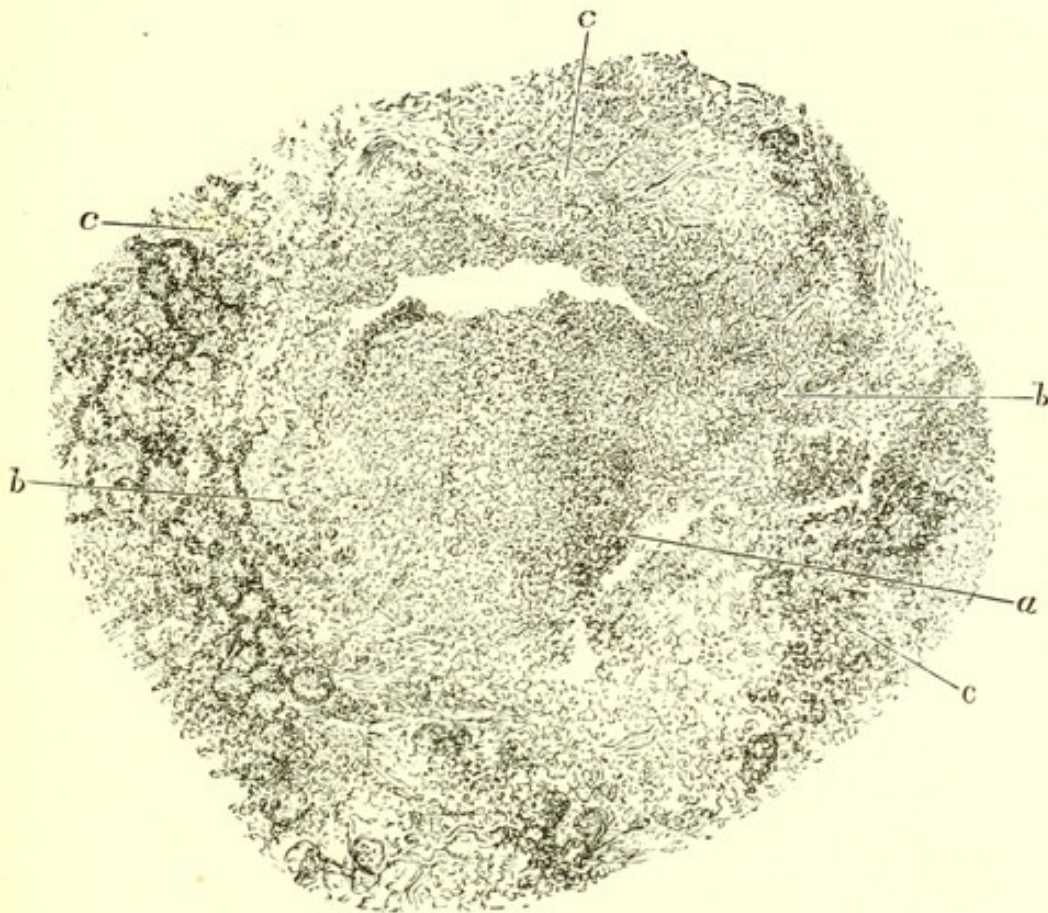


Fig. 259.—Caseous phthisis; further stage. *a*, caseous centre, including plug and bronchial wall, rendered indistinguishable by the caseation; *b*, alveoli scarcely distinguishable; *c*, alveoli further out, filled with exudation and having walls infiltrated.  $\times 35$ .

midst of the general granular appearance. A similar obscuration of the alveoli is seen in the lower part of Fig. 258. Outside the caseous area the alveoli are filled with catarrhal cells and their walls infiltrated with round cells. The caseous matter is visible to the naked eye as a yellow or white opaque substance which is somewhat brittle, and, like cheese, consists of nitrogenous matter containing finely divided fat.

The dead caseous matter may lie for a long time and do little harm, just as any inert dead animal matter may. It may even undergo a partial absorption, or with this may be combined an infiltration with lime salts, so that ultimately a cretaceous mass remains embedded in the lung. These processes, however, imply that the tuberculosis has ceased to be active and that the caseous matter is inert. It is more usual to have a disintegration of the caseous matter.

**Softening** or breaking down of the caseous matter often shows itself to the naked eye in the central parts of the caseating areas. It can



often be determined under the microscope that the softening is beginning in the situation of the bronchus, which thus again proclaims itself the centre of the process. In Fig. 259 there are indications of a crumbling of the caseous matter and a partial separation so as to leave cracks or fissures.

In this way **Cavities** are formed, and each cavity implies the death and destruction of a certain portion of lung tissue, usually involving a bronchus and surrounding alveoli. The broken-down caseous material forms a grumous turbid fluid, in which the more resisting elastic tissue of the lung may be found still retaining to some extent the form of the alveoli. This elastic tissue may be frequently recognised in the sputum of such patients by proper methods of search (see Fig. 260).

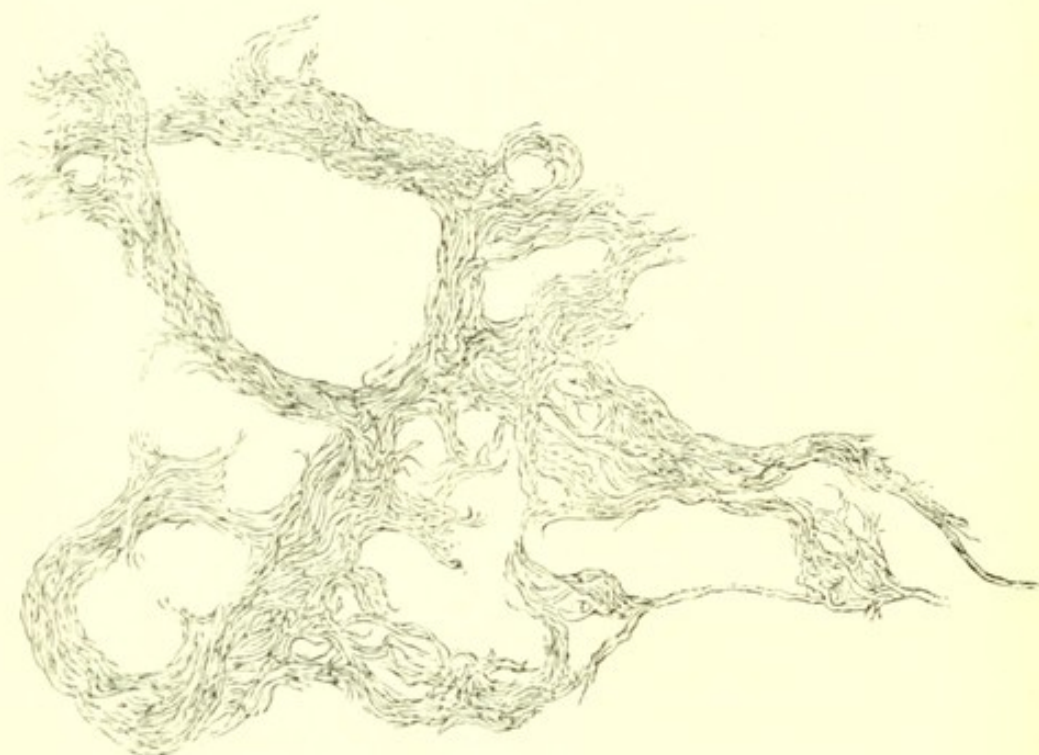


Fig. 260.—Lung tissue from the sputum in phthisis. The sputum was digested in caustic soda according to Fenwick's method, and then subjected to microscopic examination.  $\times 350$ .

At first the cavity formed is small, but by extension of the process and coalescence of neighbouring softenings, larger cavities result. These cavities are very irregular, and their walls at first ragged and ill-defined. The cavity sooner or later opens into one or more bronchi, and its contents are discharged. When the caseous material is cleared out the wall assumes an inflammatory character and secretes pus more or less abundantly. There may be ultimately a smoothing of the walls of the cavities by the formation of connective tissue, but there is not the regular lining membrane of the bronchiectatic cavity, nor the same relation to the bronchi. In the walls of such cavities, and in the



mucous membrane of the bronchi with which they are in communication, tubercular ulcers may sometimes be distinguished.

It may be that, as Hamilton suggests, the softening of the caseous matter is by a process akin to that which occurs in the ripening of cheese, in which, according to Duclaux, certain insoluble albuminates become soluble in water.

The whole process is sometimes a more acute one than that indicated above. The original exudation may approach in its characters to pus, and the process of softening may present little beyond a rapid necrosis akin to ordinary necrosis or sloughing. In these cases there is sometimes a **Purulent peribronchitis**. Suppuration occurs in the bronchial wall and in the surrounding tissue, and, these structures being broken up, the cavity partakes of the nature of an acute abscess. Some of these abscesses may be near the surface and by rapidly undermining the pleura cause it to slough. By the separation of the sloughs the cavity of the abscess may come to communicate with the pleural cavity and so pneumothorax may result. Sometimes there is even a gangrenous condition developed, and actual sloughing of pieces of lung tissue occurs.

Acute cases such as these bespeak a peculiar virulence of the morbid agent or a peculiar susceptibility of the patient. They usually pass rapidly on to a fatal issue with high fever. In some cases recovery takes place, the pus dries-in or is discharged, and the abscesses become surrounded by indurated connective tissue and contract.

(b) **The Fibroid form.**—On post-mortem examination in typical cases of this form the lung is found very firmly adherent over the affected part, which nearly always includes the apex. One often has to remove with difficulty a dense leathery cap which covers the apex of the lung (see Fig. 261). On cutting into the lung there are usually cavities, but they are not generally large and the internal surface is mostly clean and moderately smooth. A distinct membrane lines the cavity (*c* in figure). Outside the cavity the tissue is of a deep slaty colour in which opaque white spots may be occasionally visible, and it is very dense. The dense pigmented tissue may involve a considerable portion of the lung and the affected part is shrunken and contracted (see figure).

Looking away from the condensed part in which the lesion is advanced, we find, as in the caseous form, the advanced posts of the disease in the form of isolated condensations in the midst of the crepitating normal tissue. Here the **Initial lesion** is different from that in the other form. It consists in hard dark or nearly black solid bodies, scattered through the lung tissue. On running the finger over the cut surface one feels these bodies and they stand out above the general surface.



In this form as in the other a plugged bronchus will be found to form the centre of the initial lesion. This is shown in Fig. 262, which is from

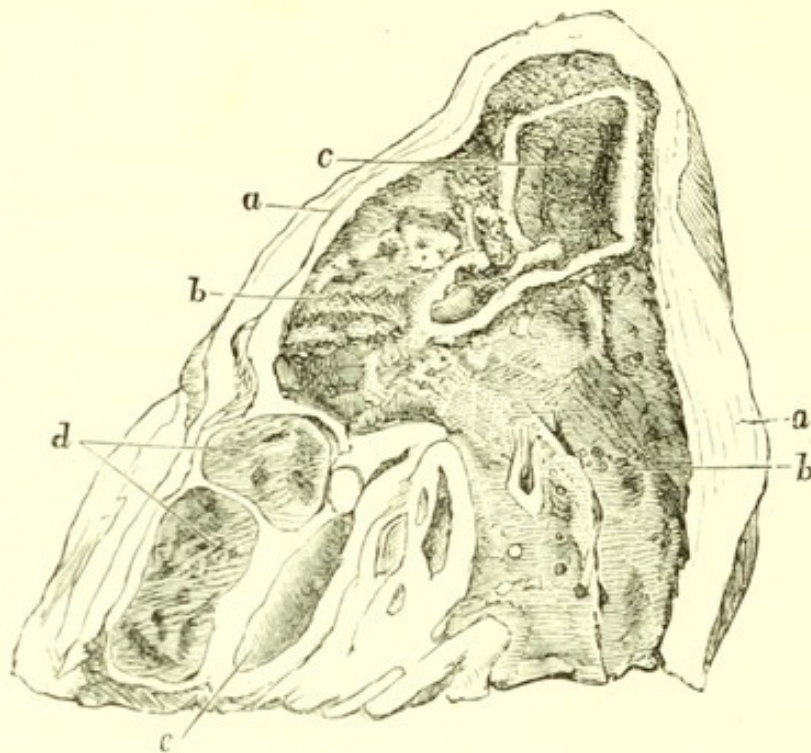


Fig. 261.—Upper lobe of lung in fibroid phthisis. *a, a*, greatly thickened pleura; *b*, condensed and pigmented tissue; *c*, cavity with distinct lining; *d*, bronchial glands enlarged and pigmented; *e*, main bronchus. The nearness of the bronchus and glands to the apex indicates the shrinking. Near natural size.

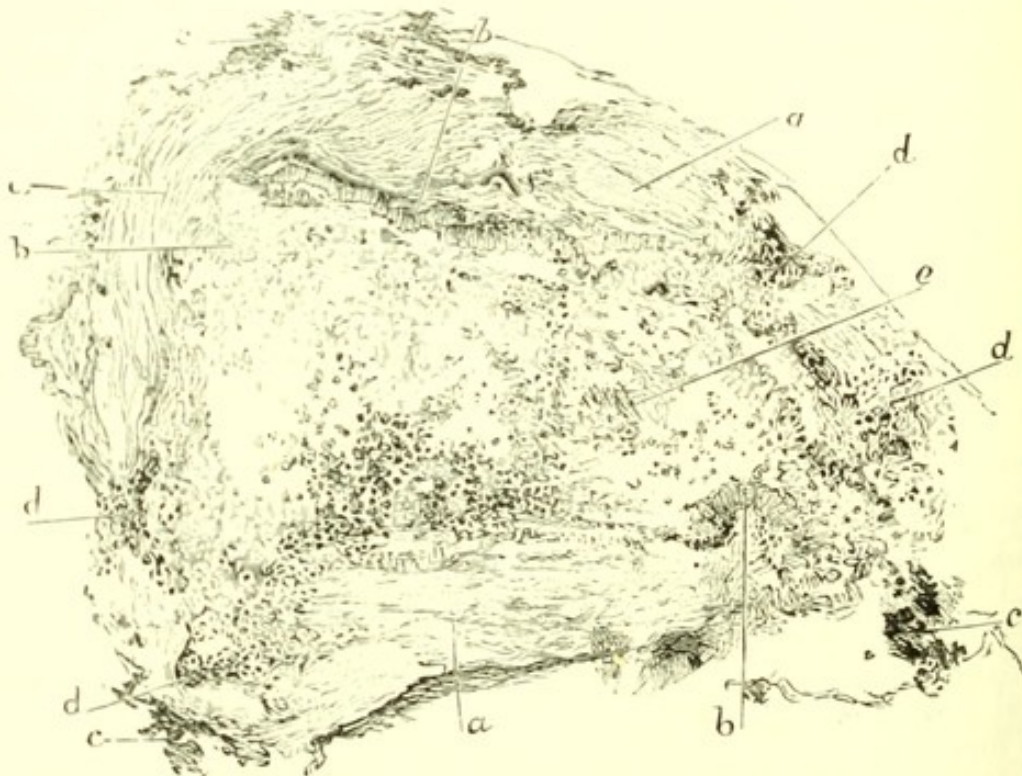


Fig. 262.—Fibroid phthisis, early lesion. *a, a*, bronchial wall; *b, b*, projecting parts still covered with epithelium; *d, d*, round cells infiltrating bronchus; *e*, exudation in calib.  $\times 60$ .

the centre of such a lesion. The bronchus contains, as before, inflam-



matory products, namely, round cells and desquamated epithelium. The wall of the bronchus is also infiltrated with round-cells, and tubercles are present in the wall and in the surrounding connective tissue. There is, however, very little appearance of inflammation in the lung alveoli, and the affection seems to advance rather by the lymphatics into the connective tissue than along the mucous surface to the alveoli.

The distribution of the **Tubercles** is indicative of the advance in the directions just mentioned. They are often grouped in the neighbourhood of bronchi, as in Fig. 263, but are also present at some distance

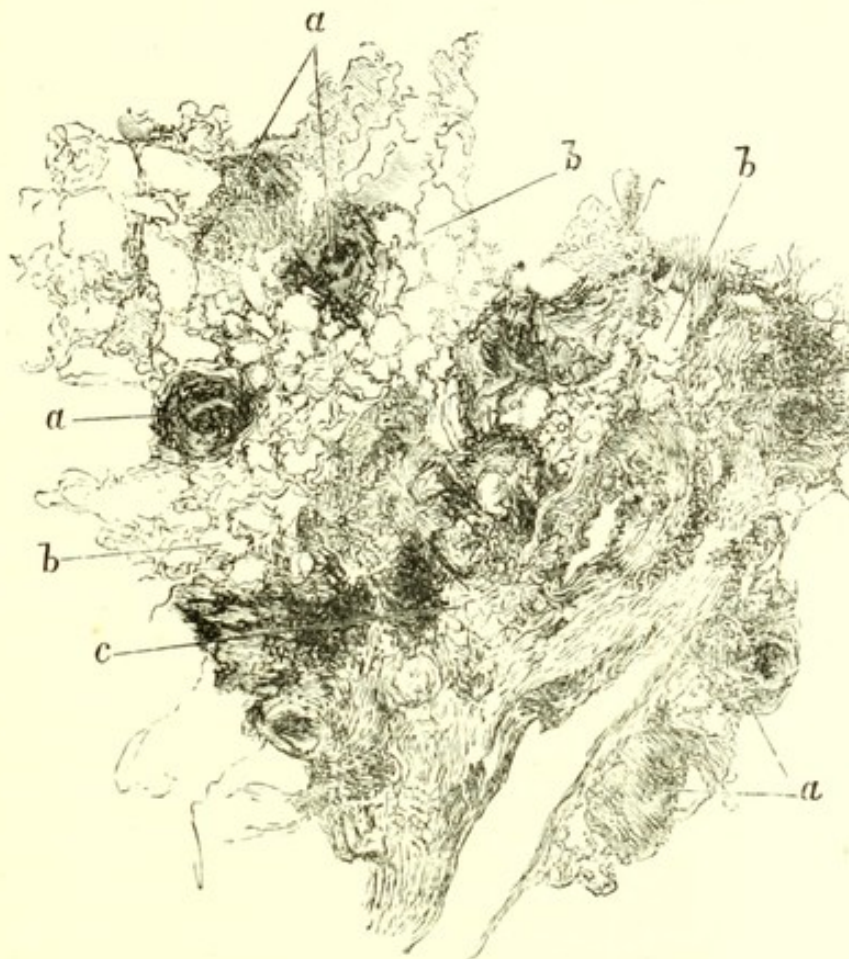


Fig. 263.—Fibroid phthisis. *a, a*, tubercles in connective tissue, some close to a bronchus, others removed; the darker centres indicate caseation. *b, b*, emphysematous lung tissue.  $\times 20$ .

(*a* to left in figure), the process travelling by the lymphatics. The tubercles present in the earlier periods the typical structure, and there are frequently several giant-cells in the midst of them.

Hæmorrhage is very common here as in the other form, and it may be sufficient to fill out the alveoli.

**Caseous necrosis** is much less a feature here than in the other form, but it is usually present in the plug which fills the bronchus, as well as to some extent in the bronchial wall. This is shown in Fig. 264, in which the central part (*a*) representing the plug is homogeneously



granular, while the walls of the tube are infiltrated with round cells. In some places (as at *c*) there is an aggregation which with a higher power is seen to be a tubercle.

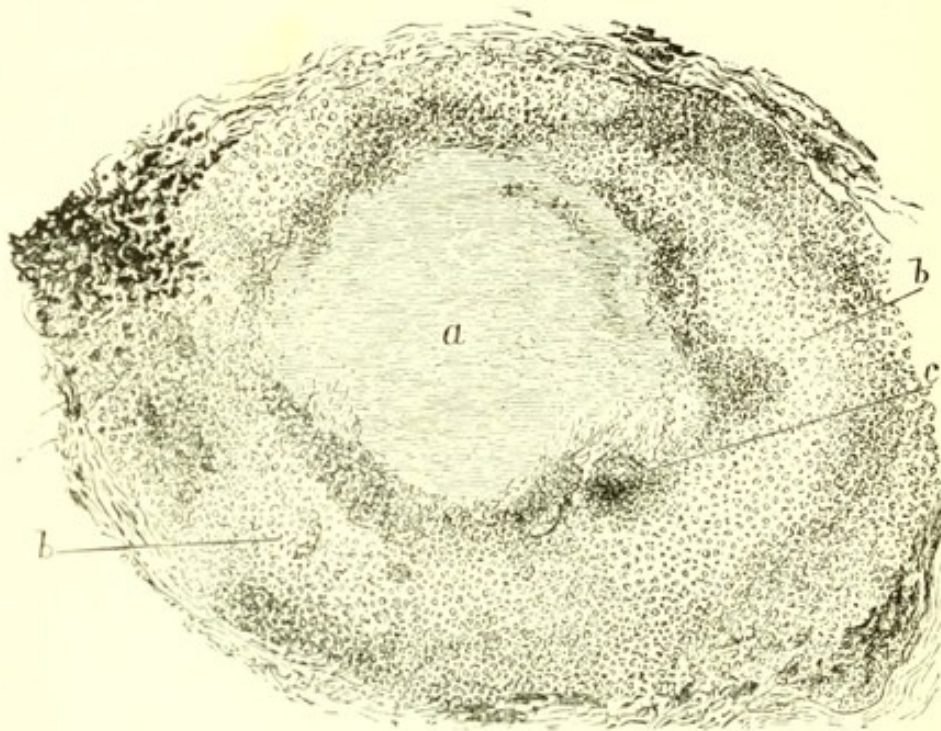


Fig. 264.—Fibroid phthisis. *a*, bronchus plugged and caseous; *b*, *b*, wall of bronchus infiltrated with round cells and tubercles; *c*, tubercle in wall.

A feature present in this form, but not in the other, is **Fibroid transformation**, which affects both tubercles and connective tissue and gives its peculiar characters to this form of phthisis. The tubercles are converted into clear structureless bodies in which all the elements of the tissues are lost except, occasionally, one or two giant-cells. These may be partly transformed, but may still be recognizable, especially as they frequently contain a considerable number of black granules.

A similar fibroid transformation affects the connective tissue around the bronchus and extends to the general stroma of the lung. There is thus great induration of the lung tissue and the fibrous tissue shrinks and produces great deformity. The shrinking of the tissue, associated as it is with adhesion of the pleura, frequently leads to drawing in of the chest wall which is often a characteristic feature, and it leads also to two lesions which are often very apparent in the lung, namely, bronchiectasis and emphysema, both of which are complementary.

**Bronchiectasis** is, in this form, the most active factor in the formation of cavities. It is partly a purely complementary process, the shrinking being compensated by the dilatation of the air-spaces. Hamilton has pointed out another mode of formation. As the chest wall forms a comparatively fixed point to which the shrinking tissue is attached by the



pleural adhesions, and as this tissue is also attached to the walls of the bronchi, the result of the shrinking will be that these two points will be approximated, the chest wall drawn in and the bronchial wall drawn out. There is a third way in which bronchiectasis occurs. The secretions may accumulate in a bronchus behind an occlusion of the tube. Such an occlusion will occur when the primary lesion has affected a bronchus of larger calibre than usual, or where the shrinking tissue has constricted a bronchus.

The **Bronchiectatic cavity** will be lined with a distinct membrane, and will usually be directly continuous with a bronchus (see Fig. 261). It may exist in the midst of crepitating lung tissue, the complementary dilatation, being in a part not affected by the tuberculosis.

**Emphysema** is a frequent accompaniment of the process. Whenever a piece of lung escapes the fibroid change, it is liable to emphysema on account of the shrinking in its neighbourhood. Hence in the midst of the shrunken tissue one often sees islands of lung tissue having a honey-combed appearance (see Figs. 263 *b* and 265).

Another occasional result of the shrinking is a formation of **Cyst-like cavities** in the pleura, as shown in Fig. 265. The shrinking of the lung, dragging the adherent pleura with it, may cause spaces to form in the pleura, or even in the interlobular septa (as at *c*), these spaces being filled with serous fluid.

**Pigmentation** is a peculiarly prominent feature in fibroid phthisis. Even the initial lesion is characterized by the almost black colour of the nodules, and the indurated tissue has a slaty or blackish colour (*slaty induration*). Perhaps the explanation of this is that the carbonaceous pigment (see further on) is retained by the affected bronchi and not swept outwards by the cilia of the epithelium.

The caseous and fibroid forms of phthisis are in general distinguishable. They have certain points in common, chiefly in respect that each begins with a bronchitis of the finer tubes, and that each is characterized by the presence of tubercles. In the one form, however, the bronchial inflammation extends to the proper parenchyma of the lung, constituting a **Lobular broncho-pneumonia**, whereas in the other it is more localized around the inflamed bronchus, constituting a tubercular **Bronchitis and Peri-**

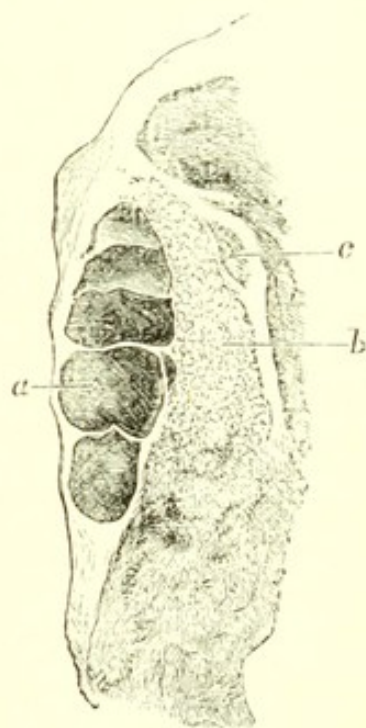


Fig. 265.—Fibroid phthisis. *a*, cysts in pleura from shrinking of lung; *b*, emphysema; *c*, cyst in interlobular connective tissue. Natural size.



**bronchitis.** There is the further distinction that in the one form caseous necrosis is characteristic, whereas in the other, while probably present in most cases, it is limited in extent and may be confined to the contents of the bronchi and the bronchial wall.

The difference is probably due to differences in the individual proclivities of the persons affected. In the fibroid form the disease is more chronic, the persons affected are as a general rule older, and it occurs more frequently in the male sex. All these facts point to the conclusion that in this form there is greater resistance on the part of the tissues to the morbid poison. This is confirmed by the resemblance which the process, in some respects, presents to that which is concerned in the healing of phthisis. It may be said that in the caseous form the tissues are directly killed by the progress of the disease, sometimes with great rapidity, whereas in the fibroid form there is a long struggle and very little palpable softening or destruction.

This being the case it may be inferred that the two forms are not absolutely distinguishable. They run into each other, and the caseous form may assume many of the characters of the fibroid, especially when it becomes very chronic or partial recovery takes place.

The **Sputum** in phthisis pulmonalis is variously composed. In the earlier stages the expectoration has the usual characters of that in catarrh, consisting of mucus, with more or less abundant leucocytes. In the sputum in this early stage are often found large epithelioid cells with one or more nuclei, such as we find in the lung alveoli in the catarrhal form of phthisis. These cells frequently present fatty degeneration. Considerable weight has sometimes been attached to the presence of such cells as evidence of the existence of the disease in its early stage, and their discovery may be useful in this respect; but as they may exist in simple catarrhs their diagnostic value may be overrated. The sputum in phthisis often contains **Elastic tissue** from the breaking down of the lung. In very rapid cases we may find this by a simple examination of the sputum, but the search is often a difficult one, because the thick mucus and pus hold the pieces of lung tissue suspended and isolated. By Fenwick's method of digestion in soda solution, pieces of lung tissue, such as that shown in Fig. 260, will frequently be found in the deposit. This method is also applicable to the sputum in gangrene of the lungs. The **Tubercular bacillus** is usually to be found in the sputum, and is sometimes of great diagnostic value. The appearances are shown in Fig. 52, p. 198, and have been described already.

4. **Extension of the tuberculosis in phthisis pulmonalis.**—In both of the forms already described it has been shown that the tuberculosis, beginning in the finer bronchi, extends, on the one hand, to the lung alveoli, and on the other, to the connective tissue. Besides that, however, there are further extensions both in the lung and beyond it.

An existing cavity, especially one arising from disintegration of caseous matter, is a great source of infection, its contents being charged with tubercular bacilli. The infective matter will be carried from the cavities



and will be partly insufflated into other parts of the lungs and partly discharged. It thus causes an extension of the disease in the lung itself, and is liable to infect the air passages as it is carried along.

Hence, **Tuberculosis of the bronchi** is very frequent in connection with cavities, the mucous membrane becoming the seat of tubercular ulcers. If the bronchial tubes be opened up, the ulcers are visible as more or less rounded erosions, sometimes with distinct white tubercles at their borders.

A further extension to **Larynx and Trachea** is very common, and from these, by way of œsophagus and stomach, to the **Intestine**.

There is also an extension by the lymphatics, so that in nearly all cases of phthisis the **Bronchial lymphatic glands** are affected. The condition here is similar in character to that in the lungs. In the caseous form the glands are caseous, resembling closely the appearances in ordinary scrofulous glands. In the fibroid form they are liable to be more or less fibrous and deeply pigmented. Tubercles are present in the glands, sometimes in the most typical form. In some cases the bronchial glands soften, but in others the caseous matter becomes impregnated with lime salts, so that it is not uncommon to meet with hard cretaceous matter at the roots of the lungs.

5. **Healing of phthisis.**—It is to be remembered that tuberculosis is due to an infective material, which usually goes on reproducing itself. In the healing of tubercular lesions generally there are two methods which may be followed, and in phthisis pulmonalis we have examples of each of these. On the one hand, the infective matter may be cleared out and the parts around become the seat of simple inflammatory processes, or on the other hand, the caseous matter may have its infective character overcome and be left as a piece of innocuous dead matter in the tissues. In either case, so far as the lung is concerned, there is implied an increase in the vigour of the parts, so that the infective character of the matter may be annulled.

When a cavity forms by the softening of the caseous matter, there is generally a further extension as shown above, but in some cases the body has attained such vigour that the further extension is hindered. In that case the wall of the cavity comes to be composed of healthy granulation tissue, which develops into connective tissue as in the ordinary cicatrix. If the cavity is so situated as that contraction can occur, then there may be a shrinking till it is completely obliterated, and a **Cicatrix** takes its place. On the other hand, circumstances may allow of only partial contraction, and a cavity or cyst may remain in the midst of the cicatrix.

In other cases the caseous matter fails to soften, the disease is checked



before cavities are formed, and in that case we have necrosed structures lying in the lung. In this case granulation tissue is formed around the dead matter, and may partly eat into it. By development into connective tissue a capsule is formed around the dead matter, and the latter will by and by become impregnated with lime salts, so that ultimately particles of lime or considerable **pieces of cretaceous matter** will be present in the midst of a cicatrix.

It is worthy of note that the connective tissue formed in both these forms of healing is deeply pigmented, so that in this respect the processes are comparable to those concerned in fibroid phthisis.

Healing by one or other of these processes is by no means uncommon. In the course of post-mortem examinations one very frequently meets with pigmented cicatrices, often with chalky particles in their midst, at the apices of the lungs, the pleura being adherent over the affected parts.

If, after healing, a return of the disease takes place, the chalky matter may afterwards be separated, and such pieces have been known to be spit up.

**6. Hæmorrhage in phthisis.**—Hæmoptysis is one of the most common manifestations in phthisis. It is necessary to distinguish between an early and a late hæmorrhage.

(a) **Early hæmorrhage.**—In examining the initial lesion in phthisis, whether in the caseous or fibroid form, it is common to meet with alveoli filled with blood, presenting an appearance very similar to that shown in Fig. 251, which is from a hæmorrhagic infarction of the lung. The blood may be so abundant and occupy so many alveoli, as to give quite the character, in some cases, of the hæmorrhagic infarction.

The blood here comes from the pulmonary capillaries, escaping by diapedesis; it does not arise by insufflation, else it would be more mixed with air and broken up. The homogeneous complete filling of the alveoli implies a regular leakage which gradually expels the air. The blood also is often in alveoli which are little altered otherwise, and it is present in a considerable group of alveoli together.

These facts would indicate a local interference with the circulation as a cause of the hæmorrhage. Such an interference is liable to occur from the proximity of the branches of the pulmonary artery to the bronchi. The arteries and bronchi run to a large extent parallel, their sheaths continuous, and the inflammatory changes in and around the bronchi which form such an important part of the initial lesion may well exercise pressure on the arteries. This view is confirmed by the fact that in general tuberculosis of the lung a similar hæmorrhage is very often present, sometimes to a very aggravated extent. In this affection the tuberculosis is usually intimately related to the arteries, so that there will be a direct interference with them.



In any case the hæmorrhage is part of the initial lesion, and it is known that the appearance of blood in the sputum is often a very early sign of phthisis.

As the hæmorrhage is associated with the earliest lesions it may occur at a time when no symptoms of disease in the lungs are present, and may be the apparent starting point, although not really so. By some the occurrence of hæmorrhage has been regarded as sometimes the origin of phthisis, a view which has been expressed by the term *Phthisis ab hæmoptoe*. This view is without adequate foundation, and the hæmoptysis is probably, in every case, evidence of the existence of the initial lesion. It is not improbable, however, that the occurrence of hæmorrhage may accelerate the progress of the disease, as the bacilli may possibly find in the blood a suitable nidus, and so exhibit a more rapid growth.

(b) **Late hæmorrhage.**—The hæmorrhage just described may occur at intervals throughout the course of the disease, but in advanced cases a much more considerable and not infrequently fatal hæmorrhage is liable to occur. In this case the bleeding arises by **Rupture of branches of the pulmonary artery** which have been partially exposed in the walls of cavities.

As a general rule the arteries in the walls of cavities are obliterated, more especially in the caseous form, but where obliteration has not completely occurred, the wall of the artery being unsupported and perhaps softened by inflammatory infiltration is liable to give way. Before the actual rupture the vessel wall usually bulges out, so as to form an **Aneurysm** (see Fig. 266). In this figure the cavity is a bronchiectatic one, and its wall was comparatively unaltered, the aneurysm having arisen purely from want of support. In the caseous form there is softening of the wall of the artery, and the aneurysm has not such a definite sac as in this case.

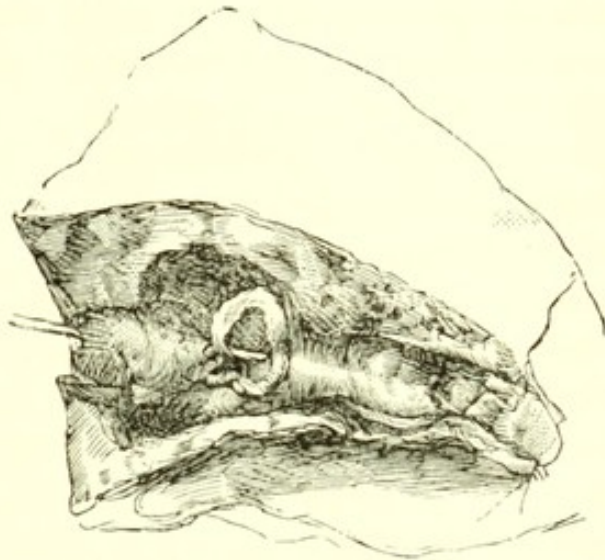


Fig. 266.—Aneurysm in a bronchiectatic cavity. A probe has been introduced into the artery, and is visible through the gaping aperture in the aneurysm. It is also indicated close to the wall of the bronchus beyond the cavity.

In cases of this kind the hæmorrhage is often very great, but even in the case of a considerable tear it may be stilled by the blood coagulating in the cavity and forming a kind of cap over the aperture.

**7. Affections of the pleura in phthisis.**—From the intimate connection of the pleura with the lung it may be expected that it will frequently be affected in phthisis.

The lymphatic system of the lung does not apparently communicate



directly with the cavity of the pleura, so that although a tuberculosis may extend to the subpleural tissue it does not directly affect the pleural sac.

The relations of the pulmonary lymphatics to the pleura are shown by the locality of the carbonaceous pigment in the lungs. This carbonaceous pigment is carried about and deposited wherever there are communicating lymphatics, and it is often abundant in the subpleural tissue; but it never penetrates into the sac itself. On the other hand, there seems to be a communication in the opposite direction from the pleura to the interlobular connective tissue. This is shown by the fact that in tubercular pleurisy, there is a certain penetration from the pleura into the lung. (See under Tubercular Pleurisy.)

While tuberculosis does not extend to the pleura, because it implies the passage of solid bodies (the bacilli), there are very commonly simple inflammatory processes, which depend on the extension of the dissolved products. Such products being present in the connective tissue of the lung may readily soak into the pleura.

**Chronic pleurisy** is a constant occurrence, giving rise, as in other cases of chronic inflammation, to newformation of connective tissue. The pleura over a tubercular lung is nearly always thickened, sometimes greatly so (as in Fig. 261), and the two layers are almost constantly adherent. The two layers are not only adherent, they have really coalesced, and their blood-vessels intercommunicate, so that if the vessels on either side be obstructed the pleura may be still nourished from the other. The thickening and adhesion are conservative processes, shutting off the diseased lung from the general pleural sac. It is when these conditions fail to occur that we commonly have the more serious pleural lesions, acute pleurisy and pneumothorax.

**Acute pleurisy** is a frequent concurrent in cases of phthisis, especially in the caseous form. It implies that the lesion in the lung has come to the surface at a place where adhesion of the pleura has not yet taken place. This will be most frequent in early periods and in acute cases.

The acute pleurisy is often connected with **Necrosis** of the pleura. The pleura is nourished by the pulmonary vessels, and as these are occluded and necrosed when involved in the caseating lesion, the pleura will be involved in the necrosis in so far as it is related to the occluded vessels. At the very outset of a caseous phthisis one of the affected areas may be immediately beneath the pleura, and we may have a necrosis before there has been time for the formation of adhesions. In this way we may explain many of the cases in which pleurisy has apparently preceded the pulmonary disease. In all acute cases we are liable to have necrosis of the pleura, and it is not uncommon to find quite a number of dead white areas visible on the surface, each indicat



ing an area of necrosis, generally concealed to some extent by a layer of fibrine, the result of the inflammation.

The mere exposure of a necrosed piece of pleura seems to induce an acute pleurisy, perhaps by allowing the penetration of irritating juices such as the living structures intercept. The acute pleurisy is of the usual kind (see further on) accompanied by fibrinous exudation, but it is usually limited by existing adhesions. Through time the acute inflammation will subside and a chronic pleurisy with adhesion result.

**Pneumothorax** also implies necrosis of the pleura, but there is, in addition, a partial separation of the dead piece and the establishment of

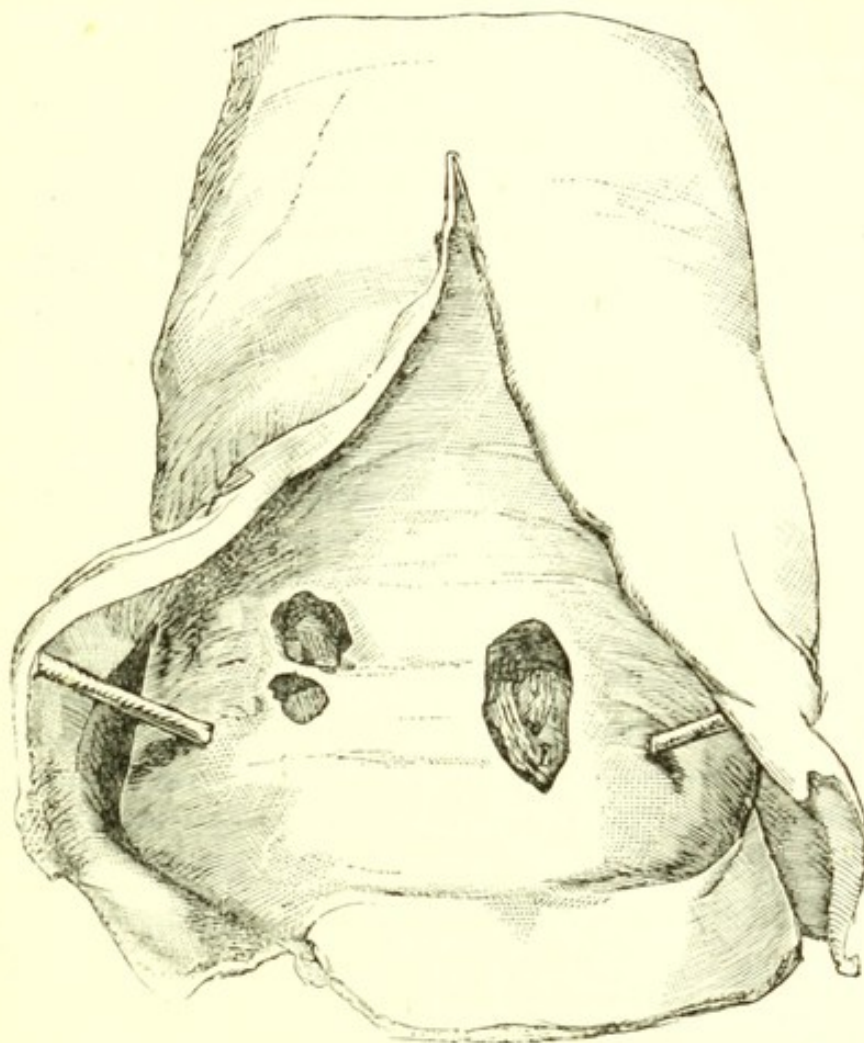


Fig. 267.—Lower part of lung and pleura from pneumothorax. There are three apertures caused by necrosis of pulmonary pleura.

a communication between the pleural sac and the air passages. This infers the existence of a cavity beneath the necrosed piece of pleura and not merely a caseous area. The necrosed piece begins to separate at one edge, and then the pneumothorax suddenly occurs. As death often occurs in consequence of a pneumothorax, one not infrequently has the opportunity of observing the condition in this stage. If the patient survive, then the whole piece is detached and apertures are left in the pleura such as those shown in Fig. 267.



Along with pneumothorax there is an acute pleurisy. If it has not occurred in consequence of the necrosis, then it will ensue when, by the separation, some of the contents of the pulmonary cavity pass into the pleural sac. These two conditions, acute pleurisy and pneumothorax, are thus closely related, both of them being connected with necrosis of pleura. They also occur in relation with the advanced outposts of the pulmonary lesion, and are hence liable to develop in connection with the less affected lung or portion of lung, and by causing serious damage to the lung on which the patient chiefly depends for respiration, they often induce very serious and even fatal dyspnœa.

**8. General effects of phthisis.**—In its general influence on the body phthisis pulmonalis is of great importance.

Its most constant effects are **Emaciation and Anæmia**, the former of which, as we have seen, was the main element in suggesting the name phthisis. These conditions have their source chiefly in fever.

**Fever** is an almost constant accompaniment of phthisis, altogether apart from any general tuberculosis. It is to be referred to the presence in the blood of the various products evolved in the course of the local processes. These consist in part of the products of the specific microbes, which will pass into the blood, even when the microbes themselves do not. There will also be the products of the disintegration of blood, and of inflammatory exudations, such as those in pleurisy. In pulmonary cavities again, various forms of decomposition, putrid and other, are liable to occur, and the products naturally lead to fever.

**Amyloid disease** occurs in about a fourth of the cases of phthisis, and is more frequent proportionally in the fibroid than in the caseous form. It usually affects the spleen, kidneys, liver, intestine, and lymphatic glands, but there are great differences in its distribution and extent. It may be absent in any one of the situations mentioned while present in the others, or it may be greatly exaggerated in one while very inconsiderable in others. (See the author's "Lectures to Practitioners.") It is typically present in the spleen in the majority of cases, and takes the form of the sago spleen.

**Disease of the kidneys** is a frequent concomitant of phthisis. It may be amyloid disease, complicated (as amyloid disease regularly is) with chronic nephritis, or it may be a tuberculosis. On the other hand there is not uncommonly a nephritis, sometimes in an acute or subacute form so as to produce enlargement with fatty epithelium, sometimes in a more chronic form, with contraction and granulation of the kidneys.

In cases of amyloid disease the mere presence of the amyloid material may induce chronic inflammation, but there is a considerable number of cases in which nephritis is present in an acute form. In these



cases we may suppose that the products from the disintegrating processes going on in the lungs lead, by their discharge through the kidneys, to an inflammation of these organs (see author, l. c.).

**General tuberculosis** not infrequently follows phthisis pulmonalis. We have seen that the disease in the lungs is a local tuberculosis, and as a general rule it extends along mucous surfaces and lymphatic channels. In a considerable proportion of cases, however, the tubercular bacilli to some extent reach the blood and are carried throughout the body. If the liver be examined microscopically there will often be found a few tubercles, generally in various stages of degeneration. There are also not infrequently a few tubercles in the kidneys visible to the naked eye. We sometimes meet even with a more considerable tuberculosis in the kidney and elsewhere, constituting a **Chronic general tuberculosis** (see p. 206), especially in children.

**Acute general tuberculosis** is a less frequent result and is generally due to a definite extension of the local tuberculosis to a branch of the pulmonary vein which has not become occluded by the advancing lesion (see p. 207). It may also arise by an extension of the tuberculosis from the bronchial glands to the thoracic duct.

**Literature.**—The literature of phthisis is very extensive. The older literature will be found in WALDENBURG, Tuberculose, Lungenschwindsucht u. Scrofulose, 1869. See further, CARSWELL, Illustrations of path. anat., 1838; VIRCHOW, Lehrb. d. spec. Path. i.; NIEMEYER, Klin. Vortr., 1867; BUHL, Lungenentz. Tuberc. u. Schwindsucht, 1872; RÜHLE and RINDFLEISCH, Ziemssen's Handb., v.; WEIGERT, Virch. Arch., lxxvii., lxxxviii., civ.; KOCH, Etiology of tuberculosis, Syd. Soc. transl., 1886; BAUMGARTEN, Tuberkel und Tuberculose, 1885; WEICHELBAUM, (Inhalation) Wien. Med. Wochenschr., 1883; TAPPEINER, Virch. Arch. lxxiv. and lxxxii.; ZIEGLER, Volkmann's Samml. No. 151, and Lehrb; HAMILTON, Path. of bronchitis, etc., 1883; EWART, (Position of cavities) Gulstonian lectures, 1882, in Brit. Med. Jour.; THOMPSON, Family phthisis, 1884; JACCOUD, Curability of pulm. phthisis, 1885; POWELL, Dis. of lungs and pleura, 1886; WILLIAMS, Pulmonary consumption, 1887; KIDD, (Distrib. of bacilli in the lung) Med. chir. trans., lxxviii., 1885; YEO, (Contagiousness) Brit. Med. Jour. 1885, i.; COATS, Lect. to practitioners, 1888; GERMAIN SÉE, Bacillary phthisis; JAMES, Pulmonary phthisis, 1888; DOBELL, Bacillary consumption, 1889.

#### IX.—DISEASES FROM INHALATION OF DUST. PNEUMOCONIOSIS.

**Carbonaceous pigment in the lungs.**—The lungs of all adults have more or less of a grey colour from the existence of a black pigment in the lung tissue. \* This pigment is absent from the lungs of children, and is undoubtedly the dust inhaled with the respired air. The air of all confined spaces, such as rooms, is loaded with finely divided particles, particularly in cities where coal is burnt extensively, and this attains its maximum in the black fogs of such cities as London and



Glasgow. The particles of dust inhaled with the air are for the most part caught by the mucus with which the surface of the bronchial tubes is moistened, and as the ciliated epithelium plays in the direction towards the larynx, the dust-laden mucus is carried upwards to the larynx, where it is either expectorated or swallowed. No doubt when the air is unusually laden with dust the mucous secretion is increased, and those who live in cities know that when the weather is thick a considerable mass of black mucus is brought up from the larynx in the morning, the busy cilia having swept it thither during the hours of sleep.

But some of the dust penetrates beyond the reach of those scavengers and passes into the lung alveoli, where it lodges. From the lung alveoli it penetrates into the lung tissue. It is to be remembered that the structure of the alveolus is somewhat like that of a serous membrane. There is a single layer of epithelium, and some have even described stomata or pseudo-stomata as existing. At any rate the dust particles penetrate through or between the epithelial cells and emerge into the lymph spaces of the alveolar wall. Having entered the lymphatic system of the lung, the dust is carried into all the communicating channels of that system, and is partially deposited and retained as it goes by the connective-tissue cells. In this way a kind of pigmentation of the entire lymphatic system of the lung is obtained, which for demonstration may serve the purposes of an injection of that system. In this conveyance of the dust particles the leucocytes which are always present in the lymphatic spaces probably play an important part. The parts pigmented are, the walls of the alveoli, the interstitial connective tissue, especially that around the pulmonary artery and the bronchi, and the subpleural tissue, which is often definitely demarcated from the pleura proper by the pigmentation. The pigment is also carried to the bronchial glands at the root of the lungs, which are more or less blackened. This pigment is a carbonaceous material consisting mostly of round particles, and is to a great extent the soot of coal.

While the light dust of the air, reaching the lungs in small quantities, does comparatively little harm, we may have damage done when large quantities or dust of a specially irritating character reach the lungs. Hence a distinct class of diseases has been distinguished as due to inhaled dust.

**Diseases due to inhaled dust.**—This subject has been very fully worked out in this country by Greenhow, and in Germany by Zenker and Merkel. The results of the inhalation depend largely on the mechanical character of the dust. If it be heavy and composed of sharp angular particles, then it is more irritating than if it be light and composed of rounded pieces.



The particles entering the lung tissue in the manner mentioned above act as foreign bodies and set up a chronic inflammation. There is great newformation of connective tissue, as in chronic interstitial pneumonia, and great shrinking of the tissue, so that considerable deformity of the lung may occur. As the irritant finds access by the air passages the lesion in its earlier stages concentrates itself around the bronchi. This is shown in Fig. 268, in which it is seen also that the

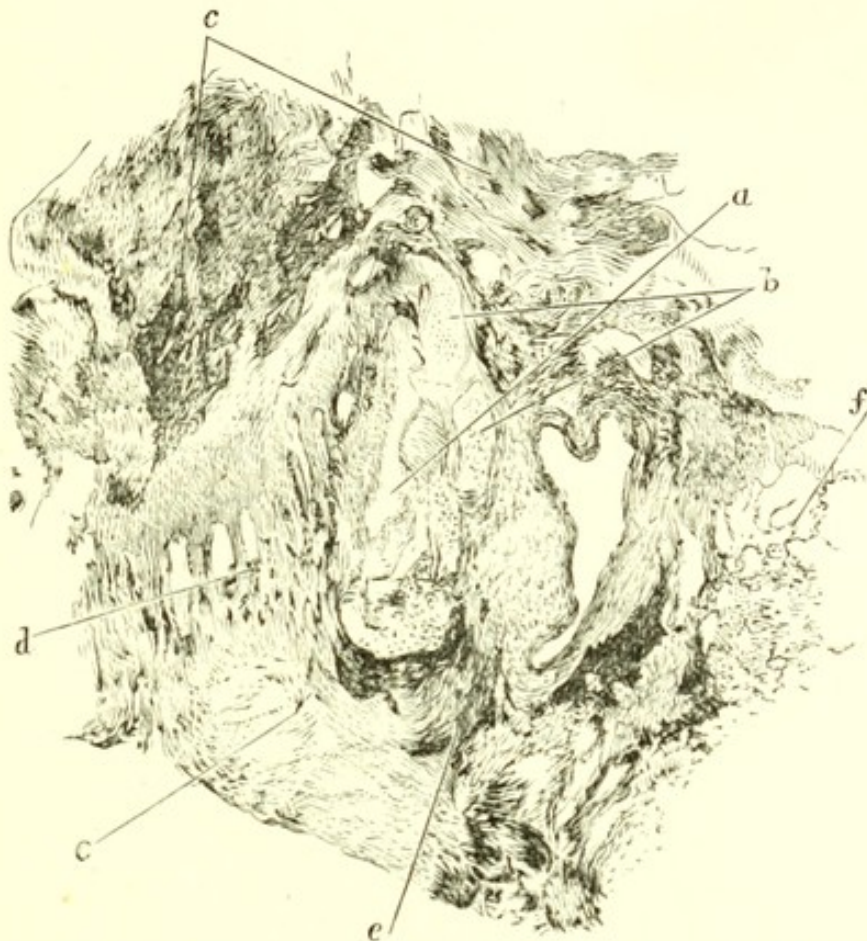


Fig. 268. —From potter's lung. *a*, Bronchus compressed and narrowed; *b*, two of its cartilages; *c*, *c*, condensed and pigmented lung tissue; *d*, lung alveoli, stretched and enlarged, some with pigment in them; *f*, lung alveoli emphysematous.  $\times 8$ .

lung alveoli are subject to great contortion (at *d*), and that in the neighbourhood they are liable to emphysema (at *f*).

The dust, whatever its kind, may be visible in the midst of the lesion in the lung. It was the observation, by Zenker, of a red pigmentation of the lung in workers with the red oxide of iron, which furnished an absolute demonstration that the dust actually finds its way into the lung tissue. We may find in the lung, coal dust, the dust of potter's clay, soot from smoky lamps, stone dust, metal dust, and dust composed of cotton or woollen fibre. In Fig. 269 a collection of silicious particles from the lung of a worker in a pottery is shown.

A peculiar feature in almost all cases of disease from inhalation



of dust is the presence of an excess of **Black pigment** in the lung. In some cases the inhaled dust is black, so that a condition of anthracosis is brought about. But even when the dust is not black, as in the potter's lung, the affected districts of lung are deeply pigmented. The reason of this is not quite apparent.

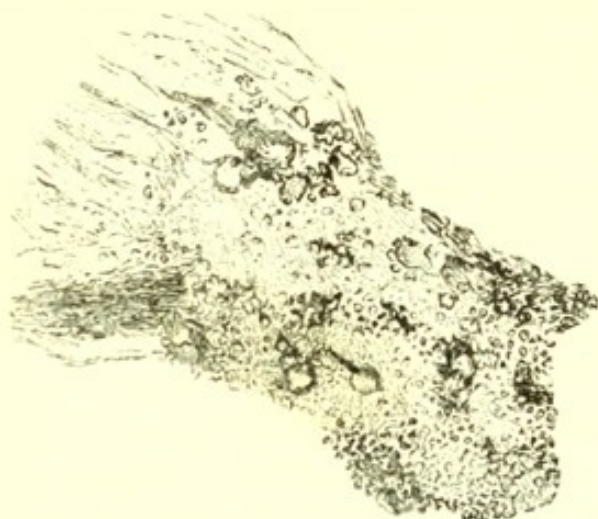


Fig. 269.—Silicious particles in potter's lung.  $\times 340$ .

when incised it yields a black juice which stains the hand. The dust here is finely divided coal and the soot from the smoky lamps used in mines. The pigment is in the connective tissue, but it is often present also in the lung alveoli, where it may be taken

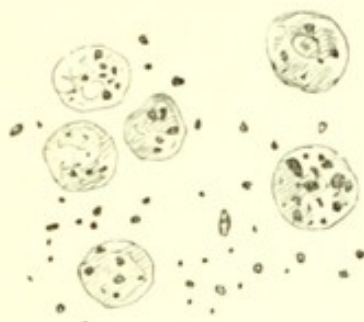


Fig. 270.—Part of the contents of a lung alveolus in anthracosis. Black particles are seen, some angular, and others rounded. The large catarrhal cells contain many particles.  $\times 350$ .

into the substance of catarrhal cells (see Fig. 270). There is not usually much induration of the coal-miner's lung, as the dust is not physically very irritating. The term **Anthracosis** is given here owing to the extremely black appearance presented.

Coal miners are subject to catarrh, and the expectoration is usually stained so as to form the **Black spit**. It is noteworthy that on a recurrence of catarrh, even years after the patient has ceased working in the pits, the black spit will return. The pigment stored up in the lung tissue will return to the alveoli and bronchi. This would indicate that a bronchial catarrh goes deeper than the mucous membrane. The agents in the return of the pigment will be the leucocytes.

**Potter's phthisis** is a term used for a very frequent form of disease amongst potters. The dust here, consisting of heavy angular particles, seems very irritating, and excites much chronic inflammation.

**Stone-mason's lung** resembles the potter's form, as does also knife-grinder's disease.

The connection of these forms of disease with phthisis is a point of some importance. A true tuberculosis may possibly be induced by the inhalation of irritating dust. There is, however, little evidence of that, and undoubtedly the great majority of cases of lung disease from inhalation of dust are quite different from phthisis. There is no formation of tubercles, no caseation, seldom any formation of cavities,



although they sometimes form by bronchiectasis. There is also the important clinical difference that persons affected with such diseases do not suffer in their general health as tubercular patients do. There is little or no fever, and the patients generally work on for many years till the dyspnoea incapacitates them (Greenhow, Coats).

**Literature.**—GREGORY and CHRISTISON, *Edin. Med. Jour.*, xxxvi. 389, 1831; HAMILTON and GRAHAM, *ibid.*, xliii. 297, 1834; THOMSON, *Med. chir. trans.*, 1840, xx. and xxi.; ZENKER, *Deutsch. Arch. f. klin. Med.*, ii. 116; ARLIDGE, On the diseases prevalent among potters, *Social Science Cong.*, 1871; GREENHOW, *Path. trans.*, 1866 to 1869; COATS, in *Lect. to pract.*, 1888, p. 150.

#### X.—ACUTE MILIARY TUBERCULOSIS, SYPHILIS, ACTINOMYCOSIS, GLANDERS.

**Acute miliary tuberculosis.**—This condition has been discussed at page 207. The tubercular virus, being present abundantly in the blood, produces very marked lesions in the lungs. As the virus is brought by the blood, the lesions are generally homogeneously distributed throughout both lungs, and are intimately related to the finer branches of the pulmonary artery.

The lungs are found studded from apex to base with innumerable small bodies, generally whitish in colour. There is in addition a general hyperæmia of the lungs.

Under the microscope it can be seen in most cases that the tubercles are formed in connection with arteries. A small artery will generally be seen at the edge or running directly into a tubercle. The tubercle has the usual structure, which is frequently obscured in the central parts by caseous necrosis (see Fig. 55, p. 207).

The lung tissue generally shows some reaction in the neighbourhood of the tubercles. The alveoli are frequently occupied by catarrhal cells as in phthisis, and there is very often blood in them. The latter is so abundant in some cases as to give a very striking character to the sections. As the cases are very acute and death occurs early, we practically never find softening of the tubercles and formation of cavities, even minute ones.

**Syphilis** is a very rare affection of the lungs, especially in adults. It is somewhat more common in new-born children, in whom it may be one of the manifestations of hereditary syphilis.

In adults we sometimes meet with gummata in the lungs, and these are surrounded by connective tissue. There may even be a softening of the gummata and formation of cavities.

**White pneumonia** is the name given to the condition met with in infants. It is a condition in which extensive tracts of lung tissue are



condensed and infiltrated by a newformation of connective tissue, the condition resembling that in chronic interstitial pneumonia.

**Actinomycosis**, when affecting the lungs, presents lesions which may resemble those of tuberculosis somewhat closely. As in tuberculosis the microbe reaches the lungs by the bronchi, and it sets up a bronchitis and lobular pneumonia. There is not a caseous necrosis of the products, but rather a more direct softening with fatty degeneration and suppuration, resulting in the formation of cavities. In other cases, however, the conditions may resemble those of fibroid phthisis, there being over an extensive tract of lung a newformation of granulation tissue which develops into connective tissue and goes on to shrinking.

The disease tends to go beyond the lung to the pleura, pericardium, muscles of chest, skin, etc.

The peculiar microbe is present, and in some cases it has been distinguished in the expectoration.

**Glanders** may affect the lung by extension from the nares and air-passages, possibly also by the blood. It leads to local inflammatory foci, which may be small or large. The inflammation is usually acute and abscesses result, but sometimes there is a more chronic affection with caseation and formation of cavities.

#### XI.—TUMOURS AND PARASITES IN THE LUNGS.

**Primary tumours** of the lungs are infrequent.

Primary fibroma, osteoma, lipoma, and enchondroma have been observed in the form of small tumours, sometimes multiple. These are unimportant.

**Primary cancer** is a form of tumour which sometimes attains to considerable size. The tumour seems to take origin in the mucous glands

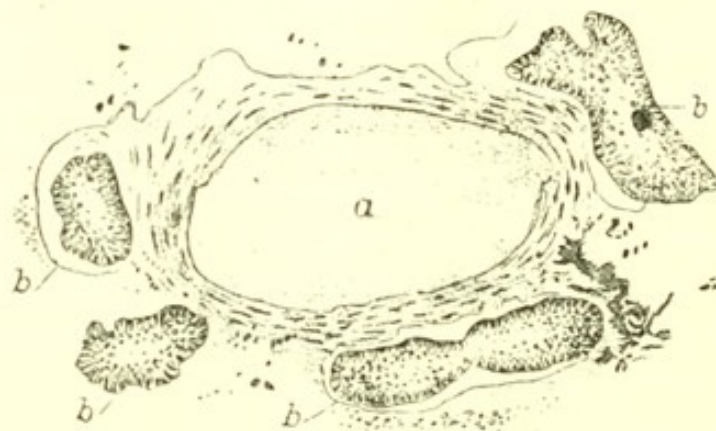


Fig. 271.—From a primary cancer of the lung, showing extension by the perivascular lymphatics. *a*, section of pulmonary artery; *b, b*, cancerous growth in lymphatics.

of the bronchi, and it usually retains somewhat of the glandular character throughout, presenting in many cases the features of the **Cylinder-**



**celled epithelioma.** The tumour may, for a time at least, confine itself to the bronchi and their neighbourhood, infiltrating the peribronchial connective tissue. But it will often extend to the parenchyma of the lung, forming in some cases bulky tumours.

In its extension the tumour will sometimes extend into the lung alveoli. On the other hand, it often forms for itself alveoli similar in size to those of the lung, but of independent origin. It penetrates into the perivascular lymphatic spaces still more than into the lung alveoli, and may often be detected filling up these around the vessels as in Fig. 271. This appearance is sometimes visible a considerable distance in advance of the edge of the tumour.

The cancer sometimes extends to neighbouring structures such as the pericardium, the wall of the heart, etc.

The cancer in the lung has a tendency to soften so as to form **Cavities**, in whose wall cancerous tissue may be found. In some cases most of the cancerous tissue from the primary tumour may be lost by this process of softening.

The author has recorded a case in which a very peculiar secondary extension occurred to the bones and to the brain. In these two situations the tumours assumed remarkable cystic tendencies, so that in the brain, they were represented chiefly by cysts (see Fig. 236, p. 584). These cysts occurred apparently by mucous or colloid change although in the primary tumour in the lung there was little such change.

**Secondary tumours** are somewhat frequent, either by direct extension from the neighbourhood or by embolism.

We have most frequently **Malignant lymphomas**, originating in the bronchial glands and extending into the lung. The newformation follows the connective tissue of the lung, and we find it penetrating in a radiating manner from the root, often burying the bronchial tubes in a sheath of new-formed tissue and partially or completely obstructing them. Sarcomas originating in the mediastinum may spread in a similar fashion, but this is rare.

Of the tumours arising by embolism, sarcoma and cancer are the chief forms, although chondromas have also been known to extend to the lungs in this way.

**Sarcomas**, as we know, sometimes penetrate directly into the veins, and portions being carried to the right heart are caught in the branches of the pulmonary artery or capillaries. The secondary tumours in the lung are multiple and they repeat exactly the structure of the primary one. Thus we have giant-celled, round-celled, and spindle-celled sarcomas. A tolerably frequent form is the pigmented sarcoma. The **Enchondroma** also not infrequently undergoes secondary development, especially in connection with tumours of the testis.



**Cancers**, as we have seen, do not readily penetrate into the veins directly. Finding an easier path by the lymphatics they nearly always first develop in the lymphatic glands. After a time the cancerous tissue may penetrate from the lymphatic sinuses in the gland into the veins, or they may possibly penetrate directly from the primary tumour into the latter, and the material is carried on to the lungs. As the cancer is usually arrested for a considerable time at the lymphatic glands, it happens that in all forms of cancer secondary tumours in the lungs are of late development. As the secondary tumours are in the lymphatic glands we may regard those in the lungs as of a tertiary order. Further, the material which produces these tumours often passes to some extent through the wide capillaries of the lung and on into the systemic arteries, so that we may have tertiary tumours occurring at the same time in a variety of organs. The tumours in the lung are multiple and they repeat the structure of the primary tumour whatever be the variety of cancer which has formed it.

That these tertiary cancers form by embolism was readily seen in a case which occurred to the author, and which may here be briefly related. It illustrates as well the tendency, which tumours even of this order show, of the cancerous masses to



Fig. 272.—Embolic cancer of lung. One of the spaces filled with epithelial cells. There is a larger space with two branches into which the epithelial cells have extended.  $\times 90$ .

penetrate into existing spaces and canals. The primary tumour was a cancer of the stomach, and secondary growths had occurred in the prevertebral glands. One of these glands was adherent to the vena cava, and several radicles of this vein emerged from the midst of cancerous glands. Actual cancerous thrombi of some of these veins were detected by the naked eye, and under the microscope it was seen that the cancerous masses had partially penetrated into the venous radicles in the glands. On examining the lungs with the naked eye it could be seen from the surface that the



newformation was largely in the lymphatic vessels which could be seen as a white network. On section the arteries were usually seen to be surrounded by new-formed tissue. Under the microscope it was seen that the finer branches of the pulmonary artery were frequently obstructed. The obstructing material was not entirely cancerous in structure. There was often a round-cell formation with now and again a distinctly cancerous appearance. In most cases the obstruction was complete. Outside the obstructed arteries there were spaces and canals filled with cancerous growths, as shown in Fig. 272. These were obviously the lymphatic channels of the sheath of the vessels into which the cancerous formation had penetrated, just as it does in primary cancer (see Fig. 271).

**Parasites in the lung.**—These are of very rare occurrence, unless we include microbes which have already been considered. A fungus of the *Aspergillus* form has been found in a few cases, in cavities, and in the expectoration in some cases (*Mycosis of the lung*).

Of animal parasites, *Echinococcus* is most frequently seen. There may be perforation of the cyst into bronchus, pleura, or peritoneum. The parasite is generally situated in the lower lobe. *Cysticercus cellulosæ* has also been observed.

Amongst rare parasites in the lung may be mentioned one case of a long round worm, the *Strongylus longevaginus*, found in a child six years of age (Diesing). It is also said that in Egypt the eggs of *distomum hæmatobium* are found in the interstitial tissue. Manson and Baelz also describe a *distomum pulmonale* as occurring in Japan. There may be twenty of the worms found in the lungs, and each in a separate cavity which communicates with a bronchus. The worms are about three eighths of an inch in length, and their eggs appear in the sputum of the patients, which also contains blood.

**Literature.**—*Tumours*—ROKITANSKY, Handbuch, iii.; VIRCHOW, Geschwülste, ii.; EBERTH, (Cancer) Virch. Arch., xlix.; LANGANS, *ibid.*, liii.; PERLS, *ibid.*, lvi.; WEICHSELBAUM, *ibid.*, lxxxv.; FINLAY and PARKER, Med. chir. trans., lx.; MACHIAFAVA, Riv. clin. di Bologna, 1874; COATS, Path. trans., 1888, xxxix. 326. *Parasites. Mycosis*—VIRCHOW, Virch. Arch., ix. and x.; FÜRBRINGER, *ibid.*, lxvi. *Echinococcus*—THOMAS, Brit. Med. Jour., 1885, ii. 692; LAVERAN, Kyst. hydat. des poumons, 1885. *Round worms*—MANSON, Lancet, 1883; BÆLZ, Berl. klin. Wochenschr., 1883.

## C.—THE PLEURA.

1. **Affections of the circulation.**—There can scarcely be any independent circulatory disturbances in the pleura. **Passive hyperæmia** exists in cases of severe dyspnœa, and this may result even in sub-pleural hæmorrhages producing **Petechiæ**, which are regarded as of diagnostic significance in death from suffocation.

**Hæmorrhage** into the pleural sac arises as a result of wounds, rupture of aneurysms, and sometimes from tubercular or cancerous new-formations in the pleura.



**Hydrothorax** is a dropsy of the pleural sac. Most frequently it is part of a general dropsy, as in Bright's disease, cardiac disease, or anæmia. Malignant tumours, especially cancers, produce hydrothorax just as they produce accumulation in the peritoneal sac. As cancers spread along the lymphatic spaces it may possibly be that the dropsy results from obstruction of these. The exudation consists of clear watery fluid. It is often limited by adhesions, which latter may also be dropsical. The lung is compressed in proportion to the amount of fluid.

**Chylous hydrothorax** is a rare form, arising from rupture of the thoracic duct (see p. 434).

2. **Inflammations of the pleura. Acute pleurisy.**—We have seen that whenever any form of inflammation comes to the surface of the lung it causes inflammatory changes in the pleura; there is acute pleurisy in acute pneumonia and in caseous phthisis, and chronic pleurisy is a constant accompaniment of all forms of phthisis pulmonalis.

On the other hand, the pleura seems to have intimate connections with the peritoneum, as there are lymphatic channels passing through the diaphragm which form communications between the two sacs.

These channels are doubtless intricate and narrow, so that the diaphragm acts to a certain extent as a barrier between the two sacs, but it is not sufficient to prevent the passage, for instance, of the tubercular virus from the one to the other. A tubercular peritonitis is nearly always accompanied by a tubercular pleurisy, perhaps limited to the lower parts of the sac. Similarly a septic peritonitis has usually a dependent pleurisy. Tumours of the peritoneum also frequently lead to similar formations in the pleura.

Besides these we may have pleurisy occurring in the course of some acute diseases, such as acute rheumatism, pyæmia, etc. In some cases there seems to be a more independent pleurisy, produced, as it is said, by cold. It is probable, however, that many cases of apparent simple pleurisy either owe their origin to tuberculosis of the lung (see ante) or are themselves tubercular in character.

There is often a localized pleurisy occurring in a limited area, just where the pleura is most directly exposed to the effects of cold, namely, in the left lower lateral region. In this position the chest is not covered by any considerable layer of muscles, as the fleshy masses of the latissimus dorsi and pectoralis major passing upwards to the arm leave, as it were, an unprotected space covered by the comparatively thin origins of the serratus magnus and external oblique muscle of the abdomen. This part is also removed from the centre of heat in the heart, and on the left side instead of the liver there is the hollow stomach. On the right side the liver is a source of heat, and renders the corresponding part on this side less exposed to cold than that on the left, but still more exposed than most other parts of the chest. It is probable that, in these localities especially, cold, acting directly on the chest wall, may cause an inflammation of the pleura by depressing the temperature.



In its **Anatomical details** acute pleurisy is closely analogous to acute pericarditis. There is hyperæmia, soon followed by a thin fibrinous deposit. This fibrinous exudation, forming a soft yellow layer, often attains to a considerable thickness, forming shaggy projections from the pleural surface especially in the region of the lower lobe. Serous fluid is also exuded, sometimes in considerable abundance. The exudation in some cases is hæmorrhagic in character.

If the inflammation goes on there is a newformation of vascular granulation tissue which may come to replace the fibrine. If two such surfaces are in contact, by absorption of the serous fluid or otherwise, coalescence occurs and complete union of the surfaces, the granulation tissue afterwards developing into connective tissue. We have already seen that the fluid accumulated in the pleural sac frequently compresses the lung greatly, producing collapse of it.

**Empyema** is a suppurative inflammation of the pleura. It may develop from an ordinary pleurisy or the inflammation may have been suppurative from the outset. In the latter case there has usually been some specially virulent irritant present in the pleura, as where a metastatic abscess in the lung has approached the pleura, or where pleurisy is one of the phenomena of septicæmia. In these cases the disease is generally rapidly fatal, and we may find remains of the fibrinous exudation mixed with abundant pus.

Where the suppuration has come on in the course of a simple pleurisy the disease is often greatly prolonged, and the pleura undergoes great thickening, being converted into a bulky layer of granulation tissue like the wall of an abscess. The granulation tissue may undergo partial transformation into connective tissue with occasional adhesion, and as the lung has been compressed by the exudation we may have drawing together of the chest to an extreme degree, should the pus be discharged or partially absorbed. The pus may dry-in instead of being discharged or absorbed. In that case it is, in course of time, impregnated with lime salts, so that considerable masses of calcareous matter may be found free in the pleura or embedded in adhesions.

**3. Chronic pleurisy and Pleural adhesions.**—We have already seen that an acute pleurisy may result in adhesion of the sac, the process of adhesion resulting from coalescence of the surfaces which have become like granulation tissue from the inflammation. Such adhesion will not occur so long as fibrine is present on the surface, and is to be carefully distinguished from the mere gluing which may occur in the earlier stages from the adhesion of the fibrinous exudation on opposed surfaces. We have also seen that in the frequent chronic pleurisy of phthisis



a similar adhesion occurs, and the method of its occurrence is similar. Without any fibrinous or serous exudation the surfaces come to have the characters of granulation tissue and coalescence with vascular communication occurs.

**3. Tuberculosis of the pleura.**—This manifests itself chiefly as an inflammation of the pleura, and hence is identical with **Tubercular pleurisy**.

The tubercular bacillus does not usually reach the pleura directly from the lung in phthisis pulmonalis, the pleurisy associated with phthisis being usually non-tubercular. On the other hand, the infection may reach the pleura by a local extension from the pericardium or peritoneum, from a tubercular lymphatic gland at the root of the lung, or from a tubercular abscess in connection with the vertebræ. It may also be conveyed by the blood.

Tuberculosis generally manifests itself at first as an acute or sub-acute inflammation, accompanied by abundant fibrinous exudation, sometimes mixed with blood. The tubercles are buried under the exudation which may be very tough, and they may escape detection unless the fibrine be peeled off. They are often best seen between the lobes of the lung, where the close contact of the surfaces hinders the deposition of fibrine, and here they are visible as closely set white nodules. The inflammation may go on to suppuration so that we may have an empyema.

The acute character usually subsides after a time, and, with chronic inflammation, newformation of connective tissue occurs, forming firm adhesions in the midst of which the tubercles may be found. As in the case of tubercular pericarditis, the adherent and coalesced pleura will show two layers of tubercles, one belonging to the pulmonary and the other to the parietal layer.

There may be a partial extension of the tuberculosis into the lung, along the interlobular septa, but it is a very superficial process.

**4. Pneumothorax.**—This has been referred to in connection with phthisis pulmonalis, and the mode of origin there indicated is that in the great majority of cases. Of the remainder there are some in which it is due to empyema—the visceral pleura having softened and the lung having become ulcerated so as to communicate with the pleura,—and others in which it is due to gangrene of the lung, or to metastatic abscesses, or to the bursting of emphysematous vesicles, or to a traumatic cause.

The air in the pleural cavity is usually at a high pressure and the cavity is much distended, so that when the chest is opened the air rushes out with some force. The pleural cavity as exposed presents a



remarkably empty appearance, the lung being compressed except where there are adhesions, which may form tense bridges across from parietal to visceral layer or may limit the pneumothorax considerably. If the patient live there is nearly always an acute suppurative pleurisy, so that the condition may be designated **Pyo-pneumothorax**.

Experimental observation seems to show that the air may be absorbed, and this is confirmed by actual clinical observation.

5. **Tumours of the pleura.**—Primary tumours are rare, but cases of osteoma and of lipoma have been recorded.

**Primary cancer** occurs in the pleura, forming, according to some, **Endothelial cancer**. In most of the recorded cases there has been great thickening of the pleura and the tumour formation was in some apparently related to the lymphatics. The pleura also presented generally the appearances of acute inflammation with fibrinous exudation. In some cases there was metastasis to other organs.

In a case observed by the author there were numerous tumours on the pleura, and also, by coalescence, larger masses, forming a layer of considerable thickness. The tumours were almost continuous over the pleura and there was no tumour anywhere else. The sac was greatly distended with a blood-coloured fluid measuring 140 ounces, which had deposited a loose brown coagulum. During life the case resembled one of acute pleurisy with great effusion. Paracentesis was twice performed, and a bloody fluid withdrawn.

Microscopic examination showed the tumour to be a superficial cancer of the pleura. There was the usual stroma enclosing epithelial cells, which were frequently fatty. The tumours were not at all deep in the substance of the pleura, there being always a layer of connective tissue beneath them. (See *Glasg. Med. Jour.*, July, 1889.)



## SECTION V.

## DISEASES OF THE THYROID AND THYMUS GLANDS.

- A.—THE THYROID GLAND.—1. *Basedow's disease*. 2. *Goitre or Bronchocele*; connection with *Cretinism*; consists of *hypertrophy and adenoma*; *colloid change* usually present; *fibrous induration and calcification*. 3. *Tumours*. 4. *Myxœdema*; an affection chiefly of skin and subcutaneous tissue, but affecting other parts; depends on *atrophic disease of thyroid*; may be from *excision of gland* (*cachexia strumipriva*); connection with *Cretinism*; producible by experiment.
- B.—THE THYMUS.—*Its various diseases*.

## A.—THE THYROID GLAND.

THE thyroid gland consists of saccules of various shapes lined with epithelium. Its function is unknown.

The gland is liable to inflammations, and has been the seat of syphilitic and tubercular newformations, but these do not present any special characters. The more important diseases are characterized by enlargement.

1. **Basedow's disease** (*Graves' disease*, *Exophthalmic goitre*).—In this the condition of the thyroid gland is only part of the morbid phenomena. The eyeballs are prominent, the heart liable to excitement, and the thyroid is enlarged and vascular. The disease seems to be due to a lesion of the vaso-motor nervous system, and the thyroid shows after death no constant lesion. It has been found hypertrophied, or abnormally vascular or cystic, but also in many cases normal.

2. **Goitre. Bronchocele**, (also called in German *Struma* and *Kropf*).—This disease consists in an enlargement of the thyroid, which, occurring as it does chiefly in certain specific localities, is ascribed to an unknown miasma. Goitre is sometimes associated with the peculiar condition of **Cretinism**, in which idiocy is associated with a stunted and deformed condition of body. By some the cretinism is ascribed to the same miasma as goitre. It may be that several morbid poisons are capable of producing goitre, and that one of them may induce cretinism as well, but there are localities where goitre occurs without cretinism. There are also cases of **Cretinism in adults** characterized chiefly by myxœdema, in which the thyroid is affected.



The lesion in goitre is primarily an increase of the normal gland tissue. This has sometimes the characters of a simple **Hypertrophy**, the saccules being multiplied (Fig. 273, *c, c*). In other cases the new-

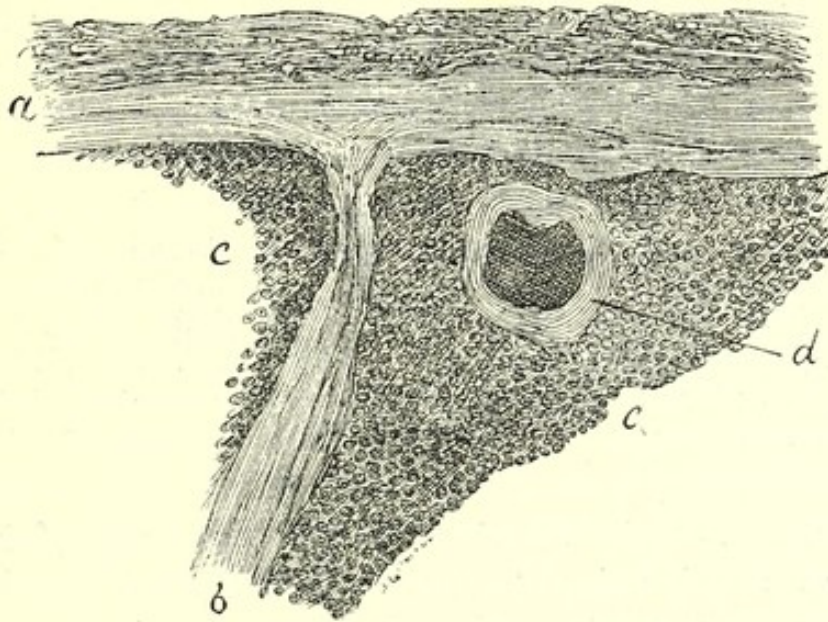


Fig. 273.—From an indurated goitre. *a*, Thickened capsule; *b*, thickened trabecula; *c*, hypertrophied gland tissue; *d*, an isolated and necrosed piece of gland tissue (adenoma).  $\times 20$ .

formed tissue is discontinuous, occurring in the form of isolable tumours in the midst of the gland. To this form the name **Adenoma** is given (see *d* in figure). In many cases hypertrophy and adenoma co-exist.

In both forms the tissue is liable to secondary changes. An almost constant condition is colloid degeneration. Some saccules nearly always contain colloid matter due to a metamorphosis of the epithelium, and in many cases there are many saccules greatly enlarged and distended with this substance. When this is sufficient to give a character to the naked-eye appearances the term **Colloid goitre** is used. Some of the altered saccules may assume preponderating dimensions so that **Cysts** develop.

The affected gland not infrequently undergoes **Fibrous induration** (as in Fig. 273); the capsule and the connective-tissue septa become thickened. In the midst of the fibrous tissue pieces of gland tissue which have undergone necrosis may be visible (as at *c* in figure). In such indurated glands calcification commonly ensues, especially in necrosed parts. A true ossification has also been observed.

The goitre is usually a simple affection, but a few cases are on record in which secondary tumours developed, chiefly in the bones. In a case observed by the author there were several such tumours in the bones of the skull, which presented the typical structure of the thyroid gland (see pp. 224 and 493).



3. **Tumours of the thyroid.**—**Cancer** is not very uncommon. It occurs chiefly as an ordinary glandular carcinoma, but also as a cylinder-celled tumour. **Sarcoma** also occurs. Wölfler has observed, besides the ordinary round-celled and spindle-celled forms, a true giant-celled sarcoma. In a case of round-celled sarcoma observed by the author there were two large cysts, and the tumour extended down into the chest as far as the pericardium. It also incorporated the anterior wall of the trachea.

**Literature.**—A very full account of Goitre and Tumours, with many illustrations by Wölfler, in Langenbeck's Arch., xxix., 1883; VIRCHOW, Krank. Geschwülste, iii.; GUTKNECHT, Virch. Arch., xcix.; BIRCHER, Der endemische Kropf, 1883. *Goitre with secondary tumours*—COHNHEIM, Virch. Arch., lxxviii., 1876; MORRIS, Path. trans., xxxi., 1880; COATS, *ibid.*, xxxviii., 1887. *Tumours*—BIRCHER, in Volkmann's Sammlung, No. 222; COATS, Catal. of Western Infirmary Museum, 1885, p. 209.

4. **Myxœdema.**—This condition was first described by Sir William Gull under the designation Cretinoid state. Dr. Ord recorded further cases, and suggested the term Myxœdema.

The disease manifests itself first as a swelling of the skin and subcutaneous tissue generally, which resembles ordinary œdema, but differs in respect that it is not influenced by gravitation, does not pit on pressure, and is obviously a more solid swelling than that of œdema. The character of the swelling seems due to the fact that mucin is present in excess in the various tissues concerned. But this is not the only change in the skin. There are also signs of irritation in the form of accumulations of round cells, chiefly in the neighbourhood of the sudariparous and sebaceous glands and the hair follicles. There is an interstitial inflammation of the skin, and it is accompanied by atrophy of the special structures. The hairs are liable to atrophy, and the secretion of the sweat and sebaceous glands is diminished.

There are less constant changes of a similar nature elsewhere, as in the mucous membrane of the mouth, and the teeth are liable to drop out like the hairs from the skin. There are also in some cases interstitial changes in the sympathetic nerves, kidneys, liver, heart, and submaxillary glands (Virchow).

Myxœdema has been found uniformly associated with **Atrophic disease of the thyroid gland**. The gland is usually diminished in size, and has a yellowish or pale appearance instead of the usual dull red colour. The condition seems to depend on an interstitial inflammation of the gland. There is an infiltration of round cells between the vesicles, followed at first by proliferation of the epithelium in the vesicles, but succeeded by atrophy. In a more advanced period the



gland tissue is replaced by fibrous tissue in which the remains of the proper gland tissue is represented by clumps of round cells.

The connection between myxœdema and destruction of the thyroid gland has been shown in various ways. Excision of the gland for disease in man gives rise to a condition virtually identical with myxœdema in a considerable proportion of cases. Kocher described the results of thyrectomy without being aware of the previous descriptions of myxœdema, and the two descriptions are very similar, and obviously refer to the same conditions. The myxœdema following excision of the thyroid has been described under the name of **Cachexia strumipriva**.

Again, **Cretinism** is accompanied by changes in the skin which much resemble those in myxœdema. In this condition, whether in the endemic or sporadic form, the thyroid gland is affected. In some cases it is atrophied, while in others it is greatly enlarged (see under Goitre).

Extirpation of the thyroid gland in animals is followed, especially in the case of monkeys, by conditions closely resembling those of myxœdema, sometimes in an acute, sometimes in a chronic form.

The chemical examination of the skin and subcutaneous tissue by Dr. Ord showed a great excess of mucin. His examination was in a case in an early stage with the full swelling of myxœdema still present. Chemical analysis of cases at a later stage shows that in them the mucin is scarcely in excess, and the swelling of the skin is due largely to an increase of the subcutaneous fat. In the experiments in animals by Horsley, in which examinations were made in an early stage, there was found an excess of mucin not only in the skin, but also in the fibrous tissues, blood and salivary glands. The parotid gland, which normally contains no mucin, presented that substance in large quantity. It showed on microscopic examination the usual appearances of a mucous gland secreting mucus, namely, gland cells in the acini distended with mucin and having the characteristic appearance of goblet cells.

In addition to the changes in the skin and elsewhere, nervous symptoms are associated with myxœdema, whether spontaneous or following extirpation of the thyroid gland. Similar symptoms have been met with in animals operated on. No changes in the nervous system have been observed to account for these symptoms.

While the connection of myxœdema with disease of the thyroid, especially atrophy, may be regarded as established, it cannot be said that it has been explained.

**Literature.**—GULL, Trans. of Clin. Soc. of London, 1873; ORD, Med. chir. trans., 1877; REVERDIN, Rev. méd. de la Suisse Rom., 1883 and 1887; KOCHER, Arch. f. Chirurgie, xix., 1883; VIRCHOW, Berl. klin. Wochenschr., 1887. A very full account of Myxœdema, with history and literature in Report to Clin. Soc. of London, 1888. In this, papers by ORD, HALLIBURTON, HORSLEY, SEMON, etc.

## B.—THE THYMUS GLAND.

The thymus gland is situated in the upper part of the anterior mediastinum, extending up in front of the trachea nearly as high as the thyroid. It attains its largest size about the second year of life,



remains stationary from that time till the fourteenth year, and then gradually atrophies. The atrophy is associated with an infiltration of adipose tissue. By some the gland is regarded as lymphatic in structure and function, but embryologically it seems to have an epithelial origin. It contains round bodies which are sometimes calcified, the so-called **Concentric bodies of Hassal**.

The thymus is liable to **Inflammations** like other glands. **Hyper-trophy** also occurs, and cases of asthma and of spasm of the glottis have been ascribed to this condition, but without sufficient evidence.

Enlargements occur in **Leukæmia** and in **Hodgkin's disease**, the relation of the organ to the spleen and lymphatic glands being thus indicated.

**Tumours** also occur, chiefly sarcomas, and also cancers.

**Syphilis** sometimes manifests itself by the presence of gummata, and also, it is said, by suppuration. **Tuberculosis**, according to Jacobi, is not of infrequent occurrence, both as a local affection and in general tuberculosis.

**Literature.**—SANNÉ, in *Dict. encycl. des sc. méd.*, Article Thymus, 1887; JACOBI, *Comptes rendus du Congrès pour l'étude de la tuberculose*, 1888, and *Anat. and Path. of thymus*, *Trans. Assoc. of Amer. Phys.*, Sept., 1888.



## SECTION VI.

## DISEASES OF THE ALIMENTARY CANAL.

- A.—THE MOUTH.—I. INFLAMMATIONS. 1. *Catarrh*, 2. *Thrush*, 3. *Special inflammations*, 4. *Cancrum oris*, 5. *Glossitis*. II. INFECTIVE AND OTHER TUMOURS—*Syphilis*, *Tuberculosis*, *Tumours proper*. III. AFFECTIONS OF THE TEETH—1. *Caries*, 2. *Inflammations about the teeth*, 3. *Syphilitic teeth*, 4. *Tumours connected with the teeth*.
- B.—THE SOFT PALATE, PHARYNX AND TONSILS. 1. *Catarrhal angina*, 2. *Phlegmonous inflammation*, 3. *Diphtheria*, 4. *Acute tonsillitis*, 5. *Chronic tonsillitis*, 6. *Syphilitic disease*, 7. *Tuberculosis*, 8. *Tumours*.
- C.—THE ŒSOPHAGUS. 1. *Dilatation, General dilatation and partial dilatation, the latter including Diverticulum, (a) by pressure, (b) by traction*, 2. *Obstruction*, 3. *Inflammation*, 4. *Rupture and perforation*, 5. *Tumours*.
- D.—THE STOMACH. *Introduction and post-mortem changes*. I. MALFORMATIONS AND CONTRACTIONS. II. DILATATION AND HYPERTROPHY. III. INFLAMMATIONS, *including catarrh and effects of corrosives and poisons*. IV. SIMPLE PERFORATING ULCER—*Characters; Effects and modes of origin*. V. HYPERÆMIA AND HÆMORRHAGE. VI. TUBERCULOSIS AND SYPHILIS. VII. TUMOURS, *principally Cancer. Structure and mode of growth of cancers. Forms*—1. *Cylinder-celled epithelioma*, 2. *Soft cancer*, 3. *Scirrhus*, 4. *Colloid cancer*.
- E.—THE INTESTINES. *Introduction and post-mortem changes*. I. MALFORMATIONS. II. EMBOLISM AND HÆMORRHAGE. III. HERNIA—*Causation; Structure of sac; Forms of external and internal hernias; Contents; Irreducible hernia; Strangulation*. IV. TWISTING. V. INTUSSUSCEPTION AND PROLAPSE. VI. INFLAMMATIONS—1. *Catarrh*, 2. *Phlegmonous and diphtheritic inflammations*, 3. *Localized inflammations, especially syphilitic*. VII. SPECIFIC INFLAMMATIONS—1. *Dysentery*, 2. *Cholera*, 3. *Typhoid fever*, 4. *Anthrax*, 5. *Actinomycosis*. VIII. TUBERCULOSIS AND SYPHILIS, *chiefly the tubercular ulcer*. IX. RETROGRADE CHANGES AND ABNORMAL CONTENTS. X. TUMOURS, *chiefly cancer*. XI. OBSTRUCTION OF THE INTESTINE—*Some unusual causes; Effects, including stercoraceous vomiting*.

**I**NTRODUCTION.—The alimentary canal is lined throughout by a mucous membrane, which consists of loose connective tissue covered with epithelium in one or several layers. In the mucous membrane and to some extent beneath it, are glands which in different parts have different structures, but everywhere secrete materials that pass into the calibre of the canal. In addition to these there are in many parts



closed follicles of a lymphatic structure which usually lie near the surface and frequently take part in changes going on there. Beneath the mucous membrane is the submucous tissue which, for the most part, is loose, and so allows the mucous membrane to go into folds or be stretched out flat according to the state of dilatation or contraction of the canal. Outside this there is a muscular coat, generally in two layers, by means of which movements and variations in calibre are effected. In most parts of the intestinal canal there is a serous coat outside of all.

In the diseases of the alimentary canal it is chiefly the mucous membrane that we have to deal with, the subjacent structures being usually subordinate and affected secondarily to it. The serous coat, it is true, is often affected independently; its diseases, however, do not belong specially to the alimentary canal, but to the peritoneum as a whole. The mucous membrane, on account of its exposure to a variety of influences derived from the varying contents of the canal, is specially liable to inflammations, and the great majority of the diseases to be considered here are inflammatory ones.

#### A.—THE MOUTH.

As the mouth is exposed in an especial manner to external influences, so its mucous membrane possesses an epithelium in many layers, and it is not nearly so liable to inflammations as most other parts of the alimentary canal.

##### I.—INFLAMMATIONS OF THE MOUTH. STOMATITIS.

1. **Catarrh.**—If we leave out of view the catarrhs of the fauces and pharynx which we consider afterwards, **Simple catarrh** of the mouth is exceedingly rare as a primary disease. It is not infrequent, however, as a secondary affection, arising from the irritation of carious teeth, from the use of mercury, the presence of ulcers, etc. It is also of frequent occurrence in the acute fevers, especially typhus, scarlet fever, small-pox and measles. In these fevers there is also a catarrh of the stomach, and the inflammation in both seems due to the action of the morbid poison in the blood.

The mucous membrane is swollen and red and there is greatly increased desquamation of the epithelium, especially inside the cheeks and on the tongue. The desquamated epithelium is mixed with leucocytes, serous exudation, and mucus in varying proportions, and on that account it has varying characters. On the tongue there is usually much epi-



thelium which lies on the surface, and so a tolerably thick layer is formed mostly of a whitish or yellowish colour—**Furred tongue**. On the cheeks and gums there is more fluid and less epithelium. In the midst of the epithelium and leucocytes bacteria and leptothrix threads are to be found. If the patient is feverish and lies with the mouth open, the catarrhal products dry-in and form a dirty brownish coating of the tongue and gums which goes under the name of **Sordes**.

Sometimes the inflammation centres especially around the mucous glands, and they may form prominent nodules or vesicles. To this form the name **Follicular stomatitis** has been given; it is frequently seen in children during dentition or after measles, and is often associated with catarrh of the stomach or intestine. The vesicles or small prominences frequently burst and leave multiple small ulcers covered by a dirty exudation.

Catarrh of the mouth rarely assumes a phlegmonous character, in this respect contrasting with inflammation of the fauces.

**2. Thrush, or Aphthous stomatitis, or Soor.**—This condition has already been mentioned as connected with the presence of a fungus.

It occurs chiefly in the mouths of young unhealthy children, but is also occasionally seen in emaciated adults, as in diabetes. The normal secretion of the mouth is alkaline, but in this disease it becomes acid, which may perhaps be the reason that the fungus develops. The mucous membrane is found beset with small white spots, which look like little bits of white curd

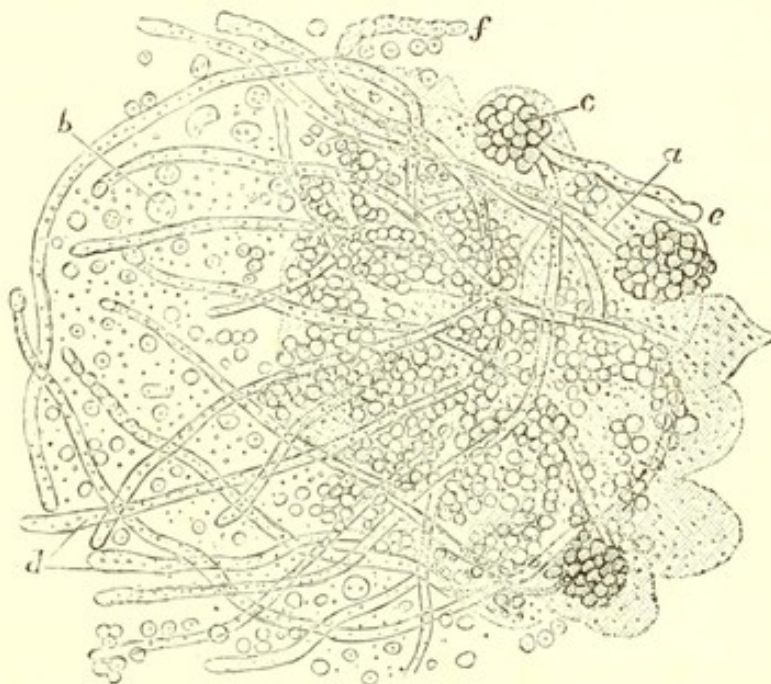


Fig. 274.—*Oidium albicans* or thrush-fungus. The constituents of the fungus are associated with flat epithelium.  $\times 450$ . (KUCHENMEISTER.)

on the surface. These patches are, however, adherent, and on removing them the mucous membrane beneath will be found red and bleeding. The white patch consists of epithelium united into a membrane by the fungus, the *oidium albicans* (Fig. 274). It consists of branching threads composed of elongated cells placed end to end and sometimes losing themselves in masses of spores. The aphthous patches frequently extend from the mouth downwards to the pharynx and œsophagus.



3. **Special forms of inflammation.**—In **Small-pox**, besides the general catarrh of the mouth already mentioned, there are vesicles or pustules analogous to those on the skin. We have first whitish patches consisting of raised and desquamating epithelium. The epithelium is soon discharged and superficial ulcers result. In **Scurvy** there is great swelling and œdema of the mucous membrane of the gums around the teeth. The gums bleed and, apparently as a consequence of this, ulcers form at the edges of the teeth and may extend down to the bone, which may undergo necrosis. Very often the teeth are loosened. As a consequence of the **use of mercury** we sometimes have a considerable stomatitis. It occurs after the medicinal use of mercury and also among workmen who employ it in their occupations. The mucous membrane is swelled, especially that of the gums and cheeks, and there is severe salivation. Ulcers frequently form, especially on the internal surface of the cheeks and lips, and on the edges of the tongue. They may be in the form of flat excoriations or deeper ulcers with a membranous covering.

4. **Cancerum oris, or Noma, or Gangrenous stomatitis.**—This disease, which is fortunately a rare one, occurs in badly nourished children, particularly when reduced by severe illness such as scarlet fever or measles. It presents itself first as a diffuse swelling of the cheek which

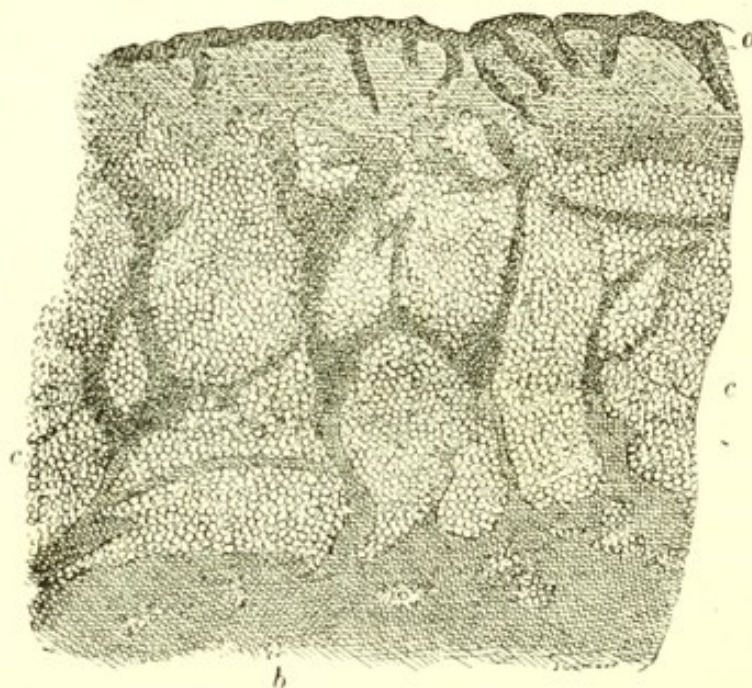


Fig. 275.—Section of cheek in case of Cancerum oris. *a*, cutaneous surface; *b*, necrosed part; *c c*, border or living tissue with abundant cells.  $\times 10$ .

is seen to be tense, red, and glistening, with one spot in the centre usually redder than the rest. On examining the inside of the mouth there is already an excavated ulcer opposite the red spot on the cheek, and the gums opposite may also be ulcerated. As the disease pro-



gresses more and more of the mucous membrane of the mouth is ulcerated away. At the same time the red spot on the cheek gets black in the centre, and afterwards extends in area. By and by a slough, which varies in size, separates and a communication forms through the cheek with the inside of the mouth. If the patient survive, still further destruction occurs, the necrosis passing on to the surrounding skin of the face, even to the ear and eyelids. There may be necrosis of the jaws. Not infrequently this disease is associated with a gangrenous pneumonia and there are general symptoms of septic poisoning.

The disease advances by a progressive necrosis, as shown in Fig. 275 (which represents a section through the cheek, stained with alum carmine). At the lower part of the figure (*b*), adipose and fibrous tissue are shown, without any differentiation of nuclei, this being evidence of necrosis. Near the middle of the figure the necrosis ends, and there is a border (*cc*) at which cells are very abundant, as indicated by the dark spots in figure. The rest of the section shows the structure as in an ordinary section, but with more abundant cells. It is apparent that the morbid agent in its advance produces necrosis, almost directly, but leads also to inflammation, which is not very great, in front of its advance.

The author found, in addition to the usual microbes of decomposition at the sloughing surface, a very remarkable growth of a bacillus into the necrosed, but otherwise unaltered tissue. The microbe was in the form of long threads, occasionally with evident spores, and it extended in large numbers to the very edge of the necrosis, and to a very slight extent beyond it into the inflamed part. The bacillus was apparently anaërobic, growing into the tissue away from the surface, and it had generally the characters of that of malignant oedema. It was readily stained by Gram's method, but not by ordinary watery solutions.

5. **Inflammation of the tongue** deserves a brief special notice. We have seen that the tongue takes part in most of the inflammations of the mucous membrane, but sometimes it becomes the seat of a special inflammation of its substance, a true **Glossitis**. This may occur as a result of wounds or irritations from without, or in the course of such diseases as erysipelas and small-pox. The tongue swells greatly in consequence of the inflammatory infiltration of the interstitial connective tissue, and this sometimes, though rarely, goes on to the formation of abscesses. The muscular fibres are swollen, pale and brittle. Usually the inflammation ends in resolution, but sometimes it becomes chronic, and there may be great increase of connective tissue with permanent induration of the tongue.



## II.—INFECTIVE TUMOURS AND TUMOURS PROPER OF THE MOUTH.

**Syphilis.**—We have various syphilitic lesions in the mouth, from slight inflammation to deep ulceration.

It is not uncommon to meet with a **Primary chancre** on the lips. In that case there is a prominent tumour which has, in more than one case, been taken for an epithelioma. On microscopic examination the tumour is found to consist of immense aggregations of round cells which may be altogether beneath the epithelium (see Fig. 47, p. 190, which is from a chancre of the lips excised under the belief that it was an epithelioma.)

The **Catarrh** which accompanies the **Secondary stage** is often very slight, but it may be accompanied by superficial ulceration of considerable extent. **Mucous patches, Warts, and Flat condylomata** are particularly frequent in the mouth, and they often break down and form superficial ulcers, especially on the lower lip and at the angle of the mouth.

The deeper ulcers arise in connection with **Gummata**, which form in the substance of the mucous membrane, and may lead to very serious loss of substance. If healing occurs a cicatrix forms, but cicatrices sometimes occur without ulceration having taken place, the gumma itself giving place to a cicatrix. The gummata not infrequently have their seat deep in the substance of the tongue, and the resulting cicatrices, with or without ulceration, may produce very marked deformity of this organ.

**Tuberculosis** is uncommon in the mouth. It may extend from the skin in **Lupus**, and in that case sometimes produces considerable ulceration. **Tuberculosis of the tongue** occurs chiefly in connection with that of the epiglottis or tonsils. There may be a somewhat deep-seated infiltration, so that tubercles are present even amongst the muscles, and interstitial inflammation causes induration of the tongue. Ulceration may occur, or there may be considerable caseous masses in the substance of the tongue.

**Leprosy** also affects the mucous membrane of the mouth sometimes, and **Glanders** occasionally manifests itself here.

**Tumours proper.**—These are of somewhat frequent occurrence and considerable variety. We meet rarely with fibromas, lipomas, and enchondromas. Rather more common are **Adenoid tumours** of the lips. These take origin in the mucous glands of the lips, and frequently become converted into mucous or colloid **Cysts**. They form prominent rounded tumours, sometimes as large as a hazel nut, and are readily enucleated.



**Warts** occur on the lips, especially at the edges. In that situation they are mostly hard, while those on the proper mucous membrane are soft. Not infrequently the wart ulcerates, and, it is said, may give origin to a cancerous tumour. Warty outgrowths also occur on the tongue. The papillæ of the tongue sometimes undergo great elongation, especially their epithelial layers, and we may have a condition as of hairs on the tongue.

The **Angioma** is met with chiefly on the lips, usually in the form of the congenital nævus. It occurs either as the cavernous or capillary angioma, and forms flat elevations or rounded prominent tumours. It occasionally occurs on the tongue.

The **Lymphangioma** is a cavernous lymphatic tumour, which occurs chiefly in the tongue, and in some cases contributes to the formation of macroglossia. The whole tongue may be permeated with dilated lymphatic vessels.

**Macroglossia** is a condition in which the tongue is greatly increased in bulk, the enlargement being nearly always congenital. It is frequent in cretins. Even at birth the tongue may be too large for the mouth and project beyond the lips. Afterwards it may increase still more, and, as the child grows, it may displace the alveolar processes considerably. For the most part there is no hypertrophy here except of the interstitial connective tissue, but evidences of newformation of muscular tissue have been observed. As a rule, the lymph spaces of the hypertrophied connective tissue are greatly enlarged, and there is even a formation of cavernous tissue, the spaces in which are filled with lymph. In this way we may speak of lymphangioma cavernosum as taking part in the condition. It will be observed that the condition here somewhat resembles that in elephantiasis, there being in both cases a great newformation of succulent connective tissue with wide lymph spaces. The resemblance is further indicated in the fact that the lips frequently hypertrophy as well (*Macrochelia*).

**Sarcomas** rarely develop in the mouth itself, although they have been met with in the tongue.

On the other hand, it is not uncommon to find a **Myeloid sarcoma** of the jaws, growing from the periosteum and projecting into the mouth. This forms the majority of the tumours called **Epulis**. This name is given generally to tumours which arise from the alveolar processes of the jaws. They are mostly composed of spindle-cells, but nearly always contain **Giant-cells**, and sometimes these are in large numbers. Osseous trabeculæ frequently pass into them from the bone beneath. As these tumours grow they push the mucous membrane of the gums before them and so form red prominences of a rounded form, behind, in front of, or



between the teeth, and are generally of dense consistence. The teeth are often considerably displaced by them as they grow, especially when they assume large dimensions, as sometimes happens.

**Cancers** of the mouth are nearly all **Flat-celled epitheliomas**. They are of very frequent occurrence on the lower lip and are also common in the tongue.

Epithelioma of the lip is almost confined to the male sex. This is frequently ascribed to the fact that in men the lip is more frequently exposed to irritation by smoking short pipes and in shaving. The tumour which has almost always its seat on the lower lip, is in the form of a superficial infiltration which soon goes on to ulceration. There may thus be great destruction of the lip structures. The structure is that of the typical epithelioma, of which this is the most frequent seat.

In the tongue the epithelioma usually forms at the edge, and it is often said that the irritation of the sharp edge of a carious tooth has been the starting point. There is here also usually a superficial ulcer, but the tumour generally penetrates deeply into the substance of the tongue. There may also be considerable irritation of the tongue and newformation of connective tissue so that the structure may resemble that of a scirrhus. As the epithelium of the tongue does not become horny, we scarcely have the typical laminated capsules of the ordinary flat-celled epithelioma.

Secondary epitheliomatous formations are liable to occur in the submaxillary lymphatic glands, and may extend to the glands of the neck.

**Ranula** is a name applied to cysts which form beneath the tongue. These mostly arise as retention-cysts from closure of Wharton's duct (duct of the submaxillary gland), or the duct of the sublingual gland, but they may take origin in the mucous glands. Before the occurrence of the cyst there is usually some inflammation of the floor of the mouth very often connected with affections of the teeth. According to the observations of Recklinghausen referred to at page 249, the cyst arises by dilatation of the duct, while the gland structure persists and furnishes the material by whose accumulation the cyst forms (see Fig. 82, p. 250).

**Literature.**—BOHN, *Mundkr. der Kinder*, 1886, and in Gerhard's *Handb. d. Kinderkr.*, iv.; HIRSCH, (Noma) *Hist. and geogr. path.* (Syd. Soc. transl.), iii. 272; WEST, *Dis. of infancy and childhood*, 7th ed., 1884; BRUNS, *Handb. d. oper. Chirurg.*, 1859; KEHRER, *Soorpilz*, 1883; PLAUT, *Soorpilz*, 1885; GRAWITZ, *Virch. Arch.*, lxx., lxxiii.; LANG, *Path. u. Therap. d. Syph.*, 1885; HUTCHINSON, *Syphilis*, 1887; HAUSEMANN, (Tuberculosis) *Virch. Arch.*, ciii.; VIRCHOW, (Macroglossia) *Geschwülste*, ii.; ARNSTEIN, *Vich. Arch.*, liv., 1872; BILLROTH, *Path. Hist.*, 1858; RECKLINGHAUSEN, *Virch. Arch.*, lxxxiv., v., 1881.



## III.—AFFECTIONS OF THE TEETH.

The hard part of the tooth consists of enamel, dentine, and cement. The **Enamel**, containing very little organic matter (only about 2 or 3 per cent.), is for the most part passive, and presents great mechanical resistance to destructive processes. In its original formation it is a superadded structure, derived from the epithelium of the surface. The **Dentine**, consisting of tubules with a hard matrix, possesses a much larger proportion of organic matter (28 per cent.), and is much more directly involved in morbid processes. The **Cement**, which covers those parts of the tooth which are devoid of enamel, consists of true bone, and is liable to changes of a similar nature to those of bone.

Turning to the soft parts, the **Pulp** fills the cavity of the tooth and has an outline parallel to that of the tooth as a whole. The pulp is a highly vascular tissue, and richly supplied with nerves whose filaments pass in part into the dentinal tubules. The pulp is so sensitive that in popular language it is called "the nerve." In the original formation of the tooth the dentine is produced by the structure which is afterwards represented by the pulp, and through life the latter retains to a considerable extent its formative power.

The tooth is fitted accurately into the alveolus by means of the cement on the one hand and the **Dental periosteum** or periodontal membrane on the other. This membrane, lying between the cement and the bone of the jaw, forms a kind of double periosteum, and is, from its position, very liable to the action of mechanical forces, being placed between two rigid structures, one of which (the tooth) is peculiarly exposed to mechanical interference. It is continuous at the apex of the root with the pulp, and at the neck with the mucous membrane of the gums on the one hand and the periosteum of the jaw on the other.

1. **Caries.**—This name is applied to a condition which is not analogous to caries of bone, except in so far as in both there is destruction of the dense calcified structure. Caries of bone is related to inflammatory processes, but caries of the teeth has no such origin. It consists of a softening, usually progressive, of the enamel and dentine and their subsequent disintegration. The process appears to begin very commonly in places where the enamel shows normally rather deep furrows, and is therefore thinner and more easily destroyed than elsewhere. When the caries reaches the dentine it advances more freely, so that the enamel may to some extent be undermined. The lime salts are first absorbed, and then the organic basis is broken down. The caries advances in the direction of the dentinal tubules, as shown in Fig. 276.



Various views have been held as to the nature of the process. At one time it was regarded as inflammatory, but this view may be set aside, chiefly on the ground that a process exactly similar occurs in teeth which have been drawn and re-inserted, as well as in artificial teeth made of the ivory of the hippopotamus. The caries consists, in fact, of a gradual solution of the lime salts, and for this solution we must infer the existence of an acid. The secretion of the mouth is naturally alkaline, but in carious teeth an acid reaction has been detected. The acidity is often connected with derangements of the

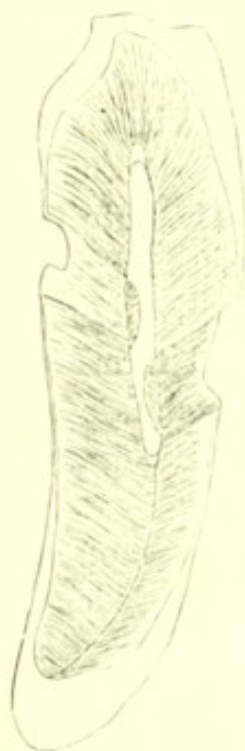


Fig. 276.—Section of tooth with caries. There are two cavities, one on each side, and in the pulp cavity a newformation of secondary dentine. The secondary dentine is not exactly on the same level as the cavities, but at the extremities of the dentinal tubules leading from these. (After SALTER.)

stomach, but it may have a more local origin, as when the secretion of the gums is abnormal, or perhaps when food is undergoing acid fermentation in contact with the teeth. It may be that before the teeth yield to an undue acidity they have already an abnormally weak power of resistance, and this may be related to personal peculiarities, inherited or otherwise.

At the advancing margin of the caries a widening of the dentinal tubes is visible, and in these widened tubes, as well as in the carious cavity, bacteria and *Leptothrix* threads are to be found. By some the process has been ascribed to the action of these organisms, especially by Klebs.

2. **Inflammation** frequently follows on caries, and the pulp is usually involved. It is almost certainly attacked if the caries causes penetration into the pulp, but before this takes place there is frequently some inflammation which expresses itself in a newformation of **Secondary dentine** inside the tooth around the point to which the caries is advancing (see Fig. 276). In this way the carious cavity may be shut off from the cavity of the pulp, and so a more serious inflammation warded off. Very frequently, however, a more acute inflammation of the pulp occurs, with redness, swelling, great pain, not infrequently also with complete necrosis of the pulp, and, in consequence, of the whole tooth.

The inflammation often extends to structures around the tooth. The root-membrane or dental periosteum, which covers the root portion of the tooth, is the structure most directly attacked. This membrane is, as we have seen, intimately connected, towards the neck of the tooth, with the submucous tissue of the gums and the periosteum of the alveolar process of the jaw. Inflammations of these structures, especially of the gums, very frequently ensue, resulting in the well-known **Gum-boil**,



which often goes on to suppuration and abscess, forming the so-called **Parulis**. The abscess usually bursts into the mouth, but it may produce extensive swelling of the gum, and lead even to penetration outwards, resulting in a fistulous opening in the skin.

The result may be even more serious if the periosteum of the jaw becomes inflamed, resulting, it may be, in suppuration which is apt to be chronic. With the periostitis there is usually newformation of bone, causing thickening of the jaw.

Inflammations of the root-membrane and periosteum of the jaw sometimes occur apart from caries, as a result, for instance, of poisoning with phosphorus or mercury, and in scurvy. In that case the inflammation is not limited to the neighbourhood of one tooth, but extends probably to a series.

**3. Conditions resulting from hereditary syphilis.**—In children who are the subjects of hereditary syphilis, the teeth show malformations first pointed out by Hutchinson. The teeth are narrow and pointed; the most obvious lesion is in the permanent front teeth of the upper jaw. The upper incisors are narrowed instead of being expanded towards the cutting edge, and the central ones have generally a crescentic notch which is very characteristic. This condition of the teeth is frequently associated with interstitial inflammation of the cornea, and is ascribed by Hutchinson to a syphilitic inflammation of the gums during the formation of the teeth.

**4. Tumours connected with the teeth.**—The cement, which, it must be remembered, is composed of bony tissue, sometimes undergoes a hypertrophy to which the name **Exostosis** or **Osteoma** is often given. This is scarcely a true bony tumour, but originates rather in a chronic inflammation of the cement, leading to a considerable newformation of bony tissue. In this way are formed prominent tuberos outgrowths from the roots of the teeth, which may be localized to one part of the fang, or cover a considerable portion of it, or even the whole root portion of the tooth. In the latter case it is as if the root were enlarged by rough accretions on its surface. These so-called exostoses sometimes offer serious resistance to the extraction of the teeth.

Another form of tumour connected with the teeth is that which Virchow has called the **Odontoma**. This tumour arises in connection with teeth retained in the alveoli by faulty development. The tumours are composed of dentine and enamel, and are of small size and rare occurrence.

**Cysts of the jaws** have already been referred to (p. 491) as usually taking origin in connection with the teeth or their rudiments, and **Cancers** have occasionally a similar origin (p. 493).



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## B.—THE SOFT PALATE, PHARYNX, AND TONSILS.

**Introduction.**—The mucous membrane here differs from that of the mouth proper, chiefly in respect that in addition to the ordinary mucous glands there are numerous **Lymphatic follicles**. The distinction between these two is not always correctly appreciated. The mucous glands are racemose glands with proper ducts, opening on the surface of the mucous membrane. The follicular glands, as they are often called in rather a confusing way, are strictly comparable with the closed follicles of the intestine. In the pharynx, soft palate, and root of the tongue they occur in the form of isolated rounded masses of lymphatic tissue, like the solitary follicles of the intestine. In the tonsils we have aggregations of these follicles not unlike Peyer's patches. Their prominence in the tonsils causes the mucous membrane to be thrown into folds, and so we have comparatively deep recesses, which are sometimes called **Crypts**, and in which secretions may accumulate, especially if their depth is exaggerated by diseased conditions.

From their exposed position these parts are peculiarly prone to irritation from agents coming to them by the inspired air. The mucous membrane of the mouth seems peculiarly resistant to external irritations, but here, where there is the transition, as it were, from the mucous membrane exposed to external influences to that protected from them, these inflammatory manifestations occur with peculiar frequency. Almost all the diseases we have to treat of here are forms of inflammation, and it is usual to describe these under the general designation of Angina.

**1. Catarrhal angina.**—In its acute form this constitutes the most ordinary sore throat, and occurs either from "cold," especially at seasons of the year when sudden changes of temperature are prevalent, or as a local symptom of a general disease such as measles, scarlet fever, small-pox. It may be said regarding scarlet fever, small-pox, and diphtheria that the throat affection in them begins with an acute catarrh, but that, especially in scarlet fever and diphtheria, it nearly always goes on to phlegmonous inflammation. In the simple acute catarrh there is redness and swelling of the mucous membrane with a mucous exudation which covers the surface and may have a tough consistence. Sometimes vesicles form on the mucous membrane.

**Chronic catarrh** very frequently follows on the acute form, especially



when there have been repeated attacks. The conditions are somewhat different in different cases, and rather complicated classifications have been introduced. As in other chronic inflammations there is here, for the most part, thickening of the mucous membrane, but in some cases the mucous membrane is indurated and atrophied. There may be a general hypertrophy of the superficial layers of the mucous membrane, and, as the blood-vessels remain congested, it is a rather succulent swelling. This constitutes what is called the **Relaxed throat**. The appearances are most pronounced in the case of the uvula, which becomes elongated, sometimes to a very marked extent. At other times the swelling is not so uniform, but with some general thickening there is an occasional localized prominence causing the mucous membrane to assume a granular appearance, hence the term **Granular pharyngitis**. This form occurs mostly among clergymen, singers, actors, and others whose occupation requires them to use their voices for prolonged periods with a loud tone.

It is not by any means certain what the exact structure of the prominent granulations in this disease is, but it seems likely that there are really two conditions included under the name. In one of them the mucous glands are involved and may be seen discharging a milky fluid. In the other the enlargement is not connected with the mucous glands, but is usually supposed to depend on enlargement of the closed follicles, although Stoerk has stated that it really consists of a hypertrophy of the epithelium. The condition is often called follicular pharyngitis from the view that the closed follicles are mainly concerned. Ulcers may form in connection with these swellings and they are called follicular ulcers.

2. **Acute phlegmonous inflammation of the fauces.**—This frequently results in the formation of **Abscesses**. It is a condition of somewhat common occurrence. There is not merely a surface catarrh, but the mucous membrane and submucous tissue are involved in an acute inflammation of a very intense kind which usually goes on to suppuration. The disease very commonly begins on one side and frequently involves the tonsil, sometimes extending thence to the posterior wall of the pharynx. The swelling and redness are very great from hyperæmia and œdema of the whole structures, and the patient may have difficulty in opening the mouth. If it goes on to suppuration the pus often collects and forms an abscess which bursts into the throat. In some cases the acute inflammation extends downwards to the base of the epiglottis and the case becomes dangerous because of the possible super-vention of œdema glottidis. It should be added that the name **Quinsy** is applied to this disease as well as to a more local inflammation of the tonsils alone.



We may infer that in this disease one of the pyogenic microbes finds access to the mucous membrane. It sometimes happens that the inflammation, after attacking one side, passes round and involves the other, and this looks like an irritant which propagates itself.

**3. Diphtheria.**—The changes which occur in this disease have already been referred to in considering its manifestations in the air passages (see p. 610). It usually begins in the fauces and has its centre there. As we have seen there is in the fauces a fibrinous exudation usually associated with necrosis of the mucous membrane. There are other signs of inflammation, chiefly in the presence of leucocytes infiltrating the mucous membrane and passing into the exudation. As a rule, there is necrosis of the mucous membrane, and this in all degrees till we come to the so-called gangrenous form. The resulting sloughs frequently look more serious than they are, but there may be considerable loss of substance, and ulcers of some depth and size may be left.

The position of the patches of exudation is very various. Sometimes they are mainly on the tonsils, sometimes on the soft palate and uvula. Extension to the posterior nares, on the one hand, and to the larynx, on the other, is very frequent, the latter, as we have seen, being particularly common.

**4. Acute tonsillitis. Quinsy.**—This inflammation of the tonsils is, as we have seen, occasionally a part of a general phlegmonous inflammation of the fauces. Occurring more independently it is accompanied by considerable swelling, consisting of an inflammatory infiltration of the lymphatic follicles of the tonsils. Sometimes it goes on to suppuration, but rarely does so when the tonsils alone are affected. The swelling in many cases does not fully subside, and gives rise to a more or less permanent enlargement such as we have next to consider.

**5. Chronic tonsillitis. Hypertrophy of the tonsils.**—Certain persons, especially in childhood or youth, are prone to repeated subacute inflammations of the tonsils, and as these recur the tonsils acquire a permanent enlargement. This consists anatomically in a true hypertrophy of the lymphatic follicles, although there is sometimes also an increase of the interstitial connective tissue. The tonsils are rarely much indurated, the soft lymphatic tissue existing so abundantly as to make the structure as a whole somewhat soft.

The enlargement is very variable, there being sometimes a rapid enlargement followed by a rapid subsidence.

The inflammation sometimes leaves behind somewhat deep pits and recesses in the tonsils. These may contain secretions which sometimes become condensed into concretions (*Calculi in tonsils*).

The contents of these exaggerated crypts often decompose and irritate the tissue around.



The hypertrophy is not always the result of chronic inflammation. There are cases of congenital hypertrophy of the tonsils, and there are still more frequent cases of a gradual enlargement without any apparent attacks of inflammation. In this way the tonsils may acquire very large dimensions, reaching the size of a hen's egg on some occasions. These conditions also are met with chiefly in children.

6. **Syphilitic disease of the fauces.**—This manifests itself in various ways. A persistent **Catarrh** having the ordinary characters of subacute simple catarrh is very common. **Mucous patches** and **Condylomata** are tolerably frequent, especially on the pillars of the fauces and on the soft palate. In the tertiary stage, with or without the formation of gummata, there may be **Ulceration** of the mucous membrane. Not infrequently the ulceration extends very deeply and causes destruction of the uvula and palate. It may extend to the epiglottis and so further. It is frequently accompanied by considerable thickening and cicatricial contraction, so that it may lead to great deformity and sometimes to stenosis of the pharynx.

7. **Tubercular ulcers.**—These are of very rare occurrence here, although so common in the larynx. The disease is associated with phthisis pulmonalis, and is really due to an extension upwards of tubercular ulceration of the larynx. It occurs in the form of superficial ulcers, originally of a circular form.

8. The **Tumours of the Fauces** are not so different from those of the mouth as to call for very special remark, and they are altogether of much less frequent occurrence. We meet with papillary excrescences, cysts, sarcomas, and epitheliomas.

### C.—THE ŒSOPHAGUS.

The diseases of this part of the alimentary canal are important from a practical point of view, chiefly because of the natural narrowness of the tube. The mucous membrane of the œsophagus is covered by a thick layer of stratified flat epithelium, and the mucous glands are comparatively few.

1. **Dilatation of the œsophagus.**—Of this somewhat frequent condition two forms may be distinguished.

(a) **General dilatation** of the tube is a result of obstruction at any part of its course. The obstruction is mostly low down in the œsophagus and may be even at the cardiac end of the stomach. With the dilatation in these cases there may be considerable thickening of the muscular coat, although this is not always present. There is also sometimes a general dilatation without obstruction, and probably due to diminution of contractility in the muscular coat.



(b) **Partial dilatation or Diverticulum** occurs in two principal forms, according as the force causing the dilatation is pressure from within the tube or traction from without.

The **Diverticulum by pressure** from within (*Pulsion-diverticulum*) may sometimes owe its origin to a slight congenital pouching of the tube. This may be due to an imperfect closure of the communication between the Œsophagus and the trachea during foetal life. If this closure be delayed, then the wall of the Œsophagus may remain bulged out in front and from this beginning a larger diverticulum may arise.

More commonly, however, diverticula form on the posterior wall and usually at the junction of pharynx and Œsophagus. It is probable that for the most part this form originates by a piece of hard food lodging in a fold of the mucous membrane, and being gradually pushed outwards, carrying the wall of the tube with it. As the sac enlarges, it hangs downwards with its mouth presenting upwards, and so it is always ready to receive the food in its passage downwards. It is probable that in most of these cases there is a separation of the fibres of the muscular coat of the Œsophagus or pharynx, and that the mucous membrane is pushed, as it were, through the muscular coat, which is thinnest at the junction of pharynx and Œsophagus. If the sac is of any considerable size and filled with food, it will press on the Œsophagus, and of itself produce an obstruction. In this way the food is prevented passing down the Œsophagus and goes readily into the sac. We have therefore, for the most part, a continually increasing enlargement of the diverticulum which may reach the dimensions of a child's head or larger. It contains the remains of the decomposing food with mucus, which is sometimes present in considerable masses. If the diverticulum is moderate in size its wall may still contain some muscular fibre, although the muscular coat is mostly absent except just at the neck.

The **Traction diverticulum** arises by cicatricial contraction of structures adherent to the wall of the Œsophagus. The contracting structure is mostly a lymphatic gland at the root of the lung, which, in connection usually with phthisis, has, after enlarging, undergone softening and cicatricial contraction. By dragging on the wall of the Œsophagus a funnel-shaped pouch is formed, whose apex is usually composed of cicatricial tissue.

This form of diverticulum is peculiarly prone to **Perforation** at the apex if hard food impinges on it. This leads to acute inflammation in the neighbourhood, resulting in an abscess-like cavity, which may perforate the trachea or a bronchus, or the neighbouring pleura or pericardium. In the former case there will probably be gangrene of the lungs, and in the latter acute inflammation of the serous membrane.



2. **Obstruction of the Œsophagus** is sometimes **Congenital**. There may be a congenital deficiency in the middle part of the tube, a fibrous cord representing the occluded tube. In some cases the Œsophagus below the occlusion opens into the trachea. Obstruction also arises as a consequence of pressure from without, by tumours, aneurysms, abscesses, but is still more frequent from disease in the tube itself. Ulcers of various kinds, by the cicatricial contraction incident to attempts at healing, may induce obstruction, as, for example, ulcers from swallowing strong acids, syphilitic ulcers, etc. Still more frequent are obstructions from tumours of the Œsophagus, and especially **Cancers** (see further on). As mentioned above, obstructions of the Œsophagus frequently give rise to dilatations with hypertrophy of the muscular coat above the seat of constriction. Sometimes the dilatation is greater in one locality so that with a general dilatation there is a partial diverticulum.

3. **Inflammations of the Œsophagus.**—The mucous membrane of the Œsophagus is formed so as to resist the action of irritants, and unless the action be peculiarly strong we have not considerable inflammation. When strong acids or alkalies, or substances at a high temperature, are swallowed, they may cause superficial necrosis and considerable inflammation of the mucous membrane. The small-pox eruption may extend into the Œsophagus, producing inflammation there.

**Strong acids or Alkalies**, when swallowed, pass rapidly through the Œsophagus, and so produce much less serious results than they do in the stomach. Acids produce whitish, yellowish, or brownish sloughs of the epithelium, sometimes penetrating to the mucous membrane itself. Caustic alkalies, on the other hand, dissolve the epithelium, producing a greyish gelatinous material which lies on the surface. If they penetrate to the mucous membrane they reduce it to a soft, brownish, half-diffuse substance. If recovery occurs, there are at first ulcers with more or less violent inflammation, and afterwards healing with contraction of the ulcers and possibly considerable narrowing of the tube.

Small-pox pustules in the Œsophagus are very similar to those in the

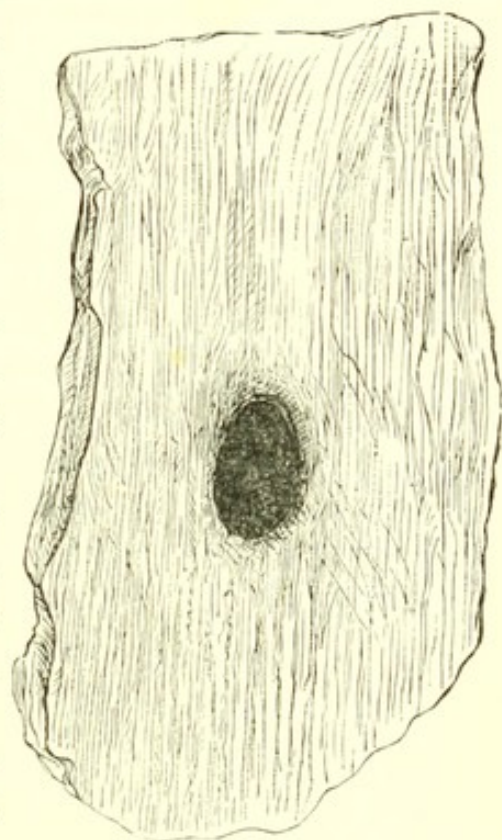


Fig. 277.—Perforating ulcer in lower part of Œsophagus. The ulcer penetrated the main bronchus of the left lung.



mouth, and, like them, they readily lose their epithelial covering and become converted into ulcers. They are accompanied by signs of inflammation of the mucous membrane generally.

4. **Rupture and Perforation of the œsophagus.**—The œsophagus may be directly ruptured by hard or sharp bodies being swallowed with the food. The author observed a case in which a fish-bone cut through the œsophagus and on into the aorta.

A more frequent cause of perforation is disease of the wall. The **Traction diverticulum** is, according to Zenker, the commonest cause of perforation. Next to that comes **Cancer** of the œsophagus, the actual perforation here being often produced by the sound.

**Perforating ulcer** sometimes occurs in the œsophagus, its causation being similar to that of the stomach (which see). Fig. 277 represents such an ulcer, which perforated into the left main bronchus, causing gangrene of the lung on account of food passing in. There have been cases in which the ulcer has penetrated into the aorta.

5. **Tumours of the œsophagus.**—Simple tissue tumours are rare. We meet with **Lipomas** and **Fibromas**, and the author has described a case of **Myoma** (Fig. 278) in which a tumour,  $4\frac{3}{4}$  inches long and 2 inches in thickness, was attached by a comparatively narrow neck, and produced death by obstructing the tube. Polypoid tumours of a similar form are met with, having a fibrous structure.

**Cancer** of the œsophagus is by far the most important form of tumour met with. The cancer is nearly always in the form of flat-celled epithelioma, and in its histological details closely conforms to cancer of the lip. The masses of epithelial cells infiltrate the spaces in the underlying connective tissue, and the tumour also projects somewhat into the calibre of the tube. Here also there is a great tendency to ulceration, the mechanical action of the food in swallowing doubtless contributing to this result.

The tumour begins at a limited part of the mucous membrane, but it has a special tendency to extend round the tube in the form of a ring. There has been considerable discussion as to the most common situation of the tumour, and the result of the comparison of various

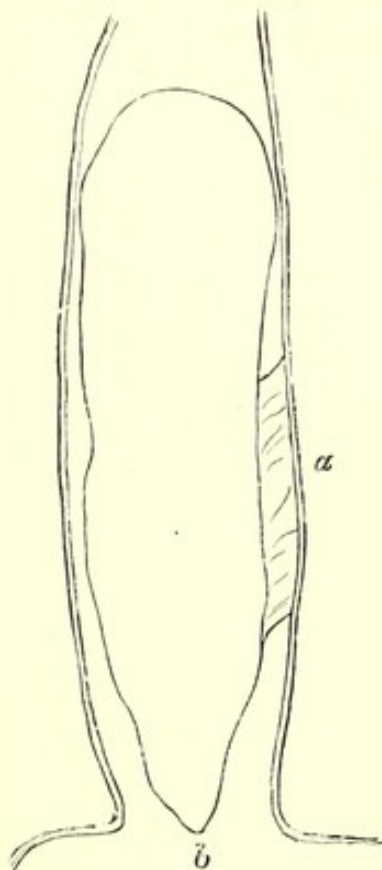


Fig. 278.—Polypoid myoma of œsophagus in section, and semi-diagrammatic. The tumour was attached by a band-like pedicle *a*. It caused considerable distension of the œsophagus, and obstructed the tube. *b*, cardiac orifice of stomach. Half the natural size.



tistics seems to be that the most frequent seat is the lower third. Scarcely less frequent than this is the middle third, and especially the place corresponding with the bifurcation of the trachea. In the upper third epithelioma is comparatively infrequent.

This form of tumour, when it surrounds the tube, frequently leads to **obstruction of the œsophagus**. On examining the œsophagus after death the seat of the tumour is often indicated externally by a narrowing of the tube, which is also more rigid here than elsewhere. The infiltration of the walls of the œsophagus, by irritating the connective tissue, causes a chronic inflammation with the usual newformation of connective tissue, which contracts and narrows the tube. Besides this, the mere rigidity of the infiltrated tube, preventing its dilatation when the morsel is being swallowed, may produce a virtual obstruction at the point concerned. The projection of the tumour into the calibre is another element, which tells especially in the earlier periods. But as ulceration occurs, this projection of the tumour usually becomes inconsiderable, and there may even be a temporary relief to the stricture by partial destruction of the tumour. The ulceration itself, however, by inducing still further cicatricial contraction, may ultimately confirm the obstruction.

The tumour sometimes extends from the œsophagus to neighbouring lymphatic glands, or to surrounding structures, and so we may have the trachea, bronchi, or lungs involved in it. Sometimes it extends to the diaphragm, pericardium, vertebræ, etc.

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#### D.—THE STOMACH.

**Introduction.**—In considering the diseases of the stomach, it is necessary to bear in mind certain points in regard to its structure and functions. The innumerable glands which exist in the mucous membrane are engaged in the formation of the **Gastric juice** for the purposes of digestion. The diseases of the stomach very readily interfere with the function of these glands, so that the secretion of gastric juice is insufficient in quantity or deteriorated in quality. In that case the



**Food** is apt to lie in the stomach undigested, and it very commonly undergoes various forms of **Decomposition**. The products of decomposition acting on the mucous membrane still further interfere with the process of secretion, and by their irritative action keep up or induce a condition of catarrh of the stomach. Gases also develop in the process of decomposition, and these, by distending the organ, interfere with its proper peristaltic movements, and so hinder the passage of the food through the pylorus.

But it is not merely local diseases, such as inflammations and tumours, which interfere with the secretion of the gastric juice. Changes characterized by **Cloudy swelling** and **Fatty degeneration** are met with in the glandular epithelium in numerous infective diseases, small-pox, typhus, septicæmia, as also in poisoning with phosphorus, etc. In these cases the secretion of the gastric juice may be almost at a standstill, and if food be introduced it is apt to lie in the stomach; by its mere presence, and by the action of the products of decomposition, it may produce still further structural changes.

The forms of decomposition which the food undergoes when it lies undigested in the stomach are various, but the chief are those characterized by the development of **Acetic, Lactic, and Butyric acids**. The agents in these changes are microbes which are introduced in abundance with the food and propagate in the stomach, unless their multiplication is hindered by the action of the gastric juice. In the contents of the stomach, when at any time they are discharged during life, swarms of bacteria are found, and in addition there are nearly always proliferating spores of fungi and sometimes large numbers of sarcinae.

It has been mentioned that the products of decomposition irritate the mucous membrane, but, in addition to that, microbes have been found in some cases to enter the glands and mucous membrane. In some of the cases the action of such microbes has produced little **Pustules** in the mucous membrane, or even larger prominences with necrosis and ulceration. It may be inferred that it is microbes of a special kind or in a peculiar state of activity which thus penetrate into the mucous membrane. Stagnation of the food in the stomach and its decomposition will occur still more when, by stricture of the pylorus, there is a **Mechanical obstacle** to the passage from the stomach.

Besides the mere local effects induced by the products of decomposition, it is to be remembered that the mucous membrane of the stomach is actively engaged in **Absorption**, and that these products may to some extent be taken into the blood. It is probable that much of the headache and other nervous symptoms occurring in dyspepsia is



due to the existence in the blood of small quantities of these poisonous agents and their action on the nervous system.

When the food remains in the stomach and accumulates there, it is usually got rid of after a time by **Vomiting**. This action is produced by irritation of a centre in the medulla oblongata, and the muscles employed are mainly those used in respiration, but in different combinations. The centre may be irritated directly, as by introducing apomorphia or tartar emetic into the blood, or by disease of the brain itself. But in the case we are considering it is irritated by reflex stimulation, the stimulation taking origin in centripetal fibres in the stomach itself. The occurrence of vomiting is dependent on the nature and amount of the irritation applied and on the sensitiveness of the centre in the individual.

**Post-mortem changes.**—After death, any remains of food lying in the stomach are apt to decompose rapidly, especially as the gastric juice in most cases is not secreted in the normal way up to the period of death, and so the decomposition is not interfered with. The decomposing juices therefore act readily on the mucous membrane, and the decomposition may even extend to the latter. The principal changes produced are **Alterations in colour**, resulting from chemical changes in the colouring matter of the blood. This may become diffused out of the blood-vessels and stain the mucous membrane of a generally red hue, the colour being specially pronounced in the neighbourhood of the larger vessels. There is often a greenish colour developed by the decomposition of the blood. Lastly, the colour may be almost black or slaty, but in many cases this deep colour is not altogether post-mortem, depending in part on a true pigmentation from chronic catarrh of the mucous membrane.

**Softening of the stomach or Gastromalacia** is also a post-mortem change. It is really a digestion of the coats of the stomach by the gastric juice. As a rule in persons near death the gastric juice is not secreted normally, but if the person die while the gastric juice is still active, then the latter may, by a process of **Digestion**, act on the coats of the stomach. This condition occurs mostly in persons who die suddenly, especially if the body is kept in a warm place, and it is more frequent in children than in adults. In the slightest degree, the mucous membrane alone is softened, and it can be removed as a soft paste from the surface of the muscular coat with the finger. Penetrating deeper, the muscular coat and even the serous coat may be half liquefied, so that on handling the stomach it may be perforated. The stomach may even **Rupture** in the body, and the contents pass outwards producing softening in neighbouring parts. In some cases the diaphragm has been softened in this way, and the stomach contents have passed partly into the pleural cavity.

These various changes occur in those parts of the stomach where the contents have been lying after death. This is generally the neighbourhood of the fundus. As the contents are usually fluid, it is often seen that the changes stop short at a definite level and the unaltered mucous membrane is abruptly demarcated from the altered part. The pyloric portion of the stomach, as it usually lies highest, is least frequently affected, and this is important, as that part of the stomach is the most frequent seat of disease.



It will be understood that many of the pathological changes of the stomach will be obscured by the occurrence of these post-mortem changes.

**Literature.**—LEUBE, in Ziemssen's *Encycl.*, vii., 1877; KUSSMAUL, *D. Arch. f. klin. Med.*, vi.; BRUNTON, *Disorders of digestion*, 1886. *Digestion of stomach*—HUNTER, *Phil. trans.*, 1772, and *Works by Palmer*, iv. 116; BURNS, *Med. and Surg. Jour.*, 1810; BAMBERGER, *Kr. d. chyl. Syst.*, 1855; VIRCHOW, *Würzb. Verhandl.*, 1850; REEVES, *Softening of stom. in children and adults*, 1867.

## I.—MALFORMATIONS AND CONTRACTIONS OF THE STOMACH.

1. **Congenital malformations** of the stomach are not of great consequence. There may be abnormal smallness either with or without other more general malformations. Sometimes an **Hour-glass form** is presented on account of the middle part of the stomach being contracted. But this malformation may be acquired by cicatricial contraction.

Another occasional malformation is **Atresia** of the stomach at the pylorus. The stomach has a blind end, and there is no open connection with the duodenum.

2. **Contractions of the stomach** may be general or local. A **General contraction** is produced when the stomach is long deprived of food. This is most directly produced by obstruction of the œsophagus or cardiac orifice, but also occurs when, for any other reason, food is not taken. There is also a general shrinking sometimes as a consequence of scirrhus cancer, and a similar shrinking may be produced by peritonitis which has led to thickening and contraction of the peritoneum generally.

**Partial contractions** are usually the result of cicatrization of ulcers. As these are mostly on the lesser curvature, the two orifices may be drawn close together. Sometimes an hour-glass contraction may be produced in this way. There is also occasionally an hour-glass contraction visible post mortem, which depends merely on an irregular contraction of the muscular coat and is of no special significance. Cancer of the stomach may produce partial contraction.

**Literature.**—WIDERHÖFER, *Handb. d. Kinderkrankh.*, iv., 1880; MAYER, (*Atresia of pylorus*) *Virch. Arch.*, cii., 1885; CARRINGTON, (*Hour-glass contr.*) *Path. trans.*, xxxiii., 1882; HUDSON, *ibid.*, xxxviii., 1887.

## II.—DILATATION AND HYPERTROPHY OF THE STOMACH.

These conditions mostly result from obstruction to the passage of food through the pylorus from contraction of that orifice. A simple weakness of the muscular coat may also allow of passive distension. **Obstruction of the pylorus**, causing an accumulation of the contents, leads in the first instance to a **simple distension** of the organ, which tells chiefly on the parts which are free to swell out. The lesser curvature is



fixed by its attachments, and it usually retains nearly its normal position except that its middle part is somewhat dragged downwards. The greater curvature, on the other hand, is carried downwards, and the stomach may virtually fill the entire abdomen, reaching as far as the symphysis pubis in some cases. Sometimes the pylorus is depressed and the duodenum correspondingly displaced.

The general result of obstruction of an orifice is compensatory **hypertrophy of the muscular coat** of the viscus, such as frequently develops in the heart and urinary bladder. But the muscular coat of the stomach has a somewhat different function to that of the heart or bladder. In the latter there is a simultaneous contraction of the entire muscle with a view to the emptying of the viscus. In the stomach, however, the contraction is vermicular, and its object is as much to move the contents about inside the stomach as to empty them into the duodenum. In the actual propulsion of the contents into the duodenum it is the pyloric portion of the stomach that is engaged, and here also the material is carried forward by a vermicular movement. Hence the hypertrophy of the muscular coat in obstruction of the pylorus does not occur uniformly in the stomach, but **localizes itself in the pyloric portion**, sometimes even with a special thickening just at the orifice, forming a **tight Sphincter**. In these cases, when the wall of the stomach is divided, the progressive thickening of the rigid muscular coat can often be distinguished as the pylorus is approached. As the muscle of the stomach is in bundles, hypertrophy produces an exaggeration of these, and on section they are frequently very prominent, especially as the connective tissue septa between them are also hypertrophied. The alternating bundles, as seen on section, have been compared with the leaves of a fan.

**Literature.**—PENZOLDT, *Magenerweiterung*, 1875; LEUBE, in *Ziemssen's Encycl.*, vii., 1877.

### III.—INFLAMMATIONS OF THE STOMACH. GASTRITIS.

**Acute inflammations** may be produced by the action of irritant poisons which have been swallowed, the inflammation here being accompanied by sloughing. **Phlegmonous inflammation** is rare in the stomach as compared with the frequency of phlegmonous angina of the fauces, or the dysenteric inflammation of the intestine. It is met with occasionally in severe infective diseases, and particularly in puerperal fever. There is great swelling and redness of the mucous membrane, terminating in a diffuse purulent infiltration of the wall of the stomach, chiefly in the submucous tissue. Sometimes abscesses have been formed in the wall of the stomach which have burst into its cavity.



Catarrh of the stomach, on the other hand, is a condition of very frequent occurrence and is met with in the acute and chronic forms.

**Acute catarrh.**—This is induced for the most part by the direct action of irritants on the mucous membrane, mainly by the use of irritating foods or drinks. When food remains in the stomach undigested it undergoes decomposition, and the irritating products may induce an acute catarrh. The mere prolonged stay of food in the stomach probably induces it, as when by exposure to cold the secretion of the gastric juice and the peristaltic action necessary to the process of digestion are interfered with.

It is seldom that the stomach can be examined after death in this condition, but from observations of Beaumont on his patient with a gastric fistula, as well as from experiments on animals, the appearances of the mucous membrane have been tolerably well made out. There is intense redness with swelling of the mucous membrane, which is covered by a layer of mucus or muco-pus, sometimes slightly mixed with blood. The appearances are most marked towards the pylorus, and sometimes confined to that region.

Under the microscope the blood-vessels in the mucous membrane, and especially in the submucous tissue, are found enormously distended, and the epithelial cells, both those of the surface and of the glands, enlarged and granular.

**Chronic catarrh.**—This often remains after one or more attacks of the acute form. It is also present in cases of passive hyperæmia of the stomach, which so frequently occurs in consequence of diseases of the heart and liver. Cancer of the stomach is also accompanied by chronic catarrh in most cases.

If the catarrh is prolonged there usually occurs a considerable new-formation of connective tissue, as in other chronic inflammations. There is thus a thickening which affects mucosa, submucosa, and muscular coat, and causes the surface of the mucous membrane to assume an irregularly folded or warty appearance, which has given rise to the designation *état mamelonné*. The mucous membrane also presents, in many cases, dark spots or a general deep slaty colour, from the presence of pigment granules in the tissue. The pigment is derived from the blood, and indicates the occurrence of hæmorrhages, probably by diapedesis. The increase of the connective tissue produces **Atrophy of the glands**, which are also considerably distorted.

It sometimes happens that the increase of connective tissue is specially great, and the wall of the stomach may be converted into a thick, hard, resistant structure. As all these processes occur mainly in the pyloric portion of the stomach, considerable **Narrowing of the orifice**



may result. The rigidity of the wall and the narrowing of the orifice induce more forcible muscular contractions, and the muscular coat therefore hypertrophies. The thickened and indurated condition may closely resemble scirrhus of the stomach, especially as the muscular coat is often hypertrophied in that disease also.

Occasionally **Mucous polypi** develop in connection with chronic catarrh, and these may develop into mucous cysts.

**Action of corrosives, caustics, and poisons.** **Strong acids and alkalies** acting on the stomach wall cause necrosis to a greater or less extent, and also produce changes in the resulting slough. As already noted the œsophagus is often but slightly involved. The intestine is frequently affected, sometimes as far down as the ileo-cæcal valve. The intensity of the action depends largely on the concentration of the acid or alkali. Sometimes the whole thickness of the stomach is dissolved, and, the contents having escaped, the action extends to the abdominal organs. The affected parts show various colours, acids generally producing a dark colour, while alkalies lead to a more tawny appearance.

If the patient survive, the sloughs will be discharged and there may be subsequently cicatricial contraction, leading sometimes to serious deformity and stenosis of the stomach.

**Carbolic acid** produces a dry stiff condition of the mucous membrane which has a brownish colour. This condition may also extend to the intestine. (See case in Western Infirmary Museum.)

**Arsenic** does not produce necrosis, but an irritation, evidenced by hyperæmia, and sometimes by ulceration. Decomposition is prevented by the presence of considerable quantities of arsenic.

**Literature.**—**ABERCROMBIE**, Researches on dis. of stomach, etc., 3rd ed., 1837; **LEUBE**, l. c.; **FENWICK**, Morbid states of stom., 1868, Atrophy of stom., 1880; **HABERSHON**, Observations on dis. of abdom., 3rd ed., 1878; **Fox**, in Reynold's Syst. of med., 1868; **BEAUMONT**, Expts. and observations on gastric juice, etc., 1833; **SILCOCK**, (Phlegmonous infl.) Path. trans., xxxiv., 1883.

#### IV.—THE SIMPLE PERFORATING ULCER.

This peculiar form of ulcer is met with only in the stomach, first part of the duodenum, and lower part of œsophagus. The duodenum is not an infrequent seat, the œsophagus a more unusual one. It is also called, sometimes; the **Round** and the **Chronic ulcer**. It is clear from the localities in which it occurs that its peculiarities are due to the action of the gastric juice.

The ulcer is usually round or oval in shape, and presents the appearance as if a conical piece of the wall of the stomach had been punched out from within, its edges being perfectly defined without any consider-



able thickening of the neighbouring mucous membrane, and the floor of the ulcer perfectly clean (see Fig. 277, p. 711). The superficial extent and depth of the ulcer vary considerably. The commonest size is about that of a shilling, but this may be exceeded considerably, and Cruveilhier has described an ulcer which was  $6\frac{1}{2}$  inches long and  $3\frac{1}{2}$  inches broad. In the smaller ulcers the floor is formed of the coats of the stomach, probably with some new-formed connective tissue. In the larger and deeper ones the tissue of neighbouring organs may be exposed, such as that of the pancreas or liver. The floor of the ulcer does not present any of the usual appearances of a granulating wound, but is clean and smooth, the actual tissue of the part being exposed, perhaps with some induration from newformation of connective tissue.

The situation of the ulcer is mostly in the neighbourhood of the lesser curvature, and nearer the pyloric than the cardiac orifice. It is more frequent on the posterior than the anterior wall. Although usually single it is not uncommon to find more than one ulcer present in the same case.

The ulcer presents a tendency to penetrate more and more deeply, from which circumstance it is named the **Perforating ulcer**. It does not appear to extend laterally to any considerable degree; it is probable indeed that at the very first the ulcer assumes its full superficial dimensions. Eating into the wall of the stomach, it may penetrate through the entire coats, and sundry accidents are liable to ensue.

One of the commonest of these accidents is **Hæmorrhage**. The ulcer penetrates one or more vessels at its base. The vessels may be small and the hæmorrhage not very considerable, but sometimes a considerable artery is laid open, and a serious, even a fatal hæmorrhage results. From the commoner situations of the ulcers the arteries most frequently penetrated are these—the coronary artery or one of its branches, the gastro-epiploic, the pancreatic, and the splenic. Sometimes the open mouth of the vessel can be seen after death in the floor of the ulcer.

**Perforation** is another result of the penetration of the ulcer. For the most part, by the time the ulcer has eaten through the wall of the stomach, the latter has already acquired adhesion to some neighbouring structure, and so actual rupture of the stomach and escape of its contents into the peritoneal cavity are not common. The adhesion may be to the liver or pancreas, or, more rarely, to the spleen, diaphragm, colon, abdominal wall. By the extension of the ulcer these structures may be eaten into and their tissue exposed. The tissue when first exposed has its normal appearance, but it usually becomes condensed and cicatricial. Sometimes the irritation of the gastric juice produces



suppuration and the formation of an **Abscess**, especially in the case of penetration into **the liver**.

**Rupture** of the stomach results if perforation occurs without previous adhesion of the wall. This will happen most readily where the wall of the stomach is liable to shift about during the regular peristaltic movements, and also where there is no solid viscus to which it may readily adhere. Both these conditions are fulfilled in the case of ulcers of the anterior wall, and so it happens that rupture most frequently occurs in this situation. The ulcers which lead to perforation are frequently very small, and the aperture in the serous coat may be as large as the ulcer itself. The result of the rupture is acute and fatal peritonitis.

The ulcer, when situated at or near the pylorus, may lead to partial **Obstruction of the pylorus**. This may be due to distortion of the parts and folding of the mucous membrane from shrinking of the ulcer. It will follow most readily in ulcers of the duodenum immediately beyond the sphincter. This situation is not uncommon, and the ulcer not infrequently partly involves the edge of the pylorus. In a case observed by the author the symptoms during life and even the appearances after death strongly suggested cancer of the pylorus.

**Healing** of the perforating ulcer is by no means an unusual occurrence. In the experiments to be referred to presently, in which ulcers were produced artificially in animals, they healed very readily. In man also they are frequently recovered from, and we often meet with cicatrices in the stomach. They may indeed heal without leaving a very obvious cicatrix at all. The author met with a case in which three weeks after a very severe hæmorrhage, presumably from an ulcer of the stomach, only an obscure cicatrix could be found. In order to healing, the acrid condition of the gastric juice, which seems to be the chief agent in their causation, must be corrected.

**The mode of origin** of these ulcers is a matter of some difficulty. It is obvious, from the shape and appearance of the ulcer, that it has arisen by the necrosis and subsequent digestion of a piece of the wall of the stomach. The funnel-shaped outline of the ulcer suggested to Virchow that the necrosis occurred by obstruction or interference with an arterial branch, and he observed as confirmatory of this that the ulcers most frequently had their seat at the point of entrance of arterial branches into the wall of the stomach. The experiments of Panum and Cohnheim confirm this view in so far as they show that ulcers may be produced by embolism of the arteries of the stomach. The perforating ulcer, however, is not met with specially in cases of embolism or thrombosis of the arteries of the stomach, but in the immense majority of instances in cases where no such disturbance of the circulation exists. It has been suggested again that a venous hyperæmia, by causing stag-



nation and even hæmorrhage (see afterwards) in defined areas of the mucous membrane, may produce such weakening of the tissue as to induce necrosis and digestion of it.

In most cases of gastric ulcer there is serious and usually prolonged **Dyspepsia**, and the persons are frequently anæmic. Some abnormal condition of the gastric juice, by virtue of which it is peculiarly irritating to the mucous membrane, seems to be an essential factor, while a weakened condition of the mucous membrane is also of consequence. In most cases the gastric juice is abnormally acid, and it has been thought that by neutralizing the natural alkalinity of the tissues it may lead to their necrosis.

On the whole it seems probable that an acrid gastric juice, taking advantage of any accidental stagnation in a defined area of the mucous membrane, may lead to its necrosis and the formation of the ulcer.

The frequency of ulcer of the stomach may be judged of from the fact that according to the results of post-mortem examinations it is said that there are ulcers or cicatrices in about one in twenty of the cases examined after death.

**Literature.**—BAILLIE, *Morb. Anat.*, 3rd ed., 1812; BRINTON, *Ulcer of stom.*, 1857; VIRCHOW, *Arch.* v.; MÜLLER, *Geschwür des Magens.*, 1860; PANUM, *Virch. Arch.*, xxv.; COHNHEIM, *Allg. Path.*, ii.; KLEBS, *Handb.*, i.; LEUBE, in *Ziemssen's Encycl.*, l.c.; HAUSER, *Das Magengeschwür, sein Vernarbungsproc.*, 1883.

#### V.—HYPERÆMIA AND HÆMORRHAGE.

**Passive hyperæmia** of the stomach is of very frequent occurrence, being brought about not only in that large class of cases in which there is general venous engorgement, but also in those in which a localized lesion in the liver obstructs the portal circulation. The mucous membrane is generally reddened in such cases, and there is usually some catarrh. There are usually also hæmorrhagic erosions visible (see below.)

**Severe vomiting** also leads to passive hyperæmia, apparently by the contraction of the muscular coat obstructing the veins. Here also there may be considerable hæmorrhage.

**Hæmorrhage** occurs under a considerable variety of circumstances. Ulcers, whether simple or cancerous, frequently cause it. It may result, as just mentioned, from passive hyperæmia, and it occurs, much more rarely, in scurvy, purpura, yellow fever, and typhus.

In the case of ulcers there may be large hæmorrhages from the rupture of considerable vessels. In passive hyperæmia there is, rather, a leakage from the superficial vessels of the mucous membrane, these being least supported, and the blood passes chiefly into the cavity of the stomach. At the same time there is some infiltration of the mucous



membrane in its superficial layers, and these parts being injured by the blood may be digested by the gastric juice. In this way arise small flat superficial ulcers, the so-called **Hæmorrhagic erosions**. These are generally present in considerable numbers, chiefly in the pyloric region. In the erosions there may be still some remains of blackened blood, and alongside them there are little areas of mucous membrane infiltrated with blood. In these cases also the mucous membrane is often generally red from the passive hyperæmia, and possibly thickened by catarrh.

The blood in whatever way arising is generally mixed with the contents of the stomach, and blackened by the gastric juice. If the hæmorrhage be very severe, as from an ulcer perforating a considerable artery, the blood may be vomited nearly in the fresh state, but usually it is tarry or like coffee-grounds. The altered blood will also pass into the duodenum and onwards.

Blood sometimes accumulates in the stomach when it has a different source, as when an aneurysm ruptures into the pharynx or œsophagus.

#### VI.—TUBERCULOSIS AND SYPHILIS OF THE STOMACH.

**Tubercular ulcers** are rare in the stomach, although very frequent in the intestine. They sometimes occur in cases of advanced phthisis pulmonalis. The ulcers are more superficial than those in the intestine, resembling rather those of the urinary bladder, although deeper than them. They have overhanging edges and granular floors.

The rarity of tubercular ulcers in the stomach as compared with the intestine may be due to two circumstances. In the first place the gastric juice will inhibit the tubercular bacilli in their passage through the stomach, and, in the second place, the stomach is devoid of the closed follicles which in the intestine are the primary seats of the tuberculosis. This fact may also account for the smaller size and more superficial character of the ulcers.

The author has only met with one case, and in it the patient had been in a state of extreme inanition for some weeks before death.

**Syphilis** is also excessively rare in the stomach, but Birch-Hirschfeld records two cases in his *Lehrbuch*, apparently of congenital origin.

**Literature.**—COATS, (Tuberculosis) *Glasg. Med. Jour.*, xxvi., 1886, p. 53; BIRCH-HIRSCHFELD, *Lehrb.*, 3rd ed., 1887, ii. 538.

#### VII.—TUMOURS OF THE STOMACH.

**Cancer.**—This is the only form of tumour which is of much practical importance, and it is of exceedingly frequent occurrence. From the statistics of a considerable number of observers it appears that



cancer occurs more frequently in the stomach than in any other situation in the body, the uterus being the next most frequent site.

The great frequency of cancer in the stomach is probably related to the fact that the epithelial structures of this organ are more exposed to various irritations than those of any other part of the body. Not only are there varieties of irritating foods, but the foods are liable, as we have seen, to decomposition, the products of which produce irritation. In cases of cancer there is very commonly a history of prolonged dyspepsia, perhaps from youth.

In this relation the simple ulcer may be again referred to. There have been cases observed in which cancer seemed to originate in the simple ulcer; but the simple ulcer is a disease mostly of youth, whereas cancer is a disease of middle life, the average age being fifty years. It is almost as if similar causes produced the simple ulcer in youth, and cancer in middle life.

In its **Structure and Mode of growth**, cancer of the stomach conforms to cancer elsewhere. It consists of epithelial masses contained in a stroma.

The epithelium originates from the glandular epithelium of the mucous membrane, and sometimes it retains to a large extent the glandular characters (epithelial cancer). In growing, the cancerous tissue first infiltrates the mucous membrane and submucous tissue, producing thickenings of them. It also insinuates itself amongst the muscular bundles, frequently separating these and replacing them. It is not uncommon to find in the muscular coat almost isolated outposts, the cancerous tissue having only a narrow connection with the primary tumour. The cancer penetrates through the muscular coat to the subserous tissue, but does not commonly involve the surface of the peritoneum except in the case of colloid cancer (see under).

In their growth the cancerous processes irritate the tissues, and there is usually a considerable infiltration of round cells. In the more chronic infiltrating forms, there is a great newformation of connective tissue, constituting scirrhus cancer.

As the cancer in its growth causes atrophy of the proper tissue, the wall of the stomach comes to be composed more or less of cancer tissue. This is less calculated to resist the disintegrating action of the food and gastric juice than the normal mucous membrane, and hence **Ulceration** is a very frequent result. This will occur readily in the softer and more superficial cancers, and as these are the commoner, it is usually a prominent feature.

The **Cancerous ulcer** (Fig. 279) is usually considerably excavated, and there may be pieces of slough in its floor. Its edges are prominent, sometimes overhanging, and they shade off into the normal mucous membrane. In its central parts the ulcer may penetrate deeply, sometimes through the coats of the stomach into neighbouring viscera, as



the liver or transverse colon. On the other hand, the floor of the ulcer may be partially cicatrized. In slow-growing epithelial cancers the ulcer may be very large and, with its overhanging edges, may occupy a large part of the stomach.

Cancer of the stomach usually begins in the **Pyloric region**, although by no means always. In extending it not infrequently takes the form of a **Ring** around the pyloric region, and by its prominence and sometimes by the contraction of the ulcer may lead to obstruction.

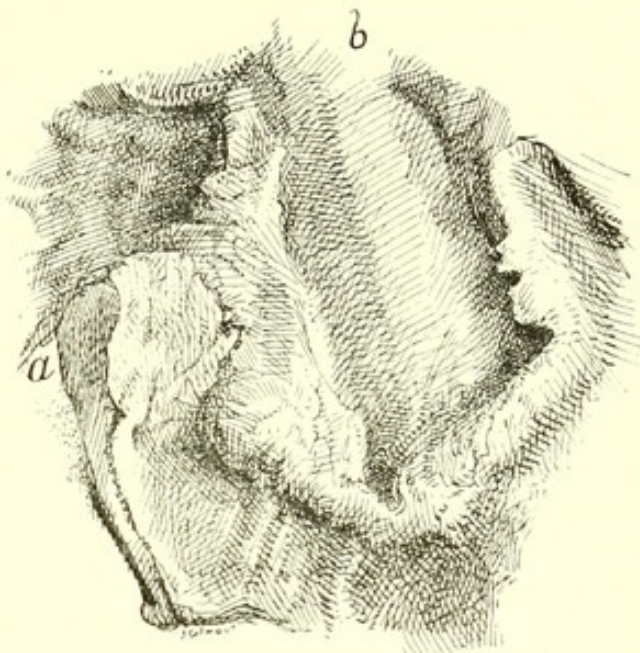


Fig. 279.—Cancerous ulcer of stomach. *a*, Pylorus; *b*, floor of ulcer, whose prominent walls are shown. Natural size.

The cancer affects surrounding parts both by **Irritation** and by **Extension** of the cancerous growth. The peritoneal surface is usually the seat of chronic inflammation, so that adhesions are present, sometimes producing great matting and entanglements around the stomach. The cancerous growth also sometimes extends by continuity into organs with which adhesions have been contracted, chiefly the liver and transverse colon.

The **Lymphatic glands** are usually affected, in the first place those immediately in connection with the stomach at the lesser and greater curvatures, but also the prevertebral glands. A very frequent extension is to the **Liver** (which see) and a less frequent one to the **Peritoneum** generally.

As the secondary extension of cancers of the stomach and intestine present many points in common, a special section is devoted to the subject further on.

The general characteristics of cancers of the stomach have been given above, and several forms have been referred to. It is possible to distinguish four different forms, which, however, are not absolutely separable.

1. **Cylinder-celled epithelioma** (*Adenoid cancer, Malignant adenoma*).—In this form there is a tolerably definite gland-like newformation, as we have in other cases of cylinder-celled epithelioma. There are spaces lined with cylindrical epithelium, but very often this regular arrangement is lost in great part, and, except in the more recently formed parts, we have more irregular masses.

The tumour is a slowly growing one, and it generally involves a considerable portion of the stomach before the death of the patient. The surface is nearly



always ulcerated, and may be considerably excavated. There are sometimes papillæ on the surface of the tumour, especially at the marginal parts, which give the surface a warty appearance.

2. **Medullary or Soft cancer.**—This form is closely allied to the preceding, but the cell masses are larger and less arranged in definite gland-like spaces, while there is a sparse and delicate stroma. The tissue is soft, and it is specially liable to bleed and to ulcerate. The latter condition is so frequent and characteristic that the tumour often presents itself as a round, shaggy ulcer with prominent edges. The bleeding may be slight but frequent, or it may be more considerable and even fatal. This form very commonly shows numerous large tumours in the liver.

3. **Scirrhus.**—In this form the newformation of epithelial cells is not very rapid or vigorous, and it is accompanied by an excessive formation of connective tissue in the form of stroma. Originating in the glandular epithelium, long processes composed of rows of cells, often with few abreast, are produced. These processes seem to be peculiarly irritating, as they give rise to the production of round cells and of dense connective tissue which are sometimes more manifest than the proper epithelial elements. These elements may, in fact, degenerate, and leave little besides condensed hard connective tissue.

The cancer mostly begins in the pyloric region, but extends inwards till, in some cases, it has involved the entire wall of the stomach, except the fundus. The wall of the stomach is converted into a stiff hard mass which may be, in some places, an inch in thickness. The surface of the affected portion of the stomach is irregular, with rounded prominences, and there are sometimes ulcers present, but there is not a general ulceration as in the case of the epithelial and medullary forms. The affected portion of the stomach is often greatly contracted in this disease, especially the pyloric region.

4. **Colloid cancer.**—The other forms, and especially the epithelioma, occasionally undergo a partial colloid degeneration, but in this form the cells have a special tendency from the first to undergo colloid metamorphosis, so that even in the more recent parts there is often already a considerable advance in the degeneration. The outlines of the cells disappear as the protoplasm becomes transformed into colloid material, and as the nuclei resist the degeneration longer, we sometimes see the peculiar appearance of oval nuclei as if floating in a clear transparent material. Finally, the whole epithelial elements are converted into colloid material, and the structure presented is a beautiful reticulated network with spaces filled with a transparent colourless jelly (Fig. 97, p. 274).

As the colloid material occupies more space than the original cells, the spaces of the alveoli are, as it were, tightly packed with the jelly, and the fibres of the stroma rendered tense and rigid. Hence, although the structure is composed mainly of a soft jelly, yet it is to the feeling hard and rigid, just as a tensely filled bladder may be.

The tumour, like other cancers, usually begins towards the pylorus, but it extends gradually till it comes to involve a large area, sometimes even as much as three fourths of the entire extent of the viscus. The wall of the stomach is converted into a transparent glancing tissue, and in the more advanced parts it is impossible any longer to distinguish the different coats, all being homogeneously replaced by the cancerous tissue. The wall of the stomach is considerably thickened, and the internal surface may present an irregular aspect with prominences; but there is little tendency to ulceration. As the thickened wall is tense and hard, the stomach when cut into does not generally collapse, but keeps its shape. There is



no tendency to contraction of the stomach as in scirrhus, but, on the contrary, the organ may be considerably enlarged.

This form has a very marked tendency to extend continuously both along the stomach and also through the stomach to the peritoneum. Hence it produces secondary tumours in the peritoneum itself much more readily than in the lymphatic glands and liver.

The remaining tumours of the stomach are of trivial consequence. We have already seen that **Mucous polypi** and **Cysts** occur in chronic catarrh. **Lipomas** and **Myomas** have been met with, as also **Fibromas** and **Sarcomas**, but they are very rare.

**Secondary cancer** scarcely ever occurs in the stomach. There may be an extension from the lower end of the œsophagus of flat-celled epithelioma, and a few cases of metastasis have been observed.

**Literature.**—*Myoma*—VIRCHOW, *Geschwülste*, iii. 126. *Sarcoma*—WICKHAM LEGG, *St. Barth. Hosp. Rep.*, x., 1874; HARDY, *Gaz. des. Hôp.*, 1878, p. 25; VIRCHOW, *Geschwülste*, ii. 325. *Cancer*—ROKITANSKY, (*Adenoma*) *Lehrb.*, iii. 155; HAUSER, *Magengeschwür*, *Bezieh. zur Entwick. des Carcinoms*, 1883; MOORE, (*Cancer in child*), *Path. trans.*, xxxvi., 1885; KÖSTER, *Die Entwick. der Carcinome*; EBSTEIN, *Volkmann's Vorträge*, 1875, No. 75; GRAWITZ, (*Metastasis of cancer, and literature*) *Virch. Arch.*, lxxxvi., 1881; COUPLAND, *Path. trans.*, xxvii., 1876, p. 264.

## E.—THE INTESTINES.

**Introduction.**—The diseases of the intestines resemble in many respects those of the stomach, but there are important differences. In structure the intestine differs from the stomach in several respects. We no longer have the specific glands peculiar to the stomach, but, on the other hand, the intestine presents numerous closed lymphatic follicles in its mucous membrane, and these are only present to a very slight extent in the stomach. These lymphatic follicles are solitary or collected into groups, in the latter case forming the well-known Peyer's patches.

After leaving the stomach the food passes rapidly through the upper part of the small intestine, occupying on an average two and a half to three hours in doing so, and it is at the same time rendered alkaline and partially protected from further decomposition by the pancreatic fluid and the bile. The movement of the intestinal contents is effected by the peristaltic contraction of the bowel, and the rapid passage of the contents through the small intestine indicates that here the peristalsis is peculiarly active, whereas, in the large intestine, it is slow. When the fæces reach the large intestine they are still fluid, and the chief function of the colon seems to be to complete the absorption of the fluid, and allow the fæces to become thicker. But if the peristaltic



action of the large intestine be increased, then there will be no time for the fæces to become thick, and fluid evacuations will be the result. This will be still more the case should the movement of the small intestine be increased, and the contents carried even more quickly than usual through it.

It will be seen that **Diarrhœa** results from increased peristaltic movement, and that the evacuations will be more fluid the higher up the increased movement begins. Certain medicinal agents produce fluid motions, and these seem to act generally by increasing the peristalsis, although some appear to produce their effects by causing a transudation of fluid into the canal (Hay). Irritating articles of food produce a like increase of the peristalsis and diarrhœa, and so may ulcers and inflammations of the intestine itself.

In the stools in diarrhœa we may expect to find chemical constituents which normally are present in higher parts of the intestine, but absorbed before reaching the rectum. If the diarrhœa arises from increased peristalsis of the colon, then we shall find material which is normal in the cæcum, such as undecomposed bile, leucin, chloride of sodium, peptones, and sugar, some of which are present in appreciable quantity in normal fæces. But if the diarrhœa has involved the small intestine, then we shall find these constituents much more abundantly, and also remains of undigested food.

We have already seen in the case of the stomach that many of its diseases are connected with the fact that the food stagnates and decomposes in that viscus. It will be seen from what has gone before that the **intestinal contents stagnate** chiefly in the **Large intestine**, and next to that in the lower part of the small intestine. It is probably due to this that we find the jejunum peculiarly free from all forms of disease; in this respect contrasting with the lower part of the small intestine, the ileum, but still more with the large intestine. Hence it is that the diseases of the large intestine resemble those of the stomach much more than those of the small intestine do. This is especially true in regard to simple inflammations, which very often are concentrated on those parts where the intestinal contents most readily stagnate, namely, the cæcum and the rectum. It is true also of cancer, which is very rare in the small intestine but common in the large, especially in the cæcum and rectum.

It is to be remembered further that the intestine is a comparatively narrow tube, and is subject to obstruction in various ways.

**Post-mortem changes.**—These are not so important as those of the stomach. After death the blood is apt to gravitate towards the more dependent parts of the wall of the intestine, and the colouring matter being dissolved out and staining the mucous



membrane, it may give rise to a deceptive appearance of inflammation. Similarly the intestine may be stained with the biliary colouring matter in the neighbourhood of the gall-bladder.

**Literature.**—COHNHEIM, *Allg. path.*, 1882, ii. 132; NOTHNAGEL, *Phys. u. Path. des Darmes*, 1884; HAY, *Jour. of Anat. and Phys.*, xvii., 1883; MARKWALD, *Virch. Arch.*, 1875, lxiv. 505.

## I.—MALFORMATIONS OF THE INTESTINE.

Congenital malformations of the intestine are of considerable frequency. The most important are those in which, from a fault of development, a part of the intestine is wanting. The colon or the rectum may be entirely absent at birth, being represented by a solid cord; or the rectum may be partially occluded, in some cases in its middle part, in others at its lower extremity. All these cases present the characters of **Imperforate anus**, but their gravity varies, the most hopeful from the surgical point of view being those in which only the extreme lower part of the rectum is defective, and the gut is separated from the anus only by a membrane (see p. 49).

There occurs also narrowness and defect of the small intestine, especially in the duodenum and lower end of the ileum. The whole intestine is sometimes deficient in length, having something like the form of the letter S instead of the usual convolutions. In such cases the absorption and digestion of food must be defective, but the persons may live on to old age.

The commonest malformation is **Meckel's diverticulum**. This consists in a finger-like projection from the intestine. It occurs in the ileum, about three or four feet above the ileo-cæcal valve in the adult, and about a foot above it in the new-born; it projects from the free convex border of the gut. It is from one to six inches long, possessing the same structure as the intestine, and communicating with the latter; it is narrower in its calibre, being of a diameter rather more than that of the finger. This diverticulum arises by the imperfect closure of the omphalo-mesenteric duct, and sometimes it is united to the umbilicus by a cord. Very rarely is the diverticulum continued to the umbilicus, and opens there, forming an **Umbilical fistula**.

Sometimes the diverticulum gets closed more or less completely at its orifice by a fold of mucous membrane or otherwise. In that case the accumulation of intestinal secretion in it may give rise to the formation of a cyst, the **Enterocystoma**.

**Literature.**—LEICHTENSTERN, in *Ziemssen's Encycl.*, vii., 1887; ORTH, *Path. Anat.*, 1887, i. 764; FITZ, (Omphalo-mesenteric remains, cysts, etc.) *Amer. Jour. of Med. Sc.*, July, 1884; ROTH, (Enterocystoma) *Virch. Arch.*, 1881, lxxxvi. 371.



## II.—EMBOLISM AND HÆMORRHAGE.

1. **Embolism of the mesenteric arteries.**—This condition has already been considered at p. 77. The embolism produces hæmorrhage and necrosis, the process being similar to that in the hæmorrhagic infarction. The hæmorrhage may be very considerable. If the patient survive, the slough after separation leaves an **Ulcer** (the *Embolic ulcer*, Parenski).

Very few cases of this kind have been examined post mortem, and in these it has been the superior mesenteric which has been plugged. There is, however, some reason to believe that embolism of the inferior mesenteric may have similar results. Both of these vessels have anastomosing communications, but they are insufficient to restore the circulation in the central parts of the area to which the vessels are distributed, although they do so at the periphery. Hence, the infarction is less in extent than the area of distribution. The subject has been very fully elucidated by the experiments of Litten.

2. **Hæmorrhage.**—Besides the rare form just mentioned, we have hæmorrhage resulting from various causes. Ulcers of various sorts lead to it, especially cancerous and typhoid. Passive hyperæmia is also not infrequently a cause, especially when it depends on obstruction of the portal system in the liver. In this case hæmorrhage is more liable to be from the large intestine than the small. Intussusception and hernia, by obstructing the vessels, may induce hæmorrhage by a local passive hyperæmia.

**Literature.**—LITTEN, Virch. Arch., lxiii., 1875; MOYES, (literature fully) Glasg. Med. Jour., xiv., 1880; GRAWITZ, Virch. Arch., cx., 1887; PARENSKI, Wien. Med. Jahrb., 1876, iii.

## III.—HERNIA OR RUPTURE.

True hernia consists in a protrusion of the intestine, omentum, or other abdominal organ into a sac formed by a prolongation of the peritoneum. The sac may project externally, or it may be contained within the abdomen, and so we may distinguish **External** and **Internal** hernias. The hernias, especially the external ones, are of so much importance in a surgical point of view that full descriptions are given in the surgical and anatomical text-books, and need not be repeated here, except in outline.

For the most part the sac is an entirely abnormal projection of the peritoneum. An exception to this occurs in the case of congenital inguinal hernia, in which the sac is formed by the persistence of a foetal condition. There is a partial exception also in the case of most internal hernias, where the sac usually arises by the exaggeration of an existing normal pouch.



**Causation of hernias.**—Hernias are usually ascribed to the abdominal contents being subjected to **undue pressure**. In severe muscular efforts, such as are involved in lifting heavy weights, the glottis is closed, and the muscles of expiration fix the chest and abdomen, the contents of the abdomen being subjected to severe pressure by the contraction of the muscles of the abdominal wall. If there is any part of the wall which is unduly weak a bulging outwards may occur here, and so give the starting point for the hernial protrusion.

In this connection the greater frequency of hernia on the right side may be noted. In violent exertions the right arm is usually more used than the left, and as the chest is bent over to the left side to counterbalance the strain on the right, the lower surface of the diaphragm faces more to the right and presses the viscera towards that side. It is clear that straining at stool or otherwise will also increase the pressure on the abdominal contents, and any excess will predispose to hernia.

The protrusion takes place where there is any **Weakness of the abdominal wall**. The external hernias occur at specially unsupported parts of the wall, while the internal ones have usually a pouch ready made as a starting point. The abdominal wall from its anatomical conformation is weak at certain points in every person, but there may be congenitally a special weakness, which in some cases seems to be hereditary. On the other hand when the abdominal contents are increased, as a result of tumours, fluid accumulation, or pregnancy, the stretched wall may be weakened. It may be so also from direct injury to the wall.

Another cause is sometimes assigned for the production of hernias, namely an abnormal elongation of the mesentery. It is supposed that such a lesion will allow the intestine to impinge unduly against the abdominal wall, especially at its lower parts.

**The hernial sac.**—The viscera nearly always push the peritoneum before them, and the proper sac is formed by the peritoneum, which shows a remarkable power of stretching. But there are cases of protrusion in which the aperture has been produced by actual rupture of the peritoneum, and in these cases the hernia may be devoid of a proper sac. These cases, however, of what may be called **False hernia**, are exceedingly rare, as an injury, although tearing the muscular wall and other tissues, will generally leave the elastic peritoneum uninjured and capable of protrusion.

In **Congenital hernias** the sac is formed of peritoneum, but there has been no actual protrusion. In congenital inguinal hernia the sac is formed by the tunica vaginalis, whose connection with the peritoneum has remained patent. In congenital umbilical hernia the peritoneum is prolonged into the umbilical cord (see p. 48).



The hernial sac usually **acquires adhesions** to the structures among which it is protruded, and it does so by a chronic inflammation. It very often happens also that the contents of the sac become adherent to its internal surface by inflammation, and in that case the hernia is irreducible.

**Forms of hernia.**—It is not necessary to enter fully into the individual forms of hernia, and of the external ones little more than an enumeration will suffice.

The **external hernias** are, (1) Inguinal hernia in the congenital and acquired forms, or, as otherwise divided, direct and oblique. (2) Femoral hernia. These two are by far the commonest forms. Of comparatively rare occurrence are, (3) Hernia of the sciatic notch; (4) Perineal hernia, protruded between the fibres of the levator ani; (5) Vaginal hernia; (6) Hernia of the foramen ovale; (7) Umbilical hernia, which is congenital or acquired, in the former case arising by protrusion into the dilated umbilical cord; (8) Abdominal hernia occurring in various parts of the abdomen, chiefly towards the edges of muscles, and arising by tearing of tendons or muscular fibres, hence frequently traumatic; its commonest situation is near the linea alba.

**Internal hernia** comes less frequently into sight, and the possibility of its existence is apt to be forgotten.

1. **Diaphragmatic hernia** is perhaps the commonest. There is a congenital form in which a sac is protruded through one of the normal apertures, or through a part of the diaphragm which by reason of defective development has given way. The protrusion is into the chest, and the sac may contain intestine, spleen, liver, stomach. There is also an acquired form, due nearly always to some injury to the diaphragm, and the hernia is frequently devoid of a proper peritoneal sac.

From a case recorded by Dr. Adams, it appears that a tumour growing against the diaphragm (in his case from the capsule of the spleen) may so weaken it as to lead to hernial protrusion. In diaphragmatic hernias from rupture, a large portion of the abdominal viscera may be protruded.

2. **Retroperitoneal hernia** includes cases in which the intestine passes into a pre-existing pouch in the peritoneum, greatly enlarging and filling it. The hernial sac hence lies behind the peritoneum of which it is an offset. There are three pouches in the peritoneum which are capable of giving rise to such hernias.

The *fossa jejunoduodenalis* is the most important. It exists just where the jejunum arises from the duodenum, and lies between the last part of the duodenum, which bounds it on the right, and the aorta, which bounds it on the left. The pouch was present, according to Waldeyer, in about 70 per cent. of the bodies which he has examined, and is generally large enough to admit the terminal phalanx of the



thumb. It is best seen when the jejunum and small intestine generally are raised and carried to the right, so that the origin of the mesentery may be exposed. The little pouch, if present, is then seen lying in the posterior wall of the abdomen with a sharp sickle-like margin. Sometimes a fold of the jejunum passes into this pouch, constituting a hernia. The pouch may be greatly enlarged by the protrusion of further portions of the intestine into it, and cases have been recorded in which the entire intestine has passed into the greatly distended sac.

The *fossa subcæcalis* has its seat between the folds of the meso-colon ascendens. Into this pouch the intestine is very rarely protruded, and the pouch itself only occurred in about 30 per cent. of the bodies examined by Waldeyer.

The *fossa intersigmoidea* is a pouch in the meso-colon of the sigmoid flexure lying between its two folds. The aperture is in the under layer. This is the commonest of these pouches, occurring in about 80 per cent. of the bodies, but from the position of the aperture it does not appear ever to become the seat of hernia.

**Contents of hernias.**—The parts protruded are usually the intestine, and, for the most part, the more moveable small intestine. Sometimes also the great omentum is carried into the sac. The urinary bladder, large intestine, or any part of the contents of the abdomen, may, under exceptional circumstances, pass into the sac.

As the contents of the sac are unduly exposed to pressure, stretching, and friction, there is apt to be a **Chronic peritonitis** set up in the wall, especially in old cases. This may unite the loops of intestine together. In the case of large hernias with wide necks there may be complete matting of the intestine so produced. It even happens that if the hernia be such as to allow successively of the descent of any part of the small intestine, the whole of the loops of the small intestine may be mutually adherent.

If the intestine be long retained the chronic inflammation may induce adhesion of the intestine to the internal wall of the sac, and the hernia becomes **Irreducible**. It may be irreducible from other causes, such as narrowness of the neck, protrusion of an excessive bulk of viscera, etc.

The mode of descent of the **Large intestine** merits special notice. The sigmoid flexure, being freely moveable, may be protruded just like the small intestine, and the cæcum or transverse colon may also sometimes pass into a sac in a similar fashion, but otherwise the large intestine having no mesentery and being only partially covered with peritoneum does not usually descend. When it does its descent is in some respects comparable with that of the testis in the fœtus. Before its descent the testis lies behind the peritoneum and is only partially covered by it. As it descends it remains with only a partial peritoneal covering, and even in the tunica vaginalis, after the sac has separated from the general peritoneum, the testis lies behind with its posterior aspect free of peritoneum. And so in a hernial sac, the cæcum may be carried down, but in its new position it remains only partially covered with peritoneum, and really forms as it were a part of the wall of the sac. This will only occur in very large hernias as a rule, but when it does occur the piece of intestine will be irreducible.



A still more peculiar condition sometimes occurs. The intestine may be protruded mainly at the part where it is uncovered by peritoneum, and instead of pushing a peritoneal sac before it, it may as it were drag one after it. As the gut is protruded it may even get more and more stripped of peritoneum, so that the hernia may be much more extensive than the sac. This, however, is a very exceptional occurrence, and it is more common to find that as an ordinary hernia advances it drags the colon into it, so that besides free loops of small intestine there may be, fixed in the wall and only partly covered by peritoneum, a piece of the cæcum, or the sigmoid flexure, or even the fundus of the bladder.

In a similar fashion to that just described, the ovary may be protruded. A large majority of cases of **Ovarian hernia** are congenital, and they appear to arise by a fault of development by which the ovary descends as the testis does normally. The ovary passes through the inguinal ring and takes a sac with it, but just like the testis it is itself attached to the wall. The sac remains open like the tunica vaginalis in a congenital inguinal hernia. The ovary in that case will be irreducible, unless, as sometimes happens, the broad ligament is so long as to allow the ovary to pass back through the neck. In this case, however, the ovary will still have its fixed attachment in the sac. Apart from this congenital inguinal form, ovarian hernias may be acquired, and these may be either inguinal or femoral.

**Strangulation and Incarceration.**—These terms express a condition in which the contents of the sac are caught tightly at the neck, so that there is not only a hindrance to their return but an excessive pressure interfering with the circulation.

This mostly occurs when, on account of some peculiarity in the situation of the intestine as it issues from the sac, there is, to begin with, a partial obstruction. If the intestine at its entrance into the sac be free, while at its exit it makes a sudden bend so as to cause a partial obstruction, then the fæces will pass readily in, but will accumulate inside as they do not find free exit. The mere loading with fæces may cause irreducibility, and if the fæces decompose the development of gas may still further increase the bulk of the contents. In this way the sac will become too full, and as the neck is narrow there will be special constriction here. Again, the intestine already in the sac, by its peristaltic movement, may drag more and more of the gut after it, till the intestine may become impacted at the neck.

In any case the neck of the sac constricts the portion of intestine concerned, and the most direct effect is **Obstruction of its veins**. This itself, by producing hyperæmia, and, it may be, œdema of the mucous membrane, leads to swelling and further constriction. The whole protruded piece becomes of a dark colour from venous engorgement and hæmorrhage. Finally the pressure may be enough to close even the arteries.

The venous obstruction alone seems sufficient, if complete, to cause necrosis of the intestine, and so **Gangrene** is an occasional result. If



the arteries are also obstructed there is still greater probability of the occurrence of gangrene.

Sometimes the obstruction is relieved before gangrene has occurred, and yet in some of these cases a severe inflammation results after the intestine has been returned to the abdomen, leading on, it may be, ultimately to gangrene of the affected piece of gut. The probable explanation of this is that during the incarceration the blood-vessels have been so damaged that, on the restoration of the circulation, they are no longer able to recover. It has been shown by experiment (in the ear of the rabbit) that if, by ligaturing the main arteries, the vessels of the part are deprived of blood for a time, and then the circulation restored by loosing the ligature, the result is active hyperæmia, acute inflammation, or the hæmorrhagic infarction, according to the time during which the ligature has acted. So in the case before us, the release of the constriction, by allowing of the re-establishment of the circulation in vessels seriously compromised, may lead to inflammation or gangrene.

**Literature.**—For external hernias see surgical and anatomical works. Wood, in Ashurst's *Encycl. of Surg.*, 1885, v.; BOWDITCH, *Diaphragmatic hernia*, 1853; GARLICK, *Path. trans.*, 1878, xxix.; BAKER, (*Pericardial diaphragmatic*) *ibid.*, 1877, xxviii.; ADAMS, *Glasg. Med. Jour.*, 1880, xiv., p. 353; WALDEYER, *Virch. Arch.*, 1874, lx., p. 65; BALFOUR, *Edin. Med. Jour.*, 1869; EVE, *Lancet*, 1885; ENGLISCH, (*Ovarian hernia*) *Stricker's Med. Jahrbücher*, 1871, p. 335; ALBERT, (*Hernia inflammata*) *ibid.* 239.

#### IV.—TWISTING OF THE INTESTINE. VOLVULUS.

This is a condition of frequent occurrence, but one which is perhaps too little borne in mind as a cause of obstruction. It occurs in the great majority of cases at the **Sigmoid flexure** of the colon. The ascending colon above the flexure, and the rectum below it, have virtually no mesentery, being fixed to the abdominal wall. The flexure therefore is fixed at its two extremities and these are near one another, while the loop forming the flexure is moveable. It is as if the loop were attached by its two ends to a fixed point, and it is easy to understand how it should sometimes twist round this as an axis. The twist is, as it were, in two half turns (see Fig. 280), and it is usually the upper limb of the loop which turns round the lower at its neck.



The twisting causes some obstruction of the intestine, but this is not usually complete. Fæces still pass into the flexure, and they may accumulate in enormous quantity. This partial obstruction with accumulation of fæces may persist for months and lead to extreme dilatation of the flexure, so that it may fill the abdomen and reach up to the diaphragm. The walls of the intestine in these prolonged

Fig. 280.—Twisting of the sigmoid flexure. The upper limb has a much sharper turn than the lower, so that a flexible tube could be passed up from the rectum.



cases may be greatly thickened, especially the muscular coat. In some cases twisting may exist without any obstruction of consequence.

While twisting is most common in the sigmoid flexure, it is liable to occur also when other parts of the intestine assume similar relations, that is to say when a free loop of small intestine becomes fixed at its extremities, and these extremities are near each other. This happens most frequently when one extremity is normally fixed, as is the case at the upper and lower ends of the small intestine, where on the one hand the duodenum, and on the other the colon, is fixed to the abdominal wall and holds the intestine down. If, by inflammation or otherwise, an abnormal adhesion is acquired so that the gut is fixed at a point near the situation of the natural fixation, then twisting is apt to occur, and probably more readily here than in the sigmoid flexure, as the small intestine is naturally more mobile.

The author has recorded a case in which a peritonitis had fixed the small intestine about a foot above the ileo-cæcal valve, so as to bring about the conditions named, and twisting occurred in consequence. A fixation may also be brought about by a Meckel's diverticulum which has retained its connection with the umbilicus (Coup-land's case).

**Literature.**—COATS, *Glasg. Med. Jour.*, xiii., 1880, p. 445; COUPLAND, *Path. trans.*, xxxi., 1880, p. 144.

#### V.—INTUSSUSCEPTION AND PROLAPSE OF THE INTESTINE.

1. **Intussusception or Invagination.**—In this condition one portion of the intestine passes into another. In order that one piece may slip inside another, the one must present active peristaltic contractions, while the other is relaxed. The portion narrowed by the violent peristalsis passes inside of the relaxed part. This usually occurs in the natural direction of the peristalsis, but it may be reversed.

The condition is most frequent in children in whom the peristalsis is very active, and in whom also the large intestine is more freely moveable than in adults.

These conditions are most frequently satisfied at the junction of the small and large intestines. The large intestine is naturally wide, and its peristaltic movements are sluggish. If then the ileum at its last part presents peculiarly violent peristaltic contractions it may pass into the large intestine. The invagination, however, is not usually a simple inversion of the ileum into the colon; in most cases the ileo-cæcal valve is carried before the advancing ileum and forms the apex of the intruded piece. This implies that the cæcum itself is carried inwards and inverted, and the orifice of the vermiform appendage is sometimes to be found near the apex of the invagination. The invagination is not infrequently very extreme, and the invaginated part may be carried right on to the rectum, so that the apex may be felt per anum.

Besides this form we also meet with invagination of the large intes-



tine itself, one part into a succeeding part. It occurs but rarely in the small intestine.

In the bodies of children, especially those who have died from cerebral or intestinal affections, we frequently meet with a form of invagination which has produced no symptoms during life, and has really occurred just at the time of death. It is usually present in the small intestine, where a small inversion of one part into a succeeding one is found. The invagination is easily reduced by slight dragging, and there are none of the secondary changes visible, such as are to be mentioned immediately as following invagination. Just at death, or immediately after it (as may be frequently seen in animals), the intestine commonly shows violent peristaltic movements, but these are irregular, and it readily happens that a much contracted part passes inside a relaxed portion.

When a piece of intestine is invaginated, there are three tubes, the outside one in its natural position, forming the sheath, the internal one, which, although abnormal in position, runs in the usual direction, and the middle one joining the other two, and with its mucous surface directed outwards. There is frequently a more complicated condition than this, the intestine being tightly packed in several layers. The mesentery is carried in with the intestine, and by being dragged on at one side, it may give the gut a somewhat oblique direction.

The packing of intestine causes considerable interference with the circulation, and this is increased by the mesentery being partly included and its vessels pressed on. The obstruction to the calibre of the intestine is thus associated with hyperæmia, œdema, hæmorrhage, inflammation, and even gangrene, just as in an incarcerated hernia. The inflammation may lead to general peritonitis; or the separation of a gangrenous part may allow of the escape of the intestinal contents and lead to a fatal peritonitis. On the other hand, permanent adhesion may occur, between the sheath and the upper end of the included part, at the proximal extremity of the invagination, and, the inner and middle tubes becoming gangrenous, they may come away in whole or in part, as a slough, and the continuity of the intestine be restored with the loss of a piece. It is not usual for complete gangrene of the included tubes to occur, but what remains becomes adherent to the sheath, and, by contraction, gradually accommodates itself, and so the calibre is fully restored.

**2. Prolapse of the intestine.**—This condition connects itself naturally with invagination. It is the protrusion of the intestine outside the body through the anus or through an artificial anus. The commonest form is the ordinary *Prolapsus ani*. This only occurs when the sphincter is greatly relaxed by catarrh or by violent pressing at stool, but when it has frequently happened the sphincter atrophies and the prolapse occurs very readily.

There are, strictly speaking, two forms of *prolapsus ani*. In the one



little more than the mucous membrane is protruded, and it is seen to be continuous with the skin at the anus. In the other form there is really an invagination as well as a protrusion. The lower part of the rectum is so fixed that the whole wall cannot be protruded, but only its mucous membrane as in the form just considered. But the upper, more moveable part, may be invaginated into the lower part and then protruded. This mostly occurs as a consequence of violent peristaltic contraction of the rectum in cases of severe diarrhœa with tenesmus.

In both forms the exposed mucous membrane becomes inflamed and is liable to bleeding. The inflammation sometimes causes through time adhesion and fixation of the bowel in its abnormal position.

#### VI.—INFLAMMATION IN AND AROUND THE INTESTINE. ENTERITIS.

We have already seen that the mucous membrane is frequently irritated by the contents being of an obnoxious nature by reason of decomposition or otherwise. In addition to that, inflammation may be produced by the action of specific morbid poisons, chiefly those of dysentery, typhoid fever, and cholera. We have therefore in the first place to consider the simple inflammations and afterwards these others.

1. **Catarrh.**—Catarrhal inflammations of the intestine are very common in children and can usually be traced to the character of the ingesta, perhaps along with exposure to cold. The common autumn diarrhœa, which is so fatal to children, is generally due to improper food. Cold, by interfering with digestion and peristaltic action may induce the intestinal contents to stagnate and decompose. The influence of the intestinal contents in producing catarrhs is shown by the localities at which inflammations are most common. The large intestine is much more frequently affected than the small, and in the large intestine the cæcum, with the vermiform appendage, is the most frequent seat, and next to that the rectum.

The catarrh may be **Acute** or **Chronic**. It is characterized by hyperæmia and swelling of the mucous membrane, with exudation of serous fluid and leucocytes. There is also increased secretion of mucus, which in the case of the colon may be very excessive. The exudation is usually **Mucous** in character, but in the more acute inflammations it may assume more or less of a **Purulent** character. If this be the case the mucous membrane is liable to be infiltrated with inflammatory cells and after a time to undergo **Ulceration**. This occurs with peculiar frequency in the cæcum, and next to it in the rectum.

**Catarrhal ulcers** are usually of considerable area and comparatively superficial. Neighbouring ulcers may coalesce so as to produce exten-



ive, variously shaped losses of substance, in the midst of which the remaining mucous membrane appears as raised patches with irregular outline. Sometimes the floor of the ulcer is so smooth, and the remaining mucous membrane so irregular from inflammatory infiltration, that it looks as if the ulcer were the normal mucous membrane and the patches of persisting mucous membrane adventitious. The ulcers of ordinary catarrh are superficial, and unless exposed to continued irritation, as by the prolonged presence of hard fæces, they do not tend to perforation, and readily heal when the cause of catarrh is removed.

In the course of catarrhs, ulcers also arise from the **Closed follicles**. There are indeed some catarrhs in which the latter are mainly affected, appearing as rounded prominences and flat elevations, corresponding to the solitary follicles and Peyer's patches. This form is called **Follicular enteritis**. The follicles may be so infiltrated with inflammatory products as to form virtually small abscesses resulting in crater-shaped ulcers which may afterwards enlarge. In this way arise the so-called **Follicular ulcers**.

A frequent result of catarrh is **Atrophy**. This occurs as part of the induration of the mucous membrane which follows chronic inflammation. The atrophy affects first the mucous membrane, its glands being specially involved, but it may extend to the submucosa and even to the muscular coat. According to Nothnagel, atrophy is of very frequent occurrence.

Another result is **Hypertrophy** of the mucous membrane such as we find so frequently in the case of the stomach. Here also there may be **Mucous polypi** and **Cysts** as in the stomach. They are most frequent in the large intestine, and if seated in the rectum they may project through the anus.

A condition of not infrequent occurrence is **Enteritis membranacea**. In this disease considerable pieces of membrane are passed at stool, sometimes in hollow cylinders as if casts of the intestine. These are composed of mucus, with epithelium entangled in them. They are generally regarded as the result of chronic catarrh of the large intestine, but Nothnagel asserts that inflammation is not necessary, and that it may be the result merely of defective peristaltic action allowing of accumulation of mucus in the folds of the large intestine. Hence Nothnagel suggests the name *Colica mucosa*.

**2. Phlegmonous and Diphtheritic inflammations.**—Some cases of catarrh have a more acute character, and assume the characters of suppurative or phlegmonous inflammation, the conditions approximating to those in severe dysentery (see further on). Such conditions are due to peculiar virulence in the intestinal contents, and may be brought about by irritant poisons such as corrosive sublimate. There may also be an acute inflammation in diphtheria. The relation of these conditions to



the intestinal contents is shown by the fact that they occur chiefly in the large intestine and in the parts where the contents are especially prone to lie, namely, cæcum, flexures, and rectum.

**3. Localized inflammations.**—It has already been indicated that certain localities are more liable to inflammation than others, and as the inflammations of certain of these regions present special points of importance they have received special names.

**Duodenitis** is usually an extension of a catarrh of the stomach, and it would not warrant any special reference except from the fact that the **Common bile duct** often takes part in the catarrh, and we have obstruction and **Icterus** sometimes resulting from this simple cause.

**Typhlitis and Perityphlitis** designate conditions which require very special notice. These two terms mean respectively inflammation in and around the cæcum, but they are frequently used so as to include inflammations in connection with the vermiform appendage.

In some cases the cæcum is very seriously inflamed and ulcerated, especially when large masses of hard fæces lodge in it. Under these circumstances the ulcers may increase in depth, and extend, in the posterior part of the wall where the gut is not covered by peritoneum, through the entire thickness of the intestine so as to penetrate into the retro-peritoneal connective tissue. The result is acute inflammation in this region, often going on to the formation of **Abscess**, a condition designated **Perityphlitis**.

The **Vermiform appendage** much more frequently gives rise to acute perityphlitis than the cæcum itself does. Inflammations are frequently caused by the presence of **Foreign bodies** in the appendage, especially in children. Any hard substance lying in the cæcum may, if small enough, pass into the vermiform appendage. In this way apple, grape, cherry, or orange seeds are said to get into it. But much more frequently pieces of hardened fæces are met with, and as these become dry and frequently assume the shape and appearance of cherry or orange stones, they are frequently mistaken for them. This occurs all the more because the inspissated fæces are often coated with phosphates which form a kind of rind.

The presence of the foreign body causes inflammation of the mucous membrane, and this, from the continued pressure and confined space, readily results in ulceration. The ulcers frequently penetrate through the appendage and lead to peritonitis. As a general rule the appendage has acquired adhesions before actual perforation occurs, and so the peritonitis is limited by the adhesions. But the inflammation is apt to recur, and not infrequently results in the formation of recurring abscesses. The bursting of one of these into the cavity of the peri-



toneum often leads to fatal peritonitis. Sometimes the suppuration in connection with the vermiform appendage is in the subperitoneal tissue. An abscess so produced may extend long distances beneath the peritoneum.

The author met with a case in which an abscess connected with the vermiform appendage extended across the brim of the pelvis to the left side of the abdomen, and thence upwards beneath the peritoneum till it reached the diaphragm. It then penetrated the diaphragm and discharged itself into the pleura.

**Proctitis and Periproctitis** are inflammations in the rectum and around it, of an equivalent character to those in and around the cæcum. If the ulceration leads to perforation and the formation of abscess, then the disease may have a very chronic course with fistulæ discharging into the rectum. Sometimes these open externally, or into the vagina, and they may retain their communication with the rectum, so forming false passages for the fæces and flatus; or the internal aperture may close, leaving merely an external fistula.

**Literature.**—*Catarrh*—NOTHNAGEL, *Phys. u. Path. des Darmes*, 1884; GOODHART, (Casts from intestine) *Path. trans.*, 1872, xxiii. 98; WOODWARD, *Med. and surg. hist. of war of rebellion*, Part. 2, vol. i., 1879; BAUR, (*Perityphlitis*) *Ziemssen's Encycl.*, viii., 1878; STEINER, *Path. Anat. d. Wurmfortsatzes*, 1882; CORNIL, *Arch. de Phys.*, iii., 1873; COATS, *Brit. Med. Jour.*, 1875, i.

## VII.—SPECIFIC INFLAMMATIONS OF THE INTESTINE.

1. **Dysentery.**—In this disease we have a violent inflammation evidently determined by the presence of an intense irritant. The disease occurs in a sporadic and an endemic form. In the former case it may be due to irritating foods of various sorts, while in the latter there is reason to suppose that a specific morbid poison is present. The morbid poison is presumably a form of microbe, and some authors have observed a minute bacillus in the mucous membrane (Ziegler).

The virus is introduced chiefly by the drinking water, but also by the air. In any case it acts on the mucous membrane directly. The locality of the lesion is apparently determined by the stagnation of the contents. It is essentially a disease of the large intestine, and is generally most intense in the rectum. It usually decreases in intensity from the rectum upwards, but not uniformly, there being more affected and less affected parts, the former corresponding usually with the flexures. In severe cases the whole colon is affected, and sometimes even the lower part of the ileum.

It is usual to divide the disease into a **Catarrhal** and a **Diphtheritic** form, but the morbid poison is the same in both, and the difference is really one of intensity. The catarrhal form may run into the diphtheritic. In the earlier stages the mucous membrane is swollen by serous exuda-



tion, soft and juicy, and it is thrown into folds, on the summits of which it is peculiarly hyperæmic. The surface is covered by a mucous or grumous material, consisting of shed epithelium with mucus and inflammatory exudation. The mucous membrane and submucous tissue are infiltrated with serous fluid and leucocytes in great abundance.

In higher degrees the mucous membrane is still more thickened and thrown into still more prominent folds. There is also considerable hæmorrhage in its substance occasionally. The summits of the folds being specially exposed to mechanical irritation very commonly undergo **Necrosis**, and the sloughs are generally distinguishable by the **Brown colour** which they assume from becoming stained with the bile pigment. The necrosis involves the mucous membrane to varying depths, sometimes very superficially, sometimes through its whole thickness, and, if the slough surrounds the gut, we may have a ring of necrosed tissue ultimately discharged by the anus.

These sloughs leave **Ulcers** behind, whose walls present great infiltration of leucocytes. But ulcers form also by processes similar to those in catarrh, namely, by inflammatory infiltration and molecular destruction of the mucous membrane, and by suppuration of and around the closed follicles.

In some cases the solitary follicles seem to be specially engaged and some authors even distinguish a follicular form of dysentery, but the follicles are probably affected in the early stages of almost all cases, and partake in the general inflammation. By suppuration of the follicles there may be the formation of numerous ulcers with small apertures.

The contents of the intestine are in severe cases formed of dark decomposing material, mixed with blood. The mesenteric glands are always secondarily affected, being enlarged and hyperæmic.

If the patient survives the acute attack, the disease very commonly passes into **Chronic dysentery**. The ulcers formed in the various ways described above show little tendency to heal, but remain as open discharging sores. Sometimes they penetrate more deeply, and lead to abscesses in the surrounding tissue, especially of the rectum (Periproctitis). The remaining mucous membrane is swollen and in a state of catarrh. The whole intestinal wall is thickened except where the ulcers exist, and it is irregularly drawn in and adherent to the surrounding parts.

In some cases a tendency to **Healing** manifests itself. If the attack has been slight and the ulceration only superficial, there may be a complete restoration of the mucous membrane, with insignificant cicatrices. But for the most part the cicatrices are of considerable superficial extent, perhaps surrounding the gut. The cicatrices of dysentery



have usually a dark, almost a black colour. When the cicatrices in the usual fashion contract, they cause narrowing of the intestine, and this, in some cases, is very considerable. Alternating with the narrowing there is very commonly dilatation of the more healthy parts, so that a very remarkable pouching of the colon may result, the narrowed parts being probably adherent to the tissues around. The obstruction produced in this way is all the greater because the contraction often throws the remaining mucous membrane into folds which may act as valves to the constricted part.

The symptoms of dysentery often continue, in the chronic form, after healing of all the ulcers has occurred. In that case along with the cicatricial contraction mentioned above, there is a very marked atrophy of the coats of the intestine, so that the wall is very thin and translucent. There will also be considerable adhesion to surrounding parts.

2. **Cholera.**—In addition to the epidemic disease known as **Asiatic cholera**, there is a condition known as **Cholera nostras**. The researches of Koch have demonstrated the connection between Asiatic cholera and the comma bacillus. It is probable that cholera nostras, and choleraic diarrhoea may be produced by microbes of allied, but not identical characters (see p. 302).

The phenomena of cholera are those of a very violent irritation of the mucous membrane of the intestine. During life the disease is characterized by the discharge of extremely fluid stools, which, from the intermixture of finely divided material, give the characteristic **Rice-water appearance**. Sometimes a patient dies without the bowels having been moved, but in that case they are found distended with the rice-water discharge. The flakes which give their character to the rice-water evacuations are not so much composed of desquamated epithelium as of little masses of mucus containing leucocytes. After death the epithelium may be found loose in the intestine, but this appears to be from post-mortem maceration, for, if examined sufficiently early, the epithelium is found present and apparently normal.

It is obvious that there is here an enormous transudation from the vessels of the intestine. The chemical character of the discharge, however, seems to indicate that it is not a mere inflammatory exudation from the vessels, but rather a secretion from the glands. Its specific gravity is low, 1006 to 1013. There is very little albumen present, and the discharge contains a ferment which has the power of converting starch into sugar.

The most marked post-mortem appearance is a remarkable rosy injection of the vessels in all the coats of the intestine, so that a red appearance is visible even in the serous coat whenever the body is opened



The mucous membrane is swollen and the closed follicles prominent. Occasionally there is a decidedly inflammatory condition present, which may be even of a dysenteric character.

The condition of the other parts of the body is directly referrible to the enormous withdrawal of water from the blood. The blood itself is thick, dark, and imperfectly coagulated. The skin, serous membranes, and all the soft tissue are shrunk, and dry, and parchment-like. The membranes of the brain are frequently injected. The kidneys present the characters of a slight parenchymatous inflammation.

**3. Typhoid fever.**—This disease is also due to the action of a specific microbe (see p. 300), and is characterized by a lesion in the intestine. The virus is sparsely present in the blood, and the general symptoms may be in great part secondary to the local lesion. The morbid poison finds entrance by the intestine, and produces irritation of the follicles and mesenteric glands in its passage.

The **Affection in the intestine** consists of an inflammatory swelling of the closed follicles and of the mucous membrane in their neighbourhood, the inflammation often going on to necrosis and sloughing. The disease has usually its point of greatest intensity at the lower extremity of the ileum, and it is often possible to see various stages of the process in proceeding from above downwards through the small intestine till it culminates close to the ileo-cæcal valve.

In the normal state, especially in the adult, the Peyer's patches and the solitary follicles are very slightly prominent. The patch viewed from the surface shows a congeries of shallow depressions separated by slightly elevated ridges, which run in from the general mucous membrane, and form a kind of network. It is in the depressions that the follicles are placed. In typhoid fever there is at first a general swelling of the patch; it is an inflammatory swelling accompanied by abundant infiltration of leucocytes which occupy both the closed follicles and the mucous membrane. The patch is obviously raised and its margins somewhat abrupt (Fig. 282, *a*). Viewed from the surface there is at this early stage simply an exaggeration of the normal appearance. The ridges of mucous membrane are swelled so that the depressions are more hidden, and an appearance is produced which has been compared to that of the convolutions of the brain in miniature. The swollen patch has a pinkish or whitish colour. At the same time the solitary follicles show themselves as elevations at intervals.

As time goes on the whole tissue gets more and more infiltrated with leucocytes (Fig. 281), and the raised patch gets more solid and its surface more homogeneous. The invasion of round cells extends to the submucous and muscular coats, even to the serous, and passes to some extent beyond the patch. A similar condition occurs in the solitary



follicles; they also present a marked increase in size, and are less defined, by reason of the infiltration around them.

On this condition follows **Necrosis** (Fig. 282 *b*). The infiltrated and

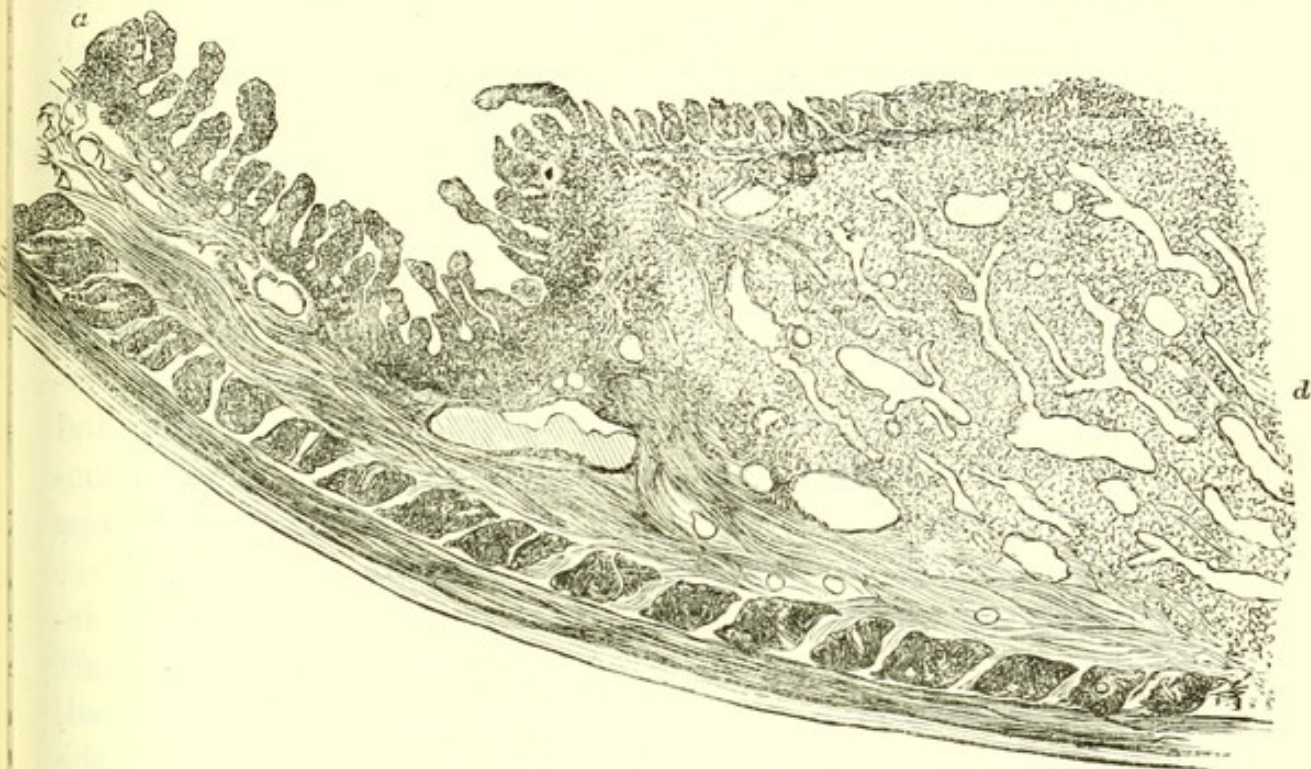


Fig. 281.—Portion of a Peyer's patch in an early period of typhoid fever; *a*, mucous membrane which becomes raised when the swollen patch is reached; *b*, internal layer, and *c*, external layer of muscular coat; *d*, swollen patch composed of round cells with dilated blood-vessels. At the right of the section the round cells are invading the submucous tissue and approaching the muscular coat.  $\times 16$ .

altered patch or solitary follicle forms a slough, of larger or smaller size. This slough remains adherent for a time, and like all sloughs in the intestine it becomes brown or yellow from the biliary colouring matter, which stains dead tissue, while the living structures are able to resist it. Generally there is a single slough on a Peyer's patch, not involving the whole patch, but of considerable superficial extent (see figure). Sometimes there are several sloughs corresponding to some of the closed follicles of which the patch is made up.

After a time the slough separates and an **Ulcer** is left with infiltrated base and margins (Fig. 282 *c*). The ulcers are confined to the patches and solitary follicles, so that they repeat their shape and locality. The walls are succulent and vascular, and considerable hæmorrhage may occur from them. The ulceration

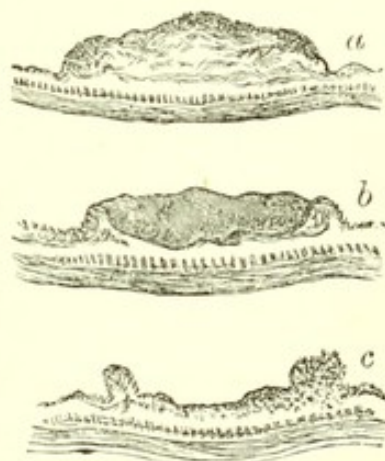


Fig. 282.—Diagrammatic representation of Peyer's patches in typhoid fever. *a*, early stage with swelling of the patch; *b*, later stage with sloughing; *c*, ulcer with infiltrated walls. (THIERFELDER.)



may extend more deeply than the mucous membrane, involving a necrosis of the muscular and even of the serous coat, so that **Perforation** may result. Perforation may occur in one of two ways. In the first place an ulcer may extend in depth till it penetrates through the muscular and serous coats. In this case the process is somewhat gradual, and there is generally an inflammation of the peritoneum sufficient to cause adhesion and prevent the intestinal contents passing into the abdominal cavity. In the second place, perforation sometimes occurs in a manner comparable to the perforation of the pleura in phthisis pulmonalis which leads to pneumothorax. The serous coat is undermined by the ulcer, and as its nutrition is cut off it undergoes necrosis. In such cases the brown slough may be visible on viewing the intestine externally. A partial separation of the slough may allow of perforation, and as this form is more acute and not so likely to be accompanied by considerable adhesions, the contents of the intestine are more apt to escape into the peritoneal cavity and produce fatal peritonitis.

When **Recovery** takes place from the fever, the process in the intestine retrogrades. According to the stage reached will be the exact process of resolution. If, in any part, ulceration has not yet occurred, then there is a gradual diminution of the patch and a return to the normal. If ulceration exists the ulcer fills up and cicatrizes. The **Cicatrix**, however, remains long, often with a slaty colour, but not usually with much puckering. It may indeed be a flat cicatrix in which the intestinal wall is merely thin and transparent. This is shown in

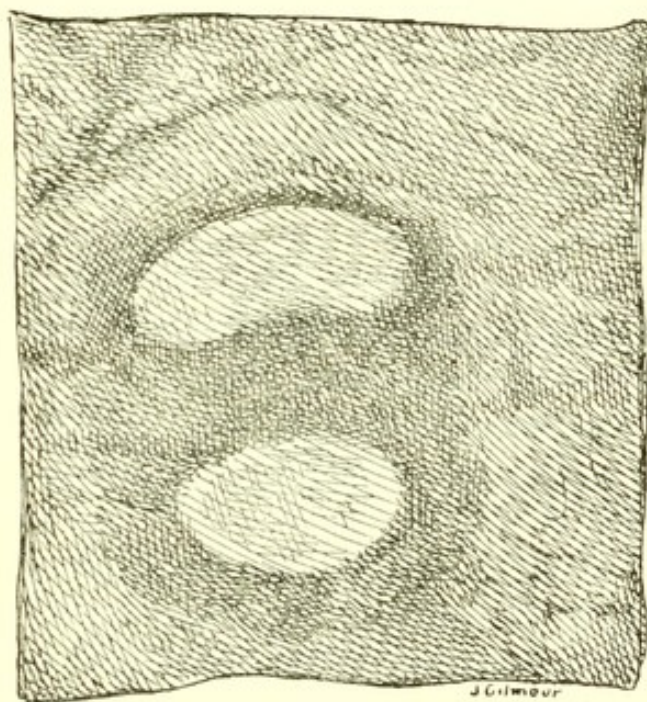


Fig. 283.—Flat cicatrices on a Peyer's patch from typhoid fever.

Fig. 283 which represents the conditions in the case of a girl who died of pneumonia a year after having passed through an attack of typhoid fever.

The **Lymphatic glands** of the mesentery enlarge, and present on section a red injected appearance and soft consistence. There may be necrosis here also, giving rise to an opaque greyish condition in the midst of the gland. When recovery takes place the glands return to the normal, and the sloughs are usually absorbed, although they may become caseous and subsequently calcareous.



It is to be remembered also that the **Spleen** is enlarged, and there is here more than in typhus fever a swelling of the Malpighian follicles, which are lymphatic in structure, and so the spleen is firmer and paler on section than normal (see p. 444).

4. **Anthrax** (*Mycosis of the intestine*).—The wall of the intestine is sometimes attacked by the anthrax bacillus. Several cases described as mycosis intestinalis belong to this group, although the name has also been used for other affections, such as septic inflammations and actinomycosis. The intestine may be the place of entrance of the bacilli, and the lesion will then be the primary one. On the other hand, the intestinal lesion may simply be part of the general infection, and concurrent with affections of other organs.

The lesion presents itself in patches situated most frequently in the small intestine, but also in stomach and colon. The patches are oedematous and infiltrated with blood, and there is generally a considerable dark slough on the prominences of the folds. The characteristic bacilli are found in the oedematous fluid and in the blood-vessels and lymphatics. The lymphatic glands are swollen, and infiltrated with blood. The bacilli are sometimes present in them in very large numbers.

Cases of this kind have mostly occurred amongst workmen who have handled the hides of animals affected with anthrax. Probably the bacilli are allowed to contaminate the food, and so reach the intestine.

5. **Actinomycosis**.—The intestine is rarely the primary seat of actinomycosis, but a case has been recorded by Chiari. The mucous membrane of the large intestine was the seat of raised whitish patches, in which the yellow granules of the parasite were visible. The mucous membrane around was swollen and red and covered with tough mucus.

**Literature.**—*Dysentery*—ANNESLEY, Diseases of India, 3rd ed., 1855; VIRCHOW, Virch. Arch., v. and lii.; HEUBNER, in Ziemssen's Cycl., i., 1875; ZIEGLER, Lehrb., ii.; MACLEAN, in Reynolds' Syst. of Med., i., 1866; BALY, Path. and treatment of dysentery, 1847; FAGGE, Medicine, ii. 175, 1886; WOODWARD, Med. and surg. hist. of war of rebellion, part ii., vol. i., 1879. *Cholera*—GOODEVE, in Reynolds' Syst. of med., i., 1866; LEWIS and CUNNINGHAM, Cholera, 1878; Report of cholera conference at Berlin, 1885, Brit. Med. Jour., 1885, i., 1011 and 1075, also separate Report of Conference; MACNAMARA, Hist. of Asiatic cholera, 1876. *Typhoid Fever*—LIEBERMEISTER, in Ziemssen's Cyclop., i., 1875; MURCHISON, Continued fevers, 2nd ed., 1873. *Anthrax*—BOLLINGER, in Ziemssen's Cycl., iii. 1875; VIERHUFF, Anthrax intest., 1885; WALDEYER, Virch. Arch., lii.; POLAND, Path. trans., xxxvii., 1886. *Actinomycosis*—ZEMANN, Wien. med. Jahrb., iv., 1883; CHIARI, Prag. med. Wochenschr., 1884.

## VIII.—TUBERCULOSIS AND SYPHILIS OF THE INTESTINE.

1. **Tuberculosis**.—In the great majority of cases this condition is second-



ary to pulmonary phthisis, and it occurs in about two thirds of the cases of that disease examined after death. It is to be accounted for by the the sputa being partly swallowed, and the virus applying itself to the intestine directly.

There seems little doubt also that tuberculosis of the intestine may be induced by the tubercular bacilli being present in the food. Experiments in animals show that this mode of infection is possible. It appears, however, that tubercular bacilli may be absorbed by the intestine without inducing tuberculosis of it. The mesenteric glands may become tubercular, or there may arise a tuberculosis of the peritoneum without any affection of the intestine. This result of experiment in animals is confirmed by observation in man.

The virus here attacks the same structures as in typhoid fever, namely, the closed follicles and Peyer's patches, and at first it produces a change of a somewhat similar kind. There is an enlargement of the closed follicles by reason of a great newformation of cells which infiltrate the neighbouring mucous membrane as well as the follicles. This primary enlargement is, however, much less uniform than in typhoid fever. On the whole, it is greatest towards the lower part of the ileum, but it presents great irregularities. Even on the Peyer's patch it generally affects a few closed follicles, and not the whole, so that there

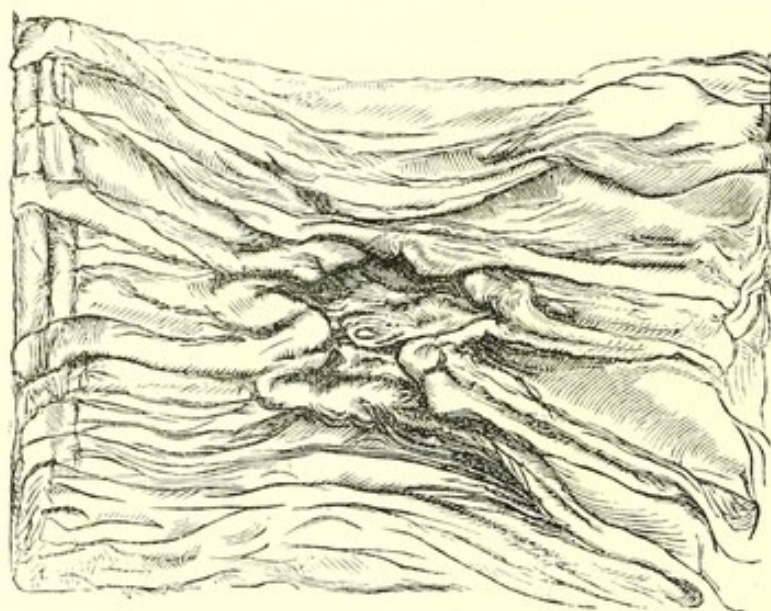


Fig. 284.—Tubercular ulcer of the intestine. Naked-eye appearance. The swollen overhanging edges are indicated.

are rounded prominences dotted over the patch. Moreover, it is not simply an inflammatory infiltration which occurs; there are tubercles and tubercular bacilli present from the first.

The enlarged follicles undergo caseous necrosis, so that very soon



they are seen to have an opaque yellow kernel, while the epithelium still covers them. In the next place softening occurs, and the caseous mass is discharged, leaving a crater-shaped ulcer. The ulcer advances and continues to have over-hanging edges, which are infiltrated with leucocytes and contain miliary tubercles (see Figs. 284 and 285). There may be at first several such ulcers on a Peyer's patch, and other solitary ones in other parts. They extend by fresh infiltration of the neighbourhood and fresh necrosis, while at the same time miliary tubercles are visible in the midst of the inflammatory infiltration. The infiltration very commonly involves the muscular coat, and even the serous, and



Fig. 285.—Section of a small tubercular ulcer. In the middle there is a crater-shaped ulcer (*a*) with overhanging edges. The mucous membrane around is infiltrated with round cells in the midst of which a few tubercles are indicated. Beneath the ulcer the muscular coat is infiltrated. At *d* there is a small sub-serous tubercle.  $\times 16$ .

tubercles are often to be seen almost free of inflammatory infiltration at a distance from the ulcers in the midst of the muscular coat, or even in the peritoneum (see Fig. 285).

The ulcer itself extends much more laterally than deeply, and in its extension it does not respect the boundaries of the Peyer's patches or solitary follicles, but passes beyond them, generally advancing more across the gut than longitudinally. In this way ulcers are, not infrequently, much elongated in a direction transverse to the intestine, and may even form a ring around it. The ulcers have usually their centres at the part of the gut opposite the mesenteric attachment, as the Peyer's patches, in which they begin, have their seats there.



The position of the ulcers is often indicated before opening the intestine by the appearance of the peritoneal surface. If the ulcer be of any considerable size there is some inflammation of the peritoneum, evidenced by redness and, very often, by elongated projections of vascular connective tissue. But above all we can generally see in the peritoneum groups of little white nodules, which are tubercles in or under the serous coat. The existence of these tubercles is sometimes even useful in determining whether an ulcer is a tubercular one or not. It is very rare indeed that the ulcer penetrates through the peritoneal coat, and if it does, it is usually protected by adhesion to neighbouring loops of intestine.

As a general rule the tubercular ulceration is most marked in the ileum, but not infrequently the colon is attacked, and there may be ulcers of very large size there.

The process is for the most part a chronic one, and the ulcers advance slowly. Sometimes, however, and usually in connection with acute phthisis, there is a rapid swelling of the follicles, and ulceration occurs by softening, without preliminary caseous necrosis. In these cases perforation is much more liable to occur.

As amyloid degeneration is a frequent accompaniment of phthisis, it may coincide with tubercular ulceration in the intestine.

**2. Syphilis.**—With the exception of affections of the anus, which may extend into the rectum, syphilitic lesions are excessively rare in the intestine. A few cases have been observed in syphilis in new-born children. There are ulcers and cicatrices in the wall of the intestine, and the coats are infiltrated. There may also be gummata and ulceration in ordinary syphilis.

**Literature.**—*Tuberculosis*—SPILLMANN, *Tuberc. du tube digestif* (with literature), 1878; ORTH, *Lehrb. d. path. Anat.*, 830; BAUMGARTEN, *Zeitschr. f. klin. Med.*, 1885, and *Tuberculose*, 1885; GRAWITZ, (*Peritonitis*) *Charité Ann.*, 1886. *Syphilis*—MRACEK (with literature), *Vierteljahrschr. f. Dermat. und Syph.*, x., 1883; FÖRSTER, *Lehrb. d. path. Anat.*, ii. 148; KLEBS, *Handb. d. path. Anat.*, i. 261; LANG, *Vorles. über Syph.*, 1885; ISRAEL, *Charité Ann.*, ix., 1884; NORMAN MOORE, in *Hutchinson, Syphilis*, 1887, p. 256; BIRCH-HIRSCHFELD, *Lehrb.*, ii. 589, 1887.

#### IX.—RETROGRADE CHANGES AND ABNORMAL CONTENTS.

**Atrophy** of the mucous membrane of the intestine is an occasional consequence of inflammations, catarrhal, dysenteric, and others. There have also been described atrophic and degenerative changes in the intestinal nerves in acute fevers and in acute affections of the central nervous system. The changes in the nerves are accompanied by changes in the muscle.



**Amyloid degeneration** is the most frequent retrograde change in the intestine. It affects the arteries and capillaries of the mucosa and submucosa, and especially the vessels of the villi, but it often extends to the whole tissue of the villi except the epithelium, and to the muscular bundles of the submucosa. To the naked eye the intestine frequently appears pale and smooth, but the condition is best detected by the addition of iodine solution. The affected villi then appear as deeply stained granules, while the closed follicles and Peyer's patches are unaffected. If ulcers be present, as is not infrequently the case, they also are unaffected and show a contrast in colour. The villi are frequently extensively lost in amyloid disease, but this may be a post-mortem effect, the brittle amyloid structures being broken off in handling the intestine.

**Gangrene** occurs, as mentioned above, in various conditions, infarction, dysentery, hernia. It also occurs occasionally in consequence of continued pressure by hard contents. This is most common in the vermiform appendage, but also occurs in the colon and rectum from hard fæces. The result is the formation of ulcers. Gangrene also results sometimes from separation of the mesentery, when this occurs close to the intestine.

**Foreign bodies in the intestine. Intestinal concretions.**—These mostly consist of the ordinary contents dried-in and impregnated with lime salts. The concretions have usually a kernel composed of a gall-stone, a fruit-seed, a mass of hairs, etc., or merely dried fæces, and such kernels are coated with phosphate and carbonate of lime. Such concretions are very common in animals, especially the horse, in which they have been found weighing 20 pounds. They are composed of the husks of corn mixed with hairs which the animals have licked from their bodies and swallowed. These are felted together and compacted into large balls.

Intestinal concretions occur chiefly in the cæcum and vermiform appendage, where they may give rise to ulceration and perforation. (See above, Typhlitis and Perityphlitis.)

**Gall-stones** also occur as foreign bodies in the intestine, and sometimes they are large enough to cause obstruction at the ileo-cæcal valve or elsewhere.

**Parasites in the intestine.**—Vegetable parasites are of little consequence if we except those considered in previous paragraphs.

Animal parasites have been described in the general section on parasites. They comprise chiefly the tape-worms, the ascarides, the oxyuris, the trichocephalus dispar, and the dochmius duodenalis.

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Virch. Arch., lxxxii., 1880; EDINGER, (Amyloid dis. and dilat. of colon) D. Arch. f. klin. Med., 1881; ZESAS, (Separation of mesentery) Arch. f. clin. Chir., 1886; SCHUBERG, (Concretions) Virch. Arch., xc., 1882; LABOULBENE, Arch. gén. de. méd., xxii., 1873; FRIEDBERGER und FRÖHNER, Lehrb. d. path. d. Haustiere, 1885; LEUBE, in Ziemssen's Cyclop., vii., 1877.

#### X.—TUMOURS OF THE INTESTINE.

**Primary cancer.**—This is by far the most frequent and important form of tumour. Cancers are almost confined to the **Large intestine**; we have already connected this with the fact that this portion is most exposed to the irritation of the stagnant fæces. This is the more evident as the most frequent localities are the **Rectum**, the **Flexures**, and the **Cæcum**. At the same time cancers are not unknown in the small intestine.

The commonest form of cancer is a very **Chronic epithelioma** with gland-like masses of cylinder cells (Fig. 286). From the resemblance of

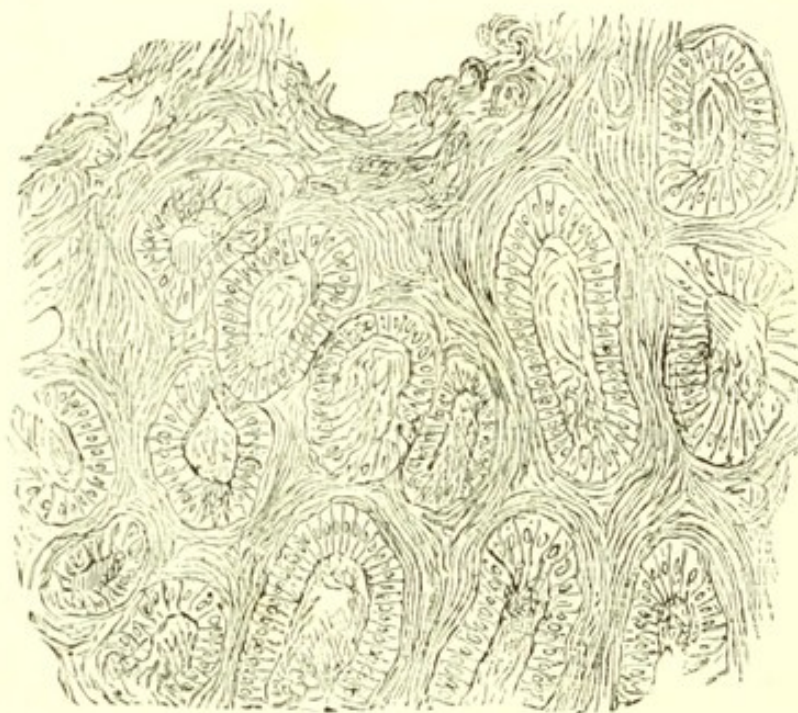


Fig. 286.—Cylinder-celled epithelioma of rectum. There are gland-like spaces containing epithelium, which at the periphery is characteristically cylindrical.

these structures to the tubular glands of the intestine, and from the fact that evidently the tumour originates from these glands, the name **Adenoid tumour** or **Malignant adenoma** is frequently used, just as in the case of the stomach.

The tumour is frequently ring-shaped, and as ulceration readily occurs, there may be little more than an ulcer with infiltrated edges. Even the infiltration of the edges may be very slight, and it may be difficult to be sure that the disease is cancerous. The ulceration is



accompanied by newformation of connective tissue in all the coats of the intestine, and while adhesion occurs to parts around, there is great contraction of the intestine. In this way strictures arise, and the occlusion of the calibre may be increased by the mucous membrane at the edges of the ulcer being thrown into folds. In fact, if we except hernias, this is probably the most frequent cause of intestinal obstruction.

It has been already noted that the proper cancerous structure is often to a large extent destroyed by ulceration, but at the edges we may distinguish masses of epithelial cells with a glandular arrangement. In the adhesions around the affected part of the intestine and at further distances enlarged glands may be found with cancerous tissue.

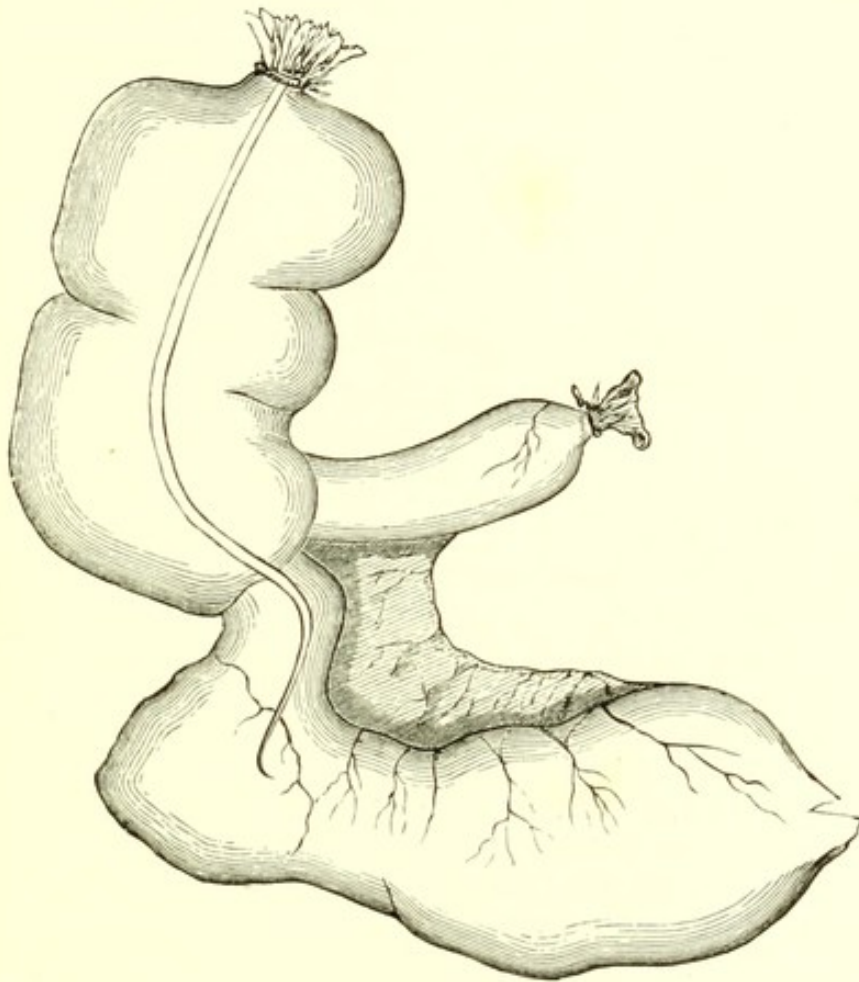


Fig. 287.—Cyst of vermiform appendage. The appendage is converted into a bulky cyst which was filled with gelatinous material.

Next to the epithelial the most common form is **Colloid cancer**. This forms a much more prominent tumour, which may involve a considerable length of the gut, and incorporate the entire coats. This form may produce obstruction by its mere bulk, although it also is liable to ulceration, and the consequent contraction of the gut added to the bulk of the tumour frequently effects obstruction. It does not so readily affect the



lymphatic glands, but more usually extends through the wall into the peritoneum.

**Soft cancer** is much less common here than in the stomach, and it resembles the same form of tumour there. It extends to the lymphatic glands and frequently also to the liver.

The remaining tumours of the intestine are of little importance. **Fibromas, Lipomas, Myomas, and Sarcomas** occur, and they are all apt to pass into the interior of the intestine and assume the form of **Sub-mucous polypi**. It is said that these polypi by being dragged on in the peristaltic movements of the intestine may produce invagination. **Mucous polypi** are also to be mentioned as occurring, especially in the rectum, in catarrh.

A peculiar form of tumour is **Cyst of the vermiform appendage**. In some cases of obstruction of the aperture the appendage becomes distended with mucus and may assume large dimensions, as in Fig. 287. The contents may remain mucous in character or become serous.

**Lymphoma** is not infrequent as a form of secondary newformation in the intestine. A general enlargement of the closed follicles occurs sometimes in leukæmia and in Hodgkin's disease, and it may be so considerable as to give the appearance of numerous prominent tumours projecting from the surface.

There is, however, sometimes a more definite tumour formation in **Hodgkin's disease**, when the primary lesion is localized in the mesenteric and intestinal lymphatic structures. There may be extensive tracts of the intestine infiltrated with lymphatic tissue which may replace the mucous membrane and the other coats, so that sometimes the enlarged intestinal tube is converted into a soft white structure composed of lymphatic gland tissue (see under Malignant Lymphoma).

**Literature.**—KLEBS, Handb., i., 1869; CRIPPS, (numerous illustrations) Path. trans., 1881, xxxii. 87; HAACK, Arch. f. klin. Chir., xxix., 1883; WEIGERT (Secondary cancer) Virch. Arch., 1876, lxxvii. 513; CARRINGTON, (Hodgkin's dis., with references to literature) Path. trans., xxxv. 385. Literature of other forms of tumour in Orth, Lehrb. der path. Anat., 1877, p. 855; COATS, (Cysts of vermiform Brit. Med. Jour., 1875, i.

## XI.—OBSTRUCTION TO THE CALIBRE OF THE INTESTINE.

This condition has been incidentally mentioned as occasioned by several of the lesions already described. It may be well here to sum up the various forms of intestinal obstruction and to describe the effect produced when the calibre of the intestine is interrupted.

Next to hernia, the most frequent cause of obstruction is the contraction of cicatrices resulting from ulceration, and in the great majority



of cases the ulcers are malignant, that is to say, they arise by the breaking down of cancers. It is to be remembered that in contractions of the calibre such as these the direct cause of the final obstruction may be the folding of the mucous membrane above the stricture, this acting like a valve to the narrowed part. Tumours also obstruct sometimes by their bulk. Similarly gall-stones, masses of fæces, collections of round worms, may occlude the calibre. Invagination is the commonest cause in children. There is also twisting or torsion. Lastly, we have obstruction by incarceration, the intestine passing into a position where a loop is caught at the neck and strangled. Hernias form the commonest examples of this kind, but there are a number of cases in which the intestine slips under a bridge or ligament, and gets, as it were, pinioned and compressed. These latter merit further consideration.

**Obstruction by a bridle or ligament** occurs so obviously to the mind that its frequency is apt to be overrated. In reality it forms a small proportion of the entire cases. The abnormal band may be formed by the adhesion of normal or other structures to various parts of the abdomen so as to form bridges. Thus the vermiform appendage may adhere by its tip and form a bridge; or a Meckel's diverticulum may similarly adhere; or the omentum may be tied down in such a way as to form a narrow bridle. But sometimes the band is itself distinctly of new-formation; there has been a local peritonitis, and two opposing surfaces have become adherent in the usual way of inflammatory adhesion, by vascular connective tissue; by the movements of the intestine the connective tissue has been dragged upon and elongated till it forms a band or ligament, which may afterwards tie down a loop of intestine.

A very unusual form of incarceration is that in which there is an **Aperture in the mesentery** through which the intestine passes. This may be produced by a tear in the mesentery. But in a case observed by the author there was a large round gap about 5 inches in diameter with rounded edges, and obviously of long standing, perhaps congenital. The greater part of the small intestine had become impacted in a complicated fashion into the aperture, and obstruction had finally resulted.

**Results of obstruction.**—The most direct effect is that the intestinal contents accumulate in the parts of the intestine above. The bowel may be greatly distended and its muscular tissue paralysed by the internal pressure; the stagnating fæces decompose, and the contents of the small intestine may acquire the characters of those of the large. This decomposition of the intestinal contents leads to several important results. In the first place there may be great formation of gas which still further distends and may produce the condition of exaggerated inflation called **Meteorism**. Then a return of the fæces in the reverse



direction may occur, and if the decomposition is sufficiently advanced there may be a vomiting of matter having the characters of fæces. This alarming phenomenon of **Stercoraceous vomiting** is sometimes referred to a reverse current extending from the colon. We can hardly suppose, however, that the fæces are able to pass the ileo-cæcal valve, and for the most part the vomited matters are really the stagnant contents of the small intestine which have undergone the fæcal decomposition. Of course there is a reverse current, else the intestinal contents would not reach the stomach. But this does not necessarily imply a vermicular movement in the reverse direction. If the intestine is distended, and is interrupted at a particular point, then the vermicular movement in the usual direction will carry the external layer of contents downwards to the point of obstruction. In order to accommodate this fresh arrival there must be a return current upwards in the centre of the tube. There is a similar condition to what exists normally in the stomach where the contents are carried to the pylorus, and meeting the obstructing sphincter return for the most part down the centre of the cavity towards the fundus.

Again, the products of decomposition may be absorbed to an abnormal extent and appear in the urine. There are two substances resulting from the decomposition of albuminous substances which are normally present to a limited extent in the urine, namely, **Indican** and **Carbolic acid**. In cases of obstruction these substances, or one of them, are increased in quantity sometimes to a very striking extent. This increase does not occur unless the stagnation extends to the small intestine where the albuminoids are present. It does not occur therefore in ordinary constipation which affects the large intestine alone, apparently because the albuminoids are absorbed before they reach the large intestine.

**Literature.**—COHNHEIM, *Allg. Path.*, ii. 147, 1882; JAFFÉ, (Indican in urine) *Virch. Arch.*, lxx. 72, 1877; SALKOWSKI, *ibid.*, lxxiii. 409, 1878; BRIEGER, *Zeitschr. f. klin. Med.*, iii.



## SECTION VII.

DISEASES OF THE LIVER, PANCREAS, AND  
PERITONEUM.

- A.—THE LIVER. *Introduction. Post-mortem changes.* I. MALFORMATIONS AND DEFORMITIES, II. DISORDERS OF CIRCULATION. 1. *Passive hyperæmia, nutmeg liver,* 2. *Thrombosis and Embolism.* III. RETROGRADE CHANGES, chiefly *fatty infiltration, amyloid disease, and pigmentation; Icterus.* IV. ACUTE YELLOW ATROPHY. V. INFLAMMATIONS. 1. *Suppurative hepatitis, including tropical and pyæmic abscesses; also biliary abscess.* 2. *Chronic interstitial hepatitis, Cirrhosis; causation; lesions; effects. Biliary and hypertrophic cirrhosis.* 3. *Perihepatitis.* VI. SYPHILIS AND TUBERCULOSIS. *Syphilitic cirrhosis and gummata. Tuberculosis, chiefly secondary.* VII. TUMOURS, chiefly *cancers; occasionally primary—chiefly secondary. Parasites, chiefly Echinococcus.*
- B.—BILE-DUCTS AND GALL-BLADDER. 1. *Gall-stones, single and multiple,* 2. *Obstruction of ducts,* 3. *Rupture and perforation,* 4. *Tumours.*
- C.—PANCREAS. *Malformations, hæmorrhages, inflammations, retrograde changes. Tumours, especially cancers. Concretions and obstructions of duct.*
- D.—PERITONEUM. *Introduction.* 1. *Malformations;* 2. *Disorders of circulation, chiefly hæmorrhage and ascites; chylous ascites and ascites adiposus;* 3. *Inflammations, septic, chronic;* 4. *Tuberculosis, tubercular peritonitis;* 5. *Tumours, including Retroperitoneal sarcoma.*
- E.—SECONDARY EXTENSION OF CANCERS OF THE ABDOMINAL ORGANS.

## A.—THE LIVER.

**I**NTRODUCTORY.—The liver is the largest gland in the body, its weight being on an average from 48 to 58 ounces in the adult male and 40 to 50 ounces in the female.

In its **Function** the liver is related to the intestinal canal and the blood. In the foetus it is probably engaged in forming the red blood-corpuscles, and probably throughout life it aids in disposing of the effete corpuscles, the bile pigment being derived from that of the blood-corpuscles. In addition it has an important function in the preparation of glycogen, whatever may be the uses to which this substance is put. It also seems to have important relations to the free fat of the



body, and under certain circumstances stores it up in its substance (see Fatty Infiltration).

The functions of the liver being thus manifold and somewhat obscure the effects of disease on it are difficult to disentangle. Many of the diseases cause atrophy of its proper tissue, but they nearly all at the same time affect either the body as a whole or the vascular arrangements of the abdomen which are so peculiarly related to the liver.

The **Circulation in the liver** is very peculiar and claims special attention, because most diseases of the organ produce important effects upon it. Most of the blood comes to the liver by the **Portal vein**, and the proper hepatic tissue is arranged in relation to the ultimate ramifications of that vessel. The interlobular veins form these ultimate ramifications, and they are in immediate connection with the hepatic lobules. **The Hepatic lobule** is a group of hepatic cells with blood-vessels, having in man a polygonal or somewhat globular shape; it measures about the twentieth of an inch in diameter. The interlobular vein lying outside the lobule sends capillaries into it, and these seeking the centre of the lobule open into the central or intralobular vein which is the radicle of the hepatic vein. Between the capillaries lie the hepatic cells, which are arranged in rows or cylinders radiating from the centre of the lobule like the capillaries. The stellate cells of Kupffer lie between the hepatic cells and the capillaries and seem to be intimately connected with both.

Blood is also brought to the liver by the **Hepatic artery** which supplies chiefly the connective tissue and walls of the blood-vessels. Its capillaries terminate in veins which open into the interlobular veins (according to Cohnheim and Litten), so that this blood also finds its way into the hepatic capillaries and on into the hepatic vein.

The **Connective tissue** of the liver is often described as if it formed a special covering to the portal vein, being called Glisson's capsule. It really forms a supporting stroma which holds the portal vein, the hepatic artery, and the hepatic duct, which all lie side by side. The lymphatic channels are also contained in it. In swine the connective tissue surrounds each lobule and defines it distinctly from its neighbour, but in man it stops short at the interlobular vein, and except where this vein is, the lobules at their margins merge into each other, and their capillaries are in common. Examination of Fig. 289, p. 764, in which the lobules are demarcated by fatty infiltration, will show how they run into one another at their peripheries. Although no proper fibrillated connective tissue is present inside the lobules, yet a fine reticulum accompanies and supports the capillaries.

The circulation in the portal vein, and especially in the capillaries of



the liver, must be unusually slow. The blood before it reaches the liver has passed through one set of capillaries, and here it passes through a second set; it has therefore lost very largely the force derived from the contractions of the heart. It is probably for this reason that the liver is so frequently the seat of secondary diseases, such as tuberculosis, abscesses, cancers, etc., and of the deposition of solid pigments. We may suppose that as the blood moves so slowly there will be time for any infective material to settle down and produce its special effects. We know that when vermilion is injected into the blood it is found largely in the capillaries of the liver. Similarly the abnormal pigment present in the blood in melanæmia is found largely in these.

The **Hepatic ducts** running in the connective tissue along with the portal vein and hepatic artery are lined with cylindrical epithelium. These are connected with the biliary capillaries, which originate inside the lobules and seem to be in part formed by the hepatic cells themselves, each cell having on its surface a groove which, with a corresponding groove in the cell opposed to it, forms a tube.

**Post-mortem changes.**—The liver is liable to local alterations in colour, in the form of pale anæmic areas, from the pressure of neighbouring structures, as the ribs, or distended loops of intestine. It frequently also assumes a blue colour from decomposition, especially where it is in contact with the transverse colon.

There are some curious cases in which **Cavities filled with gas** develop throughout the liver after death. These are due to decomposition, the agents of which have been conveyed to the liver before death. The condition may be associated with a similar appearance in the spleen, and it will occur chiefly in cases of septic wounds. It was so at least in a case observed by the author, of which the liver is preserved in the Museum of the Western Infirmary.

**Literature.**—*General works*—BUDD, Dis. of liver, 2nd ed., 1852; MURCHISON, Clin. lect. on dis. of liver, etc., 2nd ed., 1877; ANSTIE, In Reynolds' Syst. of med., iii., 1871; HABERSHON, Path. and treatment of some diseases of liver, 1872; FRERICHs, Dis. of liver; transl. Syd. Soc., 1858-61; CHARCOT, Leçons sur les malad. du foie, 1877; HANOT et GILBERT, Malad. du foie, 1888.

#### I.—MALFORMATIONS AND DEFORMITIES OF THE LIVER.

**Congenital malformations** of the liver are not common. There are cases of absence of the liver, and of defect of one of the lobes, or irregularity in the lobes. What may be called supernumerary livers have also been found in the form of isolated pieces of liver tissue in the suspensory ligament. Of more importance is congenital absence of the gall bladder, along with which there is usually a dilatation of the bile ducts.

When considerable portions of the liver are destroyed, there is liable to be a **Compensatory hypertrophy** of remaining parts. The loss of liver substance may be congenital, as in a case observed by the author in which, probably from some



injury early in foetal life, the right kidney and the greater part of the right lobe of the liver were wanting. In that case the left lobe was greatly increased in size, and the liver as a whole was of the normal weight. There may be a similar compensatory hypertrophy from destruction of liver tissue in after life. The destruction may be by pressure of hydatid cysts, syphilitic gummata, and cicatrices, or even by cirrhosis. The region of the atrophy will determine that of the hypertrophy, but sometimes the left lobe or the lobus Spigelii undergoes great enlargement.

The liver is sometimes **Transposed** along with a general transposition of the viscera. In a case recorded by Fraser, there was not only a transposition but an occasional **Dislocation** of the liver, which came down into the left inguinal region.

An **Acquired deformity** is the so-called **Stay liver** of females. This deformity is usually stated to be due to tight lacing, but although doubtless greatly aggravated by this, it is also induced by the ordinary methods of suspending the garments by a tight ligature round the waist. The effect on the liver is that it is greatly flattened, while a transverse shallow depression forms along the upper surface, affecting chiefly the right lobe. In this groove the capsule is thickened, and there is obviously considerable loss of liver tissue. Sometimes the atrophy along this groove is such that the two portions can be folded together. With the flattening there is great displacement of the liver, whose lower edge may extend as far down as the umbilicus. More or less of this deformity is to be found in nearly all female bodies, and also in the bodies of men who have been in the habit of wearing tight belts round their waists.

Sometimes the upper surface of the right lobe of the liver presents elongated depressions passing from behind forwards, which are really **Folds of the liver**, and are produced chiefly when there is some obstruction to expiration. In that case the diaphragm is depressed by the dilated lung, and at the same time the lower ribs are drawn down in violent expiratory movements by the abdominal muscles, and pressed against the liver, which is supported below by the contraction of the abdominal muscles.

The liver is very liable to **Changes of position**. Tumours or fluid in the abdomen carry it upwards. Depressions of the diaphragm press it downwards. The suspensory ligament may be elongated so that the organ is unduly moveable, and may undergo displacement downwards.

**Literature.**—THIERFELDER, in Ziemssen's Cycl. ix., 1887; LAUDAN, *Wanderleber und Hängebauch der Frauen*, 1885; FRASER, *Glasg. Med Jour.*, xx., 1883.



## II.—AFFECTIONS OF THE CIRCULATION IN THE LIVER.

1. **Passive hyperæmia.**—This is perhaps the commonest of all affections of the liver. The names **Nutmeg liver** and **Red atrophy** are sometimes given in cases of prolonged passive hyperæmia, these names indicating certain appearances presented by the tissue. Passive hyperæmia occurs when any obstruction exists in the circulation of such a nature as to interfere with the return of blood from the inferior vena cava to the right side of the heart. The commonest cause is valvular disease of the heart, especially mitral disease, but it also results from obstruction to the pulmonary circulation as in bronchitis and emphysema, etc. In these cases there is a general engorgement of the systemic venous circulation, with increase of blood-pressure in the veins. As the normal circulation in the liver is unusually slow, and the blood-pressure low, any increase in the blood-pressure in the general venous

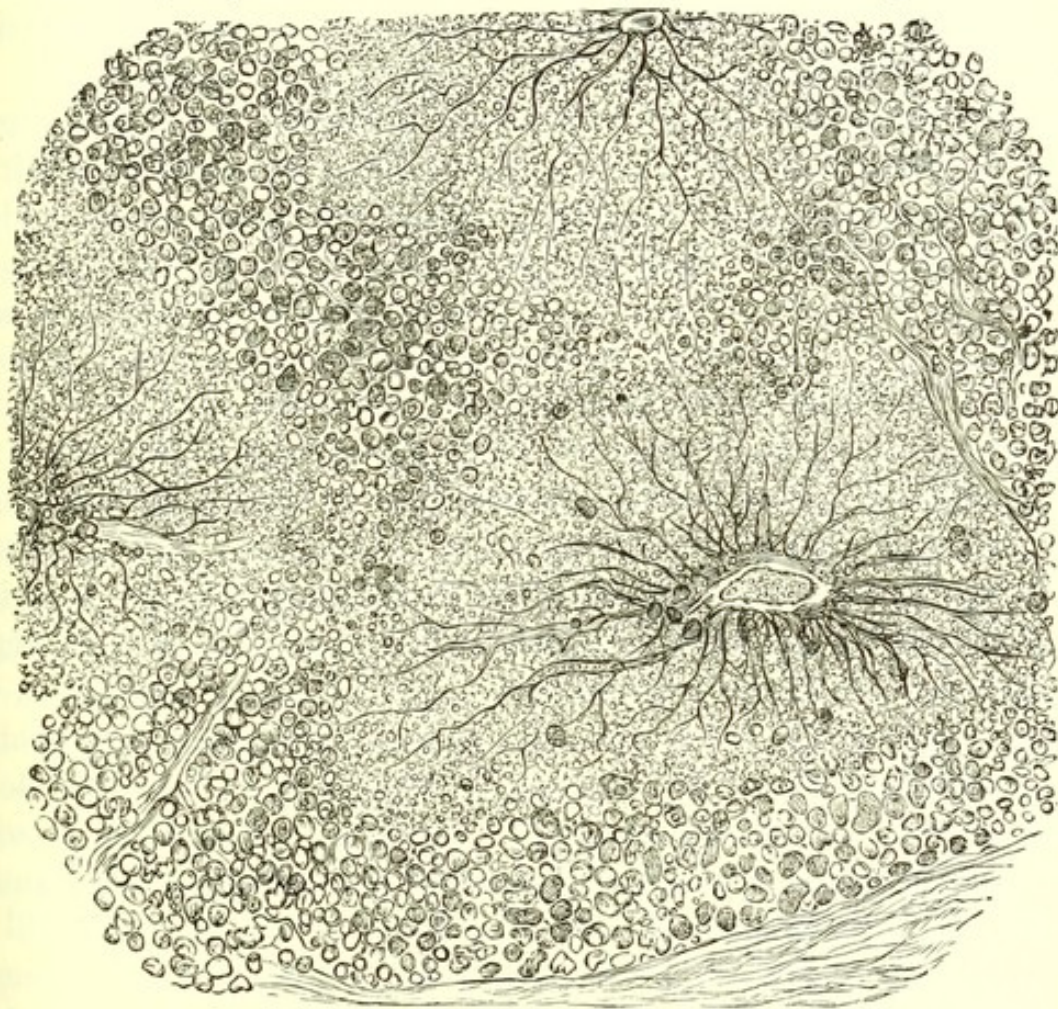


Fig. 288.—Passive hyperæmia of the liver. One lobule and parts of other two are shown. In the central parts of the lobules there is almost nothing but dilated capillaries containing blood-corpuscles; a few atrophied and pigmented hepatic cells being visible. At the peripheral parts of the lobules the hepatic cells are seen, many of them pigmented.  $\times 50$ .

circulation will tell particularly on the vessels of the liver, and we may even suppose that the blood passing up the inferior cava may in such cases regurgitate into the hepatic vein.



The result of this is very great distension of the hepatic vein and its radicles—the central or intralobular veins. The distension extends to the capillaries, which, as in Fig. 288, are sometimes found enormously dilated. The dilated capillaries by pressure cause **Atrophy of the hepatic cells**, and it often happens that the central parts of the lobules are entirely occupied by enormously dilated capillaries, while the hepatic cells are hardly visible (see figure). The remains of the hepatic cells frequently contain brown **Pigment granules**. The atrophy of the hepatic cells in these regions may be complete, only a few pigment masses representing them. Sometimes, instead of a simple atrophy we have **Fatty degeneration** of the hepatic cells, localized in the central parts of the lobules.

Sometimes in addition to these changes there is a hypertrophy of the connective tissue surrounding the lobules, but this is usually inconsiderable, and it is a mistaken view that a cirrhosis of the liver arises out of passive hyperæmia.

To the naked eye there is a general enlargement of the organ, which, on section, appears unduly red. On close examination it is seen that there are minute areas of a deep red colour corresponding with the central parts of the lobules, and these are surrounded by zones of a grey or yellowish colour. The result is that the lobules are, as it were, mapped out by these contrasting colours, and are for the most part individually visible to the naked eye. Sometimes several adjacent lobules are almost completely occupied by dilated vessels, and there is a narrow ring of normal tissue around each, so that the portion of liver has, as a whole, an almost cavernous structure and presents a deep red colour on section. It is these variations in colour and figuring, which give the cut surface the appearance of the section of a **Nutmeg**.

In parts where there has been great loss of the proper tissue, the liver may appear to the naked eye after death partially atrophied. During life the dilated veins and capillaries are distended with blood, but after death, the blood-pressure being removed, these may, to some extent, collapse. In this way the surface has sometimes a granular appearance, which may be mistaken for that of cirrhosis, but it will be noticed that the depressed or atrophied parts have a deep red colour.

2. **Thrombosis and Embolism.**—Thrombosis of the **Portal vein** is of somewhat frequent occurrence, especially as a result of cirrhosis. It also occurs sometimes by propagation of thrombosis from the radicles of the vein, or by embolism from them. The consequences to the liver itself, even of complete closure of the portal vein, are not so important as might be looked for. It is stated by some that it may lead to cirrhosis of the liver, but there is no proper foundation for this state-



ment, and cirrhosis is much more frequently the cause than the result of thrombosis. It has also been asserted that diabetes mellitus may occur as a result of thrombosis of the portal vein. The consequences outside the liver are much more important. There is dilatation of the radicles of the portal vein in the abdominal viscera, with ascites, which is generally very extreme. The results in this regard are essentially similar to those which occur in cirrhosis, and are referred to more particularly below, under that heading.

Thrombosis and embolism of the **Hepatic artery** are not usually of consequence unless the plug be of a septic nature. The hepatic artery is not an end-artery, and if some of its branches be obstructed the anastomosing communications restore the circulation. A few cases have been observed, however, in which something approaching to an infarction has occurred. If the main branch of the artery be obstructed then we may have necrosis of the liver, but such an obstruction is scarcely liable to occur in man, and its effects have only been studied in experiments in animals.

**Embolism of the hepatic vein** by solid bodies passing backwards by gravitation from the inferior vena cava has been observed in a few cases (Recklinghausen). Thrombosis of this vein also occurs in cases of cancer and of abscess of the liver.

### III.—RETROGRADE CHANGES IN THE LIVER.

1. **Necrosis.**—On account of the double blood-supply in the liver necrosis seldom occurs. In a case of rupture of the liver, however, in which fissures cut off the blood-supply from a wedge-shaped piece, the author observed an appearance like that of the pale infarction. The tissue both in naked-eye and microscopic characters showed the appearances of coagulation-necrosis. There is also necrosis in connection with the pyæmic abscess.

2. **Atrophy.**—The hepatic parenchyma is very liable to atrophy. We have a **General atrophy** in cases of emaciation and inanition, and also as a senile phenomenon. In these cases the liver is small in size and deep in colour. There is also a general atrophy in melanæmia (see further on).

**Local atrophy of the liver** is usually the result of pressure, as by tumours growing in the liver, by dilated capillaries, by the swollen capillaries in amyloid disease, or otherwise. Indeed the hepatic cells seem very readily to give way before pressure, so that one often sees in the neighbourhood of a growing cancer the hepatic cells flattened out concentrically and atrophied.

3. **Parenchymatous infiltration.**—This occurs, as we have previously



seen, in connection with many acute diseases. It is this which causes the enlargement of the liver in the acute fevers, pneumonia, erysipelas, etc. The hepatic cells are highly granular, and if the albuminous granules be destroyed by liquor potassæ it will be seen that there are numerous fine fat granules, the cloudy swelling passing into fatty degeneration.

4. **Fatty degeneration.**—This is frequently seen as a further stage of the condition just mentioned, and is a special feature in acute yellow atrophy to be presently described. It is also usually present in pernicious anæmia. A more localized fatty degeneration is met with in consequence of local affections of the liver, as in passive hyperæmia, but more especially in cirrhosis of the liver. It is characterized by the presence of minute fat granules in the protoplasm of the cells.

5. **Fatty infiltration.**—The pathology of this condition has been referred to in a former part of this work (see p. 160). It is most typically seen in phthisis pulmonalis and in some other wasting diseases. It also occurs, although in a somewhat different form, in chronic alcoholism.

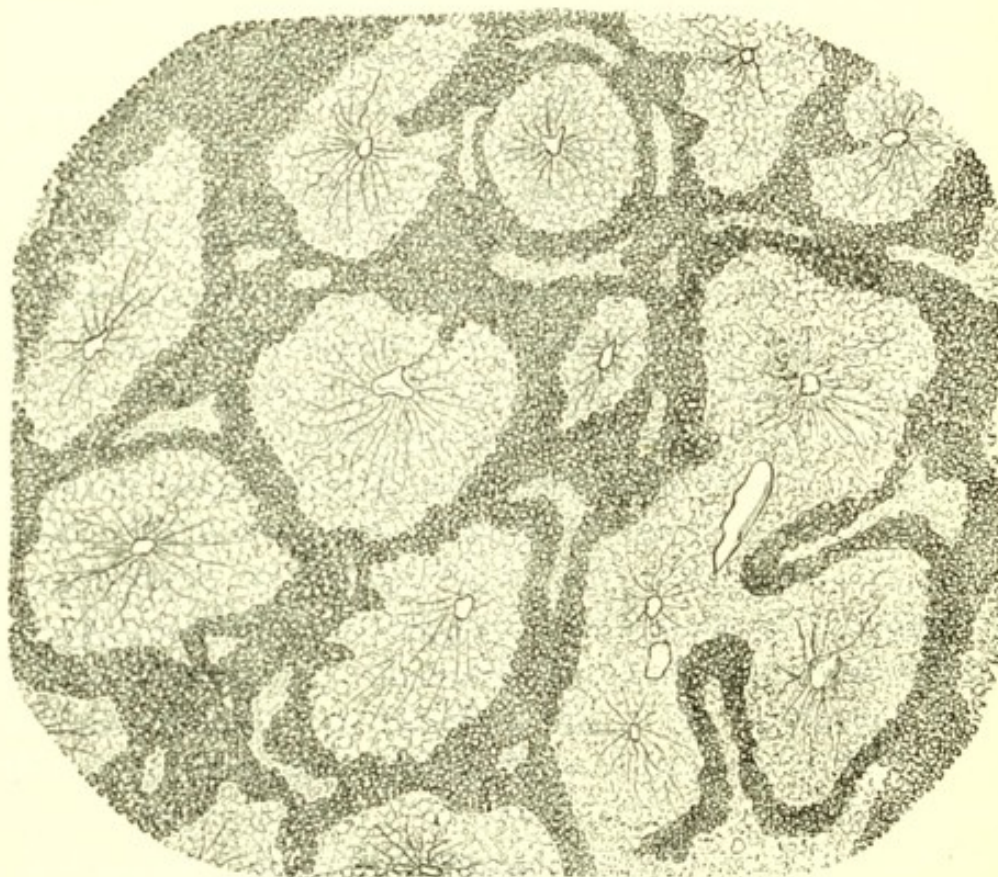


Fig. 289.—Fatty infiltration of the liver; osmic acid preparation as seen with a very low power. The peripheral parts of the lobules are demarcated by the fatty infiltration.  $\times 16$ .

In fatty infiltration fat is brought by the portal vessels and deposited in the hepatic cells. It is first deposited in those nearest the terminals of the portal vein, that is to say, in the peripheral parts of the lobules (see Fig. 289), so that at first each lobule has a peripheral zone of fatty



infiltration. This is very markedly seen when a section is examined microscopically in the fresh state or stained with osmic acid. In the latter case the fatty peripheral parts become of a blackish colour, and the section assumes a very striking figured appearance as shown in the figure. As the condition advances, more and more of the lobule is affected, and the infiltration may overtake its whole extent. Even in extreme cases, however, it usually preponderates at the peripheral parts. The fat is present in larger and smaller drops inside the hepatic cells, but the tendency is for the drops to assume a considerable size (as in Fig. 290). The protoplasm of the cells is pushed aside by the fat, but its function is not apparently interfered with. The stellate cells of Kupffer are sometimes specially affected.

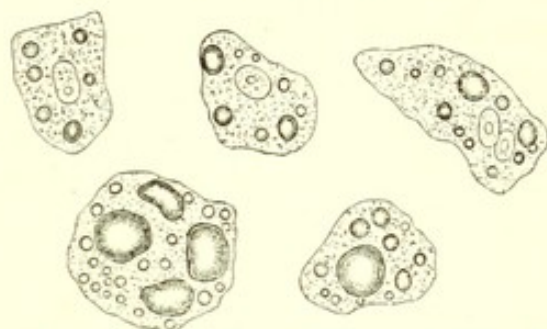


Fig. 290.—Fatty infiltration of liver. The cells are isolated, and they contain larger and smaller drops of oil.  $\times 350$ .

The fatty liver is increased in weight, and in extreme cases it may be double the normal. The organ is enlarged generally, rounded at the edges and increased in thickness. It is unduly soft in consistence and has an opaque pale colour which may be very marked, and merge towards a saffron yellow. It is nearly always possible with the naked eye to distinguish evidences of the lobular distribution of the fat. The lobules are, in fact, mapped out in the most characteristic figured manner, so that, on looking closely, we can distinguish each with perfect accuracy.

The appearance presents a considerable resemblance to that of the nutmeg liver, where also the lobules are mapped out; but in the fatty liver it is by unduly pale tissue at the peripheral parts of the lobules, which contrasts with the normal central parts; while in the other case it is the normal which is at the periphery, and it contrasts with the red central parts, and the whole cut surface is unduly red.

In **Chronic alcoholism** the fat is more generally distributed. The liver as a whole has a soft greasy appearance, and fat will be found in larger and smaller drops throughout the lobules.

**6. Amyloid degeneration.**—The liver is usually affected when amyloid disease exists in the body, but the affection shows great differences in degree. It may be very slight in the liver while very advanced in the spleen and kidney, and vice versa. It is not known how these differences are determined. In phthisis pulmonalis which is the most frequent cause of amyloid disease we have all varieties in distribution.

The degeneration appears first in the arteries and capillaries, the appearance in the latter being as if, at intervals, the wall were swelled up by a translucent material. In these earlier stages there is no diffi-



culty in making out that inside the lobules it is the capillaries which are affected. The general arrangement of the capillaries is beautifully shown by the rose-pink colour in a section stained with methylviolet, the appearance being almost that of an injected specimen. Even in the earlier periods considerable atrophy of the hepatic cells can be observed as a result of pressure from the swollen capillaries, and the atrophic cells frequently contain fat. In advanced stages, however, the cells are very greatly destroyed, so that the liver tissue is replaced by amyloid substance. The tissue still shows indications of the lobular arrangement, and also of the capillaries, although scarcely any hepatic cells remain (see Fig. 38, p. 165).

It is remarkable that, with this great loss of the secreting substance, there is actual increase in size and weight of the liver. It is not uncommon to find the liver weighing two or three times the normal. The increase in size is not proportionate to that in weight, as the amyloid substance is of greater density than the liver tissue. The edges of the organ are rounded and its consistence elastic and resistant. The surface, and especially the cut surface, is homogeneous and presents a peculiar translucent waxy appearance, which in advanced stages is highly characteristic.

**7. Pigmentary infiltration.**—The hepatic tissue is liable to a considerable number of different forms of pigmentation.

**Icterus** is the most frequent and obvious cause of pigmentary infiltration of the liver. We have here to do with hepatogenous icterus, arising from obstruction of the bile ducts. When the bile ducts are obstructed the secretion of bile does not cease, and as excretion is hindered there occurs a re-absorption of the bile. As the hepatic cells lie between the biliary capillaries and the blood, the bile must traverse the former to reach the latter, and consequently the hepatic cells are most immediately and directly stained with the re-absorbed biliary pigment. The pigment at first stains the cells generally, but when the process is prolonged, the bile is deposited in the cells as solid granules, which have a deep brown or greenish colour. It rarely forms crystals. The pigmentary deposit occurs chiefly in the central parts of the lobules. The bile sometimes condenses in the biliary capillaries and forms solid plugs or moulds of these.

This biliary infiltration of the liver occurs in consequence of an obstruction either of the main duct (ductus communis). It results also from obstruction of the finer ducts in the substance of the organ, as in cases of **Cirrhosis** and **Cancer** of the liver, having in that case a partial distribution according to the disposition of the lesion which causes it.

Infiltration with blood-pigment occurs, as already mentioned in



Melanæmia, and in pernicious anæmia. In **Melanæmia** the pigment is in the capillaries, and the hepatic cells suffer considerable atrophy, probably because of the obstruction of the capillaries. In **Pernicious anæmia** there is a reddish brown pigment deposited in the hepatic cells, which may give a general coloration to the organ as a whole. In addition, colourless granules **containing iron** are present in the liver, and may be detected by the dark green colour which they assume on the addition of  $\text{NH}_4\text{S}$  or by the blue colour with ferrocyanide of potassium and hydrochloric acid. (See under Pernicious Anæmia.)

#### IV.—ACUTE YELLOW ATROPHY OF THE LIVER.

This disease, although by its name it is connected with a definite lesion in the liver, is really a general one, due to a **morbid poison** in the blood. In some cases the poison seems to be septic in its character, the disease coming on in the course of septic infection, especially in the puerperal state or in erysipelas, but generally no definite poison can be traced. Several observers have sought to connect the disease with certain forms of microbes (Klebs, Zander, etc.). It is consistent with this view that phosphorus produces changes so similar that some observers have suggested that acute yellow atrophy is always due to poisoning with phosphorus. This view, however, is not tenable.

Although at post-mortem examination the liver is nearly always reduced in size, yet in some cases it is found enlarged, and there is reason to believe that in the earlier stages it is generally **enlarged**. The enlarged liver is of a brownish yellow or bright yellow colour, either generally or in streaks and patches. In these parts the cells are enlarged and full of fat granules. Although this is a true fatty degeneration, yet the fat is in comparatively large drops.

In the further stages the liver is found **smaller** than normal, and its tissue is soft and flabby. The capsule is often wrinkled from the atrophy of the substance of the organ, and the organ flattens out by its own weight, its thickness being greatly diminished. The diminution in size often affects the left lobe especially. On cutting into the organ the section shows usually different shades of colour. The predominating tint is **yellow**, varying from the colour of gamboge to a dark yellowish brown. Mixed with this, however, there is a red colour forming the **red substance** of Zenker. Where the colour is yellow the tissue is softest, but the red parts are the most atrophied, the red substance arising by the absorption of the degenerated hepatic cells.

On examining the yellow tissue under the microscope, it will hardly



be recognizable as that of the liver (see Fig. 291). There are no proper

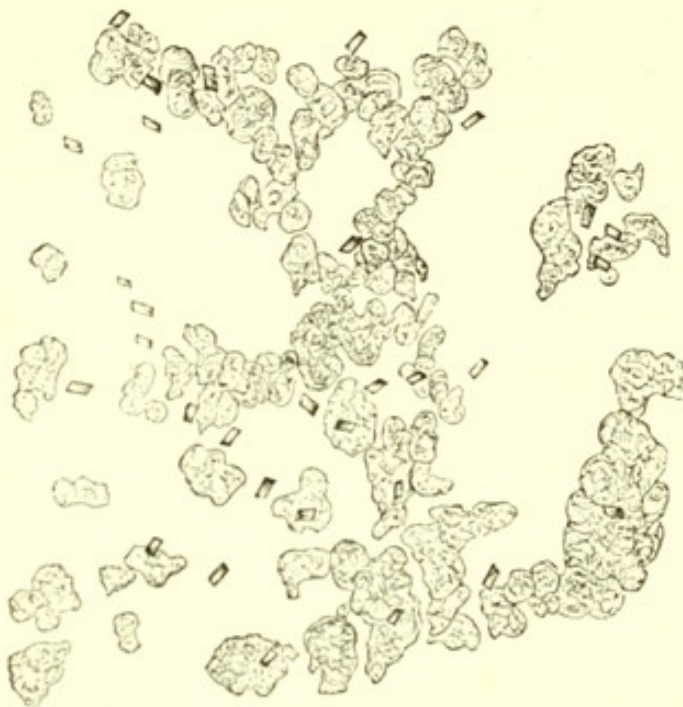


Fig. 291.—From liver in acute yellow atrophy. The hepatic cells form irregular clumps. There are numerous pigment crystals of similar shape to the crystals of haematoidin (Fig. 9, p. 84), and of a similar deep red colour, but smaller.  $\times 350$ .

hepatic cells, but instead irregular masses containing fat drops and granular debris, with here and there rhombic crystals of a reddish brown colour. The **Crystals of pigment**, which resemble those found in old blood clots in the brain, may be really formed from the biliary colouring matter. Sometimes **Leucine** is found in opaque clumps, white in colour and generally stratified. Acicular crystals of **Tyrosine** are also found.

The red substance does not present the abundant fatty debris; that has been cleared away, and there is now a fibrous material visible, merely sprinkled with fat granules. In some cases glandular-looking processes have been observed, and have been taken to indicate an attempt at regeneration of the lost hepatic substance, beginning in the capillary network of the hepatic ducts.

Besides the changes in the liver, there are lesions of a somewhat similar character elsewhere, especially in the epithelium of the kidneys and the muscular fibre of the heart. The symptoms are those of a general disease.

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#### V.—INFLAMMATIONS OF THE LIVER.

1. **Suppurative hepatitis.**—Acute suppurative inflammation will occur usually in connection with septic processes, which have been propagated to the liver.

(a) **The Tropical abscess** is scarcely ever met with in this country, except in persons who have been in hot climates, especially in India.



Much doubt exists as to the cause of the inflammation which gives rise to the abscess. In a large proportion of cases it is associated with dysentery, and it is believed by many that the abscess in the liver arises by septic matter being carried from the ulcerated intestine to the liver. This view, however, is not beyond question, and post-mortem examination shows that the tropical abscess is by no means uniformly associated with dysentery. It is possible, indeed, that an existing abscess by causing congestion of the portal circulation may predispose to inflammation of the intestine (Finlayson). It is not unlikely that certain states of the liver predispose to the development of the microbes in question, and that they may find entrance to the body without any lesion of the intestine.

A confirmation of this view is furnished by the fact, testified by Dr. David Wilkie of Simla, that in India, while the mortality ratio from dysentery among natives is not far from that among Europeans, the death-rate from hepatitis is proportionally greater amongst Europeans, probably from predisposition, brought about by their mode of living and racial characteristics. Thus in the European and native armies from 1877 to 1886 the death-rate from dysentery per 1000 men was 1·33 and 1·25 respectively (or, including diarrhœa and dysentery, 1·57 and 2·27), whereas the death-rate from hepatitis was 1·45 in the European and 0·17 in the native army. This is more strikingly apparent in the deaths in the jails of India (natives), the death-rate being, dysentery and diarrhœa, 1877-1886, 19·13; hepatitis, 0·22.

The abscess is mostly, but not always, solitary, and before death it generally grows to large dimensions, causing great enlargement of the liver. Having approached the surface, it sometimes causes a marked projection from the liver. The abscess may rupture into the peritoneum and cause fatal peritonitis, or its capsule having become adherent to the abdominal wall the abscess may by and by find its way to the surface of the body, there to discharge. It also occasionally bursts in other directions, as through the diaphragm into the pleural cavity, or lung; into the colon or other part of the intestine; or into the gall bladder from which the pus is carried to the duodenum. In these various ways the pus may be disposed of and perhaps recovery occur, but more frequently a wasting discharge from the cavity of the abscess remains.

In other cases the abscess does not enlarge, but rather dries-in. A thick connective tissue capsule forms around it like the wall of a cyst, and the pus thickens to a cheesy consistence. Afterwards the contents may become calcareous and the disease become virtually obsolete.

(b) **Pyæmic abscesses** are produced by pyogenic micrococci conveyed to the liver by the blood. They may be carried by the hepatic artery, in which case they have generally been absorbed from a septic wound and have traversed the lungs. Such abscesses will usually be associated



with similar metastatic abscesses in the lungs. On the other hand, the microbes may be conveyed by the portal vein, being associated with a septic thrombo-phlebitis in its radicles. This occurs most frequently in cases of perityphlitis.

The abscesses may occur apparently without any definite thrombosis in the veins or embolism in the liver, it being apparently sufficient that the microbes get into the blood. As the circulation in the hepatic capillaries is peculiarly slow, it is probable that such microbes will more readily lodge and propagate in the liver than elsewhere.

The microbes multiply and form zooglœa in the capillaries and interlobular veins, passing into the central veins. Wherever the colonies extend the liver tissue undergoes necrosis, the cells lose their nuclei and become individually indistinguishable. Inflammation is produced around, and there is here, as elsewhere under similar circumstances, an acute suppurative inflammation. The round cells accumulate around the necrosed portion, and gradually infiltrate it and break it down, so that an abscess forms.

The abscess may be primarily a small or a large one according as the capillaries over a smaller or larger area are invaded, but several abscesses may ultimately coalesce, and so we have them sometimes of very considerable dimensions.

The abscesses in this form are always multiple. In accordance with what has been stated the actual abscess is preceded by a grey or yellowish discoloration indicating the existence of necrosis of the tissue. There will be probably a group of such areas corresponding to necrosed hepatic lobules. Ultimately these break down and form a common abscess.

(c) **Biliary abscesses** occasionally form in connection with obstruction of the ductus choledochus. This will result when decomposition occurs in the stagnant bile and extends into the ducts in the substance of the liver. (See further on under Obstruction of Bile Ducts.)

2. **Chronic interstitial hepatitis. Cirrhosis.**—We have to do here with a chronic inflammation of the interstitial connective tissue of the organ.

**Causation.**—As the disease occurs for the most part homogeneously throughout the organ, the irritant is contained in the blood circulating in the liver. The name “gin-drinker’s liver,” frequently applied to this disease, involves the view that alcohol is commonly the irritant. Alcohol taken frequently in the form of undiluted spirits is believed to cause the disease, while beer and wine do not. But the disease may originate from other kinds of irritation whose nature is obscure. It has been produced experimentally by chronic poisoning with phosphorus



and cantharides, and it probably occurs also as a result of syphilis. It has been met with in young children, and the author has recorded a case in which a typical cirrhosis, with the usual secondary phenomena, occurred in a cat. The fact that it was a butcher's cat may indicate that it indulged in excess of eating rather than of drinking. Greenfield has also recorded two cases of cirrhosis in the cat.

It has been asserted of recent years that obstruction of the bile ducts is a cause of cirrhosis, and a special form of **Biliary cirrhosis** has been distinguished. The frequent occurrence of obstruction by gall-stones and otherwise without any of the appearances of cirrhosis, however, rather contradicts this view.

**Character of the lesions.**—In cirrhosis we have chronic inflammation of the connective tissue of the liver, resulting in newformation of a similar tissue. The interstitial connective tissue of the liver follows, as we have seen, the portal vessels, forming a frame-work in which are supported the portal vein and its branches, the hepatic artery

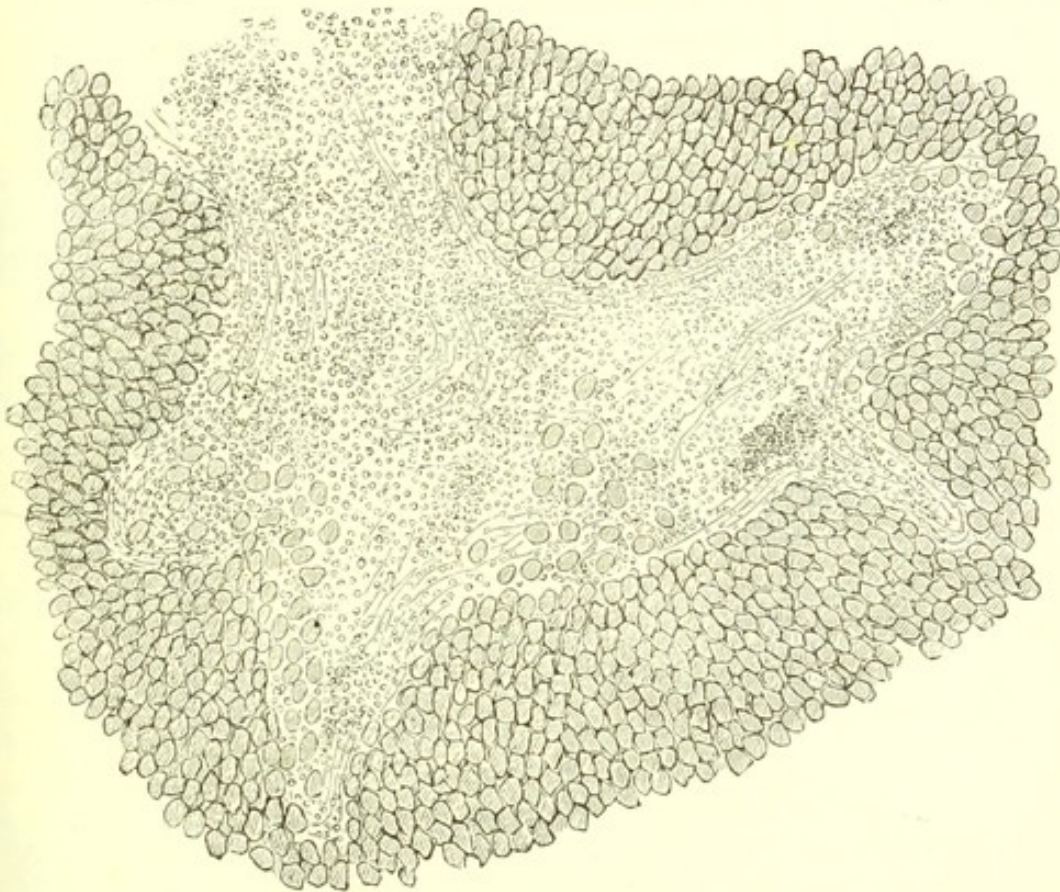


Fig. 292.—Cirrhosis of liver in earlier stage. The connective tissue is occupied by numerous round cells which are involving the peripheral parts of the lobules, the hepatic cells being here frequently isolated in the midst of the round cells.  $\times 75$ .

and hepatic duct. The newformation occurs in the great majority of cases only in these regions, that is to say, outside the lobules. In the earlier stages the affected connective tissue is abundantly cellular, like granulation tissue, and the process of newformation is evidently similar in its details to that in other chronic inflammations (see Fig. 292).



As a general rule the new-formed tissue undergoes development into dense connective tissue which has a tendency to shrink. By its shrinking it causes **Atrophy of the proper hepatic tissue**. As the cirrhosis is usually multilobular the contracting tissue isolates groups of lobules of larger or smaller size, and these, consisting of soft tissue, stand out somewhat when the liver is divided, or present themselves prominently at the surface. There are thus areas of remaining hepatic tissue which are being gradually encroached on by the connective tissue. The atrophy of the hepatic tissue takes place by the supply of blood being cut off. By the contraction of the new-formed tissue the portal vessels, and especially the interlobular branches, are narrowed. The hepatic artery is not much affected by the contracting tissue. Indeed, as the hepatic artery supplies the active connective tissue there is a new-formation of capillary blood-vessels in connection with its terminal branches. Injected preparations show that the hepatic artery is pervious, and that the new-formed connective tissue is highly vascular.

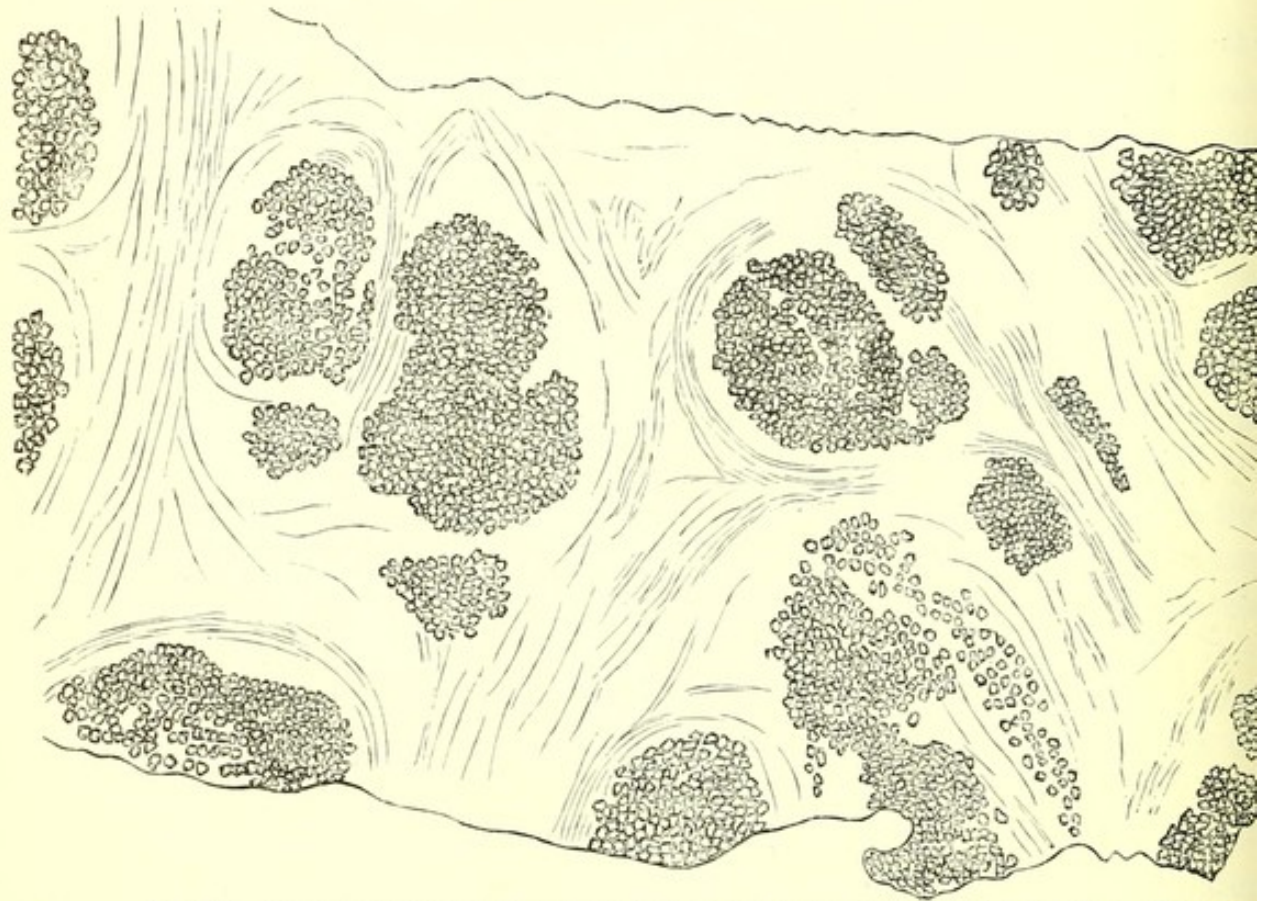


Fig. 293.—Cirrhosis of liver in an advanced stage, shown with a very low power. There are great areas of fibrous connective tissue, in the midst of which are islands of hepatic tissue having an opaque appearance from the presence of fat in the cells. 20.

The destruction of the hepatic cells takes place to a large extent by fatty degeneration, and one can often see, in the midst of connective tissue, islands representing hepatic tissue consisting of little more than collections of oil drops (see Fig. 293). The hepatic tissue is also com-



monly stained with bile pigment of a yellow or brown colour, and this we may associate with obstruction of the bile ducts by the contracting connective tissue. For the most part the pigment is biliary, but it is also to some extent blood pigment arising from the obstruction to the circulation. The connective tissue is also commonly stained, and to the naked eye the cut surface has a yellow colour, sometimes with here and there quite an orange tint. The name of the disease, cirrhosis, was originally applied from the colour of the altered organ.

A prominent feature in microscopic examination in many cases is the presence of narrow elongated canals, lined with epithelium, which are usually described as **Capillary bile ducts**. They are present in the midst of the connective tissue, sometimes very abundantly, and as no such ducts exist there normally, they are in a certain sense new-formed. By some authors (especially Charcot and Gombault) they are regarded as entirely new-formed, and their existence is taken as characteristic of a special form of cirrhosis. They are, however, to be found in all forms of cirrhosis, and they are probably to be regarded as the biliary capillaries, which have survived the destruction of the hepatic tissue. The larger bile ducts, which run in the capsule of Glisson, are unaltered, not being apparently affected by the shrinking connective tissue.

In its macroscopic appearances the liver varies considerably. In the earlier stages it is enlarged, but as a result of the changes detailed above it undergoes atrophy and distortion. The connective tissue contracts, and as it is irregularly distributed the contraction is irregular. The surface of the organ presents larger or smaller projections consisting of the less affected hepatic tissue between the cicatricial depressions. These projections may be comparatively large, forming the **Hob-nail appearance** of the surface, or they may be smaller, giving a **generally granular appearance**. They are usually yellow or brown and opaque, being fatty and stained with pigment. Sometimes there is special shrinking of the left lobe, and generally there is dragging in of the edges, so that the liver assumes more of a **compacted form**, being perhaps even thicker than usual, but reduced in superficies. It is dense to the feeling, and more tough to cut than normal. On the cut surface it may be possible with the naked eye to make out the grey connective tissue, with islands of opaque or pigmented hepatic tissue in it.

The terms **Biliary cirrhosis** and **Hypertrophic cirrhosis** are in frequent use, especially in French medical literature. The former of these was used by Charcot to indicate the mode of origin of the affection, the expression indicating that the irritation extended from the bile ducts. This form was distinguished in its causation from the hæmatogenous or venous, in which the irritation extended from the blood-vessels. The biliary form arises in connection with obstruction and inflammation of the bile ducts. In this form there are, according to Charcot and Gombault, two distinctive



features, namely, (1) that the newformation of connective tissue exists somewhat uniformly, so that the individual lobules are surrounded by it—the so-called **Monolobular cirrhosis**; and (2) that the capillary ducts already referred to are visible. To this last condition great importance has been attached.

Hypertrophic cirrhosis is a name given by another French author, Hanot, and it applies to the same class of cases. The biliary monolobular cirrhosis is characterized by hypertrophy of the liver.

It is to be acknowledged as a result of experiment that obstruction and inflammation of the bile ducts may lead to increase of the connective tissue, but further observation fails to confirm the view that a special form can be distinguished under the name biliary, or hypertrophic, or monolobular cirrhosis.

The ordinary alcoholic cirrhosis in its early stage may show considerable hypertrophy, the newformation may be monolobular in its distribution, and there may be the most marked newformation of capillary ducts.

The author is able to refer to a case which he examined post mortem. It was carefully watched for six weeks during life by Professor Gairdner. When first observed there was great enlargement of the liver and there was reason to believe that it had been greater. The patient had been addicted to alcohol for over two years. There were hæmorrhages from stomach and bowels, and marked ascites, but little or no jaundice. After death the liver was found small, weighing 43 oz., and finely granular on the surface. There was a great excess of connective tissue, which presented many round and spindle-shaped cells, and was monolobular in its distribution. The hepatic cells contained abundant fat in small and large drops. In this case the cirrhosis, although monolobular and at one period hypertrophic, was not connected with obstruction of the bile ducts.

Outside the liver the principal changes depend on **Obstruction of the portal circulation**. The connective tissue narrows the portal vessels in the liver, and this leads to a chronic passive hyperæmia in all the radicles of this vein, in the peritoneum, in the mucous membrane of the stomach and intestine, in the spleen, and so on. The most frequent consequence is ascites, but we also meet with hæmorrhages from the mucous membranes, especially of the stomach and large intestine, as well as catarrh of these. The spleen is also enlarged. Occasionally thrombosis of the hepatic vein results, and this may extend to the vena cava. A very large hæmorrhagic infarction of the lung was observed by the author as a result of embolism from a softened thrombus of this kind.

The obstruction of the portal vein often leads to widening of the venous channels which form communications between the portal radicles and the **Systemic veins**. In this way we may have great dilatation of the internal hæmorrhoidal veins (leading to piles) and of the hypogastric. The cutaneous branches of the latter often stand out prominently on the abdominal wall. The hæmorrhoidal veins also communicate with the vesical, and these latter may undergo dilatation. There may also be widening of communications with the veins of the diaphragm and œsophagus, which have sometimes been found highly varicose. Another channel occasionally met with is a small vein which runs from the portal



vein to the umbilicus. In some cases this is so much enlarged as to approach the size of the portal vein itself.

The patient generally dies from the disorders due to the continuous passive hyperæmia—the persistent catarrh of the alimentary canal, perhaps with hæmorrhages, the ascites, etc. There is great emaciation and sometimes icterus. It is not clear that the mere loss of the function of the liver bears an important part in the fatal result.

**3. Perihepatitis.**—This condition is an inflammation of the capsule of the liver, and it is always secondary to some other lesion. A chronic pleurisy of the right side often extends through the diaphragm, and causes inflammation of the subjacent peritoneum, including the under surface of diaphragm and upper surface of liver, so that firm adhesion is the result. This is often the case in phthisis pulmonalis. A general chronic or acute peritonitis usually involves the capsule of the liver as well as other parts of the peritoneum. Sometimes cirrhosis or syphilitic disease of the liver extends to the capsule.

The inflammation is in most cases chronic, and results in a thickening of the capsule. Sometimes there are shaggy papilliform projections from the capsule, and there is frequently adhesion to the parts around, especially to the diaphragm. The thickened capsule undergoes contraction like the new-formed connective tissue in other inflammations, and the result frequently is considerable deformity of the liver. The capsule contracting all round the liver doubles in its anterior edge, causing the organ to assume an approach to the globular shape. The contraction causes also atrophy of the liver tissue, and seriously interferes with the circulation in the organ.

**Literature.**—*Suppurative inflammations*—FRERICHS, l. c.; MURCHISON, l. c.; BUCKLING, 36 Fälle v. Leberabscess., 1868; KLEBS, Handb., ii.; BUDD, Dis. of liver, 2nd ed., 1852; DICKINSON, Path. trans., xxxii., 1881; FAYRER, Tropical dysentery and diarrhœa, liver abscess, etc., 1881; FINLAYSON, (with literature) Glas. Med. Jour., v., 1873. *Cirrhosis*—BUDD, FRERICHS, etc., l. c.; LIEBERMEISTER, Beiträge z. path. Anat. u. Klinik. der Leberkrankheiten; CHARCOT, Leçons sur les malad. du foie, 1877; HANOT, Cirrhose hypertrophique avec ictère., 1876; SAUNDBY, Path. trans., xxx., 1879; LEGG, (Expers. by ligature of bile duct) St. Barth. Hosp. Rep., ix., 1873; OSLER, Med. Times and Gazette, 1881; DRESCHFELD, Jour. of Anat. and Phys., xv.; BRIEGER, Virch. Arch., lxxv., 1879; KRÖNIG, (Cirrhosis from phosphorus-poisoning) Virch. Arch., cx., 1887.

## VI.—SYPHILIS AND TUBERCULOSIS OF THE LIVER.

**1. Syphilis.**—The changes in the liver in syphilis are very marked, and somewhat frequent. It manifests itself for the most part as an indurative interstitial inflammation, with or without the formation of gummata. There may be a **Cirrhosis** indistinguishable from the ordin-



ary forms of this disease, and only judged to be syphilitic from other circumstances. Sometimes in the midst of the general newformation of connective tissue there are numerous **Small gummata** about the size of peas, distinguished especially by their caseous condition.

For the most part, however, the disease is localized. To the naked eye the liver is seen to present one or several **Cicatrices** which often make deep indentations in its surface. These cicatrices are most frequent in the neighbourhood of the suspensory ligament. On cutting into the middle of the cicatrix a **Gumma** is usually divided with its central part caseous. The periphery of the gumma is not distinctly de-

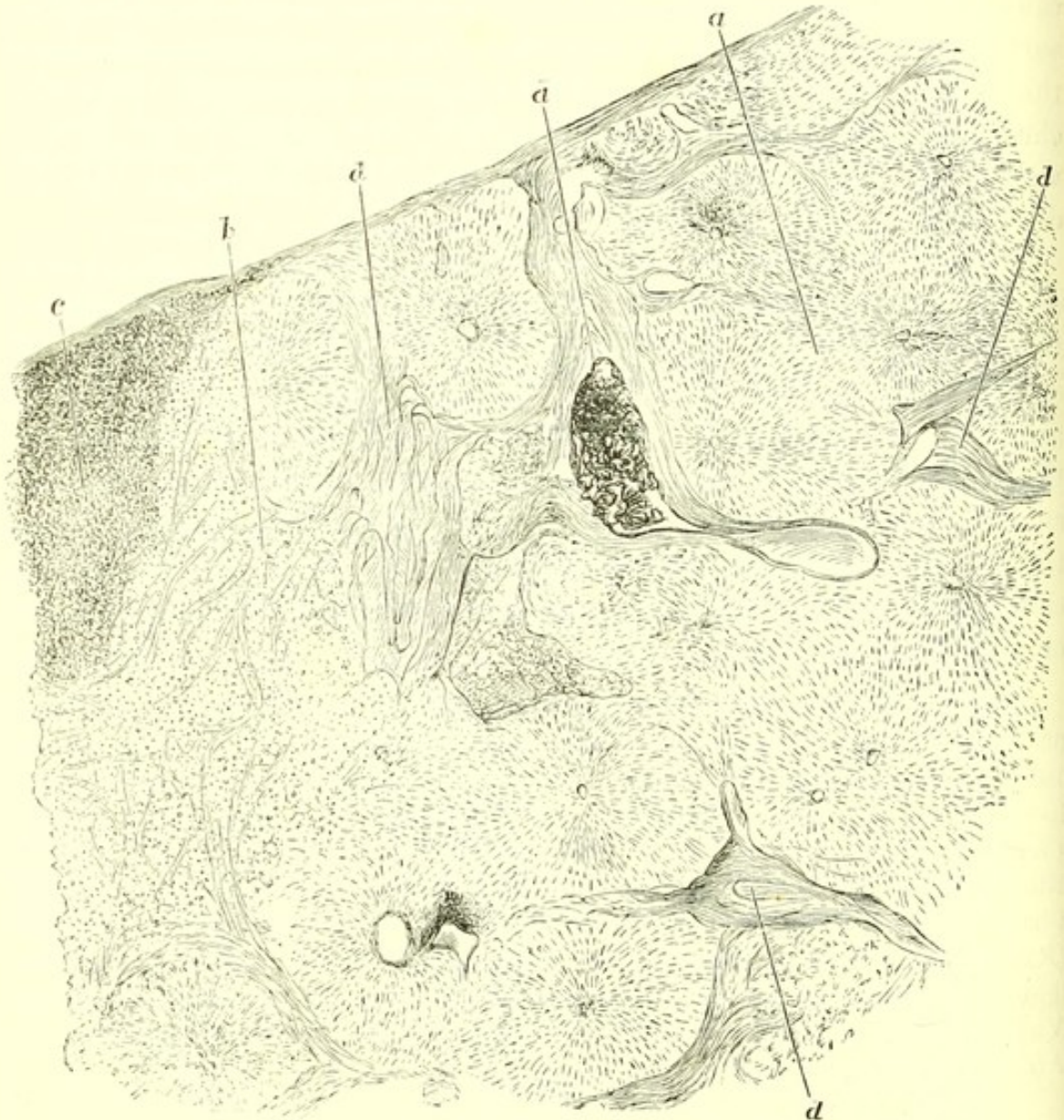


Fig. 294.—Gumma of liver. *a*, Normal hepatic lobules; *b*, recent tissue of gumma with dilated blood-vessels; *c*, caseous part of gumma; *d, d, d*, connective tissue outside gumma extending into the hepatic tissue.  $\times 16$ .

marcated from the surrounding connective tissue which extends outwards into the hepatic tissue. Other gummata may be found more deeply in the liver tissue, and these also are surrounded by cicatricial tissue.



The cicatrix is composed of fibrous tissue, and the gumma itself frequently presents irregular strands of connective tissue, but giving place to round-cell tissue. The central caseous portion contains merely granular debris (see Fig. 294). Sometimes at the periphery of the gumma there is great dilatation of blood-vessels, giving almost a cavernous appearance.

When there are a considerable number of gummata, the cirrhosis may be nearly continuous, but with special cicatricial contractions at intervals corresponding to the gummata. The liver in this way may assume a very striking appearance, as if subdivided into manifold small lobes. Be the cicatrices large or small in number, the lobed and subdivided condition is characteristic of the syphilitic liver.

The liver is sometimes found affected in infants who are the subjects of **Congenital syphilis**. It may be a general induration forming a cirrhosis, usually of the **Hypertrophic form**, with perhaps numerous small gummata; or the induration may be more localized, but never with that cicatricial condition which we find in acquired syphilis. The connective tissue formation in this case does not confine itself to the neighbourhood of the portal vessels, but extends into the lobules, so that between the cells and around the capillaries there is connective tissue, forming the so-called **Monocellular cirrhosis** of Charcot.

**2. Tuberculosis.**—Secondary tuberculosis of the liver is very frequent, while primary tuberculosis is exceedingly rare. In all forms of local tuberculosis as well as in acute general tuberculosis, tubercles are found in the liver. Hence the liver is probably more frequently the seat of tubercles than any other organ.

**Primary tuberculosis**, in the few cases which have been recorded, was in the form of solitary tubercles like those in the brain. In one recorded case the mass reached the size of the fist.

In **Secondary tuberculosis** the bacilli reach the liver by the blood, having been carried from the lungs, the bones, the intestines, or elsewhere. In ordinary local tuberculosis of these various organs a few bacilli reach the liver at intervals, and the conditions of the circulation allow of their settlement. The resulting tubercles are so small as to be scarcely visible to the naked eye. Even with the microscope search has to be made for them as they are usually comparatively few in number. If the liver is fatty, as is often the case in phthisis pulmonalis, the tubercles may be obscured by the fat. In this class of cases the tubercles are in various stages, most of them old and degenerated in their central parts. While tuberculosis is thus frequent, the tubercular growth scarcely ever attains large dimensions.

In acute miliary tuberculosis the bacilli are abundant in the blood



and the tubercles in the liver are very numerous, although still so small

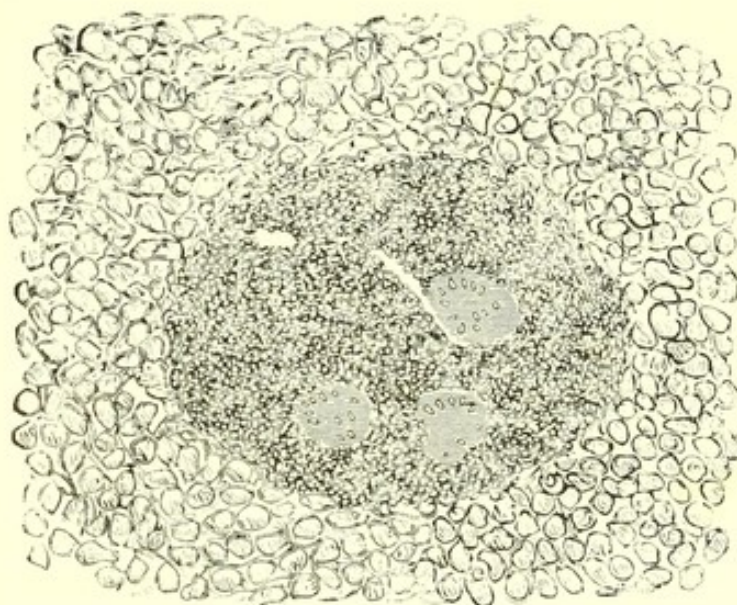


Fig. 295.—Tubercle of liver from a case of acute miliary tuberculosis. The tubercle is of a rounded form, and presents chiefly round cells with three giant-cells in the midst of these.  $\times 75$ .

as to be with difficulty seen with the naked eye. The tubercles here are usually found in an early stage and they present the typical structure, the giant-cells being generally very prominent (see Fig. 295).

A very rare form is **Tuberculosis of the bile ducts**. It occurs in the form of large caseous nodules with cavities in the centres which are filled with softened

matter coloured with bile. The wall of the cavities shows tubercles. It is possible that in this form the extension may be by the lymphatics from the porta of the liver.

Besides syphilis and tuberculosis we have **Glanders** manifesting itself in the liver, especially in horses. The nodules are moderate in size and tend to calcify. **Leprosy** produces nodules in the connective tissue. There are also newformations occasionally in **typhoid fever** and **diphtheria**.

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## VII.—TUMOURS AND PARASITES OF THE LIVER.

**Cavernous angioma** of the liver is the commonest form of primary tumour. Its structure has already been described (see p. 243 and Fig. 77). It develops apparently by dilatation of the existing capillaries and atrophy of the intervening hepatic cells, so that the tumour replaces a certain portion of hepatic tissue. When seated immediately under the capsule the tumours are visible as red blotches, and on section they have a red colour and collapsed appearance, and usually it is only after the blood is washed out that the white trabeculae become visible. The



tumours present great varieties of size and may be as large as the fist.

A case has been described by Ziegler of multiple **Fibroneuroma** in the liver on the fibres of the sympathetic, there being similar tumours in the various nerves of the body except the olfactory and optic.

**Cysts** are not infrequent in the liver, and they may be multiple. It is curious that multiple cysts of the kidneys and liver frequently co-exist, both forms being probably of congenital origin.

In **Leukæmia** the liver is very frequently the seat of newformations. This may take the form of a general infiltration of the interstitial connective tissue, but it often occurs as individual small nodules, somewhat resembling miliary tubercles but without the presence of giant-cells or any tendency to caseation. The round cells which form the nodules have been observed to show evidences of karyomitosis. The capillaries are commonly filled with leucocytes and the liver as a whole is much enlarged. Similar conditions are observed sometimes in Hodgkin's disease.

Concerning **Adenoma** of the liver there is considerable variety in statement and opinion, mainly because, on the one hand, this form of tumour is rare, and, on the other, there are tumours concerning which it is difficult to say whether they should be called cancers or adenomas. The form designated **Nodular hyperplasia** is clearly an adenoma, and the name may with convenience be confined to this form, as the other so-called adenomas have more the habit of cancers. It occurs as solitary or multiple tumours which, while perfectly defined, have the same structure as the proper hepatic tissue, the cells being usually larger and some having double nuclei. If single the tumour may reach the size of a cherry or larger; if multiple they are smaller. The larger ones are sometimes surrounded by a connective tissue capsule. These tumours are of no practical consequence and are met with accidentally. They are not uncommon in the liver of the dog.

**Primary cancer.**—This form of tumour is infrequent compared with secondary cancer, but in itself is not very uncommon. It occurs sometimes as a massive tumour involving a large part of the liver, but in addition to this large tumour there are frequently numerous secondary nodules in the liver substance. In some cases the growth seems to arise from the bile ducts at the porta of the liver, and to extend into the substance of the organ. In regard to structure several forms have been distinguished of which the principal are these:—

**Cylinder-celled epithelioma** is a rare form of tumour in the liver; it is sometimes described under the name of tubular adenoma. In its structure it closely resembles the similar tumours of the stomach



and rectum, consisting of tubular gland-like processes with cylindrical epithelium. As is usual with this class of tumours the cells in many of the alveoli are variously shaped and the typical structure is visible at the growing margins. There may be several tumours or one large one. The structure of the tumour suggests an origin from the hepatic ducts, but according to Rindfleisch the gland processes arise from the hepatic cells inside the lobules. This form of tumour is not usually malignant, but in a case recorded by Greenfield secondary extension to a lymphatic gland and to the lungs had occurred.

The primary cancer in the liver has more frequently the structure of **Ordinary cancer**, which sometimes has the dense characters of **Scirrhus**. It may involve a considerable portion of the liver, perhaps the whole left lobe, and by shrinking reduce it to small dimensions. In structure the tumour presents the ordinary characters. At the marginal parts it may be seen sending processes into the hepatic capillaries and into the branches of the portal and hepatic veins.

There are also cases of so-called **Infiltrated** or **Diffuse cancer** in which the whole liver is affected. The organ is enlarged and granular on the surface, resembling the appearances in cirrhosis. There is here also a great increase of connective tissue, but in its meshes there is cancerous and not liver tissue. This form is sometimes called **Cirrhosis carcinomatosa**. It is as if there had been a great transformation or replacement of the liver tissue by cancer.

Lastly there is the form in which the cancer originates in the **larger ducts** and extends along the capsule of Glisson. The radiating appearance of the newformation is visible in this form, and there is also obstruction of the ducts with retention of the bile and icterus.

**Secondary cancer.**—This is of very frequent occurrence in the liver. There may be a **direct extension** of a cancer from the gall-bladder or the stomach. In the latter case the organ becomes adherent and the cancer grows into the liver. There may thus be a large tumour, sometimes with a cavity communicating with an ulcerated surface in the stomach.

**Metastatic cancer** is more frequent. The primary tumour is usually in the stomach or intestine, but it may be in the œsophagus, uterus, mamma, or elsewhere. In most cases the cancer extends to the liver by the portal vessels (see further on as to Extension of Cancers in the Abdomen), but it may also occur by the hepatic artery in cases of cancer of the mamma and other external parts.

As the cancer is sown in every part of the liver, the consequence is the development of numerous tumours which may be found in all stages of growth. It is usual to find a large number of isolated tumours of circular form and pale colour (see Fig. 296). They are seated in the



liver tissue, but those near the surface produce rounded elevations which can often be felt through the abdominal parietes during life. Even to the naked eye the tumours show evidences of fatty degeneration in the appearance of an opaque yellow coloration. The absorption of the degenerated cells is also indicated by the partial contraction of the larger tumours in their central parts, producing in the superficial ones a dimpling or **Umbilication** of the rounded projections. The liver, as a whole, is sometimes enormously enlarged, weighing not infrequently as much as ten or twelve pounds.

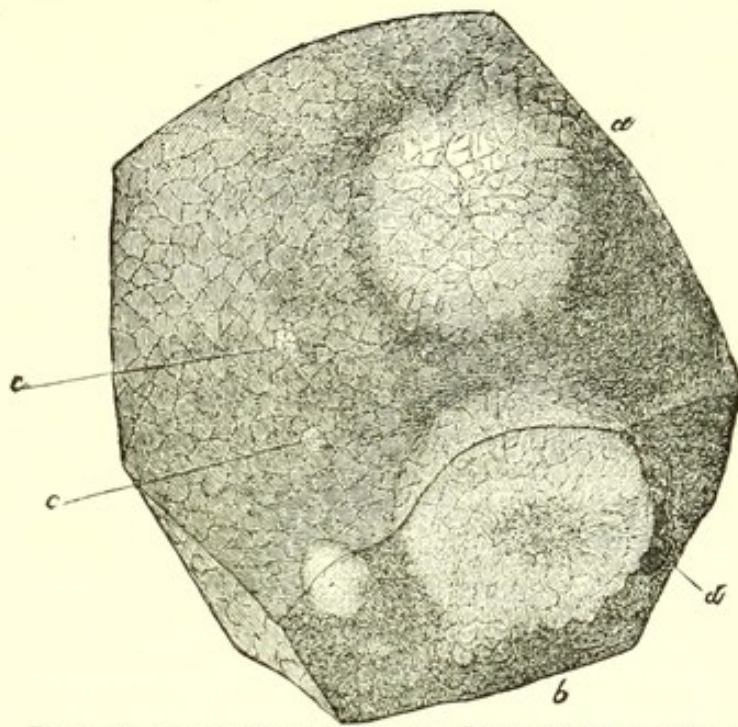


Fig. 296.—Piece of liver with secondary cancerous tumours in it. A larger one, *a*, is viewed from the surface. Another large one, *b*, in section. There are several smaller tumours, *c*, *c*. *d*, a vein in section. (VIRCHOW.)

In some cases the newformation is not so much in the form of individual tumours as a general **Cancerous infiltration**, as if almost throughout the liver a simultaneous development had occurred, and the cancerous tissue had grown vigorously, displacing the proper hepatic tissue.

In structure the tumours follow the primary ones. For the most part it is soft cancers of the stomach that form the original tumours, and we have a well-formed stroma with irregular masses of epithelial cells. But sometimes the primary tumour is an epithelioma, and in that case the tumours in the liver are usually fewer in number, firmer and more distinctly defined. When the primary tumour is in the œsophagus the structure may be that of the **Flat-celled epithelioma** with regular laminated capsules. When the primary tumour is in such a position that the cancer reaches the liver by the hepatic artery, the tumours are usually smaller in size and the liver not so much enlarged. This was at least very strikingly true in a case seen by the author, where the primary tumour was in the mamma.

The cancerous material as it is brought by the blood frequently develops **inside the vessels**, and particularly in the portal veins and the capillaries. The growing tumours cause atrophy of the proper hepatic tissue. The hepatic cells are often to be seen at the peripheral parts of



the tumours arranged in concentric layers as the pressure is exercised, and in various stages of atrophy.

The numerous developing tumours press on the bile ducts and obstruct them, and this occurs all the more as the growths arise by embolism of the portal vessels, and are, in the first place, related to them. As the ducts are in the immediate neighbourhood of these vessels, they are liable to be pressed on very soon. Hence **General icterus** is a common result, and there is also a special **Pigmentation** of the hepatic tissue (see ante).

The portal vessels suffer obstruction, not only by the pressure of the tumours, but as being the seat of their primary growth, and hence **Ascites** is a common result.

**Sarcoma** is of occasional occurrence in the liver in the spindle-celled and pigmented forms. Some cases of melanoid sarcoma have been described as primary tumours, but secondary tumours following melanotic sarcomas of the eye are more common. The liver may be largely occupied by these black tumours, and greatly enlarged. Here also the growth appears to occur in the blood-vessels and results in destruction of the hepatic tissue proper.

**Parasites of the liver.**—These have already been somewhat fully described. The most important

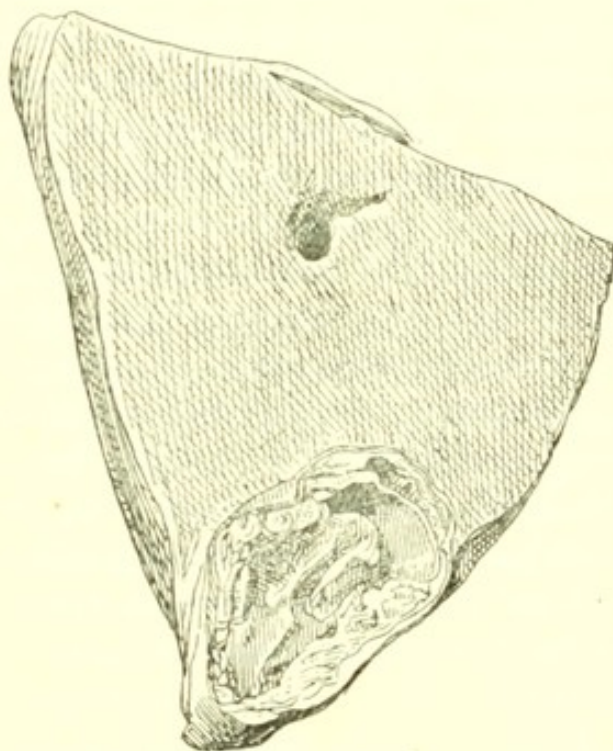


Fig. 297.—Portion of liver with collapsed hydatid cyst. Natural size.

constitutes the **Hydatids** of the liver arising from the **Tænia echinococcus**. When the liver is examined after death there is found a sac or sacs of very various size, up to that of a man's head. The larger ones produce necessarily great enlargement of the liver as a whole, and atrophy of the hepatic tissue around them. They present first a connective tissue capsule, inside which the proper wall of the cyst appears. As the vesicles are sometimes much broken down, and it may even be difficult to find hooklets, it is important to remember

that the proper cuticula of the cyst is lamellated (Fig. 134, p. 334). The multilocular and exogenous forms of hydatids are to be remembered the former resembling colloid cancer in its general appearance.



It is not uncommon to meet with the remains of a dead echinococcus in the liver. It shows a capsule which contains an indefinite debris in which pieces of the chitinous membrane are visible (Fig 297). There may be several such lesions present. In a case of this kind the author found in one of the cysts along with calcareous infiltration a production of true bone in the wall of a small collapsed cyst.

The *Distoma hepaticum, sinense*, and *lanceolatum* occur in the bile ducts, the *Distoma hæmatobium* in the portal vessels. The *Pentastoma denticulatum* also occurs in the liver.

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## B.—THE BILE-DUCTS AND GALL-BLADDER.

1. **Gall-stones.**—These are of very frequent occurrence, especially in people past middle life, and they are often found in the gall-bladder after death without their existence having been suspected during life.

The cause of their formation is very obscure. They are formed in the gall-bladder by the deposition of the constituents of the bile, chiefly cholestearine and next to that bile pigments, but also lime and magnesia salts in varying proportions. Probably stagnation of the bile in the gall-bladder at least predisposes to their formation, and this is rendered the more probable from their more frequent occurrence in old people where the actions are sluggish. In the centre of gall-stones there is frequently a nucleus composed of remains of epithelium or mucus, and it has been supposed that catarrh of the bladder may furnish this nucleus.

Gall-stones occur singly or in numbers. The **Single gall-stones** are commonly composed almost purely of cholestearine, of which there may be over 98 per cent. They are oval in shape, somewhat nodulated on the surface, and have a glistening appearance, altogether not unlike that of a sugar plum. When divided or broken they present a characteristic radiating appearance from the centre, and also sometimes a concentric stratification. The stone is very light in weight, floats in water, and has a soft almost soapy feeling.



**Multiple gall-stones** are more frequent, and although sometimes, nearly pure, they are more frequently composed of cholestearine mixed with bile pigment and lime salts. There may be two or three, but they may be present in large numbers, fifty, a hundred, or several hundreds or thousands (as many as 7800 have been counted). If few they may be comparatively large, but if numerous they are small, the bladder sometimes having the appearance of a bag filled with peas. The multiple stones are always faceted, taking their shape according as there is room, and fitting into each other. In appearance they have been compared to the macerated carpal bones, having somewhat similar facets, and often presenting a similar greyish colour, although sometimes yellow, brown, or even black. They are very light in weight, and on section present little of the glistening appearance or radiating arrangements of the single ones, being rather stratified, more deeply pigmented strata alternating with less.

Gall-stones composed almost entirely of **bile-pigment** are very rare. They are small, nodulated, and nearly black, and occur in considerable numbers at a time. Stones composed mainly of **Lime salts**, especially the carbonate, are still more rare.

The calculi may remain long in the gall-bladder, which may be found full of them, and with no bile in it. Its mucous membrane may be inflamed, and the calculi surrounded by an abundant mucous secretion.

Gall-stones frequently **leave the bladder** and pass into the cystic duct, where they may remain for a time and cause obstruction. After a time they often pass on into the ductus choledochus. The last part of this duct is narrow, and, if the stone is of any considerable size, it usually sticks here, at least for a time. By dilating the duct it may get into the duodenum, but very commonly it finds its way through by ulceration, and sometimes it ulcerates into the peritoneum, producing peritonitis. Lying at the mouth of the ductus communis it obstructs the outflow of bile and produces the results to be presently described. When it gets into the duodenum it usually passes off with the fæces, but if large, it may produce **Obstruction of the intestine** at some point. This is of rare occurrence, and can scarcely happen except in the case of large stones, chiefly of the solitary kind.

2. **Obstruction of the bile ducts.**—Obstruction occurs from various causes, of which one of the commonest is **Gall-stones**. **Inflammation** of the ducts sometimes produces obstruction, the inflammation being nearly always due to an extension of catarrh from the stomach and duodenum. As the duct near the orifice is narrow a trivial inflammatory swelling may produce an obstruction, which the bile, possessing a



low pressure, is unable to overcome. Then gall-stones themselves may produce inflammation, leading to a more or less prolonged closure of the ducts. We shall see afterwards that inflammation is not infrequently a consequence of obstruction, and these two should not be confounded. **Tumours and Inflammations around the duct** may cause obstruction. This is not infrequently the case with **Cancers of the head of the pancreas** or those involving the lymphatic glands in the portal region of the liver. We have already seen that an obstruction of the hepatic ducts in the substance of the liver occurs in cancer of the liver, in cirrhosis, etc.

The **Results of obstruction** vary according to the site of the obstruction.

If the **Cystic duct alone** be obstructed then the consequence is that no bile can get into the gall-bladder. In that case the bladder **may shrink**, and any mucus in it dry-in and perhaps afterwards become chalky. In many cases, however, there is an abundant secretion of mucus, and the bladder gets filled with it. The mucus often after a time gives way to a more fluid secretion, and the bladder may be converted into a thin-walled cyst (**Hydrops vesicæ felleæ**) which may be as large as the fist, with clear fluid contents.

When the gall-bladder is thus cut off and no longer available as a store for the bile, there sometimes occurs a **Dilatation of the larger bile ducts**, so that the bile may lie here instead of in the gall-bladder, and pass into the duodenum during digestion. This constitutes an imperfect compensation for the loss of the gall-bladder. Gall-stones may form in the dilated ducts.

In the case of **Obstruction of the ductus choledochus**, there is stagnation both in the gall-bladder and in the whole system of bile ducts. The stagnation tells first on the gall bladder, which dilates readily and stores up the bile. There may even be rupture of the gall-bladder from excessive dilatation. If the obstruction be prolonged great dilatation occurs throughout the whole system, and serious changes frequently result in the liver itself.

Next to the gall-bladder the ductus choledochus and the larger bile ducts, which are not supported by the firm liver tissue, are most liable to dilate. This dilatation may be very extreme, these ducts becoming sometimes as great in circumference as the thumb, and it may even go on to rupture.

Sometimes considerable atrophy of the proper tissue of the liver occurs, and we have the smaller bile ducts dilated and forming numerous cavities or **Cysts** throughout the liver.

There is sometimes even an **Acute inflammation** of the bile ducts



apparently from decomposition of the bile, and this may lead to **Biliary abscesses**. This occurs when the obstruction is incomplete, as when it is produced by the pressure of a tumour, and there is therefore the possibility of the propagation of septic decomposition from the duodenum to the stagnant bile. There may be numerous abscesses filled with a tenacious bile-stained pus.

3. **Rupture and Perforation**.—Rupture of the gall-bladder or of the ducts outside the liver may occur from over-distension, or it may be the result of injury. The rupture takes place into the peritoneum and the bile causes **peritonitis**. The inflammation is sub-acute causing great thickening of the peritoneum. A peculiar appearance is sometimes produced, the thickened peritoneum presenting an orange-coloured surface from biliary staining.

4. **Tumours of the bile ducts and gall-bladder**.—Of these the most important are the **Cancers**. We sometimes meet with primary cancer of the gall-bladder resembling in structure cancer of the stomach and intestine. By extension it may largely involve the liver tissue. We have already seen that some of the cancers of the liver probably take origin in the finer bile ducts in its substance.

### C.—THE PANCREAS.

The pancreas has the structure of a salivary gland, consisting of glandular acini whose ducts communicate with a main duct lying in the centre of the gland (Wirsung's duct) and opening into the duodenum in common with the ductus choledochus.

**Malformations** occur chiefly in the form of supernumerary or accessory glands. These are usually situated in the wall of the stomach, duodenum, or jejunum. Another malformation is that in which the pancreas surrounds the duodenum.

The pancreas is occasionally the seat of **Hæmorrhages** into its substance (*Pancreatic apoplexy*). These may be the result of injuries to the abdominal wall or the consequence of passive hyperæmia in diseases of the heart, lungs, or liver. Some cases of sudden death have been recorded by Zenker, Hooper, and Klebs, in which the most obvious lesion was hæmorrhage into the substance of this gland. No cause for the hæmorrhage has been apparent. It is supposed that the lesion caused pressure on the neighbouring coeliac plexus and semilunar ganglion and a consequent reflex paralysis of the heart, just as paralysis of the heart is producible by blows on the abdomen (Goltz's experiment).



**Inflammations** of the pancreas are of various kinds. We have suppurative pancreatitis like suppurative parotitis, resulting in abscesses; also indurative interstitial inflammation with loss of gland tissue, this condition being sometimes of syphilitic origin.

**Atrophy** of the pancreas is observed often as a part of general emaciation, but it has been found in a good many cases of **Diabetes mellitus**. It can hardly be held that the destruction of the function of the pancreas is the cause of the diabetes, and the probability is that, as Klebs suggests, the primary condition is disease of the celiac plexus causing vaso-motor paralysis (see p. 112).

**Fatty infiltration** of the pancreas is one of the commonest lesions of this gland. There is normally some adipose tissue in the midst of the gland, and this sometimes undergoes considerable increase, the proper glandular substance becoming atrophied, and adipose tissue taking its place, the shape and general appearance of the gland being preserved. The condition may be part of a general obesity, or it may occur in old age, and, in this latter case, it may be presumed that atrophy of the glandular tissue is the first condition, the adipose tissue developing afterwards, as in fatty infiltration of voluntary muscle. **Necrosis** sometimes occurs in the adipose tissue of the organ, producing opaque yellow patches of a striking character.

The glandular structure sometimes undergoes **Cloudy swelling** in common with that of the liver and other organs in the acute fevers. **Fatty degeneration** is also met with.

**Tuberculosis** is not common in the pancreas, but we meet with caseous masses having the characters of a local tuberculosis. More frequently the pancreas is involved secondarily in a tuberculosis arising in neighbouring lymphatic glands. Syphilitic **Gummata** have been observed.

Of the tumours of the pancreas, **Cancer** is by far the most important. It occurs most frequently in the head, rarely in the body or tail. It is mostly a dense tumour of fibrous appearance (scirrhous), but cases of soft and of colloid cancer have been seen. The cancer often obstructs Wirsung's duct, or produces still more serious results by obstructing the ductus communis. It may even by its retraction or by its prominence cause a partial obstruction of the duodenum. There may arise in this way considerable disturbances from the continuous extension of the tumour. We may also have secondary tumours in the lymphatic glands, liver, or peritoneum.

Cancers of the stomach or duodenum rarely extend to the pancreas. The gland is perhaps still more rarely the seat of secondary metastatic tumours when the disease becomes generalized.



The **Pancreatic duct** (Wirsung's) is liable to certain changes. **Concretions** occur in it, comparable to those of the salivary glands. They are mostly round or oval and white or greyish white. In size they have been met with as large as a hazel nut or larger, but they are usually small like grains of sand. They are composed chiefly of carbonate and phosphate of lime. They very often arise in dilated ducts, but if large may cause dilatation by obstructing the duct.

**Obstruction and Dilatation** of Wirsung's canal may arise, as we have seen, from calculi, cancers of the head of the pancreas, or from tumours in the neighbourhood. Dilatation also occurs secondarily to atrophy of the gland. The dilated duct forms a series of pouches, or else there is a more definitely localized dilatation so that actual cysts are formed. This latter will occur when the orifice is completely obstructed; the cysts may reach the size of the fist or that of a child's head, and these are sometimes designated **Ranula pancreatica**. The contents of the dilated duct may be simply the fluid secretion, but sometimes there is thickening of the contents and even hæmorrhage. In this way we may have coagula causing the cysts to look like aneurysms, all the more as the lining of the cyst may become the seat of calcareous plates like the internal coat of an artery in aneurysm.

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#### D.—THE PERITONEUM.

**Introductory.**—The peritoneum, which is stretched over so many different organs, and has so many recesses and pouches, is very liable to be affected by diseases having their source outside itself. It is a large lymph sac, and fluid is continuously circulating through it. The surface of the membrane is covered with endothelium, and there are innumerable apertures or stomata by which it communicates with lymphatic vessels beneath. Finely divided solid material introduced into the peritoneal cavity is very readily absorbed and carried into the lymphatics. The transudation fluid which normally passes out of the blood-vessels is doubtless absorbed by the stomata throughout the



peritoneum, but there are two localities in which its absorption presents points of peculiar interest.

From certain facts to be afterwards referred to in connection with tuberculosis and cancer of the peritoneum, it may be inferred that the **Great omentum** is specially concerned in the process of absorption. This double layer of peritoneum, lying free in the cavity, may be regarded as a **drain** by means of which the fluid is drawn off.

Again, the lymphatics of the **Diaphragm** communicate, on the one hand with the peritoneal sac and on the other with the pleura, so that fluid and finely divided solids may be carried through from one to the other. It is probable that the general course of the current is from peritoneum to pleura, although it may be reversed.

This great power of absorption is very important, especially in relation to septic processes. The products of septic decomposition are readily absorbed; sometimes in such quantity as to produce fatal results before they have had time to induce any considerable local effects. This is especially the case in rupture of the intestine, where death may occur within twenty-four hours without definite symptoms of peritonitis, and apparently from absorption of the septic poison. After death, however, signs of inflammation are usually visible in the peritoneum.

The fluid in the peritoneal sac is not at rest, but circulates, and the movements of the intestines doubtless have to do with its transportation from place to place. Hence any pathogenic agent introduced into the peritoneal cavity is generally carried to every part of the sac and produces its effects in every region. Abundant examples of this are afforded by tuberculosis of the peritoneum, inflammations, and so on.

1. **Malformations.**—The mesenteries are sometimes too long or too short. The former condition is supposed to have to do with the causation of hernias. The latter causes the intestine to be unduly controlled in its movements. In a case recorded by Lawson Tait the peritoneum passed from loop to loop without any proper mesentery. There are also dermoid cysts found in the peritoneum of congenital origin.

2. **Disorders of the circulation in the peritoneum.**—(a) **Active hyperæmia** is produced when from any cause a general relaxation of the arteries in the sac occurs. Leaving inflammation out of account, this will hardly occur except as a result of sudden removal of extra pressure from these vessels. If a large ovarian tumour be removed from the abdomen, or ascitic fluid drawn off, the arteries, which have previously accommodated themselves to the undue pressure on their walls, will relax, and hyperæmia occur. The hyperæmia may result in the occurrence of a peculiar form of **Chronic hæmorrhagic peritonitis**, comparable with



hæmorrhagic pachymengitis (see p. 594), the effused blood being here also sometimes a prominent feature (**Hæmatoma of the peritoneum**). As in the other case it may be a question whether there is an inflammation preliminary to the hæmorrhage, or whether the hyperæmia induces bleeding, the succeeding organization of the clot leading to the formation of a membranous layer on its surface. Where ascitic fluid has been drawn off many times, there may be several layers of soft membrane on the surface of the peritoneum, the innermost being the most delicate and the newest.

(b) **Passive hyperæmia** occurs as a result of obstruction to the portal circulation, either alone or as part of a general venous hyperæmia. It is chiefly important in relation to ascites, of which it is the most frequent cause.

(c) **Hæmorrhage**.—There may be unimportant hæmorrhages in the substance of the peritoneum in scurvy, hæmophilia, etc. More considerable hæmorrhages occur in consequence of rupture of considerable vessels. Aneurysms of the abdominal aorta not infrequently rupture. The blood accumulates at first behind the peritoneum, but it may also pass into the cavity. Traumatic rupture of liver, spleen, or kidneys also causes hæmorrhage. Then there are frequent hæmorrhages from the female generative organs, as from the rupture of the cyst in extra-uterine pregnancy, but also without apparent cause at menstrual periods. (See Peri-uterine Hæmatocele.) There are also hæmorrhages in tuberculosis and cancer of the peritoneum.

The effused blood is in many cases readily absorbed, but when the collection of blood is local, as in the pelvis, or the peritoneum is altered by inflammation, it may remain. If there is a considerable mass the name **Hæmatoma** is applied. The blood becomes decolorized and surrounded or encapsuled by new-formed connective tissue in the ordinary way. The blood thus sets up a chronic inflammation which, in the case of the pelvic hæmatoceles, may have serious consequences by interfering with the uterus and ovaries.

(d) **Ascites**.—This name is given to dropsy of the peritoneal cavity. It is sometimes merely a part of a general œdema, occurring in disease of the heart or lungs, in Bright's disease, and in anæmic states. But it is peculiarly prone to occur when the portal circulation is specially obstructed. This may happen by the portal vein being obstructed by thrombosis, or pressed on from without, but more frequently it is by some lesion in the liver itself, such as cirrhosis or cancer, which obstructs the portal vessels, as it were, in detail.

In these cases the ascites is from increased transudation, and it is much less common to have it occurring from interference with absorp-



tion. This may have something to do with it, however, in cases of cancer of the peritoneum, where the numerous cancerous tumours originate from material carried into the lymphatic channels and growing there. These tumours will necessarily obstruct the lymphatics to a large extent in detail, although they may also produce by their irritation a hyperæmia of the peritoneal vessels.

The fluid in ascites is contained in the general sac of the peritoneum, but where adhesions have previously existed it may be confined to particular parts, and sacculated. Sometimes also, in children, there is a dropsy in the sac of the omentum (*Hydrops omenti*).

The character of the fluid in ascites is that of ordinary transudations, a clear, slightly yellow, limpid fluid of low specific gravity. After it has stood for a time it often deposits a very gelatinous coagulum of fibrine.

In prolonged ascites the peritoneum is apt to get somewhat thickened, especially when puncture has been frequently performed. The great omentum is not infrequently gathered up so as to be thicker and shorter than usual, and in that case it will act less efficiently as a drain.

**Chylous ascites** has already been referred to (p. 434) as occurring in connection with obstruction of the thoracic duct. The fluid in the abdomen is milky, containing finely dissolved fat. Of a different nature is **Ascites adiposus**, which may be confused with chylous ascites, as in it also the fluid is milky. In this case the milkiness is from fatty degeneration of cells, either the endothelium in ordinary ascites or the cells in cancer. There is always the distinction between the two conditions that in chylous ascites the fat is free, while in ascites adiposus it is in cells.

**3. Inflammations of the peritoneum. Peritonitis.**—Inflammation of the peritoneum is very seldom spontaneous in its origin. It seems remarkable that, compared with the pleura or pericardium, this membrane is so seldom the seat of independent inflammation as a result, for instance, of the irritation of the blood in acute rheumatism, or of the more vague causes of irritation designated as cold.

The peritoneum is, however, peculiarly liable to inflammations of a secondary character, the irritant proceeding either from without as in wounds of the abdomen, or from one of the organs lying beneath the membrane.

Mere exposure to the air or the entrance of air into the abdominal cavity does not induce peritonitis, and even a somewhat prolonged cooling of the membrane, as during an operation, does not seem to lead to inflammation.



**Septic inflammations** comprise the great proportion of acute inflammations of the peritoneum. When septic matter is introduced, the peritoneal cavity forms a favourable place for the propagation of any microbes which are not inhibited by the lining membrane. Of these the pyogenic micrococci, and especially the streptococci, are the principal forms. They multiply with enormous rapidity in the warm and moist cavity, and it often happens that in comparatively few hours we may have an intense inflammation, or we may have such an absorption as to cause death by septicæmia before considerable inflammation can develop.

The septic inoculation may take place by a wound in the abdomen, made by accident or by an operation. More frequently the source is an underlying organ, as by rupture of the stomach or intestine, or by propagation of septic processes from the uterus after delivery. In regard to the last mentioned source, acute peritonitis is often a special feature in so-called puerperal fever.

The septic inflammations are pre-eminently acute, and tend rapidly towards suppuration. At first there is hyperæmia and a serous and **Fibrinous exudation**. The exuded fibrine is visible on free surfaces as a soft yellow layer, and is often present in the fluid as yellow flakes. It glues together surfaces which are in contact, such as the loops of the intestine, but the adhesions are soft and readily separated. As the inflammation goes on, the fibrinous exudation, which from the first contains very numerous leucocytes and is correspondingly soft, becomes still more infiltrated with these, and assumes the characters of pus. **Pus** may be found in some parts, while in others there is still the soft fibrinous exudation. Thus pus may be found in the neighbourhood of the original source of the inflammation, as around the vermiform appendage, the inflammation being here more intense or of longer standing. The pus, and even any free fibrine that may exist, commonly gravitate to dependent parts, and we may find a collection of yellow pus in the pelvis, especially in Douglas's pouch.

The endothelial cells of the peritoneum take part in the inflammation. They multiply and enlarge (Orth), or they are shed. The underlying connective tissue is infiltrated with serous fluid and exudation, and all the underlying tissues are altered, more especially the wall of the intestine, whose coats are often oedematous and swollen. There is not infrequently considerable tympanitic distension of the intestine from paralysis of its muscular coat. This **Meteorism** is sometimes a peculiarly distressing feature in puerperal fever.

Septic peritonitis, if general, is almost necessarily fatal. Sometimes it is localized by adhesions, and even after the occurrence of suppuration may subside and give place to chronic inflammation.



A more **localized acute peritonitis** not infrequently occurs. It may be in connection with typhlitis (see ante), or with diseases of the uterus, or may be even a result of infarction of the spleen. Such inflammations are usually survived with the result of producing adhesions.

**Chronic peritonitis**, whether developing out of the acute form or occurring in connection with disease in an underlying organ, is characterized by newformation of connective tissue, frequently with adhesion of opposing surfaces (*Peritonitis adhæsiva*). The details of this process are similar to those in chronic pleurisy; it remains here to specify some of the more common occasions of the affection.

A diffuse chronic peritonitis sometimes develops in the course of Bright's disease. There is also commonly, in cases of secondary cancer of the peritoneum, a general chronic peritonitis.

Local thickenings of the capsule of the liver and spleen are of frequent occurrence in connection with diseases in these organs or their neighbourhood. Sometimes the connective tissue is hard, almost like cartilage. Very commonly there is adhesion to the parts around, especially to the diaphragm. On the other hand, the diaphragm may be adherent by reason of the extension of an inflammation from the pleura, the irritant having passed downwards in a direction contrary to that of the usual circulation.

The peritoneum around the female generative organs is liable to very frequent local chronic inflammations (*Perimetritis*), resulting in complex adhesions and mattings of the pelvic organs. The contraction of the new-formed connective tissue may cause considerable distortion of these organs.

There is also a peritonitis from rupture of the gall-bladder or a bile duct (see p. 786). Tuberculosis is the commonest cause of Peritonitis.

**4. Tuberculosis of the peritoneum. Tubercular peritonitis.**—This disease is due to the existence of the tubercular virus in the peritoneal cavity. The virus seldom gets into the sac from tubercular ulcers of the intestine, apparently because the intestinal lymphatics are sub-peritoneal and do not connect with the interior of the sac. Tuberculosis of the vertebræ or of lymphatic glands may give rise to it, and in some cases tuberculosis of the testicle and vas deferens has extended to the peritoneum, the disease being in that case concentrated in the inguinal region where the vas deferens approaches nearest to the peritoneum. In most cases, however, no direct extension from a neighbouring locality is traceable and the tuberculosis is primary.

Having reached the peritoneum the virus is carried hither and thither throughout the sac by the regular circulation. The consequence is the formation of innumerable tubercular nodules and an **Inflamma-**



tion of the peritoneum. The inflammation is at first acute, accompanied by serous and probably fibrinous exudation as in septic inflammations. There is thus often considerable swelling of the abdomen. In some cases the inflammation is unusually acute and may even be suppurative in character.

By the time the case comes to be examined post mortem the acute stage has usually passed off and we find evidences of chronic inflammation in the form of thickening of the peritoneum and multiplied vascular adhesions in every part. The loops of the intestine are adherent to each other, and the superficial ones to the anterior wall of the abdomen, the omentum is adherent to the intestine, the liver to the diaphragm, and so on. In fact the peritoneal cavity is obliterated by adhesions. In the midst of these adhesions are numerous yellow masses of very various sizes, some as large as split-peas, and usually flat. These caseous masses are composed of groups of tubercles which have very much the character of those found in tubercular pericarditis. The caseous tubercles have developed in the usual way out of grey miliary tubercles, and examination will usually show examples in the various intermediate stages.

The condition of the **Omentum** is worthy of special mention. It is drawn together and thickened, and closely adherent to the intestine and wall of the abdomen, while in its substance numerous tubercular masses are to be found.

All these conditions indicate a chronic inflammation, accompanied as usual by the newformation of vascular connective tissue with consequent adhesion. The yellow caseous masses are collections of tubercles mostly obsolete, just as the caseous tubercles of the brain are; and here, as there, we may find recent tubercles at the margins of the caseous masses.

It has already been mentioned that **Tubercular pleurisy** often develops in association with tubercular peritonitis. There is in the pleura for the most part a serous exudation, and as the eruption is usually recent the tubercles are in the form of small white or grey nodules. They are commonly grouped mainly in the lower part of the pleural cavity, in this way indicating the source of the infective material. They may set up an acute pleurisy with fibrinous exudation.

**Healing** is probably not infrequent in tubercular peritonitis, but the resulting conditions are rarely the subject of observation. The author had the opportunity of examining a case ten years after the occurrence of the disease. The peritoneum was obliterated by soft connective tissue adhesions, which united the intestines and the various organs together. There were no tubercles visible, but here and there a small



cretaceous mass in the midst of the adhesions. When the tuberculosis had been overcome the dead caseous matter had remained. This was in great part absorbed as dead animal matter. Where, from the size of the mass or otherwise, absorption did not occur, calcareous infiltration took place. In another case the adhesions had formed several bands or bridles, under one of which the intestine had become incarcerated, thus leading to a fatal obstruction.

Tuberculosis of the peritoneum is sometimes met with in **Acute miliary tuberculosis**, but it is not frequent, and the appearances are altogether different to those of tubercular peritonitis. In the case of general tuberculosis the virus is in the blood, and the tubercles develop in connection with the blood-vessels and not on the surface. The tubercles are very small grey nodules hardly visible to the naked eye and specially abundant in the upper part of the abdomen and omentum.

5. **Tumours of the peritoneum.**—These are rarely primary. **Retro-peritoneal sarcoma** is not uncommon. The tumour, sometimes growing to a very large size, pushes the organs before it and may infiltrate them.

Sometimes we meet with bulky gelatinous tumours in the abdomen, and the recognition of the exact nature of some of them is matter of considerable difficulty. Colloid cancer of the stomach and intestine not infrequently, as we shall see afterwards, passes on till it reaches the peritoneum, and may result in the formation of bulky gelatinous masses there. But, besides that, there are primary tumours of the peritoneum which belong to the class of **Cylindroma** or **Plexiform angiosarcoma**. In these cases there is a newformation of blood-vessels in whose adventitia is produced a peculiar gelatinous tissue. These tumours may attain a large size, weighing as much as forty pounds.

**Primary cancer** occurs with similar characters to that of the pleura, and like that form it is sometimes called endothelioma. The tumour is in the form of nodules of larger and smaller size along with great thickening of the peritoneum. There is also great serous effusion and usually also fibrinous deposition. There may be blood in the exudation.

**Secondary cancer** of the peritoneum will be considered in the next section.

**Malignant lymphoma** (Hodgkin's disease) is not such a common tumour here as in the mediastinum, but it sometimes originates in the lymphatic glands of the mesentery and involves all the neighbouring structures. We may thus have bulky tumours occupying the place of



a portion of the mesentery and intestine, and repeating roughly the anatomical relations of these.

**Literature.**—*Malformations*—LAWSON TAIT, *Dubl. Jour. of Med. Sc.*, 1869; *Obstet. Jour.*, iii. *Hæmorrhages*—RECKLINGHAUSEN, *Virch. Arch.*, xxvi., 1863; CORDUA, *Resorptionsmech. von Blutergüssen*, 1877; FRIEDREICH, *Virch. Arch.*, lviii., 1873; BÄUMLER, *ibid.*, lix., 1874. *Chylous ascites*—QUINCKE, *D. Arch. f. klin. Med.*, xxx., 1882; COATS, *Museum Catalogue of Western Infirmary*, pp. 55, 56 (two cases from thrombosis of jugular), 1885; LETULLE, *Rev. de Méd.*, 1884; WHITLA, *Brit. Med. Jour.*, 1885, i. *Inflammation*—KLEIN, *Anat. of lymph. syst.*, i., 1873; ORTH, *Virch. Arch.*, lviii., 1873; FRAENKEL, *D. med. Wochenschr.*, 1884; BAUER, in *Ziemssen's Cycl.*, xiv., 1878. *Tuberculosis*—KLEBS, *Virch. Arch.*, xlv., 1868; PAYNE, *Path. trans.*, xxi., 1870; BAUMGARTEN, *Zeitschr. f. klin. Med.*, x., 1885; GRAWITZ, *Charité-Annalen*, xi., 1886. *Primary cancer*—BRISTOWE, *Path. trans.*, xxi., 1870; NEELSEN, *D. Arch. f. klin. Med.*, xxxi., 1882; BIEGER, *Charité-Annalen*, viii., 1883. *Lymphoma*—WICKHAM LEGG, *St. Barth. Hosp. Rep.* xi., 1875.

#### E.—THE SECONDARY EXTENSION OF CANCERS OF THE ABDOMINAL ORGANS.

We have seen in the study of the diseases of the stomach and intestine that the cancers of these organs very often lead to secondary tumours in the liver and peritoneum, and it may be well to consider here more systematically what paths the infective material follows in passing from the primary tumour to the seat of the secondary growths.

**The secondary growths in the liver**, in the case of cancers of the organs mentioned, form in connection with the portal vessels, and there is no doubt that the material is brought to the liver by the portal vein from its radicles. But the question remains, How does the cancerous material find its way into the radicles of the portal vein? We know that in external cancers the secondary tumours occur uniformly in the lymphatic glands, and it is only after these have been long involved that the cancerous material reaches the blood.

It seems probable that cancers of the abdominal organs form no exception to this rule, and that extension to the liver is usually a late and properly a tertiary phenomenon, although the concealed and protected position of the primary tumour and of the secondary lesions in the lymphatic glands renders this difficult of demonstration.

Supposing this view to be correct, then it follows that, if a cancer of the stomach or intestine causes secondary growths in lymphatic glands whose **Veins are not radicles of the portal**, the tumours of the tertiary order would not be in the liver but in the lungs, or beyond the lungs, in organs fed by the systemic arteries.

The author met with a case of cancer of the stomach in which, instead of the glands immediately outside its wall, as is usually the case, the pre-vertebral glands



were enlarged and cancerous. One of these was adherent to the wall of the inferior vena cava, and on opening this vein a little white thrombus was seen peeping out of a small branch which emerged from the enlarged gland into the vein. There were cancerous thrombi in other veins within these glands, and on microscopic examination it was found that the cancerous tissue in the glands had largely broken up the veins, and epithelial cells were found in them along with the blood. In this case there were innumerable cancerous embolisms in the lungs.

This case would seem to indicate that cancers in lymphatic glands, by breaking up the gland, penetrate into the venous radicles in the gland, and so pass into the general circulation. It seems a legitimate inference from this case that when the liver becomes involved in cancer of the alimentary canal, it does so by the portal blood becoming infected through the lymphatic glands.

There remains one possible difficulty in the way of accepting this view. In external cancers it is exceptional for the general circulation to become infected. The disease generally goes no further than the lymphatic glands, whereas in cancers of the abdominal organs the liver is affected in a large proportion of the cases. But in cases of external cancer the patient usually suffers from ulceration of the primary or secondary growth or of both, and dies before the infection has reached the general circulation. In the case of abdominal cancers, however, the organs themselves, and the lymphatic glands, are protected by their position, and the cancers are not so apt to interfere with the general health as external cancers are. If the history of even an extensively ulcerating cancer of the stomach be compared with that of a cancer of the mamma, the difference will be very apparent. If the cancer of the stomach does not produce vomiting or stricture of the pylorus, there may be for a long period very little disturbance of the general health, and little more than symptoms of dyspepsia. It seems probable, from the history of some cases, that a cancer of the stomach may go on for many years without causing death. The abdominal lymphatic glands are still more protected. They practically never ulcerate, and, in relation to direct injury to health, cancer in them is of little account.

It seems probable, then, that in cases where the liver is affected the disease is of much longer duration in its primary seat than is often suspected. Cases of multiple cancer of the liver are often examined after death, in which no suspicion has existed during life of the existence of a primary tumour in the stomach, and this tumour may possibly have been going on for a period whose duration cannot in any way be gathered.

**Cancerous infection of the peritoneum** also occurs by extension of cancer from the abdominal organs. The seat of the cancer may be any of the abdominal organs; the peritoneum becomes infected when the cancerous material finds its way into the cavity.

There are some cancers which have comparatively little tendency to extend along the lymphatics to the glands, but prefer to insinuate themselves among neighbouring structures, and advance by continuity of tissue. This applies especially to **Colloid cancer**, which often grows through the wall of the stomach or intestine, while the glands are hardly at all affected. We can understand that a cancer with such a rigid stroma as this form has, and with cells which so readily swell up and become transformed, will not readily allow of transportation of its



elements. But this form of cancer very readily, after growing through the wall of the stomach or intestine, infects the peritoneum, and there is no form of cancer which, in such a large proportion of cases, produces secondary tumours there.

The **Cancers of the ovary** being already very close to the peritoneum, readily produce cancerous infection, and do so in almost every form of cancer. Cancers of the **Pancreas** also frequently have a similar course for the same reason. The ordinary cancers of the alimentary canal more rarely pass through the walls and infect the peritoneum, but they sometimes do. It is besides not uncommon to meet with secondary cancers of the **Liver** which have produced an infection of the peritoneum, some of the tumours of the liver having reached the surface and extended through the capsule.

When the cancerous material gets into the peritoneum it is carried throughout it by the circulating fluid, aided by the movements of the intestine, and secondary cancerous tumours commonly spring up in the most diverse regions (Fig. 298). It is to be remembered that in the

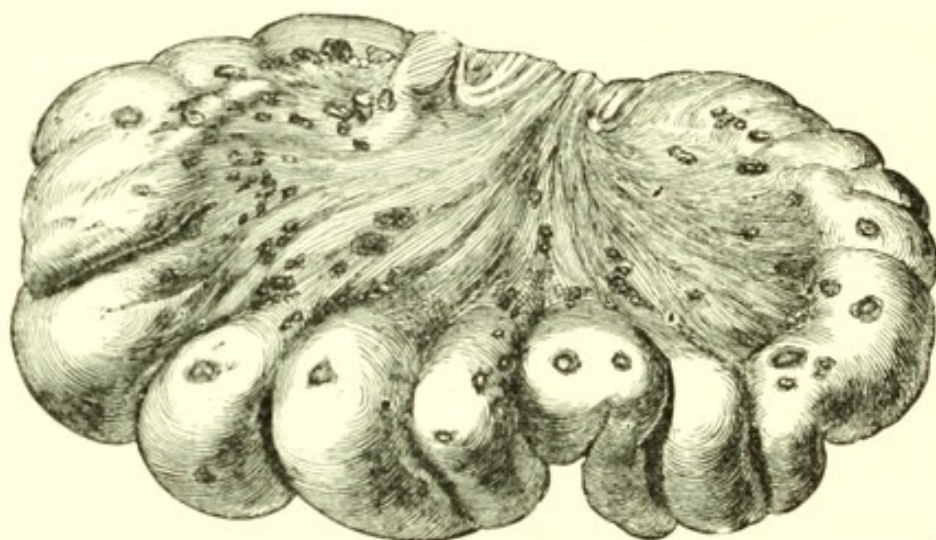


Fig. 298. —Disseminated cancer of the peritoneum, from cancer of the stomach. (VIRCHOW.)

peritoneum there are innumerable open stomata ready to absorb any finely divided solid matter that may be suspended in the peritoneal fluid. The infective material will therefore be carried from the surface into the substance of the peritoneum, or into the subperitoneal tissue, and the resulting tumours are really beneath the surface. They form usually flat growths with a smooth surface, the general surface of the peritoneum being perhaps unbroken. Not infrequently the tumours are continuous with one another in some parts of the abdominal wall, a layer of cancerous tissue appearing like a subperitoneal thickening.

The **Great omentum** is somewhat peculiarly situated in this respect.



We have seen that it probably acts as a kind of drain in the peritoneal cavity, and if this be the case it will specially absorb any material which gets into the cavity. In accordance with this there is usually in cancer of the peritoneum great newformation in the omentum. In colloid cancer it sometimes assumes the form of a bulky heavy mass, and in other forms we have it gathered up and converted into a solid tumour lying transversely in the abdomen. We may venture the statement that this fact is too little known among physicians, and that a great omentum thus altered is frequently taken during life for an enlargement of the liver, or a primary tumour of some obscure kind.

The **Appendices epiploicæ** are also, apparently, highly absorbent, and cancerous tumours are sometimes found in them.

The relation of peritoneal cancers to the **Diaphragm** presents some points of interest. We have seen that the diaphragmatic lymphatics communicate with the peritoneal sac on the one hand, and the pleural sac on the other. In peritoneal cancers the diaphragm is usually permeated with cancerous growths, and these are often in the form of cords as if following the course of the lymphatics. Through time they extend to the pleural surface, and tumours may appear there. If there are no pleural adhesions in this region the infective material passes into the pleural cavity, and numerous tumours are often found, especially in the lower parts of the pleura. A pre-existing adhesion of the diaphragm to the lung prevents this extension of the cancer.



## SECTION VIII.

## DISEASES OF THE URINARY ORGANS.

- A. THE KIDNEYS AND URETERS—*Introduction as to structure and function.* I. MALFORMATIONS AND MISPLACEMENTS—1. *Congenital malformations.* 2. *Variations in position, chiefly the moveable and the floating kidney.* II. HYPERTROPHY, *mainly compensatory.* III. DISORDERS OF CIRCULATION, *chiefly hyperæmia and embolism.* IV. BRIGHT'S DISEASE—1. *Causation.* 2. *Forms,* (a) *Parenchymatous or tubular nephritis; changes in glomeruli, epithelium, etc. The large white kidney. The contracted fatty kidney;* (b) *Interstitial nephritis; changes in interstitial tissue, glomeruli, etc.; cystic formation.* 3. *Character and origin of tube casts.* 4. *Other phenomena,* (a) *Albuminuria,* (b) *Uræmia,* (c) *Edema and dropsy,* (d) *Changes in heart and arteries; hypertrophy of left ventricle; changes in arteries—arterio-capillary fibrosis, etc. Theory of vascular changes.* V. SEPTIC INFLAMMATIONS—1. *Embolic or pyæmic forms;* 2. *Pyelitis and pyelonephrosis.* VI. HYDRONEPHROSIS AND PYONEPHROSIS. VII. RETROGRADE CHANGES—1. *Amyloid disease, its various origins; mostly with interstitial nephritis,* 2. *Atrophy, sometimes local from affection of arteries,* 3. *Parenchymatous infiltration.* VIII. CONCRETIONS AND CALCULI; *in the new-born and in adults.* IX. SYPHILIS AND TUBERCULOSIS; *chiefly local tuberculosis or renal phthisis.* X. TUMOURS—*Cysts, including cystic degeneration; Sarcomas and Cancers.* XI. PARASITES—*Echinococcus and filaria.*
- B. URINARY BLADDER AND URETHRA. 1. *Congenital malformations.* 2. *Perforation and rupture of bladder.* 3. *Dilatation and hypertrophy, including diverticula.* 4. *Disorders of circulation.* 5. *Inflammation of bladder, Cystitis, chiefly from decomposition of urine.* 6. *Tuberculosis.* 7. *Tumours.* 8. *Parasites.* 9. *Concretions and calculi; their various forms, etc.* DISEASES OF THE URETHRA, *chiefly Gonorrhæal inflammation, and stricture.*

## A.—THE KIDNEYS AND URETERS.

**INTRODUCTION.**—In studying the diseases of the kidneys, it is necessary to bear constantly in mind the general facts as to their structure, otherwise the examination of the organs will lead to confusion. The functions of the kidney must also be understood in their outlines in order to a comprehension of the changes wrought by disease.

The normal kidney measures about 4 inches in length,  $2\frac{1}{2}$  in breadth and  $1\frac{1}{4}$  in thickness. The weight varies considerably, the average for the male being  $4\frac{1}{2}$  to 5 ounces, and for the female slightly less. The surface is smooth and the capsule



although closely applied to the surface can be readily stripped off. On section the tissue is seen to be very regular, the distinction of pyramids and cortex being well defined.

When a microscopic section of the kidney, made so as to include both cortical and pyramidal substance, is examined, the contrast between these two regions is sufficiently striking in respect that in the cortical substance the uriniferous tubules have a markedly irregular and convoluted course. If attention be now confined to the cortex alone, as in

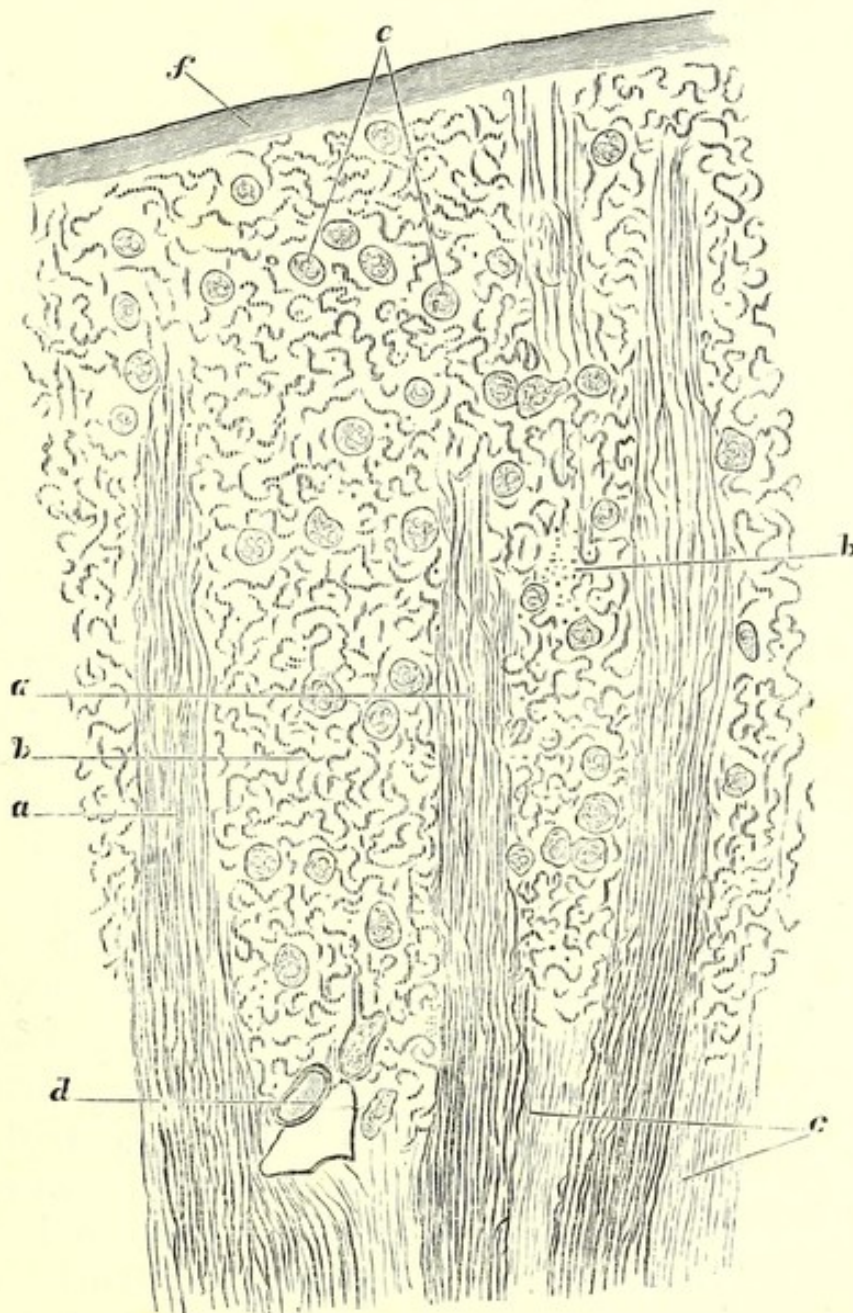


Fig. 299.—Section of normal kidney including cortex and base of pyramid with a very low magnifying power. *a*, medullary rays; *b*, convoluted tubules; *c*, region of arteriæ rectæ in pyramids; *d*, larger vessels running between pyramid and cortex; *e*, Malpighian tufts; *f*, capsule.  $\times 12$ .

Fig. 299, it will be seen that convoluted tubules are not the only kind present. There are also straight tubules prolonged up from the pyramids



in the form of tapering bundles (*a*) between which lie convoluted tubules (*b*). These tapering bundles, called medullary rays or pyramids of Ferrein, do not reach the surface, the most superficial part of the cortex presenting a continuous layer of convoluted tubules. In this way the deeper parts of the cortex present a regular division into alternating areas of straight tubules or medullary rays, and convoluted tubules.

Among the convoluted tubules lie the Malpighian capsules or glomeruli (*e* in figure). These occur somewhat frequently, and at tolerably regular intervals.

In addition to these arrangements of the tubules, the blood-vessels must receive attention. The larger arteries (*d* in figure), run between pyramids and cortex, and send up stems given off at right angles

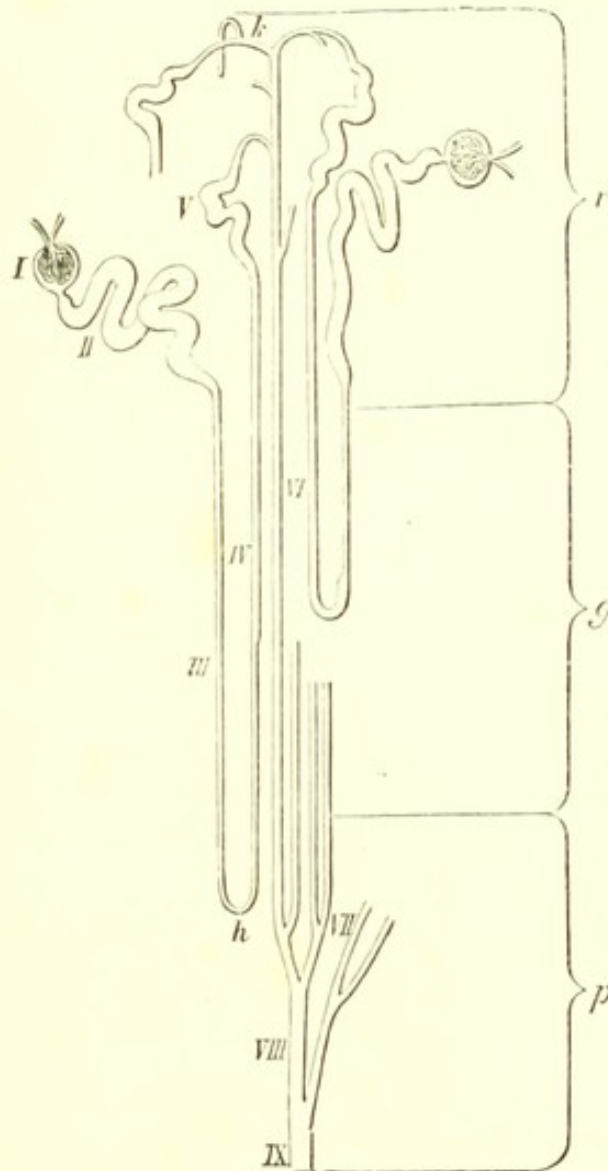


Fig. 300.—Diagram of course of uriniferous tubules from Malpighian capsule (I) to pyramid (IX). See text. (QUAIN.)

into the cortex. These pass at intervals into the region of convoluted tubules, and as they ascend they give off lateral branches to the glomeruli. It will thus appear that the areas of convoluted tubules are also the areas of the ascending arteries (which are also called interlobular arteries) and glomeruli. In the glomerulus the afferent vessel breaks up into a congeries of capillary vessels, called the tuft. These gather together to form the efferent vessel, which again breaks up into capillaries which surround the tubules with a rich network.

The large arteries which run between the pyramids and the cortex also give off arterial branches downwards to the pyramids. These arteries break up into bunches of straight arterioles (*arteriæ rectæ*, *c* in Fig. 299) which are increased by branches coming down from the afferent vessels in the deeper parts of the

cortex. These bunches of arterioles taper as they pass down the pyramids, so that they form small pyramids with their bases towards



the cortex. They correspond in position with the areas of convoluted tubules, which areas they, as it were, prolong down into the pyramids.

In studying the functions of the kidneys we have to remember in the first place the course of each uriniferous tubule, which may be followed in the annexed diagram. It begins in the Malpighian capsule (I). Issuing thence the tubule becomes convoluted (II), and then it dips down in a long loop (Henle's loop) whose bend (*h*) is usually in the pyramidal portion. Turning upwards (IV) the loop comes back to the cortex, again becomes convoluted (V), and then opens, sometimes at an acute angle, into a straight tubule (VI) which passes directly downwards, joined by other tubules (VII), till it opens at the apex of the pyramid (IX) into one of the calices.

So far as the water of the urine is concerned, it is generally agreed that it passes from the blood at the glomeruli; it filters, in fact, from the capillaries into the ends of the tubules. According to Bowman's view, it is mainly the water which passes through at the Malpighian capsules, the urea, urates, etc., being secreted from the blood by the large granular epithelium which lines the convoluted tubules.

It seems probable that Ludwig's view is correct, that the water passing through at the glomeruli is partly re-absorbed, but that the function of the epithelium is not confined to this. In experiments, in which indigo-sulphate of sodium was injected into the blood, it was found that this substance is excreted by the epithelium of the tubules, chiefly that of the convoluted tubules. The colour of the substance rendered it possible to see the seat of its excretion. As an inference from this it may be supposed that the epithelium is actively engaged in separating urea and other urinary constituents, perhaps changing some of them in transit.

The actual amount of the urine secreted will depend on the blood-pressure in the vessels of the glomeruli or on the speed with which the blood passes through these vessels. The amount secreted will be increased by increase of pressure in the vessels, as, for instance, by relaxation of the renal arteries, and it will be diminished by any cause which diminishes the pressure in these vessels. Considering the close relationship of the renal vessels to the systemic arteries and veins, it is clear that the blood in the former will be liable to considerable variations in pressure and in the speed of the current from circumstances affecting the general circulation, such as disease of the heart and lungs, etc.

**Literature.**—See full account in HEIDENHAIN, in Hermann's Phys., v., 1880; PANTYUSKI, Virch. Arch., lxxix., 1880.

#### I.—MALFORMATIONS AND MISPLACEMENTS OF THE KIDNEY.

1. **Congenital malformations.**—These are frequently such as to produce comparatively little interference with the function of the



organs. This does not apply to the extreme cases where both organs are absent or extremely small, but as this only occurs with serious malformations of the foetus as a whole, the child does not survive.

**Defect of one kidney** is not infrequently met with in well-formed adults without any of the signs of disease of the kidneys. It is mostly the left kidney that is defective, and it may be entirely absent, its vessels and a diminutive ureter ending in a piece of connective tissue. The other kidney in these cases undergoes a compensatory hypertrophy.

The kidney also not infrequently shows some trace of the **Fœtal lobulation** which in some animals is retained throughout life.

**Coalescence of the two kidneys** across the middle line is one of the most frequent malformations. Various degrees of it are presented. It may be a simple elongation of the inferior extremities of the kidneys

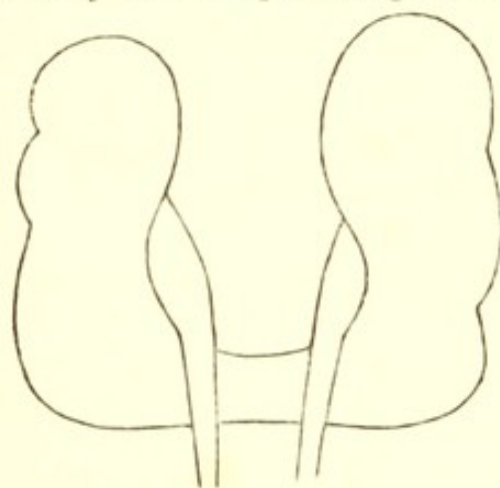


Fig. 301.—Outline sketch of horse-shoe kidney.

which are united by a fibrous band passing across the vertebræ. Or there may be a proper isthmus of renal tissue uniting the two kidneys into one and forming the well-known **Horse-shoe kidney** (Fig. 301). From this we have various grades on to complete coalescence of the kidneys into an elongated or square body across the vertebræ. In almost every case there are the regular two ureters, or they may even be increased in number, and they pass down in front of the isthmus. This form of kidney is often depressed in position, even coming as low in some cases as the hollow of the sacrum. When depressed the arteries usually have abnormal origins, as from the common iliac, hypogastric, etc.

**2. Variations of position.**—These may be congenital or acquired. In the former case and in some of the latter the kidney is fixed in its unusual situation. In **Congenital malposition** it is generally the left kidney which is concerned. It may be depressed so as to lie as low as the brim or even the cavity of the pelvis. It is not infrequently seated opposite the sacro-iliac synchondrosis. Such kidneys have usually the hilum presenting forward and are flattened, while their vessels are branches of the lower end of the aorta and of the iliac veins, or even entirely of the iliac vessels. The kidney may also lie nearer the middle line than normal or in the middle line.

The malposition may be acquired by the pressure of tumours or of the liver, by the dragging of a hernial sac, and so on.



**Moveable and Floating kidney.**—These terms designate two conditions, both of them characterized by undue mobility of the organ. The kidney normally lies behind the peritoneum, which covers it only on its anterior surface. The organ, being in the loose retro-peritoneal tissue, is surrounded by a fatty capsule as well as by its more immediate connective-tissue capsule. It is fixed to the posterior wall of the abdomen partly by its vessels, partly by the peritoneum binding it down, and partly by the pad of adipose tissue which forms its fatty capsule.

In the **Moveable kidney** the organ is unduly mobile behind the peritoneum; it may be moveable within its fatty capsule or may carry this with it in its displacement. The organ is sometimes capable of great displacement up under the ribs, down into the pelvis, and for a short distance across the middle line, although usually the mobility is limited.

The undue mobility occurs in the great majority of cases in females, and it is usually on the right side (in the proportion of 152:12, see Newman). Its frequency in women is ascribed to the disturbances produced by pregnancy, and also to the wearing of stays. The relations of the right kidney to the liver and ascending colon probably account for the greater frequency on this side. The heavy liver, especially when pushed downwards by stays, may dislocate the kidney, and the ascending colon is more loosely attached on the right side of the abdomen than is the descending colon on the left. Rapid emaciation, by diminishing the pad formed by the fatty capsule, is not an infrequent cause. It is stated that there may be a local diminution of the fat in the capsule.

**Floating kidney** is, strictly, a kidney with a mesonephron. The peritoneum covers both surfaces of the organ, and forms a mesentery which contains the vessels. This form is excessively rare, and its existence has been denied by some. It is of congenital origin. The degree of mobility is not greater than in many cases of moveable kidney, and the two conditions are scarcely distinguishable during life.

The moveable kidney is not very liable to secondary changes, although sometimes the abnormal position interferes with the flow through the ureter, and leads to hydronephrosis, or even inflammation of the pelvis. Of more importance is the fact that by dragging there may be serious nervous disturbances in the form of excruciating cramps. The kidney may, however, be moveable without any such nervous symptoms presenting themselves.

3. **Malformations of the ureters and pelvis.**—These are, in general, of little importance. The ureter may be double either in its whole course or in its upper part. The pelvis may also be double or in several divisions, each of which has a separate connection with the ureter. The ureter may arise from the pelvis at an acute angle, or there may be a kind of valve in the course of the ureter from a fold of mucous membrane.



**Literature.**—**RAYER**, *Traité des malad. des reins*, iii.; **FÖRSTER**, *Die Missbildungen*; **LANCEREAUX**, *L'union méd.*, 1880; **LANDAU**, *Die Wanderniere d. Frauen*, 1882; *Report of Committee, Path. trans.*, xxvii., 1876; **NEWMAN**, *Surgical dis. of kidney*, 1888; **BOSTRÖM**, *Path. Anat. der Niere*, 1886.

## II.—DISORDERS OF THE CIRCULATION IN THE KIDNEYS.

**Active hyperæmia** is caused by dilatation of the renal arteries. This may be from traumatic injury to the vaso-motor centre in the medulla oblongata. In a case of this kind observed by the author there was the most intense hyperæmia with enlargement of both kidneys, the injection affecting all the vessels. During the few hours that the patient survived large quantities of urine were twice removed by the catheter, and after death the bladder was again found distended by a watery urine. Again, we may have an active hyperæmia from removal of pressure, as after excision of large tumours from the abdomen or the removal of ascitic fluid, or even the removal of fluid from the pleura. Under these circumstances there is often for a day or two excessive secretion of urine lasting till the renal vessels resume their normal state of contraction.

The hyperæmia which follows on the administration of certain poisons, such as cantharides, arsenic, and carbolic acid, is to be regarded as really inflammatory.

**Passive hyperæmia** results from obstruction to the venous circulation, and is most frequently met with in valvular disease of the heart and in diseases of the lungs in which the circulation is seriously interrupted, as in severe emphysema.

If the obstruction take place suddenly there may be very intense engorgement of the renal vessels and considerable hæmorrhage from the glomeruli, so that their capsules and the tubules contain blood.

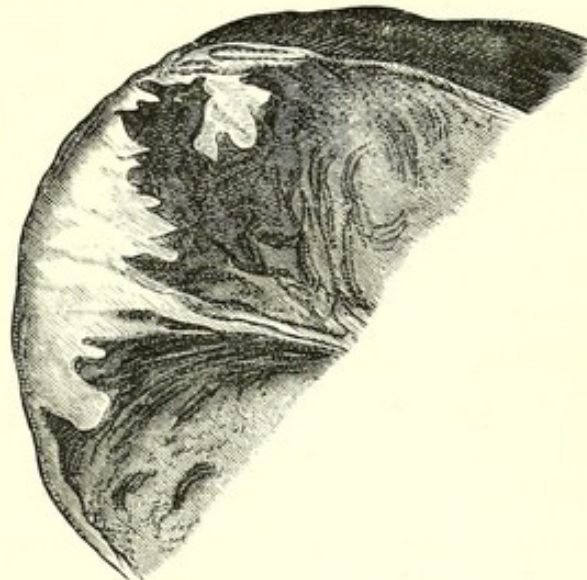
In the more usual chronic cases, such as occur so frequently in cases of heart disease, the kidneys present an increase in density due to **Cyanotic induration** (see ante, p. 62). There is also a general redness, but this is usually most manifest in the pyramids where the arteriæ rectæ often show very special dilatation, indicated by exaggeration of the red streaks which pass from the bases of the pyramids in the direction of the apices. The glomeruli are also visible in the cortex as small red spots.

Microscopic examination shows great overfilling of the vessels, accompanied in many cases with atrophy of the epithelium of the tubules, which is not infrequently fatty. There is often blood in the glomeruli and tubules, and sometimes brown pigment which has formed from blood. This pigment may be partly crystalline. The tubules also frequently contain hyaline tube-casts.



**Thrombosis** of the renal veins is sometimes a result of passive hyperæmia, but it usually occurs just before death and when the patient is very much debilitated.

**Embolism** of the kidney is very frequent. Remembering that the renal arteries are strictly end-arteries, it will be understood that when one of them is obstructed the **Infarction** virtually always occurs. The arteries of the kidney being distributed primarily to the cortex, the infarction is more or less wedge-shaped (Fig. 302), with the base of the wedge at the surface. If the wedge be of larger size it will extend also into the pyramids.



As a rule the infarction is of a pale colour and of dense consistence, the tissue having undergone **Coagulation-necrosis**.

There is not generally much hæmorrhage, but usually at the margin there is some, and if the infarction be small the hæmorrhage may extend throughout it. Around the infarction there is a zone of hyperæmia.

Fig. 302.—Embolie infarction of kidney. The white appearance and wedge shape of the infarction are represented. (After RAYER.)

The kidney tissue seems to undergo necrosis very readily when deprived of blood. Litten found that when the renal artery was ligatured for two hours the renal epithelium was already necrosed. This is probably the reason why the infarction seldom takes the hæmorrhagic form.

The infarction gradually undergoes absorption, and is replaced by a cicatrix. In this way deep depressions of the kidney may occur, and if there are several of them the kidney may assume a lobed appearance. In cases of old mitral or aortic disease it is very common to find deep cicatrices, indicating that, probably at the time of acute endocarditis, embolism of the kidney had occurred.

**Hæmorrhage** from the kidney is very frequent. It is of common occurrence in acute nephritis, and is not infrequent in chronic. It is a frequent symptom in tumours of the kidney, especially in cancers and cystic degeneration, and it results from calculi in the pelvis of the organ.

Purpura and scurvy seem to have a special tendency to affect the pelvis of the kidney, causing bleeding from its mucous membrane. A peculiar and interesting form is met with in infants, in whom a scorbutic condition has been induced by artificial feeding without sufficient



fresh milk. In this case blood in the urine may be the only direct symptom of scurvy. (See Dickinson.)

In all these cases the blood passes into the urine. Hæmorrhage into the parts around the kidney will only occur in cases of rupture from injury of the organ.

The kidney is not infrequently the seat of **Septic embolism** in pyæmia, ulcerative endocarditis, etc. The result is the formation of miliary abscesses, which will come up for consideration hereafter.

**Literature.**—COHNHEIM, *Die embol. Process*, 1872, and *Allg. Path.*, ii.; LITTEN, *Hæmorrh. Infarct.*, 1877, and *Virch. Arch.*, lxxxviii., 1882; RECKLINGHAUSEN, (*Retrograde embol. of renal vein*) *ibid.* c., 1885; DICKINSON, *Renal and urinary affections*, part iii., 1885.

### III.—HYPERTROPHY OF THE KIDNEY.

**Compensatory hypertrophy** of the kidney readily develops when one kidney is lost or congenitally defective. In the case of congenital absence of one kidney the other will be found homogeneously enlarged, and probably weighing nearly the same as the two normal kidneys together. The different regions of the kidney bear the same relations to each other, each being enlarged in its due proportion. The function of the kidneys is also completely carried out by the single one.

It has been determined by experiments in animals that compensatory hypertrophy develops after excision of one kidney in full-grown animals, although it is more complete when the operation is done in the new-born. It is remarkable how soon after such excision complete restoration of the renal functions occurs, the secretion of urea reaching its normal in one case about two days after the operation, and the animals remaining from the first apparently unaffected in health.

In the hypertrophied kidney there is newformation of tissue. There is not, however, a numerical increase in the lobules of the kidney, the glomeruli not being increased in number, although slightly in size. As the glomeruli are the expanded ends of the tubules, the latter also are not increased in number. They are not even increased greatly in diameter, and the newformation seems to be chiefly an elongation and increased convolution of the tubules.

Besides this form of compensatory hypertrophy, there may be an enlargement of both kidneys in diabetes insipidus, and perhaps also in diabetes mellitus.

**Literature.**—ROSENSTEIN, *Virch. Arch.*, liii., 1871; GUDDEN, *ibid.*, lxvi., 1876; GRAWITZ and ISRAEL, *ibid.*, lxxvii., 1879; BEUMER, *ibid.*, lxxii., 1878; COATS, *Proceedings, Med. Soc. of London*, vii., 1884.



## IV.—BRIGHT'S DISEASE. NEPHRITIS.

This is a subject of great complexity and difficulty, and one concerning which differences of opinion exist on many points.

We include under the designation Bright's disease cases in which there are undoubted inflammatory manifestations in the kidneys. The domain of Bright's disease, therefore, is not co-extensive with that of albuminuria, as this occurs in other conditions, such as amyloid degeneration and passive hyperæmia.

1. **Causation.**—In studying the causes of Bright's disease, we have to look for an irritant, and it is here important to observe in the first place that the inflammatory manifestations occur in both kidneys, and are diffused over the length and breadth of the organ. These facts indicate that the irritant is carried to the kidneys by the blood. As the blood is primarily distributed to the cortex, and as the cortex contains the more active secreting tissue of the organ, the inflammatory manifestations occur almost exclusively there.

In the various cases of Bright's disease we do not find the various constituents of the kidney tissue equally engaged. An irritant brought to the kidneys by the blood may show a predilection for the renal epithelium on the one hand, or the connective tissue on the other. It may be said, indeed, that for the most part irritants which act through a long period and with little intensity produce a chronic inflammation mainly affecting the connective tissue. On the other hand, irritants which act intensely so as to produce acute inflammation, while they induce the usual changes in the blood-vessels which we have seen to occur in acute inflammations, affect mainly the glomeruli and the epithelium of the uriniferous tubules. It may therefore be said that acute inflammations are mostly **Parenchymatous**, while chronic inflammations are mostly **Interstitial**.

As to the **Nature of the irritant** the specific poison of **Scarlet fever** frequently induces, especially in children, an acute inflammation. This may also be produced by the poison of measles and other specific febrile diseases. In adults the disease is mostly ascribed to **Cold**. Dickinson points out that cold mostly produces nephritis when the person is exhausted or asleep, and when the exposure has occurred immediately after profuse perspiration. It is as if, the functions of the skin being suspended, some deleterious material accumulated in the blood and irritated the kidneys. It appears that nephritis hardly occurs in persons exposed to cold in the arctic regions, probably because, the respiration being more vigorous, the deleterious material is carried off by the lungs. In warm climates also nephritis is uncommon, probably because the



body is less liable to sudden exposure to cold than in temperate regions.

**Chronic nephritis** sometimes remains **after an acute attack**. For the most part, however, when the disease begins as acute nephritis it remains subacute with repeated exacerbations, and the anatomical condition is a combination of that seen in acute and chronic Bright's disease. On the other hand, in a large proportion of cases, the disease is chronic from the outset and is to be ascribed to a cause which has been acting with slight intensity for a long period. Of all these causes the most definitely determined is **Gout**. The pathology of this disease is not very apparent, but there is an obvious alteration in the blood in consequence of which salts of uric acid are deposited in certain joints, generally with signs of acute inflammation. The same condition of the blood frequently induces chronic nephritis, and it is remarkable that when it attacks the kidneys it is less likely to affect the joints, and vice versa. Among the working classes gout largely arises from chronic **Lead-poisoning** (see statistics in Dickinson's work on Albuminuria), and in these cases the disease is particularly liable to attack the kidney, so much so that a large proportion of painters and others who work with lead die of chronic nephritis. Where the gout is due to the constant use of alcohol the disease is more liable to attack the joints.

Chronic nephritis is also sometimes induced by the poison of **Syphilis**. In this case it is apt to be associated with amyloid disease, but may also occur as a simple inflammation.

**Pregnancy** not infrequently leads to a chronic nephritis, in all probability by obstructing the vessels of the kidney by the pressure of the uterus. In recent cases there is extreme hyperæmia of the kidney, which with repetition of the cause, may go on to assume the regular character of chronic nephritis.

Lastly, there are cases in which there is no special cause apparent and we can only say that the person has been exposed to some influence whose nature we do not know. In many cases the disease has been very prolonged, and all through, the actual active disease at any particular time has been very slight. With this very insidious march, there may be the most serious permanent changes in the kidneys before any prominent symptoms have called attention to these organs.

2. **Forms of Bright's disease.**—Various subdivisions of Bright's disease have been made both from the clinical and the pathological point of view. The most generally accepted division is into parenchymatous and interstitial nephritis, to which some add the amyloid kidney.

Inflammations of the kidney have similar characters to those of other parts. In **Acute inflammations** the vessels are chiefly engaged at the



atset; there is hyperæmia with exudation of serous fluid, leucocytes, and red corpuscles. These find their way into the tubules and interstitial tissue, and appear in the urine as albumen, leucocytes, and red corpuscles. Death seldom occurs at the outset of the attack, and the appearances found post mortem are referable rather to secondary changes, chiefly in the secreting structures. Hence acute nephritis is included mainly under parenchymatous nephritis.

**Chronic inflammation** in the kidneys, as elsewhere, is chiefly characterized by newformation of connective tissue, and hence is included, for the most part, under interstitial inflammation.

It is to be understood, however, that as parenchymatous and interstitial changes are by no means mutually exclusive, and indeed frequently co-exist, so also the phenomena of acute and chronic inflammation are not limited to the one or the other form.

(a) **Parenchymatous nephritis** (*Tubular nephritis*).—In this form the secreting structures are specially engaged, including the glomeruli and tubules, but the degree in which these two constituents are affected varies somewhat, and an attempt has been made to distinguish a special subdivision, in which the glomeruli alone are affected.

Parenchymatous nephritis is, at the outset, usually acute, and it early corresponds with the clinical group, acute nephritis, in which the urine is scanty, highly albuminous and frequently bloody, and in which general œdema is a

characteristic feature. At acute nephritis frequently subsides into sub-acute or chronic stage, and in these, while the parenchymatous changes are still prominent, there are superadded some of the lesions of interstitial nephritis.

The **Glomeruli** show various changes. In some cases of scarlet fever there is in the glomerulus and around a great exudation of

leucocytes, which may fill the capsule and crush the tuft so as to conceal (Fig. 303). The leucocytes also over-run the neighbouring interstitial

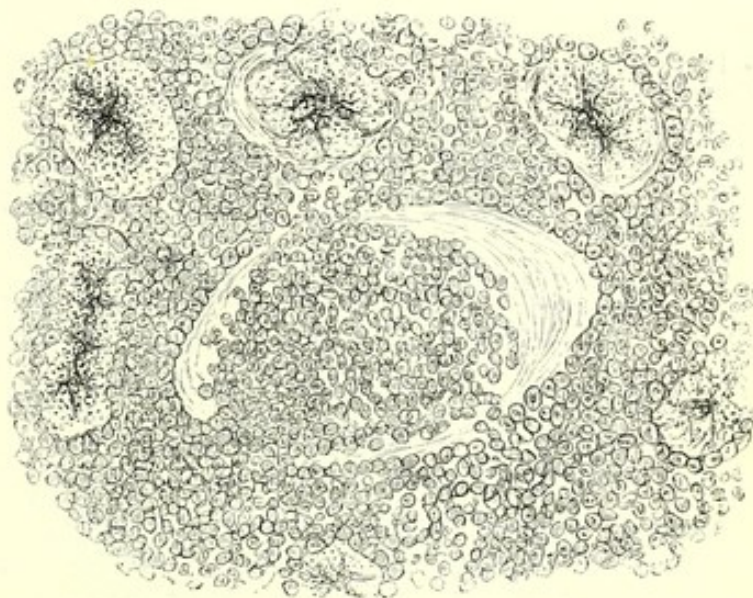


Fig. 303.—Glomerulo-nephritis from a case of scarlet fever. The Malpighian tuft in the middle of the figure is crowded with round cells, which are also present very abundantly in the interstitial substance between the tubules, whose epithelium is granular.  $\times 300$ .



tissue to a considerable extent, and the conditions may approach to those in septic nephritis. There may indeed be a septic element in many cases of scarlatinal nephritis (see below).

A much more frequent affection of the glomeruli, and one which occurs not only in scarlatina but in most forms of acute and subacute nephritis, is that in which the **Epithelium** is specially involved. There

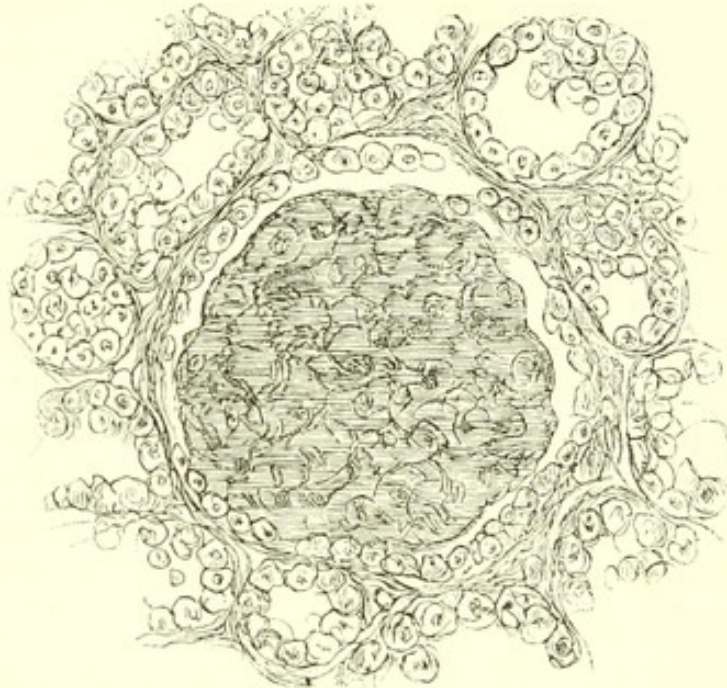


Fig. 304.—Glomerulo-nephritis in scarlet fever. The epithelium lining the capsule is unduly large and abundant.  $\times 350$ .

may be merely an enlargement of the epithelium, so that, instead of a thin layer which is usually invisible, it may form a distinct row of cells inside the capsule (Fig. 304). In many cases the epithelium multiplies and accumulates inside the capsule (Fig. 305), and the cells sometimes take on a stratified arrangement which has suggested to some authors that connective tissue is formed inside the glomerulus.

In addition to this there is very commonly in acute nephritis **Hæmor-**

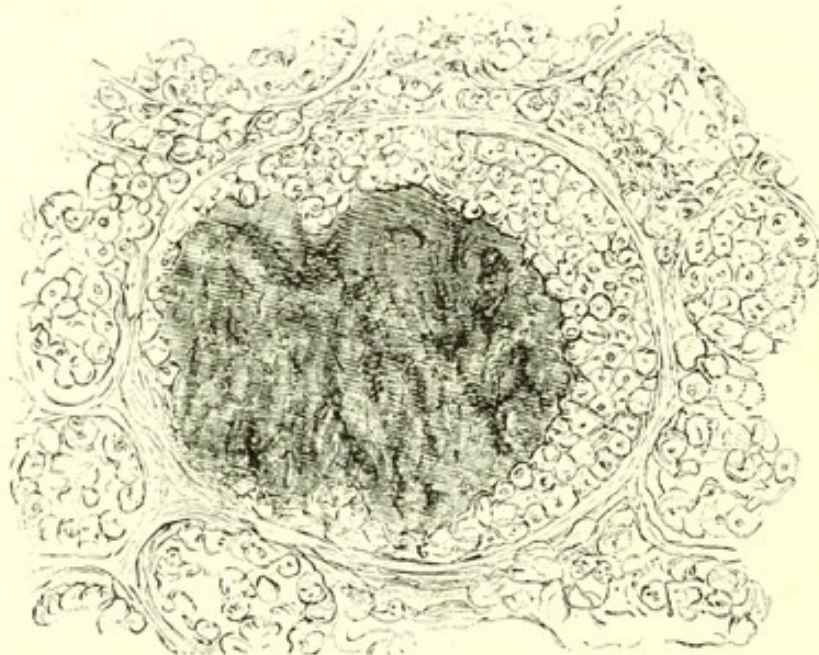


Fig. 305.—Glomerulo-nephritis in scarlet fever. The epithelium lining the capsule is greatly increased so as to crush the tuft.  $\times 350$ .

rhage from the glomeruli. Blood is usually present in the convoluted



tubules, either fresh or altered, this being generally indicated to the naked eye on examining the surface of the kidney after removal of the capsule by the presence of red or brown patches. By the aid of a lens these can often be resolved into a coil of convoluted tubules filled with blood. This is still more evident under the microscope, the tubules being visibly occupied by red corpuscles as in Fig. 306, or by brown matter representing old and altered blood. The blood comes from the glomeruli, and it is often possible, as in Fig. 307, to find glomeruli with blood inside the capsule as well as in neighbouring tubules.

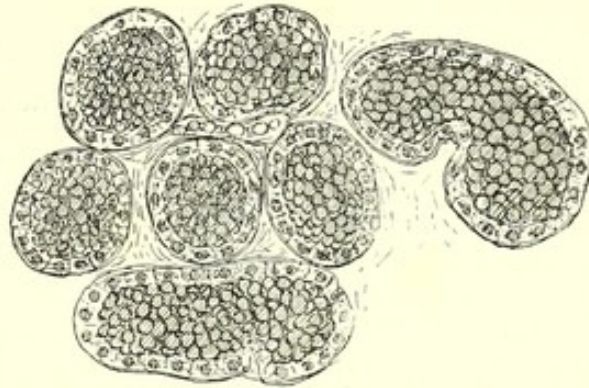


Fig. 306.—A coil of tubules distended with blood corpuscles. The epithelium is flattened against the wall.



Fig. 307.—Hæmorrhage from a Malpighian tuft in a case of scarlet fever. *a*, tuft; *b*, blood between tuft and capsule; *c*, blood in uriniferous tubule extending from tuft.  $\times 350$ .

**Scarlatinal nephritis** is usually characterized by such special changes in the glomeruli that the term **Glomerulo-nephritis** has been employed by Klebs to designate this and other forms in which the glomeruli are specially affected. In some cases of scarlatinal dropsy the kidneys are, to the naked eye, scarcely at all altered, although the patient may have died with symptoms of uræmia. The glomeruli may be visible on section as red spots, and there may be evidences of hæmorrhage but otherwise nothing abnormal. On microscopic examination, however, there will be the marked changes in the glomeruli described above. These lesions will seriously interfere with the function of the tuft. Any accumulation within the capsule will



compress the vessels and prevent the transudation of the water and other constituents. There may thus be a suppression of urine from glomerulo-nephritis.

But all cases of scarlatinal nephritis have not these characters; on the contrary, the kidneys may be greatly enlarged, and present the appearances of the large white kidney. In a case observed by the author in which the patient died on the ninth day after the onset of the fever the kidney was much enlarged, and there was a general infiltration, not only of the glomeruli, but of the whole cortex with leucocytes. It is from this case that Fig. 302 is taken.

In addition to these changes, Klein, while confirming the observations of Klebs as to the changes in the glomeruli, described minute changes in the afferent arterioles. These consist of swelling of the walls with contraction of the lumen, due to proliferation of the muscular and endothelial coats, and a hyaline change in the wall.

It is apparent that the conditions in the kidneys in scarlet fever are somewhat various, and are not all to be ascribed to the same cause. From the great frequency of the glomerulo-nephritis, which is present even in the first week of the illness, it may be inferred that it is due to the specific poison of scarlatina, which has a special affinity for these structures. But in scarlatina the condition of the throat allows of the absorption of septic microbes. Micrococci have been frequently found (by the author and others) in the kidneys both in scarlet fever and diphtheria, and certain of the forms of nephritis are probably septic. The large kidney with abundant infiltration with leucocytes is probably of this nature. (See further in papers by Crooke.)

**The uriniferous tubules** show very marked changes in parenchymatous nephritis. The changes consist in the first place in cloudy swelling of the epithelium with a tendency for the cells to become loosened and shed. The enlargement of the epithelium, occurring mainly in the cortical tubules, causes the latter, under the microscope, to present a strikingly prominent appearance. There are also parts in which the desquamated epithelium distends or chokes the tubules. Fatty degeneration soon affects the epithelium, and it is often present in a very high degree. It does not affect the epithelium uniformly, but at intervals there is a coil

of tubules with the epithelium highly fatty (see Fig. 308). The fatty epithelium may be dislodged and packed into further parts of the tube, as into the loops of Henle, so that one often sees a straight tubule occupied by fatty epithelium.

**Blood** is not uncommonly present in the form of fresh red corpuscles or a brown granular debris. This also, although chiefly in the convoluted tubules (see above) may extend into the straight ones.

**Tube-casts** are present in the form usually of translucent hyaline

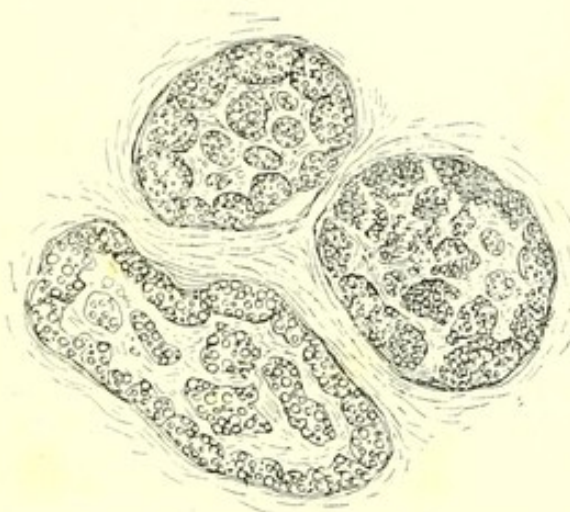


Fig. 308.—Uriniferous tubules with fatty epithelium, some of it shed into the calibre.



linders in the calibre of the tube. They are contained both in the convoluted and straight tubules, and they may be abundant in the pyramids.

As the case becomes more chronic the tubules are liable to considerable distortion, chiefly from the occurrence of interstitial changes. There may be irregular dilatations and contractions of the tubules, but the fatty condition of the epithelium remains prominent.

**The interstitial tissue** in subacute and chronic cases shows marked changes. Even in the acute stage there may be considerable infiltration with leucocytes. As the inflammation is prolonged, however, there is new formation of connective tissue and that special thickening of the capsule of the glomerulus, which is a prominent feature in interstitial nephritis. As this process goes on it produces distortion of the secreting tissue and irregularity of the surface of the kidney.

**The naked-eye appearances in parenchymatous nephritis.**—If the inflammation be mainly glomerular the general appearance of the kidney may be little altered (see above).

As a general rule in acute nephritis the organ is found enlarged, and may be more than twice its normal size. The capsule is easily removed and the surface of the organ has a generally reddish colour from injection of the vessels. Small red or brown areas are generally visible on close inspection, these being from hæmorrhages into the tubules. There are also opaque yellow patches usually visible, from fatty degeneration of the epithelium. On section the cortex is seen to be swollen and thicker than normal, and the hæmorrhages and fatty tubules will be visible as minute red and yellow markings. The latter especially produce a marked mottling of the cut surface in the cortex, and there are often elongated yellow streaks from fat in the straight tubules. The cortex, however, is usually pale as a whole as compared with the pyramids.

**The Large white kidney** is a further stage of parenchymatous nephritis. It represents a certain prolongation of the condition with partial subsidence of the inflammation into a subacute stage. The patient has been subject for months or years to intermitting attacks of dropsy with scanty albuminous urine. The kidney is large, and on the surface has generally pale appearance, with little or no irregularity. The capsule is non-adherent. On section the cortex is seen to be bulky and pale. On close examination of the surface and of the cortex it is seen that, besides the general paleness, there is an opaque mottling, representing the fatty condition of the epithelium. In this form there is considerable interstitial new formation of connective tissue, and, it may be, sclerosis of the glomeruli.



It is proper to mention that the naked-eye appearances of the large white kidney may be imitated by amyloid disease, where there is also usually some fatty mottling, and by septic nephritis, where there is a general infiltration of leucocytes.

**The contracted fatty kidney** represents the latest stages of parenchymatous nephritis. The organ may be greatly reduced in size, so as to weigh only 2 or 2½ ounces. It is somewhat irregular on the surface, but the appearance is rather of smooth rounded elevations than of granulations. The kidney is soft and flabby in consistence, and the capsule is usually non-adherent. It is generally pale and presents in addition a well-marked fatty mottling in the cortex and on the surface. The shrinking here is chiefly in the cortex, which may be very thin. The tubules are partly atrophied but partly dilated, and in the latter case their epithelium is usually fatty. The shrinking of the tissue brings the glomeruli close together and they often show marked sclerosis.

(b) **Interstitial nephritis** (*Granular contracted kidney, Cirrhosis of the kidney*).—This form of nephritis is for the most part chronic throughout, although sometimes an acute nephritis may pass into it, and it may even ensue on a scarlatinal nephritis. It is chiefly characterized by changes in the interstitial connective tissue, but there are also lesions in the glomeruli, blood-vessels, and tubules. Weigert suggests that the primary changes here may be in the secreting structures, the affection of the interstitial tissue being secondary.

**The interstitial tissue** is the seat of an infiltration of round cells, which is not uniform; it frequently occurs in wedge-shaped areas having their base at the periphery, and corresponding with the regions of convoluted tubules, but also in less definite areas in the midst of the cortex. The round cells go on to the formation of spindle cells and connective tissue so that the intertubular substance is increased by the newformation of a dense fibrous tissue such as commonly results from chronic inflammation. The shrinking connective tissue causes dwarfing of the tubules, which in the areas involved are reduced in diameter and contain small stunted epithelium.

The new-formed tissue of the cortex is continuous with the capsule and renders this adherent, so that commonly it is impossible to remove it without tearing the kidney tissue.

**The glomeruli** are strikingly altered. The first change is a thickening of the capsule (Fig. 309 a), which forms concentric fibrous layers. The tuft of vessels is subsequently occluded and converted into a dense homogeneous nodule. In this way the glomerulus is greatly contracted, being often reduced to less than half its normal size, and is represented by a solid white glancing nodule in which can still be recognized, as a rule, the thickened capsule presenting its concentric



fibrous appearance, and the central more homogeneous remains of the vessels (see figure). This condition is commonly called **Sclerosis of the glomeruli**, and the resulting appearance is very striking, especially as, with the atrophy of the tubules, the altered glomeruli are commonly brought close together.

The **Blood-vessels** take part in the inflammation, especially the **Arteries**. Their external coat, being continuous with the interstitial tissue, is thickened along with it. In addition

to that, however, the internal coat is usually very markedly thickened (see Fig. 170, p. 412). This thickening of the internal coat, which has the characters of **Endarteritis obliterans**, affects chiefly the ascending and afferent arteries of the cortex, but may also be present in the larger arteries between cortex and pyramids. It is sometimes so great as almost to amount to obliteration of the arteries.

It may be a question to what extent some of the other lesions are due to this narrowing of the arteries. We shall see further on that a primary obstruction of the arteries may lead to atrophy of the tissue and sclerosis of the glomeruli, and, according to Leyden, the lesions in the kidneys produced by lead-poisoning are due to a primary narrowing of the arteries. It may be said, however, in general that when atrophy is due to a primary endarteritis the affection is localized according to the accidental distribution of the arterial affection, whereas in interstitial nephritis the affection altogether is diffused.

The capillaries of the cortex are largely obliterated by the shrinking connective tissue.

The **uriniferous tubules** undergo alterations which are secondary to those in the interstitial tissue. In the areas in which contraction has occurred the tubules are separated by round-celled tissue or connective tissue and are dwarfed (Fig. 309). In intermediate parts, on the other hand, they are frequently dilated, there being here a kind of complementary dilatation. There is thus frequently visible a group of dilated and convoluted tubules (Fig. 310 *b*). The loops of Henle are also dilated, and by the shrinking of the cortex may be doubled up and

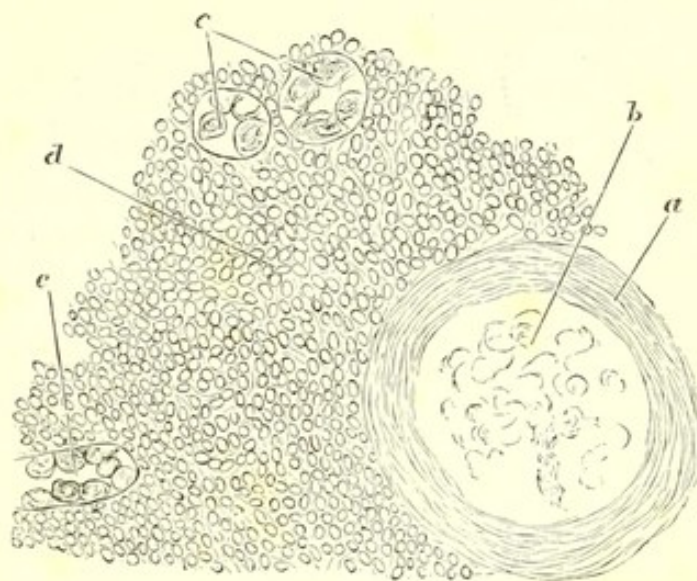


Fig. 309.—Interstitial nephritis. Sclerosis of glomerulus. *a*, thickened capsule; *b*, condensed vessels of tuft; *d*, abundant round cells in interstitial tissue; *c*, dwarfed tubules remaining.  $\times 350$ .



rendered tortuous (Fig. 310 *c*). The epithelium in the dilated tubules is

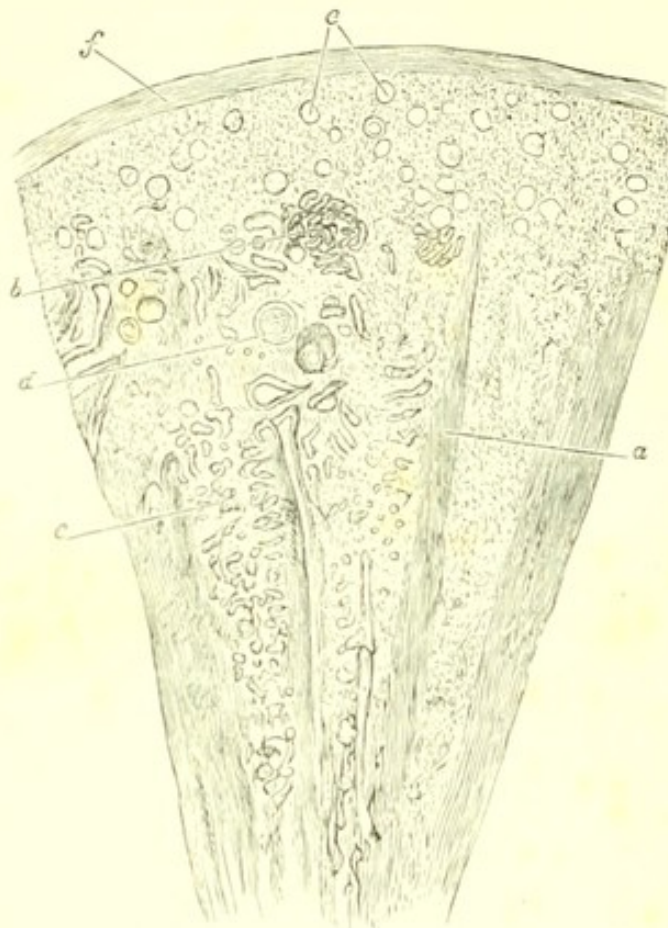


Fig. 310.—Section of contracted kidney with a very low magnifying power. (This should be compared with Fig. 299, p. 801, similarly magnified.) *a*, medullary rays; *b*, convoluted tubules crushed together and dilated; *c*, region of arteriæ rectæ and loops of Henle, the latter dilated and forming cysts; *d*, large artery between pyramid and cortex, showing endarteritis obliterans; the space between this and surface represents the cortex, and may be compared with similar space in figure 299; *e*, glomeruli, closely set and contracted, they also present a white homogeneous appearance; *f*, capsule.  $\times 12$ .

frequently fatty. **Tube casts** are also present in the tubules and they may be very abundant. They are usually hyaline in appearance and are occasionally present in the tubules both of cortex and pyramids.

**Cysts** are of very frequent occurrence in the contracted kidney. They arise mainly from the tubules by still further dilatation, and they very commonly lie together, sometimes in rows, as if from the crushing together and obstruction of a single tubule. Cysts also arise sometimes from the glomeruli, by accumulation of fluid inside the capsule. All these cysts arise by obstruction of the tubules, and the

usual contents are a watery fluid, but sometimes colloid matter is contained in them, apparently arising by secretion from or transformation of the epithelium. In nearly all cases there are cysts visible to the naked eye, and they are frequently so visible in large numbers. But many of them are microscopic in size.

By these processes great dislocation of the kidney tissue results, so that, instead of the perfectly regular arrangement of the normal kidney such as is shown in Fig. 299, p. 801, there is at once great shrinking and great distortion, as shown in Fig. 310 (drawn by the same apparatus and similarly magnified).

**Naked-eye appearances.**—The granular contracted kidney is usually small, dense, and tough. There may be in the early stages some enlargement, and a condition designated large red kidney has been described. The reduction in size varies greatly; in the most extreme



degree the kidney weighs about two ounces. The kidney feels dense, and on section it is found to be tough. The capsule is firmly adherent so that on removing it little bits of tissue come off with it. The surface has a general red colour, and is almost homogeneously granular. The prominent parts represent the less shrunken portions, and there may be some opaque markings from fat in them.

On the cut surface the regular normal appearance is greatly altered. The cortex is greatly thinned, sometimes forming only a thin rind, scarcely more than a twelfth of an inch in thickness, between the bases of the pyramids and the surface. The cortex has lost all its normal markings, and it is indefinitely demarcated from the bases of the pyramids. These changes may be partly appreciated by comparing Figs. 299 and 310, where *d* represents in both figures the vessels at the bases of the pyramids, and the cortex is the part between them and the surface. Cysts are also visible on examining the surface.

**3. Character and origin of tube casts.**—It has been pointed out that tube casts are usually present in Bright's disease, whatever the form or stage, and they are known to afford important indications in the urine passed by patients affected by this disease. But they exist in other conditions besides Bright's disease. They are not infrequently present in passive hyperæmia of the kidney, and are seen also in simple atrophic conditions.

They form cylindrical casts of the tubules, composed of a clear translucent hyaline substance, which, however, may contain in its substance, epithelium, fat, blood, and other matters which are present in the tubules, so that we may have hyaline, epithelial, granular, fatty, or blood casts, all of which may be found in the urine.

The tube casts are for the most part due to exudation from the glomeruli. They usually coincide with the occurrence of albumen in the urine, and they probably arise chiefly by coagulation of the serum-albumen which has transuded into the tubules. They are sometimes called fibrine cylinders, and their discoverer, Henle, regarded them as composed of fibrine. But they do not conform to the characters of fibrine, and as they occur in cases in which there is little or no inflammation they can hardly represent a fibrinous exudation.

Besides this origin they may be derived from the epithelium of the tubules, either by a kind of secretion from the epithelium or by a colloid transformation.

The tube casts which pass into the urine come chiefly from the straight tubules, including the loops of Henle. Those from the convoluted tubules, being thicker and having to pass the narrow loop, are less



likely to be carried outwards, although being soft and plastic they may accommodate themselves and so pass outwards.

4. **Other phenomena and consequences of Bright's disease.**—Of the remaining phenomena of Bright's disease we have here to consider, albuminuria, vascular changes outside the kidney, uræmia and anasarca.

(a) **Albuminuria.**—By this term is meant the escape of the serum-albumen along with the water of the blood, which no doubt occurs at the glomerulus. The albumen probably undergoes some changes after or during its passage (Kirk).

Albuminuria is a nearly constant symptom of Bright's disease but it is also met with in other forms of nephritis, in passive hyperæmia of the kidney, in amyloid disease, and in some acute fevers. Albumen is found even in small quantities in the urine of some persons apparently healthy, so that a condition of so-called **Physiological albuminuria** has been distinguished. There are, however, in such cases only slight traces, and the presumption is that some disturbance actually exists in the kidneys in all cases of albuminuria. It is clear from the variety of conditions which may lead to it that a comparatively slight derangement is sufficient to cause it.

The water of the urine is eliminated mainly at the glomeruli by a process which has been compared to filtration. If it were a simple filtration, albumen would be present, and some observers have supposed that albumen does pass through, to be reabsorbed by the epithelium of the uriniferous tubules. It is generally acknowledged that water is reabsorbed in the tubules and that the urine is thus concentrated, but the reabsorption of albumen is much more problematical. Such a hypothesis would imply that the appearance of albumen in the urine was due to the failure of the epithelium to absorb, and we should expect it to be associated with an excess of water. Precisely the opposite is in general the case, and albumen is more frequent in concentrated than in dilute urine. Moreover it is in acute nephritis, in which the epithelium is swollen and granular from absorption of albumen, that albumen is most abundant in the urine.

The truth seems to be that the process in the glomeruli is not a simple filtration, but a transudation through a living membrane of complicated structure. Wherever such transudations occur in the living body there is some selection of the constituents which are allowed to pass, and although the transudation fluids are albuminous they are so in very varying degrees. Thus the percentage of albumen in cerebro-spinal transudations is about 1·5 per cent., whereas in the peritoneum and pleura it may be five or ten times as great (see p. 91). The



glomerulus is as much like a secreting or glandular organ as a simple filter. The water in passing out has to penetrate the walls of the vessels and then a layer of epithelium. The capillary vessels themselves are highly cellular, being very abundantly nucleated, and the epithelium clothes the tuft completely (Heidenhain). These structures exercise a selection in allowing the passage of the constituents of the blood, giving transit especially to water and salts. It is a very interesting fact, and one entirely confirmatory of this view, that while serum-albumen is retained, egg-albumen when injected into the blood of animals is passed into the urine. (Stokvis and others).

Any derangement of the delicate glomerulus is likely to allow of the passage of albumen. Inflammation will injure it most directly, especially if the glomeruli themselves are specially affected. Hence in glomerulo-nephritis, while there may be very little water eliminated, any urine passed is rich in albumen. Amyloid disease also affects the tufts specially, and the urine, while abundant, usually contains considerable quantities of albumen. In chronic interstitial nephritis the tufts of vessels are not primarily attacked, and they are affected chiefly in the direction of occlusion. In this form there may be little or no albumen in the urine although the water is abundant. This latter fact is to be related to the atrophy of the tubules, which are unable to reabsorb sufficiently the excess of water passed, and also to an increase of blood-pressure in the vessels of the glomerulus which is characteristic of this disease. In passive hyperæmia, when prolonged and considerable, there is also liable to be damage to the kidney structures, just as, in other situations, catarrhs and œdemas are liable to result under similar circumstances.

It is not improbable that albumen may reach the urine from other sources than the glomeruli. The arteriæ rectæ of the pyramids occupy a somewhat similar position in the vascular system to that of the glomeruli. They come off to a large extent directly from the larger arteries, and the blood after leaving them also passes into capillaries. It is noticeable also that they and the glomeruli are the structures first and chiefly affected by amyloid disease. It is not improbable that in inflammations, in passive hyperæmia, and in amyloid disease there may be considerable transudation of serous fluid from these vessels, which may find its way into the uriniferous tubules.

(b) **Uræmia.**—This name is applied to a group of symptoms, the principal of which are vomiting, sleepiness, headache, convulsions, and coma, which occur in cases of Bright's disease when the excretion of the essential urinary constituents is diminished. It may occur when, from any cause, these constituents are retained in the blood, and hence



other conditions besides Bright's disease may induce it. The retention of urine may be, as in Bright's disease, due to lesions of the secreting apparatus, or it may be due to an obstruction in the outflow. According to Roberts the symptoms are different in the latter case, as if, when the kidney is healthy and secretes urine which is reabsorbed, the results were different from those which ensue when the original secretion or elimination is defective. There is thus distinguished an **Obstructive** and a **Non-obstructive suppression of urine**.

The urinary constituents retained in the blood act as poisons, but it has been found impossible to discriminate amongst the various symptoms as to how they are related to the various constituents, the chief of which are urea, creatine, and creatinine. Urea at least is by no means a vigorous poison, and it is only when in great excess that uræmic symptoms are produced. The presence of urea is often detectable in the blood and secretions of uræmic patients.

Considerable doubt was at one time thrown on the view that the symptoms of uræmia were due to poisoning by the urinary constituents. The injection of urea or urine into the blood of animals, or the ingestion of urea with the food, failed to produce the symptoms, leading only to an excessive secretion of urine. Two theories were devised to account for these apparent discrepancies. The theory of Traube, that the symptoms are due to œdema of the brain, is not now accepted. Nor is that of Frerichs, according to which it is not the urinary constituents themselves, but the products of their decomposition, chiefly carbonate of ammonia, which act as poisons. Chemical investigation shows that there is no excess of carbonate of ammonia in the blood in uræmia, and experiment indicates that carbonate of ammonia when introduced into the blood produces symptoms different from those of uræmia.

It is the accumulation of the constituents in the blood which produces, after a time, an intolerance of them. If, besides injecting urine into the blood in animals, the ureters be ligatured, then the symptoms of uræmia rapidly manifest themselves.

Defective elimination of urea and the other constituents occurs most frequently in acute Bright's disease, where the amount of urine is usually defective. But it also occurs commonly in advanced cases of chronic nephritis where the secreting structures are greatly atrophied. In these cases there may be excessive secretion of urine, which, however, is so deficient in urea that the total amount falls far below the normal. There is, however, usually a reduction in the amount of urine before the uræmic symptoms develop.

(c) **Œdema and Dropsy.**—These conditions frequently occur in Bright's disease, especially in acute and subacute cases. They have already been considered in an earlier part of this work (see p. 89). The frequent œdema of the skin is related, on the one hand, to the excess of water retained in the blood, and, on the other, to the irritation of the skin by the abnormal constitution of the blood. The irritant which acts on the kidneys will also act on the skin.



(d) **Changes in heart and arteries. Increased arterial tension.**—The nature of these changes and their relation to the affection of the kidney have been the subject of much discussion.

**Hypertrophy of the left ventricle** of the heart is one of the commonest results of Bright's disease. It occurs in subacute and chronic cases, and is much less common in parenchymatous than in interstitial nephritis. When it occurs in the former there is always some interstitial newformation. Other lesions of the kidney lead to it besides Bright's disease. Hydronephrosis with atrophy of the kidney tissue has repeatedly been observed to lead to it. Dickinson records a case in which a stone in the pelvis of the kidney produced contraction of the kidney and led to hypertrophy of the left ventricle. It may result from partial atrophy of the kidney from arterio-sclerosis, as in two cases observed by the author. Again, it has been produced in animals by inducing in them an indurative nephritis, which was done by ligaturing the renal artery for an hour and a half to two hours and then letting it go.

It is obvious from these observations that the hypertrophy of the heart is related in its causation to the lesions in the kidney, and it may be said generally that whenever there is considerable vascular obstruction in the kidneys hypertrophy of the heart is liable to occur, unless the health of the patient be too low to allow of such a formative process. It is not usually marked in amyloid disease of the kidney, as the amyloid condition frequently depends on lesions which seriously injure the health, and it is not nearly so common in parenchymatous as in interstitial nephritis, for the double reason that the vascular disturbance is much less, and the interference with the general health greater in the former than in the latter.

The changes in the arteries throughout the body consist in thickenings of their coats. This has been variously described as a thickening of the

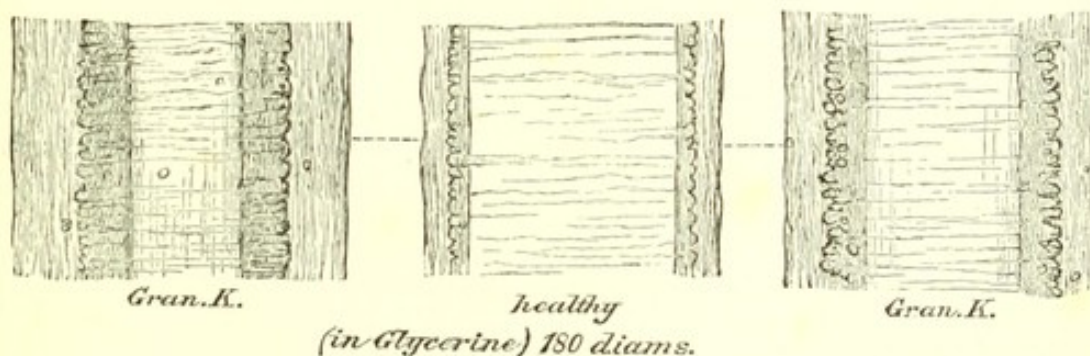


Fig. 311.—Arteries of the pia mater in case of granular kidney and in health. In the former the middle coat and, to some extent, the external are thickened. (DICKINSON.)

external coat mainly, an **Arterio-capillary fibrosis**, or a thickening of the middle coat, but it really consists for the most part of a general thicken-



ing, as shown in Fig. 311, involving both coats. It is detectable most readily in the arteries of the pia mater, which are easily examined by spreading out a small part of this fine membrane. Besides these thickenings, we have frequently atheroma in Bright's disease in people advanced in life.

These thickenings of the arteries are the natural result of the hypertrophy of the heart and the increased blood-pressure in the general vascular system. The blood entering the arteries with increased force requires greater resistance in the walls of the arteries in order to prevent dilatation, and may require also an energetic contraction to prevent an excess of blood from passing through the arteries if they remained of their ordinary calibre. The arteries subjected to increased blood-pressure are probably more liable to secondary changes such as atheroma.

The theory of these vascular changes is more difficult. The immediate cause of the increased blood-pressure is an increase in the force of the cardiac contractions, which results in hypertrophy of the muscle. Such hypertrophy is compensatory, and implies some obstruction to the circulation. According to Gull and Sutton, Mahomed, and others, the obstruction is a general one, and they regard the changes in the arteries as primary and as expressing a general disease, of which the kidney changes are only a part. They even suggest that the lesions, including cardiac hypertrophy, may exist without any kidney disease. The facts cited above, however, seem to afford sufficient evidence that the cardiac and vascular changes result from primary lesions of the kidney.

It seems difficult to believe that the obstruction in the vessels of the kidneys is by itself sufficient to induce such a cardiac hypertrophy as we frequently find, especially as excision of one kidney, implying great obstruction, does not usually lead to it. Hence a cause of obstruction has been sought for elsewhere. It has been supposed, for example, that the blood, containing retained urinary constituents, finds more difficulty than ordinary blood in traversing the capillaries, and again that the arteries contract excessively in order to prevent the impure blood from reaching the tissues too abundantly, and so produce obstruction (Johnson's "stop-cock" action). But these views imply two unwarranted assumptions, namely, that blood containing such impurities flows less freely through the capillaries than normal, and that the blood in all such cases is impure.

The changes seem inexplicable on a purely mechanical theory, and the explanations of Cohnheim as amplified by Fagge seem the most probable. In all the cases in which the cardiac and vascular changes occur there is obstruction to the vessels in the kidney, chiefly the arteries and the capillaries of the tufts. In order to the secretion of the proper amount of urine the blood must circulate much more quickly through the remaining vessels. This can only be effected by a rise in the blood-pressure in the vessels concerned, which may be brought about by a relaxation of the larger renal arteries. The activity of the renal secretion seems to be largely determined by the amount of urea and other constituents in the blood, the condition of the arteries as to contraction being regulated through the nervous system (Cohnheim and Roy). But if the local dilatation of the arteries be inadequate to cause a sufficient flow of blood, and the urinary constituents still tend to accumulate, then there can only be a relief by a general rise in the blood-pressure



brought about by increase in the force of the cardiac contractions. In this view of it the cardiac hypertrophy is compensatory, and is brought about in order to prevent the undue accumulation of urea in the system.

**Literature.**—RICHARD BRIGHT, Reports of medical cases selected with a view of illustrating the symptoms and cure of disease, by a reference to morbid anatomy, vol. i., 1827; RAYER, Traité des malad. des reins, 1840; FRERICH, Die Bright'sche Krankheit, 1851; ROSENSTEIN, Nierenkrankheiten, 2nd ed., 1870; JOHNSON, Lect. on Bright's dis., 1873, and Med. lect. and essays, 1887; GRAINGER STEWART, Prac. treatise on Bright's dis., 1868, and Clin. lect. on important symptoms, Albuminuria, 1888; CHARCOT, Lect. on Bright's dis. (transl.), 1879; AUFRECHT, Die diffuse nephritis, 1879; WEIGERT, in Volkmann's Sammlung, No. 162, 1879; RIBBERT, Nephritis and albuminuria, 1881; DICKINSON, Dis. of kidney, part ii., Albuminuria, 1877; RALFE, Dis. of kidneys, 1885. *Glomerulo-nephritis and Scarlatina*—KLEBS, Handb. d. path. anat., i. 644, 1870; KLEIN, Path. trans., xxviii., 1877; COATS, Brit. Med. Jour., 1874, ii.; FRIEDLÄNDER, Fortschritte der Med., 1883; LANGHANS, Virch. Arch., lxxvi. and xcix.; LITTEN, Charité-Annalen., iv.; CROOKE, Birmingham Med. Rev., 1886-87. *Interstitial nephritis*—TRAUBE, Gesammelte Beiträge, ii., 1871; LEYDEN, Zeitschr. f. klin. Med., ii. and iii.; ZIEGLER, D. Arch. f. klin. Med., xxv.; EBSTEIN, (Gout) *ibid.*, xxvii.; PEDELL, (Lead) D. med. Wochenschr., 1884; JACOB, *ibid.*, 1886; SAUNDBY, Path. trans., xxxi., 1880; GREENFIELD, *ibid.* *Albuminuria*—GRAINGER STEWART, l. c.; HEIDENHAIN, in Hermann's phys., v., 1880; SENATOR, Die Albuminurie, 1882; Discussion on albuminuria (by NEWMAN, ROBERTS, HAMILTON, GAIRDNER, GREENFIELD, COATS, etc.), Glasg. Med. Jour., 1884. *Cardiac and Vascular changes*—TRAUBE, Zusammenhang v. Herz- und Nierenkrankheiten, 1856; GULL and SUTTON, Med. chir. trans., lv., 1872, and Path. trans., xxviii., 1877; SOTNISCHESKY, Virch. Arch., lxxxii.; LEMCKE, D. Arch. f. klin. Med., xxxv.; HOLSTI, *ibid.*, xxxviii.; MAHOMED, Guy's Hosp. Reports, 1885; COHNHEIM, Allg. Path., ii. 349, 1882; FAGGE, Prin. and Prac. of Med., ii. 447, 1886; COHNHEIM AND ROY, Virch. Arch., xcii., 1883.

#### V.—SEPTIC OR SUPPURATIVE INFLAMMATIONS OF THE KIDNEY AND ITS PELVIS.

These are induced by the presence in the organ of septic microbes, namely, pyogenic micrococci. There are two important classes of cases; in one the source of the organisms is outside the urinary organs, the septic material being brought by the blood; in the other the source of the microbes is decomposition of the urine in the bladder, and they are propagated up from the latter. The former may be designated embolic or metastatic nephritis; the latter, being associated with inflammation of the pelvis, is included under the terms Pyelitis and Pyelonephritis.

1. **Embolic or Metastatic forms.**—In Pyæmia and Ulcerative endocarditis pieces of fibrine or other material may be carried to the kidneys and produce embolism there. In this way arise **Metastatic abscesses**. For the most part the emboli are small and are carried to the ascending arteries or the glomeruli before they stick. Hence the abscesses are



mostly in the cortical substance, although they are not infrequently seen in the pyramids along the course of the arteriæ rectæ, and there may even be large ones taking in portions both of pyramids and cortex. The abscesses are usually elongated in the direction of the arteries, and they often project slightly from the surface when the capsule is removed. They are frequently present in considerable numbers in both kidneys.

Under the microscope it can be seen that the abscesses arise by obstruction of the arteries (see Fig. 312). Where the embolism has been



Fig. 312.—From a section of the kidney in ulcerative endocarditis. An artery is plugged with dark material which contains micrococci. In the neighbourhood of the plug the wall of the artery is necrosed. Leucocytes are infiltrating the tissue around, and extending through the vessel wall.  $\times 90$ .

recent the wall of the vessel and the tissue in the immediate neighbourhood of the embolus present evidences of necrosis, while around there are multitudes of leucocytes occupying the interstitial tissue. In the embolus there are colonies of bacteria in the midst of remains of the transported fibrine. When the abscess has fully formed, these characters may be lost in the great multiplication of leucocytes. Besides in the



arteries colonies of bacteria are to be found in the vessels of the glomeruli, sometimes filling many of them out with a dark granular mass, and also in the capillaries. It is sometimes to be seen that these colonies are present in the glomeruli and capillaries without signs of inflammation around. In such cases they are of recent growth and have not had time to produce inflammation by giving off irritating products. They may, however, have produced necrosis of the epithelium in the neighbourhood, as shown by the failure of the nuclei to become stained (Fig. 313). To some extent the bacteria may multiply in the glomeruli and capillaries after death.



Fig. 313.—Micrococci in a capillary (b) of the kidney from a case of pyæmia. There is a tubule on either side, the nuclei of whose epithelium (a, a) are visible except in the neighbourhood of the micrococci, where necrosis has occurred.  $\times 650$ .



Fig. 314.—Bacteria distending a capillary blood-vessel in a case of scarlet fever.  $\times 350$ .

In scarlet fever there is rarely a true suppurative nephritis, but in some cases the kidneys are greatly enlarged and infiltrated with leucocytes. These cases are probably due to the presence of septic microbes, absorbed from the lesion in the fauces (see above, p. 814). Microbes are not infrequently found in the capillaries of the kidney in scarlet fever and diphtheria (Fig. 314), and they may also pass into the tubules.

**2. Pyelitis. Pyelonephritis. Suppurative nephritis.**—These various terms refer to inflammations which generally arise by extension from the bladder up the ureter to the kidney. Pyelitis is an inflammation of the pelvis of the kidney, while pyelonephritis is a suppurative inflammation involving the pelvis and the substance of the kidney. Pyelitis may occur independently, but pyelonephritis scarcely occurs without the ureter and pelvis being also affected.

(a) **Pyelitis.**—Inflammation of the pelvis of the kidney is not always



suppurative, as it may be produced by other causes besides the septic agents. A catarrhal pyelitis occurs in the course of some fevers, or it may be present to some extent in acute Bright's disease.

**Calculi**, by their mechanical irritation, will induce an inflammation of the pelvis, often spoken of as **Calculous pyelitis**. It is sometimes said that calculi produce suppurative inflammations, but this can scarcely occur unless there be an extension of the septic process from the bladder upwards. The irritation, however, may lead to such inflammation as to cause a considerable abundance of leucocytes in the urine, but not a true pus. On the other hand, septic inflammations, being accompanied by decomposition of the urine, often result in the formation of calculi.

**Suppurative pyelitis** is the result of the propagation of septic processes from the bladder by the ureter. It usually occurs in cases where there is some obstruction in the bladder or urethra, and a cystitis with decomposition of the urine has occurred. There is frequently some coincident dilatation of the pelvis, and there may be a proper pyonephrosis.

The mucous membrane is thickened and frequently of a bluish colour. It is not infrequently incrustated with phosphates. The contained matter is a thick brownish ill-smelling pus.

(*b*) **Pyelonephritis, Suppurative nephritis**.—This connects itself with the condition just described. There is an extension of the inflammation from the pelvis to the kidney substance.

There are two modes in which the condition may be brought about. The irritating chemical products from the pelvis may be carried into the kidney, being absorbed either by the tubules or the lymphatics, or the microbes may propagate themselves into the kidney. This latter method is not infrequent. Colonies of micrococci have been observed in the uriniferous tubules as well as in the lymphatics.

The lymphatics of the kidneys, according to Steven and Newman, are in communication with those of the ureter. When an injection is thrown with some force into the ureter, it passes into the adventitia of this canal, and thence to the capsule of the kidney. The lymphatics of the capsule communicate with those of the kidney, so that stellate vessels are visible, in these injections, on the surface of the kidney and are found to penetrate deeply into the cortex.

The resulting nephritis may be diffused so that the entire organ may be infiltrated, especially in the cortex, with leucocytes, and the appearances presented be those of an acute interstitial nephritis. This may perhaps be the case when the irritant is diffused from the pelvis in solution.

More frequently, with a certain diffused inflammation, there are actual abscesses, each abscess having a colony of micrococci as its focus. The



abscesses exist both in pyramids and cortex. They are scattered over the kidney so that the appearances may resemble those of pyæmia, but they are frequently in groups, occupying certain areas with a number of purulent centres. They are small in size but by coalescence may reach larger dimensions.

When the propagation is by the lymphatics the abscesses will be chiefly on the surface of the kidney, partly involving the capsule. It is not uncommon to open abscesses in the process of stripping the capsule. It may happen that abscesses form by this mode of extension without the pelvis being affected, the propagation occurring by the wall of the ureter to the surface of the kidney, but generally both modes of extension co-exist, and there are abscesses in the substance as well as at the surface of the organ.

When the suppuration is one-sided, **healing** may take place. The kidney is then the seat of numerous cicatrices, a condition which Moxon has named the **Cicatricial kidney**.

**Literature.**—BECK, Reynolds' Syst. of Med., v., 1879; DICKINSON, l. c.; NEWMAN, l. c.; STEVEN, Glasg. Med. Jour., xxii., 1884; MOXON, Path. trans., xxiii., 1882.

#### VI.—HYDRONEPHROSIS AND PYONEPHROSIS.

1. **Hydronephrosis.**—In this condition there is a dilatation of the pelvis and calices of the kidney, as a result of obstruction of the ureter or urethra.

**Obstruction of the urethra** may be congenital or acquired. In the latter case there is first dilatation and hypertrophy of the bladder, which may afterwards be reflected to the ureters and pelvis. In all such cases the hydronephrosis will be double, although it is not always equal on the two sides.

**Obstruction of the ureters** may occur in any part of their course. Tumours or inflammatory swellings in the pelvic organs will obstruct the lower ends, and usually both ureters will be affected. Calculi descending from the pelvis of the kidney occasionally cause obstruction. A peculiar form of obstruction is that in which the ureter does not, as in the normal condition, form a continuation of the pelvis of the kidney, but starts from it at an acute angle, passing obliquely through its wall. This is probably a congenital malformation. Of similar origin is probably the occurrence of valves in the course of the ureter. More unusual causes are, pressure from without by a renal artery taking an unusual course so as to cross the ureter; abrupt bends in the ureter, which may be congenital, or may be from the ureter making an unusually abrupt turn over the brim of the pelvis when the edge has been rendered sharper than usual by emaciation.



It appears from the last instance that a comparatively small external pressure may obstruct the ureter, and that the urine in the pelvis and ureter is at such a low pressure as to be incapable of overcoming a comparatively slight resistance.

The existence of some of these causes has been doubted, chiefly the existence of a valved aperture by the ureter arising from the pelvis at an acute angle, and the occurrence of valves in the course of the canal. The author has two specimens of the former kind in the Western Infirmary Museum (see Catalogue), and Dr. Gainsbury has published an apparently undoubted case of the latter form.

According to statistics collected by Newman, the obstruction arises, in the large majority of cases, either from obstruction of the urethra or from tumours of the pelvic organs, which may involve the ureters. These causes produced hydronephrosis in 400 out of 448 cases of double, and in 133 out of 215 cases of single hydronephrosis. Renal calculi form the most frequent cause of single hydronephrosis.

**The result of the obstruction** is dilatation of the pelvis and calices (see Fig. 81, p. 249, and Fig. 320, p. 839), but in addition there is frequently a considerable destruction of renal tissue. That is to say, the calices in dilating cause atrophy, first of the papillæ which project into them, and then of the pyramidal portion of the kidney, advancing into the deeper parts of the organ. We thus sometimes find the pyramids as if cut abruptly across transversely. But the condition frequently advances still further. The dilated calices increase in volume till, in the most extreme case, the kidney is replaced by a **Cyst** which represents dilated pelvis and calices, partitions existing in it corresponding to the divisions between the calices. When this has occurred, the cyst may go on enlarging so as to form a tumour of great bulk. The ureter is also dilated, often so greatly as to look like a piece of small intestine. It generally becomes convoluted in its course as well.

It is not in every case quite apparent what circumstances determine **the degree of hydronephrosis**. Complete obstruction leads usually to a comparatively slight hydronephrosis, the most extreme degree being reached in cases where on one side there is either an incomplete obstruction or one which gives way at intervals, such as a valved condition of the upper orifice of the ureter.

The ureter has been ligatured in animals and the processes observed. The first result is dilatation of the pelvis and of the ureter above the ligature. The tension of the urine in the ureter also rises up to a certain point, which, however, is far below the blood-pressure. When this point is reached the full extent of the distensile force derived from the secretion of the urine is attained. The secretion of the water of the urine, consisting in a transudation through the vessels of the glomeruli, is directly proportionate to the difference in pressure between the blood in the tufts and the fluid in the tubules, and when the pressure in the tubules is raised the secretion of urine ceases as soon as a state of equilibrium is brought about.



There are, however, two elements in the problem still to be considered. In cases where the ureter has been ligatured in animals the actual dilatation of the pelvis and calices has been comparatively slight, the state of equilibrium of tension being easily reached. And so in the human subject, when the obstruction is complete the pressure of the fluid in the dilated pelvis soon causes considerable obstruction of the renal vessels, and the power of secretion is reduced. But if at times an outlet is found for the urine and the pressure is suddenly reduced, there will be a relief of the vessels in the kidney and a violent hyperæmia leading to an excessive secretion of urine. As a matter of fact the sudden relief of an obstruction of the ureter has been found to be followed by an excessive secretion of urine which contained albumen. Where then there are such repeated sudden collapses and dilatations the advance of the hydronephrosis is most complete.

**The fluid contained** in the cavity is at first urine, but as the condition gets fully established the urinary constituents become absorbed and a watery albuminous fluid is found. In rare cases a colloid matter or a fatty milky fluid has been observed.

It sometimes happens that where a prolonged obstruction has existed **the external fatty capsule** of the kidney is greatly thickened, while only a moderate hydronephrosis exists, the external outline of the organ not being greatly increased. It is not unlikely that an œdema of the capsule following the obstruction of the ureter may be the cause of this great accumulation of fat by affording an extra supply of nourishing fluid, or perhaps by acting as a prolonged irritant.

We sometimes meet with cases resembling hydronephrosis, but in which some of the cysts, representing dilated calices, do not communicate with the pelvis, but form **Independent cysts**. This occurs when from inflammation there has been partial or complete obliteration of the pelvis (see below).

**Literature.**—ROKITANSKY, *Lehrb.*, iii.; COATS, *Catal. of Western Inf. Museum*, p. 119, 1885; GAINSBURY, *Path. trans.*, xxxvii., 1886; NEWMAN, *Lect. on Surg. dis. of kidney* 1888.

**2. Pyonephrosis.**—This term is used when, along with or following the hydronephrosis, suppuration occurs in the dilated structures. This is usually the result of an extension upwards of decomposition and inflammation from the bladder, such as has been referred to in relation to pyelitis. It may be followed by suppurative inflammation of the kidney, and in that case is usually fatal.

Such extension upwards will usually occur in cases where the primary condition and probably the seat of obstruction are in the bladder or urethra, hence it may be bi-lateral, and in that case is almost certainly fatal. However, it is not uncommonly unilateral, the extension having only occurred up one ureter. In that case there may be an abscess-like cavity formed, the dilated pelvis and calices being filled with pus.



If the affection of the lower urinary tract is recovered from, **healing** may take place in the kidney. One of the most constant results in this

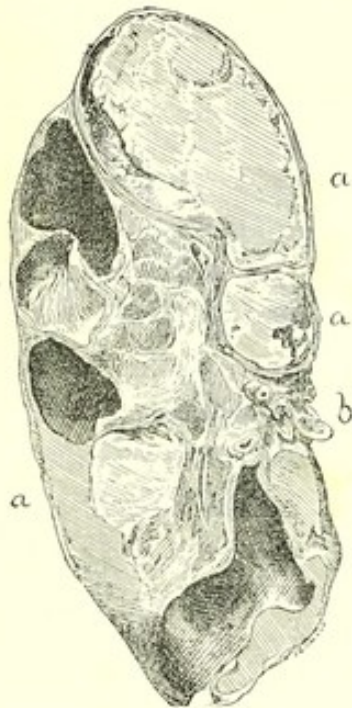


Fig. 315.—Healed pyonephrosis. *b*, Obliterated pelvis. *a, a, a*, Cavities filled with pultaceous matter. There are other cavities which contained serous fluid.

case is obliteration of the pelvis or upper part of the ureter, so that the kidney is shut off from the latter. By the obliteration of the pelvis the kidney is converted into a series of cysts, each independent of the other, and representing dilated calices (see Fig. 315). In these cysts pus is at first present, but as the corpuscles die and the fluid is absorbed, the pus dries-in and leaves a pultaceous matter in which lime salts are abundantly present (*a* in figure). It is not uncommon to meet with a kidney thus converted into cysts filled with putty-like matter, and it is remarkable that along with these there may be some in which clear serous fluid is present. Compensatory hypertrophy of the other kidney may complete the cure.

The above is believed by the author to be the course of events in cases of the kind referred to. The description is based chiefly on cases which have come under his

own observation. In one such case there was obstruction in the urethra in a female, with hypertrophy of the bladder. One kidney was found in the healed condition described above, while the other showed compensatory hypertrophy. This patient died from bronchitis without, latterly, any renal symptoms. In another case one kidney showed pultaceous masses, while the other presented a recent suppurative inflammation which had proved fatal. There was a prolonged affection of the bladder, with, apparently, an extension first to one kidney, producing a pyonephrosis partly recovered from, and then to the other, leading to a fatal result.

## VII.—RETROGRADE CHANGES IN THE KIDNEYS.

Retrograde changes are of frequent occurrence in the various forms of Bright's disease, but we have here to do with such as are met with more independently.

1. **Amyloid degeneration of the kidneys.**—Amyloid disease is of more importance in the kidneys than in other organs, both because of itself it leads to albuminuria and because its presence induces further structural changes in the organ. These facts induce some authors to include it as a form of Bright's disease. It exists along with amyloid disease in other organs, and is secondary to some disease which affects the body generally, chiefly phthisis pulmonalis and syphilis. In syphilis it not infrequently assumes more of an independent character, and the case, as a whole, may assume the aspect of one of kidney disease.



The lesion in the kidneys as in other organs affects primarily the walls of the blood-vessels. Even in very early cases, in which the amount of amyloid disease is slight, we usually find it in two distinct and separate structures, namely the Malpighian tufts and the arteriæ rectæ of the pyramids. The **Vessels of the glomeruli**, generally along with the arteries leading to them, are swelled up and hyaline. The rose colour developed with methylviolet is so striking that in sections stained with this dye it often looks as if the tufts were injected. They are also considerably enlarged, and even without the addition of iodine or methylviolet they form very prominent, transparent, and glancing lumps in the cortex. The **Arteriæ rectæ** run, as we have seen, in bunches, and their appearance with methylviolet is that of a series of rose-coloured tubes. They are also sufficiently pronounced as a rule without any reagent, appearing as pearly glancing tubes.

While these structures are first and chiefly involved, others usually follow. There is amyloid disease extensively present in the **Arteries of the cortex**, the ascending and afferent arteries. In these the lesion can often be seen to begin in the muscular coat, a distinct transverse marking being visible from the muscular fibre cells being mapped out by the staining agent. Very often there is here and there a

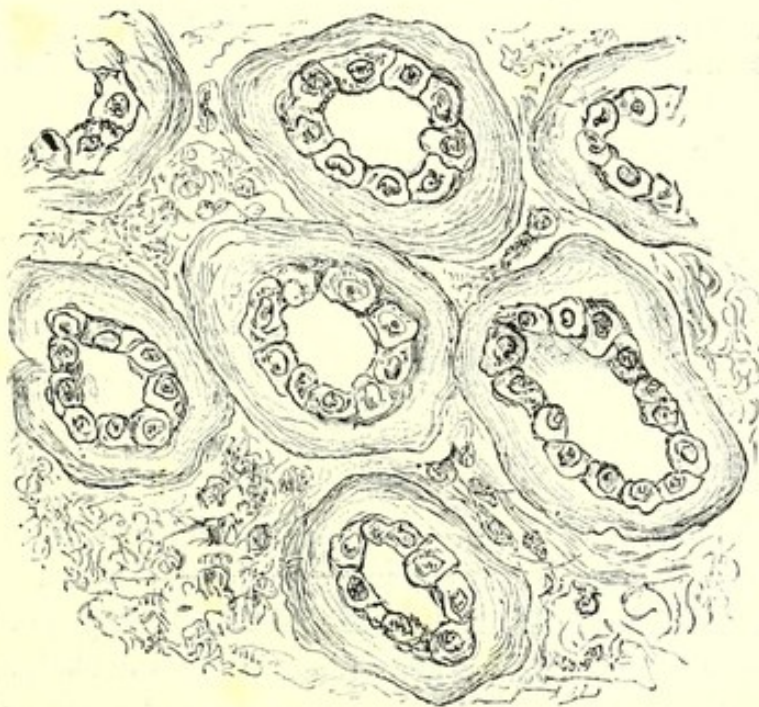


Fig. 316.—Amyloid degeneration of the basement membrane of uriniferous tubules. The thickened and translucent basement membrane is shown with the renal epithelium inside.  $\times 350$ .

capillary of the cortex affected, and sometimes these are extensively so, even in comparatively slight cases. The **Basement membrane** of the tubules frequently becomes amyloid in advanced cases, chiefly that of the tubules in the cortical substance (see Fig. 316.)



Along with the amyloid disease of the vessels there is usually **Fatty degeneration** of the epithelium of the tubules, apparently the result of anæmia from the obstruction of the vessels.

**Interstitial inflammation** nearly always accompanies the amyloid disease, and it may be so great as to produce a combination of chronic interstitial nephritis and amyloid disease, as in Fig. 317, where the

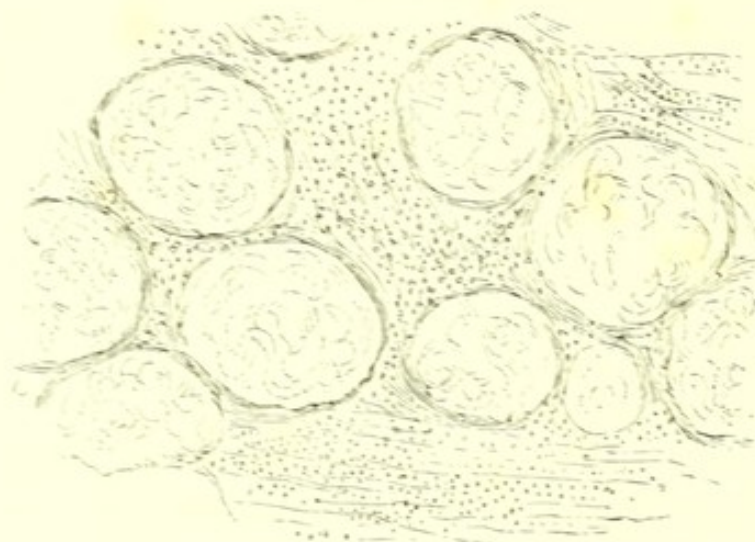


Fig. 317.—Amyloid and contracted kidney. The highly amyloid and therefore greatly enlarged Malpighian tufts are shown; their capsules are somewhat thickened. The tufts are very unduly close together, owing to contraction of the intervening tissue. Between them there is round-cell tissue with remains of tubules.  $\times 85$ .

glomeruli are brought close together by atrophy of the tubules from interstitial inflammation.

The mere existence of amyloid disease probably gives rise to interstitial inflammation, but it is not improbable that the primary disease which led to the amyloid condition may also induce nephritis. Thus in syphilis the virus affecting the kidney directly may cause interstitial inflammation, while less directly it produces a general amyloid condition. Again, in phthisis pulmonalis, and in extensive suppurations from diseased bone, there may readily occur an absorption of irritating materials which are capable of setting up inflammation.

The contracted amyloid kidney is not generally accompanied by hypertrophy of the heart, the reason apparently being that, as the blood is deteriorated, the compensatory growth of muscular tissue can scarcely occur.

In regard to the **Naked-eye appearances**, the simple amyloid kidney is enlarged, sometimes greatly enlarged, so that the organ weighs nine or ten ounces. The surface is pale and the capsule comes off easily. On section the cortex is seen to be thickened and pale in colour. The general appearance therefore resembles that of the large white kid-



ney, and doubtless many cases have been mistaken for that. But the organ in amyloid disease has a firm elastic feeling and a transparent bacony appearance on section, which is distinctive enough. The thick pale transparent cortex usually contrasts with the redder but still tolerably firm pyramidal substance, and the appearance has been somewhat aptly compared to the section of bacon ham. The name **Lardaceous kidney** is often applied from the transparent appearance of the tissue. In the midst of the transparent basis opaque yellow streaks are often visible, from the fatty degeneration of the epithelium. The addition of a solution of iodine to the cut surface of such a kidney brings out the affected glomeruli as brown dots and the arteriæ rectæ as brown streaks. If the kidney be contracted, the granulations on the surface show the usual transparent waxy appearance of the amyloid kidney, and on section the cortex, though greatly thinned, shows a similar condition.

The amyloid substance apparently allows fluid to pass through it more readily than ordinary albuminous structures. Injection of the amyloid kidney shows that the vessels of the glomeruli and the arteries of the cortex are largely obstructed. It may well be that this leads to

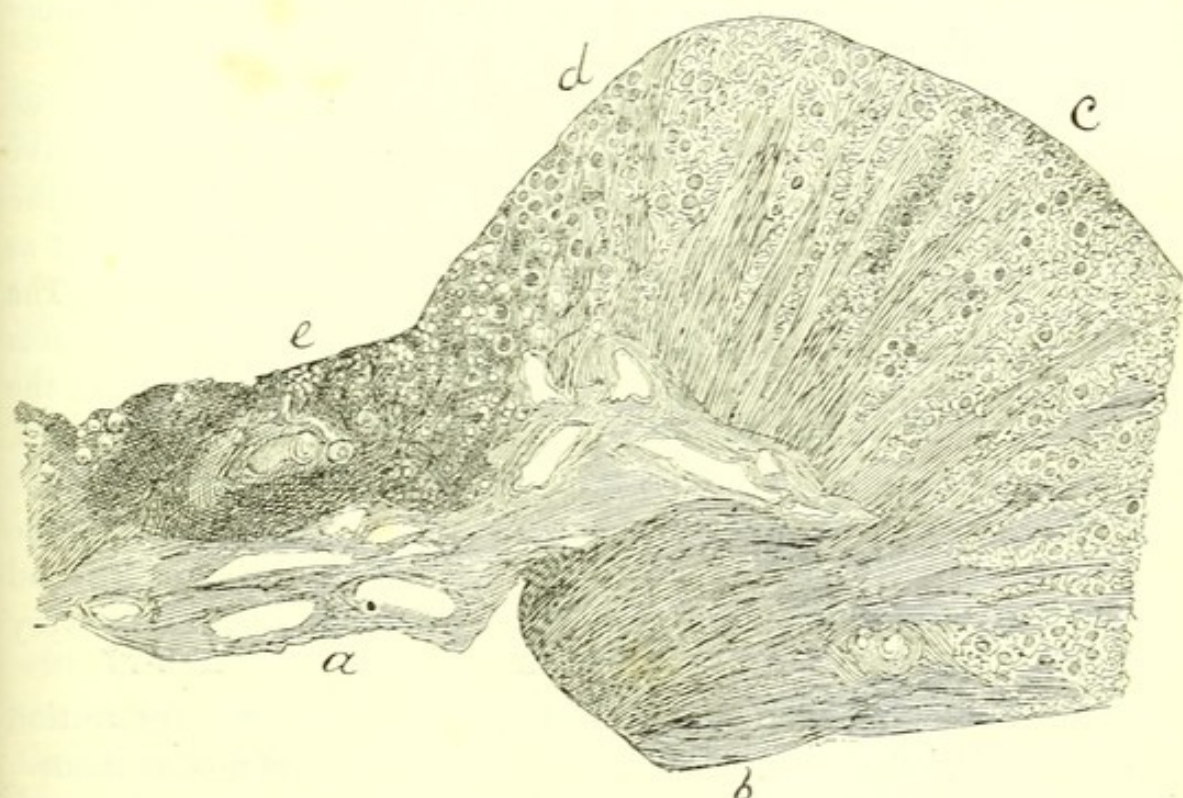


Fig. 318.—Local atrophy from endarteritis. *a*, hilum of kidney; *b*, normal pyramid; *c*, normal cortex showing glomeruli, medullary rays, etc.; *d*, atrophy beginning, the convoluted tubules atrophied while glomeruli remain. On approaching *e* the glomeruli become greatly contracted and close together.  $\times 6$ .

increased blood-pressure in the remaining arteries, but with the general narrowing of the smaller arteries this can hardly tell with much force on the glomeruli which remain pervious. This excessive secretion of



urine, like the diarrhoea which is a prominent symptom in amyloid disease of the intestine, is probably to be accounted for by a greatly increased transudability of the vessels.

2. **Atrophy.**—A local atrophy sometimes occurs from **Endarteritis** (see Fig. 318). There are flat depressions on the surface which may give the kidney a somewhat coarsely granular appearance, but sometimes these are extensive, involving considerable areas of the kidney. In this latter case the larger arteries running between cortex and pyramids are affected with endarteritis, which may be an **Atheroma** associated with a similar condition in other arteries. The thickening of the internal coat is very extreme (see above *e* in Fig. 318), and in the smaller branches amounts to absolute occlusion.

The atrophy has considerable resemblance in its details to that in chronic interstitial nephritis, but also presents considerable differences. It is not limited to the cortex, but involves the pyramids as well, and there is much less distortion but more of a simple diminution or disappearance of the tubules. The affected areas are very strikingly reduced in volume so that the hilum of the kidney comes close to the surface (see between *a* and *e* in figure). This contrasts greatly with the neighbouring unaffected parts in which the cortex is even more voluminous than normal.

By the **Atrophy of the tubules** the glomeruli are brought close together, so close in many cases as to show that the tubules have virtually disappeared. In the pyramidal portion the atrophy of the tubules causes the intertubular substance to become very prominent as a clear translucent matrix interrupted by small round areas. **The Glomeruli** are themselves in a state of sclerosis. At the marginal parts of the affected areas this condition is seen to begin by a thickening of the capsule which is less fibrous than in the sclerosis of interstitial nephritis, but has rather a hyaline appearance. The occlusion of the vessels in the tuft and the solidification of the glomerulus follow. In many parts the remaining tissue consists of little more than closely aggregated and solidified glomeruli, these being, apparently, in their altered state very persistent.

In addition to these evidences of atrophy there is some infiltration of round cells, but not any considerable newformation of fibrous tissue.

The description given above of local atrophy from endarteritis is based on the observation of two well-marked cases in which nearly half the kidney tissue was atrophied, the locality of the atrophy being chiefly in the neighbourhood of the hilum, and generally continuous, although also with isolated patches. An interesting feature in both these cases was the co-existence of hypertrophy of the left ventricle. The importance of the fact that local atrophy of the kidneys may have such a result has already been referred to.



In **Senile atrophy** the whole structures of the kidneys undergo diminution in size, but very often this affects especially the secreting epithelium of the convoluted tubules. If this be the case there may be a special shrinking of the cortex, and the kidneys may be like those of chronic Bright's disease. Sometimes also the surface is granular and the capsule adherent.

3. **Parenchymatous infiltration**, or **Cloudy swelling**, besides forming one of the lesions in acute Bright's disease, is liable to occur in acute febrile diseases. There is a general enlargement of the renal epithelium with infiltration of fine granules in the midst of which minute fat drops are



Fig. 319.—Cloudy swelling of renal epithelium. The cells are enlarged and filled with fine granules, in addition to which there are occasional minute oil drops.  $\times 350$ .

scattered (Fig. 319). The pyramidal tubules as well as those of the cortex are involved, the affection being usually homogeneously distributed. It is important to note that this condition may produce considerable enlargement of the kidney without obvious change in its structure to the naked eye.

**Fatty degeneration** of the renal epithelium is not infrequently present in cachectic conditions such as advanced phthisis pulmonalis, in which, however, it is often associated with amyloid disease or interstitial nephritis. In severe anæmias it is also met with along with fatty degeneration of the muscular substance of the heart. It occurs, too, in acute yellow atrophy of the liver.

#### VIII.—CONCRETIONS AND CALCULI IN THE KIDNEY.

Two very interesting forms of **Deposition in the tubules** are met with in **new-born children**. In the common icterus of the new-born the **Biliary colouring matter** (bilirubin) is excreted, as in the adult, by the kidneys, but in the new-born it very readily passes into the **Crystalline form**, so that round or rhombic orange-coloured crystals are found in the pyramidal tubules, sometimes in such numbers as to warrant the designation bilirubin-infarction. Similar crystals are found in the blood and tissues of the body.

**Uric acid salts** are found in the tubules of new-born children, in about half the cases of those who die within the first few weeks. It is mostly



in children who have breathed that they are met with, but they have been observed in still-born children, although very rarely. The concretions consist of salts of uric acid (mostly urate of ammonia). They are deposited mainly in the tubules near the apices of the pyramids, giving the appearance of opaque yellow or reddish streaks, converging to the apex of the pyramid. Under the microscope amorphous urates are seen in the tubules, and when acetic acid is applied these dissolve and form crystals of uric acid.

**In adults, icterus** is also accompanied by deposition of pigment in the tubules, but it is in the form of brown or yellowish granules which are found in the epithelium or forming casts of the tubules.

**Urates** are also deposited in the substance of the kidney in **Gout**. They are visible to the naked eye in the pyramidal portions of the kidney, especially towards the apices, as white streaks. With the microscope they are seen to occupy the tubules, which are dilated. There is usually a co-existing interstitial nephritis.

**Lime salts** are sometimes deposited in the straight tubules of the pyramids, especially in cases where from disease of bone there is excessive absorption of lime salts and excretion of them by the urine. The appearance presented is that of white streaks usually near the apices of the pyramids. It is to be observed, however, that a deposition of urates occurring after death in the straight tubules may produce a somewhat similar appearance.

**Calculi in the pelvis** of the kidney are of frequent occurrence. As we have seen, they may originate from chronic pyelitis, especially with stagnation of urine, but they may apparently originate in the uriniferous tubules, and having passed into the pelvis grow larger there. The calculi may be composed of uric acid, of phosphates, of oxalate of lime, or of cystine. The character of each of these is described in the section on Calculi in the Bladder.

The calculi sometimes attain to very large dimensions, moulding themselves into the shape of the pelvis and calices, so that we may have branches extending out into elongated recesses formed by dilated calices. By obstructing the flow of urine these calculi may lead to the occurrence of hydronephrosis, and as the cavity of the pelvis and calices enlarges so may the calculus. A small calculus will often pass into the ureter and obstruct it, in this way leading to hydronephrosis.

#### IX.—SYPHILIS AND TUBERCULOSIS OF KIDNEY.

1. **Syphilis** is sometimes the cause of diffuse **Interstitial nephritis**. It is undoubtedly a frequent cause of **Amyloid disease** and may by this means lead on to chronic nephritis. It is probable that many



cases of contracted amyloid kidney have this origin. Hereditary syphilis is also believed to give rise to a diffuse interstitial nephritis.

**Gummata** are very rare in the kidney. They are found in the midst of cicatrices as in the case of the liver.

**2. Local tuberculosis or Renal phthisis.**—This condition is usually associated with tuberculosis elsewhere. Thus there is commonly tuberculosis of the ureter and bladder and also, in the male, of the vesiculæ seminales, vas deferens, and testicle. It seems probable that in most cases the tuberculosis originates in the testicle and travels to the other parts (Weigert), but an origin in the kidney itself is not to be excluded. The fact that the disease is rare in females would indicate that the propagation is usually from the testicle. Renal phthisis is frequently associated with pulmonary phthisis, which is probably in most cases the primary affection.

The disease is usually very advanced in one kidney and absent or slight in the other. In the advanced cases the kidney is converted internally into a sac with irregularly ulcerated walls, and divided partly into compartments by the remains of septa. The ulcerated surface presents some adhering caseous matter and the wall is infiltrated (see Fig. 320). The capsule of the kidney is adherent.

In an early stage we find, it may be, the apex of a pyramid the seat of a small ulcer with yellow caseous walls, and with grey tubercles at the periphery in the substance of the pyramid. The lesion begins usually in the calices, frequently in those situated inferiorly, and eats its way in, attacking and destroying the kidney tissue in its advance. There is a very irregular extension of the ulceration, but usually there is some indication of the calices preserved.

The **Pelvis and Ureter** partake in the tuberculosis, and the latter is often infiltrated either in patches or throughout. Its wall is greatly thickened, and its internal surface often presents an almost continuous layer of caseous matter.

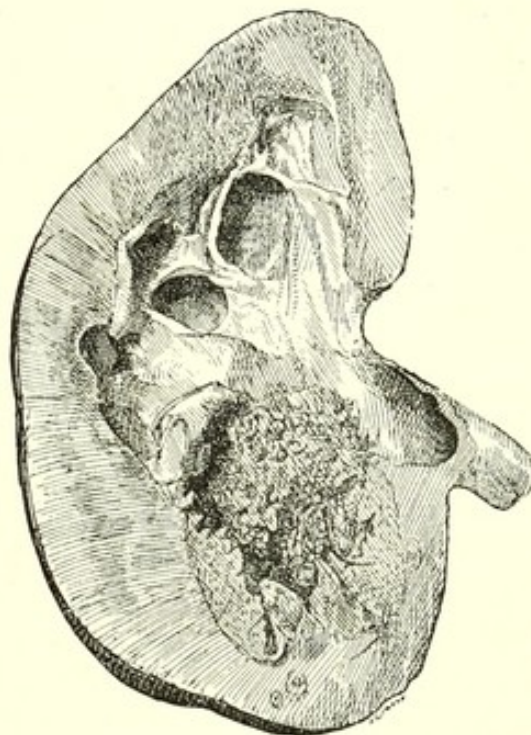


Fig. 320.—Tuberculosis of kidney with hydronephrosis. At lower part a ragged tubercular ulcer eating into kidney tissue. Two small isolated tubercular areas below. In upper part dilated calices (hydronephrosis). To right dilated ureter.



There is sometimes a partial **Hydronephrosis** (as in figure) along with the tuberculosis, from the caseous matter obstructing the ureter. It also happens sometimes that tuberculosis in one kidney is associated with hydronephrosis in the other. In such cases the tuberculosis of the bladder, affecting the orifice of the ureter, has caused obstruction of it and hydronephrosis has ensued.

The tuberculosis sometimes attacks a pervious branch of the **Renal artery**, and the tubercular virus is disseminated over the area of distribution of the artery. Hence it is not uncommon to find a larger or smaller wedge-shaped piece of kidney occupied by closely aggregated tubercles, while there is a comparatively slight tuberculosis of the pelvic portions.

3. **General tuberculosis.**—A **Chronic general tuberculosis** is not uncommon in children, in the form of yellow masses, usually softened in the central parts. This is associated with tuberculosis in other parts, usually in the lungs. In adults, phthisis pulmonalis is frequently associated with a few tubercles in the kidneys as well as in the liver, but they are of comparatively small size.

In **Acute general tuberculosis** there are numerous tubercles in the kidneys, mostly in the cortex. They are usually elongated in shape and visible to the naked eye as small pale areas. Under the microscope they are found to be caseous in their central parts while peripherally their cells infiltrate between the tubules.

**Literature.**—NÉGEL, De la syph. rénale, 1880; SEILER, D. Arch. f. klin. Med., xxix.; CURT JANI und WEIGERT, Virch. Arch., ciii. 522, 1886; HANAN, *ibid.*, ciii. 221, 1887; NASSE, *ibid.*, cv. 173, 1886; BAUMGARTEN, Tuberculose, 1885.

#### X.—TUMOURS OF THE KIDNEY.

**Fibromas** are frequently met with in the kidneys in the form of small white tumours of no practical significance, but they have also been found of large size. **Lipomas** are very rare in the kidney, but the fatty external capsule sometimes forms a bulky tumour (*Lipoma capsulare*).

**Cysts.**—Cysts of various kinds are of peculiarly frequent occurrence in the kidneys, and they probably all arise as **Retention cysts** by obstruction of tubules.

In the contracted kidney it is common to meet with small cysts which are the result of dilatation of the uriniferous tubules and Malpighian capsules from obstruction of the tubules.

**Simple cysts** are frequently met with in kidneys which are otherwise perfectly normal, and the cysts themselves do not, as a rule, seriously interfere with the functions of the organ. They are larger or



smaller well-formed cysts, which not infrequently project from the surface of the organ. They contain usually a clear fluid, but the contents are sometimes colloid in character. The wall of the cyst is composed of connective tissue lined with a proper tessellated epithelium. These cysts sometimes grow to a large size and may push aside the renal tissue to a large extent. It is probable that they are of congenital origin, arising by foetal obstruction of one or more uriniferous tubules.

**Cystic transformation** of the kidneys is also a condition of congenital origin, at least in many cases and probably in all. The whole kidney is converted into a congeries of cysts of larger and smaller size. The cysts form rounded prominences from the surface, and on section (Fig. 321) they are seen to replace the renal tissue. The wall of

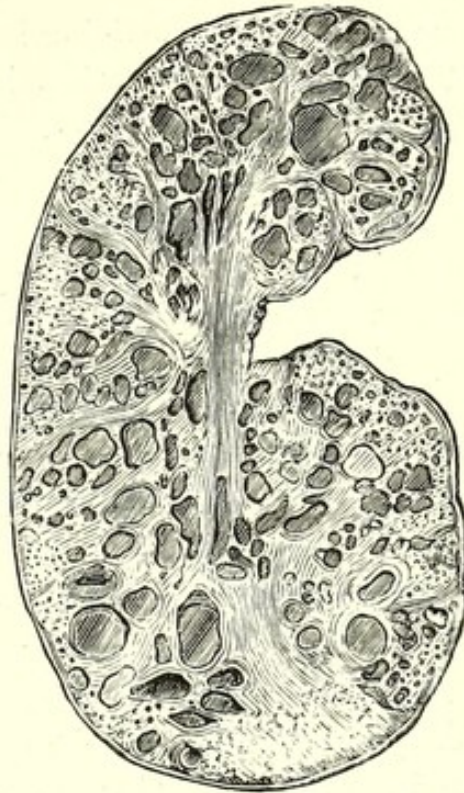


Fig. 321.—Cystic transformation of kidney. The organ is shown in section, and is made up of a congeries of cysts of various sizes. (VIRCHOW.)

each cyst is composed of a tunica propria with well-formed tessellated epithelium lining it. It contains usually a clear fluid, but the fluid may be brownish or even hæmorrhagic, and it is noteworthy that it contains the constituents of the urine, often with albumen, and sometimes throws down a granular precipitate of uric acid. Between the cysts there is very little space, but there are traces of remaining renal tissue, and the pelvis and ureter are present. The outline of the organ is greatly enlarged, measuring sometimes in the new-born child as much as eight inches by four.

Kidneys of this kind have been frequently met with in **new-born children**, and from their size they may seriously interfere with parturition, even requiring evisceration before delivery can be effected. It is of importance to note that this condition often co-exists with other congenital defects such as hydrocephalus, defective urinary bladder, and horse-shoe kidney.

But cystic degeneration is also met with in **the adult**, and it is remarkable that the functions of the organs may be preserved for a long time, although both kidneys are composed of a congeries of cysts. The author met with a case in which the patient died at the age of forty-three. For eighteen years before his death there had been



recurring attacks of hæmaturia, and he at last died with uræmic symptoms. The kidneys were much enlarged and cystic, but with some renal tissue remaining between the cysts.

The cystic degeneration in all probability arises by obstruction of the uriniferous tubules. Virchow has found in some congenital cases an obstruction of the papillæ apparently from a foetal inflammation. The fact that these cysts contain the urinary constituents is also much in favour of the view that they arise by obstruction of the tubules. Those met with in the adult are also most probably of congenital origin, although they have grown on into adult life. In the author's case there was a congenital malformation of the aortic valve, indicating perhaps a proclivity to malformations.

In a considerable number of cases the cystic kidney has been found along with multiple cysts in the liver. In some cases there were also cysts in the pancreas, in the lungs, etc. (Pye Smith).

**Dermoid cysts** are also met with in the kidneys.

**Adenoma.**—This form of tumour is described as occurring in the kidneys in the form of defined solitary growths of various sizes up to that of a walnut. They occur in the cortical substance, and arise from the convoluted tubules. According to structure two forms are described, a papillary and an alveolar, these resembling the corresponding forms in the ovaries.

The true adenoma is rare, and usually forms a small tumour of no special significance.

A form of tumour has been described as consisting of pieces of **Displaced suprarenal capsule**. The tumours lie immediately under the capsule and are distinctly defined from the kidney tissue. Under the microscope they show cylindrical masses of cells which are infiltrated with fat.

**Primary sarcoma.**—This occurs in various forms. It is stated that cases described as cancer are frequently sarcomas (Dickinson). The early age at which cancers of the kidney are stated to occur lends support to this view, as, in general, children are much more liable to sarcomas than cancers.

The forms chiefly met with are round-celled and spindle-celled sarcoma, myxosarcoma, and myosarcoma. The tumours generally grow rapidly and may assume large dimensions. They occur mostly in early life.

The **Myosarcoma** is a specially interesting form. It is composed of round or spindle-celled tissue with which is mixed striped muscle in the form either of elongated cells or cylinders. The tumour is probably congenital in its origin, being only met with in very young children



(under eighteen months). It grows rapidly and is usually bilateral. The view of Cohnheim is probably correct, that the tumours arise by foetal inclusion. The first rudiment of the uro-genital organs is close to the proto-vertebræ, and it seems probable that some of the germinal muscle cells from the latter have been included with the cells forming the rudiments of the kidney, and have afterwards formed the tumours.

Metastasis has been observed, and in one case the secondary tumours contained muscle.

**Primary cancer.**—This tumour is usually unilateral, although there have been cases of bilateral cancer. The tumour is virtually a cancerous degeneration of the organ. The kidney may be completely converted into a tumour which sometimes attains a very large size, retaining the general shape of the organ and covered by its capsule. The pelvis is also usually recognizable although the cancerous tissue may have grown into it. But in some cases only a part of the kidney is involved, and in that case, while the affected part retains the general shape of the organ, although enlarged, the remaining piece of kidney has quite its normal appearance. To the naked eye it is as if a portion of kidney were transformed, and with the microscope it can be seen, at the margin of normal and pathological, that the tumour is advancing by a conversion of the proper kidney tissue. The epithelium of the tubules is multiplying so as to form the cancerous epithelium, and is becoming irregular in form (Fig. 322), while the cancerous stroma is being formed of the connective tissue of the organ.

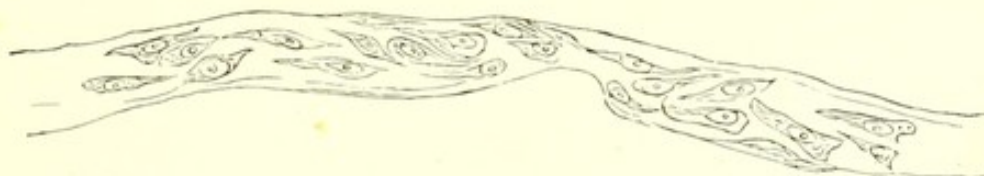


Fig. 322.—A uriniferous tubule at the border of an advancing cancer. The epithelium is undergoing transformation.  $\times 300$ .

In regard to the form of cancer, there are some cases in which the structure has been that of the cylinder-celled epithelioma, in others more that of ordinary cancer. A partial colloid transformation has been observed in a few cases. Whatever the form the tissue seems to originate from the kidney structures.

The tumour is usually very soft, but it is generally confined within the capsule of the kidney. If it passes beyond the capsule it does so more readily into the retro-peritoneal tissue than to the peritoneum, and it may thus extend along the wall of the trunk into the pleura.

The growing tumour not infrequently extends into the pelvis of the kidney and the renal veins. Hence arise hæmaturia on the one hand,



and thrombosis of the veins on the other. The thrombosis may be very extensive, the clot propagating itself in various directions.

In a case observed by the author there was a very extensive thrombosis, involving the veins of both legs. In another case cancerous elements were present in the thrombus, and some of these had been conveyed to the lungs; they were detected along with the clot inside the pulmonary artery, embolism having occurred.

**Secondary tumours.**—Metastatic tumours are not frequent in the kidneys. **Cancers** and **Sarcomas** occur. **Melanotic sarcoma** also occurs.

**Malignant lymphoma** and **Leukæmic** tumours are perhaps the commonest. As seen with the naked eye, they form rounded tumours affecting the cortex chiefly, and causing great enlargement. Under the microscope they present round-celled tissue infiltrated between the tubules.

**Literature.**—*Cysts*—RAYER, *Traité des malad. des reins*, iii.; VIRCHOW, *Ges. Abhandl.*, 871; EVE, *Path. trans.*, xxxi. 164, 1880; PYE-SMITH, *ibid.*, xxxii., 1881, and xxxvi., 1885; THORN, *Beitr. zur Genese d. Cysteniere*, 1882. *Adenoma*—STURM, *Archiv. d. Heilk.*, 1875; WEICHELBAUM U. GREENISH, *Oest. Med. Jahrb.*, 1883; GRAWITZ, *Virch. Arch.*, xciii., 1883; MARCHAND, *ibid.*, xcii., 1883. *Sarcoma*—DICKINSON, l. c.; NEWMAN, l. c. *Myosarcoma*—COHNHEIM, *Virch. Arch.*, lv., 1872, and lxx., 1875; EVE, *Path. trans.*, xxxiii., 1882; WILLIAMS, *ibid.*; RIBBERT, *Virch. Arch.*, cvi., 1886. *Cancer*—WALDEYER, *Virch. Arch.*, lxi. and lxiv., 1875; PEREVERSEFF, *ibid.*, lix., 1874; EBSTEIN, *D. Arch. f. klin. Med.*, xxx.; GAIRDNER and COATS, *Glasg. Med. Jour.*, iii., 1871; BRODEUR, *Affections du Rein*, 1886, p. 170; NEWMAN, l. c.

#### XI.—PARASITES IN THE KIDNEY.

The **Echinococcus** is of occasional occurrence, sometimes along with a simultaneous hydatid cyst of the liver. There is here, as in the liver,

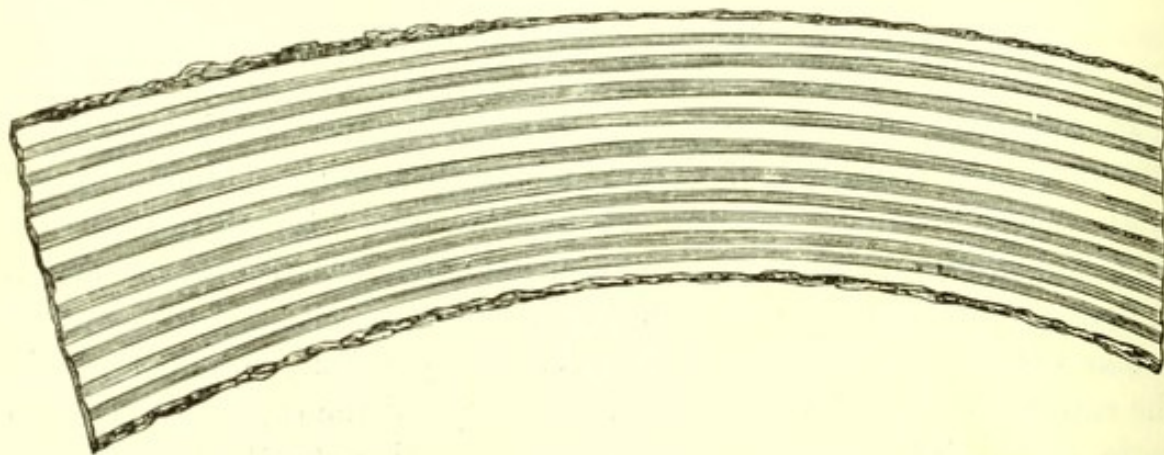


Fig. 323.—Section of a stratified chitinous membrane from an old hydatid cyst of kidney.  $\times 90$ .

a large mother cyst surrounded by a distinct connective-tissue capsule, and containing the usual daughter cysts and brood-capsules. The cyst



not infrequently bursts into the pelvis of the kidney, and the daughter cysts and heads may be evacuated by the urine. Sometimes by obstructing the ureter they lead to hydronephrosis. Rupture also occurs into the pleura.

Sometimes the parasite dies, and the cyst comes to be occupied by a pultaceous material in which the remains of the chitinous membrane are found (Fig. 323). The stratified character of this membrane is here, as in the case of the liver, of assistance in distinguishing the nature of the cyst, especially as all other trace of the parasite, even the hooklets, may have disappeared. The kidney tissue may be considerably opened up and pushed aside by this parasite.

The *Filaria sanguinis* occurs probably in the adult form in the lymphatic vessels of the kidney. The embryos have been met with in the parenchyma of the kidney and in the **Chylous urine**, which is the result of the presence of the parasite (see p. 344).

**Literature.**—ROMESTAU, Des Kystes hydatiques des reins, 1881; BARKER, Cystic entozoa of human kidney, 1856; MANSON, *Filaria sang.*, 1883; LEWIS, Memoirs, 1888.

## B.—THE URINARY BLADDER AND URETHRA.

1. **Congenital malformations.**—These are chiefly represented by **Extroversion of the bladder**, already described at page 48, and **Persistence of the urachus**, which is a minor degree of a similar malformation.

2. **Perforation and Rupture of the bladder.**—Rupture may be produced in various ways. It may be by a direct wound, by fracture of the pelvis, by injuries during parturition. Then ulceration not infrequently produces perforation, ulceration from stone on rare occasions, but most frequently the ulceration of a cancer. Lastly, the bladder may be ruptured by a blow, or by a catheter being pushed through its wall.

These conditions are important chiefly in their consequences, leading, as they commonly do, either (1) to extravasation of urine in the surrounding tissue, or (2) to the formation of fistulous communications with the surface or with neighbouring canals.

Simple **Extravasation of urine** is not in itself serious. The normal urine is a bland fluid, and it may flow from a severed ureter or a ruptured bladder into the peritoneal cavity without producing any peritonitis, the urine being absorbed by the peritoneum and again excreted by the kidney. It is when the extravasted urine undergoes alkaline decomposition that it acquires excessively irritating characters.



The urine being an exceedingly decomposable fluid, and being kept at the temperature of the body, rapidly decomposes if the proper organisms find access to it. The products of decomposition lead to the usual violent septic inflammations associated with necrosis and suppuration.

**Fistulous openings** from rupture of the bladder may be into the uterus or vagina, into the rectum, or on to the surface of the skin. From these fistulæ the urine passes involuntarily as it reaches the bladder, there being no sphincter to retain it. They occur also as a result of perforation of the bladder from without, especially from the uterus and vagina, the cause of perforation being sometimes cancer of these parts, sometimes sloughing from injury during parturition.

3. **Dilatation and Hypertrophy of the bladder.**—A simple dilatation may occur from a sudden obstruction to the urethra, or from paralysis of the muscles concerned in emptying the bladder. In this way a very extreme general dilatation may result.

**Hypertrophy of the muscular coat** is of very frequent occurrence as a result of some obstruction either at the neck of the bladder or in the urethra. The commonest cause is enlargement of the prostate leading to the prominence of the so-called middle lobe at the internal orifice of the urethra.

The muscular coat of the bladder is in the form of bundles of muscular fibre-cells which run in special directions. The muscular coat is therefore not a homogeneous layer, but more like a network of interlacing bands. It is so at least when the bladder is distended, the bands coming closer together as the bladder contracts. When hypertrophy occurs these bundles increase greatly in size, and the internal ones raise the mucous membrane into elongated prominences. As the bundles interlace, the result is that the internal surface of the bladder presents a network of prominent trabeculæ which suggest the appearance of the internal surface of a ventricle of the heart.

As these trabeculæ interlace, little spaces are left between them in the form of small **pouches**. Sometimes these pouches undergo considerable enlargement, and we may have **Diverticula** formed in this way. The diverticulum is originally formed of the mucous membrane pushed out between the thickened muscular trabeculæ. When small it will be contained in the thickened wall of the bladder and emptied during micturition. But as it deepens and projects outside the wall of the bladder it becomes free of the muscular coat, and as it possesses no muscular coat of its own the effect of contraction of the bladder during micturition will be to force the urine into it, just as it is forced into the urethra. The diverticulum is liable in this case to periodical dilatation. The urine also will stagnate in it, and if decomposition occurs, then there



will be inflammatory disturbances in the wall of the diverticulum. Newformation of connective tissue occurs, and, as this tissue is at first soft, the recurring dilatation during micturition causes it to yield so that a continuous enlargement goes on. In this way we sometimes meet with a large sac, usually behind the bladder and communicating with it by a narrow neck (Fig. 324). The sac may be larger than the bladder

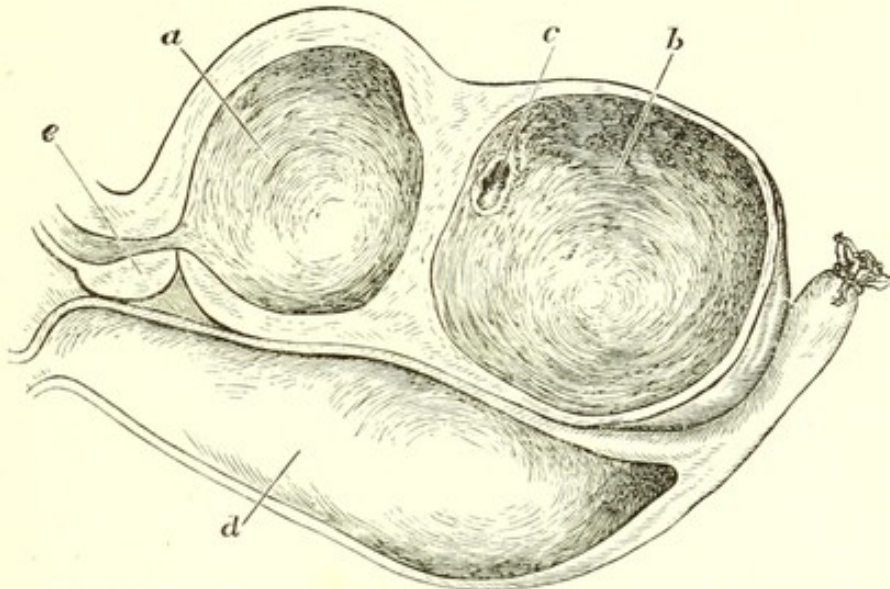


Fig. 324.—Large diverticulum of urinary bladder; *a*, bladder with greatly thickened wall; *e*, prostate and prostatic urethra; *b*, diverticulum with fibrous wall; *c*, aperture between bladder and diverticulum; *d*, rectum. (From a preparation in Museum of Western Infirmary.)

itself, and it presents a somewhat thick fibrous wall with signs of recent inflammation internally.

In diverticula of moderate or large size calculi are liable to form from stagnation of the urine. Or a calculus may slip into such a pouch and escape detection with the sound.

**4. Disturbances of the circulation in the bladder.**—Passive hyperæmia occurs in consequence of obstruction in the veins by tumours in the abdomen or otherwise. Sometimes in such cases the veins of the mucous membrane undergo great dilatation and become varicose, especially in the floor of the bladder, giving rise to **Vesical hæmorrhoids**. The dilated veins may even obstruct the orifice of the urethra, and there is sometimes hæmorrhage from them.

**Hæmorrhage** from the bladder occurs also in consequence of the irritation of calculi, from tubercular ulcers, from papillary and cancerous tumours. There may be hæmorrhages from the mucous membrane in scurvy, hæmorrhagic small-pox, etc. There is sometimes a hæmorrhage following catheterization, and probably due to the sudden relief of pressure from the emptying of the bladder, without a sufficient contraction of the muscular coat. If the bleeding into the bladder be considerable the blood may coagulate, and may be discharged with



some difficulty. Or it may mix itself with the urine, and no consistent coagulum be formed.

**5. Inflammation of the bladder.**—This occurs as a consequence of various kinds of irritation. Stone may produce it, and the extension of gonorrhœa, but by far the most frequent cause of it is alkaline decomposition of the urine.

The decomposition is due to the introduction of microbes and their propagation in the contents of the bladder. They are usually introduced by the catheter. In a healthy bladder such introduction produces no effect, the growth of the microbes being inhibited by the normal mucous membrane. In cases of paralysis of the bladder, or of dilatation, especially when combined with a certain amount of inflammation, the microbes multiply. This is particularly the case when there is stagnation of the urine, and the bladder has to be relieved at intervals by catheterization. Besides being carried in by the catheter, the bacteria may find other means of entrance, as by fistulous openings. It is not impossible that in the case of the short female urethra they may propagate along that passage from the vagina.

The products of decomposition produce the usual inflammatory manifestations with various degrees of violence. In very acute cases there is great swelling of the mucous membrane, it may be with superficial or deep sloughing. In more chronic cases the mucous membrane gets thickened and very frequently becomes the seat of ulceration, so that with thickening and ulceration there is very great irregularity of the surface, sometimes with polypoid projections. The surface is occasionally incrustated with phosphates deposited by the alkaline urine. The muscular coat is often thickened, especially when there is at the same time obstruction to the passage of the urine, and there is the usual trabecular appearance, but obscured by the thickening of the mucous membrane. The bladder may undergo great contraction in consequence of this chronic inflammation, the new-formed tissue in the mucous membrane shrinking.

The urine contains the inflammatory exudation as well as the products of decomposition. In acute cases there may be considerable quantities of pus. In more chronic cases the urine is thick and gelatinous like a mucous secretion. The toughness does not, however, depend on the presence of mucin, but is occasioned by the albumen of the inflammatory exudation being acted on by the alkaline salts in the urine. It is well known that the presence of pus in the urine may be detected by adding an alkali; the urine assumes a gelatinous character. The pus and other inflammatory products in the bladder are similarly acted on when the urine becomes alkaline, and we have the viscid character



referred to. Under the microscope the urine presents abundant pus corpuscles and epithelium, with immense numbers of bacteria and crystals of phosphates.

**6. Tuberculosis of the bladder.**—This condition is usually only a part of much more extensive tuberculosis. There is often a coincident tuberculosis of the kidney and ureter (see above). The vesiculæ seminales, vas deferens, and testicle are frequently affected, and the bladder affection is often due to propagation from the testicle.

The tuberculosis is in the form of ulcers of the mucous membrane which at first are circular, but acquire various shapes by coalescence (Fig. 325). The ulcers are mostly superficial and their edges very

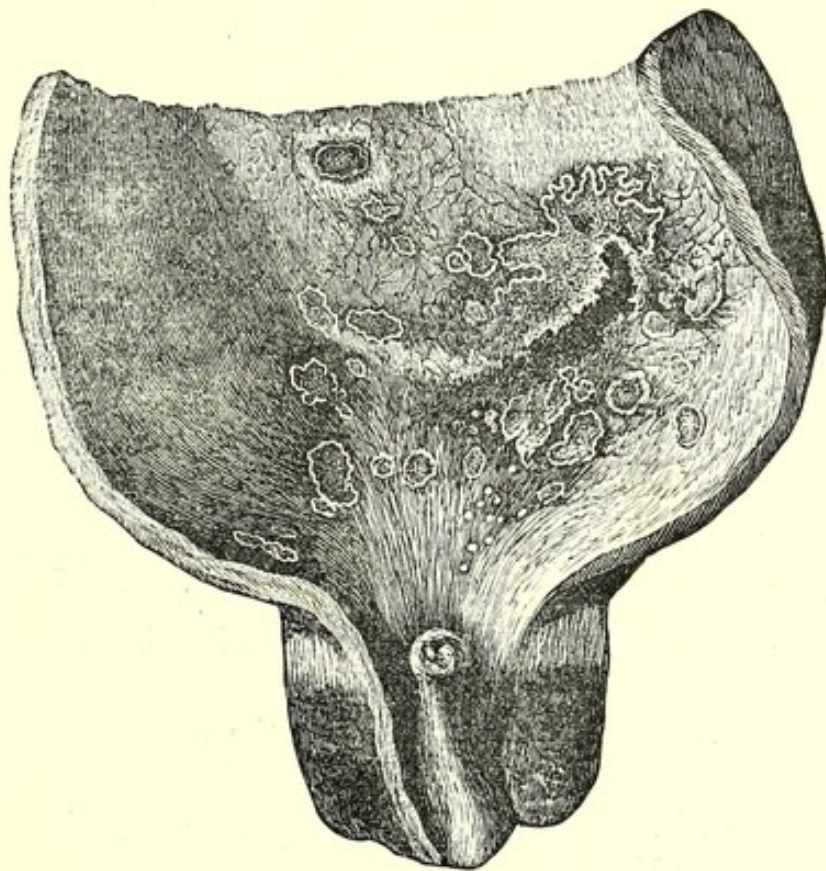


Fig. 325.—Tuberculosis of the bladder. Many round or serpiginous ulcers are shown. These are superficial, but are defined by their white edges which represent recent tubercles. There are several white spots, indicating tubercles which have not yet ulcerated. (VIRCHOW.)

slightly raised, the latter having a pale colour so that the ulcer is surrounded by a whitish zone. The disease begins by the formation of tubercles in the mucous membrane. These break down and form ulcers, which continuously extend by the further tuberculosis of their walls and the breaking down of the infiltration. The general appearances of these ulcers are very well shown in the annexed figure from Virchow's work on tumours. Their superficial character renders them liable to be overlooked.



7. **Tumours of the bladder.**—A few cases of **Fibroma** and **Myoma** have been described, but they are rare. A much commoner tumour is the **Papilloma**. This is met with for the most part in the inferior parts of the bladder, and forms either a prominent mass with projecting papillæ or else a surface covered by villi. The tendency of these villi to bleed is an important fact. Sometimes the villi break down and an ulcer forms. In any case they discharge abundant epithelium, which is to be found in the urine, and is not to be taken as evidence of the existence of cancer.

**Cancer** of the bladder is mostly met with in the form of villous cancer, the wall of the bladder beneath the villi being infiltrated with the cancerous structures. In some cases the structure gives way and a cancerous ulcer with raised edges is the result. There are also cancers without any villous projections of the surface. The cancer may extend to neighbouring structures, but it is much more common for a cancer originating in the uterus or rectum to extend into the bladder than for the reverse process to occur. We have already seen that fistulous communications occur in this way.

**Cysts** have been found, especially in the posterior wall of the bladder.

8. **Parasites.**—If we except the bacteria already referred to, parasites in the bladder are of secondary importance. We have already seen that the echinococcus may burst into the pelvis of the kidney, and portions of the parasite will pass into the bladder. Ascarides and oxyurides have been found to wander into the bladder. The distoma hæmatobium, as we have seen, by the penetration of its ova may produce considerable irritation and hæmorrhage. The filaria sanguinis is also found in the bladder as in the kidney and its pelvis.

Sarcinæ have been occasionally found in the urine. From what has already been stated as to their development from the blood, their occurrence here will be understood. They are met with in cases of inflammation of the bladder.

9. **Concretions and Calculi in the bladder.**—The most frequent constituents of urinary calculi are phosphates, uric acid and oxalate of lime. Phosphates are precipitated from alkaline urine, uric acid and oxalate of lime from acid urine. Phosphates are deposited abundantly when urine, after being passed, undergoes its usual alkaline decomposition; uric acid and oxalates are thrown down in crystals when the urine, at the time of evacuation or afterwards, is unduly acid, the deposition of oxalates occurring usually some time after the urine has been passed, so that this precipitate often, as it were, powders the surface of other deposits.

Inside the bladder or the pelvis of the kidney, phosphates may be



deposited because of undue alkalinity, especially in the case of alkaline decomposition of the urine. These often form an external coating on other stones when, from decomposition, the urine has become alkaline. Phosphates are also frequently deposited on foreign bodies which have found their way into the bladder.

In the case of **Uric acid and Oxalate of lime** it seems that sometimes their precipitation is due to an excess of them in the urine. This applies especially to the oxalates, and there are undoubtedly cases in which there is almost continually an excess of oxalate of lime in the urine (oxalic acid diathesis). But in both cases the substances are deposited from an acid urine, and in the case of uric acid it must be strongly acid. The cause of this abnormal acidity is not always clear. In some cases it may be due to excessive development of acid in the stomach.

The formation of calculi of uric acid and oxalate seems always to begin in the kidney, and even in some cases in the uriniferous tubules. In these and in the pelvis it is very common to find small concretions of uric acid, forming the so-called **Sand and gravel**, composed of aggregates of uric acid crystals, usually rounded in shape and of a brownish red colour. There may even be large concretions of uric acid in the pelvis of the kidney. In connection with the formation of the uric acid and oxalate calculi it is interesting to observe that the two bodies are often combined in one calculus, the mulberry stone especially having frequently a nucleus of uric acid, and perhaps layers of it alternating with the oxalate.

**Forms of calculi.**—We give here a brief description of the different forms of calculi, with an indication of the chemical methods for determining their constitution. As a rule it is convenient to have the calculus sawn through the middle with a lapidary's saw, so that the arrangement of its layers may be seen and the character of its nucleus.

**The uric acid calculus** is the most frequent form. It is usually a small oval stone, with rounded prominences regularly distributed over the surface. The colour varies from a light fawn to a deep brick red. It is heavy for its size, and of hard consistence. Uric acid is insoluble in water and dilute acids, but very soluble in caustic alkalies and weak solutions of alkaline carbonates. A convenient test is the *murexid* reaction. A fragment of the calculus is treated with a drop of strong nitric acid and heated. Effervescence occurs, and the heat is continued till a dry yellowish red residue remains. When caustic ammonia is added to the residue a bright violet red hue is developed.

Calculi formed of urates are rare, the salts being urates of ammonia



and magnesia. They are small soft concretions formed in the kidney, and are distinguished by their solubility in boiling water. They hardly deserve to be reckoned among the vesical calculi.

The **Oxalate of lime or Mulberry calculus** is a very important form. Sometimes small stones are discharged as gravel, forming smooth, round, greyish balls like hemp seeds. The calculus proper is mostly of an irregularly spherical shape, tuberculated on the surface like a mulberry, and of a greyish or nearly black colour. On section it is seen to be in layers, some of them generally composed of uric acid, which usually forms the nucleus. The calculus contains abundant organic material which holds the colouring matter, so that when the oxalate is dissolved out the organic basis often retains the shape of the calculus. Oxalate of lime is insoluble in the alkaline carbonates and organic acids, but soluble in nitric and hydrochloric acids. If a fragment be heated on a piece of platinum before the blow-pipe it becomes black, swells up and leaves a bulky white ash of caustic lime which gives a strong alkaline reaction with litmus.

Calculi of **Basic phosphate of lime** alone are very rare. They form comparatively small yellowish or greyish white stones, rather hard and smooth on the surface.

The **Mixed or Tribasic phosphatic calculus** is very common, at least many calculi are partly formed of phosphates, although few are so entirely. The phosphates are deposited from alkaline urine as a light, bulky, white substance, which is commonly very brittle. The salts are insoluble in water and alkalies, but are very soluble in acids. When a fragment is heated in the blow-pipe flame, the salts melt and form a hard enamel; hence this form is often called the **Fusible calculus**.

The **Carbonate of lime calculus** is rare. It forms small, round, soft stones. The salt dissolves with effervescence on adding an acid, leaving an organic matrix of the shape of the stone.

**Cystine** forms calculi in persons who are subjects of **Cystinuria**. It appears as if, by a congenital derangement of the nutritive processes, such persons form cystine, to a certain extent probably in place of uric acid; and this peculiarity occurs frequently in several members of the same family. The urine, continuously or frequently, contains flat hexagonal crystals of cystine, and these may be already present in the urine at the time of evacuation, or may be deposited after the urine has stood for a time. The cystine may begin to be deposited in the uriferous tubules so that calculi are formed in the pelvis of the kidney, or it may be deposited in the bladder. The stones are oval in shape and have a waxy consistence. The surface is brownish or greenish yellow in colour, and crystals can often be separated from it. The stone may



be buried in a shell of phosphates. Cystine is soluble in alkalies, mineral acids, and oxalic acid.

**Xanthine** calculi are exceedingly rare. The substance is allied to uric acid, and the stones are like those of uric acid but of a redder colour. On applying the murexid test to a fragment of such a stone, it is found to dissolve in nitric acid but without effervescence; the addition of ammonia gives an orange colour.

#### DISEASES OF THE URETHRA.

**Injuries** to the urethra are chiefly important on account of their tendency to lead to stricture. Falls on the perinæum when sufficiently severe to fracture the pelvis usually cause rupture of the urethra. This is followed by extravasation of urine, which may lead to serious results. In case of recovery the wound in healing draws together and leads to stricture, which may even amount to obliteration of the canal.

Injuries are also frequently inflicted from within by the passage of bougies and catheters.

**Inflammations.**—The most frequent form is **Gonorrhœa** which we have seen to result from the action of the gonococcus (p. 295). The mucous membrane in the acute stage is red and swollen and there is a purulent discharge, mixed with blood. The inflammation sometimes extends to the surrounding connective tissue or to the spongy tissue of the penis. There may be abscesses so formed, and in some cases a thrombo-phlebitis occurs, with resulting pyæmia. It may also extend to the bladder. The acute stage passes off and usually leaves a chronic inflammation which frequently results in stricture.

Other forms of inflammation are rare. There may be a simple catarrh, especially in the female urethra, propagated probably from the vagina.

**Stricture.**—Obstruction of the urethra occurs, as we have seen, in consequence of injuries. As it is mostly the membranous part of the urethra which is torn, the resulting stricture has its seat there.

Gonorrhœa is the most frequent cause. The chronic inflammation, which so frequently remains after the acute stage of gonorrhœa, commonly concentrates itself in the most dependent part of the canal, which is the point of union between the membranous and spongy portions or the first part of the spongy portion. Here the mucous membrane remains swollen, and, as the chronic inflammation continues, connective tissue is formed, both in the mucous membrane and for some distance around. The new-formed tissue is, as in other cases of chronic inflammation, dense, and possesses a tendency to contract. Its contraction narrows the canal, which may be found embedded in an ex-



ceedingly dense, almost cartilaginous tissue. There is seldom an actual obliteration of the canal, such as occurs more readily in traumatic stricture.

In cases of stricture **False passages** are frequently formed by the catheter. These have their distal aperture beyond the stricture, and after burrowing through beneath the mucous membrane, either join the urethra on the proximal side or pass on to the neck of the bladder before forming a communication.

The urethra is dilated on the proximal side of the stricture, and this dilatation may be propagated to the bladder, and sometimes to the ureters and pelvis of kidney. Sometimes the dilatation of the urethra, by widening the neck of the bladder, causes paralysis of the sphincter. Hypertrophy of the bladder is a common result of stricture.

**Tumours** are very rare in the urethra. **Carunculæ** are limited polypoid outgrowths from the mucous membrane, of very rare occurrence. **Tuberculosis** occurs in association with the same disease of the bladder, prostate and vesiculæ seminales. **Cancer** is met with chiefly in the deep parts of the urethra by propagation from prostate or bladder, and in the distal parts by propagation from the glans penis.



## SECTION IX.

## DISEASES OF THE GENERATIVE ORGANS.

HERMAPHRODITISM, *implying co-existence of both sexes, chiefly a Pseudo-hermaphroditism in various forms.*

## SUBSECTION I.—DISEASES OF FEMALE ORGANS.

- A. UTERUS, VAGINA, TUBES, AND OVARIES. I. MALFORMATIONS—1. *Defective formation.* 2. *Duplicity of uterus and vagina.* II. MISPLACEMENTS OF UTERUS—1. *Prolapse or descent.* 2. *Prolapse of vagina.* 3. *Inversion of uterus.* 4. *Flexions and versions.* 5. *Other displacements.* III. THROMBOSIS AND HÆMORRHAGE. IV. ATROPHY, HYPERTROPHY, AND DILATATION OF UTERUS. V. INFLAMMATIONS—1. *Of the uterus* (a) *Endometritis*, (b) *Metritis*; 2. *Around the uterus* (a) *Salpingitis*, (b) *Perimetritis*, (c) *Oophoritis*, (d) *Parametritis.* (*Puerperal fever.*) VI. EXTRA-UTERINE PREGNANCY, *chiefly tubal.* VII. SYPHILIS AND TUBERCULOSIS. VIII. TUMOURS OF UTERUS. 1. *Myoma*, 2. *Cancer*, 3. *Sarcoma*, 4. *Polypi and adenomata.* IX. TUMOURS OF OVARIES AND BROAD LIGAMENT. *Introduction*, 1. *Simple cysts*, 2. *Colloid ovarian cystoma*, 3. *Papillomatous cysts of ovary*, 4. *Cysts of broad ligament and Parovarium*, 5. *Dermoid cysts*, 6. *Cancer*, 7. *Sarcoma.* (*Tubo-ovarian cysts.*)
- B. FETAL MEMBRANES AND PLACENTA. 1. *Affections of decidua.* 2. *Hydatid mole.* 3. *Diseases of placenta.*
- C. MAMMARY GLAND. I. *Malformations, Inflammations, etc.* II. *Tumours.* 1. *Adenoma and Fibroma*, 2. *Myoma*, 3. *Sarcoma*, 4. *Cancer in various forms*, 5. *Cysts.* *Parasites.*

## SUBSECTION II.—DISEASES OF MALE ORGANS.

- A. TESTICLE AND TUNICA VAGINALIS. 1. *Malformations and Misplacements*, 2. *Inflammations*, 3. *Syphilis*, 4. *Tuberculosis*, 5. *Tumours, including cysts, sarcomas, enchondromas, and others*, 6. *Hydrocele*, 7. *Spermatocele.*
- B. PENIS, SCROTUM, AND PROSTATE—*Their various diseases.*

## HERMAPHRODITISM.

THIS name implies the union of the two sexes in the same individual. So far as the internal organs are concerned such a condition is rendered possible by the fact that in every foetus the embryonic structures for both sexes are present at a certain period of development; the Wolffian ducts go to form the male, and the Müllerian ducts the female



organs. It is by the subsequent retrogression of one of these and the preponderance of the other that the sex of the child is determined. In the adult female the testicle is still represented by the parovarium, while in the adult male the ovary is represented by the hydatid of Morgagni which lies beside the epididymis, and the uterus by the vesicula prostatica.

A **True hermaphroditism**, in which ovary and testicle are both represented in the same individual is, therefore, possible, and cases have been recorded in which on both sides both of these glands have been present, one of them, however, generally ill-developed. This would form a true **Bilateral hermaphroditism**. On the other hand there may be a testicle on one side and an ovary on the other, forming a true **Lateral hermaphroditism**. It is necessary, however, in such cases to be careful, and not to conclude that an organ is testis or ovary from its mere position, but to subject it to microscopic examination.

**Pseudo-hermaphroditism** is more common. In it the existence of testicles or ovaries determines the sex to be male or female, but other parts are developed so as to resemble those of the opposite sex, and so produce an apparent combination.

In **Male pseudo-hermaphroditism** the testicles are present, but the other structures, in whole or in part, resemble those of the female. Three combinations are distinguished.

(a) *Complete male pseudo-hermaphroditism* is the condition in which, while the glands are the testes, all the remaining organs, both internal and external, resemble those of the female. This arises by a persistence of Müller's ducts and an imperfect closure of the urethra. It may here be remarked that as the external generative organs arise from the same foetal structures in both sexes, they cannot really represent both the female and the male sexes in the same person. But as the male organs present as it were a further development, chiefly consisting in enlargement of the clitoris and closing in of the urinary passage to form a urethra, we may have, by arrest of development, more or less approach to the condition of the female organs.

(b) *Internal male pseudo-hermaphroditism* comprises cases in which the external organs are those of the male and the testes are present, but from the prostatic urethra a canal arises which represents an elementary vagina with a uterus at its extremity, sometimes with Fallopian tubes. The uterus may be of the regular form but small, or it may be two-horned, or with one horn and a Fallopian tube. We have already seen that the vesicula prostatica is the representative in the male of the vagina and uterus, and the condition we are now considering is an exaggeration of that pouch due to an unusual persistence of the lower part of Müller's ducts. There may be all degrees of this persistence, but the case is not one of pseudo-



hermaphroditism unless there is something that can be called a vagina and uterus, even if very rudimentary.

(c) *External male pseudo-hermaphroditism* is characterized by the external organs presenting the characters of those of the female while the entire internal organs are those of the male. The cases are to be excluded in which there is simply an opening up of the urethra (hypospadias) from arrest of development. There must, in addition, be an approach in the form of the organs to those of the female. At the same time the general form of the body is that of the female. Several such cases have been married as females, and the true sex only discovered on post-mortem examination.

**Female pseudo-hermaphroditism** is of much more rare occurrence than male and is susceptible of similar division. In all these forms there are ovaries, and the variations are in the other organs.

(a) *Complete female pseudo-hermaphroditism* presents the male form of the external organs as well as a portion of the male internal organs, while the ovaries are the glands present. In one case the male organs were complete as far as the prostate, but from this sprang vagina, uterus and Fallopian tubes.

(b) *Internal female pseudo-hermaphroditism* is that in which, with well-developed female organs both external and internal, there are male organs present from the persistence of the Wolffian ducts, tubes passing from the parovarium to the uterus or vagina. This condition is excessively rare, although in ruminating animals it is a normal condition.

(c) *External female pseudo-hermaphroditism* is the form in which the external parts have the characters of the male while the internal have those of the female. It will be understood that an elongation of the clitoris will cause the parts to approach to those of the male. Some cases present nothing more than this, while there are others in which there is a distinct penis, and the opening of the vagina is narrow and concealed. The name hermaphrodite can hardly be applied unless there is as well a type of body approaching that of the male. A case is recorded in which the real sex was only suspected when the person became pregnant.

**Literature.**—FÖRSTER, *Missbild. des Menschen*, 1861; GÜNTHER, *Commentatio de Hermaphroditismo*, 1846.

## SUBSECTION I.—DISEASES OF THE FEMALE GENERATIVE ORGANS.

**Introductory.**—The female organs are specially liable to disease from their functional relations. At the times of menstruation and pregnancy changes occur, some of which, although strictly physiological, border on



the pathological. The anatomical relations of the internal organs of generation in the female have also to be taken carefully into account in connection with the changes in position to which the uterus is especially liable.

## A.—UTERUS, VAGINA, TUBES, AND OVARIES.

### I.—MALFORMATIONS.

In treating of this subject it is necessary to bear in mind that the internal and external organs have separate origins, and that the malformations of the one may have no connection with those of the other.

#### 1. Defective formation of the female organs.—Defects of various

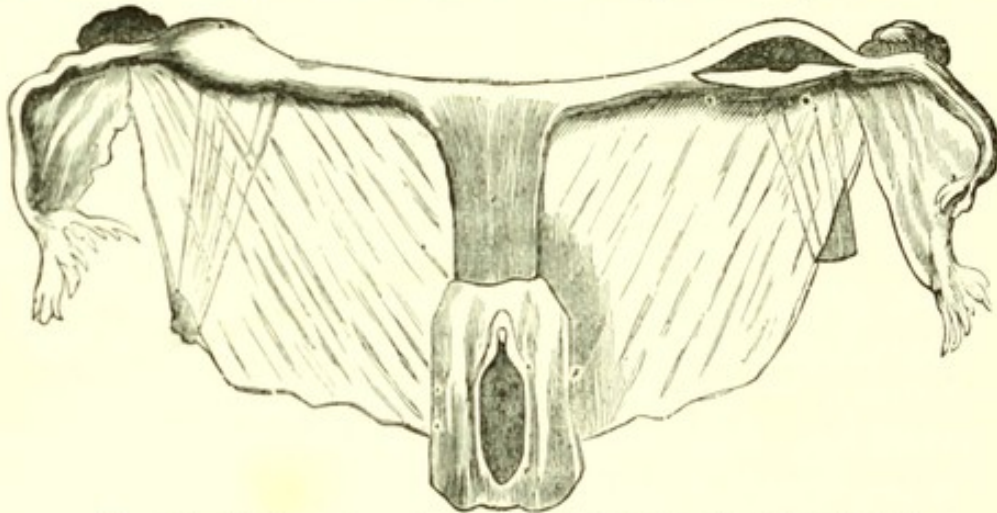


Fig. 326.—Rudimentary uterus. (GRAILY HEWITT from KUSSMAUL.)

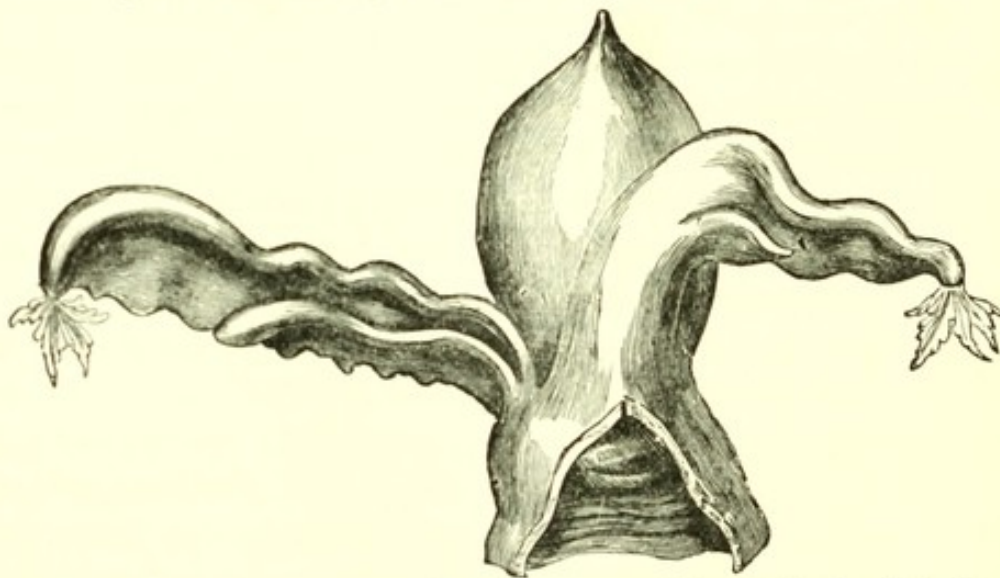


Fig. 327.—Uterus unicornis. The parts are viewed from behind, and the distended bladder occupies the background. The right horn is large, and runs into the Fallopian tube whose fimbriated extremity is shown. There is no proper left horn, the Fallopian tube and round ligament springing from the base of the right horn. (GRAILY HEWITT from KUSSMAUL.)

kinds are met with both in the internal and external parts. The ovaries may be wanting or may remain rudimentary. The uterus may be wanting, and with it the Fallopian tubes; or it may be quite



rudimentary (Fig. 326), presenting perhaps a solid rudiment, or merely two diverging horns. With this the vagina is often defective. Then the uterus may retain in the adult the foetal or infantile form. Again, the uterus or vagina may be imperforate, or the hymen imperforate. There are also various defects of the external organs, as absence of the vulva, the vagina and urethra opening by a small aperture in the region which the vulva should occupy. The hymen may be absent, or it may present fimbriated processes sometimes so large as to project externally.

The *Uterus unicornis* occurs when one Müller's duct is ill-developed. The uterus is a long thin structure which curves to one side, while the other horn is absent or rudimentary (Fig. 327).

2. **Duplicity of uterus and vagina.**—The ducts of Müller in the embryo are destined to form vagina, uterus, and Fallopian tubes, and they are at first double from end to end. The fusion of the ducts occurs soon at their lower extremities so that the vagina and lower part of the uterus early form a single canal. Up to the end of the third month, however, the uterus possesses two horns, the subsequent fusion proceeding from below upwards. In many animals this bilateral duplicity persists during life so that the adult uterus has two horns, often of great length. The duplicity persists in different degrees in the human subject presenting various forms.

(a) *Uterus separatus didelphys* is the most extreme degree of duplicity. The ducts have remained separate, so that from the fimbriated extremities of the tubes to the external orifice of the vagina there are two separate canals. The canals may be partially adherent externally, but their walls have not coalesced and they appear externally as two separate tubes. This malformation does not occur except with other deformities, most frequently with extensive fission of the abdomen, and the foetus does not survive.

(b) *Uterus duplex bicornis* (also *uterus duplex*).—In this, which is not an uncommon form, there is externally one vagina, which may or may not be divided by a septum, but there are two uteri, which may, however, be united externally in their lower parts. Each uterus has a distinct cervix and os, and each is capable of utero-gestation. Even in cases where the bodies are completely separated (as in Fig. 328) the unimpregnated uterus enlarges with the other.

\*

In the case of which Fig. 328 is an illustration death occurred a fortnight after delivery. Both organs were enlarged and almost equally so, the one which had borne the foetus measuring  $4\frac{1}{2}$  inches from os to fundus and the other  $4\frac{1}{4}$ . There had been a previous pregnancy and this had probably been in that which had not borne the foetus on this occasion. This was inferred from the fact that this uterus had adhesions around it and a hæmatocele on its posterior wall.



(c) *Uterus septus*.—In this the parts appear single externally but the cavity of the uterus is divided by a septum which may or may not be continued into the vagina.

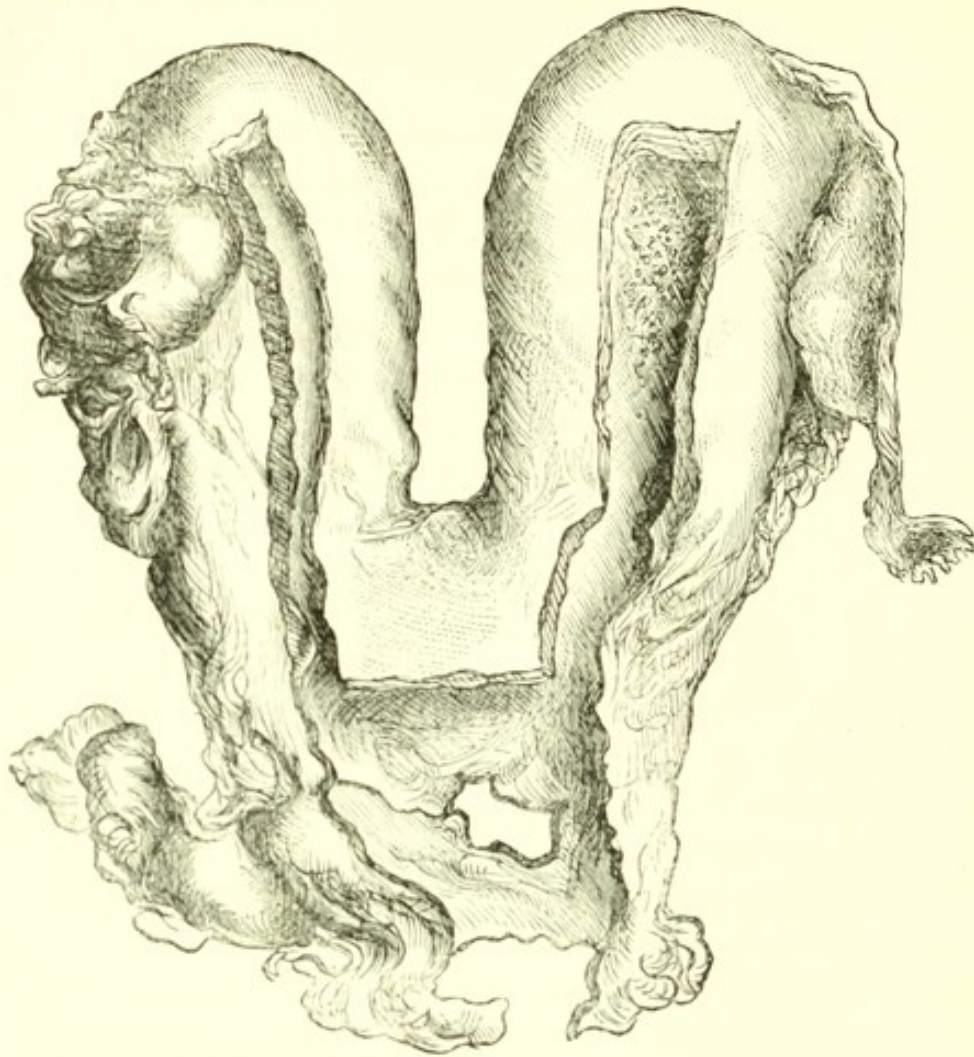


Fig. 328.—Uterus duplex bicornis. The cavities are opened from behind. Foetation had occurred in the right, but the left had also enlarged. The hæmatoecle seen attached to the left may have been from a former foetation.

(d) *Uterus bicornis*.—There is one cavity in the lower part of the uterus, but at the fundus the parts diverge. In some cases there is a mere depression in the middle of the fundus, so as to give the organ somewhat of a heart shape, in which case the term *uterus arcuatus* is used.

(e) *Uterus subseptus* is a form in which the uterus, externally single, is partially divided by a septum.

**Literature.**—KUSSMAUL, Mangel, Verkümmerng und Verdoppelung der Gebärmutter, 1859; MAYRHOFER, in Billroth's Handb. der Frauenkrank., vol. i., part 2, 1878-82.

## II.—MISPLACEMENTS OF THE UTERUS.

The arrangements by which the uterus is supported are of importance to the understanding of the misplacements of the organ. In the virgin the vagina forms a tolerably solid column, on the summit of



which the uterus is supported and so prevented from descending. The vagina is also attached, by means of the pelvic fascia, to the bladder in front and the peritoneum behind. The uterus is further supported by its ligaments, and these, especially the round ligament, assist in preventing its descent, although not so directly as the vagina. As ligaments pass off from its lateral aspects, the uterus is kept from inclining to one side or the other. While capable of very limited movement from above downwards and from side to side, the body of the uterus is very moveable, within certain limits, from before backwards. When the bladder and rectum are full the uterus will be tolerably erect. When the bladder is empty it will be inclined forward, and a certain amount of anteversion may be regarded as the normal condition with an empty bladder.

1. **Descent and Prolapse of the uterus.**—The position of the uterus is lowered for the most part as a result of the loosening of the attachments of the organ, combined frequently with increased weight from chronic inflammation. Descent may also be produced suddenly by increased intra-abdominal pressure, as in raising a heavy weight. Besides increased weight of the organ itself, a tumour by its weight may assist in dragging it downwards. As pregnancy, with its various circumstances, often tends to loosen the attachments of the uterus, prolapse is chiefly met with in married women.

The degree of descent is very various; it may simply amount to a lowering of the position of the uterus an inch or so, or it may be to any extent up to the presentation of it in the vulva or outside.

The term prolapse is, by some, limited to the condition in which the uterus is so low as to present at the vulva. When there is a distinct presentation outside the vulva then the term **Procidentia uteri** is employed.

The vagina must be inverted in proportion to the descent of the uterus, and in cases where it is completely procident the vagina will form an external covering continuous with the skin around. As the uterus descends it is held more by its posterior than by its anterior attachments, and there is accordingly a certain amount of retroversion along with the prolapse.

The mucous membrane of the uterus is mostly in a state of catarrh in prolapse, with a profuse mucous discharge, and the organ itself is enlarged (see further on). The mucous membrane of the inverted vagina is thickened, and its epithelium, where exposed, acquires characters like those of the epidermis.

2. **Prolapse of the vagina.**—This occurs mostly in consequence of pregnancy, and seems to be caused by the walls remaining hyper-



trophied when they ought to undergo the regular involution. The thickened and loose vagina is thrown into folds, and these may project outside the vulva. The prolapse is mostly of the anterior wall, and the urinary bladder is not infrequently dragged downwards with it, forming a *Hernia vesicæ* or *Cystocele*. Much less common is protrusion of the rectum or *Rectocele*.

3. **Inversion of the uterus.**—This is of very rare occurrence. The uterus, which is generally somewhat enlarged, is turned outside in, either by its own contraction or by the exercise of traction on its fundus. These conditions are best fulfilled during or after parturition, especially when the umbilical cord is pulled on while the placenta is adherent. It may also occur in connection with a tumour growing inside the uterus and attached to its internal wall. The inverted uterus projects from the vulva as a bleeding mass, the hæmorrhage being frequently so severe as rapidly to cause death. If the patient survives and the organ is not restored, inflammation results, and the uterus acquires attachments in its new situation, so that resort has sometimes to be had to amputation.

4. **Flexions and Versions of the uterus.**—Flexion is the bending of the uterus on itself, while version is the displacement of the entire organ forwards or backwards.

In flexions the bend takes place at a level corresponding with the os internum, so that the cervix is in one plane and the body of the uterus in another. The reason of this is that the uterus is specially fixed at this level by its peritoneal attachments, the body of the uterus being specially moveable. Flexions are often the result of adhesions due to perimetritis, these adhesions dragging the organ backwards or forwards, and tending to fix it there. Abnormal looseness of the organ, especially after delivery, renders it more liable to bend. When the flexion has become habitual there is apt to be atrophy in the concavity, which renders it difficult to remedy the displacement.

The flexions are divisible into the two forms—**Anteflexion** and **Retroflexion**.

In versions the uterus lies more horizontally than usual, the os projecting in one direction and the fundus in the opposite. They occur from similar causes to those which produce flexions. They are similarly divided into **Anteversions** and **Retroversions**.

These flexions and versions sometimes produce serious results in the uterus itself. It has been mentioned above that, in the concavity of the bend, the uterine tissue frequently wastes and becomes less able to retain the uterus in the upright position. Then the bend, if at all sudden, compresses the vessels, and may lead to a chronic congestion, by and by



resulting in hypertrophy. Again, the curve may obstruct the canal of the cervix, thus leading to dysmenorrhœa. The flexions and versions not infrequently predispose to prolapse. The fundus of the uterus projected backwards or forwards is apt to irritate the bladder or rectum and so induce repeated straining efforts which tend to force the uterus down.

5. **Other displacements.**—The uterus is liable to various other displacements, which, however, mostly stand in a different position to those given above. There is elevation or displacement upwards, from dragging of structures adherent to it, or by pressure from below by tumours or collections of fluid in the pelvis. It is also subject to all sorts of deviations when involved in tumours and inflammations of the pelvic organs.

**Literature.**—FRITSCH, in Billroth's *Handb. der Frauenkr.* iii., 1881; SCHULTZE, *Displacements of uterus* (transl.), 1888.

### III.—THROMBOSIS AND HÆMORRHAGES.

1. **Thrombosis of the uterine veins.**—This is an occasional result of the puerperal state, but sometimes it occurs as a result of tumours of the uterus, and even in affections of the neighbouring parts. The resulting condition is expressed by the clinical term **Phlegmasia dolens**. In the puerperal form the starting point of the thrombosis is the placental surface of the uterus, and it is most apt to occur when, through imperfect contraction of the uterus, the veins are left with gaping mouths. It may be a question whether the introduction of septic material induces the coagulation, but in the usual absence of the general symptoms of septic poisoning it may be doubted whether this has to do at least with the extension of the coagulation. Starting at the uterus, the thrombosis readily extends to the iliac veins and onwards to the femoral and its branches.

The result is often an extensive thrombosis of the veins of the legs, generally beginning in those of the left side. There is usually a hard brawny œdema of the leg.

Thrombosis here, as elsewhere, produces a chronic inflammation of the wall of the vein (*Phlebitis* and *Periphlebitis*), so that there is often considerable adhesion of the vein to its sheath and of the sheath to the parts around. The lymphatic vessels may be affected by this adhesion and partially obstructed.

2. **Hæmorrhages in and around the uterus.**—Hæmorrhage is a normal occurrence in menstruation and parturition, but it may assume pathological characters when in excess or when the blood is unduly retained.

(a) **Menorrhagia** is excessive hæmorrhage at a menstrual period. It



is induced by various constitutional conditions, but also by local lesions of the uterus, more especially tumours. Tumours of the uterus also frequently induce hæmorrhages apart from the menstrual periods.

(b) **Dysmenorrhœa membranacea** is a condition in which membranous structures are evacuated by the uterus along with the blood in menstruation. These sometimes form a complete cast of the interior of the uterus, but more usually they are in smaller pieces. The membrane is variously composed in different cases. Sometimes it is no more than condensed blood-clot or fibrine, perhaps in some cases left over from a previous menstruation. In other cases, however, it is composed of the mucous membrane of the uterus. It is chiefly the superficial layers consisting of epithelium which are exfoliated, but the uterine glands may be present in the membrane, and even the submucous tissue.

This condition is usually regarded as due to an inflammation of the uterine mucous membrane, to which the name *endometritis exfoliativa* has been given, but this view is not universally accepted. There is no doubt some inflammation, as the membrane contains round cells.

During ordinary menstruation, the superficial layers of the mucous membrane are shed, partly in small shreds and partly after previous disintegration. In the intervals there is a restoration of the epithelium by newformation from the remaining epithelium of the ducts and otherwise.

(c) **Hæmatoma of the uterus** is of some importance, as it may resemble a tumour. It consists of a polypoid mass of blood-clot attached to the internal surface of the uterus and hanging into its cavity, or even projecting into the vagina. It is sometimes called the *Fibrinous uterine polypus*. Consisting of blood-clot it has originated in hæmorrhage, but there must be some cause for the adhesion of the clot to the uterine wall. This is mostly afforded by the placenta which has been retained after delivery or abortion. The whole placenta may be retained as in Fig. 329, or it may be only a portion. On the other hand, the rough surface after removal of the placenta may induce the coagulation of blood, which if retained may grow by fresh coagulation. As the hæmatoma originates in hæmorrhage it is usually associated with the latter throughout its course. The blood mostly escapes into the vagina, but some of it may coagulate and increase the size of the polypus.

(d) **Pelvic hæmatocele**.—This name expresses an accumulation of blood in the neighbourhood of the uterus. Two forms have been described according as the blood is in the peritoneal cavity or beneath the peritoneum. A convenient nomenclature is to call the intra-peritoneal form Pelvic hæmatocele and the extra-peritoneal form Pelvic hæmatoma.

**Intra-peritoneal hæmatocele** is much more serious than the other. It



arises as a result of any hæmorrhage inside the peritoneum, such as rupture of an ectopic pregnancy, rupture of an aneurysm, regurgitation of the blood during menstruation when there is obstruction to the regular outflow, rupture of enlarged veins, and of the vessels in adhesions around the uterus.

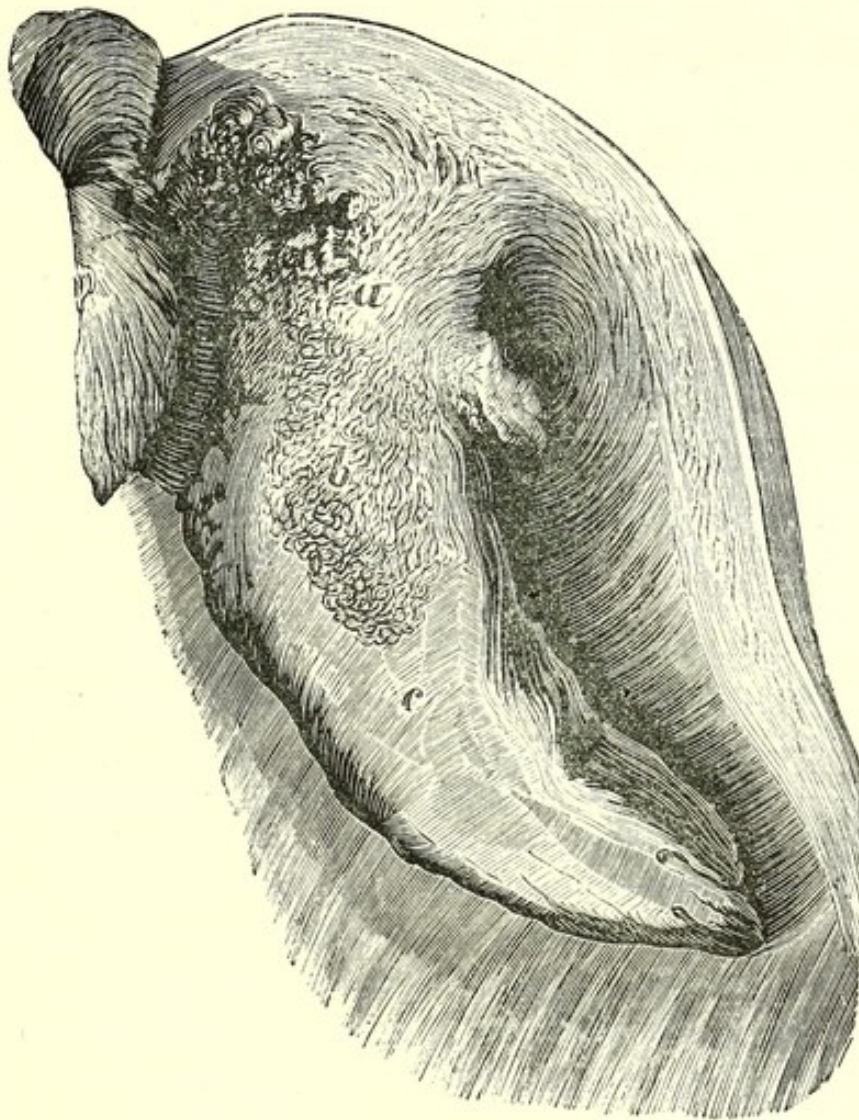


Fig. 329.—Polypoid hæmatoma of uterus after an abortion in the second month; *a*, projecting part of maternal placenta and wall of uterus; *b*, remains of foetal placenta; *c*, stratified coagula around the foetal placenta. Natural size. (VIRCHOW.)

The blood, whatever its source, accumulates chiefly behind the uterus in Douglas's pouch. Acting here as a foreign body it sets up a chronic inflammation with the usual results of newformation of connective tissue which surrounds and causes partial absorption of the blood. There may be thus considerable perimetritis brought about. In some cases suppuration ensues, and an abscess is formed, chiefly where a foetus is present from rupture in ectopic pregnancy.

**Extra-peritoneal hæmatocele or hæmatoma** also originates from rupture of the cyst in ectopic pregnancy, but it occurs also not infrequently as a kind of vicarious menstruation, and may arise in other ways. The



blood mostly accumulates in the broad ligament which it may distend into a bulky tumour. It may also pass around the uterus and rectum, sometimes obstructing the latter. The accumulated blood is generally absorbed, but it may leave thickenings and adhesions behind.

**Literature.**—BERNUTZ, Clin. memoirs on dis. of women (Syd. Soc. transl.), 1866; GUSSEROW, in Volkmann's Sammlung, No. 81; FRIEDLÄNDER, Phys.-anat. Untersuch. über den Uterus, 1870; LEOPOLD, Die Uterusschleimhaut während Menstruation, etc., 1878; BANDL, in Billroth's Handb. d. Frauenkrankh., v.; LAWSON TAIT, Ectopic pregnancy and pelvic hæmatocele, 1888.

#### IV.—ATROPHY, HYPERTROPHY, AND DILATATION OF THE UTERUS.

1. **Atrophy.**—The uterus may retain in adult life the undeveloped condition of that of the child. There may be, on the other hand, a premature atrophy, in some cases ascribed to long-continued catarrh, frequent pregnancies, pressure of tumours, etc., in which the organ anticipates the normal senile involution.

2. **Hypertrophy.**—This sometimes occurs as a result of imperfect involution after the physiological enlargement of pregnancy, and in this case the increase in size is from excess in the muscular substance mainly. Hypertrophy of a similar kind occurs from the presence of tumours in the wall of the uterus. There may be hypertrophy from congestion and chronic inflammation however caused.

A special **Hypertrophy of the cervix** has been observed in many cases. It occurs as a result of imperfect involution after pregnancy, but also as a consequence of prolapse of the vagina, the cervix being dragged down and greatly elongated. With very little descent of the uterus the cervix may be so elongated as to present externally.

3. **Dilatation of the cavity.**—This occurs in consequence of the retention and accumulation of material in the uterus. It results, therefore, from **Obstruction of the vagina** or of one of the orifices of the uterus. There may be congenital closure of the external or internal os (generally the external) or imperforate hymen. When the period of puberty is reached and menstruation begins, the blood accumulates in the uterus, and there may be enormous distension, the contents having a tarry or pulpy character. This condition is designated **Hæmatometra**. The uterus may assume the size of the pregnant organ, and there is thickening of its walls, which, however, are loose. If escape is not provided artificially the uterus may actually rupture, not usually into the peritoneum, but, after the formation of adhesions, into some neighbouring organ. The rupture is by a process of ulceration from within, unless external violence bursts the distended organ.

There may be an acquired **Obstruction of the os uteri** as from chronic



catarrh, or even from inflammation occurring after delivery. In such cases also we may have a hæmatometra from accumulation of menstrual blood. It is more common, however, to have such closure after menstruation has permanently ceased, and in that case a catarrhal secretion may accumulate and distend the uterus. After a time the contents, which are at first mucous in character, become serous, and the condition called **Hydrometra** is brought about.

Lastly, the organ is sometimes distended with **Gas**, a condition designated **Physometra**. This may be the result of decomposition of the accumulated fluid in hydrometra, or from decomposition of retained clots, etc.; but cases have occurred in which the cavity has been dilated with gas and assumed considerable proportions without apparent cause.

#### V.—INFLAMMATIONS OF THE UTERUS AND ITS APPENDAGES.

These are somewhat various, and they are differently named according to the locality specially affected. In this way we have to consider **Endometritis**, **Metritis**, **Salpingitis**, **Perimetritis**, **Parametritis** and **Oophoritis**. The condition of the uterus and vagina after delivery lays them open to the occurrence of septic inflammations, and many of the conditions here have this origin.

1. **Inflammations of the uterus**.—These are divided into inflammations of the mucous membrane and of the muscular substance.

(a) **Endometritis**.—This is an inflammation of the mucous lining of the uterus. We may have an **Acute inflammation** set up by the extension of a gonorrhœal inflammation from the vagina, or in consequence of parturition, or in the course of an acute fever. The inflammation may go on to suppuration or even to sloughing of the mucous membrane and the formation of ulcers. It is apt to extend from the uterus to the Fallopian tube, broad ligament, etc. (see further on).

**Chronic endometritis** or chronic catarrh is a very frequent disease, and, as a whitish discharge is a characteristic feature, the condition is often called **Leucorrhœa**. Apart from the excessive secretion the mucous membrane is apt to become thickened, and it may be thrown into folds or give origin to **Mucous polypi** and **Cysts**. The cervix especially is often thickened, and the os may present ulcerations. During pregnancy inflammations sometimes arise, giving occasion to abortions, adhesions of the placenta, and other lesions. The catarrh may also have its origin in a tumour, a flexion, or a version of the uterus.

(b) **Metritis**.—This name is given to inflammation of the muscular substance of the uterus. Acute inflammation, even with infiltration of the muscular substance with blood, may follow gonorrhœa.

We have also frequently a **Chronic inflammation** leading to **Indura-**



tion, it may be with enlargement of the uterus. This is frequently brought about by **Imperfect involution** of the uterus after parturition, but also results from the various causes which bring about endometritis. The condition consists in a newformation of connective tissue, and from its correspondence with interstitial inflammation in the liver and elsewhere it is sometimes called **Cirrhosis**, especially when there is great induration.

2. **Inflammations around the uterus.**—The structures in the neighbourhood of the uterus are very frequently the seat of inflammation. This occurs by extension of inflammation from the uterus itself, chiefly in cases of **Gonorrhœal** or **Puerperal endometritis**.

The extension of the inflammation occurs by two different paths. The most frequent is by the Fallopian tubes, producing in the first instance an inflammation of them (*Salpingitis*), and then passing on to the pelvic peritoneum and ovaries producing a *Perimetritis*. The other mode of extension is to the subperitoneal tissue, and the result is an inflammation in the loose tissue of the pelvis, a *Pelvic cellulitis* or *Parametritis*. It may be said that the one form usually involves a salpingitis and perimetritis, sometimes with oophoritis, and the other a parametritis.

(a) **Salpingitis** is an inflammation of the Fallopian tube, and it varies in character and intensity. There may be a simple catarrh extending from the uterine mucous membrane, or an acute septic or gonorrhœal inflammation.

A frequent result of salpingitis is **Adhesion and Occlusion of the tube**. The fimbriated extremity is frequently attached to the ovary or to a neighbouring peritoneal surface, this attachment being by connective tissue in the usual fashion of inflammations. The tube is also frequently distorted greatly by the adhesions, doubled on itself, or otherwise altered in position. The uterine orifice of the tube is so small that when the fimbriated extremity is occluded the tube is virtually closed, and the inflammation may completely occlude the uterine orifice as well.

In this case the tube frequently becomes **Distended** with various contents, and names are applied according to the different character of the contents. In simple inflammations a watery or serous fluid may collect (as there are no glands in the tube, the fluid is not mucous in character), the result being a **Hydrosalpinx**. In more acute cases pus distends the tube, **Pyosalpinx**, or blood may be extravasated, **Hæmatosalpinx**. The dilatation of the tubes is sometimes very great, so that a considerable cystic cavity may result.

(b) **Pelvic peritonitis** or **Perimetritis** frequently follows on salpingitis. It may also be that an inflammation of the peritoneum covering the uterus may occur from inflammation of the uterine wall.



The inflammation may be septic, in which case it is suppurative and may extend to the general peritoneum. But even when acute it may be limited by adhesions, and abscesses may form which remain confined to the pelvis. Such abscesses are sometimes so mixed up with adhesions that it may be difficult to distinguish whether they are in the peritoneum or outside it. They may ultimately burst into the rectum or vagina or at the cutaneous surface.

**Chronic perimetritis** is characterized by the formation of **Adhesions** and membranous newformations around the uterus. These are most frequent behind the uterus, uniting it to the rectum. The adhesions are frequently drawn out so as to form long attachments between the parts. Displacements of the uterus and abnormal fixations are frequent results of such adhesions.

The complete picture of perimetritis is that in which the uterus is buried in adhesions which abolish the pouch of Douglas behind, and completely mat together the broad ligament, tube, and ovary, so that the two latter structures are often indistinguishable. The tube may be dilated in the manner mentioned above.

Salpingitis and Perimetritis are of very frequent occurrence. According to the observations in 100 consecutive cases of post-mortem examination in adult females in the London Hospital, the tubes were affected 17 times. From his own experience the author is satisfied that this percentage is not far from the usual average.

(c) **Oophoritis** mostly occurs in connection with perimetritis. **Acute inflammation** of the ovaries is usually a sequel of the puerperal state. A septic inflammation accompanied by pelvic abscess may be associated with abscesses in the ovaries. The pus at first forms in elongated streaks from the hilum to the periphery, but after a time there are more distinct abscesses. The Graafian vesicles also frequently become filled with pus. The abscess in the ovary is surrounded and limited by adhesions like that in the peritoneum, and it is frequently difficult to say what is the actual seat of the organ.

**Chronic oophoritis** also occurs in connection with perimetritis, but it has sometimes a more independent origin. As the ovary is liable at the menstrual periods to great vascular disturbance, we may have, from checking of menstruation and otherwise, a chronic inflammation set up.

The condition has the characters of interstitial inflammation, and is comparable to cirrhosis of the liver or kidney, being, like these, accompanied by shrinking of the organ. The capsule is thickened, and the contracting tissue in the organ produces irregular depressions of the surface. The thickening is often peculiarly manifest around the Graafian vesicles, and this, with the thickening of the capsule, may



prevent the vesicles bursting. Sometimes a ripe vesicle, instead of bursting externally, ruptures into the substance of the ovary, and so produces further inflammatory disturbance. With these changes in the ovary itself there is usually adhesion of the capsule to the parts around,

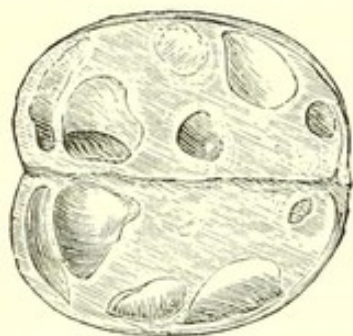


Fig. 330.—Cystic formation in ovary from dilatation of Graafian vesicles. (VIRCHOW.)

the chronic inflammation causing the formation of vascular connective tissue which unites opposed surfaces. In this way there may be displacements of the ovaries.

If the Graafian vesicles are prevented from bursting, the fluid which naturally exists in them may become augmented, and the vesicles thus be converted into **Cysts** (Fig. 330). A limited number of small cysts may thus be formed, and it is not impossible that cysts having this origin may grow to some size, having always the character of simple cysts with serous contents.

It is to be remembered that in old age the ovaries are often shrunk and the capsules thickened, but this is not to be set down as the result of chronic inflammation.

(*d*) **Pelvic cellulitis or Parametritis.**—This consists in a subacute inflammation of the pelvic connective tissue, generally occurring after delivery, but also sometimes as a result of operations on the uterus, the introduction of pessaries or the uterine sound, etc. The inflammation is no doubt septic, being in this respect comparable with erysipelas and phlegmonous inflammations generally. The inflammation extends from the uterus, finding its way apparently by the lymphatic spaces. There are the usual results of inflammation, but the exudation is here the most important. The spaces of the connective tissue get filled up with a serous exudation which may be partly fibrinous. There is in this way a great tumefaction of the subperitoneal tissue, especially of the broad ligament, but also that in front of and behind the uterus and in the pelvis as a whole. The uterus is thus, as it were, fixed in the midst of tumefied connective tissue, which may be felt as a firm swelling on examination per vaginam.

Suppuration generally ensues, but it may do so very gradually, so that it may be long after parturition before it occurs. The pus sometimes extends a considerable distance from the neighbourhood of the uterus. The inflammation may extend and the suppuration follow into the iliac or even into the lumbar region. The abscesses which result open in very various localities, into the vagina, the rectum or the bladder, or at the surface in the iliac or inguinal region. In these latter cases the condition may simulate lumbar abscess, and mistake



is the more likely as the suppuration has perhaps occurred long after the originating cause. The pus discharged has usually the characters of having been long retained, the corpuscles are largely fatty, and there is usually a faecal odour due to the proximity of the abscess to the rectum.

**Puerperal fever.**—This term does not express any single definite morbid condition, but includes cases of septic inflammation connected with the puerperal state, in which acute febrile symptoms occur. The fact that the disease occurs in a quasi-epidemic form and is communicable seems to show that it is due to a specific microbe, but it is probable that any of the more virulent pyogenic micrococci, such as that of erysipelas, may induce it. The course of the septic inflammation and the mode of extension to the general circulation vary, but much that has been stated above in regard to perimetritis and parametritis applies here.

There is, to begin with, an acute septic inflammation of the uterine mucous membrane, usually accompanied by sloughing and suppuration. From this local seat there is an extension, in the manner indicated above, by the Fallopian tubes or by the subperitoneal tissue. In the former case a **General septic peritonitis** results. The septic poison is absorbed, and we have the regular fever of septicæmia. In the other case a **Septic thrombophlebitis** may occur, and we have the phenomena of **Pyæmia** with metastatic abscesses and septicæmia.

**Literature.**—BERNUTZ, l. c.; MATTHEWS DUNCAN, *Prac. treatise on perimetritis and parametritis*, 1868; VIRCHOW, *Ges. Abhandl.*, 1856; WINCKEL, *Dis. of women* (transl.), 1887; BANDL, *Handb. der Frauenkr.*, ii., 1886; HEIBERG, *Die puerperalen und pyæmischen Proc.*, 1873. *Salpingitis, etc.*—LAWSON TAIT, *Brit. Med. Jour.*, 1887, i. 825; LEWERS, *Obstet. trans.*, xxix., 1887; POLK, (also discussion), *Amer. Gynec. Trans.*, xii., 1887; HENNIG, *Krankh. der Eileiter*, 1876.

## VI.—EXTRA-UTERINE OR ECTOPIC PREGNANCY.

By these terms is meant the development of the foetus in any other situation than the normal one in the uterus.

The causation is somewhat obscure, but if we accept the views of Lawson Tait it usually results from disease of the Fallopian tubes, which by destroying the ciliated epithelium allows the spermatozoa to pass up the tube and impregnate the ovum before it reaches the uterus.

According to this author normal impregnation always occurs in the uterus, as the ciliated epithelium of the tubes prevents the ascent of the spermatozoa. If the ciliated epithelium be destroyed the spermatozoa may pass up as far as the ovary.

According to the position in which the impregnated ovum settles various forms have been distinguished, namely, **Tubal**, **Tube-ovarian**, **Ovarian**, and **Abdominal**. The existence of an abdominal pregnancy, except by rupture of a tubal one, is denied by recent authorities (Mayrhofer, Lawson Tait), and even ovarian pregnancy if it occurs is very rare. Undoubtedly tubal pregnancy is by far the commonest. An excessively rare ectopic pregnancy is that in which, from existing



perforation of the uterus (as from a previous Cæsarian[section], the ovum has escaped into the abdominal cavity.

**Tubal pregnancy** occurs either in the free part of the tube or in the part surrounded by the uterine tissue. In the latter case the form is called interstitial pregnancy.

If an impregnated ovum settles in the tube the placenta, as it forms, adheres to the wall of the tube and forms vascular connections. As the tube does not enlarge like the uterus the growing ovum thins its wall, and rupture occurs (*primary rupture*). This always takes place before the fourteenth week. The rupture may be into the peritoneal cavity or into the broad ligament. At the time of rupture there is hæmorrhage, which in the case of rupture into the peritoneum is usually fatal.

As a result of this primary rupture the foetus usually dies, and in that case the conditions are those of the **Pelvic hæmatocele**, intra-peritoneal or extraperitoneal. In either case there may be subsequent encapsuling or absorption of the dead foetus (which will be very small) or there may ensue a suppuration of the hæmatocele.

The foetus may survive and the placenta may acquire fresh connections, so as to allow of the completion of the full term of utero-gestation. Before this, however, especially when the foetus is in the broad ligament, there may be a *secondary rupture*, resulting in some cases in a fatal hæmorrhage, but in others merely in a fresh adhesion inside the peritoneum.

If the full term be reached the foetus dies and, unless removed by operation, remains as a foreign body. As a rule general peritonitis results, sometimes with the ultimate formation of an abscess. If the death of the mother does not occur soon the abscess may come to the surface and the foetus may be discharged piece-meal.

In some rare cases the foetus gives rise to no active inflammation and after its death remains quiescent. It becomes surrounded by a connective tissue capsule inside which the mummified foetus may remain for many years, the condition usually designated **Lithopædion**. The capsule usually becomes infiltrated with lime salts so that a kind of shell is formed around the foetus. The foetus itself may be partly calcified, but its soft structures are often little altered or may be converted into adipocere. Cases are on record of a duration of life extending as long as fifty years after an ectopic pregnancy, and in some cases normal pregnancies have occurred in the interval.

Cases have been observed of molar extra-uterine pregnancy.

**Literature.**—MAYRHOFER, Billroth's Handb. d. Frauenkrankh., i.; LAWSON TAIT, Ectopic pregnancy, 1888 (gives full account and collection of cases of Lithopædion); VIRCHOW, Würzb. Verhandl. i.; SAPPEY, Comptes Rendus, Aug. 27, 1883.



## VII.—SYPHILIS AND TUBERCULOSIS.

**Syphilis** manifests itself in the vagina and vulva in the form of the hard chancre, which may even occur at the os uteri. It does not present in these situations any peculiarities distinguishing it from chancre of the skin elsewhere. Condylomata of considerable size also occur in the vulva and vagina, often forming warty projections.

**Tuberculosis.**—This does not occur by extension from the urinary organs, but on the contrary it frequently begins in the **Fallopian tubes**, the virus being absorbed from the peritoneum. This takes place in almost every case of tubercular peritonitis, but it also occurs sometimes in cases of tubercular ulceration of the intestine. It is stated by Schramm that it occurs in three per cent. of all females dying of tubercular disease.

Tuberculosis of the Fallopian tubes begins at the distal end, and this part is generally found most affected. The mucous membrane is destroyed and replaced by caseous matter which accumulates in the calibre of the tube. The wall is also thickened and infiltrated with the tubercular newformation. These two conditions lead to a great distension of the tube (Fig. 331) so as to resemble some of the conditions

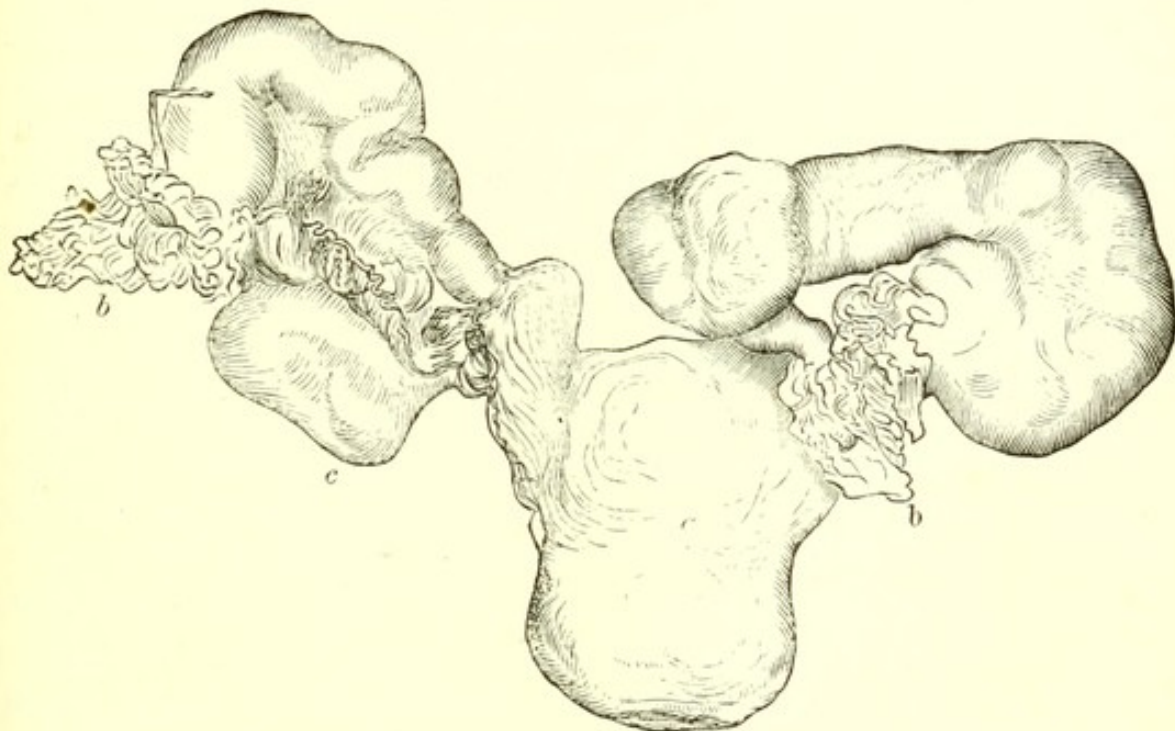


Fig. 331.—Tuberculosis of Fallopian tubes seen from behind. *a*, uterus; *b, b*, fimbriated extremities of the tubes; *c*, left ovary. The right tube is seen to be very greatly dilated and convoluted. The dilatation disappears just before the uterine termination of the tube. The left tube is much less affected, being most dilated at its distal part. There was also tuberculosis of the lungs and mesenteric glands.

resulting from salpingitis. The condition is sometimes called **Tubercular salpingitis**. There is not generally much adhesion to parts around,



but there may be, especially when the condition coincides with tuberculosis of the peritoneum.

The tuberculosis frequently extends to the **Uterus**, where ulcers form, and there may be extensive destruction of the mucous membrane; the whole internal surface is sometimes involved.

Tuberculosis of the **Ovaries** is exceedingly rare, but it occurs in the form of caseous masses.

*Literature.*—MOSLER, Tuberkulose der weibl. Geschlechtsorgane, 1883; SCHRAMM, Arch. f. Gynäcol., xix.; STEVEN, Glasg. Med. Jour., xix., 1883.

#### VIII.—TUMOURS OF THE UTERUS AND VAGINA.

1. **Myoma.**—The myoma is very infrequent in the vagina and in the cervix of the uterus, but is extremely frequent in the body of the uterus. It occurs in from 10 to 20 per cent. of women beyond 20 years of age, and in about 40 per cent. of those above 50.

The myoma presents great varieties in size, number, and in the details of its structure. There may be a single small tumour about the size of a pea, or a large growth weighing more than fifty pounds. There are frequently several tumours present, and there may be fifty attached to the same uterus. The tumour is usually hard and fibrous in appearance on section.

In structure the chief variations are in regard to the proportion of muscle, connective tissue, and vessels. **The muscle** is in bundles which to the naked eye often give a concentric arrangement to the cut surface (see Fig. 333). There is, under the microscope, the usual arrangement of the nuclei in the muscular bundles, as shown in Fig. 69, p. 237. The character of these nuclei and their arrangement are sufficiently distinctive of the tumour.

**Connective tissue** is present between the muscular bundles; sometimes it is dense and, by rendering the tumour compact, gives it a very hard fibrous character. In some cases the connective tissue increases out of proportion to the muscle and a process of induration akin to cirrhosis occurs.

**The vessels** are usually rather spare in the myoma, but sometimes great dilatation of the blood-vessels occurs, so as to give a cavernous character to parts of the tumour, a condition indicated by the name **Myoma telangiectodes**. In other cases there is a dilatation of the lymphatic vessels, leading to a condition called **Myoma lymphangiectodes**.

**Secondary changes** are very liable to occur in the myomas, especially as the tumours frequently grow to large dimensions and are liable to changes in position so as to interfere with the blood-vessels.



**Cysts** not infrequently form in them. These are sometimes from dilated lymphatics, but more frequently from softening of the tumour tissue. Large cysts are thus formed in the midst of large myomas, and may even give rise to a feeling of fluctuation.

**Œdema and Hæmorrhage** are not uncommon, the latter especially, and more particularly in the tumours with dilated blood-vessels.

**Calcareous infiltration** occurs in two different forms. From obstruction of vessels a part of a large tumour may be cut off from its blood supply and become obsolete. In that case lime salts may infiltrate it and they are deposited in all the constituents, the muscle-cells, walls of the blood-vessels, and connective tissue (see Fig. 41, p. 172).

**Separation and Transplantation** are not uncommon in the myoma. The subserous myoma, after becoming pedunculated, may acquire vascular connections with the omentum or other part of the peritoneum and ultimately become detached from the uterus. In a case observed by the author a tumour seven inches in length was attached to a very long great omentum and was very moveable in the abdomen.

The submucous forms also sometimes become detached into the cavity of the uterus, and may either be discharged or retained.

The detached or transplanted myoma is apt to undergo calcification.

In a specimen sent to the author by Dr. Chapman, of Hereford, and depicted in Fig. 332, a myoma was found lying loose in the uterus. It had externally a firm shell which had to be sawn through in order to divide the tumour. The calcification extended to intersecting trabeculæ, which divided the tissue into loculi. In these loculi soft tissue existed which had the microscopic characters of that of the ordinary myoma. It is preserved in the Museum of the Western Infirmary.

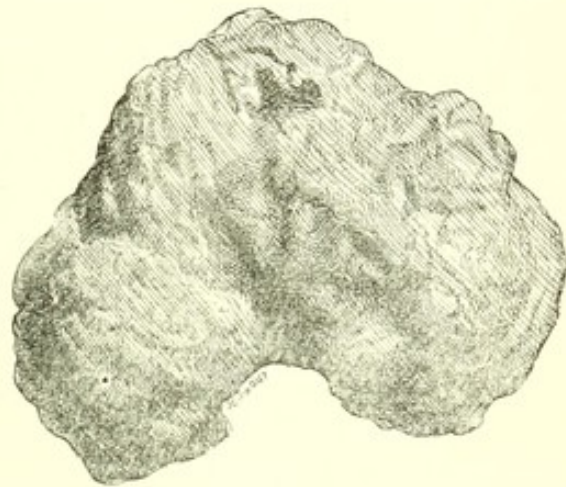


Fig. 332.—Calcified myoma of uterus found lying in the cavity. Half the natural size.

The myoma originates in the muscular substance of the uterus, and it may remain in the wall or become displaced outwards or inwards. Hence three varieties are described.

**The Subserous myoma**, originating in the external layers of the uterus, passes outwards as it grows, and pushes the peritoneal coat before it. In this way it frequently becomes **pedunculated**. The subserous form is often multiple, and as from its situation it is protected, the tumour may grow for many years undisturbed, and reach very large dimensions.



Such large myomas may be mistaken for ovarian tumours and excised as such, and this is the more likely as cysts not infrequently occur in them.

The **Interstitial or Intraparietal myoma** in its growth involves the wall of the uterus, and may cause enormous enlargement of the organ

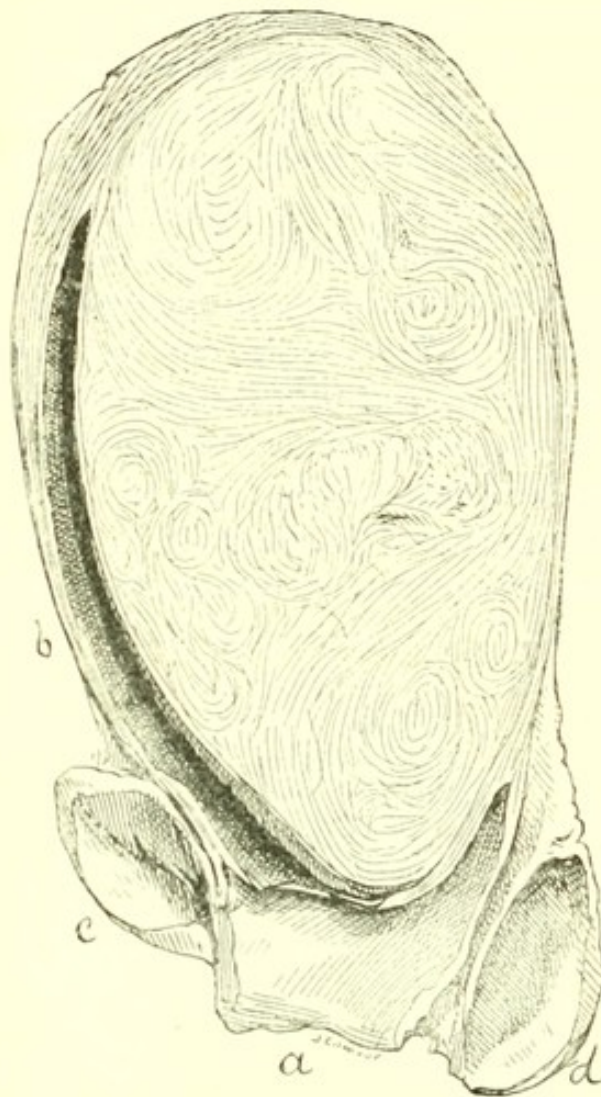


Fig. 333. — Gigantic intraparietal myoma of uterus. It occupies the posterior wall and somewhat distends the os uteri, whose lips were almost of papery thickness. It partly projected into the vagina (*a*). The greatly dilated cavity of uterus (*b*) lay in front. *c*, Urinary bladder. *d*, Rectum.

(see Fig. 333). This form develops mostly at the fundus, and usually occupies the posterior wall. The tumour and greatly enlarged uterus may form together a very bulky mass, which as a whole is liable to be mistaken for an ovarian or other tumour. The author has met with several cases in which the tumour and uterus were excised under this belief. In one of these the tumours seemed to be multiple and the uterine wall could not be distinguished from tumours, the greatly enlarged cavity of the uterus being surrounded by irregularly lobulated masses of muscular tissue. In this case it looked as if the uterus as a whole had undergone an irregular hypertrophy, or had grown into a massive tumour.

The **Submucous myoma** is the form which most frequently comes under the notice of the practitioner. Arising in the wall of

the uterus, it passes inwards, pushing the mucous membrane before it, and from the action of gravity it tends to become pendulous. The submucous myoma, therefore, very often presents itself as a polypus (the so-called *Fibroid polypus*), and it may have a very narrow neck. It very often arises at the fundus, and may grow to such dimensions as to fill the uterus, and hang down through the cervix into the vagina. The mucous membrane covering the tumour is subject to irritation, and there is frequently **Hæmorrhage and Ulceration**, even with sloughing in some cases.

2. **Cancers of the uterus and vagina.**—In the great majority of cases



the cancer begins just about the junction of the uterus and vagina, and involves both as it extends. The disease scarcely occurs before the age of thirty, and is most prevalent between forty and fifty. It appears also from the statistics of West that, contrary to what is sometimes stated, it occurs much more frequently in women who have borne children than in those who have not, and most frequently in those who have had more than the usual number of pregnancies; it is as if the disease developed most readily when the uterus is deteriorated by repeated conceptions.

The cancer mostly consists of large flat epithelial cells, and in most cases these insinuate themselves amongst the constituents of the uterine wall, while in some cases the growth is more superficial. Hence it is possible to distinguish an infiltrating from a superficial form, the latter being frequently designated Epithelioma.

**Infiltrating cancer** begins as an infiltration of a limited part of the

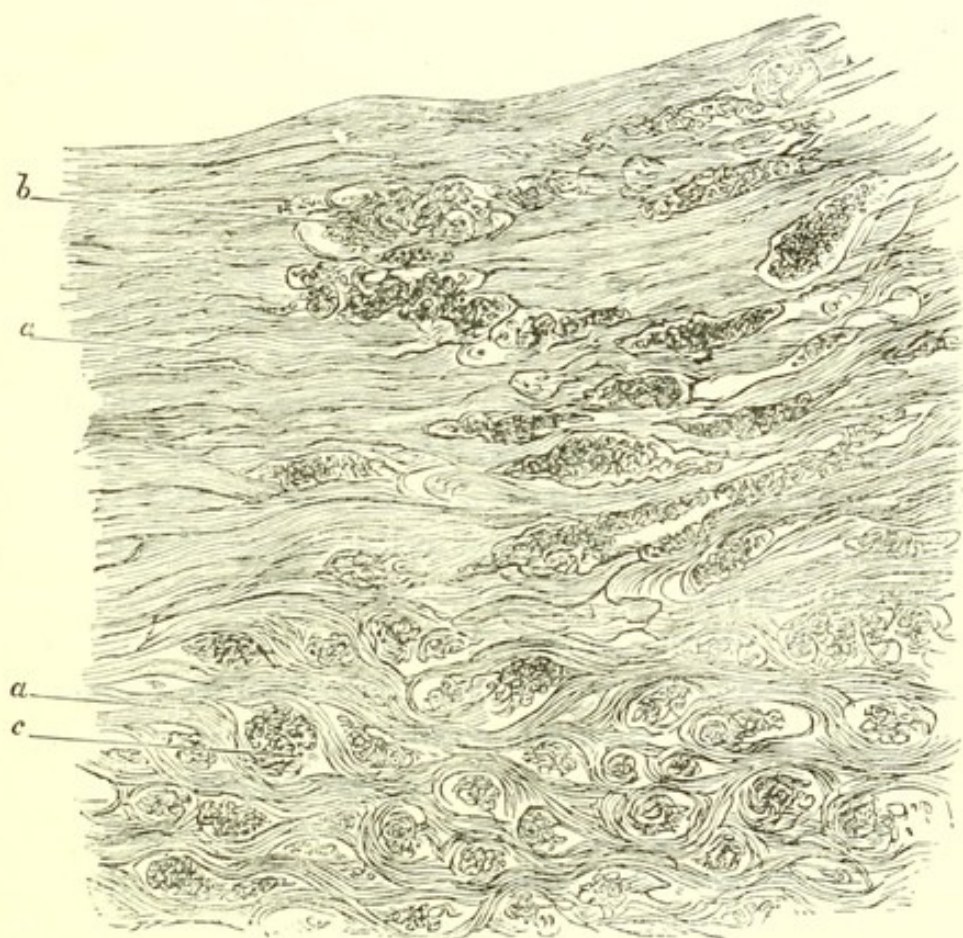


Fig. 334.—Section of cancer of uterus under a very low power, showing mode of advance into wall of uterus; *a*, muscular substance of uterus, interrupted frequently (as at *b*) by masses of cancerous structure. At lower part of figure (at *c*) the muscular substance is still more frequently interrupted and the tissue has quite an alveolar appearance, the muscle partly forming the stroma.  $\times 22$ .

portio vaginalis, and extends more or less round the external os. By and by the whole portio vaginalis is converted into a hard, irregularly



prominent tumour. At first the infiltration is confined to the mucous membrane and submucous tissue, but by degrees it spreads both deeply and laterally. It insinuates itself into the muscular substance of the uterus, separating and breaking up the muscular trabeculae, which come to form a kind of rough stroma for it (see Fig. 334). It also passes into the vagina, infiltrating its wall. Very soon ulceration of the surface sets in, and in its subsequent course there is a progressive ulceration and infiltration, the former following the latter. The infiltration passes into the body of the uterus, but does not usually reach the fundus before

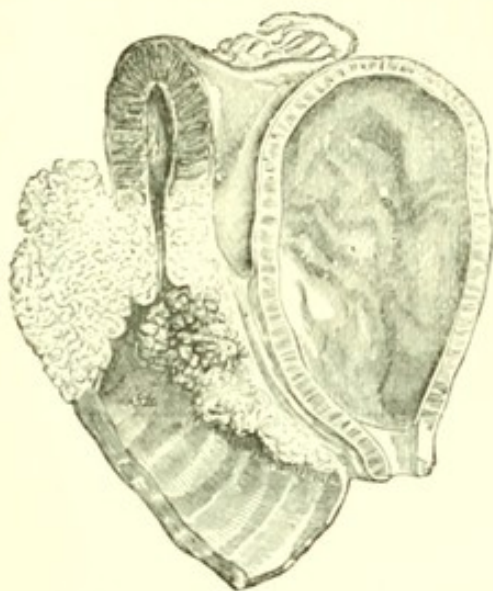


Fig. 335.—Cancer of uterus, the parts shown in section. To the right is the urinary bladder. To the left are vagina and uterus, both of them to a large extent converted into irregular cancerous masses. (GRAILY HEWITT from MARTIN.)

the death of the patient. If the parts be examined post mortem (Fig. 335), it will be seen that an irregularly excavated ulcer occupies the adjacent parts of the uterus and vagina, rendering their respective limits inappreciable. Then outside this there is the whitish cancerous tissue, which extends into the uterine substance some lines beyond the ulcer.

This disease affects neighbouring structures. There are cancerous masses usually in the ligaments and under the peritoneum. The bladder is frequently adherent to the uterine cancer and its mucous membrane red and irregular, or else it presents cancerous nodules. The ulceration even extends sometimes into the bladder, which forms thus a communication with the vagina. The rectum is much less closely related to the cervix uteri than the bladder, and it is less frequently involved. Those parts of the uterus which are not engaged in the cancerous disease are inflamed, and adhesions are formed to the rectum and urinary bladder. Although thus extending locally, the cancer has little tendency to form secondary tumours in the lymphatic glands, and still more seldom does it become generalized.

It sometimes happens that the newformation of cancerous tissue is more vigorous than the ulceration, and in that case we may have prominent ragged masses hanging into the vagina.

The microscopic examination of this form of cancer shows masses of epithelial cells, usually of considerable size, arranged irregularly in alveoli, the stroma being largely formed by the remains of the structures into which it has infiltrated.

**Epithelial cancer or Cancroid** is hardly to be distinctly delimited



from the former, but is characterized by a more superficial outgrowth of prominent warty projections, while the deeper infiltration is slower and less in degree. The disease begins, as in the other case, in the portio vaginalis, and at first there is little more than a prominent warty out-

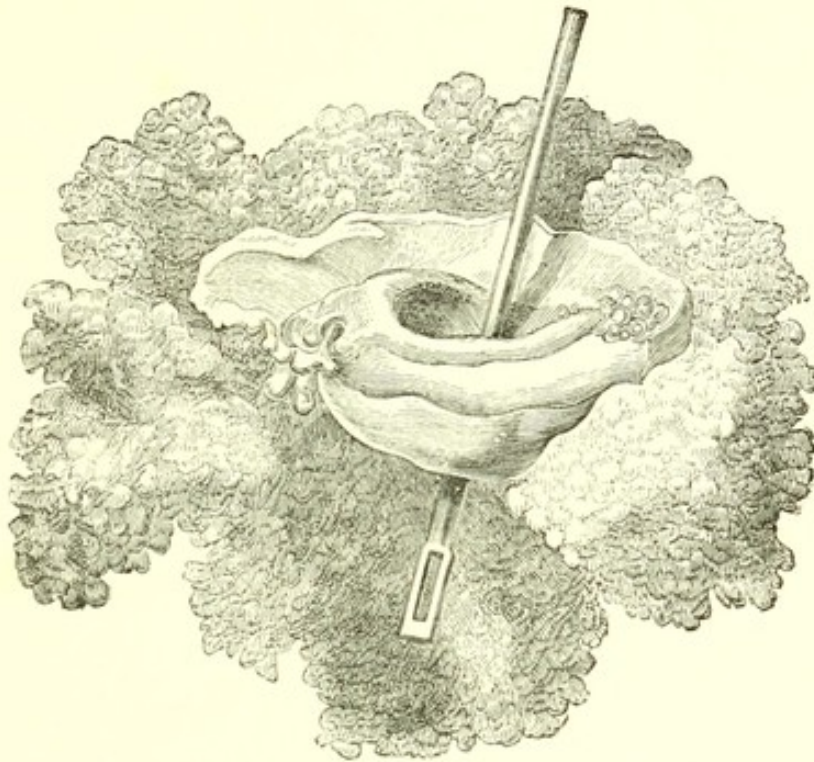


Fig. 336.—Cauliflower cancer of posterior lip of os uteri. A probe is passed through the os, and the anterior lip in front of it is seen to be normal, while a cauliflower growth projects from the posterior. (SIMPSON.)

growth. But the warty growth increases while the base becomes infiltrated till a bulky prominence results, whose surface, consisting of masses of papilliform projections, gives the character of the **Cauliflower excrescence** (Fig. 336).

This form of tumour is also liable to ulceration, and there may be a combination of ulceration with papilliform projections, although, after a time, the papillæ may be destroyed and the appearances approximate to those of the other form of cancer.

Under the microscope the structure here is more that of flat-celled epithelioma. The prominent papillæ are covered with pavement epithelium, and the deeper infiltration consists of masses of flat cells.

A few cases have been observed in which **Colloid cancer** has been the form occurring in the uterus, the situation being the same as in the other more common forms. In rare cases also a cancer may arise from the fundus uteri.

3. **Sarcoma of the uterus and vagina.**—This form of tumour is very rare. It originates mostly in the substance of the uterus, and may form



an extensive infiltration or a limited tumour. The commonest form is the spindle-celled sarcoma, but the round-celled form also occurs. Sometimes, but very rarely, a myoma assumes the characters of a sarcoma, becoming soft and loose in its structure, and its cells more detached from each other. We thus have a **Myosarcoma**.

4. **Mucous polypi and Adenomata of the uterus.**—These are met with mainly as the result of chronic inflammation of the mucous membrane, and may spring from the body of the uterus or cervix. Sometimes there is a general irregular prominence from hypertrophy of the mucous membrane, but usually there are definite polypoid outgrowths.

The mucous polypus may consist of a limited hypertrophy of the mucous membrane, the tumour being a tolerably firm one unless, as sometimes happens, it becomes soft by œdema or by the excessive development of its vessels. In this latter case we may have a tumour approaching to cavernous in character. Or polypi may consist largely of **Glandular** structures, mucous glands being apparently new-formed so as to provide the tissue of the tumour. These polypi are comparatively soft, and may grow to a considerable size, especially when they become **Cystic**. We have already seen that tumours consisting of glandular tissue are peculiarly apt to become cystic, and these form no exceptions. The larger polypi may somewhat distend the uterus.

**Parasites** are rare in the uterus and vagina. Thread worms may pass over from the rectum. Bacteria are met with in the vaginal secretion, and the fibres of leptothrix also occur frequently. In gonorrhœa the gonococcus is present. The oidium albicans is found in connection with thrush. The cysticercus cellulosa has been doubtfully seen in the uterus, and the echinococcus with exceeding rarity.

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## IX.—TUMOURS OF THE OVARY, PAROVARIIUM AND BROAD LIGAMENT.

**Introduction.**—The ovary and broad ligament are very frequently the seats of cystic tumours, which sometimes attain vast dimensions. The origin and mutual relations of the various forms are not completely disentangled, but a knowledge of the anatomical conditions is necessary as a preliminary.

The ovary lies at the back of the broad ligament. It is free on its two



sides and along its convex posterior border. It is attached by its anterior border, which presents a deep groove or *hilum* by which the vessels enter. The ovary is attached to the uterus by a dense **Ligament** (see Fig. 337) and to the Fallopian tube by the **Ovarian fimbria** of that tube (see figure).

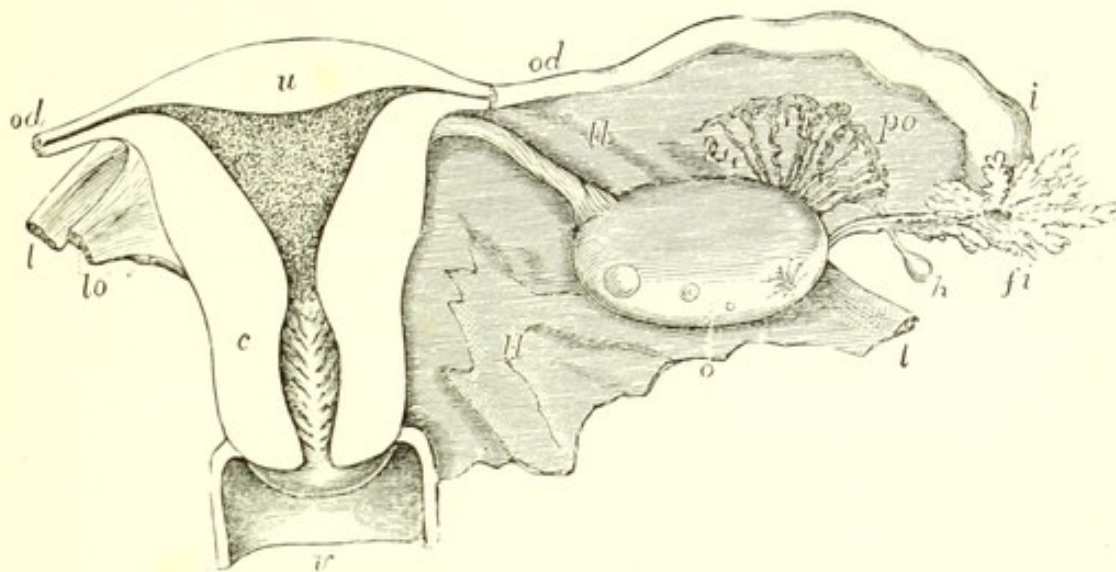


Fig. 337.—Diagrammatic view of uterus and appendages. *po*, parovarium; *od*, Fallopian tube; *fi*, its fimbriated extremity; *o*, ovary, with the ovarian fimbria to the right proceeding to the fimbriated extremity, and a ligament to the left attaching it to the uterus near the origin of the Fallopian tube. (QUAIN.)

The **parovarium**, which is the remains of the Wolffian body, is usually easily made out. It lies between the two folds of peritoneum which constitute the broad ligament, and under the arch of the Fallopian tube, between the latter and the ovary. It is readily seen in most normal subjects, by holding the parts up against the light. It then appears, as in Fig. 337, that the organ is composed of a transverse tube or duct, and of a number of vertical tubes, usually 8 or 10 well-developed. The **transverse tube** is the duct of Gärtner, and it has been traced inwards to the wall of the uterus and onwards so to end in the urethra. This tube frequently ends distally in a small pedunculated cyst as in figure. The **vertical tubes** of the parovarium pass towards the hilum of the ovary, and they partly enter into the formation of the ovary, so that some authors describe a medullary part next the hilum and containing these Wolffian structures, and a peripheral or cortical portion containing the ova.

It follows from these anatomical relations that a tumour of the ovary proper will generally grow into the peritoneum, hanging free with a narrow pedicle. On the other hand, one arising in the parovarium or in the substance of the broad ligament, being sub-peritoneal in its origin, will expand the broad ligament and be seldom pedunculated.

1. **Simple cysts** occur which are in many respects comparable with the simple cysts which are so frequent in the kidney, and like them



have no very special significance (see Fig. 330). They are produced by **Dropsy of the Graafian vesicles**, as was proved by the discovery by Rokitansky of an ovum in such a cyst.

Simple cysts may be of congenital origin, having been observed in new-born children.

As a general rule there are several cysts simultaneously developed, usually from ten to twenty, but one or a few may attain a preponderating size. The cysts have a distinct smooth lining membrane, with a single layer of epithelium. The contents are mostly clear serum, but they may be dark from hæmorrhage, or turbid from inflammation. The enlargement of the ovary is not generally great in this form of cystic disease; it rarely reaches the size of the fist, and still more rarely that of the head. If there are several cysts, they take shape by mutual pressure.

2. **Colloid ovarian cystoma**.—This form is of much more frequent occurrence and vastly more important. To this class of cysts the name **Cystoma** is properly applied, because they arise by a distinct newformation, there being first produced a preparatory tissue, which goes on to the formation of the cysts. The preparatory tissue is glandular in character, and hence the names **Adenoma** and **Adenocystoma** are sometimes applied. This group includes the characteristic **Multilocular cysts** of the ovary.

These cysts form bulky tumours, and while the tumour itself is formed of a number of larger and smaller cysts, there is nearly always, in the walls of these, more or less solid material which shows various stages in the process of cystic formation. The tumour, sometimes of enormous size, represents the **Ovary as a whole**, and is not merely something added to it, the external covering of the cyst corresponding strictly with the surface of the ovary.

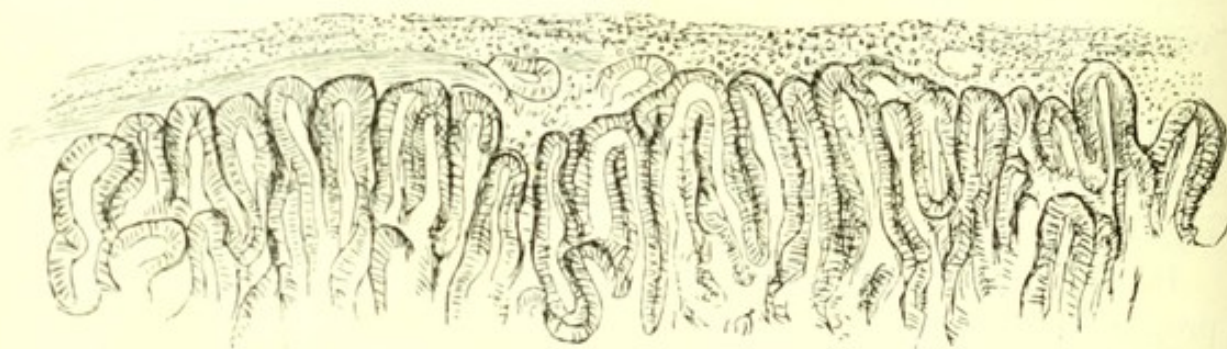


Fig. 338.—Section of tissue in wall of an ovarian cyst. There is a Congeries of glandular structures lined with cylindrical epithelium.  $\times 150$ .

The solid tissue in the wall of the cysts shows the various stages in the process of development. It shows, microscopically, a glandular structure (see Fig. 338) in the form of tubular canals lined with cylin-



drical epithelium. These gland-like structures often project into cysts, and altogether they show a very striking power of newformation. The transformation of the gland-like tissue into cysts is readily seen in many cases. The cylindrical epithelium presents very markedly the goblet form seen in mucous glands (Fig. 339), and it secretes a mucous

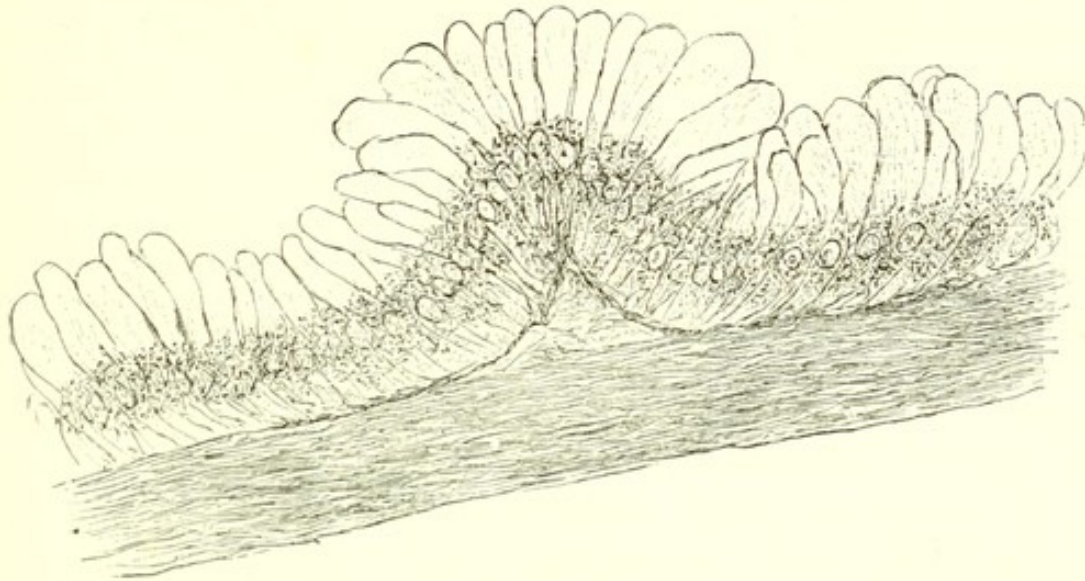


Fig. 339.—Portion of wall of colloid ovarian cyst. The lining epithelium is mostly in the form of goblet cells.  $\times 350$ .

or colloid material, which accumulates in the cavity and distends it more and more into a globular cyst. We have therefore, in the walls of the cysts, very commonly large numbers of smaller cysts which take their shape by mutual pressure, as in Fig. 340. Cysts formed in this way project into the already existing cysts as they grow, and may afterwards burst and become flattened out on the wall of the older cyst. As all the cysts have had a similar origin, they are lined with epithelium, whose function it is to secrete a colloid or mucoid fluid, and so the cysts, once formed, have an almost continuous tendency to increase.

As regards the **Origin and Significance of the glandular tissue**, authors are generally agreed that, as we have here essentially to do with glandular epithelium, its origin is to be referred to the primordial epithelial structures of the ovary. The primordial ovary consists of a layer of epithelium on the surface of a connective tissue projection. The ova form by the penetration or reduplication inwards of this superficial epithelium, which at first forms a series of communicating channels or follicles which Waldeyer compares to a cavernous tissue. By the constriction of these canals the ova are formed, it being doubtful whether the cells of the membrana granulosa which lines the Graafian vesicle are derived from this epithelium or from the connective-tissue stroma. The glandular formation in ovarian cystoma may be regarded as a pathological and exaggerated recurrence of the foetal condition.



The **Naked-eye appearances** of the colloid cystoma are generally quite characteristic. The tumour represents an ovary, and its outer covering is the outer covering of the ovary with its layer of endothelium.

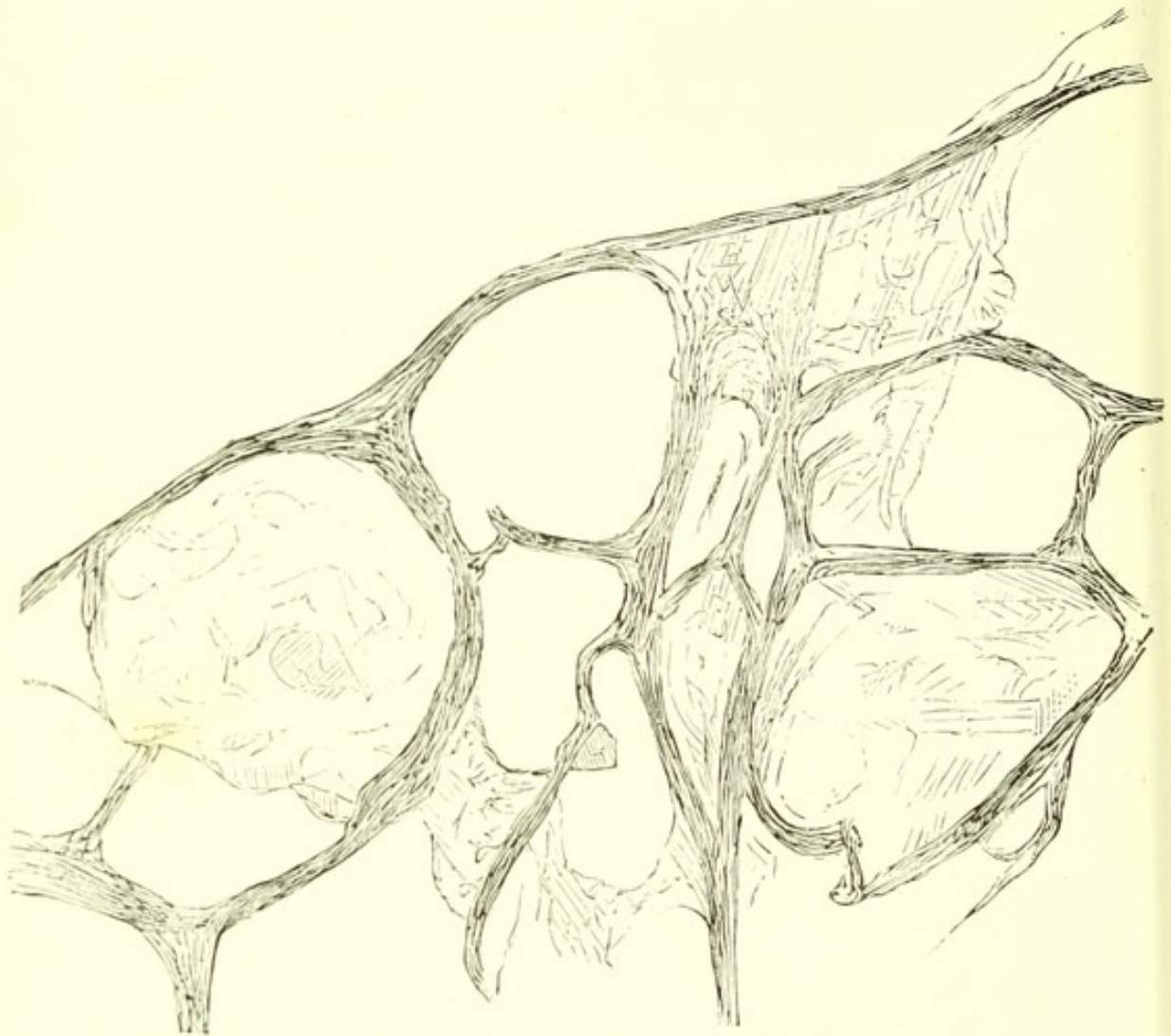


Fig. 340.—Section of portion of a projection on internal surface of ovarian cyst. It consists of a congeries of variously sized smaller cysts.  $\times 20$ .

The surface is usually smooth and very often free from adhesion to neighbouring structures. In shape the tumour is generally more or less globular, but not infrequently it is lobulated on the surface and evidently composed of several cysts. Although the cyst is for the most part obviously multilocular, yet one of the cysts sometimes attains such a preponderating size that the tumour is apparently unilocular. In these cases examination will show the existence of other cysts flattened out in the wall, or collected here and there in clusters. Towards the base of the tumour, but also in many cases at various places, there are solid or semi-solid masses in which cysts are in process of development in the way already described. The size of the tumour is frequently very great, much larger than that of the uterus at the full time. On cutting into it there escapes from the cysts a sticky brownish or yellowish fluid, which is tolerably clear unless some of the secondary changes to be presently



described have occurred in it. If, as is usually the case, there are many cysts, they take shape by mutual pressure against each other, while the general globular outline of the tumour as a whole is preserved.

As the ovary hangs free in the peritoneum the tumour also generally expands outwards freely, and preserves a comparatively narrow attachment, called the **Pedicle**. The expansion is very rarely upwards from the ovary into the broad ligament, but sometimes it is, at least partially, in that direction. If it be so, then the tumour will be more sessile and the Fallopian tube will probably be stretched over its summit.

**Secondary changes** occur in the cysts which concern chiefly their contents. It seems almost a normal condition in large cysts that the internal wall should present collections of cells in a state of **Fatty degeneration**, these cells being often in many layers. Cells also pass into the fluid, so that in all ovarian fluids cells are to be found of round shape and with oil drops in them. In addition to small cells having a few minute oil granules in them (Drysdale's cells), there are usually larger ones with more oil drops, up to large cells having the regular characters of compound granular corpuscles. Many of these cells are doubtless leucocytes which have emigrated into the cysts.

The number of these fatty cells may be greatly increased, and, as many of them in that case break down, we may have a fluid which is turbid from the presence of numerous fatty cells and free oil. The fluid is also pale like a **fatty emulsion**, and in many cases it resembles pus in its physical characters. This change is most likely to occur in old and large cysts, but it is not infrequent in younger and smaller ones.

The free fat, if long retained, forms **Crystals of cholestearine**, and in fact these crystals are frequently found in the fluid of cysts which otherwise are not strikingly altered.

**Hæmorrhage** occasionally occurs into the cysts, and this will cause the fluid to be turbid and deep brown or red in colour. There may also be masses of softened fibrine in the cavity. Sometimes, however, the fluid has a dark brown colour without hæmorrhage.

**Inflammation** of the cyst-wall is not of very frequent occurrence. There may be an acute suppurative inflammation, so that the contents become mixed with pus and assume more and more of the purulent character. With this there is generally an acute inflammation of the surface, with fibrinous exudation and adhesion to neighbouring structures. If the suppuration continues there is apt to be perforation of the pus into the abdominal cavity with resulting fatal peritonitis. A chronic inflammation is more common, causing adhesion of the cyst to neighbouring parts, and these adhesions may be very extensive and firm.



**Perforation or Rupture** of the cyst-wall is not of infrequent occurrence. It has been already stated that coalescence of cysts by rupture of their adjacent walls is a regular process in the course of growth. A rupture externally is much less frequent. It occurs by mechanical violence, by necrosis of parts of the wall from interference with the circulation, from inflammation, or from penetration of the wall by the growth of the secondary cysts.

The rupture occurs usually into the peritoneum, but it may be into the rectum or bladder, inflammatory adhesions having previously formed. The colloid matter in the peritoneal cavity sets up inflammation which may be somewhat acute. If it lasts for some time there is great thickening, especially of the omentum, which contains colloid masses, and looks like a mass of boiled sago (Doran and Olshausen).

**3. Papillomatous cysts of the ovary.**—Cysts of this kind are usually regarded as originating in the hilum of the ovary, and therefore as springing from the remains of the Wolffian body. They form large multilocular tumours with glairy colloid contents like those already described. Instead of a glandular tissue in their walls from which the cysts form, there is a papillary growth into the cavities of the cysts. The intra-cystic growth consists of dendritic papillæ covered with cylindrical epithelium which does not take the goblet characters.

This form of cystoma is sometimes combined with the other, as if the newformation had originated from both parts of the ovary.

The papillomatous cyst has much **more malignant characters** than the ordinary colloid. The growing papillæ not infrequently pierce the wall of the cyst and present themselves externally. There may be thus a considerable shaggy growth outside the tumour. Then there may be in addition transplantation to other parts of the peritoneum, and the papillæ grow where they are planted. We may in this way have shaggy papillomatous growths surrounding the uterus and other structures in the pelvis and even on the general peritoneum. The rupture of the cysts by scattering the papillæ produces serious results.

This form of tumour is likely to be **less pedunculated** than the former kind. Originating at the hilum, it may grow specially into the broad ligament, distending it and stretching the Fallopian tube, somewhat in the fashion of the cysts of the broad ligament. There are, indeed, cases in which it is difficult to say whether the cyst has originated in the ovary or broad ligament.

**4. Cysts of the broad ligament. Parovarian cysts.**—The cysts of the broad ligament occur almost uniformly in that part which lies above the ovary, and contains the parovarium. It is natural to infer that they originate in this organ, but recently doubts have been raised as to this



as the origin of all these cysts. Hence the general term cysts of the broad ligament is now more frequently in use. It is acknowledged that some of them originate in the parovarium, and, according to Doran, it is chiefly those which show papillomatous ingrowths which do so. At the same time, the general characters of all these cysts are otherwise so similar that one is led to infer a common origin.

The cysts of the broad ligament are nearly always **unilocular**. They frequently grow to large dimensions, and they have a perfectly simple connective-tissue wall, without any trace of partitions.

Growing between the two folds of the broad ligament they distend the peritoneum over them, and this **peritoneal covering**, being loosely attached to the proper cyst-wall, can be readily separated. As it has attained to considerable thickness the peritoneal coat itself makes a distinct cyst, like the outer skin of a football. In this respect the cyst differs from the ovarian forms, in which no such layer of peritoneum is distinguishable.

From the position of these tumours, lying between ovary and Fallopian tube (see Fig. 337, p. 881), they stretch these structures greatly. The figure shows that a girdle is formed round this part of the broad ligament, by Fallopian tube, ovarian fimbria, ovary and ovarian ligament. This girdle remains around the cyst but greatly elongated. **The Fallopian tube** is enlarged and elongated till it may measure fourteen or fifteen inches. The fimbriated extremity is flattened out and exaggerated, while the ovarian fimbria is stretched and thickened. **The Ovary** generally hangs from the lower part of the cyst, elongated and condensed. The ovarian ligament completes the girdle. There is thus no proper pedicle such as ovarian tumours usually have.

The contents of the cyst are usually a clear, colourless fluid, of a specific gravity not greatly exceeding 1008.

In most cases the internal surface of the cyst is smooth, but sometimes intra-cystic **papillary growth** occurs, this form being supposed to be specially parovarian. The papillomatous cysts, like the similar ones of the ovary, show malignant characters.

**Rupture** of these cysts is not accompanied by the inflammatory manifestations produced by the ovarian cysts. The papillomatous forms, however, may lead to secondary growths by rupture. The ruptured cyst may lie for years in the abdomen producing little or no disturbance, and without any tendency to refill. This was so in a case observed by Gairdner and the author, and preserved in the Western Infirmary Museum.

5. **Dermoid cysts of the ovary.**—These are sometimes regarded as belonging to the class of **Teratoma**. They are of comparatively fre-



quent occurrence in the ovary, and may be bilateral. The cyst has a

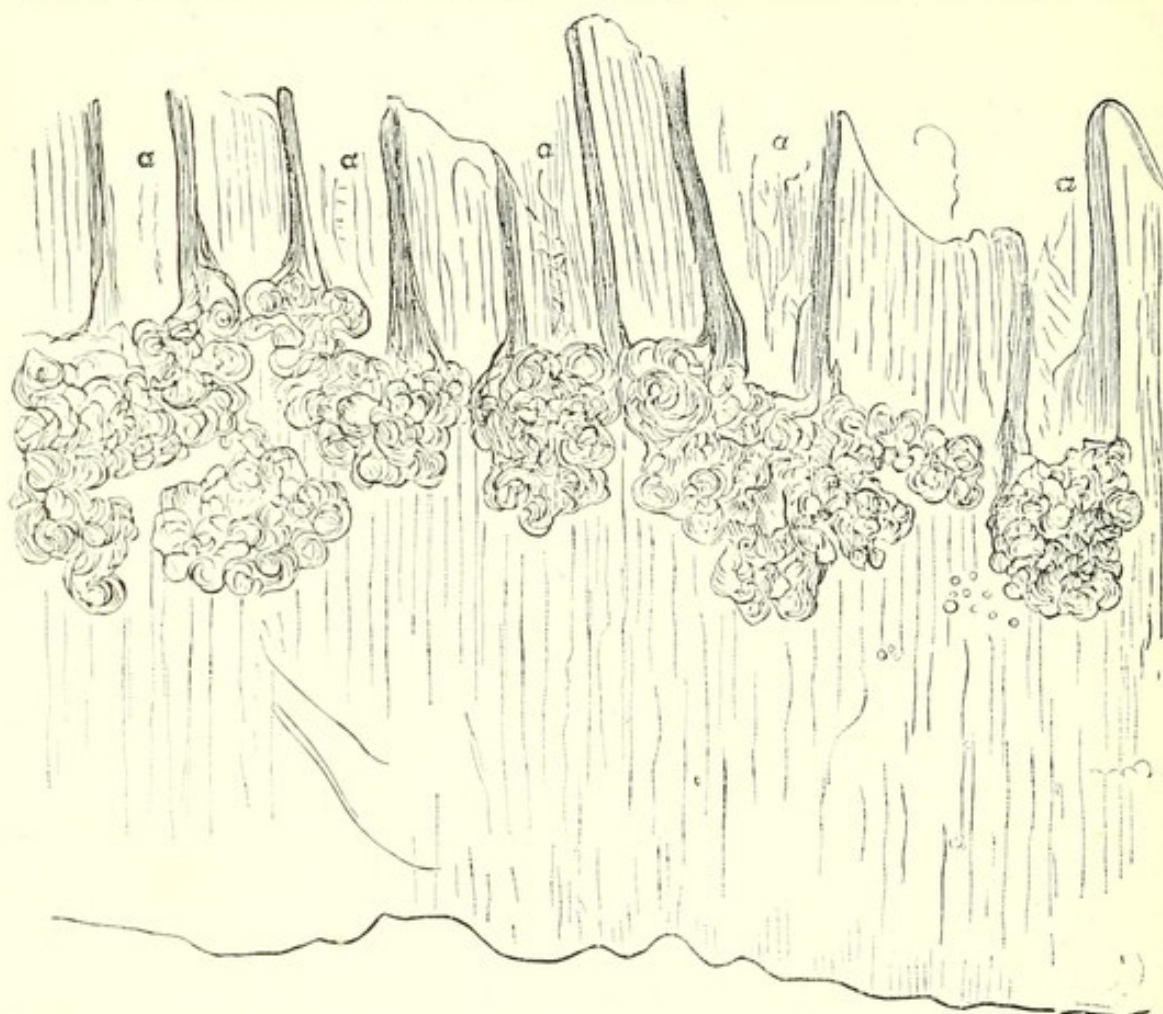


Fig. 341.—Section of wall of dermoid cyst of ovary. A congeries of sebaceous glands with open mouths (a, a, a) occupy the wall. The cyst contained large masses of buttery material with hairs.  $\times 20$ .

thick connective tissue wall, and is generally filled with yellow masses of buttery material in which numerous long hairs are entangled. Large quantities of this material may be present.



Fig. 342.—Hair in its follicle with sebaceous gland, from wall of same cyst as the former figure.  $\times 20$ .

The cyst has an internal lining of epithelium or epidermis, but in certain areas a more complex structure is developed. Thus sebaceous glands (Fig. 341), hairs (Fig. 342), bones, and even teeth may enter into the constitution of the wall.

The sebaceous glands secrete oil which is fluid at the temperature of the body, but on cooling forms the buttery mass. Hairs are shed as from the ordinary hairy parts of the skin, and these accumulate amongst the oil. Epidermic cells are also mixed



with it. Teeth are sometimes present, not only in the walls, but in the contents, sometimes to the number of a hundred.

These cysts not infrequently inflame and may rupture, most frequently into the bladder or rectum, their peculiar contents being discharged, and so revealing the nature of the case.

6. **Cancer of the ovary** occurs mostly as a **Cyst** with cancerous growths in its walls. There may be a combination of the colloid or papillomatous cyst with cancer, or a partial transformation of the former into the latter. In the ordinary ovarian cyst the epithelium has a regular and normal arrangement, and we may call the tumour in that aspect an adenoma; in the cancer the epithelium is distinctly abnormal in its arrangement, being aggregated into indefinite masses. It may happen that in the midst of the cancer the epithelium is undergoing metamorphosis, so that cysts are developing from it as from the more regular glandular tissue.

Besides these cancerous cysts we may have a **Solid cancerous tumour** in the ovary, presenting the usual characters, but, like the ovarian tumours, showing a very excessive growth.

7. **Sarcoma** is a rare tumour in the ovary. Spindle-celled sarcoma is the more usual form, but round-celled sarcomas also occur. These tumours may also assume very large proportions, reaching the size of the head sometimes, and they are not infrequently bilateral. Cysts are frequently present in the midst of them, and these may be simple serous cysts or they may have colloid contents. These latter may arise from glandular structures, and the disease may form a combination of the colloid cystoma and the sarcoma, or they may originate in a softening of the sarcomatous tissue.

**Myoma** of the ovary forms a hard fibrous-looking tumour.

**Fibroma** and **Chondroma** have also been met with, forming dense tumours.

**Tumours of the Fallopian tubes** are excessively rare, but a few cases of papilloma and of cancer have been recorded.

**Tubo-ovarian cyst** is a name given to a condition occasionally met with in which the end of the Fallopian tube is adherent to and communicates with an ovarian cyst. The tube is itself dilated in some cases, and in some the ovarian cyst discharges into the tube.

The origin of this condition is not always clear. In some cases it is evident that a salpingitis and perimetritis have caused adhesion of the tube to the ovary and occlusion of its orifice, with resulting dilatation. A chronic inflammation of the ovary coinciding may lead to cystic formation in it, by dilatation of the Graafian follicles, and coalescence may result. A colloid cystoma may coincide, but this is a rare accident. According to some authors a congenital adhesion of the tubes to the ovaries leads to the tubo-ovarian cyst in some cases.



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## B.—THE FŒTAL MEMBRANES AND PLACENTA.

The pathological changes in these structures have been very imperfectly investigated, and there are many of the lesions whose nature is not clearly understood.

1. **Affections of the decidua.**—It is generally believed that diseases of the decidua are common causes of premature discharge of the ovum, but little is known as to their nature. Atrophy and hypertrophy are stated to occur, the latter perhaps as the result of inflammation.

**Inflammation of the decidua** (*Deciduitis* or *Endometritis decidualis*) apparently leads to various local thickenings and irregularities of the decidua. It occurs when, before pregnancy, there has been a chronic inflammation of the mucous membrane of the uterus. The changes are observed in the membrane discharged along with the aborted foetus.

**Hæmorrhage. The Sanguineous or Fleishy mole.**—Hæmorrhage from the decidua is frequent in the course of pregnancy, and may be the cause of abortion.

In some cases the foetus is retained and the blood may accumulate in the chorion and the surface of the amnion. The blood coagulates, and, the fibrine condensing the structures, a fleshy mass is produced, in the midst of which the dead foetus may be no longer discoverable. This mass may remain long in the uterus, to be discharged afterwards as a fleshy mole. Sometimes it is retained so long as to become infiltrated with lime salts, a stony mole being ultimately discharged.

2. **Hydatid mole.**—This is an affection of the chorion, arising by a great newformation of the mucous tissue which normally exists in the villi.

The hydatid mole is formed of a mass consisting of the foetal membranes on the surface of which are myriads of oval or round cyst-like bodies which hang like grapes on stalks, and often hang on one another (see Fig. 343). Thus a bulky mass is formed consisting of a multitude of these berry-like cysts. When the altered membranes are opened, a



dead and blighted foetus is revealed, generally in an early stage of development.



Fig. 343.—Portion of a hydatid mole showing the berry-like masses. Natural size. (VIRCHOW.)

The villi of the chorion are branching offshoots consisting of an external layer of epidermis, and an internal basis of mucous tissue. It is by increase of the latter that the cyst-like bodies are formed. In an early stage of development the villi are present all over the chorion, while later on they are limited to a certain area forming the foetal part of the placenta. If the disease begins early the hydatids are present all over the chorion, whereas in later periods they occupy only the placental surface.

The altered chorionic villi adhere to the wall of the uterus, and sometimes are even embedded in its substance, so that when removed the internal surface may be like that of the heart, showing prominent trabeculæ. There may even be a destructive encroachment on the uterine wall.



The hydatid mole grows quickly, so that the enlargement of the uterus goes on more rapidly than in a normal pregnancy. The mole is detached at first in pieces, along with hæmorrhage, and even when the bulk of it is discharged, it may be so mixed with blood as to be with difficulty distinguished. The clear grape-like bodies afford the means of distinction.

**3. Diseases of the placenta.**—The placenta, being rapidly formed and being composed of a tissue intended for temporary purposes, is specially liable to various degenerations and to disorders of the circulation.

**Hæmorrhage** occurs in the placenta not infrequently, producing localized collections of blood, the condition being called Apoplexy of the placenta. The placental villi will be embedded in the coagulum and somewhat concealed.

**The White infarction** is a somewhat frequent lesion in the placenta. It is found in cases where the child has been still-born, and it implies grave disturbance of the circulation in the placenta. The infarction forms a solid white mass, usually wedge-shaped, and generally multiple. In it the villi are dead, and fibrine or blood lies in the spaces. The lesion has arisen by a process of **Coagulation-necrosis** like the pale embolic infarctions of the kidney and spleen.

The cause of this localized necrosis is not clear. There is probably an arterial obstruction, which Ackermann has ascribed to a periarteritis nodosa. According to Kustner serious nutritive disturbances during pregnancy are regularly followed by the formation of numerous white infarctions of the placenta.

**Inflammations of the placenta** are described as leading to fibrous thickenings. The inflammation originates in the decidua and extends to the placental tissue.

A **Periarteritis nodosa** has been mentioned above as having to do with the formation of the white infarctions. It appears indeed that these infarctions are often regarded as due to inflammations. The whole subject is in need of further elucidation.

**Syphilis** leads in the placenta to newformation of granulation tissue and gummata, causing the placenta to increase in bulk and weight. From the pressure of the new-formed tissue on the vessels, there is apt to be degeneration of the placental structures.

**Retrograde changes** are frequent in the placenta. Fatty degeneration and calcareous infiltration occur in the various structures. The placenta is sometimes speckled all over with small whitish spots, which are areas in which calcareous deposition has occurred. Along with this the microscope will reveal fatty degeneration.



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## C.—THE MAMMARY GLAND.

The mammary gland is very frequently the seat of tumours. It is also very liable to inflammations, especially during lactation.

### I.—MALFORMATIONS, INFLAMMATIONS, ETC.

1. **Malformations of the breast.**—One or both *mammæ* may be absent. It is more frequent, however, to have supernumerary breasts or nipples, sometimes three, four, or even five, instead of two. The supernumerary breasts are usually situated near the *axillæ*, or under the normal ones. But there are cases in which they have had a very abnormal situation, as in the inguinal region, on the thigh, or even on the back. They are usually small in size, but in some cases they have produced milk during lactation. Supernumerary breasts are more frequent in males than females, and they occur preponderatingly on the left side. The occurrence of additional *mammæ* is not altogether extraordinary, considering that in the lower animals the *mammæ* are usually more in number than in man.

2. **Inflammation of the mamma.**—This occurs not infrequently about the period of puberty or in connection with menstruation, but in the majority of cases it is related to lactation. In connection with lactation, and especially at its commencement, the structures in the mamma are the seat of very active processes; there is hyperæmia and an active secretion of milk. Under these circumstances inflammation is more readily induced than usual. It may happen that contraction of the arteries from accidental circumstances, such as exposure to cold, may induce inflammation. Even a general irritation of the vaso-motor centre, when the skin is exposed to cold, may, by reflex action on the highly excited vascular system of the mamma at the commencement of lactation, induce inflammation. Again, disease of the nipples, such as cracks and ulcers, often induce inflammation in the breasts. The irritant which has caused the lesion in the nipple may extend along the tubes and cause inflammation in the mamma.

The inflammation is usually an acute one, and is accompanied by exudation, the interstitial tissue being packed with leucocytes and the breast hardened. This hardening is often local, as the inflammation



is usually to some extent limited to certain parts of the breast. But very often the exudation of leucocytes goes on to actual suppuration and the formation of **Abscess**, sometimes with sloughing of the tissue. The abscess so formed may have extensive ramifications in the mamma, especially if the pus does not get vent externally. After evacuation the cavity fills with granulation cells, and finally closes, and a cicatrix is formed.

A **Chronic inflammation** resulting, after the usual manner of interstitial inflammations, in induration of the organ, has been described, but is not of frequent occurrence.

3. **Hypertrophy of the mamma.**—In some cases the mammæ undergo a progressive enlargement from the newformation of proper mammary tissue. In this way the gland may come to weigh as much as thirteen pounds. It is to be remembered that a simple enlargement of the fat around the mamma (*lipoma capsulare*) may imitate hypertrophy, which may also be simulated by diffuse tumours of the gland.

4. **Tuberculosis of the mamma.**—This, although probably rather frequent, has only recently been fully recognized although scrofulous conditions of the breast have been long known. There may be isolated caseous masses or they may be more confluent. The caseous matter undergoes softening and opens spontaneously on the surface or is punctured. The result is usually the establishment of a fistulous opening.

The tuberculosis of the mamma may be secondary to that of the lungs or other organs, but it may be primary. According to Verneuil the bacilli may find entrance by the milk ducts, and multiply in the glandular structures.

Tuberculosis of the mamma occurs not infrequently in cows which are the subjects of tuberculosis of other parts. This fact is of importance because the bacilli frequently find their way into the milk in such cases.

5. **Syphilis of the mamma.**—This is very uncommon, but cases of gumma in the breast have been recorded.

## II.—TUMOURS OF THE MAMMA.

The female breast is one of the most frequent seats of tumours, and they present considerable variety in form.

As the two constituents of the breast are the proper glandular structures and the connective tissue stroma, so tumours are derived from one or other of these, and belong to the connective tissue or epithelial forms. But the connective tissue tumours do not originate in a particular small piece of tissue, and grow so as to displace the remaining structures; on the contrary, they usually involve a considerable portion



of the gland and sometimes its whole extent. Hence they enclose more or less the glandular elements, and it may be impossible to say whether these are increased so as to form an integral part of the tumour. Moreover, these glandular elements are subject to great alterations, consisting of contortions and dilatations, which also give their character to the tumours, producing fissures and cysts in them. Again, the tumour tissue frequently grows into the cysts, producing the so-called intracystic growths, which sometimes assume a papillomatous character.

It will be inferred that the distinction between connective tissue and glandular tumours is frequently difficult, and that dilatation of the gland structures, implying, as it does, newformation of these, gives a partly glandular character to a large proportion of the tumours.

The atypical mammary tumours show a great preponderance in frequency, especially the cancers. According to statistics collected by Gross, there were in 973 tumours, 832 cancers (85·5 per cent.), 77 sarcomas, 61 fibromas, 3 adenomas, and 22 cysts.

Amongst the typical tumours, the lipoma, chondroma, and myoma are excessively rare, and require little notice.

**Lipoma** occurs doubtfully as a solid tumour in the midst of the gland. The **Lipoma capsulare** is also rare. In it the adipose tissue around the gland undergoes great increase, while the gland itself atrophies. Such tumours may assume enormous dimensions, weighing up to 100 pounds.

**Chondroma** rarely occurs as an independent tumour, but cartilage is sometimes found in sarcomas and cancers. **Bone** sometimes exists in fibromas, causing a partial conversion of them.

**Myoma** (leiomyoma) sometimes develops in connection with the nipple. Billroth observed a case in which striated muscle existed in an adeno-sarcoma.

**1. Adenoma and Fibroma. Adeno-fibroma.**—The boundaries between the adenoma and fibroma are not well-marked. They are both slow-growing, non-malignant tumours, generally of hard consistence, distinctly encapsuled, and not infrequently multiple. Many tumours which are regarded as glandular by some authors are described as fibromas by others.

The **Adenoma** is a purely glandular tumour. It occurs either in the substance of, or in the neighbourhood of the gland. It has a fleshy feel and a lobulated outline. On section it is white in colour and frequently shows cysts but without intracystic growth.

There is preserved in the Museum of the Western Infirmary a glandular tumour whose situation was behind the gland, which had to be cut through in order to remove it. The tumour was almost like a supernumerary mamma, and had to the naked eye and microscopically very much the structure of an inactive mamma (see Fig. 344). The tumour was removed from a young lady aged 21, and had been observed for two years, growing latterly more rapidly.



Under the microscope the adenoma presents elongated and enlarged acini and ducts (Fig. 344), and these commonly show dilatation, so that cysts are present in the majority of cases.

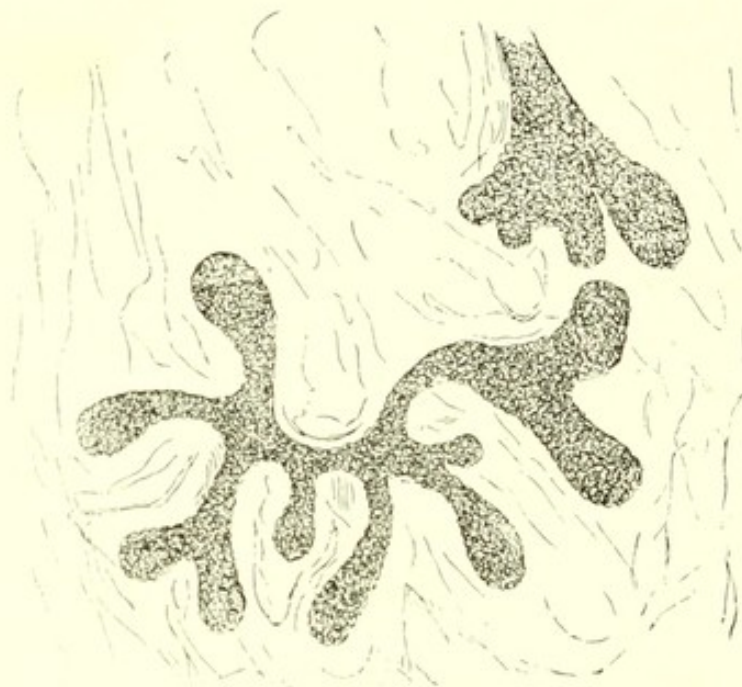


Fig. 344.—Section of an adenoma of mamma. Glandular structures of well-formed outline are shown in the midst of fibrous tissue.  $\times 90$ .

The **Fibroma** or **Adeno-fibroma** is a hard fibrous tumour, generally with a lobulated outline, and distinctly encapsuled. On section it shows a glistening, fibrous appearance, but this is rarely homogeneous. There are usually, even to the naked eye, indications of the presence of glandular structures, and these are often dilated into cysts. The whole tumour may be a congeries of cavities, which only partly contain fluid, but are largely filled up with foliaceous or dendritic structures composed of tumour tissue projecting into them.

These tumours are sometimes the seat of calcareous deposition and of osseous formation. There may be a partial myxomatous transformation leading in this way to cysts.

Under the microscope wavy fibrous tissue predominates. There are also glandular structures visible, which are either ill-developed or contorted and dilated as in Fig. 346.

2. **Myxoma**.—In this rare form of mammary tumour, the mucous tissue develops from the stroma of the gland, and it may involve one or more lobules, or the whole gland, which is then converted into a bulky tumour. Sometimes the mucous tissue grows into the milk ducts, forming the *Myxoma intracanalicular arborescens* of Virchow.

The tissue may be pure mucous tissue, but it is subject to various



modifications. It may be unduly cellular so as to approach to the sarcoma, or it may be mixed with fat or fibrous tissue. It also contains glandular tissue, which may be dilated into cysts.

This form is more malignant than the adeno-fibroma, being more liable to recur after removal.

**3. Sarcoma. Adeno-sarcoma. Cysto-sarcoma.**—The two last-mentioned names imply that the glandular structures take an important part in giving character to sarcomas of the mamma, although here it may be admitted that the connective tissue structures are those essentially engaged.

The line of demarcation between the sarcoma and the fibroma is not absolutely distinct, and there seems no doubt that the latter may develop into the former by an atypical process of growth supervening.

The spindle-celled sarcoma is the commonest form. The cells are usually small and there may be a considerable amount of fibrous intercellular substance. The tumour is usually hard and variously modified by the presence of gland tissue. The round-celled sarcoma is unusual, and generally forms a soft tumour. The cells may be small so as to be like lymphoid cells, or larger. The giant-celled sarcoma is very rare; there are smaller cells and gigantic ones. The pigmented sarcoma is also rare; the cells are usually round, but sometimes spindle-shaped.

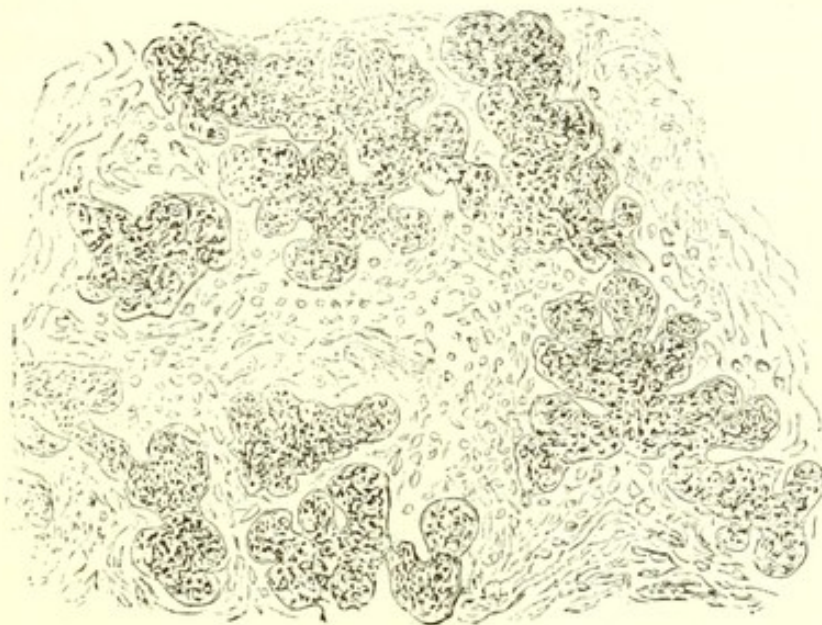


Fig. 345.—Section of an adeno-sarcoma of the mamma. Glandular structures are shown with spindle-celled tissue between.  $\times 90$ .

The sarcoma is generally solitary. It may be distinctly demarcated, occupying a small part of the breast, but it not infrequently extends so as to involve the whole mamma. It may grow slowly, but is apt, after a period of slow growth, suddenly to enlarge rapidly.



The **Adeno-sarcoma** is a tumour in which the glandular structures are specially abundant. It is usually a small, comparatively isolated tumour, somewhat resembling the adeno-fibroma, and, like this tumour, it is not infrequently multiple. Under the microscope it usually shows abundant glandular structures somewhat contorted, with a spindle-celled tissue between (Fig. 345).

The **Cysto-sarcoma** shows many gradations. There may be such simple dilatations as those indicated in Fig. 346, or there may be large cavities. Great complication is sometimes produced by intracystic growth of the sarcomatous tissue, as in the cystic-fibroma.

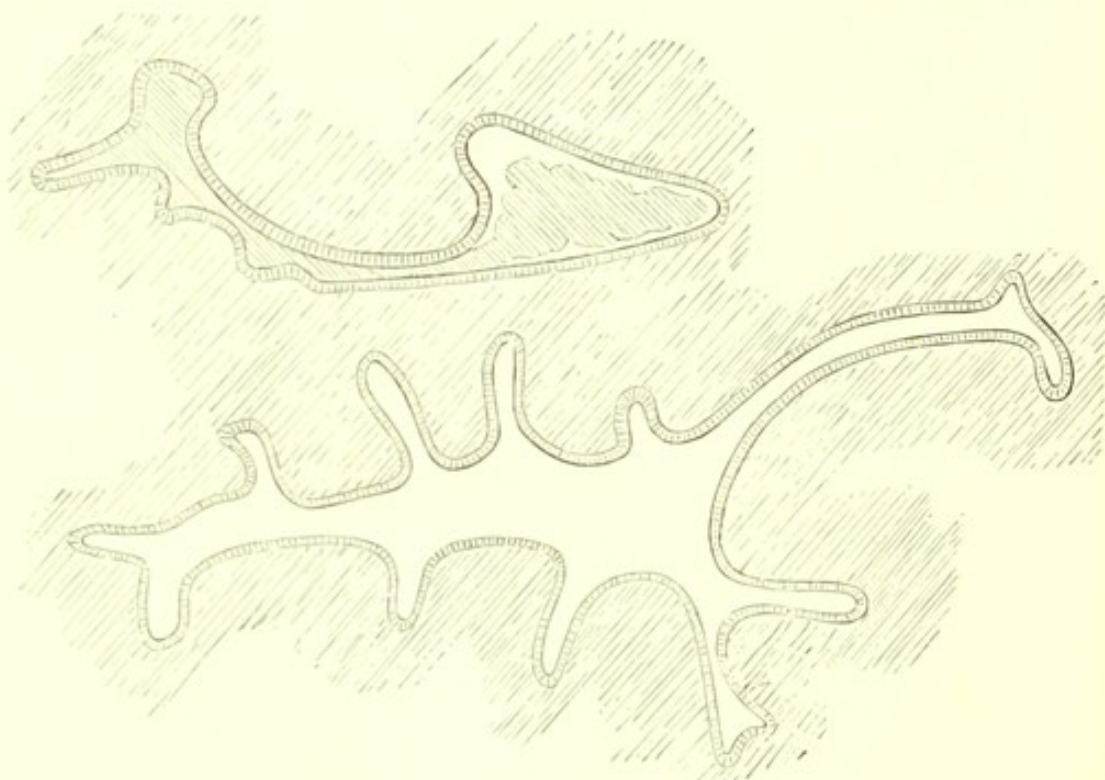


Fig. 346. —Semidiagrammatic view of dilated glandular spaces in a cysto-sarcoma of the mamma. The spaces are lined with cylindrical epithelium. The tendency to intra-cystic growth is shown, especially in the upper one. These were drawn under the camera lucida so that the outlines are correct.  $\times 62$ .

Besides these variations due to the mixture with glandular tissue, the sarcoma is subject to other modifications. It may contain **Cartilage**, as in the case from which Fig. 347 was taken. In this case the tumour contained well-formed fibrous tissue and cartilage, but also, apparently as a more recent development, spindle-celled tissue, and a tissue consisting of closely aggregated cartilage cells with little hyaline matrix. Bone also occurs occasionally.

**Mucous transformation** sometimes occurs, and cysts may arise in this way. There is also fatty degeneration, calcareous infiltration, and hæmorrhage, especially in the quickly growing and softer forms.

Sarcomas are **very malignant**. On removal they are even more apt



to return than cancers, and they also tend by metastasis to give rise to secondary tumours in internal organs, the metastasis occurring by the blood.



Fig. 347.—Section of cartilaginous part of a tumour of mamma which contained besides spindle-celled and fibrous tissue.  $\times 200$ .

**4. Cancer of the mamma.**—According to what has been stated in the general section of this work, we have, in cancer, an aberrant growth of epithelium as the foundation process. In the more ordinary cancers of the breast the process begins in the glandular acini. At the growing margin of the tumour the epithelium of the acini is seen to be proliferating, so that the acini are distended and enlarged. Along with the newformation of epithelium in the acini there is a formation of round cells in the connective tissue around. The epithelium of the acini next grows through the basement membrane, forming penetrating processes, and the round cells develop connective tissue which frequently causes great contraction of the tumour (*Scirrhus*). It is as if the epithelium, growing outwards, acted as an irritant, causing inflammatory newformation in the interstitial tissue of the gland. Where the case is very chronic the connective-tissue formation may be very pronounced, but when more acute the epithelial elements preponderate. Cancers which thus take origin in the glandular acini may be called **Parenchymatous cancers**.

But cancers may take origin rather in the ducts of the gland, forming a class called **Duct cancers** (*Thin*). These begin in the large ducts in or near the nipple. The growing epithelium distends the ducts, forming cavities which are sometimes lined with cylindrical epithelium, with more irregular epithelium inside. This process in the ducts is also associated with irritation of the surrounding connective tissue, so that there are round cells and new-formed connective tissue. The special characters of duct cancer are usually most manifest in the parts near the nipple, where it generally begins. But the cancer extends into the substance of the gland, where the appearances are more like those of the other form.



Duct cancer is sometimes associated with the condition known as **Paget's disease or Eczema of the nipple**. Eczema, as indicated in the section on Diseases of the Skin, is an inflammation of the skin, in which both the epidermis and the superficial layers of the cutis are engaged. In the case of duct cancer the cancerous affection of the ducts, which is the primary lesion, induces an inflammation of the skin of the nipple and parts around, characterized by the usual formation of round-cell tissue. The superficial or papillary region of the skin may then be replaced almost entirely by round cells with nests of epithelium in their midst, and the condition is somewhat like that of a granulating wound. The epithelium of the surface takes little part in the process, and it may be lost, so that the granulation tissue is exposed. As the cancerous process begins commonly in the nipple, this condition of eczema may be a very early one, and may precede, it may be for many months, the actual appearance of a tumour in the mamma.

It has been supposed that a primary eczema of the nipple may be the starting point of a cancer of the breast, but the observations of Thin show that the affection of the ducts is primary, although the occurrence of eczema may afford the earliest visible indication of the existence of the process. It is to be remembered also that a simple eczema may occur in the nipple, due to an ordinary irritant, and without any cancerous tendency.

In its atypical growth the glandular epithelium sends its processes into the **Lymphatic spaces** of the connective tissue. There is commonly a further extension to the lymphatics outside the gland. These are sometimes seen distended by cancerous growth, but even without this obvious involvement the infection has often spread to the axillary glands. These may be but slightly enlarged and yet the seat of distinct cancerous newformation.

**Forms of cancer of the mamma.**—The various forms are not absolutely separable one from the other. They all originate in the glandular structures, and they are distinguishable chiefly by variations in the proportion of cells and stroma, and also by the transformations of these.

(a) **Fibrous cancer. Scirrhus.**—This is the commonest form, constituting about 95·5 per cent. of the cases (Gross). The epithelial masses form elongated processes (Fig. 348) in the midst of an excessive stroma in which round cells are frequent. The epithelial cells are often atrophied so that in some parts of the tumour there is little beyond dense connective tissue. The cancer commonly forms rather a limited infiltration of a part of the gland than a proper tumour, and as the tissue contracts there is commonly an actual diminution of bulk with great induration. The gland is distorted and puckered towards the affected part.



Very often the disease is continuous with the nipple, and by the dragging of the tissue the nipple is drawn in, sometimes even forming an umbilicated depression. The tumour is very irregular in its extension in the gland, and it very often happens that in the midst of hard scirrhus tissue some **Adipose tissue** appears. In like manner pieces of the gland-tissue may crop up amidst the contracting tumour.



Fig. 348.—Section of cancer of mamma from a recent nodule. Epithelial cells in spaces formed by connective tissue; these are sometimes in single rows, and by multiplication form larger masses.  $\times 200$ . (CORNIL and RANVIER.)

The skin is frequently involved in the cancer, and it may be the seat of somewhat extensive growth. **Ulceration** often occurs, generally beginning about the site of the nipple and areola. The ulcer is crater-shaped with dense prominent walls. Sometimes after formation of the ulcer the growth of the cancer becomes more rapid.

When the mamma is cut into either the whole gland or a portion of it is seen to be occupied by a dense mass of a greyish colour on the cut surface. It is very hard to cut, and the cut surface is commonly concave, the elasticity of the dense connective tissue producing retraction. In the general grey basis there are various whitish or yellowish pieces which represent the remains of mammary or adipose tissue, or the cancerous epithelium in a state of fatty degeneration. If the cut surface



be scraped a **thickish juice** is obtained, which microscopic examination



Fig. 349.—Cells in the juice scraped from a scirrhus of mamma. They are of very irregular shape. Most of them contain several nuclei and some daughter cells.  $\times 200$ .

shows to be composed of epithelial cells and their debris. Many of the cells are large and well-formed though variously shaped, while some may contain secondary cells within them or double nuclei (see Fig. 349). There are also free nuclei which have escaped from cells as a result of the mechanical interference in the process of preparing the specimen. Many

of the cells present fatty degeneration; in fact, very often nearly all contain fatty granules, and there are some completely degenerated, showing nothing but an aggregation of fat drops like the compound granular corpuscle.

The secondary tumours in the lymphatic glands also show an excessive development of fibrous stroma, which, however, usually forms a more distinct meshwork than that in the mamma, while the epithelial cells form more definite groups.

The contraction and induration of the cancer are so great in some cases as to cause a great reduction in bulk of the mamma as a whole. This has induced some authors to distinguish a variety under the name of **Atrophying cancer**.

(b) **Soft or Acute cancer**.—This forms the opposite extreme to scirrhus, and there are all intervening grades. In the soft cancer we have a bulky tumour of soft consistence and rapid growth. There is a well-formed alveolar stroma, and the cells are somewhat loosely contained in it. The tumour involves neighbouring tissues, very readily infiltrating the skin, subjacent muscle, and even the osseous ribs and the pleura. This form is not uncommonly present simultaneously in both breasts. On section the tumour presents a grey brain-like appearance. The juice furnishes numerous cells, with prominent oval nuclei, also abundant free nuclei which have escaped from the delicate cells. As the tumour commonly extends to the skin, we may have ulcers with fungating prominence of the tissue. The lymphatic glands are early affected, and they also may ulcerate.

Many large tumours of the mamma of comparatively rapid growth do not correspond with the description either of the medullary cancer or of scirrhus. In their minute structure they are more like scirrhus, and perhaps may be designated **Acute scirrhus**. They have a remarkably firm fibrous stroma, but it forms distinct



meshes in which the epithelial masses are contained with considerable regularity. The cancer cells are also large and essentially like those in scirrhus. The tumour as a whole also is hard, this depending on the abundance and density of the stroma.

(c) **Colloid cancer.**—As compared with scirrhus, this is a rare form of tumour, constituting about 1·34 per cent. of the total cases. The colloid degeneration may affect the cells of an ordinary cancer, producing a partial metamorphosis. In true colloid cancer, however, the tissue as a whole presents, from the first, a tendency to colloid degeneration. The entire mamma is commonly affected, and it is greatly enlarged, while it presents a hard feeling like that of acute scirrhus. On section the tumour has a flickering gelatinous appearance. Under the microscope there is the usual pronounced stroma with colloid material in the meshes. In the midst of the colloid masses there are often seen groups of cells, the remains of the epithelium.

A very infrequent variety is sometimes described under the designation **Mucous cancer**. It forms a large gelatinous tumour which, on microscopic examination, shows epithelial masses embedded in a gelatinous flickering stroma.

Cysts are not of frequent occurrence in cancers, but in some cases it is reported that cystic formation is so characteristic as to induce observers to use the name **Cystic cancer**.

**5. Cysts of the mamma.**—It has been already pointed out that cysts frequently complicate other tumours, especially adeno-fibromas or sarcomas, but sometimes cancers. Cysts also occur independently, and they too, most frequently originate from the gland structures.

Most cysts of the mamma belong to the group of **Retention cysts**, arising in consequence of obstruction of the ducts. The cause of obstruction is sometimes an interstitial inflammation, and in that case the cysts will usually be small and multiple, just as they are in the kidneys in interstitial nephritis. In other cases the cause is obscure, and as the cysts may assume a large size the origin is perhaps embryonic.

(1) **Multiple cysts in old people** (*Involution cysts*) occur in connection with involution of the gland. The cysts are usually small, about the size of small shot or hemp-seed, but may be larger. When unopened they have a greenish or blackish appearance, and they contain a glairy fluid in which fat and epithelial cells are present.

On microscopic examination there are signs of irritation visible around the ducts in the form of round cells, and it is probably from contraction of new-formed connective tissue that the cysts form.

(2) **Simple cysts** may grow to a large size, forming tense thin-walled sacs. They are usually single, but if of small size they may be multiple. Their contents vary considerably, being usually serous or sero-sanguineous, but sometimes they are deep brown in colour and



contain fat and cholestearine. In some cases they contain milky fluid in which case they would be called gallactoceles.

(3) **Gallactoceles** or lacteal cysts arise usually when the gland is active. They are found chiefly in the neighbourhood of the nipple, arising by dilatation of the larger ducts. They contain milk, usually like ordinary milk, but sometimes altered so as to resemble cream or thickish oil, or curd, or butter. These cysts are usually solitary and are very rare.

(4) **Connective tissue cysts** have been recently described and are supposed to be of lymphatic origin. They have thick walls and the connective tissue may be indurated around them. Hence they resemble scirrhus in their clinical aspects. The internal surface is lined with a flat endothelium. The cysts are single or multiple.

**Parasites in the mamma.**—The only parasite of any consequence is the *Echinococcus* which forms hydatid cysts. These may be with difficulty discriminated from simple cysts, and as there is sometimes considerable induration around them, they may also be mistaken for cancers.

The *Cysticercus cellulosæ* has also been observed in the mamma (Guermontprez).

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## SUBSECTION II.—DISEASES OF THE MALE GENERATIVE ORGANS.

## A.—THE TESTICLE AND TUNICA VAGINALIS.

1. **Malformations and Misplacements of the testicle.**—The testicle may be absent on one or both sides, while the vesiculæ seminales and vasa deferentia are perfect. Or the vas deferens may be deficient while the testis is well developed.

**Imperfect descent** of the testis, or **Cryptorchismus**, is a very frequent and important condition. The descent of the testis occurs normally before birth, but in a considerable number of cases the organ has not yet appeared externally at birth. In the majority of such cases it comes down within the first week, but it is sometimes delayed for months or a year, or even till puberty. When thus delayed the descent is very often imperfect, so that the gland remains in the inguinal canal or at the ring. On the other hand, the testis may be retained throughout life in the abdomen. The testicle retained in the abdomen or on the way down is usually **imperfectly developed**, and although apparently of normal size, it does not generally contain spermatozoa in its tubules. It is also more prone to inflammation and to the formation of tumours than the normally placed testis, especially when retained in the inguinal canal where it is exposed to external violence. When it is late of descending the accompanying pouch of peritoneum, which forms the tunica vaginalis, is apt to remain open and so to induce a **Congenital hernia**.

Besides these congenital misplacements, we may have the testis descending into the crural canal, or into the perineum.

2. **Inflammation of the testicle.**—This condition presents itself under a considerable variety of forms, certain of which are difficult altogether to separate from syphilis and tuberculosis. It is to be remembered that the testicle is bounded externally by a dense fibrous capsule, the tunica albuginea, and that the products of inflammation are apt to be closely confined within this.

**Acute orchitis** is mostly secondary, although occasionally resulting from a blow or other injury. It occurs as a secondary result in gonorrhœa, the specific irritant being apparently propagated along the vas deferens; also in cases of irritation of the urethra by calculi, the passage of instruments, etc., the inflammation in such cases being called sympathetic, and probably brought about by reflex irritation of the vaso-motor centres. In mumps there is also, in a certain proportion of



cases, orchitis as a secondary result, the specific irritant attacking the testicle as well as the parotid gland.

The inflammation very often begins in the epididymis, and may remain confined to that region or pass on to the testis. There is hyperæmia and serous exudation producing great swelling, so that the tunica albuginea is tightly stretched, and the organ feels hard and tense. At the same time there is often serous fluid in the tunica vaginalis. There may be resolution without suppuration, the testis returning to the normal condition, or the disease may become chronic, or suppuration may occur. In the latter case an abscess or abscesses form in the testicle, and these may burst externally. In that case the tense albuginea often forces the tissue of the testicle through the opening so that it projects in a fungating form. This, which forms the **Benign fungus**, and is not to be mistaken for cancer, has a red colour and looks like protruding granulations. But it really contains the seminal tubules, and after the subsidence of the inflammation these may be replaced and resume their function. On the other hand, the abscess may dry-in and form a caseous mass, which becomes surrounded by a fibrous capsule. This mass may ultimately become calcareous.

**Chronic orchitis** may arise out of an acute attack, remaining after the formation of abscess, or else supervening without that. There is also a spontaneous chronic orchitis which in most cases is either syphilitic or tubercular. The ordinary chronic orchitis is characterized by newformation of connective tissue in the stroma of the organ, and consequent induration of it. The tubules atrophy and the whole organ is reduced in bulk and hardened. Generally on section it can be seen that the albuginea and the septa, which proceed towards the mediastinum, are greatly thickened. At the same time there is frequently adhesion of the tunica vaginalis which may obliterate the sac.

3. **Syphilitic disease of the testicle.**—This sometimes occurs as a general induration, having the usual characters of chronic orchitis. The albuginea is thickened and the trabeculæ are enlarged as they converge towards the mediastinum. These thickened structures may be largely composed of granulation tissue, and we may have in addition definite gummata. The gummata may undergo caseous necrosis, and by coalescence of neighbouring tumours we may have considerable masses of caseous material. In the midst of this caseous material the outlines of the seminal tubules can sometimes be made out, indicating that the necrosis has overtaken the proper tissue as well as the new-formed.

4. **Tuberculosis of the testicle.**—Tuberculosis is characterized by the formation of caseous masses, sometimes of considerable size, and these



are surrounded by a grey inflammatory tissue in which tubercles are to be found. The tuberculosis usually extends by the vas deferens to the prostate gland and vesiculæ seminales, and it may pass on to the bladder and kidney (see p. 839). There is also, not infrequently, tuberculosis elsewhere, in the lymphatic glands or lungs.

As a general rule the tuberculosis begins in the epididymis, mostly in its tubules. The result is great thickening and the formation of a caseous mass, enclosed very often in a firm fibrous capsule. Thus a firm elongated tumour may form behind and partly surround the testicle (see Fig. 350). In the testicle also caseous masses form, at first isolated, but afterwards running together into considerable conglomerates. The caseous masses are very dense, but after a time they generally break down, and having burst externally, tedious fistulæ are the consequence.

When the testicle is examined in section the caseous structure is seen to be surrounded by a transparent grey tissue, in which, as already mentioned, tubercles are present. This inflammatory and tubercular structure by its pressure destroys the tubules for the most part, and bits of necrosed tubules are sometimes expelled along with the softened caseous material.

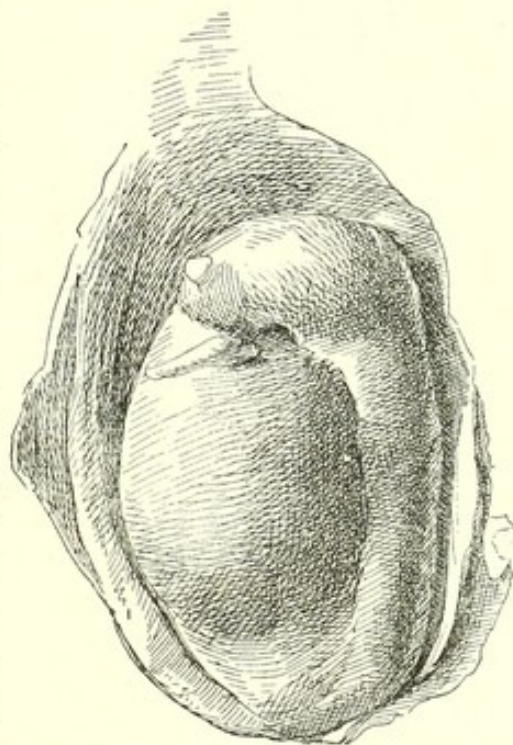


Fig. 350.—Tuberculosis of testicle. The epididymis is greatly enlarged and partly surrounds the testicle.

**5. Tumours of the testicle.**—These are mostly somewhat malignant in their characters, and there is a peculiar tendency to a mixed structure.

**Cystic tumours.**—The cystic tumours of the testicle show considerable analogies with those of the mamma and are capable of a somewhat similar division. The cysts in all cases originate in gland tissue which is believed by some to be the regular seminal tubules, but according to Eve, is rather the rudiments of the Wolffian body which are retained at the hilum of the testis, as similar remains exist at the hilum of the ovary (see p. 881). The tumours originate chiefly at the hilum and the tissue of the testis may be spread over them.

The tumours may be of considerable size, and present cysts of various shapes and sizes. The cysts are usually lined with cylindrical epithelium which is sometimes ciliated. The epithelium sometimes has the characteristic goblet form, such as that in colloid ovarian cystoma.



The stroma varies greatly in structure. It may be fibrous, in which case the tumour may be called an **Adeno-fibroma** or a **Cystic fibroma**, or it may be sarcomatous, in which case the term **Adeno-sarcoma** or **Cystic sarcoma** is used. In some cases it has the structure of mucous tissue, hence **Cystic myxoma**. It is not uncommon to find pieces of cartilage in the stroma, and pieces of striated and of smooth muscle have been found.

**Dermoid cysts** are also met with having a complex structure like those of the ovary and containing hairs and sebaceous matter.

The spermatocele is also properly a cyst of the testicle, but it is considered below along with hydrocele.

**Sarcomas** occur frequently mixed with tissue of other tumours, as the myxoma, enchondroma, etc. Even when the tumour is a pure sarcoma there may be combined the structures of various forms of this tumour, spindle cells, round cells, and mucous tissue. There may, however, be a simple round-celled sarcoma. The sarcomas generally form large tumours of soft consistence and rapid growth, and usually originate in the substance of the testicle. They occur in comparatively young persons, and are prone to metastasis, the secondary tumours occurring mainly in the lungs.

**Cancer of the testicle** connects itself with the cystic and adenoid sarcomas. It appears that not infrequently the glandular tissue in these tumours has more of the irregular atypic cancerous character than of the strictly glandular, and in particular the epithelial masses do not possess a membrana propria. These characters are also expressed in the fact that sometimes these tumours extend to the neighbouring lymphatic glands, while others penetrate more readily into the veins.

Proper cancer of the testis is mostly a large quickly growing soft tumour—**Medullary cancer**. It involves the whole organ as well as the epididymis, converting them together into a massive tumour. It may also extend along the vas deferens, and on to the lymphatic glands, inguinal, lumbar, and pre-vertebral. From these glands it may extend to the radicles of the portal vein and so produce tumours in the liver, or more frequently to the radicles of the vena cava, producing tumours in the lungs. In the midst of the tumour we may have cysts formed by mucous or fatty degeneration.

The cancerous tissue is derived from the epithelium of the seminal tubules, and the stroma from the interstitial tissue. It is necessary, however, to mention that tumours may arise in connection with the blood-vessels which have a resemblance in their structure to cancers, but which are properly plexiform angio-sarcomas.

**Enchondroma** is not infrequent in the testicle. It is usually mixed



with other forms, as mentioned above, but it may occur pure. As the tissue here is heterologous it often shows malignancy. Thus, it has been found growing into the lymphatics and blood-vessels around, and it forms secondary tumours in the lungs. For the rest, the occasional occurrence of fibromas, lipomas, myomas of the striated variety, and even osteomas, has been observed.

6. **Hydrocele.**—By this name is meant the accumulation of fluid in the tunica vaginalis. The fluid is serous and the cause of its accumulation is inflammation. It is not properly a dropsy of the tunica; a general dropsy, affecting the skin of the scrotum very greatly, causes little fluid to accumulate in the tunica vaginalis. The inflammation may be acute, in which case there is fibrine deposited on the surface as well as serous fluid in the sac. More commonly it is chronic, and there is a gradual accumulation of serum.

The fluid distends the sac, and so a bulky tumour is formed which is pear-shaped with its blunt end downwards. The fluid is usually a clear serum, but sometimes it is slightly opalescent. In the true hydrocele this is not from the presence of spermatozoa, but from the existence of fine fat drops, resulting from degeneration of the leucocytes floating in the fluid. There may even be cholestearine crystals formed in this way.

Not infrequently hæmorrhage occurs, most commonly as the result of a blow or other injury, and the hydrocele becomes a **Hæmatocele**. The blood mostly coagulates, and the coagulum through time undergoes various changes, softening into a brown pultaceous material or into a brown turbid fluid, in which are enormous numbers of cholestearine crystals. The blood seems to act as an irritant to the tunica vaginalis, causing often a very great thickening of it. The interior also is rough and sometimes presents considerable projections. The thickened cyst may contract somewhat so that the tumour is reduced in size. This thickening of the wall, even when the contents are fluid, may cause the hæmatocele to be mistaken for a solid tumour, and castration has often been performed under this belief.

A hydrocele or hæmatocele may be cured by the fluid being absorbed, the result usually being adhesion of the opposed surfaces of the tunica vaginalis and obliteration of the sac.

In the condition named **Congenital hydrocele** the tunica vaginalis retains its communication with the peritoneal cavity. The fluid may come from the peritoneal cavity, or may originate as in an ordinary hydrocele. It can be pressed into the peritoneum through the neck. It will be understood that a congenital hernia may occur along with or alternate with this form of hydrocele.



7. **Spermatocele. Encysted hydrocele.**—In this affection there is a cyst having in many cases much the external appearance of hydrocele, but containing a fluid in which spermatozoa are abundantly present. In other respects also the condition differs from that in hydrocele, for we have here not merely an accumulation in an existing sac, but a proper new-formed cyst. Hence the name Encysted Hydrocele is often used as equivalent to spermatocele.

The cyst arises in connection with the epididymis for the most part, and probably takes origin in one or more aberrant tubules which have formed blind diverticula from the seminal tubules. It usually arises near the upper end of the epididymis, but it may be at the lower end or from the rete testis. The cysts grow often to a large size, and they sometimes push themselves into the sac of the tunica vaginalis, inverting one layer of the wall against the other. As a rule, the tunica vaginalis is found below and in front of the cyst, this position being connected with the origin of the cyst in the neighbourhood of the epididymis.

The fluid from these cysts has a peculiar opalescent appearance, which is due to the presence of multitudes of lively spermatozoa. The existence of these shows that the cyst has retained its connection with the seminal tubules. The cyst is usually lined with a ciliated epithelium, but in large ones the pressure of the fluid may cause these cells to assume the paved form.

It will be observed that these spermatic cysts are comparable in their origin to the parovarian cysts.

**Other forms of hydrocele** have been described, but they require only a passing notice. There may be an encysted hydrocele without spermatozoa in the fluid. Then there is encysted hydrocele of the cord, sometimes arising by a portion of the communication between the tunica vaginalis and the peritoneum remaining unobliterated and becoming the seat of an accumulation of fluid. There is also diffused hydrocele of the cord, in which there is an œdematous condition of the connective tissue around the spermatic cord. There may even be a hydrocele from a hernial sac, which has got emptied of its contents and shut off from the peritoneum by adhesion of the neck.

## B.—THE PENIS, SCROTUM, AND PROSTATE.

In the **Penis, ulcers or chancres** are the commonest forms of disease. The simple chancre is an ulcer which has its usual seat on the glans or frænum. Sometimes we meet with a deeply penetrating or phagedænic ulcer. The hard chancre is the primary syphilitic sore, and as such we have already considered it.



Another syphilitic manifestation is the **Condyloma**, which forms a warty outgrowth sometimes of considerable size. There may be large groups of papillæ forming a cauliflower-like tumour.

**Cancer** of the penis occurs in the form of epithelioma. It begins usually in the glans, and the tumour is often covered with prominent papillæ, which give it a highly characteristic warty appearance, like the cauliflower excrescence. It may remain long without ulceration, but usually sooner or later breaks down, and there are sometimes deep ulcerating fissures or fistulæ between the groups of papillæ.

In the **Scrotum**, **Cancer** is somewhat common, and it is so often chimney-sweepers who are affected that the disease is often called chimney-sweepers' cancer. In Glasgow it has been found that workers in paraffin refineries are also liable to this disease. It generally forms a flat tumour from whose surface prominent papillæ protrude. Through time ulceration occurs and the testicle may be exposed.

**Elephantiasis** of the scrotum is referred to at p. 212, and illustrated in Fig. 59.

**Lymph scrotum** has already been referred to in connection with the filaria sanguinis. In it there is a varicose condition of the lymphatic vessels of the scrotum with the formation of vesicles in the skin. These frequently burst and discharge a fluid in which the embryo filaria is usually to be found. In some such cases the filaria is also present in the blood, but sometimes not. The lymphatic glands of the groin are indurated, and the dilatation of the lymphatics has been ascribed to the obstruction of these vessels in the glands by the embryo filariæ (see p. 344).

**The Prostate**, as we have already seen, is subject to **Hypertrophy**. The common enlargement of old age is due to increase of the muscular substance. This enlargement exists in about 30 per cent. of men above 60 years of age. It develops slowly without any apparent cause, and in this respect has the characters of a tumour. Sometimes the whole prostate enlarges, and it may reach the size of the fist. But sometimes there is a more partial hypertrophy, forming the so-called third lobe of the prostate, which projects inwards at the neck of the bladder, and is sometimes so large as to act like a valve to the orifice of the urethra. This third lobe is thus entirely new-formed, there being normally only the two lobes. The effect of enlargement of the prostate on the urethra is to be noted. If there is a general enlargement the urethra is necessarily elongated in its prostatic portion, and whereas normally this portion measures  $1\frac{1}{2}$  inches in length it may come to be 4 inches. At the same time the tube may be narrowed and even distorted. If, for instance, the central part of the prostate is specially hypertrophied,



then the urethra, being pushed upwards, has on section a crescentic shape with the convexity upwards; or if one side is larger than the other, there will be a convexity towards the opposite side.

Besides this muscular hypertrophy the much rarer hypertrophy of the glandular structure is to be mentioned. The glandular structure does not commonly increase with the muscular, but sometimes enlarges by itself, so that we have an adenoma of the prostate.

**Cancer** of the prostate is not of frequent occurrence. The gland enlarges and the disease is apt to extend to neighbouring structures.

**Tubercle** occurs not infrequently in the prostate and vesiculæ seminales in conjunction with similar disease in other parts of the urino-genital passages. There is caseous necrosis with ulceration as usual, and this may cause even perforation into the rectum or bladder.

**Concretions** are of frequent occurrence in the prostate in old persons. They are formed in the gland-ducts and are of various sizes, from very minute to the size of a grain of corn. When small they are colourless, but as they enlarge they frequently become blackish or reddish brown in colour. They are round or oval in form and frequently present concentric stratification (Fig. 39, p. 168). They have usually a central cavity. Very commonly these bodies present the character of amyloid bodies, giving a bluish or mahogany red colour with iodine. Sometimes they contain lime salts in their substance. They may pass into the urethra and escape with the urine. While in the prostate they do not appear to produce much disturbance.

**Literature.**—*Testis*—CURLING, Dis. of testis, 4th ed., 1878; KOCHER, Krankh. d. Hodens, 1874. *Tumours*—EVE (Cystic testicle, with literature), Path. trans., xxxviii., 1887; NEUMANN (Myoma striocellulare), Virch. Arch., ciii., 1886; PAGET (Enchondroma and other tumours), Surg. Path., 1870. *Tuberculosis*—RECLUS, Du tubercule du testicule, 1876; WALDSTEIN, Virch. Arch., lxxxv., 1881. *Syphilis*—RECLUS, Syph. du testicule, 1882, and Gaz. Hebd., 1883.



## SECTION X.

## DISEASES OF THE SKIN AND ITS APPENDAGES.

*Introduction, as to normal structure.* I. **HYPERÆMIA, HÆMORRHAGE, and EDEMA.** II. **RETROGRADE CHANGES.** 1. *Atrophies of skin, hair, and pigment*; 2. *Necrosis, including ulcers.* III. **INFLAMMATIONS.** *Causation.* 1. *Inflammatory skin eruptions, the individual lesions, and different forms*; (a) *Urticaria*, (b) *Erythema and Roseola*, (c) *Eczema*, (d) *Psoriasis*, (e) *Pityriasis rubra*, (f) *Lichen ruber*, (g) *Prurigo*, (h) *Pemphigus*, (i) *Acne*. 2. *Symptomatic Inflammations, in acute fevers, chiefly small-pox.* 3. *Inflammations from heat, cold, and injury.* (a) *Burns and scalds, in three degrees*, (b) *Frost-bite and chilblain*, (c) *Wounds and excoriations.* 4. *Infective inflammations.* (a) *Boil and carbuncle*, (b) *Cadaveric infection*, (c) *Phlegmonous inflammations*, (d) *Malignant œdema*, (e) *Anthrax*, (f) *Hospital gangrene.* IV. **SPECIFIC NEW-FORMATIONS.** 1. *Syphilis, in primary, secondary, and tertiary forms*; 2. *Tuberculosis, as Lupus and Scrofuloderma*; 3. *Leprosy*; 4. *Elephantiasis*; 5. *Frambæsia.* V. **TROPHONEUROSES.** 1. *Herpes*; 2. *Glossy skin*; 3. *Scleroderma and Morphaea.* VI. **HYPERTROPHIES AND TUMOURS, including Ichthyosis, Corn, Wart, Soft wart or mole, Keloid, Molluscum contagiosum, Fibroma molluscum, Xanthoma, Sarcoma, Cancer.** VII. **PARASITIC AFFECTIONS, chiefly from fungi,** 1. *Favus*, 2. *Ringworm*, 3. *Pityriasis versicolor*; *Animal parasites.*

## INTRODUCTION.

THE diseases of the skin are exceedingly manifold, and the names applied to them somewhat complicated. In this section an endeavour is made to summarize the pathological conditions and group together the various diseases according to the nature of the lesion.

**Normal structure.**—The **Corium** or **True skin** is a very vascular dense membrane composed of interlacing fibres of connective tissue, with numerous elastic fibres. It is richly supplied with nerves, and possesses bundles of smooth muscle. It has also a rich system of lymphatic vessels. In the corium we may distinguish a superficial or **Papillary layer** and a deeper one. Many of the diseases affect the papillary layer especially.

The **Epidermis** on the surface of the corium is in several layers (Fig. 351). Most superficial is the horny layer (*a*) composed of flat



cells, which are little more than scales, and have lost their nuclei. Next comes the stratum lucidum (*b*) composed of flat transparent cells. Most deeply situated is the stratum mucosum or rete Malpighii, composed of cells which, in the deepest layer, are cylindrical, but towards the surface become polygonal and flattened. In the deeper layers the

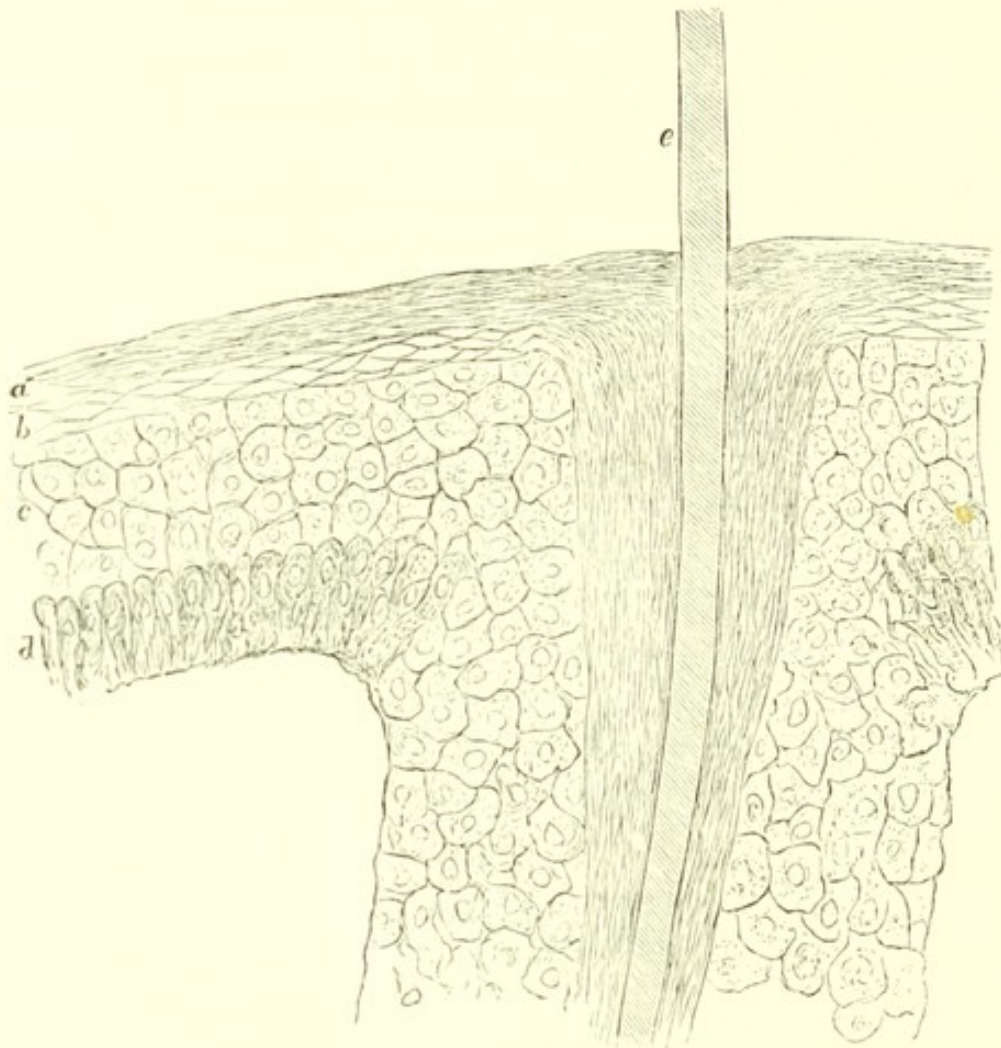


Fig. 351.—Section of epidermic layer of skin with a hair and its follicle. *a*, horny layer; *b*, stratum lucidum; *c*, Malpighian layer with (*d*) the deepest layer of cylindrical cells; *e*, hair, whose sheath presents two layers, one continuous with the horny and the other with the Malpighian layer.  $\times 350$ .

cells are serrated at the margins, so as to give the appearance of prickles by which the cells fit into each other. The Malpighian layer extends between the papillæ of the corium, forming interpapillary processes.

The **Sebaceous glands** may be regarded as modified prolongations of the Malpighian layer. They are mostly connected with hair follicles into which they open, but sometimes large glands are connected with small hairs and small glands with large hairs, while there are glands not connected with hairs at all. The **Sudoriparous glands** are usually situated beneath the skin, their ducts passing through corium and epidermis, having a spiral course in the latter. The **Hairs** are composed



of horny epidermis, and are placed in follicles. In the latter, two layers of epidermis can be distinguished (Fig. 351), the outer root-sheath corresponding with the Malpighian layer, and the inner root-sheath corresponding with the horny layer. At the bottom of the follicle is a papilla continuous with the corium, and on this is set the bulb of the hair. **The nails** are composed of compressed horny epidermis. Beneath the nail are still two layers of epidermis, a horny layer of loose cells, and a Malpighian layer covering well-formed papillæ.

The exposed position of the skin renders it very liable to the influence of agents acting from without. It is also liable to be influenced by irritants circulating in the blood, in which case the skin affection will probably be an insignificant part of a general condition. The skin again is liable to be affected by states of the nervous system. From these remarks it will be inferred that the inflammations of the skin are its most important morbid conditions, and these will call most largely for description.

**Literature.**—*General works*—ERASMUS WILSON, Dis. of skin, 1867; NEUMANN, (Pathological anatomy is very good) Lehrb. d. Hautkrank., 1880; HEBRA and KAPOSI, Dis. of skin (Syd. Soc.), 1866-1880; DUHRING, (literature very complete) Dis. of skin, 1882; McCALL ANDERSON, Dis. of skin, 1888.

#### I.—HYPERÆMIA, HÆMORRHAGE, ŒDEMA OF THE SKIN.

The skin is very liable to variations of its blood supply. An active **Hyperæmia** hardly occurs as a pathological condition except as part of an inflammation. Passive hyperæmia, on the other hand, is exceedingly common as a result of general venous engorgement, especially in disease of the heart. From the blue colour assumed by the skin in passive hyperæmia the condition is designated **Cyanosis**.

**Hæmorrhages** are frequent and of various kinds. As to their causes little need be added to what is stated at page 83. The skin is liable to hæmorrhage by traumatic rupture of its vessels, but still more, perhaps, by alterations in the state of the blood affecting its vessels. In scurvy, in purpura, in small-pox, in typhus, etc., the skin is the seat of hæmorrhage much more frequently than any other structure.

The blood escaping from the vessels collects in the serous spaces of the corium for the most part, but may pass to the subcutaneous tissue, where the fat is sometimes considerably infiltrated. When the blood has escaped from a small vessel and infiltrates a limited area so that a bluish spot is seen on viewing the surface, the term **Petechia** is given. As these small hæmorrhages depend on the state of the blood, the petechiæ are nearly always multiple. When the blood infiltrates a larger area then the term **Ecchymosis** is used. Sometimes the blood



collects between the corium and epidermis, and then a hæmorrhagic **Vesicle** is the result; but this can only happen if the deeper layers of the epidermis have been killed so as to allow of their separation from the corium. When there are numerous small hæmorrhages forming a large number of petechiæ, then it is customary to use the term **Purpura**, or to speak of a purpuric condition.

The blood effused in the skin undergoes changes like those already described at page 83. The affected area is first dark blue, and the colour does not disappear on pressure. After a time, as the blood-colouring matter is dissolved, the colour gets fainter and changes in hue, while the discoloration becomes more extended. If blood has collected between the layers of epidermis, it comes to the surface and is disposed of as the epidermis is shed.

**Œdema** of the skin, apart from inflammation, is chiefly related to disease of the heart and kidneys. Its general pathology has been considered at pages 88 and 89. The exuded fluid collects in the lymph spaces and is carried off by the lymphatics. **Myxœdema** has also been described at page 692.

## II.—RETROGRADE CHANGES IN THE SKIN.

1. **Atrophies.**—Simple atrophy is not uncommon in the skin. The most frequent example of it is afforded by **Senile atrophy**, resulting in the wrinkled skin of old people. In this case the connective tissue of the cutis loses in bulk, especially the papillary layer. The epidermis is also thinner and dry, and there is often desquamation in the form of dry scales or larger membranous pieces. The hairs are atrophied, as their papillæ have taken part in the general atrophy of the papillæ of the skin. The hair-follicles are shortened, and the sebaceous glands on this account may be brought close to the surface, almost opening directly on it. The secretion may accumulate in the sebaceous glands so as to form small or large **Cysts**.

A general atrophy also occurs in emaciated persons, and it may closely resemble the condition in senile atrophy.

There are further conditions variously named **Parchment skin**, **Xeroderma**, **Hide-bound skin**, which are of rare occurrence, and are not very well understood in their exact relations. The skin is thinned and hardened, especially the epidermis, so that the body seems covered with stiff parchment, which often looks as if too small for the surface to be covered, and the movements of the limbs are impeded. The skin is sometimes mottled with pigmented spots, sometimes white. The disease generally begins in early life and extends gradually over the skin.

Of the **Local atrophies** the most familiar is that which occurs after



**Pregnancy.** White lines are found in the abdomens of persons who have been pregnant, and similar lines occur in persons whose abdomens have been distended by tumours, by ascites, or even by accumulation of fat. The white lines have a cicatricial appearance, and they seem to owe their origin to the connective-tissue fibres of the cutis being dissociated by the stretching. Somewhat similar white lines or striæ sometimes occur idiopathically, especially over the buttocks, trochanters, pelvis and thighs.

The **Pigmentations** of the skin have been mostly referred to in previous pages. They are chiefly those in Addison's disease, in jaundice, in tattooing, and in argyria (see pp. 163, 164).

The **Hairs** are liable to atrophy, and two forms may be distinguished according as either the hair itself or its pigment diminishes. **Alopecia or Baldness** is atrophy of the hair itself. All through life a continuous falling out of the hair is occurring, and is due to an atrophy of the bulb. But the papilla remains, and a healthy new hair is produced on the site of the old one. In some people as life advances the new hairs are not reproduced of normal size, and they become gradually finer and finer, till there are only the finest woolly hairs on the bald part, or even none at all.

Besides being produced in this way, baldness may be **secondary** to syphilis, to inflammations and to certain parasitic diseases; in these cases it depends on interference with the nutrition of the sheath and papilla.

Special attention has been paid to **Alopecia areata**, in which baldness occurs in circular patches. These patches are not entirely bald, but are covered with fine woolly hairs, and the papillæ are not destroyed. The nature of this disease is obscure, some regarding it as parasitic, and others as a trophoneurosis. Its habit and mode of extension render the parasitic view very probable. It is very likely that, as Thin contends, the parasite here is a microbe (a micrococcus), which is much more difficult to detect than the other parasites of the skin, which are fungi.

**Canities or Greyness of the hair** is also for the most part a natural atrophy of advanced life. But it also sometimes comes on prematurely. It depends for the most part on a deficiency of pigment in the individual hairs at their original formation, so that the pigment granules between the cells in the cortical layers of the hair are diminished. But there may be a temporary blanching of the hair from air getting infiltrated among the cells of the cortex. Cases of sudden permanent blanching have received no satisfactory explanation.

**Albinism** is a congenital absence of pigment which affects the iris and choroid of the eye as well as the hair and skin. It is not infrequent in



negroes, the peculiar result being a white negro. The albinism is sometimes partial, so that, in the case of the negro, a piebald appearance is produced.

The **Nails** are liable to atrophy, which may be congenital or acquired. In the course of acute illnesses, such as fevers or maniacal attacks, the formation of the nails is often partially suspended, and the illness is marked for a time by a transverse depression, which, with the growth of the nail, proceeds from the root outwards, and disappears in due course. The nails also atrophy sometimes in consequence of various parasitic or inflammatory skin diseases.

2. **Necrosis**.—Death of portions of the skin occurs under various circumstances. Perhaps the commonest case is that of **Bedsore** (*Decubitus*), in which a piece of skin which has been exposed to pressure whilst lying, dies and is discharged as a slough. This occurs chiefly in weak and debilitated persons. It is met with frequently in insane persons and paralytics, and it is matter of controversy how far the affection of trophic nerves may have to do with the necrosis (see p. 146).

In all forms of **Cutaneous ulcers** there is necrosis, which may be in the molecular or in the more massive form. The margin of the ulcer is inflamed and infiltrated with inflammatory products. Enlargement takes place by a gradual molecular necrosis of the inflamed structures, and generally implies the continuous action of an irritant. The irritant is mostly an infective virus, and the majority of the ulcers will be again referred to under subsequent headings.

In the case of such **Infective ulcers** there is usually a granulating surface which is gradually destroyed by the action of the virus. When the virus has been destroyed, then the ulcer proceeds to healing like a granulating wound (see p. 135, etc.).

The **Varicose ulcer**, arising in connection with varicose veins, owes its origin to a prolonged venous hyperæmia, leading to œdema and infiltration of the skin. The nutrition of the skin is seriously compromised, and a trivial external injury may cause the epidermis to give way. The exposed and inflamed skin forms granulations, but these also, from the persistent venous engorgement, are readily destroyed by slight injuries, and the ulcer advances, while the inflammatory infiltration of the skin outside extends. The granulations will only proceed to healing in the usual way when the venous hyperæmia is removed.

The **Perforating ulcer of the foot** is a peculiar form which is believed by some to be due to a trophic lesion of the nerves. There have been found in some cases inflammatory lesions of the nerves, and the disease is sometimes associated with nervous affections, especially locomotor ataxia. It is questionable how far mere anæsthesia is sufficient to



account for the formation of the ulcer, as it seems to do in the case of anæsthetic leprosy. The ulcer is characterized by the formation of an aperture on the surface of the foot, which leads to penetrating sinuses in its substance, often attacking the bones.

**Cancerous ulcers** are incapable of healing.

### III.—INFLAMMATIONS OF THE SKIN.

These include a very great variety of diseases, and the nomenclature is exceedingly complicated. In the skin, as elsewhere, inflammation is produced by the action of irritants, and the form of the inflammation will depend greatly on the nature and source of the irritant.

**Causation.**—In some cases the origin and mode of action of the irritant is perfectly obvious, while in others it is very obscure. We have, for instance, septic inflammations, as in erysipelas, due to a definite microbe. We have also the various inflammations in connection with the acute fevers in which a specific morbid poison attacks the skin. But there are other cases in which the origin is very obscure, and in which the peculiarities of the individual play an important part. Just as there are persons especially liable to inflammations of the bronchi, of the kidneys, or of the intestine, so are there persons whose skin is prone to inflammation. Even in persons not predisposed temporary states of the nervous system apparently lead at times to a special tendency. At such times a very slight external irritation will lead to an inflammatory manifestation. There are indeed cases in which the nervous condition is the only obvious cause, as seen in a mild form in Urticaria, and in a more pronounced form in the so-called Trophoneuroses.

In considering the various forms of inflammation of the skin we shall divide them into four groups on the basis of their causation. These are (1) Inflammatory skin eruptions; (2) Symptomatic cutaneous inflammations; (3) Inflammations from heat, cold, and injury; (4) Septic and infective inflammations.

**1. Inflammatory skin eruptions.**—The affections included under this designation consist in various inflammatory affections of the skin due in some cases to local irritation, and in others to general irritation from conditions of the blood. In most of them individual peculiarities play a very important part, so that slight irritations produce eruptions in some people, while they have no effect on others. Indeed the tendency in many cases is so great that the conditions are regarded as idiopathic. There is no doubt also that nervous conditions have an important influence. The beginning of a skin eruption is often accompanied by itching, and the scratching which the itching induces not infrequently assists in the development of the eruption.



**Character of the lesions.**—The inflammation acting on the various structures of the skin produces the changes which occur elsewhere under like circumstances. There is hyperæmia, exudation of serous fluid and leucocytes, parenchymatous changes chiefly in the epidermis, and, if the inflammation be prolonged, newformation in the true skin. Various names are applied to the anatomical conditions thus brought about.

A **wheel** is a limited swelling of the skin due probably to an œdema of the superficial layers of the cutis. It is temporary, and in all cases the nervous system is concerned, as evidenced by the itching which is usually characteristic. It is produced by the local irritation of flea bites or by a general condition in urticaria. A **papule** is a small solid elevation of the skin, due generally to inflammatory exudation which infiltrates the papillary layer of the cutis and the epidermis. There is often proliferation of the epidermic cells as well. The term **Tubercle** is sometimes used, not in the histological sense, but to indicate an irregular rounded solid elevation from infiltration of the skin. In the **Vesicle** we have an accumulation of serous exudation raising the epidermis. The fluid is not between cutis and epidermis, but the separation takes place amongst the softer cells of the Malpighian layer, so that there is epidermis both below and above the fluid. The vesicles may be very small or so large as to form considerable **Blebs**, or **Bullæ**. A **Pustule** is like a vesicle but contains pus, and is due to a more intense inflammation. **Scales** or **Squamæ** are composed of epidermis which has been produced in excess and is less firmly attached than normal. The rapid production of epidermic cells is due to a parenchymatous inflammation, and the epidermic cells do not undergo the horny transformation to the same extent as normally.

The individual eruptions have received various names which only in part represent the pathological characters.

(a) **Urticaria.**—In this there is a formation of wheals in various parts of the body without any special external irritant. Such attacks are usually brought on by eating shell-fish, walnuts, or other special forms of food. These substances appear to produce their effects through the nervous system, inducing, by reflex action, a peculiarly sensitive state of the skin. The wheals occur especially in parts of the skin which are irritated by rubbing or by contact of the clothes, and there may be very striking œdema of the eyelids, for instance, when the feeling of itchiness has induced the person to rub them more than other parts.

(b) **Erythema and Roseola.**—These names are applied to mild inflammatory conditions in which the chief, and sometimes the only apparent, change is hyperæmia. The inflammatory nature of the condition is, however, often indicated by the existence of more or less swelling, due to œdema of the superficial layers of the skin (erythema nodosum and papulosum). The attacks are often followed by desquamation.

These very mild inflammations are frequently somewhat generalized, and appear



in many cases to be, like urticaria, related to special conditions of the nervous system, especially, perhaps, of the vaso-motor nerves. They may also be produced by external irritants, especially in susceptible persons.

(c) **Eczema**.—This name is applied to localized inflammations of the skin, usually of a subacute or chronic character. The mere fact of the local nature would indicate that the irritant here comes from without; but external irritation is hardly ever the entire explanation of the attacks. In no other disease do individual peculiarities and special states of the organism play a more important part. The external irritant may be very various, such as medicinal ointments, stuffs used by the patient in his occupation, as in dyeing, parasites which induce scratching, etc. The intensity of the inflammation depends on the susceptibility of the patient and the character of the irritant. The various degrees of intensity are, to a great extent, distinguished by the characters of the eruption, so that we have papules, vesicles, pustules, etc., characterizing different cases of eczema. From these differences are derived the names *Eczema papulosum*, *vesiculosum*, and *pustulosum*.

In chronic cases a condition is brought about somewhat comparable to catarrh of mucous membranes, and sometimes called *Catarrh* of the skin. The inflamed surface keeps on discharging serous fluid, not always to the same extent. The epidermis is softened, and, to a considerable extent, lost, so that the cutis is exposed or covered with irregular crusts. The cutis itself is red and it is thickened both by the serous exudation and by accumulation of round cells.

(d) **Psoriasis**.—The nature of the irritant is in this case quite unknown, but it is probably to be found in some special condition of the blood. We know, at least, that remedies introduced into the blood (arsenic, and, according to Napier, chrysophanic acid administered internally) frequently cure the disease.

The lesion is mainly of the epidermis, and especially of its Malpighian layer. The papillary layer of the cutis is hyperæmic and partly infiltrated with leucocytes; the papillæ are also described as enlarged, but these alterations are comparatively trifling, as, after death, very little indication of them is to be found. During life the hyperæmia gives the corium a red colour. In the middle regions of the Malpighian layer of the epidermis there is an abundant newformation of epidermic cells. The newformation is so great that there is no time for the cells to become horny before they pass to the surface. Hence the epidermis on the patch is entirely composed of Malpighian layer, and, these cells, being soft, as they dry they adhere much more readily into considerable scales than the horny cells do. For a similar reason, as they dry they shrink much more than the horny cells, and air insinuates itself into interstices between them. It is the finely divided air which gives the scales the peculiar silvery appearance characteristic of this disease (*Rindfleisch*). When the scales are removed the papillary layer of the skin is seen to be red and readily bleeds.

(e) **Pityriasis rubra**.—In this disease we have a more or less general affection, and it is not improbable that it depends on a lesion of the trophic centres of the skin. It is characterized by an excessive discharge of scales consisting of masses of epidermic cells. These scales pass off from slightly raised and reddened surfaces. In the earlier periods the cutis presents little more than hyperæmia with excess of leucocytes in its papillary layer, but when the disease has lasted for some time there is thinning of it with flattening of the papillæ. The sebaceous glands and hairs, as well as the sweat glands, are usually atrophied. There is often a very excessive discharge of epidermis.

(f) **Lichen ruber**.—This is a diffused papular eruption, depending like pityriasis



rubra on some obscure general cause. There are individual red papules on a red basis. As the disease goes on the skin gets infiltrated and thickened, so that the movements of the joints are hindered, and fissures are formed at the folds of the skin. The nails are usually thickened. The disease seems to consist in an inflammatory infiltration of the papillary layer of the cutis and the Malpighian layer of the epidermis. The cells of the rete Malpighii are enlarged and increased in number, and the interpapillary processes are enlarged. The papillæ show abundant accumulation of round cells, especially round the vessels. There is often accumulation of epidermic cells in the hair follicles.

(g) **Prurigo**.—This also is a diffused affection of the skin, which somewhat resembles urticaria, except that it is more permanent. Like urticaria it seems to depend on some condition of the nervous system, induced especially among children by neglect of sanitary conditions and defective nutrition. Numerous small papules appear on the skin and they are intensely itchy. The papules are due to an inflammatory thickening of the papillæ and rete Malpighii. There are round cells and a serous fluid exuded, so that when the head of the papule is removed by scratching a drop of fluid appears. The papules are often induced to bleed by scratching, and it is probably from this that they frequently leave a brown stain. The irritation produced by scratching may also lead to eczema.

(h) **Pemphigus**.—This condition is characterized by the formation of large vesicles or bullæ, filled with fluid, which is at first clear, but usually becomes more or less purulent. As the bullæ usually occur at intervals over a considerable surface, the disease is to be ascribed to some general cause, and in some acute cases the existence of an irritant in the blood is suspected. There is not here, as in small-pox, an exudation among the cells of the Malpighian layer, but for the most part the whole epidermis is raised, and the bleb is not divided by septa into loculi. If the bleb remain unruptured a layer of epidermis forms on the surface of the cutis. But if it burst, then the exposed cutis discharges for a time; by and by, however, a crust forms, under which the epidermis grows. In some cases the blebs are imperfectly formed and neighbouring ones coalesce, so that a considerable surface is affected. When these blebs burst there is little tendency to newformation of epidermis, and the skin continues discharging fluid as well as crusts produced by the fluid drying in. This form is designated **Pemphigus foliaceus**.

(i) **Acne**.—This name is applied to inflammation of the sebaceous glands and their neighbourhood, chiefly in connection with accumulation of the secretion in the gland. There are two conditions, named respectively comedo and milium, in which sebum accumulates in the glands; it is chiefly in connection with the former that acne occurs.

In **Comedo** a sebaceous follicle is filled with secretion, which forms a solid plug in the gland. The end of the plug presents at the surface, and has a black or dark blue colour. The plug consists of shed epidermis and sebum, with occasionally a small parasite, the *acarus folliculorum*.

In **Milium** there is an accumulation of the secretion with closure of the orifice of the gland. The accumulated secretion forms a rounded solid globe, which can be felt in the substance of the skin, but over which the epidermis passes unbroken. It is, in fact, a small retention cyst, and it contains the secretion of the gland mixed with epidermis, so that on cutting into it a whitish globular body is discharged. The cause of the closure of the duct is obscure.

In **Acne** there is a more or less acute inflammation of the sebaceous glands, usually in connection with comedo. There is redness and swelling around, and pus is



usually formed, and mingles with the substance of the comedo. Sometimes there is not a proper suppuration, but the substance of the plug is softened by the inflammatory exudation. The inflammation is due to the irritation of the plug, especially when by long stagnation it has become foul.

The name *Acne mentagra*, or *Sycosis*, is given to cases where there is inflammation of the sebaceous glands in connection with the hairs of the beard. There is a sycosis of parasitic origin, but in the simple form there is suppuration in and around the hair follicles of the nature of acne.

A more chronic inflammation of the sebaceous glands of the face, with special hyperæmia of the vessels around, has received the name *Acne rosacea*. There is very little tendency to suppuration, but great thickening of the skin occurs, so that sometimes there are large lobulated and red protuberances, especially on the nose, on the surface of which many comedones are seen.

**2. Symptomatic inflammations.**—This class embraces the inflammations occurring in the acute fevers. In some the cutaneous affection is very trivial. Thus the eruptions in measles, scarlatina, typhoid and typhus fevers, consist of little more than inflammatory hyperæmia with slight exudation, but also, in severe cases of typhus, hæmorrhages, forming petechiæ. The action of the irritant on the epidermis is evidenced in many cases by the subsequent occurrence of desquamation, which implies that the cells have been so injured as to lead to their premature necrosis and discharge. This occurs especially in scarlet fever and measles.

In **Small-pox** there is a much more severe inflammation. In this case the virus, consisting of micrococci which are present in the exudation or so-called lymph, evidently lodges in the skin and multiplies there, leading to pronounced local changes.

The effects of the irritant are seen mainly in the epidermis, which shows changes chiefly in the more plastic Malpighian layer. Many of the cells undergo a kind of coagulation-necrosis, their nuclei being lost and their substance converted into a hyaline material. These necrosed epidermic cells allow of spaces being formed among their layers in which exuded fluid collects. In this way a vesicle is formed, but it does not consist of a single cavity. As shown in Fig. 352, the epidermic cells or their remains form a network of fibres and partitions which divide the vesicle into compartments or loculi (*b* in figure) in which a serous fluid, containing the micrococci, is present. As the inflammation increases in intensity leucocytes are exuded, and these accumulate, as seen in the figure, both in the superficial layers of the cutis (*d*) and in the loculi of the vesicle. By the aggregation of these the vesicle becomes a pustule. In the figure it is seen that while the papillary layer of the skin (*d*) is somewhat infiltrated with leucocytes, yet it is not destroyed, and the interpapillary processes of the Malpighian layer of the epidermis (*c*) still persist. Sometimes no further



destruction occurs than this; the pustule dries in and a crust is formed and under it the Malpighian layer forms new epidermis, so that when



Fig. 352.—Section of a small-pox vesicle. Superficially there is a network formed of the altered epidermis. In some of the meshes (*b, b*) are pus-corpuscles. For the most part the Malpighian layer in its deepest cells (*c*) is preserved, but at some points, as at *d*, it is broken through and the pus is infiltrating the skin. *a*, umbilication of the vesicle, due to a sweat gland, the coils of which are distinguishable.  $\times 140$ . (After RINDFLEISCH.)

the crust is shed healing occurs without any loss of substance. But more commonly the action of the virus causes death of the superficial layers of the cutis as well as of the epidermis. This is shown in Fig. 353, where an injection thrown into the arteries runs into the surround-

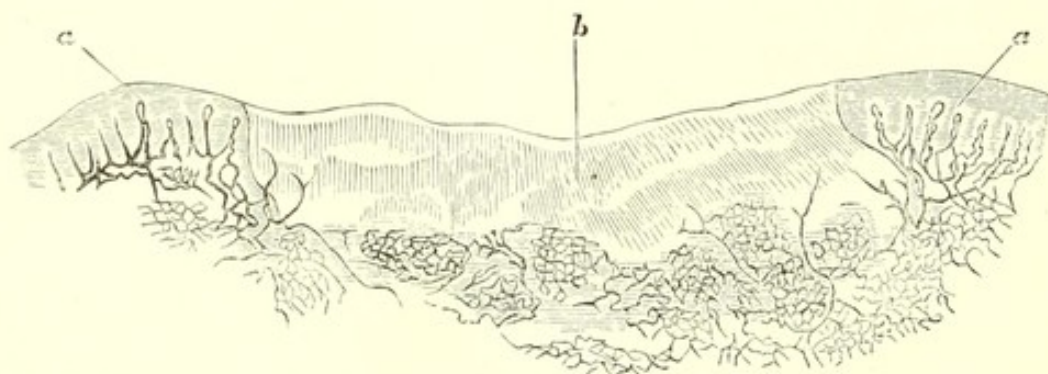


Fig. 353.—Small-pox eruption with necrosis, from an injected specimen. The normal skin at either side (*a, a*) has its vessels fully injected, while the necrosed part (*b*) is uninjected and homogeneous.  $\times 50$ . (RINDFLEISCH.)

ing skin (*a a*), but does not penetrate into the slough (*b*). In this case, when the crusts are torn, little sloughs infiltrated with pus are revealed, on the removal of which ulcers appear. By the healing of the flat ulcers we have the depressed reticulated cicatrices characteristic of small-pox (the so-called pitting of small-pox).



**3. Inflammations from heat, cold, and injury.**—(a) **Burns and scalds.**—We include here the lesions produced by the application of excessive heat to the skin, whether the heated substance applied be in the solid, liquid, or gaseous form. The heat acting as an irritant damages or kills the tissue, and so we have signs of inflammation, frequently associated with necrosis.

Burns have been distinguished, according to their intensity, into three degrees, characterized by the three phenomena—hyperæmia, vesication, sloughing.

In the **First degree** the heat is only sufficient to produce a hyperæmia of the skin. Exposure to the rays of the sun, or to hot water of a temperature of 100° F. and upwards, causes such a relaxation of the arteries as to induce considerable congestion. The inflammatory nature of this hyperæmia is shown by its continuance for some time after removal of the cause, and by the occasional occurrence of some œdematous swelling of the skin. After a time there may be desquamation.

The **Second degree** is produced by the action of a temperature of 140° to 185° F., or by the evanescent action of a still higher temperature. The effect is to cause an inflammatory exudation from the papillary layer of the skin, which accumulates in the epidermis so as to form a vesicle. The changes in the rete Malpighii are similar to those already described as occurring in the formation of the small-pox vesicle, but they are more rapidly produced and the layer of cells is more homogeneously affected. The inflammation soon subsides, and from the remaining cells of the rete Malpighii new epidermis is reproduced; there may be already a complete coating of new epidermis before the vesicle bursts. But sometimes the vesicle bursts early, and the inflamed surface of the skin is exposed; or nearly the whole of the rete Malpighii may have been destroyed, even the interpapillary processes. In these cases the covering over of the surface with epidermis may be delayed, and the surface may even become like a granulating wound, discharging pus.

In the **Third degree** there is necrosis not only of the epidermis but of the true skin. The temperature is over 140° F., and if very high may have acted for a very short time. As a general rule the necrosed tissue lies as a dry crust on the surface. An inflammatory process occurs around it, with exudation, going on usually to supuration. In this way the slough is separated, and a granulating wound is left, which heals in the usual way, leaving a cicatrix, the character of which varies with the depth of the slough.

Reference has already been made to the fact that Hæmoglobinuria sometimes follows burns (p. 96), and also that reduction of temperature may occur (p. 351). These, along with reduction of blood-pressure from dilatation of arteries, probably have to do with the occasional rapid occurrence of death from extensive burns.

If the person survives for some time after the burning, inflammations of internal organs, the lungs, kidneys, serous membranes, are frequently found; also fatty degeneration of the heart and of the liver. These are to be ascribed with great probability to the alteration of the blood.



It is much more difficult to account for the occurrence of the **Duodenal ulcer** after burns. When several weeks elapse before death, there is found, in about 20 per cent. of the cases, one or more ulcers, usually situated in the duodenum near the pylorus, but sometimes in the pyloric region of the stomach. It may be a superficial hæmorrhagic erosion or a deeply penetrating ulcer.

(*b*) **Frost-bite and Chilblain.**—These conditions are produced by the action of excessively low temperatures. We can distinguish three degrees here also.

The common **Chilblain** is an example of the **First degree**. If portions of the body are exposed to a low temperature and then heated, a chronic or subacute inflammation is induced. This occurs especially in the fingers and toes. There is passive hyperæmia sometimes amounting almost to stasis, so that the part has a livid hue. There is also serous exudation, so that the part is swollen and oedematous; there may be red corpuscles as well as leucocytes in the exudation. Sometimes ulceration occurs.

The **Second degree** includes the **Less severe cases of frost-bite**. The skin has been frozen and too rapidly thawed, the result being such an injury to the tissue as to cause an acute inflammation. The epidermis is raised in blisters which often contain a bloody serum. There may even be sloughing of the superficial layers of the cutis, and by the separation of the slough an indolent ulcer forms.

In the **Third degree**, forming the **More severe cases of frost-bite**, there has been prolonged exposure to very intense cold. The consequence is the complete freezing of part of the body, especially in regions removed from the heart, as the fingers and toes. The tissues are stiffened by freezing, and the blood coagulated, so that sometimes parts can be broken off like glass. When the parts are thawed gangrene sets in. The whole frozen part does not die, but there is partial recovery with inflammation. The gangrenous part is separated in the usual way, generally with decomposition (moist gangrene).

(*c*) **Wounds and Excoriations.**—The phenomena connected with the healing of wounds and the formation of granulations have been considered in previous pages. When merely a portion of the epidermis is removed by an injury, forming an **Excoriation**, it is restored with remarkable celerity. The cells at the margin and those of the deeper layers divide and form new cells which fill up the gap.

In the case of a deeper wound the epidermis also shows great vigour, and it sometimes advances beyond the line of the other healing processes. Dr. Macewen has pointed out to the author that the new-formed epidermis may advance over a blood-clot filling up a wound.



The epidermis here has a remarkably transparent hyaline appearance and is continuous, not with the Malpighian layer, but with the layers superficial to that. The character of the cells here and their transparency suggest that they may be derived from the stratum lucidum. It may, indeed, be suggested that this layer in the normal skin is composed of young epidermis, and that the loss on the surface is supplied by newformation here and not in the Malpighian layer.

**4. Septic and Infective inflammations.**—We include here those forms of inflammation of the skin which can be traced to the action of microbes, leaving over those which have the characters of the specific newformations.

(a) **Boil and Carbuncle.**—A boil is a localized acute suppurative inflammation with a limited necrosis of the cutis. The irritant finds access by the sebaceous glands or hair follicles and consists of pyogenic microbes. The source of these may be obscure, but in some cases they are derived from contact with the cadaver. The inflammation manifests itself in hyperæmia and exudation, so that a localized redness and swelling are the results. A piece of skin in the very centre of the inflamed area dies, and a small abscess having formed, the slough is by degrees discharged along with the pus.

The carbuncle is similar in its general characters to the boil, but a much larger piece of skin is involved. The slough is therefore of much greater extent. It sometimes happens that the whole piece of skin in its entire thickness dies and is separated as a dry leathery slough. More frequently the necrosis is less extensive in the superficial layers than in the deeper parts, and the slough is discharged through numerous small apertures.

(b) **Lesions from cadaveric poisons.**—In the juices of the dead body when in process of decomposition, there are innumerable bacteria of different kinds. Some of these may attack the skin of the dissector or pathologist, and find a lodgement there. The organisms themselves are of very various degrees of virulence. If the subject has died of a septic disease, then the organisms which have during the life of the patient been active factors in the production of the disease, may remain specially virulent after death. The fluid in the abdomen after septic peritonitis, such as that which occurs in one form of puerperal fever, teems with micrococci, and this fluid is peculiarly virulent when applied to the skin. So is it also with septic pleurisy, etc.

Besides degrees of activity in the virus, there are various degrees in the susceptibility of different persons, and of the same person at different times. Frequent exposure to the cadaveric viruses creates a certain immunity, so that a person will be more secure against them when frequently engaged in post-mortem work than



when making only an occasional examination. A state of exhaustion of the body causes a greater degree of susceptibility, and makes the inroads of the virus when once implanted more vigorous than in persons in ordinary health. The inference from this is, that the best treatment for this condition is to leave off work and at once seek rest and fresh air.

The virus very often finds access to the skin without any breach of surface. When a wound is made during dissection it is usually washed thoroughly and sucked, while bleeding is encouraged; in this way the virus may be washed away. But if it gets into the hair follicles or sweat glands it may lodge and multiply undisturbed.

The immediate effects are usually an acute localized inflammation of the skin, resulting in the formation of a small **Pustule**. The virus may have attacked at several points, and there is a crop of pustules, usually on the back of the hand or forearm where the epidermis is thin, the left hand being more frequently affected than the right, as it is most frequently exposed in separating and holding the organs. In many cases nothing more than the pustules is produced, and in the case of the

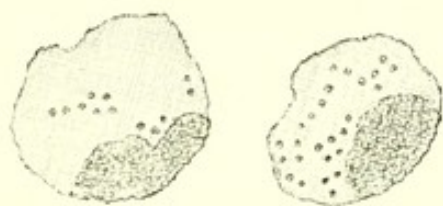


Fig. 354.—Pus-corpuses with micrococci in them from a cadaveric pustule.  $\times 1500$ .

regular pathologist a crop of pustules in the spring may be his annual experience. The pus contains micrococci, some of them free, but some in the substance of the pus-corpuses (see Fig. 354).

Sometimes, however, there are more severe local effects, and sometimes the virus travels farther inwards. Instead of dying away, the pustule, after partially disappearing, sometimes leads to necrosis and acute inflammation in a larger area around, so that a **boil** is the result. In this case there is a cicatrix left when healing occurs. In other cases it is what has been called a **blind boil** which develops after the pustule; there is a more chronic inflammation of the skin with considerable thickening and elevation but without suppuration.

The **Pathological wart** is a still more chronic lesion. It presents itself as an irregular thickening of the skin with a warty or papillary surface. This lesion is sometimes produced by the inoculation of the tubercular bacillus, forming a local tuberculosis. The bacilli have been found in the lesion.

In some cases the virus propagates itself upwards towards the trunk. It may be that an erysipelas starts from the local inflammation, extending upwards along the lymph spaces of the skin. In other cases the axillary glands become inflamed without any intervening lesions, the virus passing up the vessels and only producing mischief after lodging in the glands. Here suppuration is common. An inflammation of the



skin resembling erysipelas may spread from the glands, as the virus, checked in its upward course towards the trunk by the gland, passes outwards to the lymph spaces around. This course of events is shown in the account which Paget has given of his own case in his "Clinical Lectures and Essays." The inflammation may travel along the connective tissue of the skin and subcutaneous tissue, and may even reach the pleura. In the skin its effects may be very severe, resulting in sloughing and profuse suppuration like that in erysipelas. It may even lead to septic infection or pyæmia.

(c) **Phlegmonous inflammations. Erysipelas.**—These names are given to acute infective inflammations of the skin. In the case of erysipelas the virus depends on the streptococcus erysipelatis which extends more or less abundantly in the lymph spaces and vessels of the skin (see Fig. 355) and subcutaneous tissue. The result is a more or less severe inflammation. At first there is an inflammatory hyperæmia, which advances as the virus propagates along the lymph spaces. This, with a moderate exudation of fluid and leucocytes, may be all the phenomena. But sometimes there is a much more intense inflammation. The skin is infiltrated with leucocytes to such an extent that it is softened and opened out with pus. The epidermis is raised by exudation so that vesicles or pustules are formed. Sometimes the lymph spaces are occupied with fibrine. Besides suppuration, there is often necrosis of portions of the skin so that sloughs are separated with the pus.

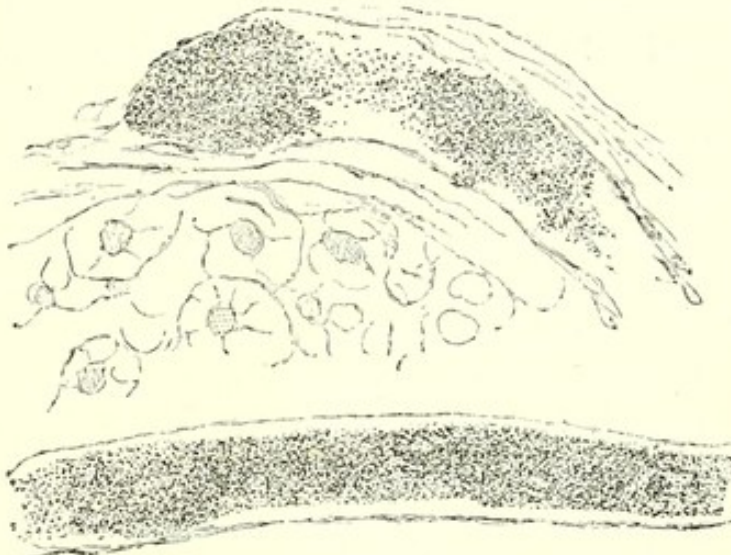


Fig. 355.—From the skin in a case of erysipelas. The upper illustration shows a lymphatic vessel at the border of a sebaceous gland, filled with micrococci. The lower shows a straight vessel similarly filled.  $\times 350$ .

A somewhat similar phlegmonous inflammation sometimes occurs around septic wounds, just as we have described in cadaveric infection.

(d) **Malignant œdema.**—This depends on the action of a specific microbe, the bacillus of malignant œdema (see p. 299). The affection usually attacks open wounds, and the most typical result is an advancing gangrene with development of gas, accompanied by acute inflammation. The author has observed a bacillus similar to this one in a case of cancrum oris. (See ante.)

(e) **Splenic fever or Anthrax.**—This disease also depends on a special



bacillus, which, however, usually extends to the blood, and the patient dies from the general specific fever (see p. 295). There is usually a local affection to begin with, and the disease may remain local. The local affection has the characters of a phlegmonous inflammation of the skin and subcutaneous tissue. Sometimes the appearances of malignant oedema are produced. From these local manifestations the condition is sometimes called **Malignant pustule**.

(f) **Hospital gangrene**.—This is an infective disease accompanied by sloughing of the skin. It is due to a specific virus which is supposed to be a micrococcus. The disease occurs in unhealthy hospitals, especially in time of war. It attacks open wounds, especially granulating wounds, and is accompanied either by sloughing or molecular necrosis. The gangrene frequently travels along the loose connective tissue and so may undermine the skin, or isolate muscles or blood-vessels.

#### IV.—SPECIFIC NEWFORMATIONS.

1. **Syphilitic lesions of the skin**.—The various manifestations in the skin are the results of the action of the syphilitic virus. The lesions, like those elsewhere, are either simply inflammatory, or they have the characters of the gumma, forming in the latter case more or less definite tumours composed of granulation tissue. Even the purely inflammatory lesions have a tendency, when they have persisted long, to present considerable newformation of granulation tissue. The syphilitic lesions of the skin may be divided into the primary, secondary, and tertiary.

(a) **The Hard chancre** has been described at p. 191, and its structure is illustrated in Fig. 47, p. 190.

(b) **Lesions of secondary stage**.—As the virus is in the blood the manifestations occur as a rule over the skin as a whole, although circumstances may determine a certain local selection. The eruptions are for the most part symmetrical, and they present a peculiar tendency to occur in round patches. These patches, by healing in the centres and extending at the periphery, frequently assume a circular or serpiginous shape.

The syphilitic eruptions of the secondary stage are somewhat similar in character to the simple inflammatory eruptions, and they are commonly designated by similar names. There is, however, more tendency in the syphilides to the development of granulation tissue in the skin, and hence the papular forms of eruption are more frequent. The syphilitic eruptions present very various characters, and there is no form which is constantly present at a particular stage. In the earliest periods there is usually a generalized hyperæmia of the skin (*syphilitic*



*erythema* or *roseola*). A more local development of elevated papules, generally in groups, constitutes the *syphilitic lichen*. In these papules, the corium, especially its superficial layers, and to a certain extent the epidermis, are infiltrated with round cells; generally there is desquamation of epidermis, and this is sometimes very prominent. Occurring in the palms of the hands and soles of the feet this desquamation is so characteristic as to give rise to the designation *syphilitic psoriasis*, the papules after a time coalescing and so merely producing a generally raised surface.

The **Condyloma** (*Plaque muqueuse, mucous tubercle*) is a further development of the papule. It occurs usually in situations where two surfaces of the skin are in contact and are thus kept moist, as in the neighbourhood of the genital organs, in the axilla, beneath the mamma, etc. We have here a very marked and extensive infiltration of the papillary layer of the skin with round cells, so that an elevated surface is formed. Sometimes there is a special elongation of the papilla so that a warty surface is produced, giving rise to the **Pointed condyloma**. Sometimes the condyloma breaks down on the surface and an ulcer forms.

The syphilitic papules sometimes develop into pustules or vesicles. The pustule forms by the epidermis being raised from the surface, while a fluid inflammatory exudation occurs between it and the corium, which latter remains infiltrated with round cells. The pustule or vesicle and its infiltrated base may enlarge so as to produce a broad bleb on a red raised base (*Syphilitic pemphigus*). The blebs often dry-in and form raised crusts (*Rupia*). In these cases the pustules are as if on the surface of condylomata. This is still more the case in **Pemphigus neonatorum**, which is one of the manifestations of hereditary syphilis. Here the corium is affected somewhat as in the condyloma, but the infiltration has more the characters of granulation tissue. In the condyloma the structure of the skin remains to a great extent, but it is infiltrated with round cells, which are mostly leucocytes. In pemphigus neonatorum, however, the tissue is replaced by a vascular tissue in which are many large cells, the condition approaching to that of the gumma. On the surface of this the epidermis is raised, forming a vesicle or pustule.

(c) **The Gumma**.—This is not of very common occurrence in the skin; it is, as we have seen, a special tertiary manifestation. It begins as a hard swelling in the cutis. It increases in size and raises the surface, sometimes forming a tumour of considerable dimensions. It is peculiarly prone to ulcerate, and after ulceration has occurred it may go on extending, the infiltrated edges gradually giving way.

2. **Tuberculosis of the skin**.—There is still some doubt as to the exact extent of this group, and more particularly whether lupus belongs



to it. Undoubtedly this affection presents all the characters of an infective disease, and the bacillus tuberculosis has been observed in it.

**Lupus** occurs chiefly in children, and attacks the face. It consists in an advancing infiltration of the cutis with granulation tissue, in the midst of which giant-cells are frequently visible and sometimes defined tubercles. These, however, are apt to be overwhelmed by the cellular infiltration. There is commonly an exaggeration of the epidermis, so that epidermic processes may penetrate somewhat among the granulation tissue. With all this there is a destruction of the specific structures of the skin. The sebaceous glands and hair follicles are destroyed, but it is said that the sweat glands, being situated beneath the cutis, are spared.

The tubercular newformation undergoes degenerative changes, which lead frequently to ulceration, but sometimes cause cicatrization without ulceration.

The appearances presented to the naked eye vary considerably. There is a local infiltration of the skin, giving an irregular raised surface. The thickening may be very great (*L. hypertrophicus*); or there may be considerable thickening and desquamation of the epidermis (*L. exfoliativa*). Ulceration is frequently manifest (*L. exedens* or *ulcerans*), and the ulcers are often very varied in shape; or ulceration may be absent (*L. non-exedens*).

Lupus is not usually accompanied by tubercular manifestations in other parts, except that the associated lymphatic glands are not infrequently affected.

**Scrofuloderma** is a name applied to tubercular affections which have not the characters of lupus, and occur mostly on the extremities, neck and trunk. There is infiltration of the cutis, sometimes in such a localized form as to produce a warty appearance. (The pathological wart or tubercle, as stated above, is sometimes of this nature.) Ulcers with infiltrated edges are frequently produced.

3. **Leprosy** has been already described at page 211.

4. **Elephantiasis** has also been considered at page 212.

5. **Frambæsia**.—This name is given to a disease evidently caused by a virus which acts on the skin. There are prominent irregular swellings produced, which have the general appearance of exuberant granulations. These have been compared in appearance to strawberries and raspberries. The local names **Yaws** (African for raspberry) and **Pian** (mulberry) also suggest these resemblances. The disease is endemic in the West Indies and Africa; it also occurs in Peru, where it is called **Ver-ruga**. It is contagious, and in many respects so much resembles syphilis that some authors have regarded it as the same disease.



It begins as a small raised spot, usually on the face, on the upper or lower extremities, or near the organs of generation. The spot enlarges and gradually assumes the red irregular appearance which has suggested the names of the disease. It may continue for months or years, and, neighbouring areas coalescing, it may affect a large portion of the cutaneous surface.

The structure of these growths closely resembles that of the syphilitic chancre, as may be seen by comparing Fig. 356 with Fig. 47, page 190.



Fig. 356.—Section of skin in frambœsia, from a part where the disease was not very far advanced. The papillary layer of the cutis is seen to be replaced by granulation tissue, while the epidermis is preserved on the surface.  $\times 75$ .

The cutis is converted into granulation tissue, whose bulky masses form the irregular prominences characteristic of the disease. The epidermis may be continuous over even the most prominent of these masses (see figure), and may even be exaggerated; but the horny layer is lost and the soft Malpighian layer alone remains. The epidermis may give way, and an ulcer form, from which a thin fluid is discharged, not usually a true pus.



## V.—TROPHONEUROSES OF THE SKIN.

We include under this heading lesions of the skin in which, by reason of disease of the nervous system, such changes occur in the nutritive processes as to lead to definite anatomical results. The existence of separate trophic nerves for the skin has not been anatomically demonstrated. But there are evidences, especially in pathological processes, that trophic centres exist, and that fibres convey impulses of this kind to the skin. The ganglia of the posterior roots of the spinal nerves and the similar Gasserian ganglion seem to contain trophic centres, and the trophic fibres are either identical with the sensory ones or run along with them.

1. **Herpes.**—This disease is typically of nervous origin. It is commonly accompanied by neuralgia, and in its site it nearly always follows the distribution of a single nerve stem. It appears, for instance, in the course of an intercostal nerve, or occupies one side of the face in the area of distribution of the fifth.

The eruption in the skin is in the form of vesicles which somewhat rapidly pass through a series of changes. There is first a group of slight elevations occupying a reddened patch of skin, and each of these rapidly develops a vesicle; in a few days the vesicle dries up into a crust, under which fresh epidermis is formed. In its details the eruption is inflammatory. There is hyperæmia and œdema of the papillary layer of the skin. The serous exudation collects in the Malpighian layer of the epidermis, separating its cells just as in the case of the small-pox eruption. The vesicle is divided by a network composed of the elongated and contorted epidermic cells, which often present clear spaces or vacuoles in their substance. In the serous contents of the vesicles there are leucocytes, and these may accumulate till the fluid approaches to the nature of pus. The papillary layer is also frequently the seat of infiltration with round cells. Occasionally there is hæmorrhage, and the papillary layer of the skin may be destroyed, so that when healing occurs by granulation a cicatrix may be the result.

The nervous origin of herpes has been abundantly established. Direct injury to a nerve, as by a gunshot or other wound, may cause it. It is to be presumed that, in this case, the inflammatory process in the wound irritates the nerve, and the cutaneous inflammation may be regarded as due to the irritation of trophic fibres. Spontaneous herpes, and especially herpes zoster, has been found associated with inflammations of the intervertebral or Gasserian ganglia, but it has been questioned whether the trophic centres have actually their seats there.

The herpetic eruption may arise from disease of the spinal cord and brain. In the case of the cord it is chiefly in locomotor ataxia that we meet with herpes, and the eruption is accompanied by the severe pains characteristic of that disease. We



know that in locomotor ataxia it is the posterior columns of the cord that are affected, and this would indicate that the cutaneous trophic centres or fibres run in these parts, while the coincidence of neuralgic pains also indicates a sensory locality. It is here again to be presumed that the lesion producing the herpes is an irritative one as indicated by the violent coincident pains.

But we may have herpetic eruptions from disease in the brain. There are cases recorded in which one half of the body was affected, and in some of these there was a hemiplegia due to a coarse lesion in the brain. It is clear that there must be trophic fibres and centres in the brain as well as in the cord, and their irritation may lead to herpes. There are also a few cases on record of universal or nearly universal herpes, as if a general centre for the whole body were irritated.

**2. Glossy skin.**—This name was applied by Paget to a condition which he observed in connection with injuries or wounds of the nerves. There is here again indication of irritation of the nerves, as there is always neuralgic pain, described by Mitchell as burning pain. The condition also occurs sometimes after an attack of herpes (Paget), and it may be associated with eczema. It may also follow disease of the spinal cord (Mitchell).

The disease occurs mainly in the hand, and especially in the fingers. The affected fingers "are usually tapering, smooth, hairless, almost devoid of wrinkles, glossy, pink, or ruddy, or blotched as if with permanent chilblains" (Paget). The most striking peculiarity is the shining, glossy appearance of the surface as if it had been varnished. The comparison with chilblains is usually applicable, but sometimes the appearance is rather that of highly polished scars, and the condition resembles that in morphœa. In the affected part the hairs mostly disappear, and the nails undergo peculiar changes. The latter become greatly curved, much more than in cases of phthisis, while the skin at the root of the nail becomes retracted, leaving the sensitive matrix partly exposed. The disease affects mainly the fingers and toes, but it may extend to the palm of the hand or the dorsum of the foot. The histological details in this condition are unknown.

**3. Scleroderma and Morphœa.**—The name *scleroderma adultorum* is given to a rare form of skin disease which affects considerable areas and is usually symmetrical. Morphœa or **Addison's keloid** has similar anatomical characters, but occurs in localized patches.

In the more diffuse form of scleroderma proper, extensive tracts of skin are sometimes involved, such as the greater part of the face, the arms, etc. In a case observed by the author, the face, chest, and both forearms and hands were affected, and dry gangrene had occurred in the fingers, which had a stiff black appearance.

The nature of these affections is not quite determined, but many facts point to a nervous origin. There is the symmetrical character of the lesion and its wide distribution, as well as the fact that lesions of the sympathetic have been believed



to give rise to it. The lesions, consisting of a slow increase of the connective tissue with atrophy of the special structures, resemble those which occur in muscles as a result of severance from their trophic centres (see p. 535).

In its anatomical details the skin in both forms presents characters which may be summarized as atrophy with chronic inflammation. The papillæ are mostly flattened, and the cutis thinner than normal; it is composed also of more homogeneous connective tissue than usual, having somewhat the characters of cicatricial tissue, and less fibrous than that of the normal skin. This condition extends somewhat to the subcutaneous tissue where the fat is to a considerable extent atrophied, apparently by the encroachment of the connective tissue. The hairs are also atrophied in the affected areas, and there is said to be alopecia when the patches extend to the hairy scalp. The evidence of chronic inflammation is the existence of abundant nuclei in the altered skin. These are usually aggregated around the blood-vessels, but they are also seen around sebaceous glands and under the rete Malpighii.

Complete recovery seems to be not uncommon in this disease, although it may be slow.

Allied to scleroderma in the general characters of the lesion in the skin is the condition commonly designated **Sclerema neonatorum**. In this disease the skin, usually of the lower extremities, is hard and swollen, so that movement of the limbs is curtailed. When it persists for some time the skin may be like parchment.

**Literature.**—*Herpes*—PICAUD, Des eruptions cutan. conséc. aux lésions traumat., 1875; MITCHELL, Injuries of nerves, etc., 1872; BÄRENSPRUNG, Charité-Annalen, ix., 1863; KAPOSI, Hautkrankh., 1882; CHARCOT, Dis. of nerv. syst., i. (Syd. Soc.), 1877. *Glossy skin*—PAGET, Med. Times and Gazette, i., 1864; MITCHELL, l. c. *Scleroderma*—HILTON FAGGE, Guy's Hosp. Rep., 1867, xv. 187; CHIARI and MADER, Vierteljahrsschr. f. Dermat u. Syph., 1878; HUTCHINSON, Lect. on clin. surg., 1878; SCHWIMMER, Die neuropath. Dermatosen, 1883; M'CALMAN, Glasg. Med. Jour., xvi., 1881; STEVEN, ibid., xxvi., 1886. *Addison's Keloid*—ADDISON, Published writings (Syd. Soc.), 1869; HILTON FAGGE, l. c.; HUTCHINSON, l. c.

## VI.—HYPERTROPHIES AND TUMOURS OF THE SKIN.

**Ichthyosis.**—This is a disease which apparently owes its origin to a congenital defect in the constitution of the skin. The disease itself is sometimes manifest at birth, but if not it begins in the earlier years of life, gradually extends over the whole body, and continues during life. It is in a striking manner inherited (see p. 19).

It consists, anatomically, in an affection of the epidermis, which is sometimes greatly thickened and irregular. The thickened epidermis forms scales of larger or smaller size, sometimes producing merely a



furfuraceous exfoliation, but in more severe cases giving rise to large plates like fish scales. From this latter appearance the name of the disease is derived. The scales do not consist entirely of epidermis, but contain sebaceous material, which is sometimes so abundant as to give the scales a polished appearance like mother-of-pearl. The large scales may be so fixed as to give an imbricated appearance. The milder forms of the disease, in which there is little more than an excessive scaliness of the surface, are often included under the designation **Ichthyosis simplex** or **Xeroderma**, while the more severe forms, in which there are prominent scales, are distinguished by the name **Ichthyosis cornea** or **Ichthyosis hystrix**.

In the milder forms the cutis is not markedly affected, but in some cases the papillæ are greatly elongated. The substance of the corium is often atrophied, and the subcutaneous fat deficient.

**Callosity.**—This is an exaggeration of the horny epidermis occurring where the parts are exposed to unusual friction. It is a true hypertrophy, being a direct provision of nature to protect the true skin from the excessive friction to which it is exposed. The best example of the callosity is in the horny hands of workmen, but it is often seen in the feet, at points which are peculiarly exposed to pressure.

The **Corn** (*clavus*) is closely related to the callosity. Here, also, there is friction, but it is associated with compression. It occurs chiefly in the foot where there is a concentric pressure from ill-fitting boots, and friction against either the boot or a neighbouring toe. There is, as in the former case, an excessive newformation of horny epidermis, but, by the concentric pressure, the layers of epidermis have their directions changed, and growing against each other, they are projected inwards towards the cutis. There is thus an ingrowing kernel of hard horny epidermis which has given rise to the name of corn. The cutis is irritated and undergoes atrophy. Around the kernel the epidermis is always thickened, the condition being like that of the callosity.

**Common wart.**—This may be variously regarded either as a tumour or a hypertrophy of the skin. The soft warts have distinctly the characters of tumours, presenting specialities of structure and growth which entitle them to that designation (see below). But the common wart, consisting for the most part of a local exaggeration of the normal structures and occurring in numbers in the same person, has more doubtfully the characters which we ascribe to a morbid growth. A group of papillæ are elongated, and the epidermis over them thickened, so that the characters are those of a hard papilloma. The **Horn** is of similar structure to the wart, but the papillæ are usually more elongated,



while the horny layer of the epidermis is compacted into a more consistent mass which forms a prominent outgrowth.

**Soft warts and Pigmented moles** (*Nævi pigmentosi*).—These are very often congenital, or they occur very early in life, and usually remain stationary throughout life. They are smaller or larger elevations of the skin, either colourless or pigmented, and frequently beset with hairs. Many of them present a somewhat peculiar structure, as shown in figure



Fig. 357.—Section of a soft wart. It is covered with epidermis and consists of a very cellular tissue. The cells were spindle-shaped. To the left a hair-follicle and sebaceous gland.

357, especially considering that they exist unchanged for many years. They are covered by epidermis which shows little difference from the normal. But instead of the normal tissue of the cutis beneath the epidermis there are masses of cells separated by connective tissue. The cells in these nests are usually large and resemble large sarcomatous cells. The presumption is that these structures are really survivals of the foetal condition, and their importance in relation to tumours has already been referred to at p. 220. In their structure it will be observed that these warts are much more like tumours than the ordinary warts, and that their tissue conforms to that of the cellular tumours, probably the sarcomas. Sometimes a soft wart is composed of a mass of adipose tissue covered by epidermis.

**True keloid.**—This is an elevation of the skin which has in its marginal parts branching processes extending out. There is evidently considerable contraction and dragging produced by the central tumour and the radiating processes, so that the skin is puckered around as in the case of a contracting cicatrix. The lesion has many of the characters of a definite tumour. It is composed, in the fully developed state, of dense connective tissue which may be unduly cellular. The newformation has its seat in the deeper layers of the cutis, and it is



stated that the papillary layer is intact. In the earlier stages it consists mainly of spindle cells, and Virchow therefore regarded it as a sarcoma. The analogy to the sarcoma is supported by the fact that the tumour recurs on removal. The most common site of keloid is the front or back of the chest, but it is found in other parts.

The name **False keloid** is sometimes used to designate a condition which occurs in **Scars**. A localized thickening, usually in the midst of a large cicatrix, as from a burn or an ulcer, presents itself, and there are raised processes passing out from it as in the true keloid.

**Molluscum contagiosum.**—This name is applied to little growths of the skin, occurring chiefly on the face and showing a preference for the eyelids, but sometimes extending so as to be almost universal over the body. It will be inferred that the growths are multiple. The individual ones have been compared to pearl buttons, which they resemble in their circular shape and central depression, but they are usually pink in colour. There is considerable elevation of the little growth, and, although generally sessile, it sometimes becomes pedunculated and may even drop off.

In the centre of the elevation a depression is visible which is probably the orifice of a sebaceous gland, and the tumour seems to consist of an enlarged sebaceous gland. A section (see Fig. 358) shows the

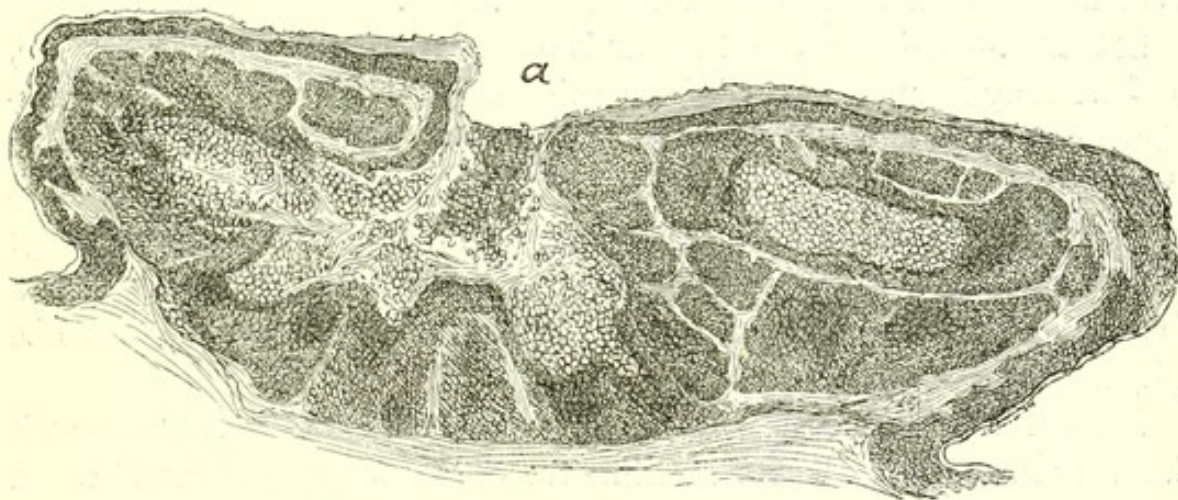


Fig. 358.—*Molluscum contagiosum*. It forms a flat elevation as shown by the position of the epidermic surface on either side. The tumour is probably an enlarged sebaceous gland, whose orifice at *a* is open.  $\times 20$

exaggerated lobules of the gland, while within there is an irregular cavity containing a white material which sends prolongations into the gland lobules. The lobules of the gland consist of epithelium, the peripheral cells of which are cylindrical. In the midst of the sebaceous material are found peculiar glancing bodies (see figure) compared to swollen starch granules. Several authors deny the origin from sebaceous glands, ascribing it rather to the Malpighian layer of the



epidermis, the virus being introduced by the hair follicles (see especially *Thin*).

The disease depends on some virus, and, as the name implies, it is contagious. It generally occurs in groups in the same person, as if spreading, and several members of the same family, or several persons living together, are usually affected simultaneously. It is usually seen in children, but occurs also in adults.

**Fibroma molluscum.**—This condition is to be carefully distinguished from that just mentioned. It has been described at p. 227.

Small **Fibromas** have been frequently observed in the skin in rheumatic cases, especially in children. They are usually temporary. Larger tumours have been observed associated with rheumatism in adults.

**Xanthoma or Xanthelasma.**—This name is applied to slight elevations of the skin of a yellowish colour. They occur mostly on the eyelids, where they appear as firm yellow patches, which are usually quite flat, but may be slightly tuberculated. They consist of firm connective tissue in which finely divided fat granules are abundantly present. The latter give the opaque yellow character to the lesion.

Of the remaining simple-tissue tumours, the **Lipoma** is common, originating in the subcutaneous tissue. The **Myxoma** is not infrequent, and the **Enchondroma** and **Osteoma** very unusual. The **Angioma** of the skin is common as the congenital *nævus* (see p. 242).

**Sarcoma.**—This form of tumour is of somewhat frequent occurrence, and we may have round-celled or spindle-celled growths, which sometimes assume considerable dimensions. Sarcomas sometimes originate from soft warts or moles, and when they do so they frequently present a similar structure to that described above, namely masses of large round cells in a stroma. To this form the name **Alveolar Sarcoma** is applicable. Sarcomas originating from moles are peculiarly apt to be pigmented.

Sarcomas rarely occur as secondary tumours in the skin, but a case is known to the author where multiple tumours composed of round-celled tissue formed at intervals in the subcutaneous tissue and some of them disappeared. These were either round-celled sarcomas or malignant lymphomas.

**Cancers.**—The **Epithelioma** of the skin forms the majority of the flat-celled epitheliomas, and it has been already considered at p. 269.

The two most interesting forms are the ordinary flat-celled epithelioma and the rodent ulcer. The former is met with most frequently in the lower lip, but also occurs not uncommonly in the penis and scrotum. In the last-mentioned seat it appears to occur with special frequency in chimney-sweepers and in workers in paraffin-refineries. The rodent



ulcer occurs in the face, especially on the eyelids and side of nose. (See p. 270 and Fig. 93.)

**Ordinary cancer** is of rare occurrence in the skin, but cases have been observed to which the names scirrhus and soft cancer have been given. **Pigmented cancers** are of occasional occurrence, and they are of importance on account of their tendency to form secondary tumours, first in the lymphatic glands and then throughout the body. They are, however, much less frequent than melanotic sarcomas with which they are apt to be confused.

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## VII.—PARASITIC DISEASES OF THE SKIN.

We have already seen that certain forms of inflammation are associated with the presence of microbes, especially the infective inflammations. We saw also that leprosy depends on a specific bacillus. These do not require further treatment here.

**Fungi** are present in three forms in the skin.

1. **Tinea favosa or Favus.**—This is a disease chiefly of the hairy scalp, although it occurs on other parts of the skin and also in the nails. On the scalp it forms crusts about the size of split peas, which, when removed, are usually seen to be somewhat cup-shaped, the convexity of the cup being next the skin, the latter presenting a corresponding depression. If the cups, which have a characteristic saffron-yellow colour, be divided with a knife they are seen to have a porous appearance, the interstices being filled with air, and they are brittle, so that they can easily be broken down in water.

The crusts are formed of fungus along with epidermic scales and their debris. The fungus is the *Achorion Schoenleinii*, in the form for the most part of short tubes (Fig. 359) with

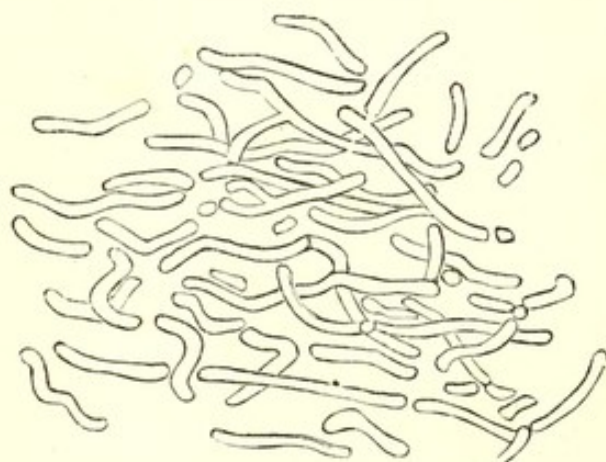


Fig. 359.—The fungus in favus. Short threads are shown.  $\times 350$ .

rows of conidia spores. The proper mycelium is not jointed, but the receptacula are. The fungus appears to begin its growth in the hair



follicles, enveloping the hair and passing into it. It also passes into the hair sheath and epidermis around, separating the cells and destroying them. It has its seat thus in the first instance beneath the horny layer which may be continuous over it. It does not invade the true skin, but the crust may cause inflammation around which may go on to supuration and ulceration. As a rule, however, it only affects the epidermic structures, the hairs being largely destroyed where the crusts are developed.

When it attacks the nail, it forms a yellow crust, the tissue of the nail being opened out and softened by the fungus penetrating between the horny cells.

2. **Tinea tonsurans** or **Ringworm**.—This is also a disease chiefly of the scalp, but the same fungus attacks the beard and the parts of the body devoid of hair. The disease is therefore divisible into three forms, **Tinea tonsurans** proper, **Tinea sycosis**, and **Tinea marginata**.

The fungus, called the **Trichophyton tonsurans**, is chiefly in the form of spores (see Fig. 112, p. 319), which infiltrate the hair and its sheath, as well as, to a less extent, the epidermis around. The mycelium is only represented by short threads. The hair shaft is greatly penetrated by the spores, which break it up.

3. **Pityriasis versicolor**.—In this disease there are patches of a yellowish or dark brown colour with a scaly surface, and occurring chiefly on the trunk in adults. If the surface be scraped and the scaly products examined under the microscope (best with the addition of liquor potassæ, both in this case and the others), there will be found epidermic cells with the elements of a fungus, the **Microsporon furfur**. There are usually abundant mycelium threads, and the spores are in rounded masses which have been compared to bunches of grapes (see Fig. 360).



Fig. 360.—Fungus in pityriasis versicolor. Threads are seen with groups of spores.  $\times 500$ . (DUHRING.)

Of the lesions due to **Animal parasites** the most important is scabies.

**Scabies**.—This disease is characterized by the formation of papules, vesicles, pustules, and other signs of inflammation due to the effects of the **Acarus scabiei** or **Sarcoptes hominis** (see p. 345). The female of this insect penetrates into the substance of the epidermis forming tunnels there.



The inflammatory conditions are the result partly of the irritation of the acarus and its embryoes, which, as they develop from the ova, begin to travel along the tunnels, and partly the effect of the scratching induced by the itching. The primary lesion is a papule on which a minute furrow is visible to the naked eye. The papule is produced by the opening up of the epidermis and swelling of the papillary layer, which is infiltrated with serous fluid and round cells. Vesicles and pustules are produced when the inflammation is more severe, and are met with chiefly in children and persons with delicate skins.

The **Acarus folliculorum** is a parasite sometimes met with in the sebaceous follicles (p. 345).

The Parasitic Insects are referred to at p. 346.

The **Larvæ of insects** are occasionally met with in the skin. The ova are deposited in the skin, and the larvæ, having developed, may traverse the skin and subcutaneous tissue for a considerable distance till they present at some part and are squeezed out or extracted. They may produce considerable inflammations in the form of boils or swellings.



## SECTION XI.

## DISEASES OF THE EYE AND EAR.

- A.—THE EYE. I. THE CONJUNCTIVA, chiefly Inflammations, catarrhal, purulent and infective (*granular conjunctivitis*). II. THE CORNEA. 1. Retrograde changes (*arcus senilis, conical cornea*); 2. Inflammations, sometimes with vascularization of cornea, *Pannus*; may lead to *Ulcers and Staphyloma*. III. THE SCLEROTIC. IV. THE IRIS. Various forms of Inflammation, serous, fibrinous, purulent. *Syphilis and Tuberculosis*. V. CILIARY BODY AND CHOROID. Inflammations. VI. LENS. Forms of Cataract. VII. OPTIC NERVE AND RETINA. 1. Optic neuritis and choked disc; atrophy of optic nerve. 2. Retinitis albuminuric, and pigmentary; 3. Embolism of retinal artery; 4. Detachment of retina. VIII. AFFECTIONS OF EYE AS A WHOLE. 1. Sympathetic ophthalmitis; 2. Glaucoma; 3. Errors of accommodation; 4. Tumours, chiefly sarcoma and glioma; 5. Parasites.
- B.—THE EAR. I. EXTERNAL EAR. 1. The auricle, chiefly hæmatoma; 2. External auditory canal; inflammations and foreign bodies; 3. Tympanic membrane. II. MIDDLE EAR. Structure. 1. Inflammations; chronic, leading to adhesions and sclerosis; acute suppurative, causing perforation of the membrana; and chronic suppurative, leading to polypi, caries and necrosis, suppurations of brain, membranes, etc.; 2. Syphilis and tuberculosis; 3. Sebaceous tumours. III. INTERNAL EAR, subject to inflammations, etc., chiefly by extension from middle ear or brain. *Ménière's disease and deaf-mutism*.

## A.—THE EYE.

THE eye, with its appendages, composed of a number of separate tissues delicately adjusted to the requirements of vision, is liable to the usual pathological changes. From the facility with which it is observed, slight differences in the appearances produced are readily appreciated and hence a considerable degree of complication exists in the nomenclature of affections of the eye.

I. Affections of the conjunctiva.—The conjunctiva is liable to the various lesions of mucous membranes, the chief being inflammation.

Conjunctivitis.—The irritant exciting to inflammation of the conjunctiva may be a very simple one, such as dust or cold, but in such cases the inflammation is usually slight and evanescent, being characterized by little more than *Hyperæmia*, with more or less serous exudation.



In most cases of conjunctivitis, however, the irritant is a morbid poison, and the disease is contracted by contagion.

In ordinary **Catarrhal conjunctivitis**, the exact nature of the irritant is not determined, but most if not all cases are contagious. The disease is characterized by hyperæmia eventuating in a mucous or muco-purulent exudation.

**Purulent conjunctivitis** (*Purulent ophthalmia*) is due to a pyogenic microbe. This is, in a large proportion of cases, the gonococcus, so that the term **Gonorrhœal ophthalmia** is nearly equivalent to purulent conjunctivitis in the adult. A purulent conjunctivitis occurring in newborn children, and hence called **Ophthalmia neonatorum**, is generally due to infection from the maternal parts during parturition. In this form microbes have been observed, in most cases the gonococcus, but there is still some doubt whether this be so in all cases.

The pyogenic virus in purulent conjunctivitis induces a violent inflammation rapidly passing on to suppuration and, if neglected, sometimes leading to necrosis and ulceration of the cornea.

**Granular conjunctivitis or Trachoma** is a more chronic inflammation of the conjunctiva which is also due to a specific virus, which Sattler asserts to be a micrococcus, but this is not absolutely determined. The disease is contagious, occurring sometimes in schools and in armies. Egyptian ophthalmia was a form observed in the French army in Egypt, and introduced by it into France and thence into this country. While due to a special virus, the disease occurs chiefly in unhealthy persons, and is hence most prevalent amongst the poor.

The lesion is in some respects like that in the specific newformations, such as tubercle or frambœsia. There is a localized formation of round-celled tissue in a delicate reticulated meshwork, so that it resembles lymphatic gland tissue. These newformations are usually defined and appear as if surrounded by a distinct capsule, in this respect also resembling lymphoid follicles. There seems little tendency towards further development of this tissue, but the follicles frequently burst and discharge on the surface. These lesions are accompanied by catarrh, and by their mere presence, especially when on the conjunctiva of the upper lids, they often produce considerable irritation of the cornea which also becomes infected by the disease. In the more chronic forms the conjunctiva may be indurated by cicatricial newformation, and this is accompanied by atrophy of the lymphoid masses. **Localized amyloid degeneration** sometimes occurs in the new-formed follicles in trachoma, this fact also tending to associate the affection with the specific newformations. The amyloid condition is found in the cells and fibres, but not in the vessels (Leber).



**Tuberculosis** is of very rare occurrence in the conjunctiva.

**Diphtheritic conjunctivitis** is a rare condition occurring along with diphtheria in the throat. There is necrosis, ulceration, and violent inflammation. Diphtheritic conjunctivitis has also been observed as an epidemic altogether apart from ordinary diphtheria, especially in certain northern lands. The virus in this case is not that of diphtheria, but produces similar effects.

II. **Affections of the cornea.**—The corneal epithelium is usually affected in the various inflammations of the conjunctiva, being exposed to the same irritations. Its peculiar structure, and especially the absence of vessels, render it somewhat liable to necrosis and ulceration.

1. **Retrograde changes in the cornea.**—A very common lesion is **Arcus senilis**, which sometimes comes on in middle life rather than in old age. It consists in a fatty degeneration of the cells and fibres of the cornea along a defined line near the periphery of the cornea, but still separated from the edge by a line of unaffected tissue.

The term **Conical cornea** or **Keratoconus** is applied to a condition in which, on account of weakness in the central parts of the cornea, the pressure from within the bulb of the eye forces out this part and so renders the cornea conical. The central part being furthest removed from the source of nutrition most readily suffers when the general condition of the body is reduced, and so yields.

2. **Inflammations of the cornea.**—The cornea, although non-vascular, is subject to inflammations even of an acute suppurative character. The inflammatory exudation passes into the canals and spaces with which the cornea abounds, and by its presence produces a clouded or opaque condition.

The cornea takes part in the acute inflammations of the conjunctiva, being liable to suppuration, softening, and sloughing. In addition, chronic inflammations begin external to, or arise in the cornea itself, some of them being due to direct irritation, and some to the presence of ulcers.

**Vascularization of the cornea** is a frequent result of chronic inflammation. It frequently occurs in the upper part in cases of granular conjunctivitis, in which case it is ascribed to the irritation of the rough conjunctiva rubbing against the cornea. The superficial layers of the cornea become the seat of newformation of blood-vessels, which frequently cover the whole surface. This condition is called **Pannus**, and it is generally recovered from when the primary affection is removed. A more profound vascularization occurs in some cases in the form of localized crescentic areas at the upper and lower borders of the cornea. This form has been ascribed to nervous lesions. Vascularization also



not infrequently attends ulcer of the cornea, a fasciculus of vessels running in from the border to the ulcer.

**Interstitial keratitis** is a name given to cases in which the cornea assumes an opaque appearance. It is met with chiefly, as Hutchinson pointed out, amongst persons the subject of **Hereditary syphilis**, in whom it is associated with a special condition of the teeth (see p. 705). The cornea has in the milder cases an appearance compared to that of ground glass. The actual histological condition of the tissue is various. There may be simply an infiltration of inflammatory products, or there may be more profound changes in the tissue, sometimes even a kind of cicatricial condition.

**Herpes** sometimes attacks the cornea along with the skin in the course of one of the branches of the fifth nerve (*Zona ophthalmica*). The eruption may lead to an ulcer or even perforation of the cornea.

**Paralysis of the fifth nerve** also leads to inflammation and ulceration of the cornea (*Neuropathic ophthalmia*). The cornea becomes cloudy and opaque and ulcers form which may perforate. These results have been variously ascribed, on the one hand to the anæsthesia, which admits of injuries to the eye and exposes it to the presence of foreign bodies without their being felt, and, on the other to a special affection of the trophic nerves, perhaps an irritation like that which occurs in herpes. If the latter be the correct view then the agent which has produced the paralysis will irritate the nerve stem or the Gasserian ganglion.

**Ulcers of the cornea** arise, as indicated in the remarks made above, under a large number of different circumstances. They may be due to injuries or to inflammations of various sorts. Ulcers heal by the new-formation of connective tissue, which fills the gap. As this has not the structure of the corneal tissue it is more or less opaque, and appears in the clear cornea as a white area. These areas, being weak, are liable if large and central, to yield before the pressure within and form a bulging outwards, a condition named **Staphyloma** of the cornea. The iris is frequently adherent to the cicatrix and is often protruded in a staphyloma.

III. **Affections of the sclerotic.**—The sclerotic is a dense connective-tissue membrane, which, on account of the paucity of vessels and nerves, and from its well-protected position, is little liable to primary disease. Inflammation extends from the conjunctiva without, but more frequently from the choroid within, and may cause softening so as to produce a staphyloma of the sclerotic.

IV. **Affections of the iris.**—The iris is a very vascular body which lies in a closed cavity occupied by the aqueous humour. The cavity is lined by a serous membrane, and the diseases are like those of serous cavities.



**Inflammation** occurs in the iris as a result of traumatic, rheumatic, or still more frequently of syphilitic irritation. The inflammation is usually acute, but its severity varies. Different forms have been described according as the exudation is serous, fibrinous, or purulent. Decemet's membrane, which covers the posterior surface of the cornea, takes part in the inflammation. **Serous iritis** is the mildest form, and leads to distension of the anterior chamber by the accumulated fluid. This fluid contains leucocytes, which occasionally render it somewhat turbid. **Fibrinous iritis** (also called *Plastic iritis*) is more severe and is accompanied by a fibrinous exudation, which appears most commonly at the edge of the pupil, but also presents itself on the posterior surface, and sometimes on the anterior. The exudation fixes the iris by forming a membrane in the pupil, or more commonly by gluing it to the lens. This adhesion to the lens (called *Synechia*) is a serious part of the lesion, as it is apt to become permanent. As the inflammation progresses the fibrinous exudation becomes replaced by the formative cells of granulation tissue, and new-formed blood-vessels pass into it. The regular development into spindle-celled tissue and connective tissue occurs, and, just as in the case of the pleura, permanent adhesions are the consequence. Sometimes the new-formed membrane fills the pupil and obscures the vision. The iris itself is the seat of newformation of connective tissue, which may cause considerable atrophy of its proper tissue. On the other hand, hypertrophy of the iris sometimes results. **Purulent iritis** is more severe. It causes an opening out and softening of the tissue of the iris, while the anterior chamber is filled with pus.

Inflammations of the iris are liable to extend to the ciliary body and choroid (*Irido-choroiditis*).

**Syphilis** has already been mentioned as a cause of iritis. It is so in the secondary stage, producing an acute inflammation usually of the fibrinous kind. A formation of gummata is an exceedingly rare occurrence.

**Tuberculosis** of the iris is also rare, but it is stated that the condition called Granuloma is really a tuberculosis. There is inflammation with development of tubercles.

**V. Affections of the ciliary body and choroid.**—The ciliary body is highly vascular, and is engaged in the secretion of the humours of the eye. Being very active it is peculiarly liable to acute inflammations, more especially as it is related anatomically both to iris and choroid, and is apt to partake in the inflammations of either.

The choroid is also a vascular structure, containing a layer of pigment cells on its inner or retinal surface. Inflammation of it is not usually acute, except in sympathetic ophthalmia (see further on), and, when



chronic, it causes disturbances of the pigment, which produce alterations in the ophthalmoscopic appearances. Sometimes a formation of true bone occurs in the choroid in chronic inflammation.

**VI. Affections of the lens.**—The only affection of the lens requiring special notice is **Cataract**. This consists in an opacity of the lens or its capsule, such as to interfere with vision. It sometimes occurs as a congenital affection, the lens being imperfectly formed. It is most frequently a senile change, but it also occurs as a result of injury and is an occasional consequence of diabetes.

At birth the lens is soft throughout, but as years go on the centre becomes dense, so as to form a hard nucleus while the periphery remains soft. The hard nucleus is not liable to change, but the soft cortex may undergo alterations. These consist in a formation of fat amongst the fibres which constitute the lens, and the loosening and separation of these fibres. The process goes on till sometimes the affected part of the lens consists of a detritus in which are fat, cholestearine and remains of fibres. The dense nucleus may sink down in this softened substance. In senile cataract there is always the dense nucleus, which may be of considerable bulk.

In **Capsular cataract** the epithelial cells, which line the capsule, undergo changes, which may be spontaneous or the result of inflammation. They multiply and cause opacity by their abundance and by retrograde changes to which they are liable.

**VII. Affections of the optic nerve and retina.**—These parts form really a prolongation outwards of the cerebrum, and are surrounded by processes from the membranes of the brain. The dura mater forms a sheath around the nerve as it passes to the eyeball. This sheath is composed of dense connective tissue, and is continuous with the sclerotic, which is composed of similar tissue. Within this sheath the arachnoid is continued as a loose covering as far as the eyeball, where also it becomes continuous with the sclerotic. The pia mater closely invests the nerve and sends processes amongst its fibres. Between the external sheath or dura mater and the optic nerve there are lymphatic spaces continuous with those of the soft membranes. The nerve itself is composed of medullated nerve fibres running in bundles and enclosed in connective tissue. On reaching the eyeball the nerve bundles pierce the sclerotic, which here forms a perforated membrane (*Lamina cribrosa*). At the same time they lose their medullary sheath and are distributed as non-medullated fibres inside the globe. From the optic nerve entrance, which forms a round disc, the fibres radiate outwards around the eyeball where they are in contact with the vitreous. They are very transparent, as the rays of light require to penetrate them in order to reach the rods



and cones which are on their outer surface. The nerve fibres, with the rods and cones and the granular layers constitute the retina.

1. **Œdema and Inflammation of the optic nerve** (*Optic neuritis, Choked disc*).—A series of lesions is included here, concerning whose pathology some questions still remain unanswered. Affections of the optic nerve are frequent in consequence of tumours of the brain, and these have partly the characters of a simple œdema and partly those of inflammation. Tumours of the brain produce these changes whatever may be their seat. Inflammations of the membranes affect the optic nerve less frequently. There are also inflammations of the nerve of obscure origin.

Tumours of the brain, and other lesions which lead to increased intracranial pressure, produce œdema of the sheath of the nerve with interstitial inflammation. The œdema of the sheath is accompanied by swelling of the intraocular termination of the nerve from venous hyperæmia and œdema. It is this œdema and engorgement which is called Choked disc. It is difficult to explain why tumours of the brain should produce this effect, but it seems likely that by increasing the intracranial pressure they obstruct the passage of the lymphatic fluid in the spaces around the nerve. The retained fluid is perhaps specially irritating, and induces a chronic interstitial inflammation.

Whether originating in tumour of the brain or not, an interstitial inflammation of the optic nerve leads to sclerosis with atrophy of the proper nervous elements.

**Atrophy of the optic nerve** arises in the manner just mentioned in cases of neuritis. It may arise also in Locomotor ataxia, by a similar process to that occurring in the cord in that disease, namely, a sclerosis characterized by increase of the connective tissue and atrophy of the nerve fibres. These changes are seen by the aid of the ophthalmoscope chiefly in the form of an abnormal whiteness of the optic disc.

2. **Inflammation of the retina** (*Retinitis*).—This affection occurs in various forms, the most familiar of which is **Albuminuric retinitis**. This affection consists of a chronic inflammation, apparently commencing in the arteries and accompanied by hæmorrhages and degenerations of the retinal and choroid tissue. The affection of the arteries consists in an inflammatory thickening or sclerosis of their wall, with diminution in their calibre. The hæmorrhages are mostly in the superficial fibrous layer of the retina. Pale white spots and streaks in the region of the macula are very characteristic of the condition, and these arise chiefly by fatty degeneration. The hæmorrhagic spots may become white, but many spots are white from the first and not hæmorrhagic.

The affection of the retina occurs in cases of chronic Bright's disease,



and its occurrence lends weight to the view that this disease is a general one affecting the blood-vessels throughout the body, and not specially those of the kidney. The argument is somewhat invalidated, however, by the circumstance that diabetes, lead-poisoning, etc., sometimes induce a retinitis essentially similar to the albuminuric form. It would seem that a grave alteration of the blood is the cause of the affection of the retina.

**Pigmentary retinitis** (*Retinitis pigmentosa*) is a form in which the choroid is involved along with the retina. It is a chronic inflammation or sclerosis, and the pigment cells seem to undergo local enlargement and increase so as to project into the retina and form dark patches. These patches follow the lines of the vessels.

3. **Embolism of the retinal artery.**—The central artery of the retina enters the eyeball in the middle of the optic nerve and then divides into branches which radiate from this centre. The artery is a branch of the anterior cerebral, and it and its branches are end-arteries. Embolism generally occurs in consequence of valvular disease of the heart. There is sudden blindness, which is usually permanent to the extent of the distribution of the occluded artery. The retina in the area affected undergoes necrosis and softening, this process being sometimes accompanied by hæmorrhage (see p. 77). Ultimately the optic nerve becomes atrophied.

Thrombosis of the retinal artery or vein is also known to occur.

4. **Detachment of the retina.**—In this condition the retina is separated over a larger or smaller area from the choroid, and the separation is usually permanent. It may be caused by shrinking of the vitreous and by effusion of fluid between the choroid and the retina. It is also produced by injuries and by tumours, but it may occur without assignable cause.

VIII. **Affections of the eye as a whole.**—Some of the diseases already mentioned affect the eyeball generally, and there are some affections which do so primarily.

1. **Sympathetic ophthalmitis.**—In wounds of the eye there is occasionally an extension of the inflammation to the opposite eye, and this is commonly spoken of as by sympathy. Inflammation, especially in the severe forms which sometimes occur, can scarcely owe its origin to nervous influences alone, and this conclusion is confirmed by the fact that the inflammation may progress after the eye primarily affected has been removed. Mackenzie first suggested that the inflammation travelled up the optic nerve to the chiasma and down the other nerve to the opposite eyeball. This view has been recently amplified and it has been asserted that microbes propagate up the sheath of one optic nerve and down that of the other so as to produce the inflammation.



This would explain how the affection of the second eye always occurs considerably after that of the first, and how removal of that primarily affected does not always stop the disease. This explanation seems generally accepted for cases of severe inflammation, but slighter inflammations and functional irritations of the eye may be produced reflexly.

The inflammation begins in the iris and ciliary body, and has generally the characters of an iritis and cyclitis. Predisposition or susceptibility seems to have considerable influence in determining the occurrence of sympathetic inflammation.

2. **Glaucoma.**—This name is given to a state characterized by increased tension of the eyeball which is over-filled and distended. The structures forming the wall of the eye suffer in consequence. The condition arises as a primary affection, but the exact causation is not well understood. There seems to be a derangement of the circulation causing an increased secretion of fluid or a diminished elimination from obstruction. Increased tension occurs also secondarily as a result of any cause which increases the contents of the eyeball, such as hæmorrhage, dislocation of the lens, sarcoma of the choroid, etc.

The increased tension within the eyeball acts on the structures of the globe. The function of the retina is interfered with, and it afterwards undergoes atrophy; the optic nerve entrance is hollowed out, the lamina cribrosa being more yielding than the sclerotic; the cornea is rendered dull and obscure and insensible.

3. **Errors of accommodation.**—Vision is brought about by the images of objects being focussed on the retina. The focussing is effected by the refracting media of the eye, chiefly the lens and the cornea. In order to accurate vision the image must fall in perfect focus on the retina. Variations in the focussing required by the greater or lesser distance of objects is brought about by the ciliary muscles acting on the lens. As life goes on the activity of these muscles and the elasticity of the lens lessen, and the power of rapid accommodation diminishes. This condition is called **Presbyopia**. The defective power of accommodation is supplied by convex glasses. Other defects of vision occur when the image falls behind or in front of the retina, or is distorted.

In **Hypermetropia** the image falls behind the retina. This is usually due to abnormal shortness of the eyeball, but may also result from the refractive structures being deficient. Vision is good for distant objects but defective for near. In **Myopia** the image falls in front of the retina. The axis of the globe is usually too long for the refractive structures (axial myopia), but sometimes the refractive structures are too powerful (refractive myopia). Vision is defective for long distances and good at a short distance. In **Astigmatism** the image is distorted, from the



refractive media, chiefly the cornea, being different in convexity at different meridians.

These various defects may lead to irritations and strainings of the eye from the attempts to bring the image into focus.

4. **Tumours of the eyeball.**—There are two common forms of tumour, sarcoma and glioma.

**Sarcoma** springs in the majority of cases from the choroid, but may take origin in the ciliary body or iris. It is usually a spindle-celled tumour and in the large majority of cases is pigmented. The unpigmented forms are usually round-celled. The sarcoma in its growth usually detaches the retina and encroaches on the vitreous humour. The eyeball is occupied by the tumour mass, which generally in course of time also extends into and through the sclerotic. The sarcoma is prone to generalization, and tumours, frequently pigmented, spring up in various organs.

**Glioma** takes origin in the retina. It consists of round cells in a fine reticulum. The cells may be so abundant as to make the tumour approach to the structure of the sarcoma. Two forms are described under the designations exophytum and endophytum, according as the tumour originates in the external or internal granular layer of the retina. The tumour retains the characters of the layer from which it has originated. The tumour grows for a time in the retina, gradually enlarging and filling the globe. After a time it may involve the coats of the eyeball and extend through the sclerotic. Occasionally it extends by metastasis to distant organs. It is characterized by a bright yellow reflection seen through the pupil.

**Epithelioma** sometimes occurs in the eyeball, usually originating at the junction of cornea and conjunctiva. It may also originate in the conjunctiva. Cancer sometimes takes origin in the lachrymal glands.

The conjunctiva is also liable to simple growths, the commonest being the papilloma, whose most frequent seat is the junction of the cornea and conjunctiva.

5. **Parasites of the Eye.**—**Cysticercus cellulosæ** (the scolex of *tænia solium*) is not very uncommon in the eye. Its usual seat is beneath the retina, which may become detached by its growth. Sometimes the parasite perforates the retina and becomes free in the vitreous where it may be seen with the ophthalmoscope as a clear cyst, from which the head of the worm is sometimes protruded. A less frequent seat is the anterior chamber, where it may be free or attached to the iris. It is capable of protruding and withdrawing its head. The cysticercus also occurs in the orbit outside the eyeball.

The *Echinococcus* occasionally has its seat in the orbit.



## B.—THE EAR.

The ear has intimate anatomical relations with various other parts, and in its diseases it is frequently associated with these parts. It is continuous externally with the skin; it is in communication by means of the Eustachian tube with the mucous membrane of the nasopharynx; it is intimately related to the bones of the skull and to the dura mater and brain. For purposes of description the diseases of the ear are divided into those of the external, middle, and internal ears, but these parts being closely related in position are frequently affected by disease together.

**I. Affections of the external ear.**—The external ear consists of the auricle, external auditory canal, and tympanic membrane.

1. **The Auricle** is liable to various **congenital malformations**, ranging from almost entire absence to simple alterations in the usual outline. Supernumerary organs, generally rudimentary, are also met with. Inflammations of the skin, such as erysipelas and inflammatory skin eruptions, especially eczema, may extend to the auricle.

A more independent affection is the **Hæmatoma of the auricle** or **Othæmatoma**. This consists in a hæmorrhage between the cartilage of the auricle and the perichondrium, the latter being raised, but generally carrying with it pieces of cartilage. It usually arises from injury, but in some cases the injury is slight or absent, and the condition is spontaneous. This condition is so frequent in insane persons, more particularly in dementia paralytica, that the name **Insane ear** is sometimes given. The concavities of the auricle are filled out with the effused blood, which generally remains fluid for a considerable time. With the absorption of the blood there is generally marked shrinking and cicatricial contortion of the auricle. Sometimes injuries induce a serous exudation instead of blood, the condition otherwise resembling hæmatoma. Small **Fibromas** sometimes form in the auricle in consequence of the irritation of earrings. **Gouty topi** sometimes occur in the cartilage of the auricle.

2. **The External auditory canal.**—Congenital contraction and closure of the canal have been observed in various degrees.

**Inflammations** are similar to those of the skin, and may be either in the form of boils, which are somewhat frequent, or of a diffuse inflammation. The former have the characters of cutaneous boils, with which they may be associated; the latter result chiefly from injuries but also from the acute exanthemata, chiefly small-pox, from erysipelas, and from ordinary skin eruptions. The disease may extend to the osseous wall of the canal and lead to caries or necrosis, but these are more frequently



the consequences of affections of the middle ear. In more chronic cases we have the usual newformation of tissue of chronic inflammations, resulting in thickenings of the cutaneous lining of the canal, and also in some cases in hyperostosis, which may greatly contract the canal.

**Exostoses** not infrequently spring from the osseous wall.

**Accumulations of cerumen** very frequently occur to such an extent as to occlude the canal and cause deafness. They are usually dry and are composed of cerumen mixed with epidermis, hair, dust, etc. The accumulation is due either to excessive secretion or to some impediment to the escape of the wax.

**Foreign bodies** and **Parasites** are met with in the meatus. The former consist of a great variety of materials introduced from without and met with chiefly in children. Insects and their larvæ are occasionally found in the ear, the latter chiefly in cases where purulent discharge exists. The *acarus folliculorum* also occurs. **Fungi** belonging to the family of the *Aspergillus* (see p. 318) have been frequently observed growing in the meatus, chiefly in its inner recesses and on the tympanic membrane. They cause obstruction of the canal and may give rise to serious inflammations.

3. **The Tympanic membrane** is liable to **Inflammations** in common with the meatus on the one hand and the middle ear on the other. Hence arise various thickenings and adhesions, sometimes with calcareous deposition, chiefly in connection with diseases of the middle ear. Hence also arise **Perforations** of the membrane, which have frequently serious consequences for the tympanic cavity. **Rupture** is of occasional occurrence, usually in consequence of condensation of the air in the external meatus as from a blow on the ear, but also from instruments thrust into the ear.

II. **Affections of the middle ear.**—The middle ear is a cavity containing air, of a very complicated outline (see Fig. 361). The central part is that immediately within the tympanic membrane, forming the tympanic cavity which is traversed by the chain of ossicula. From the lower and anterior part of this cavity passes the Eustachian tube down to the naso-pharynx. The upper and posterior part of the cavity communicates with the mastoid cells. The largest of these lies immediately behind the tympanic cavity, of which it is almost a continuation backwards. This is the antrum mastoideum whose roof, along with that of the tympanic cavity, is formed by a thin layer of bone, which alone separates these cavities from the dura mater lining the middle fossa of the skull. The remaining mastoid cells communicate directly or indirectly with the antrum. In childhood the antrum is well developed, but the other cells are very imperfect, the mastoid process being small.



The roofs of the tympanum and antrum are not infrequently deficient, especially in childhood, so that there are gaps in the osseous wall. The mucous membrane of the middle ear is, in most parts, very delicate, and is covered with a single layer of epithelium. In its structure and in the

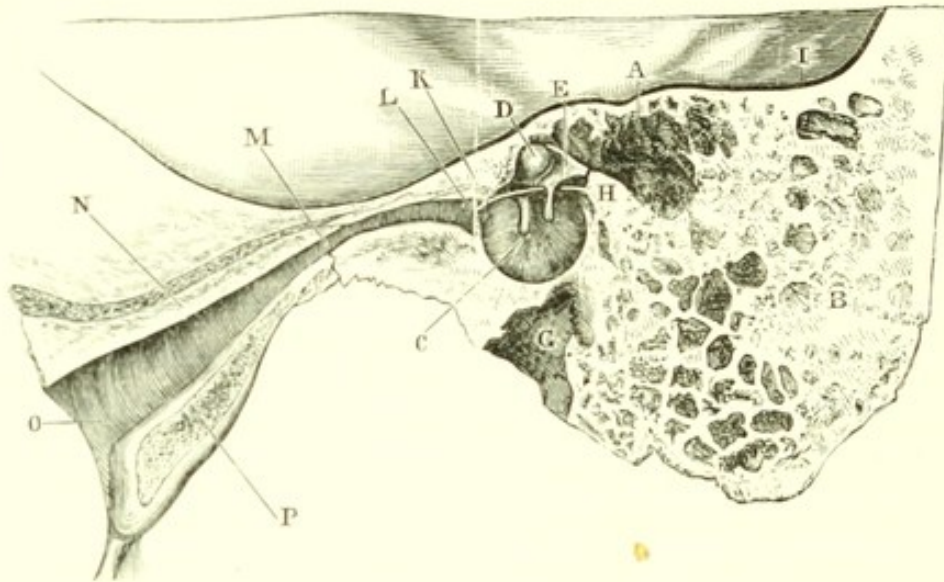


Fig. 361.—Section through middle ear. A, antrum mastoideum; B, mastoid cells; C, inner surface of tympanic membrane; G, fossa for jugular vein; H, partition between tympanum and antrum; I, dura mater; L, tympanic aperture of Eustachian tube; O, pharyngeal mouth of same. (BARR.)

character of its epithelium it resembles a serous membrane. It is particularly thin over the ossicula and the membrana tympani and in the mastoid cells. In the greater part of its extent the mucous membrane covers bone, and is inseparably united with the periosteum.

1. **Inflammations of the middle ear.**—These have in their elements the characters of inflammations elsewhere but modified by the relation of parts. **Catarrh** is somewhat common, and may be acute or more chronic. The exudation varies according to the severity of the inflammation, being mucous or muco-purulent. The catarrh may be localized mainly in the Eustachian tube, which may be obstructed by the swelling of the mucous membrane. The exudation may distend the tympanic cavity and cause bulging of the membrane. It will also interfere with the movements of the ossicula.

**Chronic inflammations** also occur which have little of the catarrhal character but rather that of the chronic productive inflammation. These conditions are sometimes described as **Chronic dry or adhesive catarrh**. These are associated with **Adhesions** and **Sclerosis** of the mucous surfaces. The inflamed surfaces are thickened, and wherever they come in contact they tend to coalesce. In this way fibrous adhesions are formed which may unite the membrana tympani or the ossicula to the opposite wall of the cavity. Like other connective tissue



formed in inflammations this has a tendency to shrink and become dense, a process expressed in the term sclerosis. The pliant mucous membrane thus becomes rigid, and stiffens the *membrana tympani*, the ossicula, and the membrane of the fenestra rotunda. There is in some cases a newformation of bone, so that thickenings and projections of the osseous wall of the cavity occur. There is not infrequently in the dense shrunken tissue a deposition of lime salts, a **Calcification** similar to that so frequently seen in chronic endocarditis. All these changes, by rendering rigid the structures concerned in propagating the sonorous vibrations, contribute to the causation of deafness.

**Acute suppurative inflammation** of the middle ear is a frequent result of scarlet fever, and more rarely of the other acute fevers. In these cases it arises by propagation upwards of an acute phlegmonous inflammation from the pharynx. In some cases the affection is ascribed to cold and ordinary catarrh of the nares or naso-pharynx, and in these the nature of the irritant is obscure. There is an acute inflammation of the mucous membrane going on to suppuration and always resulting in **Perforation of the membrana tympani**. The middle ear is, in fact, converted into a kind of abscess cavity and the pus finds its way out through the membrane. The inflammation may extend partly to the external and internal ears. The perforated membrane may afterwards heal, but in many cases there is considerable destruction of tissue, and when the inflammation subsides a permanent gap is left in the membrane, sometimes involving its entire extent.

**Chronic suppurative inflammation** of the middle ear, or **Otorrhœa**, is a common result of the acute disease. The perforation of the *membrana tympani*, brought about in the way mentioned above, has far-reaching consequences. The mucous membrane of the middle ear is exposed to the varied influences from without. It is liable to catarrhs, and, as the resulting exudation is exposed to the air, septic changes readily occur in it. The consequence of these septic changes is a suppurative inflammation, which, when once fully established, is likely to be permanent. The septic discharges lingering in all the complicated parts of the middle ear keep up the inflammation. The discharges are, consequently, in many cases exceedingly putrid. The mucous membrane is by degrees converted into a succulent granulation tissue from whose vessels the pus is exuded. Various important consequences are liable to occur in this condition, such as polypi, caries, and abscess of the brain.

**Polypi** are outgrowths of vascular tissue from the inflamed mucous membrane (see Fig. 362). They protrude through the perforation in the membrane, into the meatus, and may even present at the external orifice. They are usually single mucous polypi, consisting of a very



cellular connective tissue, in which mucous glands may be present, sometimes forming cysts. They are more rarely fibrous in structure, and in that case contain no glands. A polypoid myxoma is an exceedingly rare form.

**Caries and Necrosis** of the underlying bone not infrequently occur. There may be a gradually advancing inflammation of the bone, leading to erosion by a process of rarefying osteitis. Thus a considerable erosion of

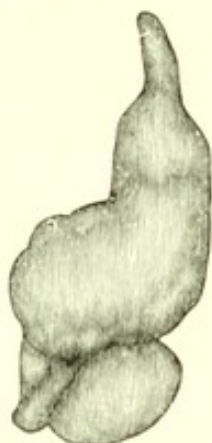


Fig. 362.—Large mucous polypus, which dilated the external auditory canal and protruded from external orifice. (BARR.)

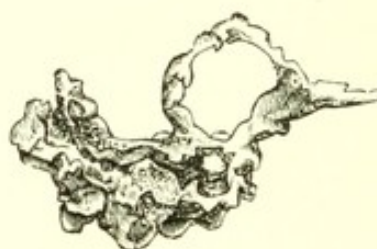


Fig. 363.—Sequesterum consisting of annulus tympanicus and part of mastoid process, removed from a child. (BARR.)

the temporal bone may ensue. In other cases necrosis of considerable portions of the temporal bone results, the dead bone being perhaps discharged (see Fig. 363). These processes are most liable to occur in the mastoid cells, where the discharges readily lie and decompose. In these



Fig. 364.—Section showing effects of prolonged otorrhoea. *a*, enlarged antrum mastoideum. *b*, mastoid cells replaced by dense bone like ivory. *c*, tympanic cavity. The patient (a man) died of purulent meningitis. (BARR.)

parts the inflammation may, after a time, become acute, so that something in the nature of an abscess may form and come to the surface of the mastoid process.

Sometimes the supuration extends outside rather than inside the mastoid process, so that there is a suppurative periostitis over this process, but without necrosis or caries.

In some cases there may be a **Formative** osteitis, so that newformation of bone occurs,



leading to thickenings, and sometimes to condensation of the cancellated mastoid structures (see Fig. 364). This may even occur coincidentally with caries and necrosis, a rarefying and a condensing otitis co-existing here as in other cases.

If **healing** occurs in this disease there are liable to be the consequences of chronic inflammation already referred to. Healing will only occur if septic decomposition is got rid of. Unless the tympanic membrane closes, which it sometimes does by cicatricial newformation, there is apt, at intervals, to be a renewal of the septic process and of the suppuration.

From the nearness of the structures concerned to the dura mater and brain, there is occasionally an **Extension of the septic process** to the parts within the skull. These consequences have already been referred to under Abscess of the Brain (see p. 566). They are chiefly **Septic thrombophlebitis**, which may result in pyæmia, **Suppurative meningitis**, which may be associated with thrombophlebitis, and **Abscess of the brain**, which may arise by thrombophlebitis but also occurs without this. Paralysis of the facial nerve occasionally results from extension to the Fallopian canal.

2. **Syphilis and Tuberculosis** of the middle ear. Syphilis is stated to cause in some cases a chronic inflammation with sclerosis as already described. Tuberculosis has been suspected in some cases of chronic otorrhœa, and the tubercular bacillus is said to have been observed in the discharge.

3. **Sebaceous tumours of the ear, Cholesteatoma.**—Under these names are included two very different conditions, one of them a true tumour such as occurs in the brain (see p. 583). This is excessively rare and, most cases of so-called sebaceous tumour, are due to nothing more than accumulated and inspissated products of inflammation, namely pus, epithelium and debris. These collections may attain a considerable size and extend into the auditory canal. There may be a deposition of crystals of cholestearine giving a glistening appearance to the mass.

III. **Affections of the internal ear.**—The internal ear, consisting of the labyrinth and the auditory nerve, is rarely the primary seat of disease, but is liable to be involved in affections of the middle ear, on the one hand, and of the brain and its membranes on the other. It does, however, sometimes suffer directly from the effects of continuous loud noises (as in boiler makers) or from blows on the head. These lead to chronic inflammations of the structures.

**Inflammations** of the middle ear by affecting the bone directly may lead to destruction, or in a less degree to thickenings, and atrophies of the labyrinthine structures. Sometimes the base of the stapes, by



pressing too firmly on the fenestra ovalis, causes increased pressure in the labyrinthine fluid and consequent atrophy of the nerve. The abnormal position of the stapes may be the result of inflammation or it may be produced by obstruction of the Eustachian tube.

**Syphilis** not infrequently attacks the labyrinth leading to thickenings and consequent injury to the nerve structures. It is said, also, that hæmorrhage and serous exudation sometimes result from syphilis. **Congenital syphilis** may attack the labyrinth, so that deafness often coincides with keratitis and iritis.

**Meningitis** (cerebro-spinal and perhaps tubercular), by extending to the acoustic nerve or labyrinth, may lead in children to such damage to the structures as to result in total deafness. This is stated to be an important cause of deaf-mutism. **Acute fevers**, especially scarlet fever and typhoid fever, sometimes cause inflammation of the labyrinth. **Mumps** has occasionally a similar effect.

**Diseases of the brain** and its membranes, such as tumours, abscesses, and inflammations, often involve the auditory nerve within the skull. The auditory centre in the brain may also be involved by local diseases such as embolism and hæmorrhage.

Affections involving the labyrinth and producing alterations in the fluid contents of the semicircular canals are apt to produce subjective sensations of giddiness and sickness. This is due to the fact that these structures apparently exercise an important function in connection with the retention of the equilibrium of the body. The group of symptoms thus produced is included under the designation **Ménière's disease**. A minor degree of giddiness and sickness may be produced by syringing the ear with cold water, especially when the membrane is perforated.

Any interference with the labyrinth leads to deafness, and this may be absolute. If this occurs in children the condition of **Deaf-mutism** results. Deaf-mutism will occur when deafness has been produced in the infant, from whatever cause, whether an affection of the middle or internal ear.



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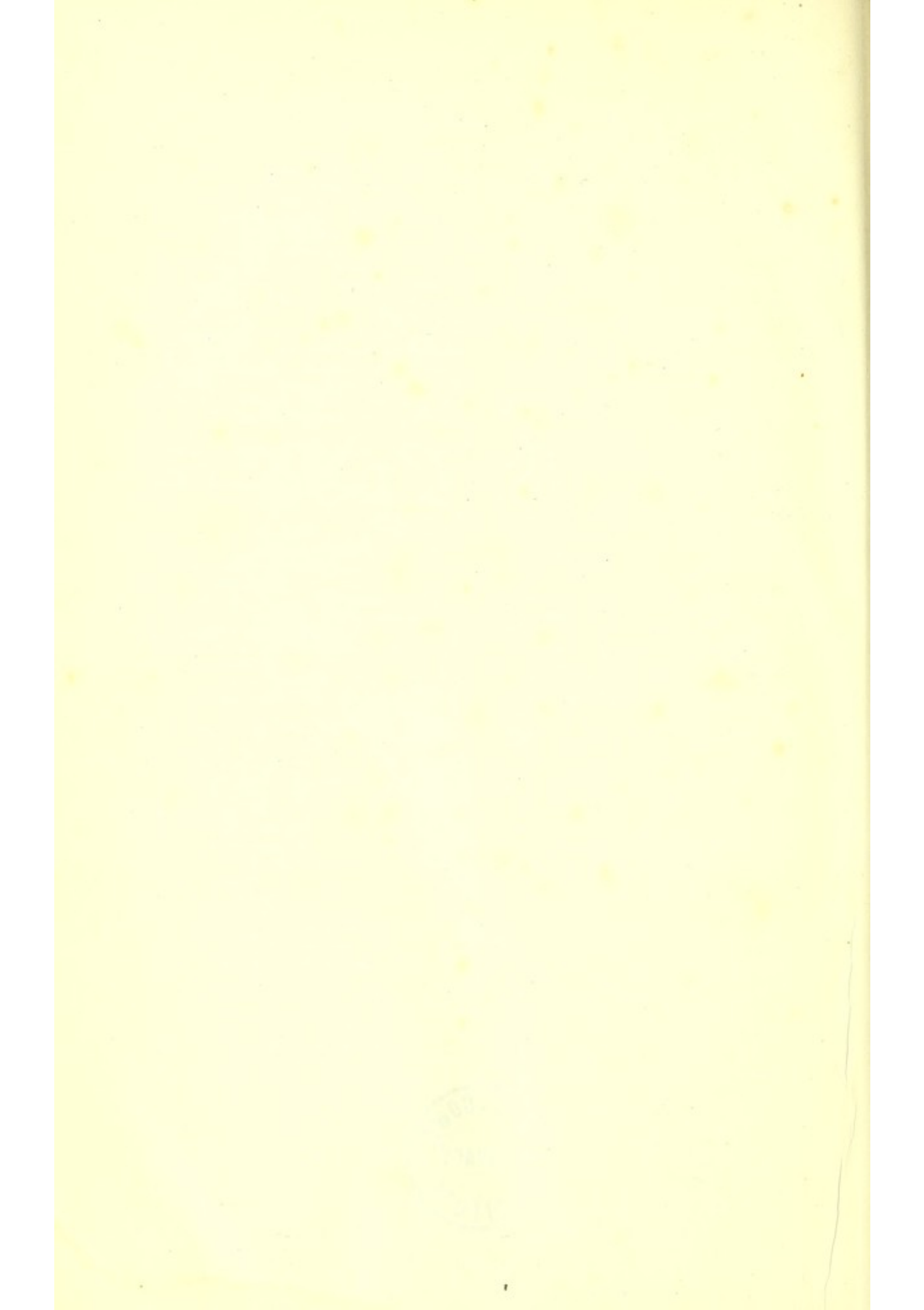
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