# Researches on the molecular asymmetry of natural organic products / by Louis Pasteur (1860).

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### RESEARCHES

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# MOLECULAR ASYMMETRY.

LOUIS PASTEUR. (1860).

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## RESEARCHES

ON THE

# MOLECULAR ASYMMETRY

OF

# NATURAL ORGANIC PRODUCTS

BY

### LOUIS PASTEUR,

Membre de la Société Chimique de Paris.

(1860).

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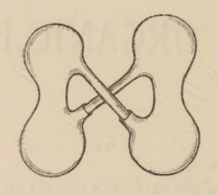
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### PREFACE.

THE two Lectures which compose this volume were published by the Chemical Society of Paris in a volume entitled "Leçons de Chimie professées en 1860" (Paris, 1861). They give in brief form an account of Pasteur's brilliant investigations on optically active compounds, and are remarkable for three things:—The fact that they describe the only ways even now known of separating optical isomers: that they contain scarcely a statement which would be changed if they were to be written to-day: and that in the theoretical speculations as to the cause of the difference between optical isomers so close an approach is made to the theory of van't Hoff and Le Bel (1874), that it seems incredible that many years elapsed before the final step was taken.

Two short paragraphs addressed especially to the Chemical Society of Paris and to its president, Dumas, have been omitted from the introduction.

The thanks of the Club are due to the Council of the Society for permission to publish this translation.

# INTRODUCTORY.

The following Lectures were given at the invitation of the Council of the Société Chimique of Paris. researches of which they form a short summary have occupied my time during ten consecutive years. thought has often occurred to me to bring together my papers, and, reviewing the details with care and with broader survey, to give co-ordination to the collection of separate memoirs. But in the life of every man who devotes himself to experimental science there comes a stage when the value of time is inestimable; a period of activity when the spirit of discovery is paramount, and when every year should chronicle some advance. To rest on our oars with what is already acquired is annoying and even dangerous, and the harm more than neutralises the pleasure or even the utility of seeing our ideas diffused abroad to the extent of our wishes.

The reluctance which I have felt to undertake the wearisome task of collecting and reviewing my researches on the molecular asymmetry of organic products occurring in nature, has only been equalled by the eagerness with which I acceded to the request of several members of the Société Chimique, that I should publish the two Lectures in which the principal results of my work have been explained. Few memoirs have had a better recep-

tion than mine, as they have appeared one after the other, and yet I have had many proofs that they are little known.

On this account, I trust that the present publication will be of some use. Besides, if it were only for the interest which the Lectures excited in the select assembly before which they were given, I should feel sufficiently repaid for my pains.

\* \* \* \*

L. PASTEUR.

Paris, 1860.

### FIRST LECTURE.

I.

A T the end of the year 1808, Malus announced the discovery that the light reflected by all objects, whether opaque or transparent, acquired new properties of a very extraordinary character, which distinguished it essentially from the light sent to us directly from luminous bodies.

MALUS gave the name polarisation to the change which the light underwent in the process of reflection. Later, the expression plane of polarisation was used to designate the plane of reflection, that is to say, the plane containing the incident ray and the normal to the reflecting surface.

Malus' discoveries in connection with polarised light were not limited to this. It had long been known that a ray of direct light was always separated into two beams of white light, of equal intensity, in its passage through a rhomboid of calcium carbonate. Thus the flame of a taper viewed through such a rhomboid is always double, and the two images are equally bright.

HUYGENS and NEWTON had already noticed that light, after its passage through a crystal of Iceland spar, no longer behaves like direct light. Thus, when we study one or other of the images of the taper, just mentioned, through a second rhomboid, we observe (1) that there is not always bifurcation of the ray; (2) that when bifurcation does take place, the two new images have not the same intensity. So that the light which has traversed a doubly refracting crystal is different from

ordinary or direct light. Assuming this, Malus showed that the change produced in light by double refraction was identical with that arising during reflection from the surfaces of opaque or transparent bodies; in other words, that the two rays, ordinary and extraordinary, given by doubly refracting crystals, were polarised rays.

From the first Malus established these fertile discoveries so clearly, using so much care and precision, both as regards fact and expression, that one would think in reading them that his memoirs had been written yesterday. But he was prevented from continuing his work, having been cut off prematurely in 1812, at the age of 37. Happily for science, two celebrated physicists, Biot and Arago, at that time young and full of activity, took up the legacy he had bequeathed, and speedily distinguished themselves by brilliant discoveries in the new path opened to science by Malus.

In 1811, Arago noticed that, when a ray of polarised light is analysed by a rhomboid of Iceland spar, after it has traversed, normally to the surface, a section of rock crystal cut perpendicularly to the axis, it always gives two images in all positions of the rhomboid; and, further, that these are coloured in complementary tints. When the thickness of the spar does not suffice for complete separation of the rays the image is white where they are

superposed.

This experiment showed a two-fold anomaly in regard to the ordinary laws of doubly refracting crystals. All other uniaxial crystals, cut normally to the axis, would have given two white, instead of two coloured images, and in two positions of the analysing rhomboid, separated by an angle of 90°, the two images would have been reduced to one.

ARAGO's conclusion was that the results of the above experiment are precisely those which should follow if it

be assumed that the two differently coloured rays from the incident beam of white light are polarised in different planes on leaving the plate of quartz.

ARAGO did not continue the study of these striking phenomena, and BIOT, from 1813 onward, enunciated the physical laws governing them, carefully isolating them from all those with which they seemed to have been confused by ARAGO.

BIOT produced the polarised ray with light from each of the parts of the simple spectrum separately, and found that the original plane of polarisation was deviated through an angle proportional to the thickness of the plate; that this angle was different for each of the primary colours and increased with the refrangibility according to a fixed law. He also made the exceedingly curious observation that of plates obtained from different crystals of quartz, some deviated the plane of polarisation to the right and others to the left, in each case according to the same laws.

But Biot's most remarkable discovery in connection with this kind of phenomena is undoubtedly that of the deviation of the plane of polarisation produced by a number of natural organic products, such as oil of turpentine, solutions of sugar, of camphor, and of tartaric acid. The first mention of this fact occurs in the bulletin of the Société Philomatique for December 1815.

To understand this Lecture it is essential particularly to recollect the existence of this rotative property in tartaric acid, and its absence in paratartaric or racemic acid, an acid isomeric with tartaric acid.

There exist, therefore, organic products which, when themselves liquid, or when dissolved in water, possess the rotative property, and resemble in this respect solid crystallised quartz. But it must be noted here that the analogy with quartz is superficial only. The deviation of the plane of polarisation is common to both, but the character of the phenomenon is quite different.

Thus quartz deviates; but it must be crystallised. Dissolved, or solid and not crystallised, it lacks the property. Not only must it be crystallised, but it must be cut in plates perpendicular to the axis. As soon as the plate is slightly inclined to the direction of the ray, the effect is diminished and finally disappears.

Sugar deviates (and what is said of sugar is true of all the other organic products), but the sugar must be dissolved, or solid and amorphous as in barley-sugar. In the crystallised state it was impossible to discover

any effect.

The tube containing the solution of sugar may be inclined. The deviation does not alter for equal thicknesses. And, what is more, active agitation of the liquid by means of clock-work leaves the phenomenon the same.

So that, from the first, BIOT quite definitely concluded that the action produced by the organic bodies was a molecular one, peculiar to their ultimate particles and depending on their individual constitution. In quartz the phenomenon is a consequence of the mode of aggregation of the crystalline particles.

These are the physical precedents, if I may so express myself, for the researches which I am about to lay before you. We must next consider their mineralogical precedents.

### II.

Hemihedry is certainly the one of the peculiarities of crystallisation which is most easily grasped in its external manifestation. Consider, for example, a mineral species crystallising in the cubic form. This species can show, as everyone knows, various kinds of forms determined by

the law of symmetry, a law so natural that it is, so to speak, a physical axiom. This law states that, if one kind of form is given, all the others compatible with it may be obtained by an artifice which consists in modifying, truncating as Romé DE LISLE says, identical parts, simultaneously and in the same manner. Identical edges are such as are formed by the intersection of faces which are severally identical and which cut at the same angle; and identical solid angles are those which are formed by dihedral angles which are identical and similarly placed. Thus in the cube there is only one kind of solid angle and one kind of edge. If one of the angles is truncated by a face equally inclined to the three faces of the solid angle, the other seven angles must be simultaneously truncated by a face of the same nature. This is what we observe in alum, in galena, and, generally, in all cubical species.

Let us imagine a right prism with rhombic base. The eight basal edges are identical edges. If one is truncated, the other seven must be truncated, and in the same manner. The four vertical edges are of another kind. In general, they will not be truncated at the same time with the basal ones, and if they are it will be differently.

These examples will suffice to make clear the law of symmetry and its application.

Nothing is simpler than to have a precise conception of hemihedry. It has long been known—in fact Haüv was acquainted with the most noted examples—that in a crystal sometimes only the half of the identical parts are modified simultaneously, and in the same manner. In such cases hemihedry is said to exist. Thus the cube ought to have its eight solid angles all truncated at once. But in certain cases this occurs only to four of them. Boracite furnishes an example of this nature. In these circumstances the modification takes place in such a way

that, if the four truncations are produced so as to obliterate the faces of the cube, a regular tetrahedron is obtained. If the modification were applied to the four remaining angles, it would give rise to another regular tetrahedron, identical with and superposable on the first, and differing from it only by its position on the cube.

In the same way let us again consider our right prism truncated on the eight basal edges. In certain species the truncation takes place on one half only of the edges, and here again the result is that, as the truncations occur on the opposite edges of each base, and alternately at the two extremities, these truncations, if produced, give rise to a tetrahedron. As in the case of the cube, two tetrahedra are possible, differently placed with reference to the prism, according as one or other of the sets of four truncations is preserved; but here the two tetrahedra are not absolutely identical. They are symmetrical tetrahedra. They cannot be superposed.

These notions will be sufficient to enable us to understand what hemihedry is, and what is understood by hemihedral faces or forms.

Now quartz, of which we were speaking a short time ago, is one of the few minerals in which Haüv discovered hemihedral faces. Everybody knows the ordinary form of this mineral, a regular hexagonal prism terminated by two six-faced pyramids. It is evident that the trihedral angles situated at the base of the pyramidal faces are identical, and, therefore, if one of these bears a face, all the others should exhibit the same modification. This is the case with the so-called rhomb-face of the mineralogists.

But Haüv was the first to observe, in certain specimens, a face very different from these, which he designated by the letter x, inclined more towards one side than the other, without being double, as the law of symmetry

would require in this case. Another strange peculiarity of these crystals did not escape crystallographers, namely, that this face x was inclined sometimes in one sense, sometimes in the other. Haüv, who liked to give suitable names to each variety of a species, called the variety of quartz bearing the face x plagihedral. The crystals in which the face x was inclined to the right, when the crystal was oriented in a given manner, were called right plagihedra; those in which x was inclined in the opposite sense, left plagihedra.

Nothing, however, is more variable than this character. Here it exists, there it is absent. On one and the same crystal there are angles bearing the face x, others which ought to bear it are without it. Sometimes both right and left plagihedral faces are found. Nevertheless, all who were conversant with crystals agreed in admitting that there was in quartz a true hemihedry in two opposite senses.

We should notice here a very ingenious association of ideas, due to Sir John Herschel, which was communicated to the Royal Society of London in 1820.

I have already said that BIOT made the remarkable observation, that among different specimens of quartz some deviated the plane of polarised light in one sense, and the others in the opposite sense, to the right and to the left respectively. This being established, HERSCHEL connected HAÜY'S crystallographic discovery with BIOT'S physical one. Experiment confirmed the idea of an actual relation between right and left plagihedra and the right and left senses of the optical deviations. Specimens of quartz bearing the face x in the same sense, all deviate the plane of polarised light in one sense.

Such is the statement of the principal facts antecedent to the researches of which I am about to give you a short account.

#### III.

When I began to devote myself to special work, I sought to strengthen myself in the knowledge of crystals, foreseeing the help that I should draw from this in my chemical researches. It seemed to me to be the simplest course, to take, as a guide, some rather extensive work on crystalline forms; to repeat all the measurements, and to compare my determinations with those of the author. In 1841, M. DE LA PROVOSTAYE, whose accuracy is well known, had published a beautiful piece of work on the crystalline forms of tartaric and paratartaric acids and of their salts. I made a study of this memoir. I crystallised tartaric acid and its salts, and investigated the forms But, as the work proceeded, I noticed of the crystals. that a very interesting fact had escaped the learned physicist. All the tartrates which I examined gave undoubted evidence of hemihedral faces.

This peculiarity in the forms of the tartrates was not very obvious. This will be readily conceived, seeing that it had not been observed before. But when, in a species, its presence was doubtful, I always succeeded in making it manifest by repeating the crystallisation and slightly modifying the conditions. Sometimes the crystals bore all the faces demanded by the law of symmetry, but the hemihedry was still betrayed by an unequal development of one half of the faces. This is seen, for example, in tartar emetic. It must be admitted that a circumstance which adds greatly to the difficulty in recognising hemihedry is the frequent irregularities of the crystals, which never develop quite freely. From this cause there arise deformations, arrestments of development in one direction or another, faces suppressed by accident, etc. Unless in circumstances of an almost exceptional character, the recognition of hemihedry, particularly in laboratory crystals, demands very attentive study. To this we must add the fact that, although hemihedry may be possible in a given form, and although it is a function of the internal structure of the substance, it may not be indicated externally, any more than one finds on every crystal of a cubic species all the forms compatible with the cube.

But however these things may be, I repeat that I found the tartrates hemihedral.

This observation would probably have remained sterile, without the following one.

Let a, b, c, be the parameters of the crystal form of any tartrate, and a,  $\beta$ ,  $\gamma$ , the angles of the crystallographic axes. The latter are ordinarily perpendicular, or slightly oblique. In addition, the ratio of two parameters, such as a and b, is almost the same in the various tartrates, whatever may be their composition, their quantity of water of crystallisation, or the nature of the bases; c alone shows sensible variations. There is a kind of semi-isomorphism among all the tartrates. One would imagine that the tartaric group dominated and stamped with similarity the forms of all the various substances in spite of the difference in the other constituent elements.

The results of this are, a resemblance in the forms of all tartrates, and the possibility of parallel orientation, taking, for example, as basis of orientation the position of the axes a and b.

Now if we compare the disposition of the hemihedral faces on all the prisms of the primitive forms of the tartrates, when they are oriented in the same manner, this disposition is found to be the same.

These results, which have been the foundation of all my later work, may be summed up in two words: the tartrates are hemihedral, and that in the same sense.

Guided then on the one hand by the fact of the exist-

ence of molecular rotatory polarisation, discovered by BIOT in tartaric acid and all its compounds, and on the other by HERSCHEL'S ingenious correlation, and yet again by the sagacious views of M. Delafosse, with whom hemihedry has always been a law of structure and not an accident of crystallisation, I believed that there might be a relation between the hemihedry of the tartrates and their property of deviating the plane of polarised light.

It is important thoroughly to grasp the development of the conceptions:—Haüy and Weiss observe that quartz possesses hemihedral faces and that these faces incline to the right on some specimens and to the left on others. Biot on his part finds that quartz crystals likewise separate themselves into two sets, in relation to their optical properties, the one set deviating the plane of polarised light to the right, the other to the left, according to the same laws. Herschel in his turn supplies to these hitherto isolated facts the bond of union, and says:—Plagihedra of one kind deviate in the same sense; plagihedra of the other kind deviate in the opposite sense.

For my own part I find that all tartrates are plagihedral, if I may so express myself, and that in the same sense; so that I might presume that here, as in the case of quartz, there was a relation between the hemihedry and the circular polarisation. At the same time the essential differences to which I have just referred between circular polarisation in quartz and in tartaric acid must not be neglected.

Thanks to the above discoveries, and to the relations which I have just enumerated, we are now in possession of a preconceived notion (for it is still nothing more than that) as to the possible inter-relation of the hemihedry and the rotative power of the tartrates.

Being very anxious to find by experiment some support for this still purely speculative view, my first thought was to see whether the very numerous crystallisable organic products which possess the molecular rotative property, have hemihedral crystalline forms, an idea which had not previously occurred to any one in spite of Herschel's correlation. This investigation met with the success which I anticipated.

I also occupied myself with the examination of the crystalline forms of paratartaric acid and its salts. These substances are isomeric with the tartaric compounds, but had all been found by BIOT to be inactive towards polarised light. None of them exhibited hemihedry.

Thus the idea of the inter-relation of the hemihedry and the molecular rotatory power of natural organic products gained ground.

I was soon enabled to establish it clearly by a wholly unexpected discovery.

### IV.

I must first place before you a very remarkable note by MITSCHERLICH which was communicated to the Académie des Sciences by BIOT. It was as follows:—

"The double paratartrate and the double tartrate of soda and ammonia have the same chemical composition, the same crystalline form with the same angles, the same specific weight, the same double refraction, and consequently the same inclination in their optical axes. When dissolved in water their refraction is the same. But the dissolved tartrate deviates the plane of polarisation, while the paratartrate is indifferent, as has been found by M. Biot for the whole series of those two kinds of salts. Yet," adds Mitscherlich, "here the nature and number of the atoms, their arrangement and distances, are the same in the two substances compared."

This note of MITSCHERLICH'S attracted my attention forcibly at the time of its publication. I was then a

pupil in the École Normale, reflecting in my leisure moments on these elegant investigations of the molecular constitution of substances, and having reached, as I thought at least, a thorough comprehension of the principles generally accepted by physicists and chemists. The above note disturbed all my ideas. What precision in every detail! Did two substances exist which had been more fully studied and more carefully compared as regards their properties? But how, in the existing condition of the science, could one conceive of two substances so closely alike without being identical? Mitscherlich himself tells us what was, to his mind, the consequence of this similarity:

The nature, the number, the arrangement, and the distance of the atoms are the same. If this is the case what becomes of the definition of chemical species, so rigorous, so remarkable for the time at which it appeared, given by CHEVREUL in 1823? In compound bodies a species is a collection of individuals identical in the nature, the proportion, and the arrangement of their elements.

In short, MITSCHERLICH'S note remained in my mind as a difficulty of the first order in our mode of regarding material substances.

You will now understand why, being preoccupied, for the reasons already given, with a possible relation between the hemihedry of the tartrates and their rotative property, MITSCHERLICH'S note of 1844 should recur to my memory. I thought at once that MITSCHERLICH was mistaken on one point. He had not observed that his double tartrate was hemihedral while his paratartrate was not. If this is so, the results in his note are no longer extraordinary; and further, I should have, in this, the best test of my preconceived idea as to the inter-relation of hemihedry and the rotatory phenomenon.

I hastened therefore to re-investigate the crystalline form of MITSCHERLICH'S two salts. I found, as a matter of fact, that the tartrate was hemihedral, like all the other tartrates which I had previously studied, but, strange to say, the paratartrate was hemihedral also. Only, the hemihedral faces which in the tartrate were all turned the same way, were, in the paratartrate inclined sometimes to the right and sometimes to the left. In spite of the unexpected character of this result, I continued to follow up my idea. I carefully separated the crystals which were hemihedral to the right from those hemihedral to the left, and examined their solutions separately in the polarising apparatus. I then saw with no less surprise than pleasure that the crystals hemihedral to the right deviated the plane of polarisation to the right, and that those hemihedral to the left deviated it to the left; and when I took an equal weight of each of the two kinds of crystals, the mixed solution was indifferent towards the light in consequence of the neutralisation of the two equal and opposite individual deviations.

Thus, I start with paratartaric acid; I obtain in the usual way the double paratartrate of soda and ammonia; and the solution of this deposits, after some days, crystals all possessing exactly the same angles and the same aspect. To such a degree is this the case that Mitscherlich, the celebrated crystallographer, in spite of the most minute and severe study possible, was not able to recognise the smallest difference. And yet the molecular arrangement in one set is entirely different from that in the other. The rotatory power proves this, as does also the mode of asymmetry of the crystals. The two kinds of crystals are isomorphous, and isomorphous with the corresponding tartrate. But the isomorphism presents itself with a hitherto unobserved peculiarity; it is the isomorphism of an asymmetric

crystal with its mirror image. This comparison expresses the fact very exactly. Indeed, if, in a crystal of each kind, I imagine the hemihedral facets produced till they meet, I obtain two symmetrical tetrahedra, inverse, and which cannot be superposed, in spite of the perfect identity of all their respective parts. From this I was justified in concluding that, by the crystallisation of the double paratartrate of soda and ammonia, I had separated two symmetrically isomorphous atomic groups, which are intimately united in paratartaric acid. Nothing is easier than to show that these two species of crystals represent two distinct salts from which two different acids can be extracted.

Using the treatment always employed in such cases, the purpose is served by precipitating each salt with a salt of lead or baryta, and then isolating the acids by means of sulphuric acid.

The study of these acids is of immense interest. I do

not know any that is more interesting.

But before enlarging on it allow me to introduce here some recollections in connection with their discovery.

### V.

The announcement of the above facts naturally placed me in communication with Biot, who was not without doubts regarding their accuracy. Being charged with giving an account of them to the Academy, he made me come to him and repeat before his eyes the decisive experiment. He handed over to me some paratartaric acid which he had himself previously studied with particular care, and which he had found to be perfectly indifferent to polarised light. I prepared the double salt in his presence, with soda and ammonia which he had likewise desired to provide. The liquid was set

aside for slow evaporation in one of his rooms. When it had furnished about 30 to 40 grams of crystals, he asked me to call at the Collége de France in order to collect them and isolate before him, by recognition of their crystallographic character, the right and the left crystals, requesting me to state once more whether I really affirmed that the crystals which I should place at his right would deviate to the right, and the others to the left. This done, he told me that he would undertake the rest. He prepared the solutions with carefully measured quantities, and when ready to examine them in the polarising apparatus, he once more invited me to come into his room. He first placed in the apparatus the more interesting solution, that which ought to deviate to the left. Without even making a measurement, he saw by the appearance of the tints of the two images, ordinary and extraordinary, in the analyser, that there was a strong deviation to the left. Then, very visibly affected, the illustrious old man took me by the arm and said :-

"My dear child, I have loved science so much throughout my life that this makes my heart throb."

You will pardon me, gentlemen, these personal recollections which have never been effaced from my mind. In our day, and with our habits, they would offend in a scientific memoir, but they have seemed to me not out of place in an oral account; and perhaps the biographical interest of such recollections will constitute one of the advantages of the kind of instruction which the Société Chimique inaugurates to-day.

Indeed there is more here than personal reminiscences. In Biot's case the emotion of the scientific man was mingled with the personal pleasure of seeing his conjectures realised. For more than thirty years Biot had striven in vain to induce chemists to share his conviction

that the study of rotatory polarisation offered one of the surest means of gaining a knowledge of the molecular constitution of substances.

### VI.

Let us return to the two acids furnished by the two sorts of crystals deposited in so unexpected a manner in the crystallisation of the double paratartrate of soda and ammonia. I have already remarked that nothing could be more interesting than the investigation of these acids.

One of them, that which comes from crystals of the double salt hemihedral to the right, deviates to the right, and is identical with ordinary tartaric acid. The other deviates to the left, like the salt which furnishes it. The deviation of the plane of polarisation produced by these two acids is rigorously the same in absolute value. The right acid follows special laws in its deviation, which no other active substance had exhibited. The left acid exhibits them, in the opposite sense, in the most faithful manner, leaving no suspicion of the slightest difference.

That paratartaric acid is really the combination, equivalent for equivalent, of these two acids, is proved by the fact that, if somewhat concentrated solutions of equal weights of each of them are mixed, as I shall do before you, their combination takes place with disengagement of heat, and the liquid solidifies immediately on account of the abundant crystallisation of paratartaric acid, identical with the natural product.\*

In accord with their chemical and crystallographic properties, all that can be done with one acid can be

<sup>\*</sup> This beautiful experiment called forth applause from the audience.

repeated with the other under the same conditions, and in each case we get identical, but not superposable products; products which resemble each other like the right and left hands. The same forms, the same faces, the same angles, hemihedry in both cases. The sole dissimilarity is in the inclination to right or left of the hemihedral facets and in the sense of the rotatory power.

### VII.

From these results, as a whole, it is evident that we have to do with two isomeric substances, whose general relations as regards similitude and molecular dissimilarity we know.

Now recall the definition of a chemical species which I gave a few minutes ago:—the aggregate of individuals identical in the *nature*, the *proportion*, and the *arrangement* of the elements. All the properties of substances are functions of these three terms, and the object of all our work is to pass, by investigation of the properties, to the knowledge of these three things.

In isomeric bodies, the nature and proportion are the same. The arrangement alone differs. The great interest of isomerism has been to introduce into the science the principle that substances may be, and are, essentially different solely because the arrangement of the atoms is not the same in their chemical molecules.

But no isomeric bodies existed whose relations in respect to molecular arrangement could be known. This desideratum was supplied for the first time by the discovery of the constitution of paratartaric acid, and of the constitutional relations of the right and left tartaric acids. We know, on the one hand, that the molecular structures of the two tartaric acids are asymmetric, and on the other, that they are rigorously the same, with the sole difference

of showing asymmetry in opposite senses. Are the atoms of the right acid grouped on the spirals of a dextrgoyrate helix, or placed at the summits of an irregular tetrahedron, or disposed according to some particular asymmetric grouping or other? We cannot answer these questions. But it cannot be a subject of doubt that there exists an arrangement of the atoms in an asymmetric order, having a non-superposable image. It is not less certain that the atoms of the left acid realise precisely the asymmetric grouping which is the inverse of this. Lastly, we know that paratartaric acid results from the juxtaposition of these two inversely asymmetric atomic groupings.

Henceforth the observation of the chemical and physical resemblances and differences, corresponding to these arrangements whose relations are known to us, offers especial interest, and gives solid foundations to molecular mechanics. It enables us to establish the connection of the physical and chemical properties, with the molecular arrangement which determines their very existence, or conversely it enables us to pass from the properties to their primary cause.

A résumé of these general relations between the properties and corresponding atomic arrangements may be given as follows:—

(1.) When the elementary atoms of organic products are grouped asymmetrically, the crystalline form of the substance manifests this molecular asymmetry in non-superposable hemihedry.

The cause of the hemihedry is thus recognised.

(2.) The existence of this same molecular asymmetry betrays itself, in addition, by the optical rotative property.

The cause of rotatory polarisation is likewise determined.\*

(3.) When the non-superposable molecular asymmetry is realised in opposite senses, as happens in the right and left tartaric acids and all their derivatives, the chemical properties of these identical and inverse substances are rigorously the same. From this it follows that this mode of contrast, and of similarity, does not alter the ordinary play of chemical affinities.

I am in error: there is a restriction to make on this last point, an important and eminently instructive restriction. Time does not permit me to develop it deliberately and suitably to-day. It will find a place in the next Lecture.

<sup>\*</sup> FRESNEL, with one of those flashes of genius, of which he had so many, had a sort of presentiment of this cause of rotatory polarisation.

He expresses himself thus, in one of his memoirs, in vol. xxviii. of the Annales de chimie et de physique, 1825:—"Rock crystal shows optical phenomena which cannot be reconciled with complete parallelism of the molecular lines, and which would seem to indicate a progressive and regular deviation of these lines in the passage from one layer of the medium to the next."

### SECOND LECTURE.

I.

HEN we study material things of whatever nature, as regards their forms and the repetition of their identical parts, we soon recognise that they fall into two large classes of which the following are the characters. Those of the one class, placed before a mirror, give images which are superposable on the originals; the images of the others are not superposable on their originals, although they faithfully reproduce all the details. A straight stair, a branch with leaves in double row, a cube, the human body,—these are of the former class. A winding stair, a branch with the leaves arranged spirally, a screw, a hand, an irregular tetrahedron,—these are so many forms of the other set. The latter have no plane of symmetry.

We know on the other hand that compound substances are aggregates of identical molecules, themselves formed of assemblages of elementary atoms distributed according to laws which regulate their nature, proportion, and arrangement. The individual, for every compound substance, is its chemical molecule, and this is a group of atoms. It is not a confused group: on the contrary, its arrangement is definitely fixed. Such is the view of all physicists in regard to the constitution of substances.

This being granted, it would certainly have been very extraordinary if nature, various in its effects as it is, and exhibiting laws which permit the existence of so many species of substances, had not presented us, among the atomic groups of compound molecules, with both of the categories into which all objects may be divided. In other words, it would have been wonderful if, in the

whole range of natural and artificial chemical substances, there had not been individuals with superposable images and others with non-superposable images.

The actual fact is just what might have been foreseen; all chemical compounds without exception likewise fall into two classes,—those with superposable images and those with non-superposable images.

### II.

It is easy to show that this is a legitimate consequence forced on us by our former discussion. To put it clearly before us I shall briefly recall the principal conditions of the decisive experiment with which we concluded the preceding Lecture.

I prepare the paratartrate of soda and ammonia from natural paratartaric acid. Beautiful crystals are deposited.

If any part of the solution of the double salt is examined in the polarising apparatus, it exhibits no trace of optical deviation; and if the acid is recovered from the crystals, paratartaric acid, identical with that used at first, is obtained. So far everything is simple and natural, and we would imagine that it was a question of crystallising an ordinary salt. Yet it is not so.

Take another portion of the same crystals and examine them one by one. You will find that half of them have the form, of which I here show a model, characterised by non-superposable hemihedry, and that the other half have the inverse form, identical with the first in all its respective parts and yet not superposable on it. If we then pick out the two sorts of crystals for separate solution, we find that one of the solutions deviates the polarised light to the right, the other to the left, and both to the same extent in absolute value.

Finally, if we extract the acids from these two sorts of

crystals by ordinary chemical means, we recognise that one of them is identical with ordinary tartaric acid, and that the other is in all respects similar to it without being superposable on it. They are related to each other in the same way as the salts from which they are obtained. They resemble each other as the right hand resembles the left: better still, they resemble each other like a symmetrical pair of irregular tetrahedra, and these analogies and differences are repeated in all their derivatives. What can be done with one can be done with the other under like conditions, and the resulting products exhibit always the same properties, with the single difference that in the one case the deviation of the plane of polarisation is to the right, in the other to the left, and that the forms of corresponding species, while identical in all their details, cannot be superposed.

All these facts, clear and decisive as they are, compel us to refer the general external characters of these acids and their compounds to their individual chemical molecules. To refuse to do this would be to disregard the most ordinary rules of logic. We thus arrive at the following consequences:—

(1.) The molecule of tartaric acid, whatever else it may be, is asymmetric, and in such a way that the image is not superposable. (2.) The molecule of the left tartaric acid is formed by the exactly inverse group of atoms. And by what characters shall we recognise the existence of molecular asymmetry? On the one hand by non-superposable hemihedry, on the other, and more especially, by the rotatory optical property when the substance is in solution.

These principles being granted, if we examine all the products of nature, or of the laboratory, we shall easily discover that a number of them exhibit at the same time this kind of hemihedry and the molecular rotatory pro-

perty, and that all the others show neither the one nor the other of these characters.

I was therefore justified in saying that the legitimate and inevitable consequence of our former discussion may

be expressed in this way:-

All bodies (I here employ the expression in the chemical sense) fall into two great classes, bodies with superposable images and bodies with non-superposable images,-bodies with asymmetric atomic arrangement and those with homohedral atomic arrangement.

### III.

Here we encounter a fact which would be worthy of our attention, even if we considered it alone and separate from the set of considerations which follows. It is this:-

All artificial products of the laboratory and all mineral species are superposable on their images. On the other hand, most natural organic products (I might even say all, if I were to name only those which play an essential part in the phenomena of vegetable and animal life), the essential products of life, are asymmetric, and possess such asymmetry that they are not superposable on their images.

Before going further, I wish to remove some objections which will not fail to arise in your minds.

### IV.

Quartz! you will say at once. We saw in the last Lecture that quartz possesses the two characteristics of asymmetry-hemihedry in form, observed by HAÜY, and the rotative phenomenon discovered by ARAGO! Nevertheless, molecular asymmetry is entirely absent in quartz. To understand this, let us take a further step in the knowledge of the phenomena with which we are dealing. We shall find in it, besides, the explanation of the analogies and differences already pointed out between quartz and natural organic products.

Permit me to illustrate roughly, although with essential accuracy, the structure of quartz and of the natural organic products. Imagine a spiral stair whose steps are cubes, or any other objects with superposable images. Destroy the stair and the asymmetry will have vanished. The asymmetry of the stair was simply the result of the mode of arrangement of the component steps. Such is quartz. The crystal of quartz is the stair complete. It is hemihedral. It acts on polarised light in virtue of this. But let the crystal be dissolved, fused, or have its physical structure destroyed in any way whatever; its asymmetry is suppressed and with it all action on polarised light, as it would be, for example, with a solution of alum, a liquid formed of molecules of cubic structure distributed without order.

Imagine, on the other hand, the same spiral stair to be constructed with irregular tetrahedra for steps. Destroy the stair and the asymmetry will still exist, since it is a question of a collection of tetrahedra. They may occupy any positions whatsoever, yet each of them will none the less have an asymmetry of its own. Such are the organic substances in which all the molecules have an asymmetry of their own, betraying itself in the form of the crystal. When the crystal is destroyed by solution, there results a liquid active towards polarised light, because it is formed of molecules, without arrangement, it is true, but each having an asymmetry in the same sense, if not of the same intensity in all directions.

#### V.

Quartz is therefore not molecularly asymmetric, and up to the present we have not any example of a mineral possessing molecular asymmetry. I have remarked that this proposition must be extended to the artificial compounds of the laboratory. Here again some scruples may be felt. It might be objected, for instance, that natural camphor, which is asymmetric, gives, artificially, camphoric acid, which is likewise asymmetric; that aspartic acid derived from asparagine by a laboratory reaction is asymmetric in the same way as asparagine, and I might cite many other similar examples. But no one can doubt that camphoric and aspartic acids owe their own asymmetry to camphor and to asparagine. This existed in the original products, and is handed on, more or less modified by substitution, from the original substances to their derivatives. In fact, a better proof of the conservation of the primitive type, in a series of products united together by a common origin, than that given by the permanence of the optical property, could not be adduced.

When I affirm that no artificial substance has yet shown molecular asymmetry, I refer to artificial substances properly so called, made entirely of mineral elements or derived from substances not themselves asymmetric. Thus, alcohol is not asymmetric. Its molecule, if we could isolate and study it, when placed before a mirror would give a superposable image. Now not a single derivative of alcohol is asymmetric. I could multiply ad infinitum examples of this kind. More than this: take any asymmetric substance whatever, and, if you submit it to fairly energetic chemical reactions, you may confidently expect to see the asymmetry of the primitive group disappear. Thus, tartaric acid is asymmetric. Pyrotartaric acid is not so. Malic acid is asymmetric. The maleic and paramaleic acids of Pelouze are not Gum is asymmetric, mucic acid is not. Artificial products have, therefore, no molecular asymmetry; and I could not point out the existence of any more profound distinction between the products formed under the influence of life, and all others. Let us enforce this point more strongly, for you will find the physiological side of these researches become more and more prominent in the remainder of this Lecture. Let us pass in review the principal classes of natural organic products.

Cellulose, starches, gums, sugars,—tartaric, malic, quinic, tannic acids,—morphine, codeine, quinine, strychnine, brucine,—essences of turpentine, of citron,—albumen, fibrin, gelatin. All these immediate principles are molecularly asymmetric. All these substances in the state of solution possess the rotative power, a characteristic necessary and sufficient to establish their asymmetry, even when, crystallisation being impossible, hemihedry is lacking as a means of recognising this property.

All the substances most essential in the animal or vegetable organism appear in this list.

There are many natural substances which are not asymmetric. But are they natural in the same sense as the others? Must we not regard bodies like oxalic acid, hydride of salicyle, fumaric acid, etc., as derivatives of natural substances properly so called, formed by interactions analogous to those of the laboratory? These products appear to me to be, in the vegetable organism, what urea, uric acid, creatine, glycocoll, etc., are in the animal organism—execretions rather than secretions, if I may so speak. It would be very interesting to follow up this point of view experimentally.

Let us add that many substances which are apparently not asymmetric, may be like paratartaric acid. We still want a word in chemical language to express the fact of a double molecular asymmetry concealed by the

neutralisation of two opposite asymmetries, the physical and geometrical effects of which rigorously compensate each other.

The double proposition to which we have been led, on the habitual asymmetry of immediate organic principles, and on the absence of this character in all the products of inorganic nature, will permit us to enlarge and render more definite our mode of regarding the subject of this remarkable molecular property.

#### VI.

In 1850 M. Dessaignes, whose ingenuity and skill are known to all chemists, announced to the Academy that he had succeeded in transforming bimalate of ammonia into aspartic acid. This was an advance which confirmed the important results obtained by Piria some years before. Piria had succeeded in transforming asparagine and aspartic acid into malic acid. M. Dessaignes, in turn, showed that conversely we could return from malic acid to aspartic acid.

So far everything connected with M. Dessaignes' observation was perfectly natural, even from the optical point of view. For, on my side, I had found that asparagine, aspartic acid and malic acid were active towards polarised light. The passage chemically from one to the other of these bodies had nothing astonishing about it.

A few months later M. Dessaignes took a further step. He announced that not only bimalate of ammonia, but also fumarate and maleate of ammonia had likewise the property of being transformed by heat into aspartic acid.

Here I saw an impossibility; or, if the facts were as M. Dessaignes stated, this skilful chemist had made a

discovery which he did not suspect. I had in fact observed that fumaric and maleic acids and all their salts were without action upon polarised light. If, therefore, M. Dessaignes had succeeded in transforming their ammonia salts into aspartic acid, he would have accomplished for the first time the production of an asymmetric substance by the aid of compounds which are not so.

But it seemed to me more reasonable to suppose that the aspartic acid of M. Dessaignes differed from natural aspartic acid in the absence especially of the molecular rotatory property. M. Dessaignes, it is true, had carefully compared the properties of the artificial acid with those of the natural acid, and stated that he had found them identical. After the example of Mitscherlich, of which I spoke in last Lecture, I knew better than anyone else what delicate things were the demonstrations of the identity of chemical species, in investigations where the greatest similarity in properties often conceals profound differences. I had therefore no hesitation in believing that the new fact reported by M. Dessaignes required confirmation.

Attaching so much importance to the elucidation of this question, and indeed foreseeing the results which I am about to have the honour of explaining to you, I immediately set out for Vendôme, where I related my suspicions to M. Dessaignes, and he hastened to supply me with a specimen of his aspartic acid. As a matter of fact I found, as soon as I reached Paris, that M. Dessaignes' acid was only an isomer of natural aspartic acid, that is of the acid from asparagine. It differed from it, as I had foreseen, in the rotatory property, which was quite absent in the artificial acid and indubitable in the natural acid. Yet all the other chemical and physical properties showed the greatest analogies, so great that

M. Dessaignes, not being put on his guard by any preconceived idea, had decided that the two substances were really identical.

What charmed me most in the examination of the new compound (which in itself shows no remarkable crystallisable compounds), was its transformation into malic acid. It is known indeed, I mentioned the fact a moment ago, that PIRIA long since showed how to pass from asparagine and aspartic acid to malic acid, and I had assured myself by the most rigid proofs that this malic acid was identical with that from the mountain ash, from apples, from the grape, and from tobacco.

I therefore applied to the new acid the treatment discovered by Piria and actually transformed it into a new malic acid, very similar to the natural acid. It was so nearly allied to the latter that a chemist would have had difficulty in distinguishing them, even if he had been warned of their real disparity. Only, this malic acid had no action on polarised light, and the same was true of all its salts.

There are certain derivatives of these two malic acids which, on comparison, do not exhibit very clearly the real mutual dependence of the molecular arrangements of these curious isomers; but there are others in which it exhibits itself to its full extent. Let us consider, for example, the ordinary active bimalate of lime, and the corresponding inactive bimalate. Their chemical composition is exactly the same and their crystalline forms are alike, with the difference that the active one has four small hemihedral faces which are always absent in the inactive one. The result is that, when the active one is placed before a mirror, the image cannot be superposed on it, while the image of the inactive one is absolutely identical with and superposable on the reality which produces it. But in all that does not concern the

hemihedral faces, the resemblance of the two forms is perfect.

Who could doubt, after that, the relation of the molecular arrangements in the two salts? Is it not evident that we have here to deal with a malic acid identical with the natural one, except for the simple suppression of its molecular asymmetry?

It is natural malic acid untwisted, if I may so express myself. The natural acid is a spiral stair as regards the arrangement of its atoms, this acid is the same stair made of the same steps, but straight in place of being spiral.

It might be asked whether the new acid is not the paratartaric member of the series, that is, the compound of the right malic acid with the left malic acid. That is improbable, for then not only would we have made an active body from an inactive one, we would have made two, a right and a left.

Besides, I have found that, just as there exists an inactive non-asymmetric malic acid, there is likewise an inactive non-asymmetric tartaric acid quite different from paratartaric acid, which cannot be resolved into a right tartaric acid and a left tartaric acid. It is impossible to doubt that we have here to do with right or left tartaric acid rendered non-asymmetric.

I have likewise discovered inactive amyl alcohol which gives rise to a whole series of inactive products corresponding to the series from active amyl alcohol.

We are now, thanks to the discovery of the inactive substances, in possession of a fertile idea. A substance is asymmetric, right or left; by certain artifices of isomeric transformation, which must be sought and discovered for each particular case, it may lose its molecular asymmetry, become untwisted, to use a rough metaphor, and assume in the arrangement of its atoms a disposition with superposable image. In this way each asymmetric substance

offers four varieties, or, better, four distinct sub-species: the right body, the left body, the combination of the right and the left, and the substance which is neither right nor left nor formed by the combination of the right and the left.

#### VII.

This general conclusion from the above-mentioned investigations throws a new light on our ideas of molecular mechanics. We discover that if the natural products elaborated under the influence of vegetable life are, as a rule, asymmetric, contrary to what we find in the case of artificial and mineral products, this disposition of the elementary particles is not a condition of the existence of the molecule,—that the twisted organic group can be untwisted, and so assume the ordinary character of artificial and mineral substances. Conversely, it seems to me logical to regard the latter as capable of exhibiting an asymmetric arrangement of their atoms after the manner of the natural products. The conditions for their production have still to be discovered.

To sum up what has been said, the groups of elementary atoms which constitute compound matter can assume two distinct states corresponding to the two general types under which every material object can be classified. The form of the group has either a superposable or a non-superposable image; but the latter type is double, for its inverse can exist equally well with itself. We must add the case of the association of these two inverse types, recalling the union by pairs of identical and non-superposable members in the higher animals. There are therefore in reality four remarkable arrangements for the groups of atoms which constitute matter. All our efforts should be bent to produce them for each particular species.

Almost all these conclusions are so rigorous that it is practically impossible to call them in question.

Thus, how can we refuse to admit that a right body has a possible left form, knowing as we do the significance of the right or left character? We might as well doubt that an irregular tetrahedron has its inverse, that a right helix has its left form, that a right hand has a possible left.

And consequently, if the mysterious influence to which the asymmetry of natural products is due should change its sense or direction, the constitutive elements of all living beings would assume the opposite asymmetry. Perhaps a new world would present itself to our view. Who could foresee the organisation of living things if cellulose, right as it is, became left; if the albumen of the blood, now left, became right? These are mysteries which furnish much work for the future, and demand henceforth the most serious consideration from science.

## VIII.

Possibly, since chemistry has been up to the present time powerless to prepare asymmetric substances, one might fear that we should remain for ever in ignorance of a way of making the inverse forms of natural organic substances. Happily this fear is exaggerated. In fact I have discovered that ordinary chemical processes, such as the action of heat, enable us to pass from a right to a left substance, and vice versa. Thus, when right tartaric acid is heated under certain fixed conditions which it would take too long to specify here, it is transformed into left tartaric acid, or rather into paratartaric acid. And conversely under the same conditions precisely, left tartaric acid becomes right.

Here are ten or twelve grams of perfectly pure left tartaric acid, which have been obtained in this manner.

Their preparation has cost me much trouble. But M. BIOT was particularly anxious to study the dispersive properties of this left tartaric acid of such remarkable origin. He wished himself to defray the cost of the operation, which was very expensive, for the transformation depends on the employment of tartrate of cinchonine or of quinine, and the base is lost because the tartrate has to be heated to a temperature which destroys it. By this means I prepared paratartaric acid enough to yield twelve grams of left tartaric acid, which exhibited, in the inverse sense, absolutely the same optical characters as tartaric acid.

We must always consider every transformation like this of a natural asymmetric body into its inverse as a great advance of organic chemistry.

### IX.

At the conclusion of our former meeting I referred to some observations to which it is now time that we devoted the attention which they deserve. These observations are connected with the comparison of the physical and chemical properties of the corresponding right and left isomers. I have already insisted on the perfect identity of all their properties, excepting always the inversion in their crystalline forms and the opposite sense of their optical deviations. The physical aspect, lustre of the crystals, solubility, specific weight, simple or double refraction, all these things are not merely alike, similar, nearly allied, but identical in the strictest sense of the word.

This identity is all the more remarkable since we shall see it replaced by a general and noteworthy contrast of the properties of the same substances when they are subjected to special conditions which I am about to indicate. We have seen that all artificial or natural chemical compounds, whether mineral or organic, must be divided into two great classes: non-asymmetric compounds with superposable image and asymmetric compounds with non-superposable image.

Taking this into account, the identity of properties above described in the case of the two tartaric acids and their similar derivatives, exists constantly, with the unchangeable characters which I have referred to, whenever these substances are placed in contact with any compound of the class with superposable image, such as potash, soda, ammonia, lime, baryta, aniline, alcohol, ethers,—in a word, with any compounds whatever which are non-asymmetric, non-hemihedral in form, and without action on polarised light.

If, on the contrary, they are submitted to the action of products of the second class with non-superposable image, —asparagine, quinine, strychnine, brucine, albumen, sugar, etc., bodies asymmetric like themselves,—all is changed in an instant. The solubility is no longer the same. If combination takes place, the crystalline form, the specific weight, the quantity of water of crystallisation, the more or less easy destruction by heating, all differ as much as in the case of the most distantly related isomers.

Here, then, the molecular asymmetry of a substance obtrudes itself on chemistry as a powerful modifier of chemical affinities. Towards the two tartaric acids, quinine does not behave like potash, simply because it is asymmetric and potash is not. Molecular asymmetry exhibits itself henceforth as a property capable by itself, in virtue of its being asymmetry, of modifying chemical affinities. I do not believe that any discovery has yet made so great a step in the mechanical part of the problem of combination.

Let us try to illustrate the cause of these identities and

differences. Let us imagine a right screw and a left screw separately penetrating two identical blocks of wood with the grain straight. All the mechanical conditions of the two systems will be the same. This will no longer be so from the moment that the same screws are associated with blocks which are themselves twisted in the same sense or in the opposite sense.

#### X.

Here is a very interesting application of the facts which have just been explained.

Seeing that the right and left tartaric acids formed such dissimilar compounds simply on account of the rotative power of the base, there was ground for hoping that, from this very dissimilarity, chemical forces might result, capable of balancing the mutual affinity of the two acids, and thereby supply a chemical means of separating the two constituents of paratartaric acid. I sought long in vain, but finally succeeded by the aid of two new bases, quinicine and cinchonicine, isomers of quinine and cinchonice, which I obtained very easily from the latter without the least loss.

I prepare the paratartrate of cinchonicine by neutralising the base and then adding as much of the acid as was necessary for the neutralisation, I allow the whole to crystallise, and the first crystallisations consist of perfectly pure left tartrate of cinchonicine. All the right tartrate remains in the mother liquor because it is more soluble. Finally this itself crystallises with an entirely different aspect, since it does not possess the same crystalline form as the left\* salt. We might almost believe that we were dealing with the crystallisation of two distinct salts of unequal solubility.

<sup>\*[&</sup>quot;Right" in original.]

### XI.

But the difference in properties of corresponding right and left substances when they are subjected to asymmetric forces, seems to me to be interesting in the highest degree on account of the ideas which it suggests to us in regard to the mysterious cause which presides over the asymmetric arrangement of the atoms in natural organic substances. Why this asymmetry? Why the one asymmetry rather than its inverse?

Carry yourselves back with me in thought to the time when, having recognised the absolute identity of the chemical and physical properties of the corresponding right and left substances, I had not any idea, not even the suspicion, of possible differences between these substances. It was, in fact, after an interval of several years that I recognised these similarities and differences.

It was then impossible for me to understand how nature could make a right substance without at the same time making the left substance. For the same forces which are at work at the moment at which the molecule of right tartaric acid is elaborated, must, it seemed, yield a left molecule also, and there would have been nothing but paratartaric compounds.

Why even right or left substances at all? Why not simply non-asymmetric substances; substances of the order of inorganic nature?

There are evidently causes for these curious manifestations of the play of molecular forces. To indicate them in a precise manner would certainly be a difficult matter. But I believe that I am not deceived when I say that we know one of their essential characters. Is it not necessary and sufficient to admit that at the moment of the elaboration of the primary principles in the vegetable organism, an asymmetric force is present? For we have just seen

that there is only a single case in which the right molecules differ from the left, the case where they are submitted to influences of an asymmetric character.

Do these asymmetric actions, possibly placed under cosmic influences, reside in light, in electricity, in magnetism, or in heat? Can they be related to the motion of the earth, or to the electric currents by which physicists explain the terrestrial magnetic poles? It is not even possible at the present time to express the slightest conjecture in this direction.

But I regard as necessary the conclusion that asymmetric forces exist at the moment of the elaboration of natural organic products; forces which would be absent, or ineffectual, in the reactions of our laboratories either on account of the violent course of these phenomena, or because of some other unknown circumstance.

## XII.

We now reach a final experiment which does not yield in interest to any of those which precede, in respect to the manifest proof which it will give us of the influence of asymmetry in the phenomena of life. We have just seen asymmetry intervening to modify chemical affinities; but the reactions involved were purely inorganic, artificial, and we know all the prudence that must be observed in the application of the results of the laboratory to the phenomena of life. On this account I kept to myself almost all the views expressed in this Lecture till the moment that I recognised in the most certain manner that molecular asymmetry exhibited itself as a modifier, no longer of the reactions of inorganic nature, but of those of physiological character, in fermentations.

Here is the remarkable phenomenon to which I refer: It had long been known, from the observation of a German manufacturer of chemical products, that the impure tartrate of lime of the works, mixed with organic matters, when left under water in summer, could ferment, giving various products.

Knowing this, I set the ordinary right tartrate of ammonia to ferment in the following manner. I took the very pure crystallised salt, dissolved it, adding to the liquor a clear solution of albumenoid matter. One gram of dry albumenoid matter was sufficient for one hundred grams of tartrate. Very often it happens that the liquid ferments spontaneously when placed in an oven. I say very often; but it may be added that this will always take place if we take care to mix with the liquid a very small quantity of one of those liquids with which we have succeeded in obtaining spontaneous fermentation.

So far there is nothing peculiar; it is a tartrate fermenting. The fact is well known.

But let us apply this method of fermentation to paratartrate of ammonia, and under the above conditions it ferments. The same yeast is deposited. Everything shows that things are proceeding absolutely as in the case of the right tartrate.' Yet if we follow the course of the operation with the help of the polarising apparatus, we soon discover profound differences between the two operations. The originally inactive liquid possesses a sensible rotative power to the left, which increases little by little and reaches a maximum. At this point the fermentation is suspended. There is no longer a trace of the right acid in the liquid. When it is evaporated and mixed with an equal volume of alcohol it gives immediately a beautiful crystallisation of left tartrate of ammonia.

Let us note, in the first place, two distinct things in this phenomenon. As in all fermentation properly so called, there is a substance which is changed chemically, and correlatively there is a development of a body possessing the aspect of a mycodermic growth. On the other hand, and it is this which it is important to note, the yeast which causes the right salt to ferment leaves the left salt untouched, in spite of the absolute identity in physical and chemical properties of the right and left tartrates of ammonia as long as they are not subjected to asymmetric action.

Here, then, the molecular asymmetry proper to organic substances intervenes in a phenomenon of a physiological kind, and it intervenes in the rôle of a modifier of chemical affinity. It is not at all doubtful that it is the kind of asymmetry proper to the molecular arrangement of left tartaric acid which is the sole and exclusive cause of the difference from the right acid, which it presents in relation to fermentation.

Thus we find introduced into physiological principles and investigations the idea of the influence of the molecular asymmetry of natural organic products, of this great character which establishes perhaps the only well marked line of demarcation that can at present be drawn between the chemistry of dead matter and the chemistry of living matter.

## XIII.

Such, gentlemen, are in co-ordinated form the investigations which I have been asked to present to you.

You have understood, as we proceeded, why I entitled my exposition, "On the Molecular Asymmetry of Natural Organic Products." It is, in fact, the theory of molecular asymmetry that we have just established, one of the most exalted chapters of the science. It was completely unforeseen, and opens to physiology new horizons, distant, but sure.

I hold this opinion of the results of my own work with-

out allowing any of the vanity of the discoverer to mingle in the expression of my thought. May it please God that personal matters may never be possible at this desk. These are like pages in the history of chemistry which we write successively with that feeling of dignity which the true love of science always inspires.



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