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RURAL CHEMISTRY.
RURAL CHEMISTRY:

AN

Elementary Introduction

TO

THE STUDY OF THE SCIENCE IN ITS RELATION TO AGRICULTURE AND THE ARTS OF LIFE.

BY

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PREFACE TO THE FIRST EDITION.

The following pages formed the substance of a short series of articles on Chemistry, which originally appeared in the columns of the Gardeners' Chronicle. The interest which they excited in the readers of that journal, has led to their republication in a separate and more complete form. It would have been easy to have greatly increased the size of the book; and indeed it was frequently very difficult to select from the mass of information which exists those facts which appeared most worthy of notice. The original object of the Author was to give such an elementary sketch of the science as should enable those ignorant of the subject more readily to comprehend the works of the various authors who have written on Agricultural Chemistry.
As a general rule, care has been taken, as much as possible, merely to give well-established facts, or, when doubtful theories are mentioned, to state distinctly that they are more or less problematical.
PREFACE TO THE THIRD EDITION.

In accordance with numerous suggestions, very considerable additions have been made to this little book, in preparing it for a new edition; several important practical matters, not treated of in the former editions, having been introduced. Brief descriptions of the more important of the domestic arts, such as wine and vinegar making, brewing, the manufacture of spirits, baking, cheese-making, cookery, &c., have been added; together with some account of the scientific principles involved in those arts. Numerous recent analyses of agricultural crops have likewise been given, and the whole has been carefully revised and corrected.

E. SOLLY.

Tavistock Square, London,
Nov. 1, 1850.
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INTRODUCTION.

No branch of knowledge has made more rapid progress, during the last hundred years, than chemistry. From being merely a confused collection of marvelous facts and incomprehensible phenomena, it has become a definite and methodical science: no longer mysterious and full of uncertainty, but based on clear and simple laws, the knowledge of chemistry at once gives us the key to a great number of natural changes and phenomena, which without it, would be quite unintelligible.

Chemistry is intimately connected with all other sciences; for it embraces the study of the various forms and conditions of matter, their nature and properties, and the changes which, either from natural or artificial causes, they undergo. The study of chemistry is of the greatest importance in relation to the arts of life, which all depend more or less on
chemical principles. Hence, a knowledge of this science enables us more readily to understand the processes of the manufacturer, points out the best and most economical mode of effecting his objects, and teaches how that which was before useless and of no value, may be converted into sources of wealth and happiness.

It is needless to point out examples of the influence which the progress of chemistry has had on our manufactures; for every one of them owes, more or less directly, its present improved condition to the labors of the chemist. Such being the case, it becomes interesting to inquire, What has chemistry done for agriculture, the most important, because the most necessary, and most extensively practised, of all the arts? It is remarkable that agriculture should have received infinitely less assistance from the labors of chemists, than any other art; but this ceases to be surprising, when we consider how little attention was paid by the ancient chemists to that subject.—The chemistry of the earthy and metallic substances presented to them easier and more attractive objects of inquiry; they were led away by the visionary hope of discovering a mode of making gold; and they consequently neglected everything, in order to try all sorts of experiments, in the vain idea
of converting lead, iron, &c., or the base metals as they called them, into gold. Hence, it is not to be wondered at, that the chemistry of the metals was studied and investigated, long before the nature and properties of vegetable and animal substances were examined. The labors of the alchemists brought to light many valuable discoveries respecting the uses, nature, and properties, of earthy and metallic substances; but it was only in the last century, when the constitution of the air and other gaseous bodies was discovered, that anything approaching to correct ideas respecting the nature of animal and vegetable substances was entertained.

The important discoveries of Dr. Priestley led the way to a complete revolution in the science, and may almost be said to have laid the foundation of agricultural chemistry. It is true that, before his time, there had been many careful and accurate observers, and multitudes of laborious and valuable experiments have been made on plants, by such men as Van Helmont—Evelyn—Boyle—Hales, and others; but, before the time of Priestley and his contemporaries, Bonnet, Ingenhousz, Henry, and Percival, little progress had been made in studying the chemical changes on which the growth of plants depends. We are indebted to Hales for much curious information
respecting the rise and motion of the sap in plants, the perspiration, or evaporation, which is constantly going on to a greater or less extent, from the surface of the leaves, and the effects of various substances on plants; his chemical speculations, however, are for the most part, of little value, though he was often apparently on the point of making important discoveries. The discovery of carbonic acid gas, or fixed air, by Dr. Black, and the beautiful experiments of Priestley, opened a new field of inquiry and research: he observed that plants possessed the property of purifying the air; in fact, that they were able to decompose the carbonic acid gas which it always contains in small quantity; appropriating the carbon, and restoring back to the atmosphere the oxygen, or vital air, so necessary to the processes of respiration and combustion. The knowledge of this great fact, necessarily led to many minor discoveries respecting the growth of plants, and the sources of their food. After this period, Organic Chemistry began to attract a large share of the attention of chemists, the composition of vegetable substances was carefully investigated, new modes of analysis were discovered, and an immense mass of curious and useful facts was collected. A great number of chemists occupied themselves with researches in Vegetable Chemistry, but
for the most part they were employed in examining the innumerable substances which plants produce; whilst the great questions as to the food of plants, their growth, and nourishment, were left very nearly in the same state which the experiments of Priestley and Ingenhousz had brought them to.

At the end of the last century, and in the commencement of the present, Organic Chemistry made rapid advances; the labors of Hassenfratz, Humboldt, Berzelius, Saussure, Senebier, Einhof, and Davy, contributed to throw light on many parts of the subject; whilst the investigations of Gay-Lussac, Hatchett, Lampadius, Lavoisier, Marcet, Prout, Thompson, Vauquelin, Thénard, and others, in all parts of Europe, led to a more complete and accurate knowledge of the nature, composition, and properties of organic matter.

The first chemist who wrote on agriculture appears to have been J. G. Wallerius, who, in 1754, published a book on the Cause of Fertility. Even before this time, however, several books had been written on agriculture, in which attempts were made to explain the operations of farming on chemical principles; such, for example, were The Rational Farmer, 1743, a curious book, containing numerous accounts of rude chemical experiments, together with a num-
ber of sound practical facts: the author, however, was evidently not a chemist. Wallerius was succeeded by several other authors, amongst whom ought to be mentioned Cullen, Pearson, Gyllenborg; De Beunie, Rückert, Einhof, and Dundonald; but the speculations of these authors (though ingenious) were, for the most part, crude and incomplete. The writings of Einhof were valuable for the numerous accounts of careful experiments which they contain; the analyses, though not carried to that degree of minuteness which subsequent discoveries led to, were trustworthy and accurate, and, as such, will always continue of value. Humboldt's *Sketch of the Chemical Physiology of Vegetation*, which appeared at this time, is a book of far higher talent than those just mentioned, and contains enlarged views and cautious generalizations, which the subsequent progress of science has in most cases confirmed. At the commencement of the present century, when Organic Chemistry was rapidly advancing, Berzelius and Davy endeavored to apply the conclusions to be derived from chemical experiments, to agriculture. If the deductions which they made were not always correct, and if the plans which they proposed did not always produce the effects which the authors anticipated, it must be remembered that they were
amongst the first to take the subject up, and that though they did so under far more favorable circumstances than their predecessors did, yet that even then the science of organic chemistry was in many respects very imperfect and incomplete.

During the last forty years, many important additions have been made to this department of Chemistry; improved and more accurate modes of chemical investigation have enabled more exact analyses to be made of the different varieties of organic matter; the composition of those substances which constitute the bodies of animals and plants has been accurately and carefully ascertained. At the same time, many valuable observations have been made respecting the functions of plants, the conditions requisite to germination, the formation of flowers and seed, the chemical changes attendant on the ripening of fruit, the office performed by roots and leaves, and a variety of other important subjects of inquiry. The names of Liebig, Schübler, and Sprengel, in Germany; Braconnot, Boussingault, Chevreul, Colin, Chaptal, Dumas, Edwards, and Payen, in France; and of Daubeney, Fownes, Johnston, Pepys, Turner, Christison, and Way, in our own country, deserve especial mention.

Even in so short a sketch of the subject as this, it
would not be right to omit altogether the name of Grisenthwaite, whose book on the theory of Agriculture (1819) is remarkable for the enlarged and extended views which it contains. It is true that the author falls into many errors; but, at the same time, he was the first who entertained correct views of the importance of nitrogen as an element of manure, and of the necessity of supplying phosphates, as well as substances containing nitrogen, to plants, like wheat, chiefly cultivated for the sake of the azotized principle which renders them valuable as food.

Intimately connected with the progress of Vegetable Chemistry, is the study of Vegetable Physiology: a knowledge of the one is essential to a perfect comprehension of the other; for it is impossible well to understand the chemical changes going on in the organs of plants, if we are wholly ignorant of the forms and structure of those organs; and, on the other hand, the most complete knowledge of the anatomy of vegetables could never lead any one to sound and correct conclusions respecting the nutrition of plants. It is rather to be regretted that both Chemists and Physiologists have appeared to avoid availing themselves of the advantages which each might have derived by studying the results that the others had obtained; it is only by comparing
together the observations of both, that correct conclusions can be formed. The observations made by the older physiologists, like those of their chemical contemporaries, were mostly imperfect, and the deductions they formed, were, in consequence, very frequently erroneous; the modes of examination, and the instruments which they employed, were far less perfect than those which have been used in more recent times. Nevertheless, the observations recorded by Grew, Malpighi, and Duhamel, are of considerable value; they may be said, indeed, to have laid the foundation of Vegetable Physiology. As the study of Botany itself advanced, greater care was bestowed in examining the structure and anatomy of plants; and, from the labors of many zealous and careful observers, there has resulted a tolerably complete system of Vegetable Physiology. In recent years, De Candolle, Brongniart, Decaisne, Dutrochet, and Mirbel, in France; Link, Mohl, Meyen, and Schleiden, in Germany; Amici, in Italy; and Brown, Griffiths, Henslow, Knight, and Lindley, in England, have, besides many others, made valuable additions to Vegetable Physiology. The relations of plants to climate, and the influence of heat, light, and moisture, have also been studied; especially by Daniell and Royle in our own country.
Amongst the names of those who have contributed to the science of Agricultural Chemistry, that of Liebig stands pre-eminent. The thanks of all are due to him, both for the valuable and laborious experiments he has performed, and likewise for the exertions which he has made to remove the many doubts and uncertainties that surrounded the very elements of the subject. But little has been done since the days of Priestley and Ingenhousz to prove how plants obtained their food, what were the sources whence they derived the elements of organic matter, and the nature of the office performed by manures. Chemists, and likewise physiologists, had formed many ingenious speculations; but they had not employed the only real mode of getting at the truth, namely, well-selected and carefully performed experiments. Liebig, in his *Organic Chemistry applied to Agriculture and Physiology*, has strongly drawn attention to these important questions; he has exposed the fallacy of many of the theories which had been formed to explain them, and has established, on good evidence, the simple chemical rules which regulate the growth of plants.

Although the experiments of Priestley and Ingenhousz had shown that plants possess the power of decomposing Carbonic Acid; and, although they had
advanced numerous arguments to prove that plants derive the carbon which they contain from the decomposition of that gas; yet this doctrine, although admitted by many physiologists, was by no means universally believed by chemists. M. Hassenfratz, in particular, opposed these views, asserting that plants did not derive their carbon from the decomposition of carbonic acid existing in the air, but absorbed it direct from the soil, in a state of suspension or solution; he gives the name of carbon to the brown substance left on the evaporation of dung water, and, in fact, to the various modifications of decaying organic matter, subsequently described under the general name of Humus. Few experiments, indeed, were made to show that the explanation of Priestley and Ingenhousz was improbable; but it was conceived that plants must derive their carbon from the soil, and many theories were formed to explain the mode in which they might be supposed to obtain it. These theories have been rigidly examined by Liebig, and the results of his investigation have shown, that the old views put forth by Priestley and Ingenhousz were in truth correct.

It has long been known that plants consist of Carbon, Oxygen, Hydrogen, and Nitrogen, and also that they invariably contain a small quantity of inor-
ganic, or earthy and saline matters. The presence of Nitrogen was formerly greatly overlooked, in analyses of vegetable substances; it is contained in less quantity than the other three elements of organic matter, and was very commonly regarded as being merely accidental, and not a necessary constituent of plants. Improved modes of analysis have proved that nitrogen always exists in the same proportion, in certain constituents of plants; and, as it appears that these substances are also those which form the most valuable part of food, it becomes a question of the first importance—Whence do the plants derive their Nitrogen? They obtain it, principally, if not wholly, from the air; they do not absorb it in the free and uncombined form, but they absorb it combined with Hydrogen and with Oxygen, in the states of Ammonia and Nitric Acid. The importance of the earthy substances in plants was, likewise, greatly overlooked formerly. It has been proved, by repeated experiments, that these substances are of the greatest importance in the growth of plants, being quite essential to their development.

Although much has been done, and although chemists have labored to remove the perplexities which encompassed the subject, there is still a very great deal which requires investigation; many important
INTRODUCTION.

points are as yet imperfectly, or even not at all, explained, and many questions must be satisfactorily settled before a complete system of Agricultural Chemistry can be established. Till these difficulties are removed, it is premature to expect that Chemistry can be of more than partial assistance to Agriculture; for, whilst many of the fundamental laws of Agricultural Chemistry are still scarcely understood, all attempts to apply them to practice must be incomplete, and liable to error.

The composition of the principal varieties of organic matter is well known; the substances which, by combining together, form the various constituents of plants, have been ascertained. The food of plants, the great sources whence they derive it, and the manner in which they absorb it, are known. The various changes which organic matter undergoes, the conversion of one substance into another, and the influence which these changes have on the growth of plants, is likewise easily understood; nearly all the purely chemical operations which are concerned in their nutrition, can be explained by reference to simple chemical laws; but there are many most important phenomena which are as yet wholly in the dark. Thus, for example, the manner in which wood is formed; and, indeed, all those natural operations
in which cellular or organized matter is generated under the influence of light and heat, are but very imperfectly explained. A knowledge of the chemical composition of soils, and the various substances employed as manures, enables us to comprehend the mode in which the latter act; and a knowledge of the nature of those substances which plants require, points out the best and most economical methods of restoring to the soil, by manures, those substances which plants remove from it: but our knowledge of this part of the subject is very far from being complete; for although it is certain that, in addition to the great elements of organic matter, which plants derive from both air and soil, they likewise absorb small quantities of inorganic or mineral substances from the soil exclusively, the office performed by the latter in the vegetable economy is not yet well understood. Many theories, indeed, have been formed respecting their use, but very little is positively known on the subject.

Although Agricultural Chemistry is in this imperfect state, and though much still remains to be done in that branch of science, yet it is so far advanced as to be able to render substantial assistance to the practical agriculturist. It can teach him the principles which govern the growth of plants, and, conse-
quently, guide him in the application of artificial means to produce the most beneficial results. He must, however, not expect too much from the aid of Chemistry, nor give himself blindly up to speculations or theories. Whilst he gives due credit and belief to well-authenticated facts, he must always receive theories with caution and doubt.

Perhaps the most important advantage which a practical man may at present derive from a knowledge of Agricultural Chemistry, is connected with the use of manure. If he knows what it is that gives the fertilizing powers to manure, and is aware of the nature of those substances, he will soon learn the best method of preserving and using them; he will then understand how to make the most of the various sources of manure at his disposal, and he will be enabled readily to save much, that, for the want of such knowledge, would otherwise be lost.
RURAL CHEMISTRY.

CHAPTER I.

COMBINATION—DECOMPOSITION—AIR—WATER.

1. The object of Chemistry is to determine accurately the properties of all natural substances, to study the changes which are going on in Nature, to find out the rules which govern them, and the manner in which these natural operations are influenced by circumstances.

2. In pursuing these inquiries, the chemist is obliged to proceed slowly and with great caution; it is quite impossible for him to predict beforehand the result of a new experiment; he must try it, and then, if it has been properly conducted, it always furnishes him with a new fact, for he is sure that, on repeating it in the same manner, he will obtain the same result. Hence Chemistry is purely an experimental science; every fact is the result of careful experiment, and every theory is deduced from the study of such facts. The greatest care must be taken to distinguish facts from theories; the former are well-established and
unquestionable truths, the latter are plausible conjectures, to which we are led by the attentive study of facts. When a chemist has made a number of experiments, or has observed many phenomena, he endeavors to ascertain the causes of the effects he has been studying. He selects the most probable explanation, and adopting it as a theory or view, to be confirmed or disproved by future experiments, proceeds to try in all possible ways the truth of his conclusion. By thus forming a theory he is enabled to arrive at the truth more easily than if he were merely to continue making experiments at random. It is by reasoning on the results of thousands of experiments that chemists have been enabled to reduce the science into a useful form, as they have thus been led to discover certain great leading laws, which govern all chemical changes or operations.

3. Nearly all the changes which are going on in Nature may be classed under two heads. The one kind of change is that which takes place when two substances come together which have, as it were, an attraction or affinity for each other. As a familiar example of what then happens, we may take the common process of soap-boiling. When an alkaline or caustic lye is boiled with tallow or fat, soap is formed. The alkali which is contained in the lye has an attraction for the fat; the two become thoroughly mixed, and combine or unite together, and form a new substance, quite different from either the fat or the alkali, which new substance is called soap.
4. This kind of action is quite distinct from simple mixture. When we mix together two substances—such as, for example, brown sugar and sand, no change takes place, however long they are kept together, or in whatever way they are treated, for they have no affinity or attraction for each other; and, therefore, if boiling water is poured upon the mixture, it will soon dissolve out all the sugar and leave the sand, and neither the sugar nor the sand will be at all altered by having been mixed.

5. When we mix two substances which have an attraction for each other, they are both changed, and the new substance formed by their union is quite different from either; and when two substances are thus united or combined together, they are not so easily separated as when merely mixed, because they require the exertion of some attraction more powerful than that which made them combine, to cause their separation. In the case of the soap just mentioned, the compound of fat and alkali does not resemble either of its components; it is different from the ley in not being caustic, and differs from the fat in being easily soluble in water.

6. It is a rule which holds good in all cases, that whenever two substances combine or unite together, and form a new substance, the properties of the new substance are quite different from those of either of its components; but when two substances are only mixed, the properties of the mixture are intermediate, or half-way, between those of its two components:
thus, in the mixture of sand and sugar, we may easily recognize both substances, for the characters of neither are altered by being brought together.

7. Another common case of affinity is observed when we slake quicklime. Quicklime has a strong affinity for water, and when it is wetted, it becomes very hot; the lime combines with a quantity of water, and when is has cooled, we find that the lime is much altered, having to a great extent lost its strong caustic properties, and become slaked, as it is termed. Here again we observe that the properties of the compound differ remarkably from its components. Dry caustic lime, in combining with water, forms a dry compound of lime and water, the water becomes solid, entering into the composition of a dry solid powder, whilst the lime no longer possesses the power of heating when water is poured over it, and has become less caustic (235).

8. It may perhaps seem as if these two examples of the change produced by attraction or affinity were processes of Art, and not of Nature. They will, however, serve as examples of what is going on in a great many natural operations; and as we proceed with the subject, it will be evident that this kind of change, by which two or more different substances unite and form one new substance, is exceedingly common throughout Nature.

9. The second kind of change which we shall have to consider, is that which goes on whenever anything decays. This change is quite opposite in its nature
DECOMPOSITION.

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to that which we have just been describing. It takes place whenever any substance is separated or divided into its component parts. Thus, to return again to the quicklime, which is made by burning chalk or limestone, we say that the chalk or limestone is decomposed, when, by burning or heating it in a very hot fire, whatever it contains which can be roasted out by fire, is driven off, and the lime only remains (118).

10. The decomposition of a substance is also effected when it is mixed with anything which has a very strong attraction for one of its components. Soap is made by the attraction which the alkali has for fat (3); but if we add to a solution of soap in water, anything which has a stronger attraction for the alkali than the latter has for the fat, we shall decompose the soap: there are many substances which have the power of doing this, but it is sufficient now to mention one. If vinegar is poured into a solution of soap, the soap is decomposed; the fat is separated and floats on the surface, and the vinegar combines with the alkali of the soap.

11. This kind of change is always going on when anything decays or putrefies, and therefore is of considerable interest in connection with manures; but in fact, combination and decomposition are almost always going on at the same time in most natural changes, for when a compound consisting of several different substances is decomposed, it is generally found that
these substances again combine together, one with another, to form various other compounds (105, 145).

12. Combination takes place whenever substances are brought together which have an affinity for each other, under suitable circumstances; chemical action then takes place, and a compound is formed. Decomposition of a compound is caused either by the influence of some external power, the presence of some substance capable of acting on one of the elements of the compound, or some influence able to weaken the chemical affinity which binds these elements together.

13. Combination is often modified and controlled in a very remarkable manner by circumstances, accordingly as they are favorable or unfavorable to the union of the substances brought together. Fine division, or any other method whereby the particles are enabled readily to mix or come in contact, generally assist combination. When two solid lumps are placed in contact with each other, they are only able to touch by a very few points, and hence in many cases do not combine even though they have an affinity for each other. If they are finely powdered and well mixed they are much more able to combine; and for the same reason if they are fusible, heat, by rendering them fluid, and thus enabling the particles more easily to mix, assists their combination.

14. A similar effect is produced by solution in water. In a common Seidlitz powder or saline
draught, we have two solutions mixed together, which are able to act on each other, but cannot do so in the dry state; when water is added they are brought thoroughly in contact, and chemical action at once takes place. In many cases combination is greatly assisted by heat, which exalts the chemical affinity that the substances have for each other (35).

15. Precisely in the same way decomposition is often curiously affected by various circumstances. Some compounds decompose spontaneously; they cannot be kept any time, and without any external influence they undergo decomposition; this is especially the case with many animal and vegetable substances (354). Decomposition is frequently caused by the influence of light (187, 295, 699), or heat (9, 119, 233). The decomposition of many compounds, likewise, is caused by the presence of particular substances. A great number of different organic substances are decomposed when a small quantity of some other substance in an active state of decomposition is mixed with them; this may be called decomposition by example; it is a very singular form of decomposition, and is termed fermentation (365). A similar change is sometimes caused by the mere presence of a particular substance, even though it is not itself undergoing any change whatever at the time (361).

16. It is a common saying that there are only four elements; air, earth, fire, and water; and many
people believe that all things are composed or made up of these four elements. This is very incorrect, because there are many substances which do not contain any of these so-called elements; and they are, besides, themselves compounded of many different substances. The term elements, in the sense in which it is used by chemists, means a certain set of simple substances, which, by combining or uniting together, form all the various matters that occur in Nature.

17. To return once more to the example of soap, we may say that the elements of soap are alkali and tallow; but then the question will arise, what are the elements of tallow and alkali? which can only be ascertained by chemical experiments. In this way, then, we may analyze, or as it were pull to pieces, different substances, till at last we find that we are unable to separate or decompose them any further, and the substances which then remain are called elements, or simple substances. It is possible that chemists may hereafter discover that some of the substances now called elements are really compounds, and of course it is impossible to prove that they are not so; all therefore that is meant by the term element, or elementary substance, is that chemists have not yet been able to prove them to be compounds.

18. There are upwards of sixty of these elements, but it will not be necessary to study the nature and properties of the whole series, because many of them are of very rare occurrence, and found only in small quantities.
19. The following is a list of all the substances which are at present considered as being simple, or elementary:

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<td>Erbium</td>
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<td>19.</td>
<td>Fluorine</td>
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<td>20.</td>
<td>Glucinum</td>
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<td>21.</td>
<td>Gold</td>
<td>42.</td>
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<td>22.</td>
<td>Hydrogen</td>
<td>43.</td>
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<tr>
<td>23.</td>
<td>Iodine</td>
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<td>24.</td>
<td>Iridium</td>
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<td>25.</td>
<td>Iron</td>
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<td>26.</td>
<td>Lantanum</td>
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<td>27.</td>
<td>Lead</td>
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<td>28.</td>
<td>Lithium</td>
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<tr>
<td>29.</td>
<td>Magnesium</td>
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<td>30.</td>
<td>Manganese</td>
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<td>31.</td>
<td>Mercury</td>
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<tr>
<td>32.</td>
<td>Molybdenum</td>
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<tr>
<td>33.</td>
<td>Nickel</td>
<td>54.</td>
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<tr>
<td>34.</td>
<td>Niobium</td>
<td>55.</td>
</tr>
<tr>
<td>35.</td>
<td>Nitrogen</td>
<td>56.</td>
</tr>
<tr>
<td>36.</td>
<td>Osmium</td>
<td>57.</td>
</tr>
<tr>
<td>37.</td>
<td>Oxygen</td>
<td>58.</td>
</tr>
<tr>
<td>38.</td>
<td>Palladium</td>
<td>59.</td>
</tr>
<tr>
<td>39.</td>
<td>Pelopium</td>
<td>60.</td>
</tr>
</tbody>
</table>

20. We will commence with those substances which are of the greatest importance, whether simple or compound, and gradually go through them, before entering upon the Chemistry of Vegetation. Foremost in importance, of the substances whose properties we are about to study, stands the Air.

21. We are too apt to think of the AIR as being merely empty space; we move about through it with-
out feeling any resistance, and, from its being invisible and totally unlike anything else we know, many forget its existence altogether. The fact is, that every part of the surface of the globe is surrounded by air, which floats on its surface almost like water.

22. It is easy to prove that the air is really a substance. When we try to squeeze together the sides of an inflated bladder, the mouth of which is tightly tied up, we feel that the bladder is full of something which resists the pressure; this something is the air which it contains, and which, though so easily displaced, or pushed aside, by anything moving through it, resists strongly any force applied to it when thus confined in a limited space; and if whilst we are pressing the bladder we prick a hole in it, the air then rushes out, we feel that the resistance is gone, and the sides of the bladder are easily squeezed together.

23. And again, when working a pair of bellows, it is the resistance of the air which we have to overcome by the force of the arms, which constitutes the labor of working the bellows; and if the nozzle of the bellows is stopped up, we presently find that it is impossible to go on working the bellows any longer, because having forced in as much air as it can hold, the natural tendency of the air to resist compression prevents any more from entering.

24. Although we are so forgetful of the very existence of the air, it is of the greatest importance to all our daily occupations, and even to life itself. Without air nothing could burn; we could have neither
fires nor lights; and, indeed, without air neither animals nor plants could live, for it is just as essential to the life of animals as it is to the growth of plants and the burning of coals and candles.

25. If a lighted candle is put into a large glass bottle, and the mouth of the bottle is then stopped up, the candle soon gets dim, and in a short time goes out: the air is no longer able to keep it alight. If we put a second lighted candle into the bottle it will go out immediately. Were a living animal substituted for the lighted candle, after living for a certain time in a confined portion of air, it would die, and a second animal placed in the air would immediately expire.

26. The question now will naturally arise: Is the whole of the air, then, spoiled or used up; and if it is, why does not fresh air enter the bottle and supply its place? The truth is, only a small portion, about one-fifth of the bulk of the air, is able to feed the flame of a candle; the remainder, which cannot feed flame nor the life of an animal, is of a different kind from the air which can; and we find that the common air which we breathe is a mixture of two kinds of air, or gas, as it is called by chemists;—the one kind, which we might call good air, which supports the life of animals, and is essential to the burning of fires and candles: and the other, or bad air, in which animals cannot live, and which immediately puts out fire and lights.

27. Chemists call the good air Oxygen, and the
bad air Nitrogen, or Azote: but we must not suppose that, because the nitrogen appears thus useless, it is really so; for it is, in fact, of very great importance, as we shall hereafter see. In the experiment just mentioned, of burning the candle in a large bottle, the oxygen is all combined with the elements of the tallow, for which they have a strong affinity, whilst the nitrogen is left unchanged because it cannot combine with them; for the same reason, also, it puts out a fresh-lighted candle plunged into it.

28. Oxygen, when obtained pure, and separate from any other substance, is a gas like common air in its ordinary characters, but remarkable for the very brilliant manner in which all kinds of combus-
tible matter burn in it. Oxygen may be breathed with safety, but it causes all the functions of the ani-
mal system to be carried on with great vigor and rapidity, so much so, that an animal breathing pure oxygen would be soon destroyed, from the very powerful effect which the gas would have on its organs. Oxygen has a strong affinity for most of the other elements, and combines with them to form a numerous and important series of compounds.

29. Nitrogen, though it resembles oxygen in ap-
pearance, yet differs from it very remarkably in chemical characters; it extinguishes flame, and cannot support the combustion of any substance; it is irrespirable, suffocating animals if they attempt to breathe it pure, and seeming to have very little affinity for other elements; at least under ordinary air-
cumstances, it shows very little tendency to combine with them; under particular circumstances, however, it does form compounds (147, 163), and some of them are very curious and important. Common air consists of one part of oxygen to four parts of nitrogen: it is a mixture, not a compound.

30. When a candle burns, it gradually disappears; it grows shorter and shorter, and at last, when all the tallow is burnt, the candle goes out; but we must not therefore suppose that it is utterly destroyed. A change has taken place; the tallow, or rather its elements, have combined with the oxygen of a portion of air, and two new compounds, one of which is a gas or kind of air, are produced. If we put a piece of salt into water, it will get less and less, and at last will disappear altogether, having wholly dissolved; but the salt is not destroyed, it is only dissolved in the water.

31. Now we may compare the burning of a candle to dissolving a piece of salt; for all the solid matter of the candle remains diffused throughout the air, after it is burnt, just as the salt remains dissolved in the water; but with this difference, the salt is dissolved in the water, but not combined with it. The elements of the tallow are dissolved in the air, but they have combined with a quantity of oxygen, because they have a strong affinity or attraction for it. If the solution of salt is left for some time in a warm place, the water evaporates, and we get the salt again unchanged; but in the case of the candle its elements
have combined with oxygen, and they cannot be separated again from it except by the action of something which, having a more powerful attraction for the oxygen than it has for the elements of the candle, causes it to relinquish them.

32. There are substances which have sufficient attraction for the oxygen to effect this: we cannot get back the tallow, it is true, but we may obtain its elements, or the simple substances of which it was composed. What has been said with regard to the burning of a candle is equally applicable to the burning of wood, coal, or in fact any combustible matter. In all ordinary cases they burn in consequence of their affinity for the oxygen of the air, and they are never destroyed when burnt, for their elements may always afterwards be found combined with oxygen, in the air in which they have been burnt.

33. It must also be remembered that when the candle goes out for want of air, it does not do so because all the oxygen is burnt, but because the elements of the candle having combined with all the oxygen of the air, or, as it were, saturated it, there is no more free oxygen left to keep up the combustion of the candle.

34. The changes occasioned by chemical action frequently proceed slowly and quietly, but in many cases, and especially when substances combine together which have a strong affinity for each other, a great deal of heat is given out. Sometimes, as soon as two substances are brought together, they combine di-
rectly, and become very hot; this is the case in the slaking of lime; but it most usually happens that the mere bringing together of two substances, even though they have an affinity for each other, is not sufficient to cause them to combine.

35. In these cases combination cannot take place until the substances are heated up to a certain point. Thus, charcoal has a strong affinity for the oxygen of the air, yet it cannot combine with it whilst both are cold; but as soon as a part of the charcoal is heated redhot, combination commences, and this very act evolves so much heat that the surrounding parts of the charcoal soon begin to burn, and thus the combustion, or combination of the charcoal with oxygen, continues and increases, until either the charcoal is all burnt, or the oxygen in the surrounding air is saturated with carbon, and therefore unable to cause the combustion any more.

36. The common operation of lighting a fire is a daily illustration of this. The fuel contains carbon, or charcoal, ready to combine with the oxygen of the air, but unable to do so, until, by applying a light to it, we heat a portion up to the point required to commence combination: after which the heat given out by the chemical action going on, keeps it alight, and causes the combustion to spread to the surrounding fuel.

37. The atmosphere is composed chiefly of two different gases, called oxygen and nitrogen: but besides this, it also contains a small quantity of a
third gas or kind of air, which is not simple, like oxygen and nitrogen, but a compound of charcoal (by chemists named CARBON) with oxygen, and called CARBONIC ACID GAS (108).

38. It is known that all things containing carbon will produce a quantity of this gas whilst burning; and hence we can have no difficulty in accounting for its presence in the air. Indeed, we might at first suppose that it must be always increasing in quantity; this, however, is not the case, for we always find exactly the same quantity in any portion of air that we analyze. The cause of this is, that all plants contain substances which have a very strong affinity for carbon, but which cannot combine with it in its solid forms, because they are unable to come in contact with it; but which, when the carbon has combined with oxygen and become a part of the air, are able, in consequence of their having a more powerful attraction for it, to seize upon the carbon of the carbonic acid gas thus diffused throughout the air and cause it to relinquish the oxygen, with which it was previously combined (697, 708).

39. These facts show us a new use of plants, for we learn that the objects which we have only admired for their beauty, or valued for their utility as producing articles of food; that even weeds themselves, and things we usually consider as wholly useless, are all constantly, by the agency of attraction or chemical affinity, decomposing carbonic acid gas, and thus keeping the air in an uniform and healthy state (745).
40. The air then always contains a regular proportion of carbonic acid gas, which is constantly produced by the burning of combustibles, and in many other ways, and as constantly decomposed by the action of plants. As we are now only considering the properties and nature of the air, we will, for the present, pass over further consideration of carbonic acid gas, to which we shall shortly return (108) when studying the nature of carbon, and merely mention now that it is of the greatest importance to the life of plants, being the principal source from whence they derive the carbon necessary for their growth.

41. The air always contains dissolved in it some water, or rather vapor, which varies in quantity according as the air is hotter or colder. When it is hot, a larger quantity of water is evaporated or converted into vapor, and dissolved in the air, which in consequence becomes more damp; whilst, on the other hand, when the air becomes cold, the vapor in the air is condensed, returning to the state of water, and the air becomes drier.

42. This, of course, is modified according to circumstances; thus, in dry barren countries, where the ground contains but little moisture, the air, when it becomes hot, remains comparatively dry; whilst in moist or swampy countries, under similar circumstances, the air becomes damp from the abundance of vapor given off; and thus some of the principal differences of climate depend mainly upon the quantity of water dissolved in the air.
43. The solution of water in the air may easily be seen, by observing the steam issuing from the spout of a teakettle. When the water boils strongly, and there is a large volume of steam coming out of the spout, we observe that just where it comes out, the steam is almost invisible; at a little distance it becomes white and cloudy, and when it gets further out into the air it soon disappears and is again invisible. The reason of this is, that hot steam is quite colorless and invisible; like air; and it only becomes apparent to us when it is partly cooled by rushing out into the cold air, and therefore is beginning to condense, and it would fall to the ground in a shower of little drops like rain, if it were not dissolved and carried away by the air, as fast as it issues from the teakettle. When a large quantity of steam is quickly cooled, as in escaping from the funnel of a steam-boat, it is condensed, and falls in the form of water.

44. The quantity of moisture in the air is also rendered apparent to us, whenever a cold substance is exposed to it; this cools the vapor in the air so much, that it is condensed and appears again as water, in little drops on the cold surface: thus a bottle of cold water brought into warm damp air, speedily becomes covered on the outside with dew, or water thus condensed from the air.

45. The substance next in importance to air is water, which exists naturally in three different states: namely, in the solid state as ice; in the fluid state in its ordinary condition; and lastly, as vapor
or steam. These three states of water are familiar to every one, but few are aware what causes the great difference between them.

46. When ice is placed before the fire, or exposed to the sunshine, or in any other way warmed, it absorbs heat, it melts and becomes water; and when water is heated, it assumes the form of steam or vapor. The difference between these three forms of water is entirely caused by the quantity of heat they contain; and we may truly say that water is a compound of ice and heat, and that steam is a compound of water and heat.

47. Although this seems very like chemical action, it is really quite different, and must not be confounded with it. Chemical action can only take place between material substances, or those that have weight; now heat is not a substance—it is not a thing we are able to weigh, like all the chemical elements, and consequently when it combines with any substance it only alters the appearance and outward characters of that substance, but does not at all change its chemical properties or nature.

48. When heat is thus combined with a substance, it is said to be latent (hidden), which means that it is not sensible to the feel. This will be easily understood from a very simple experiment. If we put some water in a kettle on the fire, we find that it will soon begin to feel warm to the hand if immersed in it; the warmth which we then feel is called free or sensible heat: but if we put some ice into the kettle in place
of water, it will not become warm so soon; the ice melts, but the water thus formed will remain ice-cold until the whole of the ice is melted, because all the heat supplied to it by the fire is absorbed or combined with the ice in melting; and therefore as the heat so absorbed does not make the melting ice any warmer, it is called latent. When all the ice is melted, the water will begin to get warm.

49. In the same way heat is absorbed, or rendered latent, when water is converted into steam. If a kettle full of cold water is placed on the fire it rapidly becomes hot until it begins to boil; but as soon as that is the case it remains constant at the same temperature. The fire of course continues to give out as much heat as it did before, but the water does not become any hotter; the only change is that a small portion of it is converted into steam, and this steam is not apparently any hotter than the boiling water itself was. All the heat of the fire, therefore, becomes latent, and is combined with the water in thus changing it into steam.

50. When steam is condensed, all that heat which became latent during its formation, is given out again in the free and sensible form. This fact is well shown in all stills, in which we see how large a quantity of cold water is necessary to cool and condense a comparatively small quantity of any steam or vapor. If a gallon of water is converted into steam, the steam formed will not be sensibly any hotter than boiling water, yet it contains so much latent heat that, if it is
condensed into the liquid form again, we shall obtain a gallon of boiling hot water, and heat enough will be given out during its condensation to raise seven gallons more of cold water to the boiling point.

51. The very large proportion of heat which steam therefore contains in this latent state, shows why steam is such an excellent means of conveying heat about from place to place, as in the arrangements for warming buildings, heating coppers, &c. It is quite as hot as so much boiling water, and, in addition, has nearly seven times as much heat thus stored up in the latent form, which becomes sensible, however, as soon as it is condensed.

52. The general effect of heat upon substances, whether solid, liquid, or gaseous, is to expand or make them larger; thus whenever we heat a portion of water or any other fluid, it increases in size: upon this fact the construction of the common thermometer depends, which consists of a bulb and tube containing a certain bulk of mercury, or quicksilver; when this is heated it becomes larger, and when cooled, the mercury shrinks, or occupies less space than it did before.

53. When a substance expands, or becomes larger, it of course becomes lighter. If ten measures are expanded by heat to eleven, it follows that ten measures of the heated substance must weigh one eleventh less than ten measures of it when cold. For example, air, when heated, becomes lighter, and consequently rises, because, becoming larger, it weighs comparatively, bulk for bulk, less than the cool air around.
54. Air, like water and solid substances, has weight or gravity; it presses downwards towards the centre of the earth; and, as the bulk of the air is very great, the whole weight of it pressing on the surface of the globe must be enormous. It is important to distinguish between the weight of air, and the weight of the atmosphere. By means of a light glass globe, an air-pump, and a good pair of scales, we can readily ascertain the weight of a cubic foot of air, and we find that, under ordinary circumstances, it weighs thirty-one grains.

55. The weight of the atmosphere is, however, the accumulated weight of many thousand cubic feet of air, for the air surrounds the earth to a height of many miles, floating upon its surface, and pressing downwards in consequence of its weight or gravity. The weight of the column of air which rests upon any given space or surface, may be known by the use of the barometer: and is found to be about two thousand pounds on the square foot.

56. The barometer consists of a glass tube nearly a yard long, closed at one end, filled with mercury, and inserted in a cup of the same heavy liquid. The height which the mercury stands in the tube, is an exact measure of the weight of a column of air of the same size as the diameter of the tube, but as high as the whole atmosphere. The mercury in the tube being very heavy, presses downwards, and tends to fall in the tube; but the air without, which also presses downwards, resting on the mercury in the cup,
exactly counterbalances the weight of the mercury in the tube, and so keeps it from falling.

57. When from any circumstance the weight of the atmosphere is for a time diminished, it counterbalances a less weight of mercury, and the column of mercury in the tube of the barometer falls; on the other hand, when the weight of the air increases, the mercury rises. These fluctuations in the weight of the air are constantly going on with the changes of the weather, and consequently the barometer is a most valuable instrument, because it renders evident to us the changes in the air, which being invisible we should not otherwise know of.

58. The higher we ascend in the air, either by going up a hill or by means of a balloon, the less weight of air shall we have above us, and consequently the shorter will be the column in the barometer. Hence the use of that instrument in measuring heights. The mercury falls about one inch for every thousand feet as we ascend from the level of the sea.

59. We do not feel the weight of the air at all, because it presses equally in all directions, and completely surrounds us. The weight of a column of air resting on a table four feet square, is about fourteen tons; but this weight does not press upon the table, it is met by a corresponding pressure upwards, from the air below the table, due to the weight of all the surrounding air.

60. In moving through the air, therefore, we merely displace it, or move it aside; but as we do not disturb
this equilibrium, or the equal pressure in all directions, we do not feel its weight. On plunging the hand into a pailful of water, we feel very little of its weight, only in fact that of the small portion which we displace, because it is much heavier than air is; but if we attempt to lift up the whole pailful, we then feel the weight of the whole bulk of water.

61. Flame seems to ascend in the air; this is not because heat has any tendency to ascend, but because it expands the surrounding air, makes it larger, and therefore specifically lighter than the cold air is, and the latter, therefore, displaces it and causes it to rise. Hot air, though it rises in the air, nevertheless has weight, though it is not so heavy as cold air. A piece of cork falls through the air towards the surface of the earth, it has weight; yet the same piece of cork rises upwards through water till it reaches the surface. As the cork does not fall downwards in water, but rises upwards, so heated air ascends in the atmosphere; not because it has no weight or gravity, but merely because, bulk for bulk, it has less weight than the surrounding cold air. It is for precisely the same reason that a balloon rises in the atmosphere.

62. It is, in fact, a general rule that, when substances are heated, they expand; when water is heated, it becomes larger and lighter, and consequently rises through the cooler portions above it. Just the reverse of this happens when substances are cooled; they then become smaller and heavier.

63. There is one remarkable exception to this rule
in the case of water; when water is cooled it contracts, and this goes on till very near the freezing point; but then the water begins to expand, and in place of continuing to contract, as all other liquids do, it becomes larger; this leads to a very important result in Nature. When the air above the surface of a lake or pond becomes cold, as towards the end of autumn, it gradually cools the surface of the water; the upper part becoming cold, shrinks, and consequently becomes heavier, it therefore sinks through the warmer water. This circulation or gradual sinking of the cooled water goes on, if the air continues cold, until the whole of the water is very near freezing, but then it stops; because if the surface still goes on cooling, the water begins to expand, becoming larger again, and consequently lighter; the surface, therefore, gets colder and freezes, whilst the lower part of the water remains considerably above the freezing point. If it were not for this curious fact, water would continue to become colder, until the whole of it froze together.

64. In passing from the liquid to the solid state, some substances contract, such as melted lead, for example, whilst others expand; thus ice-cold water, in freezing, expands very considerably, and therefore ice is even lighter than the water on which it is formed. It is for this reason that ice floats on water, and if the ice did not expand in forming, the curious fact just mentioned would not prevent lakes and ponds from freezing entirely, because the ice, if it contracted in
forming, would then sink through the water to the bottom, and thus soon cool the whole mass of water.

65. This expansion of water in the act of freezing, takes place with immense force, giving rise to the bursting of water-pipes and vessels full of water, in cold weather. It is often supposed that this effect is occasioned by the thaw, and not by the frost. This is a mistake; the mischief is caused by expansion at the moment of freezing, though we only discover it on the approach of warm weather, when the ice begins to melt. Another, and very important natural result of this power, is the disintegration or breaking up of rocks, stones, and soils by frost, during winter. A few drops of water, in freezing, are able to break asunder the hardest rocks, and this effect year after year, gradually destroys them, causing them to crumble down into powder (647).

66. It has already been stated that water is not an elementary or simple substance; it is a compound, and consists of two gases. This fact at first seems incomprehensible, for we can hardly believe it possible that a hard and solid substance like ice, or a weighty fluid like water, is composed of colorless and invisible gas. The difficulty, however, greatly diminishes when we remember that heat alone, without adding anything to the weight of ice, converts it into water, and that a little more heat will convert that water into an invisible colorless vapor; for, as has already been said (43), pure hot steam is quite invisible, and only becomes visible to us when partly
condensed by the contact of the colder air, which deprives it of the heat necessary to keep it in a state of vapor (43, 73).

67. The consideration of these facts makes the composition of water appear far less wonderful; for we have little difficulty in believing that steam is composed of two gases, and we know that steam, water, and ice, are, chemically speaking, the same.

68. One element of water is oxygen gas (28), that part of the air which is so essential to life and combustion: it constitutes eight-ninths of the weight of ice, water, and steam. One thousand parts of water, therefore, consist of

<table>
<thead>
<tr>
<th>889 parts of Oxygen</th>
<th>111 &quot; Hydrogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>1000 &quot; Water.</td>
<td></td>
</tr>
</tbody>
</table>

69. The other element, or the remaining one-ninth, is called hydrogen gas, or inflammable air, because it is very combustible, being the basis of the common coal gas used for lighting the streets, and entering into the composition of the inflammable air or firedamp of mines, and many other combustible substances (82).

70. Water is not, like common air, a mere mixture of two gases: it is a compound, and therefore is quite different in its properties from either of its two elements. The very inflammable gas, hydrogen, having combined with a certain quantity of oxygen, which
is the great promoter of combustion, forms water, a compound which we always regard as the greatest enemy to fire or combustion.

71. The purest kind of water which occurs naturally is rain-water, for all others, such as spring, river, or sea-water, are more or less contaminated or rendered impure by substances dissolved in them. Thus sea-water contains, along with other matters, a large quantity of common salt, which in some places is procured from it by exposing it in shallow pits to the heat of the sun: this causes the water to evaporate, and leaves the salt behind.

72. The composition of sea-water from different parts of the world is found to vary slightly. The following table shows the composition of 100,000 parts of sea-water from the English Channel. (Schweitzer.)

<table>
<thead>
<tr>
<th>Substance</th>
<th>Parts of 100,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>96,474</td>
</tr>
<tr>
<td>Common Salt</td>
<td>2,706</td>
</tr>
<tr>
<td>Chloride of Potassium</td>
<td>76</td>
</tr>
<tr>
<td>Chloride of Magnesium</td>
<td>366</td>
</tr>
<tr>
<td>Bromide of Magnesium</td>
<td>3</td>
</tr>
<tr>
<td>Sulphate of Magnesia</td>
<td>229</td>
</tr>
<tr>
<td>Sulphate of Lime</td>
<td>140</td>
</tr>
<tr>
<td>Carbonate of Lime</td>
<td>3</td>
</tr>
<tr>
<td>Iodine</td>
<td>traces</td>
</tr>
<tr>
<td>Ammonia</td>
<td>traces</td>
</tr>
</tbody>
</table>

100,000

73. Water may be artificially purified by distillation; when heated and raised into vapor, all the
impurities are left behind, and accordingly condensed steam is perfectly pure water; there are numerous contrivances for thus purifying water. The common still, which consists of a vessel to generate steam in, and a pipe, passing through a tub of cold water, to condense the steam, is a familiar example.

74. This explains why rain-water is much purer than other sorts of water, because when the heat of the sun evaporates water from the surface of the earth, all the impurities which it contains are left behind; and of course when this vapor is cooled and falls down in the form of rain, it must be very nearly pure.

75. Springs, which rise from the ground, always contain earthy matters dissolved in them, which vary in nature and quantity, with the soil through which the springs rise. The presence of these impurities in water in any quantity gives to it that peculiar character which is termed hardness. Sometimes springs contain a small quantity of iron or sulphur, and other substances, which constitute the many varieties of mineral waters. These matters, like the more common earthy impurities, are all derived from the beds of stone, sand, or clay, through which the springs rise; because the source of all springs is rain-water, which, falling pure from the clouds, becomes contaminated by filtering through the earth, and collects in holes and cavities, or porous beds of sand, constituting springs and wells.
76. The quantity of saline and earthy matter in spring-water varies from about 20 grains to 1800 grains in the gallon; when above 100 grains per gallon, it constitutes a mineral water. The average quantity in ordinary spring-water is from 20 to 80 grains. The most common salts are Sulphate and Carbonate of Lime (230), Sulphate, Muriate, and Carbonate of Potash and Soda.

77. Thames-water contains usually from seventeen to twenty-four grains of earthy and saline matter per gallon, and of this at least fifteen grains consist of carbonate of lime. The same quantity of New River water contains about nineteen grains of solid matter, and that of the River Lea nearly twenty-four grains. The chief constituent in both these waters is also carbonate of lime.

78. The proportion of solid matter is almost always greater in well-water than in that of rivers. A great number of other substances besides those just mentioned are occasionally found in mineral springs; amongst these are silica, alumina, oxides of iron, and manganese, salts of baryta, strontia, magnesia, ammonia, &c.

79. The presence of phosphoric acid in some waters has recently been discovered (194). The following analysis of the deep well-water from below the London clay, shows the presence of a considerable quantity of phosphoric acid. Ten gallons of the water contained five hundred and sixty-four grains of saline matter, consisting of
Hydrogen.

80. Besides these saline and earthy substances, water always contains atmospheric air dissolved in it. This is essential to the life of fishes, and to the growth of water-plants, which could not exist if they were not thus supplied with common air.

81. Water is essential to the existence of all plants and animals: it constitutes a large proportion of all animal and vegetable substances, it is the principal component of the blood of animals, and the sap of plants, and is of the greatest importance, as being the means of introducing into their systems many soluble matters, necessary for their healthy growth.

82. Hydrogen, the inflammable element of water, is a substance of considerable interest; it is true, it is never found in Nature in a pure and separate state, but its compounds are abundant, and some of them very important; when pure, hydrogen is an invisible transparent gas, like the air; very combustible, burning readily when once inflamed, and remarkable for being so much lighter than common air, that a thin bladder, filled with this gas, would rise through the
air in the same manner that a bubble of common air rises through water. Balloons are sometimes filled with pure hydrogen gas, but more commonly carburetted hydrogen is employed, which, as it consists in great part of hydrogen, is much lighter than common air.

83. The most important of its compounds are water, which is formed by its union with oxygen; ammonia, a gas which it forms by combining with nitrogen; and carburetted hydrogen or coal-gas, an inflammable gas consisting of hydrogen and carbon. The two latter will shortly come under notice (131, 148).

84. As hydrogen is the lightest gas known, it is often employed by chemists as a standard of comparison in expressing the relative weight of all other gases. By weighing a thin glass globe filled with hydrogen, and then having pumped that gas out of it by means of an air-pump, and filled the globe again with any other sort of gas, a second weighing gives us the comparative weight or specific gravity of the gas as compared with hydrogen. Suppose, for example, that the globe held exactly ten grains of hydrogen and twenty-five grains of the second gas, then it is plain that the latter is twice and a half as heavy as hydrogen is; or, that taking equal volumes of both, that of the gas would weigh twice and a half as much as the hydrogen would. We should say, then, that the specific gravity of that gas was $2\frac{1}{2}$ or 2.5, taking hydrogen as the standard of unity.
85. The following table shows the weight in grains of several gases which a bottle containing one hundred cubic inches, or about three pints, would hold; and also the specific gravity or relative weight of the same gases compared to hydrogen, and also to common air, as a standard:

<table>
<thead>
<tr>
<th>Gas</th>
<th>Weight in grains of 100 cubic inches</th>
<th>Specific gravity compared to hydrogen, taken as 1000</th>
<th>Specific gravity compared to air, taken as 1000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen</td>
<td>2</td>
<td>1,000</td>
<td>69</td>
</tr>
<tr>
<td>Common Air</td>
<td>31</td>
<td>15,200</td>
<td>1,000</td>
</tr>
<tr>
<td>Oxygen</td>
<td>34</td>
<td>16,000</td>
<td>1,109</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>30</td>
<td>14,000</td>
<td>971</td>
</tr>
<tr>
<td>Carbonic Acid Gas</td>
<td>47</td>
<td>22,000</td>
<td>1,520</td>
</tr>
<tr>
<td>Ammonia</td>
<td>18</td>
<td>8,500</td>
<td>589</td>
</tr>
<tr>
<td>Chlorine</td>
<td>76</td>
<td>36,000</td>
<td>2,470</td>
</tr>
</tbody>
</table>
CHAPTER II.

CARBON—NITROGEN—SULPHUR—CHLORINE—PHOSPHORUS.

86. We now come to the consideration of an elementary or simple substance, which has been already more than once alluded to, namely, carbon or charcoal. Carbon is the name applied to the pure element, but common charcoal is so nearly pure, that we may consider the two words as meaning the same thing; it is an essential part of all kinds of fuel or combustible substances, during the burning of which it combines with oxygen, and forms carbonic acid gas, the substance before adverted to as always existing in the atmosphere (37).

87. The forms of carbon which we are accustomed to see, are almost all black, like common charcoal; but this is not the case with all the varieties of carbon, for we know that the brilliant transparent gem called the diamond, is really pure carbon, there being no chemical difference between that gem and common charcoal.

88. There are many substances in Nature which exist in two or more different states, presenting very dissimilar appearances, but being really chemically
the same. Thus, chalk and marble are very different looking substances, but they are composed of precisely the same elements; the difference between them is not caused by heat, like the difference between the three states of water, but is wholly of a mechanical nature. The particles composing a piece of chalk are much smaller than those composing a piece of marble, which are in fact compound particles, consisting of many joined together, and hence a piece of marble appears made of many little grains, whilst chalk is composed of particles so small that we are unable to distinguish them, and therefore appears to be a uniform substance.

89. The power which holds together the little particles composing the piece of marble or chalk, or any other substance, is called cohesion, and this power varies in strength in different substances; thus it is far stronger in marble than in chalk, and hence a piece of marble requires a much harder blow to break it, than a piece of chalk. In the same way, we say that the particles composing the diamond are held together more firmly by cohesion than the particles composing a piece of charcoal. Cohesion is, of course, quite independent of chemical attraction, for it holds the different particles of a substance together, constituting its mechanical strength; whilst chemical affinity binds together particles of two different substances, forming a compound substance, but does not in any way affect the strength of the compound to resist mechanical force applied to it.
90. Under common circumstances, carbon is one of the most unchangeable things we know; neither air, water, nor any of the substances commonly found in nature, have any action upon it; and hence the practice of charring the lower parts of wooden posts, which are intended to be driven into the ground; the coat of charcoal thus formed, protects the wood from decay for a much longer period than if not charred (882). When, however, carbon is heated, its chemical affinity for oxygen is greatly increased, and it no longer appears to be the unchangeable substance which it is whilst cold.

91. Carbon has been already stated to be a necessary element of plants, which, though so various in form and color, are nevertheless composed of very few elements. They consist almost wholly of oxygen, hydrogen, nitrogen, and carbon, which, by combining in different proportions, form all the principal parts of both plants and animals.

92. The greater number of vegetable substances consist wholly of oxygen, hydrogen, and carbon; whilst animal matters mostly contain, in addition to these three elements, a quantity of nitrogen: some of the compounds of plants, however, resemble animal matter in containing nitrogen. When wood is charred, or decomposed by heat, its elements are separated from each other; the hydrogen and oxygen combine together and form water, whilst the carbon is left behind. When wood is burnt in the open air both its hydrogen and carbon combine with oxygen, caus-
ing flame; the combination of carbon with oxygen proceeds slowly and steadily, the carbon continuing to glow until all is consumed; hydrogen, on the other hand, being a gas, mixes and combines with oxygen more rapidly, burning at once, with a flame.

93. All organic substances burn with a flame, and this alone is a proof that they contain hydrogen, because under ordinary circumstances the other elements of vegetable matter could not cause flame; and whenever the flame is bright, we are sure that it contains a good deal of carbon, for the flame of pure hydrogen is very pale indeed, and the brightness of a flame, such as that of a candle, is entirely due to the carbon contained in the tallow, which is burned at the same time with the hydrogen, both combining with oxygen.

94. As the products of the combustion of a candle are carbonic acid gas and water, it would be natural to expect that a cold substance held over the flame of a candle would take from the vapor of water thus formed, the heat necessary to its existence in the state of vapor, and consequently condense it into the fluid form. This is an experiment which may very easily be made, for we have only to hold a cold glass at a little distance above the flame of the candle, and we shall soon find it lined with a fine dew of water condensed in this manner.

95. There are several important facts to be observed connected with the combustion of a lamp or candle. The wax, tallow, or other combustible sub-
stance, is undergoing decomposition; and its elements, the combustible substances of which they are composed, are combining with the oxygen of the surrounding air to form new compounds; heat is necessary to both of these two changes, and this heat is evolved by the very changes themselves.

96. When a candle is lighted a portion of the fuel is melted, drawn up by capillary action through the wick, and decomposed and converted into combustible vapor; this vapor burns as its elements combine with the surrounding oxygen, and the heat which results from this chemical action renders the process continuous, by causing the decomposition of fresh portions of the tallow, &c., and the consequent production of more combustible vapor.

97. In ordinary flame, therefore, several things are necessary; air and heat are of course quite essential. The heat evolved by the flame itself, causes a circulation of air (61), provides a sort of natural ventilation, and insures a constant supply of fresh air to the burning vapors; on the other hand, this circulation of air, by bringing fresh oxygen to the burning combustible vapors, causes the evolution of heat enough to insure the combustion of the fresh vapors, which are about to be given out.

98. Every one knows well how necessary fresh air is to perfect combustion; the importance of heat is not so evident, though it is really quite as essential. The flame of a candle may be extinguished by a coil of wire, or by bringing a piece of cold metal in
contact with its outer edge, and is frequently "snuffed out" merely in consequence of its being cooled. A candle, moved quickly through the air, flares and smokes, and its combustion becomes imperfect, because it is cooled.

99. It is a necessary consequence of this fact, that flame cannot pass through a piece of wire-gauze, and the miner's safety-lamp merely consists of a lamp surrounded with a cylinder of wire-gauze. A lamp thus protected, may safely be taken into an explosive mixture of air and carburetted hydrogen, the flame may be put out, the explosive gas may burn within the cylinder, but the flame cannot pass through the wire gauze to set fire to the explosive atmosphere without, because it is too much cooled by contact with the wire gauze.

100. It is evident that the combustion of a flame can only take place on its outside, or at that part at which it is in contact with the air, and, as a necessary consequence of this, it follows that flame is hollow. It is only the outside of the flame of a candle that gives out light, the inside is dark, because no combustion is there going on. In a large flame this is easily shown, because, as no combustion is going on, no heat is given out, and consequently the inner part of such a flame is comparatively cool; a small piece of wood or paper may be held in the centre of such a flame, and it will hardly be singed, or a small spoon containing a portion of gunpowder may be
placed in the centre of the flame, and the powder will not be fired.

101. The wick of a common candle requires constant snuffing, because, being in the very centre of the flame, it is in fact deprived of two necessary conditions to combustion, namely, heat and fresh air; composition candles with twisted wicks, on the other hand, do not require snuffing, because, as the candle burns, the wick constantly twists outwards towards the edge of the flame; it is thus brought into the hottest part of it, and at the same time is supplied with fresh air, which causes it to burn.

102. Smoke is merely combustible matter which is unable to burn, because it cannot come in contact with free oxygen, or does not reach it till it is too cold to burn. The production of smoke from a fireplace or furnace, may always be prevented, and in so doing a loss of heat is obviated. A great deal is often said about the burning of smoke; this is a very difficult thing; it is far more easy to prevent its formation than to burn it when once formed. It requires, however, constant care and attention to prevent the production of smoke; there is no reason to doubt that at least nine-tenths of the dense black smoke which contaminates the air of large towns might be altogether prevented.

103. Carbon is unable to combine with oxygen at common temperatures; it requires to be heated before it can enter into combination with that substance; but when a compound substance containing carbon is
exposed to the air, it usually happens that, if the other elements which it contains combine with oxygen, the carbon also is then able to combine with oxygen, and forms carbonic acid gas.

104. Thus when a plant dies and decays, its elements separate and form new compounds; but the carbon is not set free in the form of black charcoal—it is slowly combined with oxygen to form carbonic acid gas (767).

105. Hence we see that decay is very like burning, similar effects being produced by both; only that the change which is effected by combustion in a short time, and accompanied with a great deal of heat, is very slowly and gradually effected by decay, very little heat being at the same time evolved. In both cases carbonic acid gas is produced.

106. This explains one great use of decaying vegetable substances in manures; they, of course, contain carbon, which is slowly combining with oxygen, and therefore always supplying the growing plants with carbonic acid gas, which is essential to their growth, as they, being able to decompose it, thus obtain carbon (698, 708).

107. Carbonic acid gas is produced in large quantity by the breathing of animals. A constant supply of fresh air is requisite for the support of life, and we know that if an animal is prevented from breathing or inhaling fresh air, it will very soon be suffocated. The chemical action which goes on in the lungs of an animal, is just the reverse of that which
takes place in the organs of plants: in the latter case, carbonic acid gas in the air is decomposed, and the carbon which it contained is appropriated by the plant; whilst in the lungs of an animal, carbonic acid is formed; for the oxygen of the air is found, on being expelled from the lungs, to have combined with carbon, and become converted into carbonic acid gas. In fact, the process of breathing is very similar to that of combustion, the same results being produced in either case (605).

108. Carbonic acid gas is invisible and transparent like common air, slightly soluble in water, and remarkable for being much heavier than air, and for extinguishing flame and destroying animal life; it is called an acid, although it certainly is not acid or sour to the taste, like vinegar and the other common acids we are acquainted with: it will, therefore, be proper, before proceeding any further, to explain why it is called an acid, and in fact what is meant by that term.

109. There exists in Nature a numerous class of substances which are called bases; amongst which are potash, soda, and lime, &c. Now an acid is a substance that has a strong affinity for these bases, and which in combining with one of them forms a neutral compound, possessing none of the properties of either.

110. Some acids are exceedingly sour, and very corrosive substances, like oil of vitriol, and aquafortis; but when poured upon a base, such as potash or soda, they combine with it directly, and both the
NATURE OF SALTS.

67

acid and the base lose their caustic and corrosive qualities.

111. These compounds of acids and bases are usually called SALTS, or saline compounds, and are very numerous; for there are many acids, and many bases. Most of the acids combine with bases in two or three different proportions, forming sub-salts, neutral salts, and acid, or super-salts. In the sub-salts, the proportion of acid is not enough to more than half neutralize the base; in the super-salts it is twice as much as is required to neutralize the base; and in the neutral salts, as the name indicates, the acid and base are combined in single proportionals, or perfectly equal quantities (139, 141).

112. There is also a great number of salts called double salts: these are compounds of two similar salts; thus there are double sulphates, like alum, the sulphate of alumina and potash (257); and double phosphates, like the phosphate of ammonia and magnesia (253). These double salts are distinct compounds of the salts of which they consist, and possess perfectly different properties from either of their constituents; they may generally be made by merely mixing together solutions of their two components. The number of double salts is very considerable, but there is also an immense number of salts which do not thus combine together.

113. Acids have a curious property of changing certain vegetable colors. The greater number of vegetable blue colors are by acids changed to red,
and this property is therefore made use of by the chemist, to detect the presence of a free acid; for this power of acid ceases immediately, when they are combined with bases: because their powers are then neutralized (110).

114. A very common illustration of this change of color may be seen in the pickling of red cabbage. Every one knows that red cabbages, as they are called, are really dark purple or blue, whilst growing, and they only become red by the action of the acid vinegar employed in pickling them; the same effect would be produced if any other acid was employed.

115. If now we add to some cabbage thus reddened by acid, a little alkali, either potash, soda, or ammonia, or indeed a portion of any base, the color will soon be restored to its original blue; because the acid is neutralized by the base. And if the base employed be one of the three alkalies, or their carbonates, such as common pearlash, which is the carbonate of potash, the solution of cabbage will become green, because free alkalies, and likewise their carbonates, have the power of turning vegetable blues green.

116. Alkaline solutions have also the property of turning certain vegetable yellows red, such, for example, as common turmeric; these tests, as they are termed, are very easily applied, and papers stained with blue or yellow vegetable colors, are consequently most useful indicators of the presence either of free acid, or free alkali, in a solution.
117. Although carbonic acid, being a gas, is not perceptibly sour to the taste, like the strong acids just mentioned, it nevertheless combines with bases, and in so doing neutralizes, or at least weakens to a considerable extent, their caustic properties; but as it has a far less powerful attraction for bases than most other acids have, it is very easily expelled from compounds containing it, by the action of another acid, which seizes upon the base, and sets the carbonic acid gas at liberty.

118. We have already spoken of the conversion of chalk into lime by heat, in which process the chalk is decomposed, certain matters being expelled or driven off, and the lime left caustic. Chalk or limestone is a compound of carbonic acid gas and caustic lime, and is called a carbonate of lime (233).

119. The carbonic acid is combined with the lime by so weak an attraction, that heat alone is sufficient to expel the acid, together with a small quantity of water which the chalk always contains. If a piece of chalk is put into some vinegar, or indeed into any sour liquid, the chalk will be decomposed, and the carbonic acid will bubble through the fluid, until the vinegar or other acid is fully combined with lime, and its acid powers entirely neutralized. The same will happen with any other carbonate.

120. In consequence of this gas being considerably heavier than common air (85), it frequently collects in caverns, cellars, and other similar situations, and often occasions fatal accidents, suffocating those who
unfortunately enter the places thus filled with carbonic acid. Its presence in such places in the air, in any quantity, may always be easily ascertained, by letting down a lighted candle into the well or cellar: if the candle continues to burn, we know that there is enough oxygen present to support the life of an animal; whilst if the candle is extinguished, we are certain that the place is full of carbonic acid gas, and therefore that it must not be entered until the heavy gas has been dispersed by proper ventilation.

121. Carbonic acid is also evolved in large quantities during the process of fermentation, and occasions the pricking taste and effervescence of cider, bottled ale, and other liquors. Common ale is allowed to ferment in open vessels, so that nearly all the carbonic acid formed during that process (424) is dispersed; but bottled ale being confined in close vessels before its fermentation is completed, much of the carbonic acid evolved subsequently is pent up in the liquor, and escapes from it in innumerable small bubbles, when the cork of the bottle is removed.

122. Carbonic acid, then, is constantly being formed in several different ways; it is produced during the combustion of all substances containing carbon, during the respiration of animals, during the decay of almost all vegetable and animal substances, and during the process of fermentation. It is likewise, in many situations, naturally given out by the earth in large quantities.

123. To counterbalance all these sources of in-
crease, there is only the power of plants already mentioned of decomposing it, by abstracting its carbon and setting free the oxygen again (106, 698, 710, 745).

124. When two substances combine together and form a compound, they unite in definite and invariable proportions. A given weight of carbon, in burning, always combines with a uniform quantity of oxygen, to produce a certain weight of carbonic acid gas; and this rule holds good in all cases of chemical combination, for it is one of the distinctions between mixture and combination, that we are able to mix two substances together, in any relative proportion we like; but we are only able to make substances combine in certain fixed proportions.

125. Compounds do not always consist of equal parts of their elements, for they can consist of one part of one element, and one, two, three, or more parts of another element; and, indeed, there can frequently be formed several different compounds, by the union of two elements in various proportions. When, however, we mix together two substances which can unite together, they always combine in one of these fixed proportions; and if there is more of the one element than is requisite to form the compound, it is left unaltered.

126. Thus we know that every six grains of carbon, or pure charcoal, require sixteen grains of oxygen to burn them perfectly, and convert them into carbonic acid; and exactly the same quantity will be
required whether the carbon is burnt in a few seconds, or slowly combined with oxygen by the gradual process of decay. If we were to try to combine six grains of carbon with twenty grains of oxygen, we should find that only sixteen grains of the oxygen would combine with the charcoal, and the remaining four grains of oxygen would be left unchanged.

127. When we try to burn charcoal so that it shall get less oxygen than is requisite to convert it into carbonic acid, we find that it is possible to do so, but that the gas produced is not carbonic acid. Carbon and oxygen are able to combine together in more than one proportion; and consequently, though when carbon is burnt in the air, or where it can get abundance of oxygen, it always forms carbonic acid, yet when burnt so that it cannot combine with a sufficient quantity of oxygen to form that gas, it forms a different compound containing less oxygen, which is called carbonic oxide.

128. This compound is a transparent colorless gas, like carbonic acid, and resembles it also in being totally unfit to support animal life; indeed, it appears to be far more dangerous than carbonic acid when taken into the lungs, even though considerably diluted with common air. It extinguishes flame, as might be expected, but it is itself combustible, burning with a pale blue flame, and at the same time is converted into carbonic acid, in consequence of having acquired more oxygen from the air in which it has burnt.
129. We frequently see on the top of charcoal or coke fires a pale blue flame, quite different in appearance from the usual bright flame of wood or coal; this is occasioned by carbonic oxide, which is formed in the midst of the mass of burning fuel, where the carbon, being unable to get enough oxygen to form carbonic acid, produces this gas, which, when it reaches the top of the fuel, meets with fresh air, and combines with a further quantity of oxygen. When, however, a charcoal fire is burning slowly, a quantity of carbonic oxide escapes into the air without being burnt into carbonic acid; and its poisonous nature is often shown, when persons have foolishly placed a pan of burning charcoal in a close bedroom. The charcoal is not only abstracting the oxygen of the air, and converting it into carbonic acid, which cannot support life, but is also forming quantities of the highly poisonous gas, carbonic oxide, the presence of which in a room, in any considerable quantity, is sure to destroy life.

130. Besides these two compounds of carbon, there is yet a third, intermediate in composition between carbonic oxide and carbonic acid, though very different from either in its properties; this substance is oxalic acid, a strong, and very poisonous acid. Oxalic acid occurs in many plants, and may be easily formed artificially (503); it is a white solid substance, soluble in water, in which it forms a very sour solution; and has a strong affinity for bases. It has never been formed direct from carbon and oxygen.
131. Although when carbon burns in the air it only combines with oxygen, it can, under some circumstances, combine with nitrogen and also hydrogen. Thus when vegetable matters decay under water, we find that a gas is given off in bubbles which consist of hydrogen and carbon, and is therefore called carburetted hydrogen.

132. This gas is, as may be supposed, inflammable; burning in the air with a tolerably bright flame, and forming, by the combustion of its two elements, water and carbonic acid. This gas is found in very large quantity in coal mines, where it is called firedamp, and occasions violent explosions, when a light is incautiously brought into a mixture of it and common air.

133. In these cases the gas is mixed with a quantity of atmospheric air, but the affinity of the carbon and hydrogen, of which it consists, for the oxygen of the air, is not powerful enough to cause combination. When, however, a lighted candle or lamp is brought into the mixture, that part is immediately raised to the temperature at which combination can take place, the mixture takes fire, the flame spreads with very great rapidity, and in a few seconds the mixed gases are changed from air and carburetted hydrogen, into carbonic acid gas, steam, and nitrogen. At the moment of explosion the gases are very greatly expanded by the heat of the flame, and subsequently they are suddenly condensed, as the steam is cooled and converted into water. Carbonic acid and nitrogen alone
being left, many of the dreadful accidents which occur in coal mines are less caused by the violence of the explosion, than by the suffocating effects of the after-damp, as the foul air left in the mines after the explosion is termed.

134. There are also many other compounds of carbon and hydrogen, in which different proportions of the two elements give rise to a great variety of different substances: one of the most important of these is the common coal gas, obtained by distilling or roasting coals in close iron vessels, and which is used for lighting the streets; it differs from the fire-damp of mines, in containing rather more carbon. India-rubber, gutta-percha, coal-tar naphtha, oil of turpentine, &c., are also compounds of carbon and hydrogen.

135. In expressing the composition of any substance, chemists are in the habit of saying that it consists of such and such proportions of its elements; whatever quantity they may have taken for analysis, they generally calculate the proportion which a hundred or a thousand parts would consist of. Thus, for example, 550 grains of pure carbonate of lime contain 308 grains of lime, and 222 grains of carbonic acid; hence 1000 grains must contain 560 grains of lime, and 440 grains of carbonic acid; and 100 grains of carbonate of lime contain 56 grains of lime, and 44 grains of carbonic acid.

136. This is a very simple example, but it constantly happens that the composition of substances is not so easily expressed. 1000 grains of dry gypsum, or sulphate of lime, consist of 412 grains of lime, and
588 grains of sulphuric acid. The composition of 100 grains of such a substance is represented thus:—

<table>
<thead>
<tr>
<th>Lime</th>
<th>41.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulphuric Acid</td>
<td>58.8</td>
</tr>
</tbody>
</table>

This merely means that one hundred grains contain 41 and two-tenths of a grain of lime, and 58 and eight-tenths of a grain of sulphuric acid: hence there is no real difference whether we put the dot or not: if it is used, the figures behind are known to be fractions; if not they are whole grains.

137. Sulphate of lime may be expressed either as:

<table>
<thead>
<tr>
<th>Lime</th>
<th>0.412</th>
<th>4.12</th>
<th>41.2</th>
<th>412</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulphuric Acid</td>
<td>0.588</td>
<td>5.88</td>
<td>58.8</td>
<td>588</td>
</tr>
</tbody>
</table>

1 grain. 10 grains. 100 grains. 1000 grains.

In all the analyses given in the following pages, no fractions are used; the composition of everything is given as it would be obtained if 10,000 or 100,000 grains were analyzed.

138. Chemists are constantly in the habit of speaking of atoms, proportions, combining numbers, and similar terms; it will be well briefly to describe what is meant by these words. It has been already stated that the composition of all compound substances is definite (124); that a certain weight of carbon, for example, can only combine with a fixed quantity of oxygen, to form carbonic acid (126).

139. Let us observe what relation there exists between the quantity of different substances which can
combine together. It is found that one grain of hydrogen (for hydrogen, though a gas, can be readily weighed) will combine with eight grains of oxygen, to form nine grains of steam, and this relative proportion between the two elements of water is perfectly invariable. One grain of hydrogen can combine with exactly sixteen grains of sulphur, to form seventeen grains of sulphuretted hydrogen (182), a compound shortly to be described, or, with six grains of carbon, to form seven grains of carburetted hydrogen (131).

140. These numbers, then, express the quantity of each of these substances which can combine with one another, for, of course, it is perfectly the same whether we take a grain, an ounce, a pound, or any other weight. But this is not all; the number thus found for carbon, namely six, is not merely the quantity of that substance which could combine with one of hydrogen, but represents the quantity of carbon which can combine with eight parts of oxygen, to form carbonic oxide (127), or twice eight, 16, parts of oxygen to form carbonic acid; and, again, eight parts of oxygen is not merely the quantity which can combine with one part of hydrogen, or with six parts of carbon, but is exactly the quantity which is able to combine with a definite proportion or equivalent, of any other substance.

141. The numbers which are in this manner obtained, are called combining weights, proportionals, equivalents, &c.; they express the relative proportions in which substances combine together. Some com-
pounds consist of a single proportional of each of these elements, but more commonly they contain one of one element, and two, three, or more of the other; organic substances, for the most part, consist of nearly ten or a dozen proportionals of each of their elements (316, 369).

142. The following table shows the proportions or combining weights of the most important simple substances.

<table>
<thead>
<tr>
<th>Substance</th>
<th>Combining Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen</td>
<td>8</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>1</td>
</tr>
<tr>
<td>Carbon</td>
<td>6</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>14</td>
</tr>
<tr>
<td>Chlorine</td>
<td>35</td>
</tr>
<tr>
<td>Phosphorus</td>
<td>31</td>
</tr>
<tr>
<td>Sulphur</td>
<td>16</td>
</tr>
<tr>
<td>Iron</td>
<td>27</td>
</tr>
</tbody>
</table>

143. When a compound is formed by the union of two elements, the equivalent or combining proportion of the compound is exactly the sum of the equivalent of its elements; thus, for example, carbonic acid consists of one equivalent of carbon, the number of which is 6, and two equivalents of oxygen, weighing 16; the sum of 6 and 16, is 22; this, then, is the equivalent of carbonic acid, the quantity which will combine with an equivalent weight of any base, for example, with 28 parts of lime, 47 of potash, and so on.

144. In the following table the combining weight of some of the most important compounds is exhibited:
145. When plants or vegetable substances consisting of oxygen, hydrogen, and carbon, decay, their elements form new compounds, the principal of which are carbonic acid and water. We must now consider what are the products resulting from the decay of animal matters, and of those vegetable substances which, like them, consist of oxygen, hydrogen, carbon, and nitrogen; these are water, carbonic acid, and ammonia.

146. Nitrogen or azote differs from most other substances in appearing to be remarkably inert; it seems to have little or no affinity for any other substance. It is always mixed with oxygen in the air, but it appears to have no inclination to combine with it;
and when carbon, or substances containing it, are burnt, they combine only with the oxygen, and never with the nitrogen of the air: so that it would appear as if the chief use of nitrogen in the air was to dilute the oxygen, and prevent it from combining too rapidly with carbon, and other substances.

147. Under some circumstances, however, nitrogen does combine with other elements, and its compounds are amongst the most curious and important substances we know. When animal or vegetable matters containing nitrogen decay, we find that it, like the carbon, is not set free in its simple and uncombined form, but that during decay it combines with a portion of hydrogen.

148. Therefore, in addition to water and carbonic acid, the two principal substances arising from the decay of ordinary vegetable matters, we find a pungent, strong-smelling gas, composed of nitrogen and hydrogen, which is called Ammonia.

149. This substance, though a transparent invisible gas, is, like potash and soda, a base; like them it has a strong affinity for acids, and when combined with them neutralizes their powers; and, therefore, as it is evolved at the same time with carbonic acid, it combines with that acid and produces a solid salt, which is called a carbonate of ammonia, just as the compound of carbonic acid and lime is called a carbonate of lime.

150. Ammonia is always produced when animal or vegetable substances containing nitrogen are decom-
posed, whether by combustion or by decay; in both these cases carbonate of ammonia is formed, which is a very volatile salt, and consequently is carried away by the air, as fast as it is formed. Now water absorbs large quantities of ammonia, which, as well as all its compounds, is easily soluble in that fluid; and consequently the ammonia and salts of ammonia, formed during these processes of decomposition, are never retained in the air, but are dissolved and brought down to the surface of the earth, by rain.

151. Ammonia was formerly called the volatile alkali, because in many properties it closely resembles the alkalies potash and soda, which are distinguished from all other bases, by that name. Ammonia, however, differs from the two other alkalies in being a gas, whilst they are both solid, and almost all its compounds are more or less volatile, or capable of being converted into vapor by heat; whilst the compounds of potash and soda are all fixed, or quite incapable of being raised into vapor by any ordinary application of heat.

152. Ammonia has also a less powerful affinity for acids than the two other alkalies have, and consequently it is easily expelled from its compounds, by either potash or soda, as well as by lime and several other bases, which set free the ammonia in the gaseous form, and unite themselves with the acid previously in combination with the ammonia.

153. Ammonia, in its pure state, is a colorless, invisible, and very pungent-smelling gas, readily soluble
in water; like potash and soda, it has a very strong affinity for acids, and is, therefore, hardly ever met with in its free state, but always in combination with carbonic, or some other acid. It is far less abundant or common than the other alkalies, potash and soda; for its principal source is the decay of animal and vegetable substances, which contain nitrogen.

154. The most important of the salts of ammonia are the carbonate, the muriate, the sulphate, and the phosphate: the carbonate, which is the salt produced during the decay of organic matters, was formerly called salt of hartshorn, because it was mostly procured by burning horn; it is now, however, obtained from the tar-water and refuse ammoniacal liquors of the gas-works. A considerable quantity of ammonia is formed during the distillation of coal, which always contains a little nitrogen, and, therefore, gives off a small quantity of ammonia; this combines with a portion of the carbonic acid formed by the burning of the carbon of the coal, and collects in the hydraulic main, or first condenser, of the gas-works; hence, also, carbonate of ammonia is always an ingredient of soot.

155. Carbonate of ammonia is not obtained direct from the gas liquor, as it would then be impure from the tar, and other substances which exist in the gas-water. The ammoniacal water is generally first converted into a sulphate or muriate, and this is decomposed by heating it with chalk; carbonate of ammonia, which is volatile, rises in vapor, whilst sulphate or
The muriate of lime is left. Carbonate of ammonia possesses the peculiar pungent smell of pure ammonia, though far less powerfully; because, being but partially neutralized by carbonic acid, and united to it by but a feeble affinity, it is constantly gradually escaping from its compound, and evaporating into the air. Like the carbonates of potash and soda it possesses the property of reddening many vegetable yellows, though less powerfully than the pure alkali.

156. The muriate of ammonia, commonly called sal-ammoniac, which used also formerly to be obtained by burning animal substances, is now procured from the refuse of the gas-works, which contains carbonate and other salts of ammonia. They are first of all converted into a sulphate, by the action of oil of vitriol, which expels the carbonic and other weaker acids; and the sulphate thus obtained is decomposed by mixing it with common salt, and heating the mixture.

157. The nature of this operation is very simple. Sulphate of ammonia, water, and common salt, or chloride of sodium, are heated together, and are decomposed, because sulphuric acid has a stronger affinity for soda than it has for ammonia; a portion of water is decomposed at the same time with the salt, supplying oxygen to the sodium to form soda, and hydrogen to the chlorine to form muriatic acid; the former unites with the sulphuric acid, whilst the latter combines with the ammonia.

158. Muriate of ammonia, like many other of the salts of ammonia, is volatile at a certain heat; that
MURIATE OF AMMONIA.

is to say, it becomes a vapor, and resumes the solid form when cooled again: in the process for obtaining the muriate of ammonia, therefore, it is converted into vapor by the heat applied to the mixture, and is condensed in the solid form again in the upper part of the vessel in which it is heated, which is kept cool on purpose. The uses of sal-ammoniac in the arts are numerous; its principal consumers are the dyers and workers in metals.

159. Ammonia has a strong affinity for sulphuric acid; the sulphate may either be formed directly by mixing together sulphuric acid and caustic ammonia, or its carbonate, as in the process just described for preparing the muriate from gas liquor; or it may be obtained by substituting for the sulphuric acid, added to the gas liquor, a sulphate in which the acid is held by a less powerful affinity than that which it has for ammonia; when this is done, the ammonia seizes the sulphuric acid and causes it to relinquish the weaker base with which it was previously combined, and which thus unites with the carbonic acid, of the carbonate of ammonia.

160. When, for example, a solution of carbonate of ammonia is mixed with a quantity of sulphate of lime, they are both decomposed; the ammonia combines with the sulphuric acid, and the lime takes the carbonic acid. Sulphate of ammonia has a strong saline taste, but no smell: it is a perfectly neutral salt, and may be kept any time without undergoing change. It is readily soluble in water.
161. Phosphate of ammonia may be formed by adding a solution of ammonia, or its carbonate, to phosphoric acid (194); it is very soluble in water, and decomposes when heated, the ammonia passing off in the gaseous state, and leaving the phosphoric acid pure. A rough salt may be made by mixing the impure phosphoric acid obtained by acting on bone, earth, by oil of vitriol, with gas liquor, and evaporating the solution; it is then, however, apt to contain lime, tarry matter, and other impurities. It is sometimes employed as a manure (793).

162. Ammonia often seems to combine with substances for which it really has no affinity: this is merely a mechanical effect. Such substances have the power of absorbing a large quantity of the gas, which is condensed or packed up in the pores of the substance, but not combined with it, and is ready to be given off again on the application of heat. Thus, freshly burnt charcoal, and, indeed, most porous substances, absorb or condense a large volume of this gas. The value of charcoal as a disinfecting agent, and purifier generally, chiefly depends upon this power of absorbing gaseous matter. But it also possesses bleaching powers.

163. Ammonia is not the only compound containing nitrogen, which is formed during the decomposition of animal substances; for although nitrogen and oxygen appear to have no affinity for each other, as they exist together in a state of mixture in the air, and do not enter into combination, yet when a com-
pound already containing nitrogen is decomposed, we find that, besides the compound of nitrogen and hydrogen just described, a compound of oxygen and nitrogen is also in some cases formed, which is possessed of very extraordinary powers, being a strong acid, and commonly called aqua-fortis, or nitric acid.

164. Nitrogen is able to combine with oxygen in five different proportions, giving rise to as many distinct substances; four of these are of comparatively little importance, and may therefore here be passed over; the fifth, or that containing the greatest quantity of oxygen, is the only one at present worthy of consideration.

165. NITRIC ACID is never found in its free or uncombined state, but always in combination with some base; being a very powerful acid, it is easy to understand that, as soon as formed, it seizes upon some base to combine with, and forms a neutral salt. The commonest salts containing nitric acid, or nitrates, as they are called, are the nitrates of potash, soda, and lime, which are found native abundantly in different places.

166. The nitrate of lime is very frequently to be met with on old walls, near which organic substances containing nitrogen have decayed and formed nitric acid, which has combined with the lime of the mortar; and in the same way nitrate of lime may be formed artificially, by mixing lime with decaying substances which can yield the acid, and, after having allowed
them to remain some time together, by pouring water over the mixture: this dissolves the nitrate of lime, and forms a solution, which leaves the salt in question, when the water is evaporated by heat.

167. Although nitric acid is a very strong acid, and has a very powerful affinity for bases, yet there are even more powerful acids, which are strong enough to expel it from its combinations, and we are thus able to procure the pure acid: this is usually done by what is called oil of vitriol, or sulphuric acid (179), a substance consisting of sulphur, or brimstone, combined with oxygen. When this acid is poured on a nitrate, it expels the nitric acid in the form of a very acrid or sour and biting vapor; this, condensed in cold water, constitutes the intensely sour liquid called nitric acid, or aqua-fortis, and was before combined with the base of the nitrate.

168. Nitric acid acts very strongly on all organic bodies, decomposing them sometimes with great violence, and causing their elements to enter into new combinations: thus when strong nitric acid is mixed with oil of turpentine, it sets it on fire. This is caused by the large quantity of oxygen which nitric acid contains, which enables it to burn substances, or at least makes them combine rapidly with oxygen, which we know is the cause of ordinary combustion.

169. When we mix nitric acid with substances which, having a strong affinity for oxygen, are able to decompose the acid, we obtain some of the other compounds of nitrogen and oxygen before alluded to,
and by various processes we are able to obtain compounds of nitrogen, with most other elements, setting out with this acid as a source of nitrogen; for it seems that nitrogen hardly ever will combine with an element, except when in the nascent state; that is, when being evolved or set at liberty by the decomposition of a substance, in which it was already combined with some other element (769).

170. There are three other elements deserving notice, which, like nitrogen, are of little interest in themselves, but which, by combining with other substances, form important compounds; these are sulphur, chlorine, and phosphorus.

171. The properties of sulphur, or brimstone, are familiar to every one; it occurs native, both pure and also in combination with various metallic substances; entering into the composition of many ores, such as those of copper, lead, zinc, mercury, &c. Sulphur is a pale-yellow, solid substance, insoluble in water, having hardly any taste, but a faint and peculiar smell, melting easily when warmed, and catching fire very readily, burning with a pale-blue flame, and giving out at the same time a peculiar and very disagreeable smell. The compounds which sulphur forms with oxygen, and also with hydrogen, are the principal which demand attention.

172. Sulphur, like nitrogen, is able to combine with oxygen in several proportions, and hence it forms several distinct compounds. It is only necessary to describe two of them—sulphurous acid, and sulphuric acid or oil of vitriol.
173. When sulphur is burnt in the air, it forms sulphurous acid, a transparent, invisible gas, possessing a very strong and suffocating smell; it is easily dissolved by water, and the solution then obtained has a strongly sour taste.

174. A portion of sulphurous acid is formed whenever we light a common brimstone match; and the peculiar suffocating smell then perceived is occasioned by this acid, and not by the sulphur alone, which has no smell, but which is used to tip the matches because it has a strong affinity for oxygen, and requires less heat to enable it to commence combination than the wood itself; but when it has once fairly got alight, it then produces heat enough to fire the wood.

175. Sulphurous acid is a transparent gas, rather heavier than common air, readily soluble in water, very poisonous when breathed, and extinguishing the flame of combustible matters; it does not form any important compounds when combined with bases, and its principal interest consists in its bleaching powers. It has long been used to destroy color in things desired to be bleached, which are merely hung up over some burning sulphur, and exposed to the fumes of sulphurous acid thus formed. Articles thus bleached, however, frequently retain an unpleasant smell of sulphurous acid.

176. It is likewise used in fumigation, being considered destructive of contagion; and, therefore, sulphur is burnt in houses where persons have died of an infectious disorder, and in other circumstances
where evil effects are feared from the communication of disease. In all these cases sulphurous acid has really a useful influence, as it seems able to destroy a minute quantity of poisonous matter diffused throughout the air; but it must be remembered, that it can have no power of improving air which is unwholesome, in consequence of the presence of a large quantity of carbonic oxide or carbonic acid, for in these circumstances, burning sulphur would only tend to make the air worse, both by removing oxygen, and also by adding sulphurous acid.

177. The bleaching power of this acid may be very well shown, by holding a lighted brimstone match near a dark purple or blue flower, the color of which will be immediately more or less destroyed, and we may easily thus, either in part or wholly, render the flower white.

178. Sulphurous acid contains less oxygen than the sulphuric acid; in fact, it stands in the same relation to that acid, which carbonic oxide does to carbonic acid. Sulphurous acid is formed whenever sulphur is burnt in the air, because, under those circumstances, it is not able to combine with enough oxygen to form the more powerful acid.

179. SULPHURIC ACID is made by burning sulphur mixed with nitrate of potash, which supplies it with more oxygen than it could get by merely burning in the air; because the nitric acid in the nitre, when decomposed by the burning sulphur, gives rise to the formation of another compound of nitrogen and oxygen con-
SULPHURIC ACID.

180. It is a heavy, very corrosive, poisonous fluid, although its elements are only oxygen and sulphur, the one being that part of the air necessary to support life, and the other a tasteless, yellow, solid substance. Sulphuric acid in its free state is chiefly remarkable as being one of the strongest acids we know, destroying vegetable and animal substances: that is to say, abstracting and combining with some of their elements, and causing the others to enter into new combinations; and having a strong affinity for bases, with which it forms a class of compounds called sulphates: some of these are of very great importance, and must be considered hereafter (159, 213, 224, 237.)

181. Sulphuric acid is a most valuable substance to the chemist; being one of the strongest known acids, it enables him to expel or drive out most other acids from their combinations, and thus through its agency many other acids are obtained pure and separate, from their compounds with bases (167, 188, 195).

182. Under certain conditions sulphur combines with hydrogen to form sulphuretted hydrogen—a transparent colorless gas, remarkable for the exceed-
ingly disagreeable smell which it possesses. It is to the presence of this gas the nauseous, disgusting smell of putrefying animal matter is due. Sulphuretted hydrogen is very poisonous, and it would probably cause many fatal accidents, were it not that its offensive odor warns us of its presence, and induces us to take measures to disperse it by ventilation and other means.

183. Sulphuretted hydrogen is soluble in water to a considerable extent, and the solution possesses the strong and nauseous smell of the gas itself. The gas is inflammable, and in burning produces water, and sulphurous acid gas. Sulphuretted hydrogen has sometimes been called an acid, for it possesses feeble acid powers, combining with some bases and to a certain extent neutralizing them.

184. Sulphur has a strong affinity for most metallic substances forming a numerous series of compounds, which are called sulphurets: many metallic ores are sulphurets, and in the process by which the metals are obtained, the first operation consists in roasting the ores, or exposing them to a dull red heat, when the sulphur, being a volatile substance, is gradually driven off.

185. Chlorine is an element which is never found free, but only in combination with some other element; it is a very poisonous gas, causing very great irritation when breathed, even though it is largely mixed with air; it is transparent, has a bright-yellow color, and is remarkable for possessing in a very great degree
the power which sulphurous acid has, of bleaching or destroying colors, and also contagious matters. Chlorine is, therefore, now largely used in all cases where it is wished to destroy or remove color: it is however, a very powerful agent, and therefore must be used with great care, as otherwise if linen, for example, be bleached carelessly by means of it, the texture is destroyed, and the linen becomes rotten and useless. Common writing-paper is often found to be quite decayed and useless, the rags from which it was made having been too strongly bleached with chlorine.

186. Small quantities of this gas are used with very beneficial effects in removing foul smells, and otherwise rendering sweet and wholesome the air of large buildings, such as hospitals. Chlorine readily combines with the metals, and most other of the elements, to form a series of compounds called chlorides; thus corrosive sublimate is a chloride of mercury, &c. It is unnecessary for us further to occupy ourselves with this element than to know that when combined with hydrogen it loses all these peculiar powers, and forms a strong acid, the muriatic, which, by combining with bases, forms a series of salts called muriates.

187. When chlorine and hydrogen are mixed together in equal volumes, an explosive mixture is formed, which, on the application of a flame, detonates loudly, and is converted into muriatic acid gas. An explosion is also caused if the mixture is exposed in a glass vessel to sunshine, but if kept in the dark, no combination takes place; if merely exposed to day-
light, but not to the direct rays of the sun, the gases
do not explode, but combine slowly and quietly.

188. The process by which muriatic acid is procured
is very simple: a quantity of oil of vitriol or sulphuric
acid is poured on some common salt, acid fumes arise,
they are made to pass through cold water, which ab-
sorbs them, and in time a very sour corrosive liquid
is obtained, which is the muriatic acid, or the spirit
of salt of the shops. In order to explain the action
which takes place in this process, it is necessary to
say a few words respecting the nature of common
salt.

189. Common salt is a compound substance, con-
sisting of two elements; the one is chlorine, the cor-
rosive yellow-colored gas just described, and the other
is a white silvery-looking metal called sodium, a sub-
stance having a very strong affinity for oxygen, eager-
ly combining with it and forming the alkali soda
(215).

190. The substances, then, concerned in the process
for making muriatic acid, are common salt, or, to
speak scientifically, chloride of sodium, sulphuric acid,
and of course a portion of water, as oil of vitriol is a
solution of dry sulphuric acid in water. Both the
chloride of sodium and the water are decomposed, the
oxygen of the water combines with the sodium of the
salt, whilst the hydrogen of the water combines with
the chlorine of the salt, and thus soda and muriatic
acid are formed; the latter is given off or expelled in
the form of gas or vapor, whilst the soda combines
with the sulphuric acid to form a neutral salt—the sulphate of soda. This is a very good example of combination and decomposition. In consequence of the powerful affinity which sulphuric acid has for soda, both the salt and water are decomposed; and the chlorine and hydrogen being free from their combinations, unite together to form muriatic acid (216).

191. Chlorine is procured from muriatic acid by the action of some substance capable of abstracting its hydrogen; the most convenient for this purpose are those which contain much oxygen. The usual process is to heat together a mixture of muriatic acid and oxide of manganese, a substance which is rich in oxygen; the oxygen which is thus given off, takes the hydrogen of the muriatic acid, with which it forms water, and sets free the chlorine.

192. Another and yet cheaper process is to heat a mixture of common salt, oxide of manganese, and sulphuric acid; in this case the sulphuric acid decomposes the salt, and the muriatic acid formed, is decomposed as fast as it is set free, by the oxygen of the oxide of manganese.

193. IODINE, bromine, and fluorine, are substances hitherto undecomposed, and therefore called elements, which in many respects resemble chlorine; like it, they form acids by combining with hydrogen, and unite to the metals to form a numerous series of compounds resembling chlorides, which are called iodides, bromides, and fluorides. None of these substances are of much importance; those most deserving of notice are the iodide of sodium, which exists with the chlo-
ride in sea-water, and the fluoride of calcium, which exists in small quantity in the bones of animals.

194. Phosphorus in several respects resembles sulphur; like it, it is a readily fusible and very combustible solid. It is of a white color, about as hard as wax, and has so strong an affinity for oxygen that it takes fire in the air with the greatest facility as soon as it is a little warmed. Like the preceding element, chlorine, it is never met with in a separate state, but always in combination. Its most important compound is phosphoric acid, the substance which is formed when phosphorus burns in the air or in oxygen; it is a white solid substance, very soluble in water, and eagerly combining with bases to form a class of salts called phosphates (161, 242).

195. Phosphoric acid is generally obtained by decomposing one of its compounds by sulphuric acid. The most abundant compound of phosphoric acid is that in which it is united to lime, called the phosphate of lime (242): this substance is an ingredient of the bones of animals, and of most organic substances. When sulphuric acid is mixed with phosphate of lime, the latter is decomposed, and its phosphoric acid is set at liberty. The acid may be artificially made by burning phosphorus in the air, from which it abstracts the oxygen, and leaves the nitrogen unaltered; just in the same way that carbon when burnt in air, forms carbonic acid: only that in this case the product of combustion is a gas, like the air; whilst the result of the combustion of phosphorus, is a white solid acid.
CHAPTER III.

METALLIC ELEMENTS—THE ALKALIES—THE ALKALINE EARTHS, AND THEIR SALTS.

196. In classifying the elementary substances, the first and most obvious arrangement seems to be that of dividing them into non-metallic and metallic; the former including oxygen, chlorine, &c., or the supporters of combustion as they are sometimes called, together with the non-metallic simple combustibles, such as carbon and sulphur, &c.; whilst the latter includes iron, gold, lead, tin, and all the ordinary metals, as well as a number of very scarce metals; some of which, however, are very abundant in a state of combination, though they are hardly ever met with in a pure and separate form.

197. In the pure state, the metals are chiefly important for their strength, hardness, malleability, and other mechanical properties; their interest to the chemist is for the most part confined to their combinations, and the different compounds which they form by combining in various ways, with the non-metallic elements. The oxides, or the substances which the metals form by uniting with oxygen, are
perhaps the most important of all their compounds, a very great number of them are bases (109).

198. The class of substances called bases, or those with which acids can combine, is very numerous. The commonest, and therefore the most important, are the three alkalies—potash, soda, and ammonia, together with a few substances called earths, because they are the principal ingredients which enter into the composition of the earth or soil. The nature and properties of ammonia have already been considered (148).

199. Potash never occurs pure in a free state; its most abundant source is the vegetable kingdom, and the simplest method of procuring it is by burning wood. When this is done, after all the combustible matter is burned, there remains a quantity of a white or gray substance called ashes, which when put into water forms a caustic solution. This is caused by the presence of potash, which, not being altered by the heat occasioned by the burning of the wood, is left in the ashes.

200. In this way, however, we do not obtain the potash pure, because, being a powerful base, and having a strong affinity for acids, it combines with some of the carbonic acid formed by the burning of the carbon contained in the wood, and therefore the caustic substance found in the ashes of the wood is an impure carbonate of potash. It is obtained tolerably pure by pouring water on wood ashes, straining off the clear liquor, and evaporating it; a white salt
is left, which is the carbonate of potash separated from most of the other matters which were contained in the ashes, and when thus purified it is called *pearl-ash*.

201. Pure potash is readily obtained from its carbonate, by means of quicklime; that substance has a stronger affinity for carbonic acid than potash has, and accordingly it decomposes the alkaline carbonate and leaves the alkali in its pure state: when free from any acid, it is found to be a very caustic solid substance, which has a strong affinity for acids, and is difficult to keep perfectly pure, as when exposed to the air it rapidly acquires carbonic acid from it, and becomes converted into the carbonate.

202. Potash is not a simple substance; like soda, it is a compound of a metal and oxygen (190). By the action of charcoal at a very high temperature it may be decomposed, for at a white heat the affinity of charcoal for oxygen is so great that it is even able to take it from potassium, as the metal of potash is termed. Potassium is of course never found in a native state, for its affinity for oxygen is so great that when exposed to the air it instantly absorbs a quantity of that gas, and is soon converted into potash. It may readily be proved that potash, soda, and the earths are all compound bodies, and that they consist of peculiar metals combined with oxygen; but as the affinity which they have for that element is very great, and as they are never decomposed under ordinary circumstances, it is quite unnecessary to
study the nature of potassium, sodium, &c., further than to know that they are white silvery-looking metals, which by combining with oxygen form potash and soda. For all practical purposes we may think of potash, soda, and the earths as elementary substances.

203. Potash has a strong attraction for water; so that, when dry, pure potash is exposed to the air, it very soon becomes moist, and in a short time has attracted so much water from the air as to appear changed into a liquid. This power of attracting water from the air is common to most of the salts of potash, as well as to many other saline compounds; it is termed deliquescence, and such salts are said to be deliquescent. Pearlash, if left exposed to the air, rapidly becomes moist, and finally runs into a liquid, which is a very strong solution of that salt in water; pearlash, therefore, must always be kept in a dry place, and in well-closed vessels.

204. In the same way, we often find that common salt, when kept in a damp place, becomes very moist: this is not occasioned by any attraction which the salt itself has for the moisture in the air, but by certain saline impurities usually contained in the salt, which have this power in a high degree (252).

205. Chloride of potassium may be formed by heating the metal potassium in chlorine, or by acting upon potash by muriatic acid (190); it is not a salt of much interest; it has been occasionally recommended as a manure (891).
206. One of the most important of the salts of potash, is the nitrate, or, as it is commonly called, saltpetre, or petre, which is found native in many places as a white powdery matter on the surface of the ground, and is procured pure by washing the earth, and evaporating the solution of nitre thus obtained.

207. Nitre is a perfectly neutral salt, and is in that respect very different from the carbonate of potash, which, although far less caustic than pure potash, has still very considerable caustic powers, in consequence of which it is so commonly used in washing, and for other household purposes. Carbonic acid, in combining with the alkalies, does not seem able to neutralize them, or destroy their alkaline powers so completely as most other acids can; and therefore it must be remembered that the carbonates of the alkalies still possess some of the powers of the alkalies themselves. The other compounds formed by the alkalies with the more powerful acids are quite neutral, and in them no traces of the alkaline nature of the bases can be found.

208. One of the principal uses of nitre is in the manufacture of gunpowder, which consists of nitre, charcoal, and sulphur, finely powdered, and very carefully mixed together. The chemical action which takes place during the burning or explosion of gunpowder is very simple. The nitre consists of potash and nitric acid, the latter substance contains a large quantity of oxygen, combined by a comparatively...
weak attraction with a portion of nitrogen. The finely-powdered charcoal has a very strong attraction for oxygen, and when sufficiently heated is able to decompose the nitre, seizing upon the oxygen of the nitric acid, and forming carbonic acid, a small portion of which combines with the potash of the nitrate, and forms carbonate of potash.

209. The use of the sulphur in gunpowder is similar to its use in the lighting of a common sulphur match; it very easily catches fire, and the flame thus produced instantly heats the charcoal so much that it is able to decompose the nitre.

210. The mechanical force of gunpowder when fired, is occasioned by the instantaneous production of a large quantity of gas from a small bulk of solid matter: as the whole of the nitrogen, and the greater part of the carbonic and sulphurous acids formed, are given off in the gaseous state, and comparatively little is absorbed by the potash.

211. During the burning of gunpowder, a small portion of sulphuretted hydrogen is commonly formed, which occasions the bad smell (182) commonly perceived: the hydrogen necessary for the formation of this gas is derived from a little moisture which gunpowder generally contains; pure and perfectly dry gunpowder contains no hydrogen, and hence, in its firing, no sulphuretted hydrogen can be found.

212. Nitre is one of the few compounds of nitric acid which are found native, and, indeed, it was till lately almost the only source of that acid. It is also
used as a manure, and will therefore come before us again on a future occasion (898).

213. Sulphate of potash is a soluble salt, readily formed by acting on any of the ordinary salts of the alkali, such as the carbonate or nitrate, by sulphuric acid; it is chiefly used by the manufacturers of alum (257).

214. Salts of potash are met with in a great many plants in small quantities; they are likewise almost always to be found in the soil, and potash is by no means uncommon in stones; indeed, it is evident that the salts of potash found in the soil must have been derived from the gradual breaking down and decomposition of stones and rocks containing compounds of potash.

215. The second alkali, soda, is in its general characters similar to potash; it is a very caustic, solid, white substance, has a powerful attraction for acids, and is consequently never found in a pure and separate condition, but always in combination with an acid, or some other substance. The most abundant source of soda is common salt, which exists in very large quantities in sea-water, and is likewise found native in the earth, when it is called rock-salt, to distinguish it from the salt obtained by the evaporation of sea-water, and called sea or bay salt.

216. Common salt has already been stated to be a chloride of sodium (189), a compound of chlorine with a white brilliant metal called sodium. It used formerly to be called a muriate of soda, because it was
known that sulphuric acid poured on salt expels muriatic acid, whilst, on the other hand, there are substances which can take away the acid and leave caustic soda. It is now known that dry salt contains nothing but chlorine and sodium; and when it is decomposed by sulphuric acid, or in any other way, the muriatic acid and soda evolved were not contained in the salt, but are formed by the decomposition of a portion of water, the oxygen of which unites with the sodium to form soda, whilst the hydrogen combines with the chlorine to form muriatic acid. Hence, though strictly speaking common salt is a chloride of sodium, yet for practical purposes it may be considered as a muriate of soda; because whatever change it undergoes, or in whatever manner it is decomposed, soda and muriatic acid are given out just as if they were really the components of salt. Soda is an oxide of sodium.

217. The only other compound of soda, which occurs native in any quantity, is the nitrate or cubic nitre, as it is commonly called. It is found native in large quantities in certain parts of South America, and is used to a considerable extent as a manure. When it was first brought over to this country, some few years ago, it was proposed to substitute it for the nitrate of potash in the manufacture of gunpowder, and in the preparation of nitric acid. It was, however, found that it possessed the property of attracting moisture from the air; and hence, being always more or less damp, it could not be used by the gun-
powder-makers. When first imported its price was low, and it was accordingly advantageously substituted for the more expensive nitrate of potash in the manufacture of nitric acid; but as the price of nitrate of soda soon rose, it was no longer found profitable to use it for this manufacture, and now almost the only purpose for which it is employed is as a manure. It appears to resemble the nitrate of potash in its effects on growing plants, and like it to exert considerable influence on the fertility of certain kinds of land (901).

218. The "soda" of commerce is, as will be readily supposed, not the pure alkali soda, but is a carbonate, and, like the carbonate of potash, possessed of considerable caustic properties. It was formerly for the most part procured by burning seaweed, the ashes of which contain a large quantity of carbonate of soda, and immense quantities of seaweed were annually collected and burnt, for the purpose of obtaining weed-ash, kelp, or barilla, as the crude salt was called.

219. Of late years, however, means have been discovered of obtaining the carbonate of soda by decomposing common salt; from which it is now manufactured so cheaply that it has not only superseded the use of kelp or barilla for all purposes where carbonate of soda is required, but has also from its low cost been substituted to a great extent for potash and pearlash; and hence the carbonate of soda, or "soda," as it is commonly called, is very largely used
in the arts of glass and soap-making, and other processes where an alkali is employed.

220. In order to obtain carbonate of soda by the decomposition of common salt, it is first of all necessary to convert the salt into a sulphate; this is done by mixing it in certain proportions with oil of vitriol or sulphuric acid, and heating the mixture in a furnace. Muriatic acid is expelled, and the sulphuric acid, which has a powerful attraction for the soda, combines with it and forms sulphate of soda (190).

221. The second process is to mix the sulphate of soda thus formed with chalk and pounded coal, and to strongly heat the mixture in a furnace; the sulphate is decomposed at a high heat by the charcoal or carbon of the coal, which takes oxygen both from the sulphuric acid and from the soda, forming a sulphuret of sodium. This in turn is decomposed by the chalk, and a carbonate of soda is produced, which is of course impure, containing charcoal, and an insoluble compound of lime; so that, by pouring water on the crude salt after it is taken out of the furnace, a clear colorless solution is procured, yielding on evaporation a clean and nearly pure carbonate of soda.

222. This "soda" contains a large quantity of water, although apparently dry; and when left exposed to the air, it does not, like the carbonate of potash, attract water from the air, but, on the contrary, parts with it and becomes drier, at the same
time crumbling down to a fine powder, which is found to be less caustic than it was before it underwent this change. The reason of this is, that when the carbonate of soda, by exposure to the air, parts with a quantity of water, and consequently becomes much lighter, it at the same time gradually combines with a second dose of carbonic acid, which it obtains from the air, and its caustic qualities become more fully neutralized than they are in the common carbonate.

223. The salts of soda are for the most part white, and easily soluble in water; and, like the compounds of potash, are widely diffused throughout nature. They are found, besides the sources already mentioned, in many rocks and soils: a great many plants, more especially those which grow near the sea or in salt marshes, contain large quantities of salts of soda; and a good many of the compounds which this alkali forms with different acids are found native, though in far smaller quantities than the nitrate.

224. The sulphate of soda is also found native in Spain and other countries, and is frequently met with in mineral waters; it is also called Glauber's salts, and Cheltenham salts, as it is the principal ingredient in the mineral springs of Cheltenham. Sulphate of soda is left after the process for obtaining muriatic acid (190): it is occasionally employed as a manure.

225. The earth, or soil, consists of a mixture of various ingredients; but the greater part of it is composed of certain substances which are called earths. Some of these are bases, and resemble the alkalies;
whilst the remainder are not bases: that is to say, they seem to have no affinity for acids, and cannot combine with them to form salts. The most important of the earths which are bases, are called lime, magnesia, and alumine; the two former are called alkaline earths, to distinguish them from the last, as they possess the property of reddening vegetable colors, though in a far less powerful manner than the alkalies, whilst alumine does not. Silica is not a base.

226. **Lime,** in its pure condition, is a highly caustic whitish-gray solid substance, slightly soluble in water, but far less so than the alkalies are. It has a very strong attraction for acids, being even able to decompose the salts of the alkalies, in consequence of the superior attraction which it has for the acids contained in them; and it is for this reason that it is customary to mix quicklime with wood-ash, which contains comparatively little free potash, as the greater part is combined with carbonic acid (200), when it is desired to make a very strong caustic ley: the quicklime decomposes the carbonate of potash, combines with the carbonic acid, and becomes converted into carbonate of lime; whilst the potash, being no longer combined with an acid, is able to exert its caustic powers. Lime is the oxide of a metal called Calcium (202).

227. In consequence of its strong affinity for acids, lime never occurs native in a pure and simple state, but always in combination with some acid; the most common compounds of lime are the carbonate, and
CARBONATE OF LIME.

the sulphate; the former of these, especially, is a very abundant substance, existing naturally in immense quantities in the different forms of chalk and limestone; it is also very commonly an ingredient of the soil, and constitutes a considerable part of the bones of animals, and shells.

228. As has already been stated (119), lime has a less powerful attraction for carbonic acid, than it has for any other acid; and therefore when we pour a quantity of any acid over carbonate of lime, great bubbling, or frothing is occasioned by the escape of the carbonic acid, which is set at liberty, when the lime combines with the new acid (117).

229. This bubbling, or effervescence, as it is called, enables us very easily to ascertain whether a stone be limestone or not, and likewise whether any particular soil contains carbonate of lime; for if any acid, such as vinegar, does not occasion any effervescence, it is certain that no carbonate of lime is present; and if, on the other hand, it does expel bubbles of gas, we may be quite sure that a portion of some carbonate is present, and by far the most common of these is the carbonate of lime.

230. All the forms of this substance are tasteless, and quite insoluble in pure water; but rain-water, which commonly contains dissolved in it a very small quantity of carbonic acid, has the power of dissolving carbonate of lime: hence we commonly see the surface of chalk or limestone appearing to be gradually corroded, or worn away by the rain; and for the same
reason, buildings of freestone or limestone are much less durable than those which are built of granite, a stone which does not contain carbonate of lime, and is, therefore, not thus acted on by rain-water.

231. In this kind of action, no chemical change is effected: the carbonate of lime is merely dissolved by water holding in solution carbonic acid: hence when, either from heat or any other cause, the carbonic acid is expelled from the water, it is no longer able to hold the carbonate of lime in solution, which is therefore deposited again unaltered.

232. Spring-water, for example, which often holds carbonate of lime thus dissolved, becomes milky or turbid when boiled, and not unfrequently deposits fur in the inside of boilers and kettles; such water is of course not fit for domestic uses, cooking, &c.; but by simple boiling it is at once rendered far better, the chalk being then separated.

233. The uses of carbonate of lime are very numerous: one of the most important is for burning into lime. This is a very curious process, for heat generally increases the attraction which two different substances have for each other; yet in this and some other cases it rather diminishes it, and the strongest heat is unable to drive off carbonic acid from its combination with potash or soda, though lime, which parts with it so readily when heated, is able when cold to take it from either of the alkalies.

234. When quicklime is mixed with vegetable or animal substances, it renders the process of putrefac-
tion far less noisome than it is under ordinary circumstances; because the lime absorbs sulphuretted hydrogen, carbonic acid, and many of the products of decay.

235. When quicklime is slaked with water, it crumbles down to a powder, and is found to have combined with a quantity of water; although it appears quite dry, for the water is chemically combined with the lime: if this lime thus slaked is left exposed to the air, it combines with carbonic acid, and becomes changed into carbonate of lime, and at the same time parts with the water with which it had previously combined. Slaked lime is less strongly caustic than quicklime, because part of the action of lime on organic matter is caused by its strong affinity for water; nevertheless, even slaked lime possesses strong caustic properties.

236. Common mortar, which consists of slaked lime mixed with sand, chopped hair, &c., is a hydrate of lime; its setting depends chiefly on the absorption of carbonic acid from the air, which forms again with it a kind of chalk or limestone. Hence mortar gradually improves, becoming harder after a time, when it has absorbed a sufficient quantity of carbonic acid from the air to be wholly converted into carbonate.

237. Sulphate of lime is less common and abundant than the carbonate; it is, however, a frequent ingredient in the soil, in spring and mineral waters, and is found sometimes in large beds, constituting what is called plaster-stone, gypsum, and alabaster; these
are all compounds of lime and sulphuric acid, and precisely similar in composition. Common plaster of Paris is dry sulphate of lime, or gypsum, deprived by heat of the water which it naturally always contains, and, when mixed with a small quantity of water, it recombines with it, and becomes a hard, dry, solid substance.

238. The burning of gypsum to make plaster of Paris, is quite different from the burning of chalk to make quicklime: in the former case the native sulphate of lime, which contains water, is heated and the water thus expelled; whilst in the burning of lime not only is the water expelled which the chalk usually contains, but also the acid itself, with which the lime was combined.

239. Sulphate of lime is dissolved in small quantity by water, and accordingly we find it almost always present in spring-water, which derives it from the soil through which it rises; it is the presence of this substance in spring-water which gives it that hardness which distinguishes it from common river-water, and renders it less pleasant to use than soft water; in fact, one of the great distinctions found between hard and soft water is the presence or absence of sulphate, or carbonate of lime, which, though contained in but small quantity, greatly influences its usefulness for domestic purposes (232).

240. Nitrate of lime is a white salt, easily soluble in water, and readily formed by acting on lime or its carbonate by nitric acid. It is also formed whenever
organic matters containing nitrogen are suffered to putrefy in contact with lime; when animal manure and lime are left together, nitrate of lime is formed; the same salt is frequently found in the mortar of old walls, particularly those of stables and similar out-houses. When mixed with a solution of carbonate of potash, both salts are decomposed; nitrate of potash and carbonate of lime being formed. By this process, nitre has often been made in what are called nitre-beds.

241. Muriate of lime, or, more properly speaking, chloride of calcium, is a very soluble salt, easily formed directly from lime and muriatic acid, and remarkable for its great attraction for water (203). It exists in sea-water, and consequently is often found in common salt. Mixed with a solution of carbonate of ammonia, both salts are decomposed, chalk and muriate of ammonia being formed; this is remarkable, because those two salts, when mixed together dry, and heated, form muriate of lime and carbonate of ammonia (155).

242. There is only one other compound of lime of much importance, and which has already been mentioned; this is the phosphate. It is found in small quantity in most plants, and forms a great part of the solid matter of the bones of animals (594, 836, 876).

243. Phosphate of lime is by no means abundant in nature, but it exists in small quantity in a great many substances. Traces of phosphoric acid are found in a great number of rocks and stones, in the
soil, in almost all plants, and in animal matters. It never occurs free, or uncombined, but always in combination with a base, and this is very commonly lime. Thus we always find phosphate of lime in wheat, and all these vegetable substances which constitute part of the food of man and animals; and we find it in a very considerable quantity associated with carbonate of lime, in the bones of all animals, who obtain from it all the substances on which they feed. Phosphate of lime is insoluble in water, but readily dissolves in solutions containing a little free acid.

244. Phosphate of lime is easily decomposed by the more powerful acids, such as sulphuric acid and muriatic acid; in the former case an insoluble sulphate will be formed, and phosphoric acid left in solution; and in the latter, a clear solution will be obtained containing phosphoric acid (195) and muriate of lime, a salt which is also very easily soluble in water.

245. Sulphuric acid does not wholly decompose phosphate of lime, though it takes from it the greater part of the lime which it contains; the phosphoric acid is still combined with a small quantity of lime, constituting a bi-phosphate of lime (840); the results of the action of sulphuric acid, therefore, are sulphate of lime, a super or bi-phosphate of lime, and free phosphoric acid; the two latter may be readily separated from the gypsum by the addition of water, they being very soluble, whilst the latter is very little soluble in that fluid. A mixture of phosphate of lime
and sulphuric acid is frequently called super-phosphate of lime, and is much used as a manure.

246. **Magnesia** is an earth somewhat similar to lime, but far less common and abundant; like lime, it is the oxide of a very combustible metal, which, exposed to the air, soon takes oxygen and forms the earth. The base of magnesia is called magnesium (202). Magnesia occurs usually in connection with lime, forming what is called magnesian limestone, which is a mixture of carbonate of lime and carbonate of magnesia.

247. The commonest forms in which magnesia is found are the carbonate, the sulphate, the muriate, and the phosphate. The carbonate is sometimes found native, pure, but by far more frequently it is mixed with carbonate of lime or limestone, the properties of which it considerably modifies. When carbonate of magnesia is strongly heated, it parts with its carbonic acid, and the pure earth magnesia is obtained.

248. This earth is almost insoluble in water, and far less strongly caustic than lime; it also has less affinity for carbonic acid, so that when left exposed to the air, it does not, like lime, rapidly become converted into carbonate, by absorbing carbonic acid from the air, but remains for a long time caustic; and hence, when a magnesian limestone is burnt in a limekiln, the quicklime obtained of course contains caustic magnesia, which, if exposed to the air, will remain caustic long after the lime has become converted into carbonate; it is for this reason that lime-
stone containing magnesia cannot be used for some purposes, where the caustic powers thus retained are objectionable (873).

249. The carbonate of magnesia, like the carbonate of lime, and indeed all other compounds of that acid, is easily decomposed by any more powerful acid (117).

250. When magnesia or its carbonate is acted on by sulphuric acid, sulphate of magnesia is formed, which is a salt easily dissolved by water, and distinguished by a peculiar sweet-bitter taste. It is found in many mineral waters, and exists in unusually large quantity in some springs near Epsom, in consequence of which it is commonly called Epsom salts. A solution of sulphate of magnesia is readily decomposed by the alkaline carbonates; carbonate of potash, soda, or ammonia, throwing down carbonate of magnesia.

251. The compound of muriatic acid and magnesia, or chloride of magnesium, is also frequently found in mineral waters, and exists in some quantity in sea-water; it is very soluble in water, and therefore, when sea-water is evaporated in order to procure the common salt which it contains, it is never wholly evaporated; but when the greater part has been driven off by heat, and a large quantity of common salt is deposited, the remaining liquor, commonly called bittern, which contains a little common salt, and nearly all the muriate of magnesia, and other easily soluble salts, is drawn off.

252. The muriate of magnesia, like the muriate of
SALTS OF MAGNESIA.

lime, is a very deliquescent salt; that is to say, it has a very strong attraction for water, and hence, when exposed to damp air, it soon becomes moist, and at last we find, instead of the solid salt, a very strong solution of it in water. It is to the presence of a small quantity of these two salts in common sea-salt that its deliquescent properties before alluded to are due (204).

253. The phosphate of magnesia, like the carbonate, is seldom found alone, but usually mixed with the phosphate of lime: thus we find it associated with that substance in small quantity in many animal and vegetable matters, and likewise in the soil; like the phosphate of lime, it is insoluble in water, but readily soluble in dilute acids. Phosphate of magnesia has a considerable tendency to form double salts (112); thus a double phosphate of magnesia and ammonia is very frequently found in plants and animals; it is readily formed artificially by adding phosphate of ammonia to a solution of sulphate of magnesia; it is slightly soluble in water, and has been strongly recommended as a manure. The uses of magnesia are limited, and the most important purposes to which its compounds are applied are as medicines.
CHAPTER IV.

METALLIC ELEMENTS—THE EARTHS—METALS—METALLIC OXIDES, AND SALTS.

254. Alumina, or pure clay, is a very abundant and widely-diffused substance. It occurs native both pure and in combination, but it is most usually met with mixed with another earth, called silex, and combined with a quantity of water. It is sometimes found pure and free from water or any acid, and constitutes the hard gems called ruby and sapphire; but these are very rare, and seldom met with. Alumina is the oxide of a metal called aluminum (202, 225).

255. The properties of alumina, in the ordinary state in which it occurs in the soil, are very different from those of the bases described in the last chapter; it combines with acids to form salts, but is quite insoluble in water, has no caustic powers, and does not absorb carbonic acid from the air. It has a strong attraction for water, and when thoroughly wet, it appears in the form of a very tenacious paste, remarkable for its great plasticity, and the ease with which it may be moulded into any form. All the varieties of clay derive their tenacious property
from the large quantity of this earth which they contain; and the whole of the art of making pottery, bricks, tiles, &c., is in great part dependent on this property of alumina.

256. This earth is able to combine with acids, but the salts which it forms are mostly of very little importance, and we need not inquire into their characters. The only one which is found native is the phosphate, but this is by no means of common occurrence; it is only found in certain districts, and in very limited quantity.

257. The sulphate of alumina is largely manufactured from certain kinds of slate or shale, which contain alumina and sulphur. When sulphate of alumina is mixed with sulphate of potash, the two salts combine and form a double salt, the sulphate of alumina and potash, or common alum. If a little potash is added to a solution of alum, the sulphate of alumina will be decomposed, pure alumina will be separated, and sulphate of potash alone remain in solution (112).

258. Silica, or silex, is the only other earth of much importance besides the three already mentioned; it is found abundantly, both pure and mixed or combined with alumina and other substances, constituting, in a nearly pure condition, quartz, sand, flint, &c., and, when associated with alumina, forming clay and a numerous series of stones; being, in fact, an essential ingredient of the greater number of the hard stones with which we are acquainted.

259. Silica differs from the preceding earth, alu-
mina, in not being a base—in being unable to combine with acids; indeed, it has rather the character of an acid, for it is able to combine with the alkalies so much in the manner of acids, that it is very frequently termed silicic acid. Silica is the oxide of a peculiar substance having many of the properties of metal, and to which the name of Silicon has been given (202).

260. Silica, in its common forms, is quite insoluble in water, unacted on by the air, and, under ordinary circumstances, a very unchangeable substance: when, however, combined with the alkalies, it is easily dissolved in water, and the compound thus formed is sometimes present in very small quantity in mineral waters.

261. When silica is obtained by the decomposition of any of its soluble compounds, it appears in the form of a transparent jelly, which dries into a very fine white powder, like flour; when freshly precipitated from a solution, this jelly is slightly soluble in water and in dilute acids, a property, however, which it loses by being thoroughly dried.

262. The ordinary forms of silica or silicic acid, such as sand or flint, are very little acted on by potash; but when finely powdered silica is mixed with potash or soda, and strongly heated, they melt and form a clear transparent substance, which is in fact glass. Indeed, so strong is the affinity which silica has for potash and soda, that, if fine sand or pounded flints are mixed with the carbonate of either of those
bases, and strongly heated, the carbonic acid is expelled, and the silica and alkali unite to form a glass. Such a compound is called a silicate.

263. Common glass always contains other substances, but the basis of all good glass is this compound of silica and alkali—either silicate of potash or soda. The proportions taken of the two ingredients are always such that the glass obtained shall be perfectly unacted on by water; but if more alkali be employed than is requisite to form a good glass, a silicate will be obtained which is readily soluble in water. A solution thus made is easily decomposed by any acid, as the potash has comparatively but a weak attraction for the silica, and hence that substance is separated from its solution on the addition of almost any acid.

264. Though potash is scarcely able to combine with silica at a common temperature in its usual states, yet when the silica is in an exceedingly fine powder, the alkali is able to dissolve a small quantity; but this action is far slower and less perfect than when the two are strongly heated together.

265. Silica is almost always an ingredient of the soil, and exists there not only in its solid and insoluble form of sand, but also in the soluble condition of silicate of potash or soda; it will easily be seen that as many stones contain silica, they will, whilst gradually decomposing and crumbling down by exposure to the air, constantly add to the soil silica in a very finely divided state, and therefore well adapted to
combine with either potash or soda; and besides, as many stones contain silica in combination with potash or soda, so these stones, in crumbling away, present a constant source of soluble silica.

266. Silica is found in many plants, such as, for example, corn and grasses, the stalks of which mainly derive the strength requisite to enable them to grow erect from the silica which they contain. Plants derive this earth from the soil, and are only able to absorb by their roots the silica, which, by having combined with alkali, has become soluble in water, and is consequently able to enter into the structure of the plant.

267. Silica, or silicic acid, combines with lime, magnesia, and alumina, to form silicates, as well as with the alkalies, potash and soda; these silicates are for the most part insoluble in water, and constitute, either pure, mixed, or combined together, an immense variety of different stones; the action of the atmosphere and other natural circumstances combine to effect the gradual decomposition of such compounds.

268. Besides the three earthy bases already spoken of, and a few others of far less importance, as they are very rare, and only to be found in particular places, there are a numerous series of bases called metallic oxides, several of which are of great importance in the arts, and two of which are almost always present in small quantity in the soil.

269. Most of the common metals, such as iron, lead, and zinc, gradually tarnish, and become rusty
when exposed to the air: the reason of this is that they have a strong affinity for oxygen, and under these circumstances they gradually become covered with a film of an oxide, or compound of the metal with oxygen. Many of these metallic oxides are bases, and form with acids a very numerous series of salts.

270. Modern discoveries have shown that both the earths and alkalies, are, in fact, the oxides of peculiar and very oxidizable metals (202). Hence the earths, alkalies, and ordinary metallic oxides, are all classed together under the general term, base: they combine with acids to form salts; thus gypsum, or sulphate of lime, is a compound of lime, which is the oxide of a peculiar metal, and sulphuric acid. Green vitriol, or sulphate of iron, is a compound of oxide of iron and sulphuric acid; and Cheltenham salts, or sulphate of soda, consists of soda (the oxide of sodium) and sulphuric acid.

271. To speak correctly, green vitriol should be called sulphate of oxide of iron, but such a system would be very inconvenient; it is therefore customary, when speaking of the salts formed by the oxide of a metal, merely to call them by the name of the metal. Hence, when chemists speak of sulphate of iron, and carbonate of lead; they always mean salts of the oxides of those metals; the metals themselves, not being bases, could not combine with the acids to form salts.

272. The salts formed by the combination of the different metallic oxides are called just as if they were
salts of the metals themselves, because the oxides of the ordinary metals have no special names, like potash, and soda, &c.; thus the sulphate of the oxide of lead, for example, is simply called sulphate of lead. When there are two separate oxides of a metal, both of which form salts with acids, that which contains least oxygen is called a protoxide, and that which contains most, a peroxide; the addition of proto or per to the name of a salt, shows whether it is a salt of the protoxide, or of the peroxide; thus the proto-sulphate or per-sulphate, means a sulphate of the protoxide, or peroxide.

273. The most widely diffused and abundant of all the metallic oxides, as well as that which is the most important and valuable in the arts, is the oxide of iron, which exists in different quantities in a great variety of stones, is very common in soils, and is constantly present, though only in small quantity, in the blood of animals, and in the juices of plants.

274. Iron is very rarely indeed found native in its pure metallic state, but is usually met with in the form of an oxide, either pure or combined with carbonic acid, and mixed with alumina and silica. Thus the rich black and red iron ores of Cumberland and other places are nearly pure oxide of iron, whilst the common clay iron-stones, as they are called, of Staffordshire and Wales, are either carbonate or oxide of iron, mixed with various proportions of alumina and silica.

275. The important art of smelting iron is entirely
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a chemical operation, and depends mainly upon the fact that, at a high temperature, carbon has a stronger affinity for oxygen than iron has; and hence, when the native oxide of iron is heated with coal or charcoal, it is decomposed, and carbonic acid gas and metallic iron are the results of the process.

276. When those ores are smelted which consist principally of oxide of iron, they are at once heated with carbon; but when the clay iron-stones are used—and they are the ores most commonly employed—they are first submitted to a preparatory process, something like the burning of lime, in order to expel the carbonic acid gas which they contain; and when thus converted into oxide of iron, they are mixed with carbon and lime, the use of the latter being to combine with the silica or silicic acid and alumina, and form with them fusible silicates called the slag, which greatly assists in the melting and running together of the newly-reduced iron; and, besides, by covering the metallic iron with a glassy coat, they protect it from further oxidation from the oxygen of the air.

277. Iron is able to form two distinct compounds with oxygen, according to the quantity of that element with which it combines: when it is combined with two-sevenths of its weight of oxygen, it constitutes a black substance, which is called the protoxide, and when combined with three-sevenths, forms a brownish-red substance, called the peroxide. These oxides are both bases, and each forms a distinct series
RUSTING OF IRON.

of salts by combining with acids; but the salts formed by the protoxide have always a tendency to absorb oxygen from the air, and thus become converted into the salts of the peroxide.

278. The color of a great many stones and soils is principally caused by the presence of a small quantity of either the peroxide of iron, or of a mixture of both its oxides.

279. The rusting of iron, which proceeds so rapidly when iron is exposed to damp air, is caused by the attraction which the metal has for oxygen. It is very remarkable that iron is unable to combine with the free oxygen always in the air, but is able to take it from water, its compound with hydrogen; for we find that in dry air, iron remains clean and bright for a long time, but very rapidly rusts when exposed to the joint action of air, carbonic acid gas, and moisture, under which circumstances water is decomposed, and oxide of iron formed.

280. The rust of iron is not a pure oxide, but commonly a mixture of both oxides with a portion of carbonate, or compound of the protoxide with carbonic acid. Rust generally contains a considerable quantity of ammonia, for which substance oxide of iron has a strong attraction; when oxide of iron is thrown down from a solution which contains it, by ammonia, it is very difficult to expel the whole of the ammonia from the precipitated oxide even by strongly heating it.

281. Although iron cannot combine with the free
oxygen of the air at ordinary temperatures, yet when strongly heated it rapidly absorbs oxygen, and then becomes converted into a black scaly oxide; whenever a piece of iron is heated in the fire, a quantity of a brittle, black oxide is formed on its surface, which easily rubs off the iron—and hence this metal is gradually worn away by exposure to fire; it is from this reason that the iron pokers, fire-bars, and other things much exposed to the fire, gradually get thinner and thinner, from the constant oxidation of the surface whilst hot, and the removal by rubbing of the brittle coat of oxide thus formed.

282. When a piece of iron is very strongly heated, it at last begins to burn; that is to say, the combination of the external part with oxygen goes on so rapidly, and evolves so much heat, that the whole mass of iron is kept sufficiently hot to continue this process of combination, and in consequence the iron glows brightly, gives off abundance of sparks, and runs down in drops of the melted oxide, for some time after it has been removed from the fire in which it was heated. In fact, iron, when thus strongly heated, would catch fire and continue to burn like charcoal, if it were not that the crust of oxide formed protects the metal from further oxidation, and soon stops its combustion.

283. One of the most common and abundant of the ores of iron is called pyrites, which is a compound of iron and sulphur; it is not used in the manufacture of iron, because it is very difficult to
separate the sulphur completely from the iron, and
the native oxides and carbonate are far more con-
venient sources of the metal.

284. Pyrites, or sulphuret of iron, is, however,
a substance of considerable importance in the arts,
being one of the sources of sulphur, which is ob-
tained by heating pyrites in an oven, so constructed
that the sulphur, which is expelled in the state of
vapor from the pyrites, is cooled and condensed
into the solid form in a different part of the oven.

285. Pyrites, when exposed to the air, soon crum-
bles down, and undergoes a very curious change, in
consequence of absorbing and combining with oxygen.
Both the iron and the sulphur combine with oxygen,
and form oxide of iron and sulphurous acid; and hence
the result of this action is sulphate of iron, or com-
mon green vitriol, a salt much used in the arts for a
variety of purposes.

286. Pyrites is most abundantly found in the form
of variously-shaped balls imbedded in chalk; to
which the common name of "thunderbolt" is very
improperly applied. As the chalk-hills on the sea-
side gradually wear away, from the action of the sea
and weather, these balls of pyrites get exposed to
the air, and fall down on the beach, whence they are
collected for the use of manufacturers. Many springs
of water contain a small quantity of iron, in conse-
quence of which they have a peculiar inky taste;
this is usually derived from the gradual oxidation of
SULPHATE OF IRON.

129

sulphuret of iron; and from the same reason soils also sometimes contain traces of this salt.

287. Sulphuret of iron is likewise very commonly found in coal, being sometimes dispersed throughout it in the form of little yellow shining particles, and sometimes as layers or lumps of the solid sulphuret; its presence in coal is for some purposes highly objectionable; because, whether the coal is burnt in its crude state, or after being converted into coke, a quantity of sulphurous acid gas is always formed by the imperfect combustion of the sulphur; and that nauseous-smelling gas causes serious mischief in several operations in the arts, and consequently in such cases, coal free from sulphuret of iron can only be employed. It is the presence of this substance in coal that causes the strong suffocating smell of sulphurous acid sometimes given out by coal and coke fires (173).

288. The oxides of iron are quite insoluble in water, but many of the salts of iron, like the sulphate, are readily soluble in it; the solutions of these salts are all decomposed when alkali is added to them; this combines with the acid, and the oxide of iron is separated as an insoluble powder.

289. The most important of the salts of iron is the sulphate, or common green vitriol: it may be formed by acting upon iron by dilute sulphuric acid; when this is done, a large quantity of hydrogen gas is given off, in consequence of the decomposition of a portion of water; the oxygen combines with the
iron to form oxide of iron, which unites with the acid to form proto-sulphate of iron, whilst the hydrogen escapes. Green vitriol is, however, made on a large scale, principally from pyrites, in the manner just described (285).

290. Sulphate of iron, when pure, is a green transparent salt, wholly soluble in water; exposed to the air it becomes brown and earthy-looking, being partially decomposed, owing to the absorption of oxygen and formation of peroxide; when strongly heated, it is wholly decomposed, water and sulphuric acid being given off, and oxide of iron left. When pure proto-sulphate of iron is decomposed by an alkali, a gray or black precipitate is formed of the protoxide; after a short time this precipitate becomes red, having absorbed oxygen, and become peroxide.

291. The number of different metals known to chemists is very considerable, amounting in all to forty-six; but of these the greater number are comparatively rare, and of little importance, not being used for any practical purpose, and consequently chiefly interesting in a scientific point of view. It will, therefore, be sufficient to consider briefly the leading characters of eight of the most important of them (19).

292. Gold is found native in a pure and separate state; it has very little affinity for oxygen, never tarnishing or showing any tendency to oxidize in the air either at common temperature or when strongly heated. It is insoluble in acids, except in a mixture
of nitric and muriatic acid, which dissolves it; such an acid contains free chlorine, for the oxygen of the nitric acid takes the hydrogen of the muriatic acid, and sets free the chlorine; hence, we learn that the only solvent of gold is, in fact, a solution of chlorine.

293. When a solution of chloride of gold, thus obtained, is mixed with a solution of potash or soda, a dark-colored precipitate falls: this is an oxide of gold; when heated, it parts with the oxygen which it contains, and pure gold is left. The soluble compounds of the metal gold are all very easily decomposed, because the metal has but a very feeble affinity for oxygen, chlorine, &c.

294. Silver, like gold, occurs native in its pure metallic state; but it more commonly is found combined with sulphur, as a sulphuret. It resembles gold in having a feeble affinity for oxygen, so that it does not oxidize either at common temperatures, or when heated, and most of its compounds are easily decomposed, the silver reassuming the metallic state. Silver readily dissolves in nitric or sulphuric acid, and the solutions obtained yield oxide of silver, when decomposed by an alkali; the oxide of silver is a salifiable base, and combines with acids to form salts.

295. The compounds of silver are nearly all decomposed by mere exposure to light. So feeble is the affinity which it has for most other substances, that the greater number of its compounds are decomposed or reduced by mere exposure to sunshine. This fact is interesting as an illustration of the chemical
powers of light, and, further, as giving us the foundation of that very beautiful art of producing pictures by means of light—the Daguerreotype. The salts of gold, also, and some of the compounds of mercury are very easily decomposed, when exposed to the action of light.

296. Silver has a very strong affinity for chlorine, and the chloride of silver is quite insoluble in water; hence silver cannot be dissolved in muriatic acid; and if muriatic acid, or any solution containing chlorine, either free or in combination, be added to a solution of silver, the whole of the metal will be thrown down as insoluble white chloride. A solution of nitrate of silver is, therefore, a very useful test to ascertain the presence of chlorine in any solution.

297. Silver has also a strong affinity for sulphur, combining eagerly with it and forming a shining gray brittle substance. The tarnish which we see on old silver is a thin coat of sulphuret, formed by the sulphuretted hydrogen which generally exists in the air of towns. Gold and silver are frequently called noble metals, from their having no tendency to oxidize when exposed to the air.

298. Mercury, or quicksilver, likewise, is sometimes found native in the metallic state, but by far most commonly as a sulphuret; it is obtained from its ore by heating it with a mixture of iron filings and lime; these substances combine with the sulphur; and the mercury, being a volatile metal, is obtained by a process of distillation.
299. Mercury is fluid at all ordinary temperatures, but, by exposure to very intense cold, it may be frozen into a brilliant hard solid, looking like silver; when heated nearly to redness it boils, rises in vapor, and may be distilled, just like water. When mercury is kept for some time at a heat very near its boiling point, it slowly absorbs oxygen, and becomes converted into a red earthy-looking oxide. Mercury is easily oxidized and dissolved by the strong acids; it forms two oxides, and both of them are salifiable bases—the protoxide is black, the peroxide red; when strongly heated, these oxides are decomposed into metallic mercury and oxygen gas.

300. Chlorine acts strongly on mercury, and forms two chlorides, corresponding in composition to the two oxides; the protochloride, or calomel, is a powerful and valuable medicine—the perchloride, or corrosive sublimate, a violent poison. The former is insoluble in water—the latter soluble; they are both easily decomposed by alkaline solutions—calomel yielding the black protoxide, and corrosive sublimate the red peroxide of mercury.

301. Corrosive sublimate has been a good deal used to prevent the dry-rot of wood, cordage, &c.; it combines with some forms of organic matter, and renders them less prone to change. Sulphur and mercury easily combine, and form a beautiful red compound, vermilion, or sulphuret of mercury; it occurs native as cinnabar, the chief ore of the metal.

302. Copper exists naturally in the pure metallic
state, but chiefly as a sulphuret, constituting copper pyrites; it is obtained from this ore by roasting, when the sulphur is gradually driven off and an impure oxide of copper left, which is subsequently strongly heated with charcoal, to reduce it to the metallic state. Copper has a considerable affinity for oxygen, which it absorbs from the air at common temperatures. Oxide of copper is a black substance, readily obtained by heating copper in the air, or by decomposing any of its salts, such as the sulphate, by an alkali.

303. The salts of copper are mostly of a blue or bluish-green color; they are all decomposed by alkaline solutions, a blue hydrated oxide of copper being precipitated; if ammonia be employed, no precipitate is obtained, or, if formed, easily dissolves, because the oxide of copper is soluble in solution of caustic ammonia, forming a very beautiful deep-blue liquid: this property is useful in testing for the presence of copper.

304. Sulphate of copper, blue or Roman vitriol, may be formed directly. It is manufactured on a large scale, like the sulphate of iron (285); by exposing the roasted sulphuret to the air, it absorbs oxygen, and is converted into the sulphate. It is a bright-blue salt, easily soluble in water, and used for several purposes in the arts. It is frequently employed as a steep for seed-corn.

305. ZINC is never found in the metallic state; its ores are calamine, which is a carbonate, and blende,
a sulphuret; it is obtained by roasting the ores, which in the one case drives off the carbonic acid, and in the other dissipates the sulphur; the roasted ore is then mixed with charcoal and distilled; the metal is volatile at a very high temperature.

306. When zinc is strongly heated in the air, it burns with a bright flame, and is converted into a white oxide, which may also be obtained by acting on the metal by an acid; the metal easily oxidizes and dissolves, forming a salt from which the oxide may be obtained, on the addition of an alkali. Sulphate of zinc, or white vitriol, is a white salt, very easily soluble in water, and made either from the metal and sulphuric acid, or from the native sulphuret.

307. Zinc has a strong affinity for chlorine; by dissolving the metal in muriatic acid, a solution of the chloride is obtained. It is very soluble in water, and has been much employed to preserve wood and canvas from decay.

308. Tin occurs native almost entirely as an oxide, from which the metal is obtained pure by merely heating with charcoal. Heated in the air it easily oxidizes, and by the action of acids a protoxide and peroxide may be procured. Peroxide of tin, the same substance which occurs native as tin-stone, is artificially made, as a polishing powder, being called "putty powder." Oxide of tin has a remarkable affinity for coloring matter, and hence is much used by dyers in fixing colors. The bisulphuret of tin is
of a beautiful golden yellow color, and is employed for various ornamental purposes in the arts, under the name of mosaic gold.

309. MANGANESE, like tin, is found only as an oxide; it is a metal in many respects considerably resembling iron, but having a much stronger attraction for oxygen, and consequently obtained in the metallic state with very great difficulty; it forms several oxides, only one of which, however, is a base; some of its salts are employed in dyeing; and the peroxide is much used in the process for obtaining chlorine (191). Manganese is very often found associated with iron in rocks and stones, and not unfrequently exists in minute quantity in soils; it is comparatively speaking, however, a rare metal.

310. LEAD is obtained almost exclusively from the native sulphuret; it never is found in the pure metallic state. The sulphuret is roasted, by which the sulphur is gradually driven off, and an impure oxide formed, which remains mixed with a large quantity of sulphuret; this is then smelted with small coal or other carbonaceous matter. Lead has a strong affinity for oxygen, in consequence of which it tarnishes slowly at common temperatures, and quickly when melted. At a red heat, lead is gradually converted into a yellow substance called litharge—this is an impure protoxide; if still longer exposed to the air and heat, it absorbs more oxygen and becomes red lead or minium; besides these two, there is yet a third oxide of lead, which, however, cannot be formed
by further heating red lead under the influence of air, but which is easily made by acting on red lead by nitric acid; the lead then acquires a third portion of oxygen and becomes a dark brown peroxide.

311. Of these three oxides, only one, the protoxide, is a base; the other two oxides, when acted on by acids, part with a portion of their oxygen, and pass into the state of protoxide before they can combine with the acid. The most important of the salts of lead is the carbonate, or white lead, a substance better suited than any other which is known for the manufacture of white paint; it is made either by decomposing a soluble salt of lead by an alkaline carbonate, or by exposing lead to the action of the vapor of vinegar and carbonic acid gas. The acetate of lead, likewise, is manufactured on a large scale, and used for various purposes in the arts.

312. Lead has a strong affinity for sulphur, and in consequence of this most of the salts of lead are decomposed by sulphuretted hydrogen; the blackening of white paint is due to this cause. As the sulphuret of lead is decomposed by chlorine, white paint thus blackened may be cleaned by muriatic acid; this converts the black sulphuret into a white chloride, though it never looks so white as the carbonate did before.

313. Lead is acted on by pure water in a very remarkable way, being oxidized and dissolved with great facility; this is not the case with common water containing salts of lime, &c. (75). Rain-water, or very
pure water, kept in leaden cisterns, or passing through leaden pipes, often dissolves so much lead as to become unwholesome, or even poisonous; and for the same reason leaden covers to cisterns are equally objectionable, because the water which condenses on the cover, being of course pure (73), corrodes and dissolves the lead, and dripping down again into the cistern, contaminates the water, which otherwise might have remained pure and wholesome: a great deal of disease is probably caused by want of attention to these facts.

314. Many of the metals, when melted together, combine to form what are called alloys, or mixed metals; some of these appear to be regular definite compounds, though others are obviously mere mixtures. The most important of the alloys are gold and copper, and silver and copper, which are harder than gold or silver alone; these alloys are used for plate, coin, &c. Zinc and copper, or brass; tin and copper, or bell-metal; tin and iron, or common tin plate, which is often supposed to be merely tin, though it really consists of thin plates of iron, alloyed on the surface with tin, so as to have the strength and stiffness of the iron, together with the freedom from rusting of the tin. Zinc and iron, or “galvanized iron,” as it is frequently called, is iron alloyed or covered on the surface with a film of zinc, which greatly protects it from corrosion; and, lastly, lead and tin, or pewter, and common solder.
CHAPTER V.

ORGANIC MATTER—THE NATURE AND COMPOSITION OF VEGETABLE SUBSTANCES.

315. We have now very briefly described most of those substances which are of importance in studying vegetable chemistry. Before explaining the action which they have on the growth of plants, it will be proper to go a little more into detail respecting the nature of organic matter; that is to say, the various compound substances which constitute the bodies of animals and plants.

316. Most of the substances hitherto described, such as water, ammonia, carbonic acid, common salt, &c., consist wholly of two elements, and are therefore sometimes called binary compounds. On the contrary, all animal and vegetable substances consist of three or four elements, and are consequently termed ternary or quaternary compounds. It has been already stated that plants and animals are composed of carbon, oxygen, hydrogen, and nitrogen (92). It is very important to understand clearly the nature of the compounds formed by these elements.

317. When we endeavor to analyze a plant, that
is to say, to ascertain of what it is composed, we find that the greater part of it consists of carbon, oxygen, hydrogen, and a small portion of nitrogen, combined together. When we burn it, or in any other way weaken the affinity which the elements have for each other, they separate, and, by combining together, generally form water, carbonic acid gas, and ammonia.

318. When a plant is boiled in water, it is found that part of the plant dissolves in the water, whilst part remains insoluble, and we are unable by long-continued boiling to make the whole of it dissolve in the water. These, then, are two great divisions of vegetable matter—that which is soluble in water, and that which is not. By very simple operations of this kind it is easy to discover that plants are composed of a variety of different compound substances, readily distinguished from each other by the different properties which they possess. Of those which are usually found in all plants, the most abundant are called lignin, starch, gum, sugar, gluten, and albumen. The four former consist of carbon, oxygen, and hydrogen alone, whilst the two latter contain, in addition to these elements, a portion of nitrogen.

319. **LIGNIN**, or pure woody fibre, exists in almost all plants; it constitutes the greater part of the stem, wood, bark, and branches of trees; and is present, though in smaller quantity, in the leaves and flowers of trees, shrubs, and succulent plants. It is the most solid constituent of plants, giving strength to those parts in which any quantity of it exists. It may
WOODY FIBRE.

readily be separated from the other matters with which it is associated, by bruising and long-continued boiling in water and spirit; by these means the softer or more soluble substances may be separated, and pure lignin is left. In the process for preparing flax, the stems of the flax plant are allowed to remain in water for some time; the green soft parts decay, and at last nothing but the lignin or woody fibre is left.

320. Pure lignin is a white, tough, fibrous substance, composed of an infinite number of very fine threads or fibres, perfectly insoluble in water, and not at all altered by keeping in dry air. When heated in the air it soon turns brown, being partly decomposed: if it be still further heated, it takes fire and burns with a bright flame, the results of its combustion being water and carbonic acid gas. Its composition is—

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321. The woody fibre of plants is not pure lignin. It consists of cells and tubes more or less incrusted and filled up with resinous and other matter, which renders them stiff and hard. The fibre of fine linen or cotton may be taken as an example of tolerably pure lignin; because the foreign matters originally associated with it have been almost entirely removed,
by various processes which the fibre has undergone, in the different operations of the manufacturer.

322. Pure lignin, or cellulose, as it is sometimes termed, is scarcely at all acted on by acid or alkaline solutions, either hot or cold, unless they are very concentrated. Strong sulphuric acid converts it into gum (359). Strong nitric acid produces a very remarkable effect on lignin, and changes it into gun-cotton, or xyloidine. In this action the lignin is partially decomposed, and a portion of its oxygen and hydrogen, in those proportions which would form water, is replaced by some of the oxygen and nitrogen of the nitric acid.

323. Gun-cotton is best formed by steeping pure clean cotton wool, quite free from all oily matters, either in very strong nitric acid, or in a mixture of nitric and sulphuric acids; the addition of the latter is merely made for the purpose of rendering the nitric acid stronger by extracting the water which it contains. The cotton increases considerably in weight, and when well washed and dried is found to be highly explosive. It detonates at a heat a very little above that of boiling water, with a bright flash, and is wholly converted into carbonic acid, nitrogen, steam, &c.

324. Woody fibre has a strong affinity for various coloring matters, and also for some particular metallic oxides, especially alumina, and the oxides of iron and tin; solutions of these substances are consequently much used by dyers. The cotton to be dyed is first impregnated with the mordant, as these substances
are called, and subsequently dyed by being immersed in the colored solution (542).

325. When wood is distilled, or roasted in close vessels, various substances are formed; amongst these are vinegar or pyroligneous acid (488), and wood or pyroxylic spirit. The latter is a volatile, pungent-smelling liquid, which burns easily, with a pale flame like spirit of wine. It is much used by hatters, varnish-makers, and others, as a solvent for resins.

326. Starch is almost always found in considerable quantity in all parts of plants. When pure it is a white powder, insoluble in cold water, but readily dissolving in that fluid when boiling hot. There are many different varieties of starch, distinguished from one another by some peculiar property, and which have received various names, according to the plant from which they are obtained. Common wheat starch, which exists in large quantities in the seed of wheat, is a good example of the general character of this substance.

327. When starch is dissolved in hot water, it forms a thick, viscid, semi-transparent liquid, which if evaporated leaves a yellowish, transparent, horn-like substance, which readily swells and softens in cold water. Hot water, though it greatly changes the appearance and properties of starch, does not in any way alter its chemical composition.

328. Starch is obtained from wheat by steeping it in water, and subsequently squeezing and washing the softened grain, and then allowing the milky
liquid thus obtained to stand for some time. The cells and membranes which inclose the starch are thus broken and destroyed, and as the starch is insoluble in cold water, it is then easily washed out, and separated from the husk and other insoluble parts of the grain; the water is then allowed to settle, the starch falls to the bottom, and is collected and dried.

329. This softening and destruction of the membranes of the seed is greatly assisted by the presence of a small quantity of certain acids in the water, which dissolves the gluten; a small quantity of lactic acid is always produced when grain is thus steeped in water, and this is essential to the manufacture of starch. A similar effect may be produced by a weak alkaline solution, and accordingly a dilute solution of caustic soda is employed by starch-makers to soften particular sorts of grain, such as rice, Indian corn, &c. (377).

330. When grated potatoes are placed on a sieve, under a stream of water, a very large quantity of starch may be washed out. The starch will soon settle to the bottom of the water, the soluble matters of the potato will be dissolved, and at last there will remain on the sieve little else beside the lignin or fibre which the potatoes contained, together with a quantity of starch which cannot be separated from the fibre by mere washing.

331. Potato-flour, arrowroot, tapioca, and sago, are all varieties of starch; similar substances are likewise obtained from Iceland moss, the seed of the
chestnut, and many other plants. They all agree in general chemical characters with wheat-starch, and, like it, consist of carbon, oxygen, and hydrogen, rather less than one-half of their weight consisting of oxygen. Starch consists of—

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332. When starch is examined by the help of a magnifying glass, it is found to consist of variously shaped transparent little grains; marked in a very peculiar manner. These grains vary in shape and size according to the plant from which they are obtained; this fact renders it possible to ascertain whether any particular sample of starch is arrowroot, potato starch, wheat starch, or a mixture of several kinds.

333. The quantity of starch obtained from different plants varies very much. Good wheat generally yields from 70 to 75 per cent.; barley contains nearly 80; oats and rye, 60 to 65; beans 40 to 45; peas about 50; potatoes 15 to 20; arrowroot, 20 to 25. Almost all the seeds and grain used as articles of food, such as wheat, barley, oats, rye, maize, rice, millet, &c., contain a large quantity of starch. Semolina is prepared from hard Italian wheat; it is the hard granular particles which escape the action of the mill-stones. Macaroni is the dried paste of hard
wheat, which is pressed out from a box through apertures, which gives it its hollow form. The polenta of the Italians is prepared by roasting maize when the seed is only half ripe.

334. There is a peculiar modification of starch which exists in the tubers of the Jerusalem artichoke, dahlia, and many similar plants, and is called inulin; it contains a rather smaller proportion of carbon than common starch does.

335. Under the name of GUM are included several substances, which differ considerably in their nature and properties. They all agree in being tasteless, or nearly so; but some dissolve readily in water, and form a clear, transparent solution; whilst others do not dissolve, but merely soften and swell up in water, forming a gelatinous mass. Gum Arabic is a good example of the first kind, whilst common cherry-tree gum belongs to the insoluble variety of gum.

336. Gum is a natural exudation from many plants, appearing on their surface in the form of transparent drops or tears, which dry and harden in the air. Most fruits contain a considerable quantity of gum, and exudations of gum are frequently found on many fruits, particularly the plum. There is a peculiar substance resembling gum, to which the name of pectine is given, and which exists in most fruits; it gelatinizes with water, and gives to fruits the property which they have of forming imperfect jellies. Black currant and apple jelly consists chiefly
of pectine. The same form of gum is found in the carrot, parsnip, and many similar roots.

337. A solution of gum in water is called mucilage; comparatively a small quantity of gum renders water thick and slimy: such a solution feels sticky to the fingers; it dries slowly, and leaves a small quantity of gum on the surface of any substance over which it has been spread. Gum is insoluble in spirit of wine; hence mucilage is precipitated, and the gum it contains thrown down as an insoluble curd when mixed with spirit. The juices of many plants are sticky from the quantity of gum which they contain.

338. Gum consists of the same elements as starch, namely, oxygen, hydrogen, and carbon; but it contains a rather larger proportion of oxygen than starch does. Gum consists of—

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<td>Hydrogen</td>
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339. Associated with gum in many plants, there is found a substance which dissolves easily in water, and like gum, forms a sticky thick solution; it is distinguished readily from gum, however, by possessing a sweet taste, whilst gum is insipid or tasteless; it is called sugar or saccharine matter, under which names a considerable variety of different substances are included.
SUGARS are divided into two classes: those which are crystallizable, and those which are uncrystallizable: that is to say, those which, when their solution in water is evaporated, are obtained in the form of regular-shaped little grains, like common cane sugar, and those which under these circumstances do not form regular grains, but remain thick viscid-liquids, like treacle. The crystallizable sugars are divided into two classes—cane sugar, and grape sugar; these two varieties differ slightly in composition, the latter containing rather more hydrogen and oxygen in proportion to its carbon than cane sugar; they differ very considerably in chemical properties, grape sugar only being capable of undergoing fermentation (365).

Sugar exists in a great many vegetables, but it very frequently happens that there are so many other substances present, that the sweet taste of the sugar is quite hidden. It is only in those plants which contain a very large proportion of sugar, or which do not contain any strong-tasted substances, that we are able to recognize sugar by its sweet taste. Most ripe fruits contain a large quantity of sugar; it is likewise found abundantly in the sap of a good many trees, from some of which, like the sugar maple, it is procured in such quantities as to be extracted as an article of commerce.

The composition of sugar is similar to that of starch: the proportions in which its three elements, oxygen, hydrogen, and carbon, are united together
are very nearly the same as in starch. The varieties of sugar contain slightly different proportions of these elements; thus the composition of cane sugar is not precisely the same as that of sugar obtained from grapes, beet-root, or other plants. Cane sugar consists of—

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The composition of grape sugar is—

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<tr>
<td><strong>Total</strong></td>
<td><strong>100%</strong></td>
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343. The manufacture of sugar from the sugar-cane is chiefly carried on in tropical countries. The canes are cut in pieces, and crushed in a rolling mill, so as to squeeze out the juice. The juice thus obtained is mixed with a small quantity of lime, and rapidly heated to the boiling point. The scum which separates is collected and removed, and when the juice is sufficiently evaporated, it is allowed to cool that the sugar may separate. The uncrystallizable syrup, or molasses, is drained off from the crude sugar thus obtained, and which is termed raw or Muscovado sugar.

344. Sugar is refined by dissolving the raw sugar
in a proper quantity of water; a portion of albumen is added, either blood or white of egg; the solution is then heated, and the albumen, as it coagulates and separates, entangles and removes all the solid impurities which the sugar contained. The solution is purified and bleached by filtration through charcoal (162), carefully evaporated, and then allowed to crystallize. The evaporation of the cane juice must be carried on as rapidly as possible, but at the same time, exposure to a high temperature is very objectionable, because the higher the heat, and the longer the juice is exposed to it, the greater will be the quantity of molasses formed; a number of ingenious arrangements have consequently been devised to facilitate the evaporation of the solution of sugar at the lowest possible temperature.

345. Sugar is readily soluble in water, and also dissolves, though less easily in alcohol. When a strong hot syrup is suffered to cool slowly sugar is deposited in large and regular crystals. In the manufacture of sugar-candy, as these crystals are termed, the syrup is left quiet, and strings are suspended across it, which assist in the deposition and formation of large and clear crystals. In ordinary sugar-refining, the formation of crystals is not desired, and therefore the solution is continually stirred whilst it is cooling. When sugar is melted, it loses its crystalline character, and becomes barley-sugar: if a very high heat is employed to melt the sugar, it becomes converted into caramel, or burnt sugar; this
ALBUMEN.

has a dark brown color, and is used to give color to brandy and other similar spirits.

346. **ALBUMEN** and gluten in many respects are very similar; they exist in plants in smaller quantities than the three substances already described; but they are nevertheless most important, and of especial interest, as their presence in vegetables is essential to their value as food (597).

347. When the clear juice of any plant is boiled, there usually collects on the surface a thick green scum, which may easily be separated by straining the liquor through linen. This scum consists almost entirely of albumen and gluten, the former being a distinct proximate vegetable principle, the latter a mixture of two separate principles, fibrin and gliadine, with oil; they are associated together, and exist in greater or less quantity in almost all plants. In general, the seeds of plants contain even a larger relative proportion of these substances than the plants themselves. A modification of these substances, which is found in many plants, is called legumine or vegetable caseine, because it is almost identical with the peculiar principle of milk, called caseine (568).

348. The flour of wheat contains a considerable proportion of gluten; it may be readily separated from flour by tying a portion of thick paste in a piece of linen, and then kneading the paste thus enclosed in linen under a stream of cold water; by this process all the starch will gradually be washed out,
and at last there will remain in the linen nothing but gluten.

349. Gluten, when thus obtained, is a grayish white, soft, solid substance, elastic and tough, and almost resembling a piece of animal skin in appearance; it may be dried by carefully warming it, till all the water which it contains is evaporated; and when dry may be preserved for a long time without its undergoing any change. By boiling gluten in alcohol it is separated into three substances—pure vegetable fibrin, an oil, and a tough glue-like substance called gliadine. Vegetable fibrin is insoluble in water, soluble in dilute alkaline liquids, very prone to decompose, and apparently identical with animal fibrin in nature and composition.

350. Albumen, fibrin, caseine, and gliadine, contain nitrogen; and all organic substances containing nitrogen have a great tendency to putrefy. Albumen consists of—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>5501</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>723</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>1592</td>
</tr>
<tr>
<td>Oxygen with Phosphorus and Sulphur</td>
<td>2184</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10,000</strong></td>
</tr>
</tbody>
</table>

351. Vegetable fibrin consists of—

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<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>5460</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>730</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>1581</td>
</tr>
<tr>
<td>Oxygen with Phosphorus and Sulphur</td>
<td>2229</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10,000</strong></td>
</tr>
</tbody>
</table>
ORGANIC CHANGES.

352. It is certainly a very surprising fact that so many different substances should be formed by the combination of the same elements in different proportions. Nothing can well be more dissimilar than oil and sugar, flax and starch; yet it is easily proved that they all consist of the same elements—carbon, oxygen, and hydrogen.

353. The knowledge of this might naturally lead one to suppose, that if the whole difference between such substances consists in the relative proportions of carbon, oxygen, and hydrogen which they contain, it might be possible, by some chemical operation, to take away a small portion of carbon or hydrogen, and thus, by altering the relative proportions of the elements, to convert starch into gum, or flax into starch. Now this can really be done; and strange as it may appear, it is nevertheless true, that, by very simple means, it is easy to change gum, starch, and lignin, &c., into each other.

354. The various vegetable substances, though so different in properties, are very similar in chemical composition, and may for the most part be readily converted or changed into each other, by simple means. They may, when pure, be preserved unchanged for an unlimited time, if quite dry; but when exposed to air and moisture they sooner or later begin to decompose. Those which consist of carbon, hydrogen, and oxygen alone, are far less prone to decompose than those which contain in addition nitrogen; and these latter, when decaying, pos-
sessed the singular property of causing substances which do not contain nitrogen to decompose or change likewise.

355. When vegetable substances, such as lignin or sugar, are burnt in the air, water and carbonic acid are produced; precisely the same substances are formed by decay, as by combustion. When gluten and albumen are burnt, water, carbonic acid, and ammonia are formed.

356. The transformation of one organic substance into another, is a very interesting and remarkable operation. It is plain that such changes must be constantly going on in the organs of plants, and hence everything which throws light on the nature of these transformations is of interest.

357. When the chemical affinity which unites the elements of any compound substance is feeble, and especially when the compound consists of several elements, there is always a considerable tendency to decompose, and the elements are very prone to arrange themselves in simpler, and more permanent combinations. Very slight circumstances therefore are able to determine the decomposition of such compounds, and the change may be brought about in various ways.

358. Some of these transformations are spontaneous, some require the presence of a particular substance, which however has no direct chemical action on the compound itself, or on any of its elements; and many are induced, or caused by the example of
some other compound, which is itself in an active state of decomposition.

359. When strong sulphuric acid is poured over lignin, it rapidly changes it into gum; after a few minutes the fibrous character of the lignin disappears, and a thick slimy substance is formed; if the acid is then diluted with water, and chalk be then added to neutralize the acid, sulphate of lime will be thrown down, and a solution of gum is obtained. If the acid is left in contact with the gum it soon begins to char it, and finally completely decomposes it.

360. Starch may also be converted into gum by a very simple process; when starch is heated considerably above the boiling point of water, it is changed into a kind of gum; large quantities of starch are thus made into gum by roasting, and used by the calico-printers under the name of British gum; by chemists it is called dextrine. The same change may be effected by boiling gelatinous starch with dilute sulphuric acid; it soon loses its gelatinous character, becomes sticky, and is converted into gum. Starch also undergoes this change when acted on by diastase (685); a small quantity of this substance, added to a considerable quantity of gelatinous starch, rapidly converts it into gum.

361. If, having converted a portion of lignin into gum by the action of strong sulphuric acid, we add a quantity of water so as to dilute the acid, and then boil the whole together for some hours, the gum will gradually be changed into sugar; by this process, then,
we may convert flax, hemp, or woody fibre of any kind, even old rags, for they consist chiefly of lignin, into sugar. When the change is complete, the sulphuric acid may be neutralized by the addition of a quantity of chalk; sugar, and a little sulphate of lime, will then be left in solution. By continuing to boil starch with dilute sulphuric acid, in the manner just described, after it has all been converted into gum, sugar is also obtained.

362. In these very curious transformations, the sulphuric acid employed does not undergo any alteration whatever: none of it is decomposed, it retains all its powers unimpaired, and it is easy to recover, at the end of the operation, the whole of the acid originally employed. It does not abstract anything from the substances with which it is boiled, nor does any part of the acid unite or combine with them.

363. All the effect which the acid produces is, that its presence under these circumstances causes a change in the chemical nature of the substances with which it is boiled. In the case of starch, it is thus enabled to take up and combine with about one-sixth of its weight of water, or rather of oxygen and hydrogen, in those proportions which would form water, and in so doing the starch is converted into sugar. The same effects may be produced with many other acids besides the sulphuric (373).

364. Sugar, spirit of wine, and vinegar consist of the same elements: they contain oxygen, carbon, and hydrogen; but the elements are united in different
proportions: the properties of these three substances are as opposite as they well can be, and yet, the whole difference is in the proportions of their elements. It is well known that, when a solution of sugar ferments, it is changed into spirit; it parts with a portion of carbon and oxygen, and the elements left constitute spirit; hence, during fermentation, carbonic acid is given off. Again, when spirit and water is exposed to the air and moderate warmth, it soon changes into vinegar: this change is wholly effected by the absorption of a little oxygen from the air.

365. Fermentation is a very singular process, and a knowledge of the effects which it produces, enables us to understand many changes which would otherwise appear incomprehensible. Under ordinary circumstances, pure sugar, dry or dissolved in water, may be kept for a long time without its undergoing any change; when, however, it is mixed with a small quantity of certain decomposing matters containing nitrogen, it ferments, and is changed into spirit. The substance added, does not combine with the sugar or its elements, but, whilst itself decomposing, it causes the sugar also to change.

366. Fermentation, then, is the spontaneous decomposition of a substance, occasioned by the presence of a small quantity of decomposing matter. The yeast, or ferment of beer, possesses the power of inducing the decomposition of sugar and similar substances when mixed with them. Common yeast, and
all substances which possess the power of causing fermentation, contain nitrogen; they are compounds of carbon, oxygen, hydrogen, and nitrogen, and are accordingly very liable to decompose. During the fermentation of any liquid, the azotized matters present, which constitute the ferment, and are in an active state of decomposition, gradually separate, either rising to the surface or falling to the bottom. This ferment, yeast, or barm, may then be removed and added to any other solution, the fermentation of which it is desired to excite. In most cases, there is more azotized matter present than is required to induce the fermentation of all the sugar, and hence there is generally a quantity of ferment left over, when the operation is finished.

367. Most varieties of sugar are capable of undergoing fermentation, but it appears that, in all cases, they previously pass into a state of grape sugar. When a solution containing sugar and ferment is kept for some time at a moderate temperature, the sweet taste of the sugar gradually disappears, much gas is given off, the mixture froths up from the escape of carbonic acid, and, when the fermentation is complete, spirit is found in place of sugar. When such a fermented liquor is distilled, the spirit passes over before the water, being more volatile; at least, the first portions of the liquid which pass over contain the greater part of the spirit.

368. Common spirit of wine is a mixture of alcohol and water; by repeated distillation with substances
which have a strong affinity for water, pure alcohol may be obtained; it is much lighter than water, and boils at a lower temperature than that liquid. Alcohol consists of carbon, oxygen, and hydrogen; but contains in proportion less carbon, and much less oxygen, than starch or sugar do.

369. In expressing the composition of starch, sugar, and the other principles of plants, the proportion of the different elements, per cent., has been given; for some reasons, however, it is more convenient to calculate the number of equivalents of each constituent; and this is not difficult to do when the composition of the substance is once known.

370. For example: ten thousand parts of sugar is found to consist of carbon 4198, hydrogen 643, and oxygen 5159: now, the equivalents or combining weights of these three elements are respectively 6, 1, and 8. If, then, we divide the numbers just given by these combining weights, we shall obtain 699 equivalents of carbon, 643 of hydrogen, and 644 of oxygen; and on equally reducing these, so as to obtain the lowest number of equivalents, preserving this ratio, we shall at last find that sugar consists of 12 equivalents of carbon, 11 of hydrogen, and 11 of oxygen.

371. This mode of stating the composition of a substance, is very convenient in explaining the various transformations which organic matter undergoes. When a solution of sugar begins to ferment, the first thing is, that the cane sugar passes into the state of
grape sugar; each equivalent of cane sugar takes up and combines with three additional equivalents of water. This change may be thus expressed: 1 equiv. cane sugar contains 12 equiv. carbon, 11 equiv. oxygen, 11 equiv. hydrogen. In passing into grape sugar it takes up 3 equiv. of water, or 3 equiv. oxygen, and 3 equiv. hydrogen, and so becomes 1 equiv. of grape sugar, containing 12 equiv. carbon, 14 equiv. oxygen, 14 equiv. hydrogen.

<table>
<thead>
<tr>
<th>Carbon</th>
<th>Oxygen</th>
<th>Hydrogen</th>
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</thead>
<tbody>
<tr>
<td>1 equiv. Cane Sugar, containing 12 11 11</td>
<td>and 3 equiv. Water &quot; 0 3 3</td>
<td>forms</td>
</tr>
<tr>
<td>1 equiv. Grape Sugar &quot; 12 14 14</td>
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</tbody>
</table>

372. In the second stage of fermentation, this grape sugar is decomposed and separated into alcohol, carbonic acid, and water: 1 equiv. of grape sugar containing 12 equiv. carbon, 14 equiv. oxygen, 14 equiv. hydrogen, during fermentation is resolved as follows:

<table>
<thead>
<tr>
<th>Carbon</th>
<th>Oxygen</th>
<th>Hydrogen</th>
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<tbody>
<tr>
<td>1 equiv. Grape Sugar, containing 12 14 14</td>
<td>forms</td>
<td></td>
</tr>
<tr>
<td>2 equiv. Alcohol &quot; 8 4 12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 equiv. Water &quot; 0 2 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 equiv. Carbonic Acid &quot; 4 8 0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

373. In the same way, the changes which occur during the transformation of starch into grape sugar, may be similarly explained: when this happens, each equivalent of starch takes up four equivalents
of water, and forms one equivalent of sugar: it follows from this fact, that the quantity of sugar formed always weighs considerably more than the starch from which it is obtained.

374. One equivalent of starch contains 12 equiv. carbon, 10 equiv. oxygen, 10 equiv. hydrogen; when acted on by diastase, or boiled with a dilute acid, it takes up in addition the elements of 4 equiv. of water, or 4 equiv. of oxygen, and 4 equiv. hydrogen, and is converted into 1 equiv. of grape sugar, consisting of 12 equiv. carbon, 14 equiv. oxygen, 14 equiv. hydrogen.

<table>
<thead>
<tr>
<th>Starch</th>
<th>Water</th>
<th>Grape Sugar</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 equiv. containing 12</td>
<td>0</td>
<td>12</td>
</tr>
<tr>
<td>and 4 equiv.</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>forms</td>
<td></td>
<td>14</td>
</tr>
</tbody>
</table>

375. A liquid which has already undergone the vinous or alcoholic fermentation, is still able to undergo another change, namely, the acetic fermentation; which occurs where weak spirit of wine, in contact with some ferment, is exposed to the air. Alcohol and water together, whether exposed to the air or not, undergoes no change whatever; when a small quantity of decomposing azotized matter is added, then the acetic fermentation commences, and vinegar is formed (476).

376. In the formation of vinegar from spirit, the presence of air or oxygen is necessary. One equivalent of alcohol consists of 4 equiv. carbon, 2 equiv. carbon, 10 equiv. oxygen, 10 equiv. hydrogen.

14*
oxygen, 6 equiv. hydrogen (372); in passing into vinegar, it takes up 4 equivalents of oxygen, and forms 1 equivalent of acetic acid.

<table>
<thead>
<tr>
<th>Carbon</th>
<th>Oxygen</th>
<th>Hydrogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 equiv. Alcohol, containing</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>and 4 equiv. Oxygen</td>
<td></td>
<td>4</td>
</tr>
<tr>
<td>forms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 equiv. Acetic Acid</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>and 3 equiv. Water</td>
<td>0</td>
<td>3</td>
</tr>
</tbody>
</table>

377. When azotized matters are beginning to decompose, they are at first not able to excite the true alcoholic fermentation in solutions of sugar; it is necessary for this that their decomposition should be tolerably active and advanced. But even in this early stage they are able to effect a very important change in the nature of sugar, and cause it to undergo a peculiar acid fermentation, the result of which is the formation of lactic acid. The relation of this substance to sugar may be learnt from the fact that it contains 6 equiv. carbon, 5 equiv. hydrogen, and 5 equiv. oxygen (445, 570).

378. It sometimes happens also that, in place of lactic acid, another acid substance, called the butyric, is produced. This is especially the case when a solution of sugar is left in contact with caseine, in the first stage of decomposition. Butyric acid is a colorless, sour, pungent-smelling liquid, readily soluble in water and alcohol, and consisting of 8 equiv. carbon, 7 equiv. hydrogen, 3 equiv. oxygen.

379. There are several other forms of fermentation
besides these, though the alcoholic and the acetous are by far the most important. It is evident, however, from the great facility with which starch, gum, sugar, &c., are converted into each other, that all of these forms of organic matter may be readily made to undergo any of these kinds of fermentation, and likewise that the nature of the change thus brought about will depend entirely on circumstances.

380. Organic matter, placed under proper circumstances, may be made to pass through several successive stages of fermentation; the last of these is usually the putrefactive fermentation, the results of which are chiefly carbonic acid, ammonia, and water. When it is desired to excite any particular kind of fermentation, care and attention are always requisite to prevent it from proceeding too rapidly, or too far, as, when this is the case, it very commonly changes into the putrefactive fermentation, and, when this has once commenced, it is hardly possible to stop its progress.

381. A number of important arts are wholly based upon these facts; and many of the arts of life are more or less dependent in principle upon the effects just described. The manufactures of wine, spirit, beer, vinegar, bread, cheese, starch, leather, &c., may be mentioned as examples.

382. Grapes contain sugar, and a small quantity of azotized matter, capable of acting on the sugar as a ferment; but no fermentation can take place, so long as air is altogether excluded. The azotized
matter present will not enter into an active state of decomposition, unless free oxygen has access to it. Consequently whole grapes, or those in which the skin remains perfect and entire, may be dried and converted into raisins; but if the skin is once injured, a little air gets in, and fermentation soon commences. The fermentation of grape juice will not take place if the grapes are pressed in a jar containing carbonic acid gas, or nitrogen.

383. When grapes are pressed, which is the first process of the wine-maker, they are generally bruised, and then allowed to remain some little time in the pressing-vat, before the operation is completed. This allows fermentation to commence, for it begins as soon as the juice comes in contact with the air; and this fermentation, besides softening the skins and cellular tissues of the grapes, also dissolves a portion of the coloring and astringent matter which the skins, seeds, and stalks contain. The pressing is then completed, and the juice separated from the residue, which is termed the murk, or the mark.

384. A certain degree of warmth is always necessary to fermentation. The fermenting liquid must not be lower than 51° or higher than 85°. A temperature of 70° is that most favorable to rapid fermentation. When the liquid is cooled, its fermentation suffers a proportionate check, and if cooled too much, it ceases altogether. When this is the case, it can only be brought on again by the careful application of heat.
385. The must, as the expressed juice of the grape is called, being left undisturbed in the fermenting tun, soon passes into a state of active fermentation; the azotized matter decomposes, causing at the same time the decomposition of the sugar; the liquid becomes turbid and froths up from the escape of carbonic acid. After a time these appearances cease, the liquid clears, and fermentation becomes very feeble.

386. The fermentation of the wine, however, is not yet completed; the half-formed wine is racked off into casks, and allowed to continue slowly fermenting for some time, the casks being kept quite full, so that the froth and ferment which rises to the open bunghole, easily escapes as fast as it rises. When it is judged that this second fermentation has continued long enough, the casks are closed and left for some time.

387. As the quantity of alcohol which is thus produced depends wholly on the proportion of sugar which existed in the must, so, when the grapes are not sweet, and the quantity of sugar is small, the wine formed will be very weak. In such cases, therefore, either the must is concentrated by evaporation, or a portion of sugar is added to augment the proportion of fermentable matter in the must. Much better wine is formed by thus increasing the sugar in the must previous to its being fermented, than by the addition of spirit to the wine when its fermentation is complete.

388. It is generally better to add sugar than to
concentrate the must by boiling, because, in the latter case, a portion of the azotized matter necessary to its subsequent fermentation, is rendered insoluble and useless. Care must, however, be taken not to add too much sugar, as, when this is done, fermentation proceeds very slowly; the proportion of sugar should in no case be greater than one-eleventh of the weight of the must.

389. During the two stages of fermentation through which the must passes, the greater part of the azotized matter undergoes decomposition, becomes insoluble, and falls down, constituting, together with the other insoluble matters, the lees of the wine. The sugar disappears, carbonic acid gas is given off, and alcohol formed. A small quantity of certain peculiar volatile oils is also formed, to the formation of which the wine owes its flavor, or bouquet. A considerable quantity of tartar or bitartrate of potassa likewise is thrown down during the fermentation of the wine. This salt, which is soluble in water, but almost insoluble in a mixture of water with spirit of wine, exists naturally in the juice of the grape; as the fermentation proceeds, and alcohol is formed, the tartar being consequently no longer soluble in the liquid, is deposited in the form of crystals on the side and bottom of the cask (498).

390. The fermentation and ripening of wine continues for a long time in the wood; a small quantity of undecomposed azotized matter remains in the wine; this acts as a very slow ferment, and hence the wine
continues to improve, more spirit being formed, and consequently more tartar thrown down. Wine also becomes stronger when kept in the wood, because, under these circumstances, a greater proportion of water passes off by evaporation, than of alcohol. This is always found to be the case when a mixture of spirit and water is kept in a slightly porous vessel, the water evaporates faster than the spirit.

391. Finally, the wine is either racked off into a clean cask, the lees being left behind, or it is clarified, or fined, with albumen or isinglass, and then bottled. Wines which are intended to be sparkling or effervescing are bottled before the fermentation is quite finished, so that the carbonic acid subsequently evolved, remains stored up in the liquid.

392. In making what are termed home or domestic wines, the juice of various fruits, &c., is mixed with sugar and fermented. A good deal of skill is required in this operation, and care must be taken so to proportion the different ingredients, as to insure a proper fermentation. It is a very common error to use too much sugar, and not add enough ferment to cause its due fermentation; for this purpose the yeast or ferment of wine is best; that obtained from the fermentation of beer injures the flavor of the wine.

393. When the vinous fermentation of must is quite complete, it should not be suffered to remain longer in contact with the lees, because when this is the case, especially with weak wines, many circumstances may tend to commence the acetous fermentation, giving
the wine an acid flavor. When this has once commenced, and the wine is what is technically termed pricked, it is very difficult to render the wine wholesome again; wines are subject to many diseases, as they are called, and some of them may be prevented or remedied. The chief cause of them is, the presence of a minute quantity of ferment, of some kind or other. When wine is not perfectly clear, it is evident that it contains "ferment," and is, consequently, peculiarly liable to ferment and turn sour.

394. There are many other fermented liquids which are prepared in the same way as wine, and, like it, contain alcohol, derived from the fermentation of sugar. Thus cider obtained from apples, perry from pears, and mead from honey, derive the alcohol which they contain from the sugar of fruit or honey, and owe their flavor chiefly to the volatile oils formed during its fermentation.

395. In the preparation of these liquids, especially cider, it is often necessary to put a stop to the fermentation when it is arrived at a certain point; this is usually effected by the fumes of burning sulphur; a small quantity of sulphuric acid gas arrests the fermentation of a large quantity of fermenting liquid. Fermentation is arrested or altogether prevented in a very remarkable manner by the presence of various substances, such as some of the strong acids, certain salts, and some of the volatile oils, even in very minute quantity. No substance possesses this power to a greater degree than sulphuric acid gas (173).
396. When wine is distilled, the liquid which passes over is brandy, and not a pure spirit of wine; a small portion of the volatile flavoring matter of the wine passes over with the alcohol and water. By repeated rectifications this flavor may be removed, and a pure spirit of wine, or alcohol, obtained.

397. Good wine brandy, such as Cognac, when obtained by distilling wine, is always colorless like water; but old brandy, or that which is kept some time in the wood, acquires more or less of a brownish color, in consequence of its dissolving a portion of the coloring matter of the cask. The manufacturers of brandy, who prepare a spirit sold under the name of brandy, but really derived from some other source than the juice of the grape, color their spirit with oak-wood, burnt sugar, treacle, or even more objectionable substances.

398. Spirit is obtained in a great number of ways, and varies in flavor according to the mode in which it is prepared. Sometimes, as in the manufacture of liqueurs, which are, in fact, flavored brandies, some substance is purposely added to alter the flavor. Maraschino and Kirschwasser are brandies distilled from the fermented juice of crushed cherries, and derive their agreeable flavor from the volatile oil and prussic acid which exist in the kernels of the cherry; but maraschino is likewise frequently flavored by the addition of cloves, orange, or cinnamon. Noyeau is flavored with the kernels of bitter almonds.
399. The more common kinds of spirit, though all, in fact, obtained from the fermentation of sugar, are yet in the first instance obtained from starch; this substance, however, in all cases, is first converted into grape sugar, and then undergoes the vinous fermentation.

400. Distillers either employ malted grain, or a mixture of malt with raw grain. In the first case, a very large portion of the starch which the grain contains is already converted into sugar; and when raw grain is used, its starch is gradually changed into sugar in the process of mashing (413).

401. The malt, or mixture of malt and grain, is ground, and then stirred or mashed into warm water, in the mash-tub; in this operation, the soluble matter of the grain is extracted, and the starch converted into sugar. The wort, as this infusion of grain is called, is then cooled, drawn off, and transferred to the fermenting tuns; a quantity of good yeast or ferment is added, and the mixture is left to ferment for several days.

402. The fermentation of the sweet wort is allowed to go on as long as it can be safely left, without fear of the acetous fermentation being brought on; for it is the object of the distiller to convert the whole of the saccharine matter which it contains into spirit. When this is done, the fermented wort, which is then termed wash, is transferred to the still, and heated. The product is a dilute, weak spirit, and requires to be rectified or distilled again.
403. During the fermentation of the wort, as in all similar operations, a quantity of a peculiar volatile oil is produced, some of which passes over with the spirit when it is distilled; that which is obtained towards the end of the distillation is generally most contaminated with this oil, and is termed faints. The wash usually yields rather more than one-tenth of its volume of pure alcohol, or one-fifth of strong spirit, half of which is pure alcohol;—for the strongest spirit which can be thus obtained, always contains a considerable quantity of water (368).

404. As there is an excise duty on the manufacture of spirit, it is necessary to estimate the strength and quantity of spirit which the distiller obtains. In doing this the spirit is stated to be over or under proof. Proof spirit consists of equal parts of water and alcohol. It was formerly the custom to pour a small quantity of the spirit to be tested, over gunpowder, and then set fire to it; if the spirit was strong, the gunpowder fired, and then the spirit was called proof spirit; but if it contained too much water, the gunpowder was not fired.

405. This method of testing the strength of spirit is highly inaccurate. As alcohol is much lighter than water, the strength of spirit is easily ascertained by weighing a portion of it in a bottle, the capacity of which is known; this gives its specific gravity, or weight; compared with an equal bulk of water, and from that it is easy to calculate its strength.

406. The flavor of the grain spirit, or whiskey, thus obtained, depends a good deal on the ferment
used in its manufacture. The whiskey obtained by means of beer or porter yeast has always an inferior flavor to that prepared in the Highlands and in Ireland, where no brewer's yeast can be obtained. Whiskey is sometimes artificially flavored; thus, in the old Usquebaugh, nutmegs, cloves, cinnamon, and other similar substances were added; and Geneva, or gin, was originally prepared by bruising juniper berries in the mash-tun, along with the grains and malt.

407. In the manufacture of common gin, a small quantity of oil of turpentine is frequently substituted for the juniper berries; as the flavor thus communicated to the spirit greatly resembles that derived from juniper. The best Dutch gin, Hollands or Schiedam, is chiefly obtained from rye, mixed with about one-third its weight of malted barley, and fermented; the weak spirit first obtained by the distillation of this liquid is then rectified once or twice off juniper berries, by which means it loses the crude flavor which it originally had, and acquires the agreeable one of the junipers.

408. Other spirits are similarly prepared from various substances—in fact, almost all vegetable substances, containing either starch or sugar, may be made to yield spirit by fermentation. Very large quantities of spirit are manufactured from potatoes, which are grated, and the pulp mashed in water with a small quantity of ground malt, in an active state of fermentation. Great care is required in the manage-
ment of potato wash, as the wort is very liable to pass into the acetous fermentation. The spirit obtained possesses a peculiar, and by no means pleasant flavor, which, however, may be removed by rectification.

409. Rum is obtained chiefly in the West Indies, from the fermentation of molasses, or uncrystallizable sugar, mixed with a portion of sugar-cane juice, and a few crushed fragments of the cane itself; if these are not added, the spirit has very little flavor. The fermentation of the liquor is assisted by the addition of skimmings from the sugar-boilers, and some of the refuse from a previous distillation, called dunder, and which is a tolerably active ferment. When first distilled, rum has often a harsh and disagreeable flavor; it loses this after being kept some time in casks.

410. In India, a great variety of spirits, which are described under the general name of arrack, are prepared. The chief sources are rice, sugar-caneces, and the sweet juice of various palm and other trees, abounding in sugar. This juice, which is called toddy, ferments very easily; and if care be taken that it does not pass into the acetous fermentation, yields a pure and well-flavored spirit.

411. In the manufacture of beer, the object of the brewer is, to obtain a solution containing the nutritive matters of the grain in a soluble and partly fermented state; part of the sugar being already converted into spirit, whilst the fermentation of the
rest has been prevented, and its tendency to ferment altogether destroyed.

412. The first operation is to convert the insoluble starch in the grain into soluble gum and sugar; this is effected by malting. The grain is moistened and allowed to germinate, and, in so doing, it becomes sweet, starch disappears, and sugar is formed; if germination were allowed to go on, the sugar thus formed would be absorbed by the young embryo plant, and the labor of the maltster consequently lost; as soon, therefore, as the required chemical change is effected, he dries the malt, destroys the vitality of the seed, and prevents further loss (690).

413. Barley generally contains about 8 or 9 per cent. of sugar and gum; after being converted into malt it contains 30 per cent. of those substances, and of this at least half is sugar. In the process of malting, some of the azotized matter originally contained in the seed is lost (685).

414. Although barley is generally used in the manufacture of malt, yet most other grains may be rendered sweet by a similar process. In various countries, rye, wheat, oats, maize, rice, and millet, are malted.

415. In the manufacture of beer, as in the preparation of distillers' wash, the malt is first ground, or rather crushed, and then mashed or infused in warm water, at a temperature of about 160°; so as to extract all the soluble matters which it contains. The malt is kept constantly stirred in the mash-tun for
about three quarters of an hour, the tun is then covered, and the whole is left quiet for about the same time, or rather longer.

416. Two objects are attained in the process of mashing; the soluble matters in the malt are extracted, and a further portion of the starch is converted into sugar and gum, by the influence of the diastase which the malt contains (685); and hence, if a quantity of raw grain is mixed with the malt, the starch which it contains will be found to be converted into sugar and gum in the mash-tun.

417. It is of the first importance to use water of a proper temperature in the mash, tun; a good deal of heat is given out in the process of mashing, and this must be allowed for. If the water is too hot it converts the starch into a jelly, which interferes with its conversion into sugar; this is called setting the mash. If on the other hand the mash is too cool, the conversion of the starch into sugar will likewise be very slow and imperfect, and the wort will have a great tendency to turn sour.

418. When the mash is complete, and the wort has become clear, it is run off from the spent grains; which are then mashed a second time with fresh warm water, to dissolve the soluble matters still left in the grains after the first mashing. The product of the first mash alone is used for ale, while that of the subsequent operations is employed for common or small beer. Sometimes the grains are subjected even to a third mashing.
419. The sweet wort is then run into the copper, and boiled, and the sooner this is done after it has been drawn off, the better. Sweet wort contains a much larger quantity of vegetable albumen than is necessary to insure the requisite degree of fermentation, and this, if left in it, would be almost sure to cause it to pass into the acetous fermentation. By boiling the wort, a considerable quantity of this albumen is rendered insoluble, and precipitates in flakes. The wort must, however, not be boiled too long, or the whole of the albumen will be thrown down.

420. When the wort has been boiled a certain time, a quantity of hops is added; the hops are not allowed to boil with the wort the whole time, or too much of their rank flavor would be thus extracted, and the long-continued action of heat would drive off some of the fragrant volatile oil which they contain, and the presence of which is essential to the flavor and goodness of the beer.

421. This part of the operation, as indeed are all which in any way depend on the regulation and management of fermentation, is very delicate, and requires much care; the boiled wort is very prone to pass into the acetous fermentation; it must be rapidly cooled, and brought down to the temperature best suited for alcoholic fermentation.

422. The wort must be cooled to about 60° or thereabouts, and if this is not quickly done the wort begins to decompose; it acquires a rank and disagreeable flavor, and becomes slightly mouldy; this, which
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is termed foxiness, can never be completely remedied if it has once commenced. It is peculiarly common in moist, close, warm weather.

423. The cooled wort is then run into the fermenting vat or gyle-tun; if left to itself, fermentation would soon commence; but it is far better to add at once some good yeast from the last brewing, stir the whole up well together, and then, having covered the vat, leave it quiet; in a few hours the whole will be in an active state of fermentation.

424. In three or four hours, bubbles of gas will be seen rising from all parts of the liquid, a ring of froth forming at first round its edge, and gradually increasing and spreading till it meets in the centre, and the whole surface becomes covered with a white creamy foam. The bubbles of gas then rise and break in such numbers, that they emit a low hissing sound, and the white foam of yeast continues to increase in thickness, breaking into little pointed heaps, which become brownish on the surface and edges.

425. The yeast gradually thickens, and at last forms a tough, viscid crust, which, when the fermentation begins to slacken, would break and fall back again into the vat, if it was not removed. The brewer skims it off and removes it as soon as he judges that the fermentation is complete; but if he thinks that the fermentation has ceased too soon, he stirs the whole up again so as to mix the yeast with the wort. This, which is called rousing the vat, pro-
longs the fermentation, or rather causes it to recommence.

426. Whilst the fermentation of the wort is proceeding, it gradually becomes clearer; the ferment added, as well as that which is generated in the wort (for all the azotized matter present is soon brought into the same condition of active fermentation), is gradually separated; meanwhile the sugar is decomposed, spirit is formed, carbonic acid given off, and a considerable amount of heat is generated. Where the quantity of wort is large, this rise in temperature amounts to ten or fifteen degrees.

427. In making ale and beer, it is the object of the brewer to leave a portion of the sugar in the liquid, he therefore removes the yeast so as to put a stop to further fermentation; but in brewing porter he does not remove the ferment so soon, but allows the fermentation of the wort to continue for a longer time. It is sometimes found convenient to add a small quantity of sugar to the wort.

428. After the first fermentation is finished and the yeast removed, the liquor is racked or drawn off into casks, when it undergoes a second fermentation far more slow and protracted than the first. This is termed cleansing, and, whilst it goes on, a considerable quantity of mucilage, spent ferment, and other impurities subsides, constituting the lees or dregs. When this second fermentation is complete, the liquor is drawn off into casks, the bungholes of which are then well closed.
429. The proportion of alcohol in different kinds of ale, &c., varies greatly; small beer usually contains from one to two per cent.; ale from four to nine; and porter from four to six per cent. of alcohol. It is commonly calculated that a bushel of good malt should make about twelve gallons of ale, or twenty-four gallons of beer; but if the ale is intended to be kept, a greater proportion of malt must be taken.

430. The quantity of hops taken varies according to the strength of the liquor, and many other circumstances. Strong ale, which is intended for keeping, must have more hops than that which is intended to be used at once. From three-quarters of a pound to a pound and a quarter is the quantity used to each bushel of malt. Occasionally, other bitters are used together with hops; there are several which may be very well used for this purpose, such as the bitter of the chamomile flower; but there are others occasionally employed which are highly objectionable, like quassia. The bitter principle of this wood does not answer the same objects as that of the hop, and is besides very unwholesome.

431. Hops are used in brewing, for several purposes. The aromatic bitter which they contain greatly improves the flavor of the drink; their volatile oil prevents further fermentation, and renders it less likely to turn sour; and the tannin and astringent matter of the hops help in precipitating the mucilaginous and albuminous substances which the wort
contains, and which, if left in the beer, would be very apt to cause its fermentation.

432. The goodness of ale and porter depends a great deal on the water with which it is brewed; it is sometimes supposed that the water must be soft, but this does not appear to be really the case, because the water employed in many of the largest and best breweries is decidedly hard, and contains a very considerable quantity of sulphate of lime (239).

433. Porter is essentially ale brewed with a portion of scorched or high-dried malt, and the fermentation of which has been allowed to proceed so far that nearly the whole of its sugar has disappeared. It was formerly brewed entirely from high-dried malt, but as in that case a large quantity of the sugar was needlessly destroyed, it was found better to use common malt, mixed with a small proportion of high-dried malt; this gives the desired color and peculiar flavor of porter or brown stout. Sometimes other coloring matters are used for the same purposes; these for the most part must be regarded as adulterations.

434. The slow, continued fermentation which malt liquors undergo whilst kept ripening in the casks, is essential to their preservation; because the carbonic acid thus generated protects them from the influence of the air; hence, too, the casks must be kept closely bunged up to prevent the escape of this gas. As soon as this fermentation ceases, and air gets access
to the liquor, oxidation commences, and it begins to pass into the acetous fermentation(476).

435. For the same reason, also, when a cask has once been tapped, it should be used at once; the liquor is brisk at first, being charged with carbonic acid, which renders it pleasant to the taste, and preserves it from further change; ere long the carbonic acid escapes, the liquor becomes flat, then the oxygen of the air begins to act upon it, and it soon turns sour.

436. Malt liquor, which has once become clear and fit for use, must never be shaken; as ferment is always formed and deposited, even during the very slow fermentation which takes place in the cask: agitation not only renders the liquor thick and turbid by diffusing the spent ferment throughout it; but is very likely to induce a fresh fermentation, which would probably pass into the acetous one.

437. In the Bavarian mode of brewing, the fermentation of the wort is allowed to proceed very slowly, exposed to a low temperature in large open vessels. Though slow, this fermentation is very complete, the whole of the ferment is deposited at the bottom of the vat, whilst very little rises to its surface, and there is hardly any frothing of the wort. Beer thus made has very little tendency to enter into a new state of fermentation, even by the action of the air.

438. Bread-making depends in great part on fermentation; a small quantity of sugar which exists in
dough is converted into carbonic acid and spirit, by the action of the yeast or ferment mixed with it; the carbonic acid thus evolved by dividing the dough, and forming little holes and cavities throughout its mass, renders the bread light and porous.

439. There are two distinct objects to be attained in the manufacture of bread, the one is the chemical union and transformation of the starch and other components of the grain, the other the formation of a uniformly light, spongy, and easily digestible mass; the latter is merely a mechanical effect, but both objects are effected by chemical means.

440. The flour of wheat and other grains used in bread-making, consists chiefly of starch, together with a much smaller proportion of sugar, gum, and azotized matters. These substances may be made to undergo various changes, and these changes may be brought about in many different ways; the most common and convenient, is to excite in them a state of vinous or alcohalical fermentation.

441. There are two distinct modes by which this is effected, namely, the use of leaven and of yeast or barm. Leaven is dough which has been kept till it has turned sour; when this is mixed with a quantity of fresh dough, it has the power of communicating to the whole of the latter its own state of fermentation.

442. A quantity of flour is kneaded up into dough with a suitable quantity of water, and then left in a warm place; it will soon become sour, and a greater or less degree of fermentation will be established in
it, but it will be irregular and very uncertain. If, however, a small piece of this sour dough is taken and thoroughly kneaded up with a large lump of fresh dough, the whole of the latter will very soon enter into a uniform state of fermentation; and if a small piece of this is set aside for the next baking, it will be in a fit state for use by the time that it is required.

443. The leaven may be formed either of the same kind of flour as the bread is intended to be made with, or any other sort; it is most commonly the custom to employ bean, pea, or some similar flour, which passes very easily into fermentation, or at least to mix a portion of it with the flour used in forming the leaven. When well made, leaven may be kept fit for use for weeks and months, and by adding a portion of dough to the leaven as large as that removed for the use of the bread-maker, the stock of leaven is always kept up.

444. In this country, however, the use of leaven is almost wholly superseded in favor of yeast. This substance can usually be obtained from a neighboring brewery, and a small quantity of it mixed up with the dough, brings it rapidly into the required state of fermentation, without communicating to it the sour taste which leaven always gives to bread. In many parts of Europe, however, and when yeast cannot be had, all the bread is made with leaven.

445. Leaven added to dough excites in it a true alcoholic fermentation, but it also produces a portion
of lactic acid (377), and frequently vinegar likewise; the latter is for the most part driven off in the subsequent process of baking, but the former remains in the bread.

446. The baker usually commences by mixing together in a suitable vessel a proper proportion of yeast, warm water, common salt, and flour. The salt is first of all dissolved in warm water, which is then allowed to cool down to about 80° or 90°, after which the yeast and then the flour is added. The quantities taken vary slightly. The third part of a sack of flour, a pailful of warm water, four or five pounds of salt, and three or four pints of yeast, are about the usual proportions.

447. This mixture, which is called the sponge, is worked up to the consistence of stiff batter, and then left in a small trough for an hour or two, covered over with a cloth. During this time a tolerably active fermentation is commenced, and as the mixture is tenacious and viscid, the carbonic acid generated does not escape in bubbles, but is retained in the dough, causing it to swell up to about twice its original size.

448. If the sponge is made too thin, the gas thus generated will escape as fast as it is evolved; if of the right consistence, the whole of it will be retained. If the fermentation is allowed to continue too long, the sponge will become sour, and pass into the state of leaven; the baker, however, does not permit this, but when he judges that the fermentation has pro-
ceeded far enough, he adds about twice as much more flour as he originally took.

449. The sponge and flour are then very fully and carefully kneaded together, so as to insure the complete mixture of the half-fermented sponge with the fresh flour; this is a very laborious part of the operation, but it is quite essential to the success of the process, for if it is not very thoroughly attended to, the ferment will not be equally and uniformly diffused throughout the whole of the dough.

450. The dough is then left for about an hour and a half; it is then kneaded a second time, divided into pieces of proper size and weight to form loaves, and then set aside in a warm place for a short time. Lastly, the loaves are put into the oven and baked, during which operation they are still more expanded and enlarged. In baking, the loaves generally rise or swell up to about twice the size which they previously had.

451. The object of all these different processes is to decompose the sugar which the flour contains, for the purpose of generating carbonic acid, and thus rendering the dough light and porous. The sugar, therefore, which the flour contains is essential to bread-making; but though the whole, or the greater part, of this sugar is of necessity thus decomposed, by the action of the yeast, yet when bread is analyzed it is found to contain very nearly as much sugar as flour does.

452. When the yeast is mixed with the flour, it
immediately begins to act on the sugar, causing its decomposition into alcohol and carbonic acid; but at the same time it also acts upon the starch, and produces in it a change analogous to that which diastase effects in the germination of a seed. It consequently happens, that though the sugar which originally existed in the flour is almost entirely decomposed, yet a quantity of sugar, nearly equal in amount, is produced or generated from the starch (360).

453. In baking the bread a further portion of the starch is decomposed; more gum and sugar are formed, the azotized matters uniting with them and the modified starch, make a uniform substance, which is far more digestible than flour or unbaked bread. At the same time that these chemical changes are effected, the fermentation is altogether stopped, a considerable quantity of water and the alcohol formed during the rising of the bread are driven off, and the bread is at the same time expanded and rendered still lighter, by the action of the heat of the oven on the gas contained in its cavities (62).

454. The necessity of attending to the proportions in which the several ingredients of bread are mixed, as well as the need of mixing them very thoroughly, is evident. If the dough is badly mixed, the carbonic acid, in place of being regularly diffused throughout the substance of the dough, will be collected into large irregular bubbles, and then the bread will be full of large holes and cavities, in place of being uniformly light and spongy; and the loaf,
though perhaps as large as it would have been if well made, will be close and heavy.

455. The goodness of bread depends a good deal on the proportion of water which it contains. The usual quantity in well-made bread is about 44 per cent., but it varies somewhat according to the fineness and goodness of the flour. The percentage of water is also modified in a remarkable manner by the presence of various saline substances. The use of common salt improves bread in several ways; it not merely improves its mechanical texture, and renders it more wholesome and digestible, but it also increases its retentive power from moisture.

456. When inferior or damaged flour is used, it is generally found that the bread rises slowly and imperfectly, and when baked is close, and has a bad color. Saline matters are found to correct this to a considerable extent, and some salts far more powerfully than others; alum is sometimes employed for this purpose. No object is gained by adding it to good flour, but in the case of inferior flour its use decidedly improves the bread; and though it is unquestionably an injurious adulteration, yet it is harmless compared with some of the other salts which have been employed for the same purpose.

457. A very small quantity of alum greatly improves the color of the bread, and also causes it to rise better, thus making inferior bread equal to the best. As far as the mere alteration of color is concerned, its use only serves to please the eyes of the
customer, or rather to deceive him into the belief that he is buying better bread than is really the case. The increased lightness which it causes is, however, really an improvement.

458. Some bakers have even added sulphate of copper to the dough. This salt produces, in very minute quantity, the same effects on bread as alum does. Being a highly poisonous salt, its use is most objectionable; fortunately, it is an adulteration very easily detected.

459. Carbonate of ammonia, which is sometimes mixed with slightly damaged flour, is a harmless addition. It does not improve the color of the bread, but renders it lighter, and removes any sour flavor which the bread might otherwise acquire, either from the inferior quality of the flour, or the nature of the ferment employed (462).

460. When bread is made with leaven, it always has a distinctly sour taste; this is partly caused by the leaven itself, and partly by an acid fermentation which it induces in the flour or dough. It sometimes happens also that bread which is made with barm or yeast has a similar sour taste; this is usually caused by want of skill or care in the preparation. It is occasionally caused by the use of sour yeast; when the latter has been kept too long or is spoilt, and then of course the bread will have a sour taste, the same as if it were made with leaven.

461. In places where a fresh supply of new yeast cannot always be had, sour yeast must sometimes be
used; but there is no need for the bread to be bad in consequence; because it is quite possible to remove the sour taste of the yeast before using it. The addition of a small quantity of carbonate of lime, soda, or magnesia, will effect this; but, of course, care must be taken not to use more of either of these salts than is necessary to neutralize the acid matter present.

462. The sour taste which bread acquires when it is badly made, or when damaged flour is used, is less easily corrected. Nevertheless, it may be greatly diminished, if not altogether prevented, by the addition of a small quantity of carbonate of soda or ammonia. The latter salt may often be employed with great advantage, and, being a volatile salt, it is raised in vapor by the heat of the oven. Any excess of it, therefore, is driven off by the heat, and in escaping, assists in making the bread lighter.

463. Yeast is not unfrequently bitter, and then communicates a very unpleasant flavor to the bread made with it. This may be derived from hops, especially when the yeast is collected from beer which is strongly hopped. But yeast sometimes also acquires a bitter taste from keeping, which is quite independent of that derived from the hops; this also may be remedied to some extent in various ways. One way, which is said to be very effectual, consists in throwing into the yeast a few clean cinders freshly taken from the fire, but allowed to cool a little on the surface. When cold, the cinders fall to the bottom, and the yeast is poured off. This operation appears to de-
pend in principle upon the singular property which freshly-burnt charcoal has of absorbing gases, and removing coloring and odorous matter generally from substances to which it is added (162).

464. When, however, there is any great difficulty in procuring fresh and good yeast, it is generally better to use artificial yeast, or to prepare a ferment on purpose; this is by no means difficult, and then the inconveniences just mentioned are wholly avoided. There are a great number of different ways of making artificial yeast. The object usually is to obtain a quantity of paste or dough in an active state of fermentation, and then, by removing the water, to check further change until the ferment is wanted. By carefully washing and pressing beer yeast, so as to separate the moisture as much as possible, it becomes far less liable to spoil than it is in its fresh state.

465. A tolerably good kind of dry yeast may be made by mixing together a decoction of wheat and bean flour with a small quantity of brewer's yeast; when the whole is in a state of active fermentation, enough wheat or barley flour is added to form a thick dough, which is then thoroughly kneaded, formed into small cakes, and carefully dried. A very good ferment may at any time be obtained from one of these cakes, by crumbling it into warm water, and leaving the mixture a few hours quiet in a warm place.

466. Yeast may also at any time be prepared direct from malt. When a thick paste or dough of wheat flour is left in a warm place, it soon begins to
ferment and emits a sour smell; at the end of a week, however, it loses this sour smell and acquires a vinous one, and is then able to act powerfully on sugar as a ferment.

467. If a small quantity of dough in this state is diffused in some warm water, and added to a strong decoction of malt and hops, such as brewers make, or even of malt alone, it soon brings the whole into active fermentation, and in a few hours a quantity of good fresh yeast will be deposited. When the alcoholic fermentation is complete, the clear liquor may be poured off, and the yeast is fit for bread-baking.

468. The object to be attained by the use of ferment or leaven in the manufacture of bread being a purely mechanical one, and as the process of fermentation is always uncertain and troublesome, a great number of attempts at different times have been made, to make bread without its undergoing any fermentation whatever, though hitherto with but partial success. With care and attention, however, excellent unfermented bread may be made.

469. It is sometimes stated by those who recommend the use of unfermented bread, that, in the ordinary mode of bread-making, a large portion of the most valuable part of the flour is destroyed by fermentation; this is not really the case. Very little, indeed, of the azotized matter of the flour is lost during the fermentation of the dough; the chief effect produced is the loss of a portion of sugar, but, as a nearly equal quantity of sugar is at the same time
formed from the starch (462), the real effect of the fermentation may be said to be principally the loss of about five per cent. of starch. The weight of the baked bread is always much greater than that of the flour from which it is made, owing to the large quantity of water which is incorporated with it. Two parts of flour make about three parts of bread.

470. The best of the various chemical modes of making bread, is that in which dry carbonate of soda and muriatic acid are employed. A small, but definite quantity of dry carbonate of soda is thoroughly mixed with the flour; enough pure muriatic acid to neutralize it perfectly is dissolved in the proper quantity of water, and the flour then added; whilst the water and flour are being mixed, the acid acts on the carbonate of soda, decomposes it, expels its carbonic acid, and forms chloride of sodium, or common salt, with its base.

471. Hence, in this process the same effect in the end is produced as in the ordinary mode of bread-making, but with this difference, that it has not undergone any sort of fermentation, and that nothing has been lost. A light and spongy dough is produced, containing abundance of bubbles of carbonic acid; and the only residue of the process, the common salt formed, so far from being objectionable, is, in fact, necessary to the formation of good bread.

472. Another point in favor of this process is, that it takes much less time than the old methods do, because there is no need to leave the dough to rise by
fermentation; in fact, the less time the dough is left to itself, and the more rapidly the operation is carried on, provided the dough is thoroughly mixed, the better will the bread be.

473. The most serious objection to this mode of bread-making is, that, unless great care be taken, there is considerable risk of introducing poisonous substances into the bread if the acid is not perfectly pure, even when all reasonable precautions are taken to avoid adulterations. It is, of course, necessary to take care that the carbonate of soda is most thoroughly mixed with the flour, and great attention must be paid to the proportions of the several ingredients.

474. In toasting bread, two very different effects are produced—the modified starch in combination with gluten, which the bread contains, is rendered still more soluble, and a fresh portion of gum is formed, whilst, at the same time, a part of the bread is more or less charred or carbonized.

475. In making toast and water, it is sometimes recommended to burn the toast well, so as to make it quite black; this is often very good advice, for the porous charcoal thus formed is a powerful purifying agent, and when water has a bad flavor, the addition of a piece of blackened toast does not merely cover the flavor, but removes it, and renders the water more wholesome. The soluble matters, and also the finely powdered charcoal held in suspension, both contribute to render toast a very useful addition to water.
476. Acetic acid, or vinegar, is formed when a mixture of spirit and water, together with some ferment, is exposed to the air; a portion of oxygen is absorbed, and acetic acid is formed. This change takes place when any liquid containing a portion of spirit and some organic substance, like gluten, is exposed to the air and kept moderately warm; the acetous fermentation, as it is called, then commences, and vinegar is formed.

477. When a liquid containing spirit, water, and some decomposable organic matter is exposed to the air, unless the temperature is too low, it soon begins to turn sour, and forms vinegar. When a solution which has fully undergone the alcoholic fermentation is thus kept at a temperature of 70° to 85°, it slowly and gradually undergoes this change; but unless care is taken it is very apt to become mouldy and putrefy (380).

478. The acetous fermentation is facilitated by the addition of a little ferment; hence in the manufacture of vinegar from wine or beer, yeast or ferment of some sort is usually added. When a small quantity of wine which has turned sour during its fermentation, is added to a much larger quantity of properly fermented wine, it induces the whole to pass into the acetous fermentation, and good vinegar is formed. For this reason also, better vinegar can be made in an old cask or vat which has been often used, than in a new one.

479. Wine-vinegar, however, is commonly made in
vats provided with false bottoms pierced full of holes, a layer of vine-stalks and mark is first placed in the vat, covering the false bottom, and the vats are then nearly filled with wine. The vine-stalks and mark act as a ferment, and soon bring the wine into a state of active fermentation: air is absorbed, heat given out, and vinegar is formed (383).

480. Care is always taken to check the rapidity of this fermentation, or to prevent the liquor from becoming too warm; if this were permitted, the spirit of the wine, as well as the vinegar formed, being both volatile, would be driven off, or at least a considerable portion of them would be lost.

481. Any liquid containing spirit may be fermented into vinegar; and, consequently, as starch and sugar may be converted into spirit, so, solutions containing those substances may readily be turned into vinegar, by being fermented: when this is the case, the process is not stopped when the alcoholic fermentation is complete, but is allowed to proceed as long as possible.

482. In countries where wine is made, vinegar is chiefly made from the grape; in other places malt is usually employed. A similar form of vat is used for that purpose as in the manufacture of wine-vinegar, the only difference being, that a sweet-wort is substituted for the wine. Grape-mark, which is commonly called rape by the vinegar-makers, is generally used as a ferment.

483. As soon as the wort has acquired a certain
degree of sourness, and, in fact, before the acetous fermentation is quite complete, it is racked off; this is necessary to prevent it from passing into a state of putrefaction, which would be sure to happen if it were left too long in contact with the ferment (380).

484. Vinegar, whether prepared from wine or malt, has a great tendency to spoil by keeping, in fact to putrefy; this is merely caused by the impurities which it contains. As made from malt, it always contains spirit, sugar, gum, and ferment, in greater or less quantity. It is thick and turbid, and must be clarified before it is fit for use or can be kept; this is done by the use of isinglass or some similar fining. Sometimes also it is purified by filtration through powdered charcoal.

485. When vinegar putrefies, the change is not confined to the impurities alone which it contains, the acetic acid also is decomposed, and hence, as soon as the putrefactive fermentation commences, the liquid begins to lose sourness and becomes flat and insipid. The impurities of the vinegar may of course be wholly separated by distillation. Distilled vinegar has no tendency to spoil by keeping; it is quite as strong and sour, or even stronger than it was before, but its flavor is not quite so agreeable.

486. Vinegar is sometimes conveniently strengthened by the addition of sugar. In this case the impurities which it contains, carry on the slow fermentation of the sugar, and cause its conversion into vinegar. The sugar must be added from time to
time, and the liquid should be kept at a temperature of from 70° to nearly 90°.

487. Very pure vinegar is sometimes made from a mixture of spirit of wine, and water, exposed to the air at a proper temperature. A large cask is filled with wood shavings which have been steeped in vinegar, and the mixture of alcohol and water is allowed to trickle through it at the same time that a current of air passes through the cask. The large surface which is thus exposed to the air, and the influence of the vinegar contained in the shavings greatly assist the action; oxygen is absorbed, the mixture becomes warm, and after passing once or twice through the cask is wholly converted into vinegar.

488. Another source of acetic acid is the destructive distillation of wood. When wood is burnt, only water and carbonic acid are formed; but when wood is distilled or roasted in close vessels out of the contact of air, it is decomposed, and several new substances are formed by the recombination of its elements; the most important of these substances is vinegar, or, as it is called when thus procured, pyroligneous acid (325).

489. Acetic acid cannot be obtained perfectly dry. The substance obtained by the above-mentioned processes, is a solution of acetic acid in water; this may be concentrated, but if we attempt to obtain the acid free from any water, it is decomposed. Acetic acid forms numerous salts, some of which are important in the arts, such as sugar of lead, which is an acetate
of lead, and pyrolignite of iron, which is an impure acetate of iron; the salts made with the crude wood-vinegar, or pyroligneous acid, are commonly called pyrolignites.

490. The strongest acetic acid is usually obtained by neutralizing vinegar with carbonate of soda or lime. By evaporating the liquid, a dry acetate of soda or of lime is procured, and these salts, when acted on by strong sulphuric acid, give off acetic acid in the form of a very pungent acid vapor, which may be easily condensed by cold. A small portion of water distils over with the acetic acid, and is condensed with it, forming a very strong solution.
CHAPTER VI.

VEGETABLE PRINCIPLES—NATURE AND COMPOSITION OF ANIMAL SUBSTANCES.

491. It is well known that plants possess very various properties: some are noted for their fragrant smell, others for the brilliant colors they produce; some yield oils, others resins, and a great many are valued for their peculiar medicinal qualities: all these various properties are caused by the presence of a certain quantity of some organic substance, some peculiar compound of oxygen, hydrogen, carbon, and nitrogen, which is formed by the plant.

492. The number of these organic substances which have been discovered in plants is very great: it is quite unnecessary to describe or even enumerate them all; it will be sufficient to mention very briefly the most important of them, arranged under three or four heads.

493. Amongst the substances formed by plants, is a numerous series of organic acids: substances composed of oxygen, hydrogen, and carbon, which possess acid powers, and combine eagerly with bases to form neutral salts. In general, they do not exist in
plants in the free state, but are combined with various bases derived from the soil; sometimes they are found in the state of perfectly neutral salts, but more frequently they form acid salts (111); that is to say, there is more acid than the base is able to neutralize; such plants have a sour taste.

494. When the salts of organic acids are burned or strongly heated, the acids are decomposed; in these cases, the base, which was previously combined with the organic acid, is found, after its destruction, combined with carbonic acid, in the state of a carbonate (546).

495. The most important of the organic acids, are the acetic, citric, tartaric, malic, and oxalic. There are few plants which do not contain a small portion of one or other of these acids, either in the free state, or else combined with a base. Unripe fruit contains a considerable quantity of malic and tartaric acids; and the leaves and stems of many plants, such as sorrel and rhubarb, have a strong, sour taste, from the presence of oxalic and malic acids. The nature and properties of acetic acid have already been described (476).

496. Citric acid exists in a free state, abundantly in the juice of limes and lemons, and in unripe gooseberries, currants, and other similar fruits; it is easily procured from these sources, in white crystals, having an agreeable, sour taste, and readily dissolving in water. Citric acid is used in considerable quantity by dyers: none of its salts are of much importance.
TARTARIC ACID.

497. Pure citric acid is obtained from the juice of limes or lemons, by a process resembling that used in the preparation of pure acetic acid (490). The sour juice is saturated with powdered chalk, which forms with its acid an insoluble citrate of lime; this is well washed with pure water, to remove the mucilage and other foreign matters present in the juice of the fruit. The pure citrate of lime is subsequently decomposed by dilute sulphuric acid, sulphate of lime is formed, and, as the clear solution cools, the citric acid is deposited in crystals.

498. TARTARIC ACID is obtained from the juice of grapes, pine-apples, and several other fruits; its principal source is wine, from which it is deposited as a super-salt of potash. The bi- or super-tartrate of potash is slightly soluble in water, but almost insoluble in a mixture of spirit and water; hence, after the fermentation of grape juice and consequent formation of spirit, the bitartrate of potash previously dissolved, is thrown down as a crystalline precipitate, constituting argol, or cream of tartar (309).

499. Tartaric acid is manufactured from tartar, by the use of lime and sulphuric acid. When tartar is boiled with chalk, the excess of wine in the tartar combines with lime, and forms an insoluble tartrate of lime. This salt, decomposed by sulphuric acid, yields pure tartaric acid.

500. The tartar deposited by red wine, has always a pink or red color, and contains some of the coloring matter of the grape. By boiling it with a portion
of powdered animal charcoal, the color may be removed. By merely exposing the crystals of tartar to the sun and air, they become bleached to a considerable extent; for this purpose they are spread out upon large sheets of canvas, and left some time thus exposed, after which, they are dissolved and recrystallized.

501. Tartaric acid is a white crystalline solid, like citric acid, easily soluble in water, and uniting to bases to form salts called tartrates. It is chiefly used in dyeing, and calico-printing; it is also used in medicine, and in the manufacture of saline and effervescent drinks.

502. Malic acid occurs associated with citric acid in apples, pears, and in the stems of common rhubarb. It forms salts called malates, but neither the acid nor any of its salts are of much importance; they are not used for any practical purpose.

503. Oxalic acid exists in considerable quantity in rhubarb, sorrel, and a great many other plants; it is, however, generally made artificially from sugar. When sugar, starch, lignin, &c., are heated with strong nitric acid, violent action takes place, and the organic substances are decomposed, the whole of their hydrogen being abstracted, and the carbon and oxygen left in those proportions which form oxalic acid (130).

504. The commonest salts of oxalic acid in plants are oxalate of lime and super-oxalate of potash. The former is a white insoluble salt, readily obtained by adding oxalic acid to a solution of lime. In conse-
quence of the strong affinity which lime has for oxalic acid, and the fact that the compound which they form is insoluble in water, oxalic acid is a very valuable test for the presence of lime and all its soluble compounds.

505. The super-oxalate of potash is a soluble crystalline salt; it is found in sorrel, rhubarb, and many similar plants, giving them a sour flavor; it occurs in many fruits associated with malic and tartaric acid. This salt, which is termed salts of sorrel, and sometimes also (though most improperly) salt of lemons, is used in taking out iron-moulds, or stains in linen caused by oxide of iron. This use depends upon the fact, that the oxalic acid has a strong affinity for oxide of iron, and forms with it a readily-soluble salt, whilst the acid used has not corrosive powers, and therefore does not in any way injure the texture of the linen.

506. The oily substances formed by plants are likewise very numerous; they are of two kinds, fixed and volatile; those which when exposed to the air remain without diminishing their bulk, and those which when so exposed evaporate. Olive oil, castor-oil, and cocoa-nut oil, belong to the former class, whilst oil of lavender, oil of cloves, and oil of lemons, are volatile oils.

507. All vegetable oils, whether fixed or volatile, contain carbon and hydrogen; in addition to which the greater number contain a portion of oxygen. They are all combustible, and when burning form
water and carbonic acid gas. Oils are found in all parts of plants, but rather more abundantly in the seeds and fruits than in other parts. The seeds of some plants, such as the poppy, linseed, rape, and mustard, contain frequently nearly half their weight of oil.

508. Fixed oils are divided into fat oils, and drying oils: the former remain fluid when exposed to the air, like olive oil; the latter gradually harden, and dry up into a kind of varnish, like linseed oil. Fat oils are used for soap-making, in wool-spinning, to burn in lamps, and in diminishing friction; those like cocoa-nut and palm oil, which remain solid at common temperature, are used for the manufacture of candles. Drying oils are employed in preparing paints and varnishes; for this purpose they are usually boiled, which causes them to dry and harden even more rapidly than in the fresh state.

509. The odors of plants are for the most part occasioned by the presence of volatile oil, in greater or less quantity. The volatile oils are obtained by distilling the plants which contain them with water; fixed oils are procured by pressing the seeds in which they are found, either cold or after a slight roasting; the former process gives the purest oil.

510. The most important of the non-drying fat oils, are those of the olive, cole-seed, rape, poppy, almond, mustard, and till, or sesamum. The chief drying oils are, linseed, hemp, nut, and castor. These
oils are all lighter than water, fluid at common temperature, and very combustible.

511. Amongst the solid oils may be mentioned the oils of palms and cocoa-nuts, and the various vegetable butters. The different waxes, too, of which there are a great number, are closely connected in nature with these solid vegetable oils. These substances all consist of carbon, oxygen, and hydrogen.

512. The volatile oils, such as those of turpentine, lemons, orange-peel, bergamot, sabine, and pepper, &c., are pure hydrocarbons. They contain no oxygen; while those of cloves, peppermint, caraway, and lavender, &c., contain oxygen. Camphor may be said to be an oxidized volatile oil; its composition is the same as oil of turpentine, with the addition of one equivalent of oxygen.

513. When the fixed oils are acted on by bases, they are decomposed, and resolved into certain fatty acids, and a peculiar sweet, uncrystallizable substance called glycerine. Most of these oils are mixtures of three distinct, fatty substances, distinguished by different properties; these are termed stearine, margarine, and elaine or oleine, and, when acted on by a base, they are respectively decomposed into glycerine, and stearic, margaric, or oleic acid.

514. Stearine is a white, friable, crystalline solid, which exists in many of the solid oils; it melts at a temperature of $130^\circ$; when saponified or heated with a base, such as potash, it is decomposed, and yields a soap, or salt of stearic acid and potash.
This soap is readily decomposed by a stronger acid, and then yields pure stearic acid, which resembles stearine in appearance and properties. It is, however, a distinct acid, and melts at 158°.

515. Margarine closely resembles stearine, but it is more fusible, melting at 116°. When saponified, it yields margaric acid, a substance very similar to stearic acid, but more fusible, melting at a temperature of about 140°.

516. Elaine, or oleine, constitutes the fluid part of oils; it remains liquid at all common temperatures, and, treated with alkalies, yields a fatty substance, oleic acid, which greatly resembles oleine in properties.

517. These substances may to some extent be separated by mechanical means alone. When semifluid oils, those which consist of stearine and oleine, are cooled, and then strongly pressed, the whole of the fluid oil or elaine may be squeezed out, and nearly pure stearine is obtained. The stearine, or mixture of stearine and margarine thus procured, is an excellent material for candles, whilst the fluid oil which is pressed out may be used for various other purposes not requiring solidity.

518. Stearic acid, however, being harder and still less fusible, makes even better candles than stearine; hence it has become a considerable manufacture. The stearine is boiled with alkali or lime, and the salt formed, then decomposed by a stronger acid.

519. Candles made of stearic acid are hard, bril-
liant, and nearly equal to wax: the acid, however, has a great tendency to crystallize, which interferes with its use for the manufacture of candles. At one time a small quantity of arsenic was added to the stearic acid, as it was found to counteract this tendency. It has since then been found that other substances will produce the same effect, and thus the addition of this very objectionable substance is obviated. A great number of different mixtures of stearine, stearic acid, wax, spermaceti, and other solid fatty substances, in various proportions, are used in the manufacture of candles.

520. Soap may be stated to be a salt, or rather a mixture of several different salts; namely, the mar-garate, stearate, and oleate of potash and soda. Soap is either hard or soft; the former is made with soda alone, or a mixture of soda and potash, whilst the latter is made exclusively with potash.

521. In making soap, all that is necessary is to boil together caustic alkali with the oil or fat to be saponified; carbonated alkali will not combine with the fat acids. The first thing, therefore, is to prepare a strong caustic alkaline lye; for this purpose the kelp, barilla, or other crude alkali, is mixed with lime, which decomposes the alkaline carbonates, and sets free the caustic alkali; the mixture is then lixiviated with water, and a strong alkaline lye is obtained.

522. The oil, fat, or tallow, is next heated in a suitable boiler, and the alkaline lye is then gradually
added, the mixture being kept constantly heated and well stirred. A cream-like liquid is thus formed, and when the proper quantity of alkali has been added, and the mixture has been boiled long enough, a quantity of common salt is stirred in, and the whole is allowed to cool, the fire being withdrawn.

523. Soap is easily soluble in pure soft water, but it is insoluble, or nearly so, in a strong solution of common salt; consequently, when salt is added to the newly-formed soap in the boiler, it causes the separation of the soap from the lye. The cream-like liquid quickly separates into two parts, the lower one being clear and transparent, and containing common salt, glycerine, and the impurities of the lye (513), whilst the upper part is thick and opaque. As it cools it solidifies, and forms hard soap.

524. The addition of common salt to a thick hot solution of soda soap has no other effect than that just described; but when it is added to a potash soap, there is produced in addition another effect which is purely chemical. The salts which the fatty acids form with potash are decomposed by the chloride of sodium; a mutual exchange of acids takes place; and hence, when a potash soap is mixed with a solution of common salt, both the soap and the chloride of sodium are decomposed, and soda soap and chloride of potassium are the results.

525. When the spent lye has been separated from the soap, the latter is heated a second time so as to melt it, well stirred up, and then poured into wooden
frames or moulds, to cool and harden. Good soap contains generally about one-third of its weight of water; the exact proportion depends upon the mode in which it is made. It is possible to prepare a solid soap containing as much as two-thirds of its weight of water.

526. Such soap looks very well whilst fresh, but it soon begins to shrink and shrivel up when kept. The retailers of soap generally store up their soap in a damp place, where it shall lose as little weight as possible by evaporation; sometimes they keep it in strong brine, and many devices are adopted to increase the weight of soap, by causing it to take up a larger proportion of water.

527. Common yellow soap is made in the same way as white or curd soap; but it contains, in addition, a quantity of resin. This is a useful addition; and such soap is both good and cheap. The yellow soap is softer and more soluble than hard white soap; there is consequently more of it wasted in washing.

528. Black, or soft soap, is made entirely with potash, and is generally manufactured from the commonest and cheapest oils; it is semi-transparent, and has usually a dark greenish color, speckled throughout with little white spots. These spots are caused by the addition of tallow, which, though it does not really in any way add to the goodness of the soap, is generally mixed up with it, because many persons imagine that the soap is not good, unless it exhibits this peculiar appearance.
529. Soap forms a clear solution in pure water; it will not dissolve in saline solutions, and is decomposed by a great many salts of lime, and, indeed, nearly all earthy and metallic salts decompose soap, and throw down insoluble compounds of the fatty acids with the bases of these salts. It is for this reason that hard water is not fit for washing; the soap is decomposed by it, and rendered insoluble. The slimy scum which is formed by soap in such water, is a compound of the fatty acids of the soap, with lime.

530. Water which is hard in consequence of its containing carbonate of lime (230), deposits earthy matter, and becomes softer by mere exposure to the air; the same effect is at once produced by boiling the water for a few minutes, so as to drive off the carbonic acid. Water which is hard from the presence of other salts of lime, may be improved by the addition of a little carbonate of soda; this will throw down some of the lime; and, rendering the water softer, will make it much fitter for washing, and prevent a great waste of soap.

531. Soap is easily soluble in spirit of wine; such a solution affords a very good test of the relative hardness of waters; the white precipitate it forms, is a measure of the quantity of earthy matter present in the water. When such a solution of soap is evaporated, the spirit is driven off, and the soap is obtained in a transparent state; various perfumed soaps are thus made. A solution of soap in spirit, to which
a portion of camphor has been added, constitutes opodeldoc.

532. The resins, which are not less numerous than the oils, resemble them in chemical composition and combustibility; they are formed in large quantity by many plants, and are very frequently seen adhering in large transparent drops to the bark of trees which produce them. Resins frequently look very much like gum, but they may readily be distinguished from gum by the fact that they are quite insoluble in water, whilst gum easily dissolves or softens in that fluid (335).

533. Most of the resins as they are obtained from the trees which yield them are fluid, or at least soft, and contain a portion of volatile oil; this is separated by heat, and then the pure resin is left. Common turpentine, which is a natural exudation, consists of resin and oil of turpentine; it is distilled, and the oil of turpentine is thus procured, whilst the resin called colophony is left. A healthy pine-tree yields from six to twelve pounds of turpentine annually.

534. When resinous woods are distilled or burnt, they yield tar. The trees are commonly cut down and burnt, so that the heat given out melts the resin in the wood; and that which escapes burning, runs down and is collected below.

535. In the manufacture of tar, the billets to be burnt are piled into a stack, and then covered over with turf; the combustion of the wood is thus smothered and kept down, and the loss of tar is prevented.
In France, the wood is generally distilled in large brick ovens. Tar differs from common turpentine in consequence of the mode in which it is obtained; it is darker colored, and contains less volatile oil, as a good deal of the volatile oil is injured and decomposed, during the extraction of the tar, by the heat of the fire.

536. When tar is heated strongly and boiled, the volatile oil which it contains is gradually driven off; there then remains a brittle, black, shining, fusible solid, called pitch.

537. There is another variety of tar, which is obtained during the manufacture of coal gas, and formed by the destructive distillation of bituminous coal. This substance resembles vegetable tar in nature and properties. It contains a volatile oil called naphtha, or coal-tar oil; and when this substance is driven off by distillation, there remains a kind of pitch. Coal naphtha resembles the native rock oil, or mineral naphtha; it is very combustible, and forms a valuable solvent for caoutchouc and some of the resins. It consists of hydrogen and carbon—the same elements, therefore, as oil of turpentine.

538. Most of the simple resins, like common rosin or colophony, the residue of the distillation of turpentine, are brittle, fusible, and very combustible solids, which easily dissolve in alcohol, or pyroxylic spirit; but are quite insoluble in water. They are found to consist of mixtures of several distinct acid substances. When resin is boiled in an alkaline
solution, it readily combines with the base, and forms a soluble salt—a compound of the acids of the resin with the alkali.

539. This property of resins of dissolving in alkalis is practically employed by the paper-makers, who prepare a size for the commoner sorts of paper by dissolving resin in a solution of carbonate of soda. Such a solution is immediately decomposed by any of the strong acids, which combine with the soda, and separate the resinous acid in an insoluble state. For the same reason, too, resin is used as a constituent of common yellow soap.

540. There are likewise found a number of gum-resins, as they are called, which have some of the properties of gum, and yet resembles resins; these are mixtures of gum and resin. They dissolve, though but imperfectly, in both water and spirit of wine.

541. Plants contain an immense variety of different coloring matters, all of which, however, when analyzed, are found to consist of nothing but oxygen, hydrogen, and carbon, or these three with the addition of nitrogen. The nature and properties of these substances are as various as their colors. They are all destroyed by heat, which burns them, like all the other varieties of organic matter. When burnt, the results are water, carbonic acid, and frequently ammonia.

542. Some vegetable colors are tolerably permanent, whilst others are very fugitive, fading in a short time when exposed to air and light, &c. Most vege-
table colors are soon bleached by sulphurous acid, and rapidly destroyed by chlorine. The art of the dyer consists in staining wool, linen, and cotton of various colors; some of the colors he employs are metallic, but the greater number are of vegetable or animal origin. Dyes are divided into those which at once permanently stain the fabric to be dyed, such as indigo; and those which require the use of some mordant or substance to fix them. The most important mordants are alumina, oxide of iron, and oxide of tin. These substances have a remarkable attraction for coloring matters, and also for the fibres of cotton, wool, and silk; they consequently assist in dyeing those materials of various colors.

543. Lastly, there are a number of peculiar vegetable substances, to the presence of which in plants their medicinal properties are in many cases due. Chemists have obtained from various plants a great number of substances called "active principles," some of which are highly poisonous, whilst others in small quantities are valuable medicines. Amongst this class of substances are the vegetable alkalies, as they are called; these are compounds of oxygen, hydrogen, carbon, and nitrogen, which possess the power of combining with acids and forming neutral salts. They are, of course, destroyed by heat, and, whilst burning, give off the usual results of the combustion of organic matter—water, carbonic acid, and ammonia.

544. All plants contain a small quantity of inor-
ganic matter. Besides the lignin, gum, sugar, starch, gluten, and other similar substances which consist of carbon, hydrogen, oxygen, and nitrogen, and which burn in the air when heated, there are always found substances which cannot burn, but which are of an earthy nature; these substances remain as ashes, when plants are burnt.

545. The inorganic substances found in plants are by no means numerous; the most common are salts of lime, potash, soda, magnesia, oxide of iron, and silica. These substances are found in the ashes of plants, combined with carbonic, sulphuric, phosphoric, and some other acids.

546. The ashes of plants very frequently contain a considerable quantity of carbonate of potash (494); this salt does not exist in the growing plants, but is formed during their combustion. Potash is found in a very great number of plants, sometimes in combination with sulphuric acid, or silica, but more commonly united to tartaric, oxalic, or some other organic acid, constituting a tartrate, oxalate, or other salt of potash. When plants are burned, all the organic acids are burned, as well as the lignin and similar substances which they contain; consequently, although in living plants the potash is combined with organic acids to form neutral or even acid salts, yet, as these acids are destroyed when plants are burned, caustic potash is left (199), which, of course, combines with carbonic acid to form carbonate of potash.

547. In the same way, carbonate of soda is fre-
quently formed by burning plants which contain salts of soda united to various organic acids. Before the mode of obtaining carbonate of soda from common salt (220) was discovered, the chief source of that salt was the ashes of sea-weed, and other marine plants.

548. The ashes of many plants contain silica (266), generally in combination with either potash or soda. Canes, and almost all plants allied to the grasses, are completely coated over with a thin film or varnish of silica. Sometimes even, as in the case of the bamboo, solid lumps or concretions of silica, called tabasheer, are found in the joints of the plant.

549. When a stack of hay or straw has been burnt, so that all the volatile or combustible matters have been dissipated, there is found a large quantity of a dark-colored glassy-looking substance, which consists principally of silicate of potash previously existing in the plants, and which, being unaltered by the heat given out during their burning, is merely melted, together with other substances contained in the plants, into a glass. This may also very easily be observed, by burning a piece of straw in the flame of a candle: abundant white ashes will be left, which continue to retain nearly the form of the straw, and which, if kept in the flame of a candle, melt into an imperfect sort of glass; this is principally silicate of potash.

550. The quantity of this substance which exists in the stems of different plants, varies much; some, such as the plants used for polishing, commonly called
Dutch rushes, contain even more than hay or straw; and cane contains so much silica, that it appears completely coated with flint.

551. Lime is almost invariably found in the ashes of all plants, and frequently constitutes a large proportion of the earthy substances present in them. It sometimes exists in combination with sulphuric acid, as sulphate of lime (237), but far more commonly united to phosphoric acid, or in the state of phosphate of lime (242). It is likewise very frequently found in considerable quantity in plants, combined with organic acids, particularly as oxalate of lime; hence the ashes of plants commonly contain a portion of carbonate of lime, which is formed in a similar manner to the carbonate of potash already described (494). The quantity of oxalate of lime in some plants is so large, that, upon cutting them through with a knife, multitudes of little crystals of this salt may be seen by means of a common pocket lens.

552. Magnesia is very often found associated with lime, more especially in the state of phosphate of magnesia (253), and as a double phosphate of magnesia and ammonia; but as this salt is decomposed by heat, the ammonia being volatile, only phosphate of magnesia is found in the ashes.

553. The ashes of almost all plants contain a trace of oxide of iron; and occasionally of the oxide of manganese also; and in one or two rare instances a minute quantity of copper has been found, especially in certain fungi.
554. **Animals**, like plants, consist almost entirely of four elements, but these are grouped or combined together to form various different compounds. The bodies of all living animals contain a considerable quantity of water: when this is evaporated by heat, there remain certain substances, which may be readily separated from each other in consequence of the different properties which they possess, just in the same way that we can separate the various substances which compose plants (318).

555. The constituents of the bodies of animals may be naturally divided into the organic substances of which the softer parts of animals are composed, and the earthy or inorganic matters, such as the bones.

556. Animal substances, for the most part, are rather more complicated in their nature than vegetable substances; they almost all consist of oxygen, hydrogen, carbon, and nitrogen; in consequence of the presence of this latter element, they are more liable to decompose than those vegetable substances which do not contain it.

557. The principal substances which compose the bodies of animals, and which are, therefore, called proximate animal principles, are: albumen, fibrin, caseine, gelatine, and fat, or oil; besides these, chemists have detected a multitude of other substances; but these five are the most important, or those which constitute the greater bulk of animal matter.

558. There are many varieties of albumen and
ALBUMEN.

fibrin, which, in consequence of peculiar properties, or slight differences in composition, have received different names; it is unnecessary to study the nature of all these substances; we may include them under the general terms, albumen and fibrin. Albumen consists of—

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559. ALBUMEN is a white, solid substance, which swells up, and slowly dissolves in water, forming a clear, transparent solution. Albumen is separated from its solution in water by the addition of certain acids, and also by the action of heat; when a clear solution of albumen is boiled, the albumen separates in the form of a white curd or scum; if the solution contain much albumen, the whole becomes solid.

560. This substance exists in many parts of the animal system. The white of the egg consists of hardly anything else; when a fresh egg is boiled, the albumen separates as a white curd; it coagulates, or becomes insoluble in water. Albumen occurs in a state of solution, in blood, and many of the liquids and softer parts of animal bodies. It is also found in a dry or solid form. Hair consists of albumen, together with a small portion of gelatine.
561. Dry albumen may be kept for a long time without undergoing decomposition; but in solution, or when moist, it is very liable to change. When heated, it burns; and the results of its decay, as well as of its combustion, are water, carbonic acid, and ammonia; it leaves a small quantity of white ash, which consists principally of phosphate of lime.

562. When perfectly pure, albumen appears to be nearly insoluble in pure water; but it readily dissolves in a weak alkaline solution, a very small quantity of which, gives water the power of dissolving albumen. Albumen is also soluble in acetic and phosphoric acid; these substances, therefore, do not throw it down from its alkaline solutions, but it is precipitated by sulphuric, muriatic, and most other acids, and the albumen thus thrown down is found to have combined with a portion of the acid used to precipitate it.

563. Most metallic salts also precipitate solutions of albumen, forming with it insoluble compounds; it is for this reason, that white of egg is recommended in certain cases of poisoning, as, for example, with corrosive sublimate. Albumen is used in many cases as a fining material; when mixed with any solution which it is desired to purify, and then precipitated, it separates in a curdy form, and, in so doing, entangles and separates the solid impurities previously suspended in the solution. When there is a large quantity of free alkali, or of acetic acid present, a solution of albumen cannot be coagulated by heat alone.
564. **Fibrin**, when pure, is a white elastic substance, insoluble in water, whether hot or cold; but soluble in acid and alkaline solutions. The muscle or flesh of animals consists principally of fibrin, associated with albumen and gelatine; when dry, fibrin may be easily preserved; but when moist, like albumen and gelatine, it soon begins to decompose. The composition of fibrin is almost identical with that of the albumen; they consist of the same elements, united together in nearly the same proportions. Fibrin contains—

<table>
<thead>
<tr>
<th>Element</th>
<th>Quantity</th>
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</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>5456</td>
</tr>
<tr>
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<td>690</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>1572</td>
</tr>
<tr>
<td>Oxygen, with</td>
<td>2282</td>
</tr>
<tr>
<td>Phosphorus and</td>
<td></td>
</tr>
<tr>
<td>Sulphur</td>
<td></td>
</tr>
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<td>10,000</td>
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565. Fibrin exists in the bodies of animals in two states; solid, and in solution: in the former state, it occurs in muscle or flesh; in the latter state, in the blood; the soluble fibrin of the blood, however, soon becomes insoluble after it has been taken from the body, and separates as a clot or coagulum. Blood consists of water holding in solution albumen, fibrin, a peculiar red coloring matter, fat, and various inorganic substances, including chlorides of sodium and potassium, carbonates, sulphates, and phosphates of potash, soda, lime, and magnesia, and also a portion of oxide of iron.
566. The blood which is sent out from the heart through the arteries all over the body, differs slightly in chemical nature, from the blood which returns through the veins to the heart, undergoing an important change in the course of its circulation; when blood leaves the heart, it is of a bright red color, and contains oxygen, which has been absorbed in the lungs; when it returns to the heart, it is black, and contains carbonic acid in place of oxygen (606). A pound of blood contains nearly two ounces of solid matter; of this about ninety grains is inorganic, the rest organic.

567. The fibrin, which exists in the arteries, is slightly different in chemical properties from that which is obtained from the veins. The latter approaches very nearly in character to albumen, it liquefies under the influence of a warm solution of nitrate of potash, and forms a solution which is coagulated by heat and metallic solutions. This kind of fibrin is almost identical with the fibrin of flesh. When fibrin is burnt, it leaves a light-colored ash, varying in amount from one to two and a half per cent.; it consists chiefly of phosphates of lime and magnesia.

568. Caseine is a substance which very closely resembles albumen, in its properties and composition; it is insoluble in pure water, but dissolves in dilute acid or alkaline liquids, in the latter far more perfectly than in the former: it exists in milk, in which it is dissolved by a small quantity of alkali; the addition of acid to milk, by neutralizing the alkali, causes the separation of the caseine as a curd.
569. Caseine differs from albumen in not being separated from its solutions by boiling; in chemical composition they are almost identical, the only difference being that caseine contains no phosphorus. Caseine consists of—

<table>
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<th>Element</th>
<th>Percentage</th>
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</tr>
<tr>
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<td>715</td>
</tr>
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<td>Nitrogen</td>
<td>1580</td>
</tr>
<tr>
<td>Oxygen</td>
<td>2173</td>
</tr>
<tr>
<td>Sulphur</td>
<td>36</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10,000</strong></td>
</tr>
</tbody>
</table>

Caseine contains about one quarter per cent. of inorganic matter.

570. Milk, like blood, consists principally of water holding organic matter in solution or suspension; milk contains caseine, fatty matter, a peculiar kind of sugar, and small quantities of saline substances, particularly phosphate of lime and various alkaline salts. When milk is kept at a certain temperature, it soon begins to change, a portion of the caseine turning into a kind of ferment, commences putrefaction; this induces an imperfect fermentation of the sugar, and converts it into lactic acid, and this acid by combining with the alkali present in the milk, causes the coagulation or separation of the rest of the caseine. Perfectly fresh milk is always feebly alkaline; but it frequently happens that a small quantity of lactic acid is formed almost as soon as it is exposed to the air, which combines with and neutralizes the alkali originally present in the milk.
The sugar of milk, when pure, cannot undergo the alcoholic fermentation; but by certain means it may be modified and converted into grape sugar, which can ferment. The Tartars, by causing milk to undergo the alcoholic fermentation, prepare an intoxicating drink which contains alcohol.

An imperial gallon of fresh cow's milk contains about one pound seven ounces of solid matter. Of this five ounces are fat, nearly eight ounces caseine, and about seven ounces sugar. The saline matters present in the milk include about eighteen grains of common salt, and one hundred and sixty grains of phosphate of lime.

Milk derives its opaque white appearance from the presence of innumerable minute globules of fatty matter, which are suspended throughout it; it is, in fact, a natural emulsion, or a solution holding particles of oil in suspension.

Cream, or the lighter part of milk, consists principally of these minute globules, or drops of oil, separated from each other by the solution of caseine, &c., in which they float; being lighter than water, they rise to the surface, when fresh milk is allowed to stand. The operation of churning consists merely in agitating the cream, under the influence of a moderate degree of warmth; the particles of oil then unite and collect together in masses, forming butter.

It is difficult to remove the whole of the caseine from butter, and its presence is highly objectionable from its great proneness to putrefy; by care-
ful and complete washing the greater part may be removed. Salt is generally added, as its presence greatly checks the tendency of the caseine to change, and consequently helps to keep the butter sweet.

576. In making butter there are two distinct methods employed. The one consists in churning the entire milk, the other in churning the cream alone; the latter is said to give the richest and finest butter, though the former is considered to give a larger yield. In either case a certain amount of incipient fermentation appears to be essential to the process of churning, and a moderate degree of warmth, as well as the contact of atmospheric air, are necessary.

577. The milk or cream intended to make into butter, is not used perfectly fresh, but is always allowed to stand some little time, in order that it may pass into this state of incipient fermentation, and a small quantity of lactic acid may be formed. The same state may be immediately brought on by the addition of a little sour cream, which acts on the rest as a ferment. There is some danger in doing this, however, that the whole may pass into the putrid fermentation (380).

578. In consequence of the extreme facility with which caseine undergoes decomposition, and the rapidity with which it passes into the state of putrid fermentation, the utmost care and attention must always be paid to insure complete cleanliness in the dairy. The least taint of impurity in the vessels employed,
is sufficient to bring on this sort of fermentation, and of course render the products useless.

579. However well butter is made, and however thoroughly it has been washed, it always retains a small portion of water and caseine; and in consequence of this it is apt to turn rancid and acquire a bad flavor. If butter be carefully melted, these impurities will fall to the bottom, and may be separated; but though butter so treated keeps better, yet it is always found that its flavor is slightly injured. Clarified butter may be kept sweet for weeks, and even months, without the addition of salt.

580. The yellow color of butter made in summer, is derived from the grass and plants on which cows feed. Winter butter is pale, or nearly colorless, and often has a tallowy appearance. Such butter is very frequently colored yellow by the addition of annatto, carrot-juice, or some other yellow coloring matter. These substances, though not unwholesome, can only be regarded as adulterations, and are decidedly objectionable.

581. The tendency of butter to turn rancid is much diminished by the use of a small quantity of common salt. It must however, be remembered that, though the bad flavor of butter which is very slightly turned, may to some extent be covered or concealed by the addition of salt, yet that its rancidity is not thereby at all corrected or remedied.

582. Cheese varies considerably in composition, according to the mode in which it is prepared. When
made from fresh milk, cheese consists of caseine and fatty matter; when made from skimmed milk, it contains little or no fat: in either case, during the pressing and curing of the cheese, it undergoes a peculiar kind of fermentation, which gives rise to several complicated chemical changes in its composition. Cheese is generally made from milk, by the action of rennet, the lining membrane of the stomach of the calf, which has the property of causing the coagulation of the caseine. Cheese which contains butter, becomes soft and viscid when toasted; whilst that which contains none, becomes hard and shrivels up almost like horn.

583. The curd of milk may be separated in a great variety of ways; any substance capable of entering into fermentation, and the greater number of acids, organic as well as inorganic, cause its coagulation. As the caseine of milk is held in solution by a small quantity of alkali, which causes fresh milk always to give a slightly alkaline reaction to colored vegetable test-papers; so the addition of an acid, or of any substance which by inducing fermentation can generate lactic acid, neutralizes this alkali, and consequently precipitates the caseine.

584. The most powerful of all the ferments which are thus able to cause this separation of the caseine, is rennet, which is in fact a membrane in a state of slow putrefaction. When a piece of this substance is put into milk, the temperature of which is then slowly raised to 120°, or thereabouts, a slight degree
of fermentation will be caused, and the caseine will be separated as a curd.

585. The whey, as the residue of the milk is called, after the separation of the curd, contains nearly all of the sugar; nearly the whole of the fatty matter which it contained is entangled and separated by the coagulation of the caseine.

586. The variety of cheeses is very great, depending upon a number of circumstances. One important distinction is caused by the state in which the fatty matter is associated with the caseine. In double Gloucester, for example, the oil and caseine are intimately mixed together; whilst in Cheddar, the oil is not so thoroughly mixed with the caseine, but is chiefly collected into little drops or globules, many of the little cavities in the cheese being, in fact, filled with oil.

587. Stilton cheese is made from entire milk, to which an additional quantity of cream has been added; Parmesan is made from skimmed milk; and Cheshire cheese is made from fresh milk without any alteration whatever. The difference between single and double Gloucester is caused by the fact, that the milk for the former is partially skimmed so as to remove about half the cream, whilst that for the latter is left entire. Cream cheese, properly speaking, is not really a cheese; it is rather cream, from which the watery parts have been allowed to drain.

588. A very strong and high-flavored cheese is sometimes made, by allowing milk to become sour
spontaneously, and then collecting and pressing the curd thus formed. The color of cheese is very commonly heightened by the use of annotto, or some other yellow dyestuff. In using these substances, care should always be taken to use only pure materials.

589. Gelatine, the fourth great principle of animal matter, is a tough, colorless substance; in cold water, it very slowly softens and dissolves; in boiling water, it dissolves more readily, and forms a solution, which becomes a jelly as it cools. The skin, horns, and hoofs of animals, consist principally of hard, dry gelatine; and it likewise occurs in many of the softer parts of the body, associated with albumen.

590. Common glue and isinglass consist almost wholly of gelatine; the former is prepared by boiling the clippings of skin, refuse horns, hoofs, and similar matters, in water; a strong solution of gelatine is thus obtained, which, as it cools, becomes a jelly, and is then termed, size; this, when cut in slices, and dried in the air, is the glue of the shops.

591. Isinglass is the lining membrane of the swimming bladder of the sturgeon; but inferior sorts of isinglass are obtained from other fishes. Gelatine is insoluble in spirit. In the dry state, it may be preserved unchanged, for any length of time; but when moist, or dissolved in water, it very soon begins to change; it becomes mouldy, and putrefies. Gelatine consists of—
Carbon . . . . . . . . . . . 5077
Hydrogen . . . . . . . . . . . 715
Nitrogen . . . . . . . . . . . 1832
Oxygen with
Sulphur and
Phosphorus . . . . . . . . . . . 2376
                      10,000

592. One of the most remarkable properties of this substance, is its strong affinity for tannin; when any astringent solution, which contains tannin, is added to a solution of gelatine, they combine together, and form an insoluble, elastic compound, which is remarkably stable, and does not putrefy or ferment. The skins of animals, which consist chiefly of gelatine, are converted into leather, by tanning, or steeping them in solutions of tan.

593. Gelatine is used as an article of food, as in the preparation of soups and jellies; by whitewashers and paper-makers in the form of size; in the manufacture of glue; and, as a fining material for clarifying wine, beer, &c., when they are turbid. This latter use is a purely mechanical one; a jelly, commonly called finings, is prepared by the action of very weak vinegar on isinglass, and a quantity of this is diffused through the liquid to be clarified; it of course precipitates any tannin which may be present, and as it subsides carries with it all the solid impurities which were previously suspended throughout the liquid, rendering it turbid.

594. The fat of animals is perfectly similar in
nature to vegetable oil: some kinds of fat are solid, others fluid, at common temperature; but they all become fluid when made sufficiently hot; animal oils, like those of vegetable origin, contain no nitrogen—they consist of carbon, oxygen, and hydrogen, and when burnt form carbonic acid and water; like vegetable oils, also, they consist of margarine, stearine, and oleine, united to a peculiar base (513), and consequently they form soap when boiled with alkalies.

595. The bones of animals contain a very large proportion of earthy matters; and, indeed, derive their strength and solidity principally from the quantity of those substances which they contain. When bones are burnt, there remains, after the combustion of all the organic matter which they contain, about three quarters of their weight of earthy substances; this is phosphate of lime, together with a small portion of carbonate of lime; bones consist of phosphate and carbonate of lime, cemented together as it were with gelatine and a little albumen; they also contain a small quantity of oil. Ivory and the teeth of animals are composed of the same substances as bone. Hoofs and horn likewise contain phosphate and carbonate of lime, but in far less quantity; they consist principally of gelatine (589).

596. The same remarkable similarity of chemical composition which is found amongst vegetable substances, is likewise observed amongst those of animal origin; the various proximate elements which constitute the bodies of animals, are, for the most part,
almost identical in composition, and, like vegetable substances, they appear more or less convertible into each other. A very slight alteration in the relative proportion of the elements of which they consist, causes very great differences in their nature and properties. As in consequence of the very peculiar nature of nitrogen, all substances which contain it are exceedingly liable to change, therefore all those forms of animal matter which contain any nitrogen, very soon pass into a state of decomposition.

597. On comparing together the various substances which constitute animal and vegetable matter, it is observed that lignin, gum, sugar, and starch, are all perfectly distinct in nature and properties from any of the substances usually found in the bodies of animals: a remarkable similarity, however, exists between the fibrin, albumen, legumine, and oily substances of plants, and certain forms of animal matters.

598. The albumen, fibrin, and legumine of plants closely resemble the albumen and fibrin of animals; indeed, some of the varieties of these substances may be said to be absolutely identical; they consist of the same elements, and possess the same properties: thus, for example, there is no chemical difference between that variety of albumen which exists in peas, beans, and other leguminous seeds, which is called legumine, and that form of albumen which is found in milk, cheese, &c., and called caseine (347, 569).

599. When albumen, fibrin, or caseine, are acted on by a solution of potash or soda, they soon dissolve,
forming a clear solution; and if an acid is added to this, so as to neutralize the alkali, a precipitate falls, which is precisely the same from whichever of these three substances it is obtained; to this precipitate the name of protein is given. Protein consists of oxygen, hydrogen, carbon, and nitrogen, like the substances from which it is obtained; but it contains no sulphur or phosphorus, whilst they always contain a small quantity.

600. The knowledge of this fact, that the fibrin and albumen, &c., of plants, are identical in composition with some of the most common forms of animal matter, throws great light on the nutrition of animals. It shows that the gluten and albumen of plants used as food, may immediately enter into the system of the animal; whilst gum, starch, sugar, &c., must undergo a change, before they can constitute a part of the body of an animal.

601. It has already been stated that vegetable and animal oils consist of the same three elements; many of these oils contain precisely the same proportions of carbon, oxygen, and hydrogen: hence by some chemists it has been supposed, that the oil which exists in those vegetables used as food, might contribute directly to the formation of fat, without undergoing any change; though, on the other hand, many facts might be quoted to show that this view is improbable; and that the fat of animals is formed from starch, gum, and sugar, by a kind of fermenta-
tion in the animal system; and that the fatty matters
of the food are not directly appropriated by the ani-
mals which feed on them.

602. Bearing in mind, then, that the strength of
man and animals depends mainly on muscle, and
that the formation of muscle is greatly dependent on
the amount of organic subject containing nitrogen in
their food, it becomes a matter of the first importance
to study the mode of increasing the quantity of these
matters in food.

603. Some animals feed entirely on vegetable
food; others feed entirely, or in part, on flesh: in
either case they derive their nitrogen, or the sub-
stances containing it, from plants. Animals do not
appear to have any power of absorbing nitrogen from
the air; all the albumen, fibrin, &c., which they con-
tain, is therefore either directly or indirectly obtained
from plants.

604. The most important of the chemical functions
of animal life may be classed under the two great
heads of nutrition and respiration; and, consequently,
food also may be divided into those kinds which con-
tribute to the one or other of these two objects.
The changes which ordinary food undergoes in pass-
ing through the stomach of an animal, are briefly
these: mechanical division, effected by chewing, &c.;
chemical division or digestion, effected in the stomach;
chemical transformation, conversion of starch, &c.,
into animal matters; absorption of azotized matters
identical in composition with animal matters, and
CIRCULATION OF THE BLOOD.

which are passed directly into the blood; and lastly, separation of useless matters as excrementitious.

605. The chemical office of food is, to supply to the body albumen and fibrin, the elements of blood; in order to counterbalance the waste continually going on in the system, by the constant addition of all those matters which enter into its composition; and secondly, to contribute to the formation of animal matter, by the transformation of starch and other substances of vegetable origin.

606. Respiration has already been described (107) as being, chemically speaking, a mere process of combustion, in which carbon and hydrogen are burned at the expense of the oxygen of the air; this process of combustion is carried on through the medium of the blood, and goes on in all parts of the body. In the lungs, the blood is exposed to the contact of a quantity of atmospheric air; oxygen is absorbed, and carbonic acid given off; the blood thus charged with free oxygen is sent, by the action of the heart, to all parts of the body, in the innumerable minute bloodvessels which terminate the arteries; carbon and a portion of hydrogen are taken up, and the oxygen, which leaves the heart free, returns to it through the veins, converted into carbonic acid and water; the former is at last given off from the blood, and expired from the lungs, previous to the absorption of a new quantity of fresh oxygen from the air.

607. Two great objects are effected by the circulation of the blood: the one is the removal of carbon
and hydrogen, by means of free oxygen, which, by combining with those elements, produces heat, and keeps the body at a uniform temperature; the other is the addition of new matter, to replace that which is removed. The digested food, or chyme, as it is called, on passing from the stomach into the smaller intestines, becomes mingled with a portion of bile, a secretion of the liver, consisting of soda and a peculiar fatty acid. In passing through the smaller intestines, the chyme is separated into two portions—one containing the elements of blood, called chyle, which is absorbed and carried into the blood, the other containing rejected matters, which are passed from the system as excrementitious.

608. The true nature of the change produced in food, by the action of the gastric juice in the stomach of an animal, is by no means satisfactorily understood. This remarkable secretion contains a notable quantity of free muriatic acid, and possesses great powers of reducing organic matters to a state of solution; especially, when aided by the temperature of the body.

609. The excess of fluid taken into the system with the food, is conveyed away from the body through the medium of the kidneys, as urine; a secretion which also contains the rejected azotized matters of the blood, in the form of two peculiar substances, to which the names of urea and uric acid have been given; these substances consist of carbon, oxygen, hydrogen, and nitrogen, and are very prone
FOOD OF ANIMALS.

237

to decompose, especially when mixed with other forms of organic matter.

610. Urea has been made artificially, by chemical means, but uric acid has not yet been so obtained: its only source is the animal system. When urine is kept in a moderately warm temperature, it soon begins to decompose; when fresh, it is generally slightly acid, but after a short time, it becomes alkaline from the decomposition of urea, and formation of ammonia. The half solid urine of birds and serpents contains a large quantity of uric acid, in combination with ammonia. Guano, the decomposed excrement of sea-fowl, likewise contains a considerable portion of this salt.

611. It is evident that food of all kinds may be classed under two great divisions; according as it yields the elements of flesh, or contains substances capable of being at once transferred to the blood, and so carried to all parts of the body; or as it merely contains substances capable of undergoing transformations, or of supplying the waste caused by respiration.

612. The quantity of food required by an animal for either of these objects, varies greatly, and depends entirely on circumstances: when the waste going on in the system is great, a large supply of blood-making food, that which is rich in the elements of nutrition, will be required. When, however, the body is exposed to cold, or to violent exercise, the loss must be met by a proportionate increase in food
rich in the elements of respiration. It happens, however, that the food of animals, for the most part, is rich in both forms of nutriment.

613. A large accumulation of fat in an animal can never be considered as healthy; but, on the other hand, leanness or the absence of fat is also unhealthy, because, if at any time exposed to cold, hunger, or violent exercise, the tissues of the body itself will be consumed and converted into elements of respiration, whilst in an animal supplied with a reasonable proportion of fat, the latter will be consumed first, before the tissues of the body will be thus acted on.

614. Some animals fatten far more easily than others do; this depends partly on their general construction, and partly on the circumstances to which they are exposed—such as temperature, abundance and kind of food, exercise, &c. In man, too, the greater or less activity of the mind exerts a very remarkable influence on all the functions of the body; and, therefore, amongst others, on the formation of fat.

615. Setting aside individual peculiarities of construction, the size of the lungs, &c., the circumstances most favorable to the formation and secretion of fat, are warmth, little exercise, abundance of food, and the absence of all worry and irritation. Under these circumstances, an animal is supplied with as much of the elements of flesh as suffices to keep up the healthy state of all the tissues of the body; having an excess of the elements of respiration, there is a natural
tendency to store up fat, which is that form of matter which accumulates in animals, as a provision against future demands, just as plants form and store up starch. Hence the effect of stall-feeding, upon cattle.

616. In cold climates a larger quantity of the elements of respiration are required, or the tissues of the body will begin to suffer. In carnivorous animals there is always more or less of this waste of the body going on, and hence the demand for azotized food. In the case of herbivorous animals, or those which feed wholly on vegetables, there is very little of this waste. In cold countries a larger quantity of food is required, and can be digested, than in hot ones. A greater quantity of the elements of respiration is needed to generate the proper amount of heat; and at the same time, as the air is much colder, and therefore more condensed, a larger quantity of oxygen is taken in at the lungs by each inspiration (52, 107).

617. The art of cookery, or the preparation of food, is a very important one, and has been yet only partially brought into a systematic and intelligible form. A great many of the processes of the cook can be explained and regulated on known chemical principles; but, at the same time, there are also many which appear to depend on facts not hitherto recognized or explained.

618. The object of cookery is to render digestible and palatable the various substances used as food;
many of which, without such preparation, would be wholly indigestible and useless. The chief agents employed are heat and moisture. The real object of the cook should be to render wholesome food digestible and palatable, and not, as is too often the case, to render unwholesome things agreeable to the palate.

619. In boiling meat two things are effected. It is exposed to a heat of 212°, and, as far as it is in contact with the water, the soluble substances which it contains are extracted. These are, in fact, very different operations, though they are commonly confounded together.

620. The albumen of flesh, which is to a considerable extent in a fluid state, is coagulated and brought into the solid form by exposure to a heat of boiling water; albumen begins to coagulate at a temperature of 168°. Now, when meat is plunged into boiling water, the albumen in the outer parts is immediately rendered hard and insoluble, and the passage of heat to the inner parts of the meat proceeds very slowly. If, therefore, the joint is of any size, the inside will not be thoroughly boiled, until the whole has been boiled so long that the outside is quite overdone.

621. In order to boil meat well, it should be put into cold water, and then gradually heated up to the boiling point; a slow and gradual application of heat is that best fitted to render meat tender and digestible. The time of boiling must, however, of course depend on the size of the joint to be cooked. Meat
should never be boiled rapidly, not only for the reason just given, but also because, when the water boils strongly, the steam carries off a large portion of the volatile matters of the meat, and so renders it less savory and palatable. The best effect is produced by only allowing the water to simmer or boil very gently. In a large kitchen, boiling by steam is better, more manageable, and more economical, than with an open fire.

622. The gelatine of flesh, on the other hand, is softened by the action of the hot water; it is chiefly in a solid form, and softens and gradually dissolves in the heated water. In steaming, by the long-continued action of a gentle heat, the whole of the gelatine is softened and brought into an easily digestible condition.

623. When meat is made into soup, if simply simmered in water, the gelatine only will be dissolved. The fat either remains entangled in the fibre of the meat, or melts and rises to the surface of the water. By adding vegetables, or any substance containing starch, which acts as a thickener, a sort of emulsion is formed, and a large quantity of the oil remains divided and suspended throughout the soup, much in the same way that the oil or butter is naturally diffused in milk (573).

624. Generally speaking, gelatine is more easily digested than albumen, though the latter is the more nutritious. It is a mistake to suppose that the jelly of meat is the most nutritious part of it; the value
of gelatine as a part of food is commonly overrated. Its value consists chiefly in its being easily soluble, and therefore more readily digestible, than albumen or fibre.

625. In boiling vegetables, very similar results are produced; the solid parts are softened and rendered more soluble, whilst the albumen is coagulated. It should always be borne in mind, that there is nothing gained by heating the water very strongly so as to make it boil rapidly; but, on the contrary, it is highly objectionable. Water which simmers, is very nearly quite as hot as that which boils rapidly, and does not carry off so much of the volatile principles of the food.

626. When it is desired to soften and dissolve food as much as possible, pure soft water is best; but when it is designed only to soften, but not dissolve it, hard water is preferable. In general, the solvent powers of hard water are much less than those of soft water. For this reason, pure soft water is best for making soup, whilst hard water is best for boiling joints of meat. Salt is constantly added to the water used for cooking; this diminishes its solvent powers, rendering it harder, and therefore better fitted for those operations which are intended to soften the food, but not dissolve it.

627. In roasting meat, the chemical changes produced are not very dissimilar from those effected by boiling. A good deal depends on the management of the fire; if it is too hot at first, the outside of the
meat will be scorched and burnt before the inside is properly warmed. By the use of the spit, or by otherwise causing the turning of the joint, the action of the fire is rendered more slow and gradual; and by basting, or continually pouring the gravy which drips from the roasting meat, over it again and again, the evil effects of too much heat on the outside are guarded against.

628. The effect of roasting meat, is to harden the albumen; whilst the gelatine is liquefied, and the fibre becomes softened and rendered easily digestible. If meat is over-roasted, so that it becomes dried up, more harm than good is done; because, when the heat has driven off the natural juices of the meat, its continued action, instead of softening it more, renders it hard and less soluble, and therefore less digestible.

629. Fat, taken in conjunction with other substances, is a valuable part of food, provided it enters the stomach in a proper state. Fat should always be divided and mixed up with other substances, so that the mere application of heat may not at once cause its separation. The oil in seeds is so divided by the starch and other matters with which it is associated, that it is wholly digestible; ground up with water, oil seeds furnish an emulsion resembling milk.

630. The oil used in cookery should, as far as possible, be brought into this state; if it is permitted to come into the greasy state, it is not only far less
digestible itself, but likewise produces a similar effect on the substances with which it is mixed. These two conditions may be well observed, on comparing together good melted butter, with that which has been allowed to "oil."

631. Common salt is a necessary ingredient of all good food. Its use in the preparation of food is evident from the preceding remarks; it is also valuable in its preservation, acting as an antiseptic, and preventing it from undergoing fermentation or change. Salt is further important as aiding digestion, to which both the soda and muriatic acid it contains are necessary (608).

632. The effects of various condiments and spices on the appetite are very remarkable; stimulating it, and sometimes in a very beneficial manner. The principles upon which these substances act is very ill understood, and cannot be satisfactorily explained by reference to ordinary chemical facts; indeed, it can hardly be otherwise till the chemistry of digestion is more fully understood.

633. The phenomena of digestion in some respects resemble those of fermentation. The substances used as food, are for the most part all very liable to undergo fermentation; and the nature of the change thus brought about depends wholly upon circumstances. Any derangement in the process of digestion, may permit some other change to commence, and the system must then be immediately thrown out of order. A very great number of diseases may,
either directly or indirectly, be traced to some disturbance in the functions of digestion.

634. Very little is known respecting the mode in which medicines act, or the effects which they produce when taken into the stomach. The influence of some substances may, to a certain extent at least, be explained, such, for example, as dilute sulphuric acid, which it is evident must react upon any common salt it meets with, generating muriatic acid and sulphate of soda. Again, the manner in which one form of indigestion, arising from the incipient fermentation of food, is arrested by certain volatile oils, or by saline draughts charged with carbonic acid, may be explained; but the mode in which bark, opium, and indeed nearly all other drugs act, is quite unknown.

635. It is true that chemists have ascertained that these substances derive their active powers from the presence of minute quantities of various peculiar principles which they contain. These substances have been separated, analyzed, examined, and named; but the knowledge thus obtained has not thrown much light upon the real mode in which they act. It has merely shown the nature of the active agent, but neither the way in which it acts, nor the principle upon which its activity depends (513).
CHAPTER VII.

THE FOOD OF PLANTS—ITS NATURE AND SOURCES.

636. Having shortly described the elements of vegetable matter, and enumerated the principal compounds of those elements which are found in plants, we may at once proceed to consider the sources of the food of plants; that is to say, the means naturally provided to insure them a due supply of the various substances necessary to their growth; namely, oxygen, hydrogen, carbon, nitrogen, and the various earthy and saline substances which are always found in plants.

637. There are only two sources whence it is possible for plants to derive these matters; namely, the air and the soil; let us inquire what substances they can obtain in this manner, and how they avail themselves of the food thus offered to them.

638. It has been already stated, that the air at all times contains a small quantity of carbonic acid gas (37); it likewise always contains a still more minute proportion of ammonia, which is constantly being formed by decay. Here, then, we see that the air contains the four elements of organic matter; and
when, in addition to these facts, we remember that the air is always more or less damp, it is easy to understand that plants can derive from the air, alone, the greater part of the substances which they require (41, 150).

639. Although the air contains so large a proportion of oxygen, and although that substance is in a free state, that is to say, not combined with any element, but ready to combine with any substance for which it has an affinity; yet it does not seem that plants derive the oxygen which they contain directly from the air.

640. In the same way, there is no evidence to show that they are able to absorb nitrogen from the air. It might have been supposed that plants would obtain the nitrogen which they require, directly from the air, which contains nearly four-fifths of that gas; but there is very good reason to believe that this is not the case, and that plants can only obtain nitrogen, or assimilate it, as chemists say, by absorbing it in combination with some other substance.

641. What has just been said with regard to oxygen and nitrogen, is equally applicable to carbon and hydrogen: the former is a solid substance, and therefore, as one might rightly conclude, plants cannot absorb it in the separate state; when combined with oxygen in the form of carbonic acid gas, and possibly also when in the form of carburetted hydrogen (131), it can be absorbed by plants. Hydrogen has never been found in the air, except in a state of combina-
tion; the commonest compounds of hydrogen, and indeed almost the only ones from which plants could obtain that element, are water and ammonia. It may be laid down as a rule, that plants can only absorb oxygen, hydrogen, carbon, and nitrogen, in a state of combination, and moreover that those compounds, to be absorbed, must be either fluid or gaseous.

642. The soil consists of silica, alumina, lime, magnesia, oxide of iron, small quantities of various alkaline and earthy salts, and a portion of decaying organic matter. It likewise contains water, and the small quantity of ammonia and carbonic acid which the rain has brought down from the air. Plants cannot derive the elements of organic matter from the earthy constituents of the soil, or from the organic matter which it may contain, unless there is air present; by the action of air these substances decay, and are gradually changed into gases, which plants can absorb.

643. It is commonly supposed that plants derive the whole of their food from the soil; but this is an error: it is a fact well ascertained by chemical experiments, that plants derive the greater part of their nourishment from the air, although the soil is equally essential to their growth.

644. The earthy substances contained in plants are principally obtained from the soil: it is true that the air contains exceedingly minute traces of various earthy and saline substances, which are suspended in
it in the form of dust, and carried about by the winds; but the quantity which plants can derive from this source is comparatively small. The air near the sea-shore, and even to a great distance inland, is frequently loaded with saline particles derived from the sea: after a storm at sea a quantity of salts thus suspended in the air is very considerable.

645. The fact that some solid substances can be thus carried along in the air in a state of very fine powder is important, and well worthy of being remembered, as it explains many apparently mysterious phenomena. It is exceedingly difficult to detect the very minute quantity of solid matter contained in the air, but there is no doubt that it often does exist, although we are not aware of its presence.

646. Setting aside the small quantity of earthy matters which plants may derive from the air, it is evident that the great source of the lime, potash, and other similar substances contained in plants can only be the soil: hence, the chemical composition of the soil must exert great influence on the plants which are cultivated in it.

647. The soil or earth is essential to the growth of plants in several distinct ways. It enables them to fix themselves firmly, as, from its open porous nature, it allows the roots to extend in various directions, and obtain a secure hold, so that the plant can grow erect into the air, without danger of being blown away by the winds. The soil likewise supplies them with substances essential to their growth; such as
carbonic acid and ammonia, either generated by the decay of substances which the soil naturally contains or absorbed from the air, and also yields them the earthy and alkaline salts which they require.

648. As may be supposed, the soil is very variable in composition; its nature is generally a good deal dependent on the subsoil and stony matters which are buried beneath the surface, many of which are slowly decomposing or crumbling away, and adding to the soil the substances of which they consisted.

649. The origin of all soils appears to be the disintegration or gradual crumbling down of rocks, from the action of frost, and various chemical and mechanical agents; and, therefore, all soils contain innumerable little fragments of different rocks, which are slowly but constantly becoming smaller, as the chemical combinations of which they consist are broken up and destroyed.

650. This gradual decomposition of stony particles in the soil is caused by the action of the air. Many of the common rocks are compounds of several different earthy and alkaline substances, in which silica, united to lime, alumina, potash, and soda, forms silicates of those or similar bases (267).

651. Silicates of this kind, or natural compounds containing silica in combination with several earthy and alkaline bases, are quite insoluble in water, and are scarcely acted on, even by the strongest acids; nevertheless, they gradually decompose when exposed to the air. Under the joint action of the moist-
ure and carbonic acid in the air, these compounds are disintegrated, carbonate of potash or soda is formed, and in part washed away by the rain, whilst the insoluble earthy bases are left, in the form of a very fine powder.

652. The most abundant constituent of soil is commonly silica, which frequently forms nearly nine-tenths of the whole of its weight; but this is by no means always the case, for in calcareous or limestone countries, we frequently find soils containing a very large quantity of lime; whilst others, again, contain a large proportion of alumina. These differences in the proportion of the earthy components of the soil, give rise to the varieties of light or free, and stiff or clayey soils, which are also modified by the presence of a greater or less quantity of organic substances.

653. Silica and alumina, which are generally the two principal constituents of soils, differ very greatly in their respective uses. The former is of importance, both chemically and mechanically. Chemically, as forming a soluble compound with alkali, and thus being absorbed by the roots of plants, and conferring strength and stability to their structures; and mechanically, by diminishing the extreme closeness and tenacity of alumina, and thus, by making the soil more porous and open, allowing the passage of air and water through it, and enabling the roots of plants more easily to penetrate it than they could were it wholly composed of alumina.

654. On the other hand, the use of alumina is
principally mechanical, tending to keep the soil moist, from its attraction for water, and likewise being highly useful in absorbing ammonia, both from the air and from all decaying substances evolving it in the vicinity, by virtue of that property which many porous substances, and more especially charcoal, possess, of absorbing or condensing that gas (162); which, as we shall shortly show, is of great importance in the growth of plants.

655. Soils differ greatly in their mechanical as well as in their chemical nature. The same substances constitute a soil possessing very different properties, according as they are in the form of little grains like sand, or in very fine powder. This state of mechanical division is of great importance for several reasons, and most particularly in relation to water. A soil containing a large quantity of alumina is generally known by its stiff, tenacious character, and is remarkable for its great retentive power for water; whilst those consisting principally of silica, and more especially those in which it exists in the form of sand, are generally light and porous soils, and far less retentive of water.

656. Again, a soil containing alkaline silicates in the form of little grains, always contains free alkaline matter in a soluble state, set free by the decomposition of those silicates, and this separation of alkaline matter continues so long as there is any of the solid silicates left; this effect would cease in a short time if all the silicates were very finely powdered; they
would soon undergo decomposition, and the whole of
the alkaline salts would then be washed out by the
rain (651).

657. The best soils are those in which the earthy
constituents are so proportioned that the light,
porous qualities of the one are balanced by the close,
retentive properties of the others; for they are then
most uniformly suitable to vegetation.

658. The silica and alumina in soils are of course
almost wholly free and uncombined with any acid, as
the former is not a base, and the latter has hardly
any affinity for the weaker acids, such as the carbonic.
Small quantities of silica are almost always found in
soils, combined with either soda or potash, forming
those curious compounds before alluded to, in which
the silica seems to play the part of an acid (259);
soils never contain more than a very small quantity
of these substances; but it is evident that plants,
such as grasses, which contain silica, must obtain it
from the soil in a soluble form, by gradually absorb-
ing it in combination with alkali, dissolved in water.

659. Silica also exists in soils in combination with
several bases together, such as lime, potash, soda,
magnesia, and alumina, constituting the natural
rocky silicates just spoken of; the nature and com-
position of these compounds, as well as their tend-
ency to decompose, varies considerably in different
soils.

660. Lime and magnesia, both of which have a
powerful affinity for acids, are never present in the
soil except in combination with some acid, and this is most commonly the carbonic; the former substance is also not unfrequently found combined with sulphuric acid, constituting gypsum, or sulphate of lime (237).

661. The oxides of iron in the soil are usually uncombined, as they have not sufficient attraction for carbonic acid to combine with that gas, which is always present in the air. They in great measure occasion the variations of color observed among soils; for according as the iron is in a state of protoxide or peroxide, it gives to the soil a black or brownish-red color.

662. It must not be supposed, however, that the color of soils is wholly dependent on the iron which they contain, or that the blackness of any particular soil is indicative of the presence of oxide of iron. The decomposing vegetable substances, which all soils contain in greater or less quantity, are usually of a brown or black color, and therefore not unfrequently give a very dark color to a soil which only contains a very small portion of oxide of iron.

663. Sulphate of iron is also sometimes present in soils in very small quantity, being formed by the gradual oxidation of sulphuret of iron in the manner previously mentioned (286). A very minute quantity of this salt of iron confers upon the soil peculiar properties, which render it appropriate for particular plants; but a slight increase of its amount is attended with bad results, for, unless in very minute
quantity, it is poisonous to plants; and hence such soils as contain much sulphate of iron are invariably barren, or incapable of supporting healthy vegetation.

664. The saline matters in the soil are principally the sulphates, muriates, nitrates, and phosphates of the alkaline and earthy bases. The nature and quantity of these substances vary considerably, but they never constitute more than a very small portion of the soil; their influence upon vegetation is, however, considerable, for certain plants seem to require particular saline substances, and do not grow well in soils not containing them.

665. Thus all kinds of grass and corn contain silica and phosphoric acid; the former substance requires the presence of an alkali, either potash or soda, to render it soluble, and to enable it to enter the roots when it is dissolved in water; whilst the latter is also always associated with a base, which is usually either lime or magnesia. In the same way, all plants are found to contain small quantities of similar saline and earthy matters.

666. The relative proportion of the different ingredients of soils vary very greatly. The chief constituent of nearly all soils is generally silica, in the form of sand; and in fact there are comparatively but few good fertile soils in which this is not the case. The sandy matter of most silicious soils consists of nearly pure silica, but in some cases it contains alumina, lime, &c. (625). The following ana-
COMPOSITION OF SOILS.

lyses, taken from Sprengel's book on the composition of soils, will serve as examples of the general nature of ordinary soils; they are examples taken from lands in the north of Germany.

Silica and fine Sand 94,908 92,980 96,721
Alumina 610 820 370
Oxides of Iron 1,080 1,666 480
Oxide of Manganese 263 188 trace
Lime, chiefly combined with Silica 141 748 5
Magnesia 208 168 80
Potash 56 65 trace
Soda 44 130 36
Phosphoric Acid 80 246 trace
Sulphuric Acid 41 trace trace
Chlorine (in common Salt) 4 trace 58
Humic Acid 400 764 800
Humus, containing Nitrogen 2,070 2,225 1,450

100,000 100,000 100,000

667. The nature and quantity of the organic substances in the soil have great influence upon its character and fertility. They act in several ways. By slowly decaying, they present a constant source of carbonic acid, and likewise of ammonia, if they contain nitrogen; whilst, by rendering the soil open and porous, and at the same time augmenting its absorbent powers, they assist in keeping it uniformly moist. They also increase the means which the soil possesses of absorbing and condensing ammonia and carbonic acid from the air.

668. It is a remarkable fact, that a mixture of two or three earths, such as lime, silica, and alumina, is
better fitted to absorb moisture and gaseous matter, than either of the earths taken singly; and in the same way, the addition of organic matters to the soil has the effect of increasing this power still further; and, therefore, as was before said, the best soils are those containing a quantity of each of the three earths, mixed with a moderate proportion of decaying organic matters.

669. Some soils contain a very large quantity of organic substances, and indeed not unfrequently consist almost wholly of the remains of vegetables: the different varieties of peat are of this character. In such soils we observe the bad effects of the presence of too much organic matter in a soil. It is thus rendered tough, porous, and spongy, so that it is soon saturated with water, or completely dried, by changes of weather. Besides all this, it frequently contains a peculiar acid, produced during the decay of vegetable matter, which is hurtful to the growth of plants.

670. Another use of the organic constituents of the soil is, that they furnish to the plants growing on it the saline substances which they naturally contain; and which not being subject to decay are gradually washed out by the rains during the decomposition of the organic substances, and are ready to be absorbed by the roots of growing plants.

671. The decay of the vegetable remains in the soil, never proceeds so far that the whole of the elements which they contain are converted into carbonic

22*
acid and water. The first change which they undergo, when exposed to air and moisture in circumstances favorable to decay, is a diminution of the quantity of oxygen and hydrogen, and likewise, though in a smaller relative proportion, of the carbon which they contain.

672. The result of this is a change of color somewhat similar to that produced by slightly burning or singing them by heat: they become more or less brown, according to the degree to which this change has proceeded. Whilst in this state, they constitute what is called humus, the name by which chemists distinguish all kinds of vegetable matter when in a state of decay, and more especially woody fibre, which indeed usually constitutes a very large proportion of the bulk of plants.

673. This process of decomposition proceeds slowly for a considerable time, the relative proportion of the oxygen and hydrogen to the carbon gradually decreasing, carbonic acid and water being all the time formed, until, at last, the humus, consisting of carbon with only a small proportion of oxygen and hydrogen, ceases to undergo any further change; it then is termed mould by chemists.

674. In order to understand this process, which is one of great importance in connection with the growth of plants, it must be remembered that woody fibre and most vegetable substances consist of carbon, oxygen, and hydrogen. Though the elements of these substances are united together in nearly the propor-
tions to form carbon and water, we must nevertheless clearly understand that they are not composed of carbon and water, but that they are distinct compounds of the three elements; and therefore, of course, that the water produced during their decay is not merely set at liberty by a process of decomposition, but is formed by the hydrogen which is given off during decay, entering into combination with oxygen.

675. When woody fibre and similar vegetable substances are exposed to air and moisture, they decay; the chemical affinity which binds their elements together seems to be weakened, and the elements acquire a tendency to form new and more simple combinations. The oxygen and a portion of the carbon unite and constitute carbonic acid; whilst a portion of the hydrogen combining with oxygen from the air, gives rise to the formation of water.

676. This explains the use of air in facilitating the decay of these substances; for it is evident that if free oxygen be required to combine with the hydrogen, this process cannot go on without a constant supply of fresh air. The rapidity of the decay of vegetable matters in the soil, therefore, depends greatly on the porosity or closeness of the soil, permitting a more or less perfect action of air.

677. In soils containing a large proportion of clay or alumina, the decay of vegetable matter or humus proceeds very slowly, because the close, dense nature of the soil greatly retards the free access of air; though in other respects, such soils are decidedly
favorable to decay, inasmuch as they remain long moist, and a certain degree of moisture is essential to decay.

678. It is necessary to acquire a clear conception of the nature of what is called humus, and the office which it performs in the nutrition of plants. It was formerly supposed that plants derived their carbon from the organic matter of the soil, that the brown decaying substances called humus were directly absorbed by plants, and their carbon appropriated: it is now known that this is not the case. The real use of humus is, that it constantly generates carbonic acid.

679. When water is poured on humus or decaying vegetable matter, very little of it is dissolved, and the water acquires a pale yellow color: if, however, the water contains potash or soda, either caustic or in the state of carbonate, it dissolves far more humus, and acquires a deep brown color; the cause of this is, that the alkalies facilitate the decay of that substance. Under the influence of the alkalies, a peculiar acid is formed, which has accordingly been termed the humic. The brown solution obtained under these circumstances contains humic acid in combination with potash or soda.

680. It has been supposed that this acid might be formed by the action of alkalies on the humus or decaying matters which the soil contains, and that the humates, or compounds of humic acid, thus formed, being absorbed by plants, might supply them with
carbon. There is, however, no proof whatever that this really is the case; on the contrary, as has already been stated, there is every reason to believe that decaying vegetable matters merely yield carbon to growing plants, by generating carbonic acid gas.

681. The food of plants, then, is carbonic acid gas, water, and ammonia, partly derived from the air, and partly from the soil; and certain earthy and saline substances, derived almost wholly from the soil. Let us consider the circumstances under which plants are able to absorb and assimilate these substances, and what other conditions are requisite to their growth.

682. The important influence which water has on the changes which vegetable substances are liable to undergo, has already been alluded to, when speaking of albumen, fibrin, and the other similar substances which enter into the composition of seeds. All these substances, which under the joint action of air and moisture, are so liable to undergo various changes, become comparatively fixed and unchangeable, when well dried and protected from the action of water. During the germination of seeds, a great change takes place in the nature of the substances composing them. Carbonic acid is formed by the combination of a portion of the carbon which they contain, with the oxygen of the air. The conditions requisite to the germination of seeds are, therefore, moisture, a certain degree of heat, and the presence of air.

683. Light appears to be prejudicial, and, there-
fore, darkness may be said to be also necessary to the perfect germination of seeds; and these conditions are all secured by a good soil. The office performed by moisture is, in the first place, principally mechanical, serving to soften and swell up the dry matters composing the seed, and by assisting in the introduction of air to facilitate the changes requisite to germination; it likewise acts chemically, its presence being essential to the conversion of starch into gum and sugar. It is also possible that water may sometimes be decomposed during germination, its oxygen combining with carbon to form carbonic acid.

684. When seeds germinate in a confined portion of air, we find that the air does not increase in bulk at all; the nitrogen is not at all affected, but the oxygen is found to have combined with a quantity of carbon, and to be converted into carbonic acid, and, in consequence, the insipid and comparatively insoluble starch contained in the seeds, has changed into more easily soluble, sweet, and mucilaginous, or gummy substances, fit for the nourishment of the embryo plant, before its roots and leaves are sufficiently developed to enable it to derive support from the different sources of food presented to it.

685. When the access of air is prevented, whilst at the same time seeds are exposed to moisture, they are unable to undergo this change; germination, or the growth of the embryo, cannot take place, and the seeds decay; a portion of the water is decomposed,
and the oxygen and hydrogen which it contained unite with the elements of the vegetable matter, and form carbonic acid, ammonia, carburetted hydrogen, and other substances.

686. The first chemical change which takes place in a germinating seed, is the change of a portion of its azotized constituents into diastase (360), a substance which does not exist previously in seeds, but is formed by the process of germination: it appears to be a peculiar modification of vegetable albumen. This change cannot be effected without the presence of free oxygen, which removes the excess of carbon. The azotised matter converted into diastase immediately begins to act as a kind of ferment, causing the starch and gum of the seed to pass into the state of sugar. During the earliest stage of germination, a small portion of vinegar or acetic acid is formed. Alkalies tend to assist germination, because they combine with and neutralize this acid.

687. The chemical changes attendant on the process of germination, are very different from those which go on during the growth of a complete plant. Before the formation of leaves, and in the very earliest stages of its growth, a plant requires only warmth, moisture, and the presence of air; but subsequently, it requires, in addition to these, carbonic acid and light.

688. It is the oxygen of the air alone which is essential to the germination of seeds, for the nitrogen being unable to combine with carbon, under or-
ordinary circumstances, is quite useless in diminishing the quantity of that substance contained in the seeds: its presence in the air is, however, very useful, serving to dilute the oxygen and to prevent its acting too rapidly. Seeds are found to germinate very quickly in pure oxygen gas, but the plants produced are weak and unhealthy.

689. The germination of seeds may be readily effected in water, although they are for the most part unable to grow and vegetate under that fluid. The change which, under ordinary conditions, is effected in seeds by the oxygen of the atmosphere, is, under these circumstances, caused by the oxygen of the small quantity of common air always dissolved or held in solution by water. In no case can a seed germinate unless free oxygen is present, or some other means exist by which a portion of the carbon in the seed can be removed, so as to cause the change in the constituents of the seed before described.

690. In the process of malting, or converting raw grain into malt, the object to be attained is to change a large portion of the starch which the barley contains into gum and sugar. This is effected by steeping the grain in cold water, and then heaping it up together on the floor of the malt-house; it is thus placed in the most favorable conditions for germination, and consequently the chemical changes attendant on that process immediately commence. The seeds lose carbon, and at the same time convert the oxygen of the air into carbonic acid; the embryo
or young plant begins to increase in size, and the grain becomes warm from the heat evolved by the chemical action. In malting barley about 20 per cent. of starch is changed into sugar and gum (412).

691. If this were suffered to continue, the sugar and gum formed would be consumed by the growing young plant, and the malt would then become useless. When, therefore, the process has proceeded to a certain extent, it is stopped by the gradual application of heat, which, by driving off the water and drying the grain, destroys the conditions requisite for the further growth of the young plant, and preserves the grain in the state most proper for the uses of the brewer and distiller.

692. Malting essentially consists of four distinct processes, which are respectively called steeping, couching, flooring, and kiln-drying. The barley is first of all steeped in stone vats or tanks filled with water; it swells, and at the same time gives out a small quantity of carbonic acid, together with some coloring matter. The grain is allowed to remain in the steeping tanks for from forty to forty-five hours; sometimes the water is renewed; during this time it has expanded considerably, and increased in weight from fifteen to twenty per cent. The malster judges that it has been steeped long enough, when it is so soft that the two ends of the grain can be easily squeezed together, between the thumb and finger.

693. The water is then drained off, and the steeped grain is spread over the floor in square heaps, about
SEED STEEPING.

thirty inches deep, in which state it is allowed to remain about twenty-six hours. After this, the grain begins to heat, and, if left to itself, would ere long become so hot that it would be injured. This is prevented, however, by the malster; who takes care to turn over and gradually diminish the thickness of the heaps of malt about twice a day. When the steeped grain is first couched, the surface grains soon dry up a little; but as soon as the heap begins to heat a little, these grains again become moist; this is termed the sweating of the grain, and shows that germination is commencing.

694. The time required for malting varies according to the temperature and the kind of malt it is desired to make; about fourteen days is the average time. Winter is far better suited to malting than summer, because in the former it is much easier to regulate and keep down the temperature than it is in summer. The malster judges of the progress of his operation entirely by the growth of the young roots, which the seed puts forth.

695. In drying malt, the great object is to drive off water and destroy all further growth, without exposing the malt to so high a temperature as would risk its injury or decomposition; as, in that case, the sugar which had been formed would be destroyed and lost. The application of heat, therefore, must be very gentle and gradual; in fact, the less heat the malt is exposed to, the better will it be. When thoroughly dried it is screened, so as to remove the
dried rootlets, which are of course brittle, and hence easily detached from the grain. After this, the malt is once more spread out and exposed to the air, in order that it may mellow and become soft and mealy.

696. There are several chemical means by which germination may be accelerated, but in general very little benefit is derived from such processes. Thus, for example, substances which have a strong affinity for hydrogen assist germination, because they tend to decompose water, and set free oxygen. For this reason, solutions containing free chlorine are found to cause seeds to germinate more rapidly than they would otherwise do.

697. When seeds are sown in the earth, they are placed in a situation where they are protected from the action of light, and exposed to the influence of air and moisture; and provided they are not too cold, that is, above the freezing point of water, they will germinate and produce plants. Below that temperature seeds cannot germinate, because, as has been already said, the action of water is in the first instance to soften the seed, and this it cannot effect when so cold as to assume the solid form itself. If buried too deep in the soil, the free access of air to the seeds will be retarded, or even perhaps wholly prevented, and under these circumstances of course they cannot grow.

698. The nature of the soil, too, is of considerable importance: very clayey soils allow a less perfect
action of air than those which are of a more porous and open structure, and in the former, therefore, the process of germination proceeds more slowly.

699. In the next stage of the growth of plants, or when they have leaves, they begin to absorb carbon from the air instead of parting with it; and this they do by decomposing the carbonic acid always present in the air, a power which they possess when exposed to the influence of light.

700. The effects of light in increasing and diminishing chemical affinity are highly curious, and but very imperfectly understood: there are a good many substances which, although they have an affinity for each other, cannot combine in the dark. In the same way, that peculiar chemical change which goes on when plants grow, cannot proceed well in the dark; the decomposition of carbonic acid and water, and the combination of their elements in order to form lignin or starch, &c., goes on very imperfectly without light.

701. The effect of Light, in causing chemical combination and decomposition, is quite independent of its brightness or illuminating power. The rays of light which reach the earth from the sun, possess three distinct classes of properties; those which give light, those which give heat, and those which cause chemical action. In passing through different substances it is found that one or other of these properties is lost; the heating rays, for example, passing
through, whilst those which give light are stopped (187, 295).

702. Chemical action is caused in the cells of the leaves by the agency of these chemical rays of light, which enable the vital power as it is called, to form organic matter: hence it is frequently said that Light stimulates vegetation. Chemical action is also augmented by Heat, which, though it cannot alone effect those changes usually produced by light, considerably assists them, and is therefore classed with light as a stimulus.

703. Electricity also, that marvellous power, in many respects so similar to light and heat, the effects of which we frequently see in lightning and thunder storms, is believed to exert great influence on vegetation; but though there certainly appears much probability of the truth of this, it is as yet not proved by direct and unexceptionable experiments.

704. When the young plants appear above the surface of the ground, their mode of growth is changed: they then require air and light, and being no longer preserved by the soil from the action of light, are immediately able to effect the decomposition of carbonic acid.

705. The roots of a plant require little more than moisture. The water which they obtain from the soil contains those saline and gaseous matters which the plants want, whilst the leaves require in addition to these matters the influence of light, and all these they obtain by growing up into the air.
706. When plants are shaded from the light, or covered up, either artificially or by the leaves of surrounding plants, the action of light is impeded, and they are unable to effect the proper decomposition of carbonic acid. Every one knows that, under these circumstances, plants become unhealthy and send up long weak shoots, endeavoring to reach the light, and to place their leaves in a situation where they will be exposed to its influence.

707. The existence of most plants may be divided into four periods: firstly, germination, or the development of the young plant from the embryo; secondly, the growth of the plant to maturity; thirdly, blossoming, and the formation of seed or reproductive parts; and, fourthly, decay, or a cessation of vitality, and consequent decomposition of the organic structure.

708. In each of these four periods different chemical changes are going on, and therefore different conditions are requisite for the perfection of those various processes. In the commencement, water, air, a certain degree of warmth, and the absence of light, appear nearly all that is required for the growth of the embryo, which obtains food from the matters stored up in the seed, until it has so far increased in size as to have acquired roots and young leaves, and in other respects has become fitted to derive nourishment from external sources.

709. During the growth of the perfect plant, it is observed that, in addition to water and a certain
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degree of warmth, carbonic acid, ammonia, and certain earthy and saline substances, are required; whilst light, which in the first stage of its growth appears prejudicial, is now required to assist in those complicated chemical changes which are going on, when the compound organic substances, such as woody fibre, gum, starch, gluten, &c., are formed in the organs of the plant by the combination of the elements of water, carbonic acid, and ammonia.

710. Hence during the day, or whilst plants are exposed to the influence of light, the carbonic acid absorbed by them is decomposed, the carbon alone being retained in their structure, whilst the oxygen is given off.

711. During the night, or when light is withdrawn, this process of decomposing carbonic acid and acquiring carbon goes on very slowly, and accordingly some of the carbonic acid then absorbed by the roots is given off again unchanged, by other parts of the surface of the plants, in place of undergoing decomposition. For plants are at all times receiving from the soil carbonic acid, which must necessarily enter their system with the water they absorb from the soil through their roots; but they can only derive nourishment from it when, in consequence of the influence of light, they are able to effect its decomposition, and so, by retaining the carbon, to increase the quantity of organic matter which they contain.

712. In the third stage of the existence of a plant, or during the formation of seed, the same general
conditions are required which are necessary during the second; for the formation of seeds and fruit is governed by very nearly the same laws which regulate the production of leaves and woody fibre. Those causes which are most influential in maintaining the healthy growth of the leaves, such as temperature, a due supply of food, and more especially light, are likewise the most important in the growth and perfection of the seed.

713. The principal chemical use of the leaves is to assist in the formation of the organic substances existing in plants, which they do by exposing the crude juices to the action of light, and thus effecting those peculiar changes dependent on the combination of carbon, nitrogen, hydrogen, and oxygen, under the influence of light, on which the growth and increase of plants depend.

714. The organic substances thus formed in the leaves are either stored up in the stem of the plant, or in other ways appropriated to its increase and perfection; and in most cases but little of them remains in the leaves. The office of the fruit or seed-vessel is different; for though, like the leaves, it is able to assist in the production of organic matter, the substances so formed are retained and stored up in itself, and the fruit or seed-vessel, not only returns nothing to the plant on which it grows, but in addition attracts to itself a large portion of the vegetable matter formed, by the combination of carbon, oxygen,
and hydrogen, under the influence of light in the leaves.

715. Thus, whilst the leaves are continually adding new matter to a plant, the fruit, on the other hand, by absorbing that matter, checks the growth. Hence, also, the practice of pruning fruit-trees with a view to improve the fruit; the young fruit thus gets an additional supply of organic matter, which would otherwise have gone to the formation of fresh leaves had not the branches been removed. It must not, however, be supposed that the formation of flowers and seeds is similar to the formation of woody fibre, or leaves; very different changes take place, but the same general conditions are required by plants in both cases.

716. Flowers do not, like leaves, possess the power of decomposing carbonic acid under the influence of light; on the contrary, flowers appear at all times to give out carbonic acid. From this it is evident, that they must consist principally of substances containing in proportion less carbon than the ordinary proximate principles, such as gum and lignin. There is a common belief that plants in blossom deteriorate the air, and, therefore, that their presence in bedrooms is highly objectionable. It is true that flowers generate carbonic acid; but it is probable that the evil effects occasionally produced by flowers in sleeping-rooms, are occasioned by a minute quantity of volatile oil, to the presence of which the smell of flowers is to be attributed; and many of which, even in very
small quantities, act powerfully on the animal system.

717. When fruits are first formed, they act very like leaves; they absorb food from the air, and under the influence of light form organic matter, whilst at the same time they collect and appropriate much of the organic matter generated by the surrounding leaves. At this period of their growth fruits have very little taste, and in composition they somewhat resemble leaves; when they have attained a certain size, they undergo a new change and ripen, during which they acquire a sweet, or slightly acid taste.

718. In the first stage of the ripening of fruit, acid matter is generated, in consequence, apparently, of the conversion of some of the tasteless constituents of the fruit into malic, tartaric, and other organic acids. During the second period of ripening, the greater part of these acids is converted into sugar, and a portion of coloring matter is at the same time formed. The formation of sugar and coloring matter takes place quite independent of the plant; for fruits may be ripened after removal from the plant which produced them.

719. In the first period of vegetation, when a seed has been placed in the conditions requisite to germination, the embryo plant has no power of obtaining food for itself, either from the air or the soil; it is entirely dependent on the seed for a supply of those matters which are necessary to its growth. A seed, then, is a magazine, or store of food, prepared for
the use of the young plant; and accordingly a large quantity of the matters secreted by a plant, are necessarily employed in the formation of seed.

720. In the case of all plants which shed their leaves annually, at the end of the season buds are formed. In some respects these resemble seeds; they consist of growing points, surrounded by small scales, which in time would develop into branches and leaves; but they are closely wrapped up in several layers of a brown scaly matter, which protect them during the winter from the action of cold and other injurious influences.

721. When spring comes on and the weather is milder, the buds undergo a chemical change a good deal resembling germination. It frequently happens that the buds open before the winter is really over, or that there is a return of cold weather, which kills them, and thus gives the tree a very material check. Buds may be formed at any time during the year; but they are, generally speaking, and of course with the exception of flower-buds, formed only at the end of the year, and at the time when the leaves fall off. In cold and uncertain seasons it is common, however, to see two or even more sets of buds formed in the same year.

722. There is a great distinction between organic and organized matter. The former term signifies, as has already been explained, sugar, starch, or similar compounds; whilst by the latter name is meant the cellular and fibrous parts of plants. The chemist
may by artificial means readily make some forms of organic matter, and convert them into others; but he cannot make organized matter; the latter is a product of vital action, and has never been formed by artificial means.

723. The first change effected by plants, is to convert carbonic acid and water, under the influence of light, into starch and similar forms of organic matter; the second operation is to form these substances into the various cells and tubes which compose the structure of plants. The former is a purely chemical operation; the latter requires something more, namely, the exertion of vital power.

724. As starch, though it is very easily convertible into other substances under the influence of a ferment, is nevertheless one of the most stable of all the products which a plant forms; so it is also that one best fitted to serve as nutrition to a young plant, and is the substance always prepared in plants, to act as a store of nutriment for any future growth. Whenever a growing point or embryo is formed, a quantity of starch also is, at the same time, deposited.

725. Thus starch is found in all seeds and buds, as well in those which are formed on underground stems and tubers, as in those which are formed at the extremities of the branches. Whenever an embryo or point of growth begins to develop, the chemical action, in the first instance, is the same. The azotized matter begins to change, diastase or some similar substance is generated; this acts on the
store of starch, and sugar and other soluble principles are formed. For this reason all kinds of tubers, such as potatoes, for example, become sweet as soon as the eyes or buds begin to grow: it is an effect similar to the malting of grain.

726. The embryo of a seed possesses vitality, but has no power, at first, of assimilating the elements of organic matter directly from the air; it is, therefore, able to convert starch, gluten, and other proximate principles, into organized matter, and consequently grows and increases in size, by feeding on the store of organic matter prepared for its use in the seed.

727. The means provided to insure the reproduction of plants, are very various. In some plants, seed only is formed; whilst in others we observe tubers or collections of food, prepared and preserved in underground receptacles, for the nourishment of the young plants of the next season.

728. Many plants, such as wheat, barley, beans, &c., form seeds which consist merely of an embryo, or growing body, surrounded with a supply of organic food sufficient for the use of the young plant, until it has grown large enough to feed itself. The seeds of other plants are more complete, being surrounded with an additional quantity of organic matter, which, by decaying, insures to the roots and leaves of the young plant, a due supply of gaseous food as soon as it is able to feed itself by those organs.

729. The seeds of all plants which bear fruit, are of this kind. When a peach, for example, falls from
the tree which produced it, the soft pulpy matter of
the fruit soon decomposes, and by the time that the
seed begins to germinate, the fruit is wholly converted
into humus; hence the roots of the young plant are
placed in a soil rich in carbonic acid, as soon as it is
possible for them to absorb it.

730. During the whole existence of a plant, from
the formation of its first leaves till its death, it is
constantly absorbing moisture from the soil through
the roots, and as constantly parting with it by the
leaves. The water thus collected from the soil, con-
tains carbonic acid, ammonia, and such saline matters
as are present in the soil. If the leaves of the plant
are exposed to the action of light, the carbonic acid
and ammonia are decomposed, together with a portion
of the water, and organic matter is formed.

731. During the night, or in the absence of light,
this change takes place but imperfectly; and hence,
then, the water absorbed by the roots, which is still
given off by the leaves, carries with it a considerable
portion of the gaseous matter collected from the
soil.

732. The exact office in the nutrition of plants,
performed by the saline substances they absorb, is as
yet but imperfectly known. Some plants appear to
require particular substances, and it is known that
certain substances cannot be formed by plants, unless
the soil contains particular saline matters. All plants
contain more or less lime, as well as other bases, such
as potash and soda; these, of course, are never in
the pure state, but combined either with organic or inorganic acids, or else with chlorine. Albumen, gluten, caseine, and fibrin, are always found to contain a small quantity of certain phosphates, particularly those of lime and magnesia; hence these substances appear to be essential to the formation of those principles in plants.

733. It is not known whether plants have any power of selection by their roots; that is to say, whether they are able to absorb from the soil only those substances which they require, or whether they absorb all the soluble matters present in it. To a certain extent, they seem to have some power of selection, but at the same time they are very frequently injured by the absorption of poisonous matters from the soil.

734. When the roots of a healthy young plant are plunged into a vessel containing water, in which two different saline substances are dissolved in equal proportions, the plant will not take up both salts in similar quantity. If, after continuing this experiment some days, we then evaporate the remainder of the solution, we shall find that the plant has taken up the two salts in very different proportions; perhaps half of the one salt has been absorbed, but only one-third of the other.

735. The quantity of inorganic matter found in plants is various at different periods of their growth. In general, young plants contain a larger proportion than old ones; and as seeds contain a supply of all
that young plants require, a quantity of these salts is necessary for the ripening of seed; hence plants require a large supply of inorganic matter from the soil, during the third period of their existence, or during the formation of seed.

736. These are, in very general terms, the conditions requisite to the growth of plants; but there are many special conditions appropriate to particular species, and many important circumstances, which must not be overlooked in studying the chemistry of vegetation. The differences of climate, and their influence on the growth of plants, are not less remarkable than those of the soil itself; and, indeed, the very same soil would possess a very different degree of fertility in different climates.

737. In the hot moist regions of the tropics, plants grow with far more rapidity, and vegetation is more vigorous, than in temperate regions. In tropical countries, decay proceeds far more rapidly than it does in our own country; carbonic acid and ammonia, the food of plants, are produced in greater quantity than here; whilst, from the greater power of the sun, plants are able to assimilate more of those substances than they can in colder countries.

738. The same circumstances which are favorable to the putrefaction of organic matters, are likewise those which facilitate the decomposition of inorganic compounds in the soil (651). Consequently, in warm tropical climates, a more copious supply of the requisite earthy matters is continually being set free,
ready to be absorbed by the plants; proportionate, in fact, to the augmented vigor of the vegetation.

739. Hence, in endeavoring to introduce into one country the plants of another, it is necessary to consider the conditions under which they naturally grow; and care must be taken to imitate as closely as possible those conditions, not merely as regards soil, but likewise in respect to temperature, moisture, and light. These, however, are practical matters, into which it is unnecessary here to enter.

740. In the last stage of the existence of a plant, or when, either from excessive cold, disease, or merely old age, the vitality becomes extinct, all that curious series of changes by which organic matters are generated under the influence of light in the cells of plants, ceases; decay commences, the organic compounds of the plant begin to decompose, and their elements rearrange themselves into simpler forms. The greater part of the hydrogen combines with oxygen, and is gradually given off in the state of water; the nitrogen and hydrogen combine, and form ammonia; whilst the carbon is slowly dissipated in the air in the state of carbonic acid, and at last little remains beside mould, or charcoal in combination with a small quantity of oxygen and hydrogen (673), and the insoluble earthy matters which the plant may have contained.

741. A plant dies, when, at the end of the season, it has passed through the various stages of its existence, fulfilled the office for which it was created, and
formed seeds or reproductive particles, which will insure a fresh growth of the same kind of plant next year; after this, vitality gradually declines in energy, the formation of fresh organic matter ceases; and that already formed begins to decompose.

742. In following out the chemical changes which occur during the life of a plant, we find all throughout, that there is a constant struggle going on between vital force and the ordinary chemical affinities of the elements of vegetable matter. The influence of vital energy tends to form complex combinations, whilst the natural affinities of the elements tend to form simple ones. So long as the former is the more powerful, the plant grows and flourishes; as soon, however, as the latter gets the upper hand, the plant withers and fades—disease, and perhaps death, follow.

743. This kind of struggle is well seen in the annual fall of the leaf. During the whole of the summer, whilst the vital energy is great, lignin, starch, sugar, and other similar compounds, are formed in the leaves, and through their agency; towards the close of the year, when they have fulfilled the objects for which they were formed, these effects cease, the vital power diminishes, and ordinary chemical agency gets the ascendant. The first effect of this is oxidation; the leaves begin to absorb oxygen, their green coloring matter oxidizes and becomes yellow, the leaves wither, fade, die, and fall off. But even then chemical power continues to act on them; they
remain subject to its influence till they are wholly decomposed, or nothing but a little mould remains.

744. Disease is occasioned by many causes; but independent of numerous minor sources, such as the attacks of insects, &c., the most important are deficiency of light, deficiency of vital energy, and deficiency of heat; any of these alone, or a combination of them, induces disease in plants, and often leads to death, by enabling the elements of organic matter to commence those changes which constitute decay and decomposition.

745. As has already been observed, plants are constantly employed in preserving the purity of the atmosphere: they feed on the foul or vitiated air produced by respiration, combustion, and decay; and their vigor and luxuriance are always in proportion to the impurity of the air. It must, however, be clearly understood, that there is a limit to the quantity of carbonic acid plants are able to decompose; an excess would kill them; whilst, if there were none of it in the air, they could not live. The vegetation of the whole globe is just enough to keep the air in a uniform state of purity (106, 123, 710).
CHAPTER VIII.

ARTIFICIAL SOURCES OF THE FOOD OF PLANTS—ACTION OF MANURES.

746. When we remember, that although plants derive a large portion of their nourishment from the air, yet at the same time they absorb considerable quantities of saline matters from the soil, it is evident that all plants must more or less impoverish the soil, by taking away that which causes its fertility.

747. The natural vegetation of any country enriches rather than deteriorates the soil, because nothing is carried away from its surface; the plants which grow on it return to the soil, during their decay, all the earthy and saline substances which they had absorbed from it during their growth, whilst they add to it a considerable quantity of the carbon they had collected from the air.

748. Very different, however, is the condition of cultivated lands; on them large crops are raised year after year, which are removed and carried away to a distance, to form the food of men and animals. Such land is gradually impoverished; because with the crops a large quantity of inorganic matter, ne-
cessary to the fertility of the land, is removed. It remains for us to inquire, what are the best means of restoring these matters to the soil; and likewise, in how far it is possible to increase the fertility of the soil, by augmenting the natural quantity of any of its constituents.

749. In practice there are three courses adopted to maintain the fertility of the soil; namely, fallowing, a rotation of crops, and the application of manure.

750. In order to comprehend the effect produced by fallowing, it is necessary to remember the mode in which soil is formed. As has already been stated, soils consist principally of small fragments of different rocks and stones, which, from a variety of causes, have gradually crumbled down to powder. The sand or fine particles of stone in the soil are constantly decomposing (649) and adding to the soil, potash, soda, and very finely divided earthy matters; it is from sources of this kind that most of the saline substances present in the soil, are derived.

751. The soluble salts formed in this manner, are dissolved by the rains; and in great part washed away from the surface; a portion, however, always remains in the soil, and is absorbed by plants. When a crop of some plant, requiring, for example, a large quantity of potash, is raised in such a soil, it often happens that the crop takes away nearly all the soluble potash the soil contains; and in consequence, it would be impossible to raise a second crop of that
752. If, however, the soil is left for some time fallow, if no crop at all is raised on it, the soil has time to renew itself; by the action of the air, the further decomposition of the silicates and other similar stony compounds in the soil, is effected, and a fresh supply of potash is provided.

753. The same end is obtained by a system of rotation. In place of sowing a second crop of the plant which requires so much potash, some other plant is taken, which does not require potash, but which requires other substances that the soil contains.

754. In both these cases we restore, by the slow effects of the air, those substances which we remove with the first crop. In manuring, we at once add to the soil that which the plants require.

755. Generally speaking, wheat cannot be grown on the same land year after year, and accordingly various systems of rotation are adopted. It is probable, however, that as soon as we know what are the substances required by wheat and other crops, we shall at once know how to restore to the soil those substances which each crop removes, and thus be enabled to obtain crops of the same plant, for any number of years in succession, from the same soil. The labors of chemists are, however, hardly far enough advanced for this at present.

756. The period of time over which the rotation
is made to extend, varies greatly in different countries and in different soils. The ordinary period in England is four or five years, but it is sometimes extended to nine, or even more. In the ordinary four or five course system, wheat is only grown once in four or five years, manure being given only once, at the commencement of the course, when a crop of turnips being grown, is fed off by sheep; the land thus receiving a rich dressing of sheep's dung (812).

757. With respect to the natural sources of saline matters in the soil, it must be observed that, although their nature and quantity are in a great measure influenced by the composition of the subsoil, and stony substratum; yet it frequently happens that, from want of mixture, the soil is deficient in some of those very substances which the subsoil is rich in; and in consequence the soil becomes considerably improved by mixture, or by spreading over the surface some of the decomposing stones dug from below.

758. We are apt to think that a soil must of necessity contain portions of those substances which enter into the composition of the substratum; but this is by no means always the case. Thus the surface-soil of a chalk district is not unfrequently nearly wholly destitute of calcareous or chalky matters; and soils in such situations are, therefore, often improved by spreading lime over the surface.

759. It is consequently of high importance to know the nature and composition of the subsoil, as well as the surface soil; for by a judicious use of the
former, the soil itself may frequently be very much improved, and probably some saving effected in the quantity of manure required for the land.

760. The system of subsoil ploughing, so successfully practised in many parts of the country, is connected with this subject, and depends mainly upon the advantages derived from admixture of the subsoil with the surface. In this operation, a plough is employed to break up and pulverize the soil to a considerable depth below the surface, without bringing the soil so disturbed up to the top; by this process the permeability of the soil to air is greatly increased, and in consequence, the subsoil is soon brought into a fit state to be mixed with the ordinary soil, by the use of a suitable plough.

761. Part of the benefit derived from this operation is doubtless due to the greater openness conferred upon the soil, which permits a more perfect access of air and moisture, and allows the roots of the growing plants more easily to penetrate through it; but at the same time, advantage is also derived from the greater supply of saline and soluble earthy matters which are thus made available for the use of the plants.

762. A very great variety of different substances is included under the general name of manure. Some of the substances added to the soil act principally in a mechanical way, improving its texture; others are chiefly valuable as sources of carbonic acid and ammonia; whilst many are useful as supplying saline and earthy matters in which the soil may be
deficient. It would be easy to divide all manures into these three classes; but it is more convenient to divide them simply into the organic and the inorganic; because many manures act in all these three ways at once, improving the texture of the soil and supplying carbonic acid, ammonia, and saline matters.

763. The Chinese, who are very economical of their manure, apply it in small quantities at several distinct periods to their plants, when they judge it to be wanted; it may in fact be said that they manure their plants, whilst in Europe it is the custom to manure the soil; this must of necessity be the case, where labor costs less than manure.

764. Organic manures may in general terms be described as consisting of any kind of organic matter in a state of decay or putrefaction. Setting out with the knowledge that organic matter consists of carbon, oxygen, hydrogen, and nitrogen, it is not difficult to understand the changes produced by its decomposition.

765. The complicated changes which organic matters undergo in decomposing are generally divided into four separate classes, namely: Fermentation, or the formation of new compound substances by the partial decomposition of a compound, the change being induced or commenced in consequence of the presence of some other decomposing matter; Putrefaction, or the complete decomposition of organic matter and its conversion into different inorganic compounds, such as water, ammonia, carbonic acid,
sulphuretted hydrogen, &c.; Decay, a slow process of oxidation, almost analogous to combustion, differing from putrefaction in being dependent on the presence of free oxygen or air. This change is always accompanied by the evolution of heat; Mouldering, a change intermediate between putrefaction and decay, taking place in organic matters exposed to the action of water, but not to that of air.

766. Organic manures consist of mixtures of various organic substances in a state of putrefaction or decay. The more changeable substances at once enter into putrefaction, sometimes communicating their own state to those which are less changeable, sometimes only inducing decay in them; thus, for example, urine soon putrefies by itself, and, if mixed with straw, it gradually causes the latter to heat and decay.

767. Whilst describing the different varieties of organic matter, attention has constantly been drawn to the results produced by their putrefaction. It has been repeatedly stated that the substances formed by the putrefaction of organic matter are water, carbonic acid, and ammonia. The nature and rapidity of the change is, however, greatly modified by circumstances.

768. Organic matter containing nitrogen changes far more rapidly than that not containing this element (350). It might have been supposed that under such circumstances the nitrogen would be given off in a free and uncombined state; it is therefore re-
RESULTS OF PUTREFACTION.

markable to find that, at the moment of escaping from one compound, it enters into combination with another element. This fact is important, and deserves a little further consideration, for it is found that, at the moment of separation, substances have a peculiar tendency to enter into fresh combinations.

769. Substances in this state, or whilst being set at liberty by the decomposition of compounds which previously contained them, are said to be in the nascent state. It is always found that substances which have an affinity for each other, but under ordinary circumstances cannot combine, are able to do so when in the nascent state.

770. All attempts have failed to cause the combination of a mixture of hydrogen and nitrogen gases; but when these two substances are in the nascent state they combine readily, and ammonia is formed.

771. The rapidity with which animal substances decay or putrefy depends on moisture, warmth, and air. It has been stated that substances of this kind may be preserved a long time if dry; water is necessary to their putrefaction. It is well known that meat may be preserved fresh a long time if frozen, or kept at a temperature below the freezing point of water, whilst, on the other hand, if kept in a warm place, it soon begins to change.

772. The decomposition of moist animal matter may likewise be accelerated by the mixture of potash, soda, or lime, &c., with it; under these circumstances nitric acid is formed (166), though all the ammonia
formed at the same time is lost (152). It appears that the mixture of these bases assists in the formation of the acid, in consequence of the strong affinity which they have for it, and that therefore they increase the natural tendency which nitrogen has to separate from the carbon, oxygen, and hydrogen with which it is united in organic matter.

773. Whilst animal matters are putrefying, they commonly emit a very offensive smell: now, as neither water, carbonic acid, ammonia, nor nitric acid can cause this, it is evident that some other substance must, at the same time, be formed. The fact is, that the nauseous odor given off under these circumstances is occasioned by the formation of sulphuretted hydrogen (182).

774. Sulphur and hydrogen, though hardly able to combine with each other under common circumstances when brought together, unite readily when the hydrogen is in that peculiar nascent state above adverted to. Thus, when organic substances containing small quantities of sulphates decay, a portion of the hydrogen, whilst set at liberty, combines with some of the sulphur contained in the sulphates, and sulphuretted hydrogen gas is formed.

775. This gas is evolved in considerable quantity during the decomposition of nearly all animal substances, and likewise those vegetables, such as cabbage, &c., which contain both nitrogen and sulphur; though its odor is not unfrequently almost over-
powered by the strong pungent smell of the ammonia evolved at the same time.

776. When such decomposing substances are exposed to the air, the sulphuretted hydrogen gas formed is dissipated, and carried away in the atmosphere; but at the same time a large quantity of the ammonia also formed is lost, and this is of course highly objectionable. The value of these kinds of manure depends in great part on the quantity of nitrogen which they are able to supply to the plants; and everything tending to assist in retaining that substance in the soil, and rendering it available to the growth of plants, is advantageous; whilst, on the other hand, everything increasing its loss, or dissipation in the air, lessens the value of the manure, and renders its application of less service.

777. Animal manures, to a slight degree, modify the mechanical texture of the soil; though from their great proneness to decompose, the effects of this kind which they produce are comparatively transitory. Their principal value consists in the ammonia and carbonic acid, or nitric acid, which they yield, and the earthy and saline compounds which they contain. In addition to this, they are also of value by facilitating the decomposition of vegetable substances employed as manure (767).

778. As perhaps the most useful part of these kinds of manure is the volatile substances formed during their putrefaction, great care should be taken to suffer
as little as possible to escape, and be lost. It is probable that the best method of preserving the greater part of such manure, is to dissolve it in water. The employment of liquid manure is gradually coming more and more into use, and the prejudices against its application are vanishing, as the beneficial results produced by its use become known.

779. Organic manure is of comparatively little value in a chemical point of view, until it has undergone decomposition; plants do not seem able to absorb and appropriate organic matter; hence, in all cases, whether applied fresh or decomposed, it must undergo decay or putrefaction, before it can constitute food for plants. It appears further, that plants cannot absorb solid matter; it is, consequently, also requisite that these matters should be soluble in water, or capable of entering into the organs of plants in a fluid state. When water is added to organic manure in a state of decomposition, putrefaction is assisted, the escape of gases is diminished, and the useful part of the manure obtained in that form best suited to the wants of growing plants.

780. When plants are growing very vigorously, and are abundantly supplied with manure, it appears that they sometimes do absorb a small quantity of organic matter together with the inorganic products of putrefraction. Vegetables forced with abundance of strong animal manure, such as pig's dung, are frequently found to have acquired a bad taste, from the presence of a minute quantity of some
substance, which they have absorbed undecomposed, from the manure.

781. The greatest attention should be paid to the modes of collecting, preserving, and economizing animal manure. There can be no doubt that a very large quantity of manure is constantly wasted under the old systems of using it, and the least consideration will convince any one of this.

782. The food of plants consists principally of certain volatile or gaseous substances, produced, amongst other ways, by the decomposition of organic matter; we add organic matters to the soil, to supply plants with more food than they could otherwise obtain. If, then, previous to using these substances as manure, they are exposed for some time to the air, sun, and rain, a considerable portion of the volatile products of putrefaction must be lost; and yet this is constantly done.

783. Manure should be carefully collected—none should be neglected or suffered to go to waste; it should be preserved in proper receptacles, sheltered from the sun and rain, so that the volatile products of its decay may neither be lost by the heat of the former, nor washed away by the latter; and lastly, by the action of water it should be softened, and as much as possible reduced to a fluid state, so that it may be easily and uniformly applied to the land. The formation of liquid manure-tanks deserves the most serious attention of the farmer.

784. In the application of these kinds of manure,
it has to be remembered that although carbonic acid and ammonia are the principal food of plants, yet that there are limits to the quantity of these substances they can absorb, and that too great a supply of such food is not only useless, but objectionable, and even highly injurious. It is useless to give a plant abundance of carbonic acid and ammonia, if we do not at the same time give it those earthly and saline substances which it requires (554). Even if both are supplied to plants, it is impossible for them to absorb more than a limited quantity. Excess does harm (745).

785. As animal matter is either directly or indirectly formed from vegetable substances, it is natural to expect that animal substances must contain, in addition to the four elements of organic matter (556), the same earthy substances which plants contain, and this is really the fact (595). The phosphate and carbonate of lime of the bones is obtained from plants, and the alkaline and other salts found in the various parts of the animal body are derived from a similar source.

786. The knowledge of this fact necessarily leads to the conclusion that all animal matter must be valuable as manure. The flesh and softer parts of animals, as well as the bones and other solids of the body, are composed of the same substances, both inorganic and organic, as plants; and during their putrefaction yield them up again in a fit state to be absorbed by plants.
787. The vegetable substances which constitute the food of animals contain more earthy and saline matters than animals require, and they are accordingly passed from the body as excrementitious. The food of animals in great part goes to supply the waste occasioned by respiration. In this process, carbonic acid is formed by the oxidation of carbon in the body, by the oxygen of the air; hence, in the air expired from the lungs, it is found that the oxygen is more or less combined with carbon, and converted into carbonic acid (107, 606). The heat evolved by the combination of this carbon with oxygen keeps up the warmth of the body. The waste of organic matter in the body thus occasioned, is supplied by food, the organic part of which supplies that consumed by respiration; but as the greater part of the inorganic substances contained in food are not required for this purpose, the excess is voided in the solid and fluid excrements.

788. There are few things of greater value as manures than these offensive and apparently useless substances, which consist of a mixture of organic and inorganic matters; the former, in consequence of the nitrogen they always contain, ready to decompose, and furnish carbonic acid and ammonia; the latter, those very substances which we require to add to the soil, being the very substances which are removed with crops. In consequence of the volatile nature of the products of their decomposition, every means must be employed to prevent their loss.
789. When animal substances are left for some time exposed to the air, they undergo decomposition, and there at last remains nothing but the more fixed substances which they contained, together with a quantity of salts of ammonia, formed during the decomposition of the easily putrefiable matters. This residue is a valuable manure, though in forming it a very large portion of the ammonia produced is lost by evaporation. The guano or huano of South America, which has of late excited so much attention, is the remains of the excrements of sea-fowl, which has partly undergone this change (813).

790. There are other modes besides that already adverted to (162), whereby the loss of ammonia may be prevented. Both ammonia and its carbonate are volatile, and though by the addition of water it is easy to retard their escape, it cannot be entirely prevented. It has therefore been proposed to add to liquid manure a small quantity of some acid, which shall combine with the ammonia to form a neutral salt, and so prevent further loss from volatility. The trials which have hitherto been made appear perfectly satisfactory, and the only question is—which is the best method of thus neutralizing or fixing it?

791. The simplest method of fixing ammonia is to add to the manure a small quantity of a weak solution of any acid; but we may also fix it by the addition of any salt containing an acid united to a base by a less affinity than the acid has for ammonia;
when this is done, the salt is decomposed, the base is set free, and the acid combines with the ammonia.

792. Ammonia may be fixed by the addition of a small quantity of sulphuric, nitric, muriatic, phosphoric, or any other acid. Sulphuric is, generally speaking, the cheapest, and therefore the most convenient to employ; the quantity to be added of course depends on the quantity of ammonia in the manure; when enough acid has been used, all smell (153, 155) of ammonia disappears. A slight excess of acid does no harm, as it is certain to become neutralized by the bases always present in the soil.

793. It not unfrequently happens that, from local circumstances, a large supply of some other acid liquors may be obtained, more particularly in the vicinity of manufactories; all such substances may be used with advantage. The comparative value of the different salts of ammonia is as yet unknown; perhaps the best acid, if it could be procured cheap enough, would be the phosphoric. Phosphate of ammonia forms a most valuable manure.

794. Various salts have been proposed for the purpose of fixing ammonia; amongst these the best appear to be gypsum, or sulphate of lime (237), and copperas or green vitriol, the sulphate of iron (289). When either of these salts is mixed with solutions containing ammonia in the state of a carbonate, both are decomposed, sulphate of ammonia is formed (159), and carbonate of lime or oxide of iron is left, as an insoluble powder. For several reasons, acid is
preferable to fix ammonia; it is more easily used, and requires far less care and attention in mixing with the solution containing ammonia.

795. Animal substances must have undergone decomposition before they can serve as the food of plants. It has already been stated, that plants appear only able to absorb nitrogen in a state of combination (640); no form of organic matter is suitable to the nourishment of a healthy plant. It is the office of the leaves to convert carbonic acid, ammonia, and water into organic matter, and hence organic matter must be resolved by decomposition into these substances, before it can be absorbed by plants.

796. There are some plants, however, which seem to be exceptions to this rule, and which appear able to feed on organic substances. Certain fungi and parasitical plants, or those which grow upon others, probably feed by directly absorbing organic matter. The small fungi which constitute the various forms of mouldiness are of this description; they flourish in the dark, and grow on any kind of dead organized matter. All plants which have leaves, and require the influence of light, feed on gaseous matter, and never on organic compounds, except during the first period of their growth (719).

797. Liquid manure, consisting chiefly of urine and other waste animal manures diluted with water, is never so valuable in its fresh state as it becomes after a time, when the organic matter is chiefly con-
verted into carbonic acid, water, ammonia, &c. The
time required for this change varies from four to
eight weeks, or even more, according to the season
of the year; putrefaction being much more rapid in
hot than in cold weather.

798. The substances containing nitrogen are those
which first begin to undergo putrefaction, and ac-
cordingly ammonia is one of the earliest products of
the putrefaction of liquid manure; at this time much
of the ammonia is in the caustic state, and hence it
is not so fit for manure as it is subsequently, when
combined with carbonic acid in the completely putre-
fid liquid. In consequence of the great volatility
of caustic ammonia, much of it is lost by evaporation,
in the early stage of putrefaction; it is for this rea-
son that close tanks for liquid manure are so much
recommended, and that the addition of fixers is found
advantageous.

799. Amongst the principal animal substances
employed as manures are urine, and dung of all
kinds, the flesh and blood of dead animals, fat and
oily matters, hair, wool, skin, and, lastly, horns,
hoofs, and bones. These substances are all more or
less valuable, as yielding to the soil a large quantity
of the substances which constitute the food of plants;
they putrefy, and their elements form new combina-
tions with great rapidity. Those which change most
readily, of course yield ammonia and carbonic acid
most rapidly, and these constitute the most powerful
manures: those which decompose more slowly are less powerful, but more lasting in their effects.

800. Urine, dung, and the decomposing carcases of all animals are excellent manure; they are for many purposes considered to be too strong, and means are adopted to diminish their power. These manures evolve, during their rapid decay, a very large quantity of ammonia, carbonic acid, &c., far more, in fact, than plants require or can absorb. This excess is hurtful, and must be prevented.

801. There are two ways by which this may be effected; the one is to mix strong manure with a considerable quantity of some substance far less prone to decomposition, so as to dilute it, or check its putrefaction; the other, and certainly infinitely the worse way, is to expose it to the air for some time, and not to use it until a great part of the ammonia evolved by its decomposition is dissipated, or combined with acids; what remains is then sufficiently mild to be used with safety.

802. In order to preserve as much as possible the valuable parts of these manures, they should be mixed with a certain quantity of vegetable refuse manures, such as sawdust, weeds, &c. This addition tends to check their too rapid decomposition, and prevents the great loss which ensues when manure putrefies too rapidly and becomes hot from the effects of its own decomposition; and at the same time the vegetable matters added are gradually brought into a state which renders them also a most valuable ad-
diction to the soil. Farmyard dung is a mixture of this kind.

803. In mixing weeds with strong animal manures, it must be borne in mind that their seeds are in many cases not destroyed, although all the rest of the plants may be completely decomposed. Rapid and active putrefaction will no doubt destroy the seed also, but this is not desirable; because, when the manure ferments very strongly, it heats, and a considerable portion of volatile matter is sure to be lost. Many seeds have so hard a husk that they will not be at all injured by exposure to mere ordinary putrefaction, and retain therefore their vitality through the decomposition of the dung. Some seeds will even pass uninjured through the stomach of an animal, and not have their vital powers impaired even by the process of digestion.

804. In most cases it is a waste of manure to burn weeds, and whenever it is practicable it is better to form them into dung by rotting them with other substances; but when they are full of seeds, or it is not possible to destroy their vitality by so doing, it is doubtless better economy to burn them than to run the risk of a fresh crop for the next season.

805. As the hedges, banks, and ditches round the manure-yard or mixed heaps are always sure to be well manured, and to get a good deal of the exhalations which are lost during the fermentation of the manure itself, so the weeds which grow in such situations are of course peculiarly luxuriant and abundant.
The farmer should take particular care to destroy
them, and never allow them to come to seed, in such
situations. It is unnecessary to describe the bad ef-
fects produced by the abundant crops of weeds which
too often are left to seed in the immediate vicinity of
the manure-heaps.

806. A very bad custom, too, is often followed of
leaving manure-heaps too long untouched, so that
they become in fact covered with a mass of weeds,
which are left to flower and seed. The labor saved
in not turning over and trimming the heap, is not a
tenth part of that required to remedy the evil done
in thus rendering the land foul.

807. One of the greatest obstacles to good farm-
ing is the practice of keeping wide and foul hedges;
they serve not merely as harbors for vermin, but as
nurseries for weeds. The preservation of game is, in
fact, the bane of the farmer; for his farm is at least
diminished in value by one half, from the various
evils which are thus introduced.

808. Those, however, who advocate the total re-
moval of hedges certainly go too far; for hedges are
useful as shelter, and in other ways. A small and
well-trimmed hedge, if constantly attended to, does
not, after all, require very much labor to keep it in
good order.

809. The extremely offensive odor of night-soil is
a great obstacle to its collection and use as manure.
In its fluid state the expense of conveying it to any
distance is of course great, and accordingly various
plans have been adopted to reduce it to the solid form, both to diminish cost of conveyance, and also to admit of its being used with the drill; in these, however, a portion of the volatile constituents is generally lost, though what remains is undoubtedly a valuable manure.

810. In the manufacture of some of the best of these "disinfected night-soils," charcoal-powder, burnt clay, and other similar porous substances, are used to absorb the gases evolved (162).

811. The most economical mode of using night-soil is, probably, to allow it to putrefy, diluted with water, and mixed with a considerable quantity of vegetable matter, and to employ it as liquid manure. Of course this cannot be done when it is intended subsequently to carry it to a distance.

812. The dung of different animals varies considerably in its value as manure, depending, in great part, on the substances which constitute their food; that of those which live on animal food being of course richer in nitrogen than that of vegetable feeders. The value of those kinds used as manure are in the following order: Pig's dung, night-soil, sheep and rabbit's dung, horse dung, and cow dung; the first being that of most value.

813. In manuring land by feeding off with sheep, not only does the land receive a large quantity of valuable manure from the dung, urine, and perspiration of the animals, but it likewise has its mechanical
texture materially modified by the constant treading of the sheep.

814. Guano, the residue of the putrefaction of the excrement of sea-fowl, consists chiefly of various salts of ammonia, inorganic compounds, and undecomposed organic matters, analogous in nature to horn and wool (818). The salts of ammonia dissolve easily in water, and are at once absorbed by plants, whilst the undecomposed organic matter, gradually undergoing decomposition, continues for some time to yield a regular supply of ammonia. In guano, there exists all the chemical elements of an excellent manure; but of course it does not produce the same mechanical effect on the soil which yard dung does.

815. In using guano as liquid manure, it must be remembered that the solution formed by pouring water over it only contains the ammonia and about one quarter of the phosphates, the rest of the phosphates, and the organic matter, being almost insoluble in water; hence the residue is nearly as valuable a manure as that which is dissolved; and in order to derive the whole benefit from the manure, the insoluble part must, by agitation or other means, be kept suspended in the liquid whilst it is being spread over the ground.

816. Analogous in nature and value to these substances are dead fish, and the refuse matters left in curing and preserving fish; sugar-refiners' waste, which is a mixture of charcoal, blood, and the various
organic matters containing nitrogen, which exist in crude sugar; and wool, soap, an abundant refuse from the wool-mills, consisting of putrid urine, in which the wool is boiled to remove the grease which it naturally contains, and which is consequently rich in ammonia.

817. Strong animal manures are doubly valuable; for not only do they contain abundance of those matters which constitute the food of plants, but they also assist in rendering useful materials which would without them be of far less service. Whether these manures are employed in the liquid or in the solid form, some means should be adopted to retain the ammonia, either by the addition of acids or by some other method. (788).

818. These remarks are far less applicable to animal manures, such as fatty substances, horn, wool, bones, &c., which change but slowly. These substances for the most part act slowly and gradually, and in a very different manner from the softer parts of animals.

819. In general, the strength of animal manures is in proportion to the quantity of nitrogen which they contain; those which contain much of that element decompose rapidly, and evolve a considerable quantity of ammonia, whilst those containing little or no nitrogen change slowly, and, though valuable, and perhaps more lasting in their effects as manures, are far less powerful for the time.

820. Woollen rags, hair, feathers, clippings of horn and skin, greaves, the refuse of the tallow-melters, the
refuse of glue manufactories, and similar matters, form valuable manures; because, though they do not decompose so rapidly as other animal manures, they slowly and gradually decompose, and continue to give out ammonia, &c., for a very long time.

821. The substances of this sort employed as manures are very numerous; including as they do the refuse of all those processes of the arts in which animal matters are used, their value in all cases chiefly depends on their slowly decomposing, and affording a continual source of ammonia, &c.

822. Although oil or fat of all kinds contains no nitrogen, and therefore exhibits very little tendency to change when pure, yet when mixed with earth, the large surface which is exposed to the air assists in its decomposition, particularly if other animal matters also are present; and it accordingly forms a valuable constituent of mixed manures.

823. The dregs of oil, waste blubber, and refuse oils of all sorts, are excellent manures; soap, likewise, in which oil or fat is combined with alkali, is used with advantage whenever it can be obtained at a moderate price. The waste soap of the wool factories has already been mentioned as of great value (816).

824. The water which runs to waste from many manufactories in which soap is employed might be used with great advantage for manure; it often holds in solution a considerable quantity of valuable matter, and might be well employed in the formation of liquid manure.
825. The drainage-water of towns, generally containing, as it does, urine, refuse from the kitchens, soap, &c., is an excellent manure; the most valuable substances which it contains exist in a state of solution; the mud which falls to the bottom when it is left to stand at rest, is comparatively speaking of little value.

826. The last animal manure which it is necessary to speak of is bones, though, as their chief value consists in the inorganic matters which they contain, they might almost be classed amongst the inorganic manures.

827. Bones consist of earthy matters together with a quantity of gelatine and fat; the value of bones as a manure consists in their mechanical effect on the texture of the soil, the earthy substances which they supply (876), the ammonia given out during the decay of the gelatine or glue, and the carbonic acid and hydrogen supplied by the fat (595).

828. The quantity of putrefiable matter in bones is small in proportion to their whole weight, and its decomposition is retarded by the large quantity of earthy matter they contain; hence bones form a good and lasting manure.

829. Boiled bones form a better manure than raw or green bones. This appears to be caused by the removal of fat by boiling; the fat which bone contains retards the decomposition of the gelatine, and probably also renders the phosphate of lime less soluble than it is when the oil is removed.
830. In stating that boiled bones form a better manure than raw or unboiled ones, it is of course evident that the operation of boiling cannot in any way increase the absolute quantity of manure which they can supply, but merely renders them more rapid in their action; as they become less lasting in their effects, in exact proportion to the rapidity of their action when first used.

831. As phosphate of lime is insoluble in pure water, and but slightly soluble in water containing carbonic acid, and as the most valuable component of bone is phosphate of lime, we should be led to expect that they would form a lasting manure, producing a beneficial effect so long as any of the phosphate of lime remains unabsorbed. As the quantity of this substance which plants are able to obtain from bone in one season is comparatively small, it has been proposed to mix with bones some substance which shall render the phosphate of lime more soluble.

832. The cheapest and most convenient substance for this purpose is sulphuric acid or oil of vitriol (179, 245); this forms with bones a compound called "superphosphate of lime," one of the most valuable of the artificial manures yet proposed.

833. Crushed bones, either burnt, or in the raw state (the former, however, being preferable), are mixed in a leaden vessel with sulphuric acid; after a short time the acid is found to have completely decomposed the bones and reduced them to the state of a moist solid, which, however, may be easily rubbed
to powder, and used in any way as manure, either alone, or mixed with dry soil.

834. The effect thus produced by the sulphuric acid is very simple; burnt bones consist of phosphate and carbonate of lime, the latter in a far smaller proportion than the former; when sulphuric acid is poured over burnt bones the carbonate of lime is wholly decomposed, being converted into gypsum or sulphate of lime (245); the phosphate of lime is partly decomposed, the sulphuric acid takes from it the greater part of the lime forming gypsum, whilst the rest of the lime being still combined with the phosphoric acid, constitutes a very sour and easily soluble salt, a biphosphate or superphosphate of lime.

835. As the most valuable part of this manure is soluble in water, it may easily be used as a liquid, being dissolved in water. Whether used dry or in solution, it is most advantageous to use with it some manure containing ammonia, such as yard dung, guano, or any of the strong animal manures.

836. It is frequently the custom to mix a considerable quantity of earth or soil with putrefying animal manure. This to some extent is a good practice; the earth added prevents too rapid putrefaction, and retains some of the products of decomposition; a considerable quantity of ammonia, which would otherwise be lost, is absorbed by the soil (162). The addition of burnt clay or charcoal is useful for a similar reason.

837. The use of lime, on the other hand, is de-
cidedly objectionable; it increases the rapidity of decomposition, and tends to the formation of a portion of nitric acid, but its use causes the loss of a large quantity of ammonia, in consequence of its power of decomposing all the salts of ammonia (152), combining with the acid which they contain, and expelling the ammonia in the state of gas. The addition of lime renders putrefaction a far less noisome process, as the lime absorbs the sulphuretted hydrogen evolved (182), which would otherwise escape into the air.

838. Wood ashes mixed with putrefying animal matters act in a similar manner to lime; the alkali which they generally contain frequently accelerates decomposition, but occasions loss of ammonia.

839. A good deal of what has been said respecting animal manures is equally applicable to those of vegetable origin; but in general the decay of vegetable manures is far less rapid than that of animal substances, and they are more valuable for their mechanical effects, and the carbonic acid and saline matters which they supply; whilst, in addition to these, animal manures yield ammonia, the substance of which is indispensable to the formation of gluten, albumen, fibrin, and caseine, the most important of the constituents of vegetables (602).

840. The use of decaying vegetable matters has already been described whilst speaking of the nature of humus, and the organic constituents of the soil
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(667). A moderate quantity is very desirable, but too much is decidedly hurtful.

841. A soil containing a very large quantity of humus or vegetable matter in a state of decay is always full of carbonic acid; seeds do not germinate well in such a soil; and the excess of carbonic acid is even hurtful to plants themselves (745).

842. The principal vegetable substances employed as manure are straw of all kinds, leaves, sawdust, bran, oilcake, seaweed, and green manures, or crops which are merely sown to be ploughed in, and thus afford food to a second crop, of some more valuable plant.

843. All these manures when mixed with soil slowly decay, and yield carbonic acid and small quantities of saline and earthy matters. They are most advantageously used when employed in combination with some kind of animal manure; this is the case in farm-yard manure.

844. Straw alone decays but slowly, but when mixed with the dung and urine of cattle, it soon begins to change, and in a short time the whole is brought into a state of decomposition (802).

845. In this case, a sort of putrid fermentation is caused; the animal manure decomposes rapidly, and causes a similar change to take place in the vegetable substances with which it is mixed (766); decomposition proceeds rapidly, heat is evolved, and if the bulk of the mixture is large, this action becomes so
energetic that the value of manure is seriously injured, by the high temperature to which it is thus exposed.

846. The decay of some vegetable manures may be facilitated by the addition of lime; for the objection which applies to the mixture of lime with animal manures is not applicable to the ordinary vegetable manures. The latter for the most part contain but little nitrogen, their value principally depending on their mechanical action, and on the formation of carbonic acid.

847. Vegetable manures decay more or less rapidly in proportion to the quantity of nitrogen which they contain; green manures contain a notable quantity of gluten and albumen, and accordingly decompose rapidly, whilst sawdust, which consists principally of woody fibre, and contains hardly any nitrogen, decomposes slowly. Sawdust is, therefore, a most excellent substance to mix with the excrement of animals, and other strong animal manures.

848. Wood sawdust is valuable as manure in proportion to the facility with which it decomposes, and the inorganic matters which it contains; that obtained from young trees decomposes with more facility than the sawdust of old wood.

849. The wood of those trees which contain much resinous matter decays less rapidly than other woods, and is therefore not so valuable as a constituent of mixed manures (802). Those woods which, when burnt, yield a large quantity of ashes rich in alkaline
salts (199), are useful additions in the state of sawdust to manures rich in ammonia.

850. There are a few manures of vegetable origin which contain a considerable quantity of nitrogen, and which consequently approach very near in nature to those formed of animal substances: such, for example, is yeast or barm; this consists of matter rich in nitrogen in a state of incipient putrefaction (366), and is a very strong manure.

851. So also is oilcake, the residue left after expressing the oil from seeds like linseed, which are rich in oil; it contains much gluten and albumen, and is for that reason a strong and valuable manure.

852. Green manures, as they are called, always contain enough nitrogen to insure their own decomposition, though it goes on far slower than does that of animal manures; the value of green manures is shown in the ploughing up of an old pasture; and even more strongly in the case of those crops which are sown, merely to be ploughed in, when they have formed a quantity of organic matter.

853. Some plants decompose with far more readiness than others; for the most part, water plants, both sea and land, decay less easily than land plants; they notwithstanding are useful manures. Seaweed, employed alone, decays but slowly; it is, however, valuable for its mechanical effects, and likewise for the alkali which it contains; it may be used with great advantage together with strong animal manures (802).
854. Green manures, which contain but little nitrogen, may also be used very advantageously together with urine and similar animal substances, which soon bring them into a state of decomposition. A similar effect is also produced by steeping these matters in the ammoniacal liquor of the gas-works (154, 886); the whole of the vegetable matter is soon reduced to a pasty decomposing mass, which is a very good manure.

855. The dry and less changeable kinds of vegetable matter are chiefly valuable for their mechanical effects, the influence which they exert upon evaporation, and the inorganic matters which they contain; they likewise may, however, be brought into a state of decomposition by the addition of animal matters (802).

856. The number of inorganic substances employed as manures is very great, and their use is naturally much dependent on the nature of the soil. In many cases, great benefit is derived from mixing soils; either mixing a portion of the subsoil with the surface-soil (761), or by adding to the soil earth brought from a distance, and possessing a different nature and composition.

857. The earthy matters thus added to the soil can hardly be called manures, although in truth they considerably resemble manures in the mode in which they act. By draining, deep ploughing, mixing, and by the use of mineral manures, either the mechanical
texture of the soil is improved, or substances are added to it in which it was previously deficient.

858. It is almost unnecessary to say anything respecting the importance of draining, and the change which it produces in too retentive soils, and in soils which from their position are naturally wet and swampy. The facts which have previously been stated show that too much moisture in a soil is hurtful, preventing the free circulation of air throughout it, and in several ways interfering with the healthy growth of plants; this is remedied by draining.

859. By draining, and subsoil ploughing, the access of air is facilitated, and the decomposition of silicates and similar stony compounds in the earth is assisted. In mixing, or by adding mineral manures, we increase the quantity of certain inorganic components of the soil.

860. Amongst the modes of fertilizing the soil and augmenting the growth of crops, the subject of irrigation must not be omitted. The effects produced by flooding grass-land are well known; they depend on several distinct causes.

861. Very dry parched land is therefore much improved by irrigation, when from the heat of the climate the greater part of the natural moisture of the soil is evaporated; under such circumstances, the artificial application of water is necessary for the successful cultivation of plants, and accordingly it is a constant practice in tropical countries, in the cultivation of rice and similar crops.
862. Water is of service in temperate climates as a solvent of saline matters, as assisting the decay of organic matters in consequence of the air which it holds in solution, as bringing saline substances which are useful to plants, and as affording a supply of organic matter capable of affording ammonia, nitric acid, &c., by its subsequent decomposition.

863. In flooding land, the saline substances contained throughout the soil, and formed by the decomposing agency of the atmosphere (651), are dissolved, and consequently brought into a condition fit for absorption by growing plants; but quite independent of this, almost all water contains a considerable quantity of inorganic matter (76), which in many cases is a valuable addition.

864. In those cases where drainage-water is allowed to mix with that employed for irrigation, of course the organic matters and saline substances which the former contains, produce their effect in addition to that of the latter. The benefit is produced almost entirely by the liquid, and the substances which it holds in solution; the fine mud which it leaves on the surface of the ground is of comparatively but little value.

865. Road-drift and the scrapings of ditches are often used as manures with very beneficial results: their value of course depends in great part on their composition, and the nature of the soil to which they are applied; the scrapings from roads made of stone,
such as whinstone, are more valuable than those from grit or sandstone.

866. This difference depends entirely on the alkaline matters, &c., which the whinstone contains; but besides this, road scrapings are often rich in animal matters, derived from the dung and urine of cattle. Road-drift is seldom used alone, but forms an excellent addition to more concentrated manures, such as alkaline salts, guano, super-phosphate of lime, &c.

867. The fine mud deposited in stagnant waters, and on the banks of many rivers, is of value as manure, though by no means so valuable a substance as is frequently supposed; its composition varies considerably, and sometimes, particularly in the vicinity of large towns, it contains a considerable quantity of animal matter, phosphoric acid, and ammonia; but, generally speaking, the most valuable part of the manure is carried away by the water (826). Even in those cases when it is rich in animal matters, its weight, and the heavy expense of carrying it to any distance, very seriously interfere with its use.

868. The mode in which many inorganic matters act is very imperfectly understood, and must of necessity remain so until the office performed by saline and earthy substances in the nutrition of plants is explained. It is, however, certain that the addition of an earthy substance to the soil is of no use, if the soil already contains abundance of that particular compound; and consequently that the applicability
of any such manures, is wholly dependent on the nature and composition of the soil.

869. When reading accounts of experiments with various inorganic manures, it must always be borne in mind that on one soil salts of potash, on another nitrate of soda, and on a third phosphate of lime, may be the best manure, because the soil happens to be deficient in alkaline salts or phosphates.

870. Plants almost invariably contain salts of the alkalies, and lime, or magnesia; sometimes combined with organic acids, sometimes with sulphuric, muriatic, or phosphoric acid. Phosphates of lime and magnesia, in particular, are very commonly met with in plants. It will be proper to enumerate briefly the artificial sources of these inorganic substances, and to consider the best method of supplying them to the soil.

871. Lime is a very common ingredient in plants, and is also found in almost all fertile soils; the addition of lime or calcareous matter, therefore, to soils which are destitute of it, or contain but very little, invariably improves them. Lime is added to the soil pure, in the form of quick and slaked lime; as a carbonate in the various forms of chalk, limestone, marl, shell-sand; as a sulphate in gypsum and plaster-stone; and as a phosphate, in bones, native phosphates, and the various organic manures already referred to.

872. When quicklime is spread over the surface of land, other effects are produced besides the mere
addition of carbonate of lime. The quicklime soon absorbs moisture, and crumbles to powder, it then gradually absorbs carbonic acid from the air, and returns to the state of carbonate (235). The chalk which is thus formed is in a state of very fine powder, and becomes more thoroughly mixed with the soil than it could be by ordinary means; but at the same time the quicklime has greatly assisted in promoting the decomposition of inorganic and organic matters in the soil, and probably caused the formation of a small quantity of nitrate of lime, a salt of some value as a manure.

873. Caustic lime possesses the power of gradually effecting the decomposition of earthy and alkaline silicates in the soil, combining with alumina and silica, and setting free the potash which they contain; consequently, one of the most important of the chemical effects produced by the action of lime upon the soil is that of rendering useful a portion of the potash which it previously contained in an insoluble, and therefore useless state.

874. There is great difference in the value of different kinds of lime for this purpose. All limestones which consist merely of carbonate of lime yield pure caustic lime when burnt (119), but many limestones contain a portion of carbonate of magnesia, and these when burnt yield a mixture of caustic lime and magnesia. Such lime is in general objectionable as manure. It is probable that the tendency which magnesia has to remain caustic for a long time, ab-
sorbing carbonic acid but slowly from the air, is the cause of this (248). Limestones which contain much alumina and oxide of iron do not burn into good lime, because the lime and clay partly unite and form a kind of slag.

875. Chalk, shell-sand, and indeed shells themselves, which consist of carbonate of lime and a small quantity of animal matter, are useful additions to all soils which contain but little lime. Quick or caustic lime is especially useful in soils rich in humus. Soils of this kind generally contain a small quantity of acid, which greatly interferes with their fertility; this acid is neutralized by the addition of lime. Both lime, and likewise its carbonate, are valuable additions to soils containing sulphate of iron (289, 663). That salt which is hurtful to vegetation, is decomposed by lime; gypsum is formed, whilst the iron remains as an oxide or carbonate.

876. Sulphate of lime, or gypsum, is likewise a very useful addition to all soils which do not naturally contain it. It is partly useful as supplying lime and sulphuric acid, and partly as serving to fix ammonia from the air (794), and thus yielding plants more of that substance than they could otherwise obtain. Gypsum is sometimes employed after being burnt, and sometimes unburnt; the only use of the burning is that it renders it very easily crushed, the gypsum in fact falling to powder when burnt. This operation is rather roasting than burning, because,
if too strongly heated, it loses its property of again absorbing water.

877. Phosphate of lime is found native as a mineral in Spain and other countries; it certainly forms a most valuable manure for poor soils. As phosphate of lime is found in nearly all plants, all substances containing it are useful as manures (732). The native phosphate of lime has, as yet, hardly come into use at all as a manure, in consequence of the expense of collecting and carrying it to any distance. Phosphate of lime exists abundantly in bones (595), and in smaller quantity in all organic manures, and in the ashes of plants. A minute quantity of phosphate of lime is found in many rocks, and in particular kinds of chalk and marl, which are consequently valuable additions to the soil. The coprolites, and other forms of fossil manure which have been so much talked of, of late, contain a considerable quantity of phosphate of lime; large deposits of these substances exist in various parts of the country.

878. There are no mineral manures which contain phosphate of magnesia; it exists in many soils, and in a good many organic manures. Bran contains a considerable quantity of this salt. Magnesia, in its pure and caustic state, appears to be hurtful to plants, but some of its salts are useful; thus, sulphate of magnesia, or Epsom salts, has been found a valuable manure for potatoes, &c.

879. Ashes of all kinds constitute an important class of manures. They are of value for the char-
coal, lime, phosphoric acid, and alkaline salts which they contain. The ashes of organic substances, such as the ashes of wood and vegetable matters, consist principally of those substances which plants require. Kelp, or the ashes of seaweed, consists of similar saline compounds, generally containing more alkaline salts than the ashes of common land plants. Soap-maker's ash, which consists of the insoluble part of wood-ash, contains a considerable quantity of phosphate of lime, and is consequently a valuable manure (890).

880. The ashes of turf, peat, and weeds, consist of pretty nearly the same substances as the ashes of trees and other plants; they therefore constitute valuable manures. The good effects which have been produced by paring and burning, are in great part due to the charcoal and saline matters which are spread over the soil in the form of ashes. The earth which adheres to the roots and plants burnt in this process acquires considerable power of condensing ammonia from the air (162). This is because burnt clay, or clay which has been strongly heated, has its mechanical properties very greatly altered, and acquires the property of absorbing ammonia in large quantity. When dry burnt clay, which has been exposed to the air for some time, is moistened or breathed upon, it gives out a very perceptible smell of ammonia.

881. The ashes of coal are of less value as manure than most other ashes; they do not contain alkali or
phosphoric acid, but consist of silica, alumina, oxide of iron, and a small quantity of sulphate of lime. Their chief value appears to consist in the charcoal and sulphate of lime which they contain.

882. Although charcoal has been spoken of as undergoing no change whatever under ordinary circumstances (90), and as having no tendency to oxidize or form carbonic acid when exposed to the air, yet in some conditions when exposed to moisture, warmth, and air, the more porous kinds of charcoal do slowly and gradually form carbonic acid; and this action appears to go on more rapidly when charcoal is mixed with humus (672) and other decaying organic matters.

883. Waste charcoal of all kinds is useful as manure, particularly in close and retentive soils; hence it is often the custom to burn or char sawdust, brushwood, and dry vegetable rubbish: a mixture of ashes and charcoal is thus obtained, which is more valuable than that which is left when such matters are entirely burnt to ashes. For various purposes in the arts, such as the manufacture of gunpowder, that charcoal is best which contains the least quantity of inorganic matter, but for manure it is just the reverse (162).

884. In connection with ashes, soot, and the refuse of gas-works may be mentioned, although very different in their nature. Soot consists principally of finely-divided charcoal or carbon, but it contains a considerable quantity of salts of ammonia; hence it
is a powerful and valuable manure. If a little quick-lime is mixed with soot, the salts of ammonia will be decomposed, and the strong pungent smell of ammonia becomes evident. For this reason lime and soot should not be used together as manure: the objection of course does not apply to the use of chalk and soot.

885. Soot contains a good deal of the ammonia formed during the combustion of coal, but, in consequence of its volatility, a considerable portion is lost. In the process for making gas, where coal is roasted in close iron vessels, all the ammonia is saved and is condensed by means of cold water. The gas is made to bubble through a quantity of water: this condenses the ammonia, and constitutes what is termed gas liquor, or the ammoniacal liquor of the gas-works. It is a solution of ammonia, partly caustic, and partly combined with carbonic acid and sulphuretted hydrogen (183).

886. Ammoniacal liquor is a strong and excellent manure, resembling in some respects the manures formed by the decay of animal substances. As obtained from the gas-works, it is almost always too strong to be used as a manure for grass-lands or crops, and must be diluted with four or six times its bulk of water. Gas liquor may also be conveniently used, mixed with green vegetable manures, the decomposition of which it greatly facilitates (854).

887. In consequence of the volatility and caustic nature of free ammonia, it is found advisable for
most purposes to fix the ammonia of gas liquor. This may either be done with an acid, with gypsum, or sulphate of iron (792, 794): a cheap acid like the sulphuric is, generally speaking, the most convenient. Gas liquor resembles putrid urine, inasmuch as it is very rich in ammonia; but it differs from it in not containing the fixed alkaline and earthy salts which that manure does.

888. As the strength of gas liquor mainly depends on the salts of ammonia which it contains, and as the relative quantity of these salts varies considerably, it is difficult to state the exact quantity of any acid necessary to neutralize the ammonia; the average quantity of strong sulphuric acid requisite for this purpose is from 20 to 25 lbs. for every 100 gallons of gas liquor; a little excess of acid hardly ever does any harm, as it immediately becomes neutralized by lime, &c., in the soil.

889. Potash and soda, as has been several times stated, are very commonly found in plants, and consequently are important constituents of manure; they both exist in most soils, though present but in small quantity, and are generally combined with silica and other earthy substances; these compounds are slowly and gradually decomposed by the carbonic acid of the air (651); their decomposition may be artificially assisted by the action of lime (873); the principal salts of the fixed alkalies used as manures are the carbonates, muriates, nitrates, and sulphates.

890. The ashes of some plants contain a very large
quantity of carbonate of potash: in fact, the potash of commerce, the chief source of the alkali, is the ash of trees; whilst kelp, the ash of sea-plants, consists chiefly of carbonate of soda. Both pot-ashes and kelp are good manures in moderate quantity, but their value depends as much on the earthy phosphates which they contain as on their alkaline salts.

891. Muriate of potash or chloride of potassium is sometimes used as a dressing for grass-land. The petre salt of the nitre-refinners contains a large quantity of this salt. It used formerly to be obtained in large quantities by the soap-boilers, who mixed salt with their potash soap; the salt was decomposed and a hard soda soap and chloride of potassium in solution were the results. The soap-maker's spent lees were considered a valuable manure for grass-land, and were believed to destroy moss (524).

892. Chloride of sodium or muriate of soda (common salt), is frequently recommended as a valuable manure; and in many soils, particularly those containing little chlorine, it produces very marked effects, when applied in moderation. It is by many stated that cattle thrive better on plants which have been manured with a small quantity of common salt, than on those not so manured. Common salt is not unfrequently added to dung with a view of facilitating its putrefaction; it is very improbable that any such effect should be produced by salt, which in fact is more likely to retard decomposition.

893. Common salt is sometimes recommended to
be added to liquid manure for the purpose of fixing ammonia, because by particular processes, and by employing very strong solutions, carbonate of ammonia and common salt may be made mutually to decompose each other; under all ordinary circumstances, however, the two salts have no action on each other, and it is quite useless to add common salt to animal manures, as a fixer for ammonia.

894. Salt is of course of no use as a manure on land near the sea-coast, or exposed to winds loaded with the salt of sea spray. On other land it is frequently employed, either alone or mixed with earth and lime; when used in the latter manner, it is common to make a compost, by mixing salt, earth, and lime, some time before it is required, and allowing the mixture to remain some time, sheltered from the rain, the whole being occasionally turned: such a mixture contains a portion of carbonate of soda.

895. When solutions of chloride of calcium (241) and carbonate of soda are mixed together, they are both decomposed, chloride of sodium (common salt) and carbonate of lime (chalk) being produced. From this fact it is evident that, under ordinary circumstances, chalk and common salt cannot decompose each other; nevertheless, when chalk or lime is mixed with salt and moist earth, a considerable quantity of carbonate of soda and muriate of lime are produced, an effect which does not take place if a mixture of the two salts is thrown into a quantity of water.
896. A mixture or compost of lime and salt is a good manure; it contains carbonate of soda and chloride of calcium, both of which are useful to plants. It is possible that salt when applied to land is always more or less decomposed in this manner, and that, according to the extent in which this takes place, salt produces a beneficial effect.

897. Common salt is a necessary addition to the soil in the cultivation of marine plants, or such as naturally grow near the sea-shore. It is also frequently used to kill insects. For all these purposes the most impure salt may be used; the impurities for the most part rather increase the value of the substance as manure (252).

898. Nitrate of potash has long been used as a manure, being found to produce beneficial effects on most soils, in quantities from one to three hundred weight per acre. The exact nature of the chemical effect produced by alkaline nitrates on plants is not accurately known; but as both nitric acid and fixed alkali are separately valuable as manure, or at least are found to assist vegetation, we might reasonably expect that the nitrates should be good manures; it must always be remembered, however, that though they contain both nitrogen and alkali, they cannot well be used alone on ordinary soils, and are best employed in conjunction with some other substance containing phosphoric acid.

899. A great many rich and fertile soils are found
to contain a small quantity of nitrate of potash, soda, or lime, which appears to produce nearly the same effect as the salts of ammonia, rendering vegetation vigorous and dark colored. The way in which these salts are formed will be easily understood when we remember that, whenever substances containing nitrogen decay in the neighborhood of lime or alkaline salts, a portion of nitric acid is formed (166, 837). Under these circumstances, the ammonia which would otherwise be produced is oxidized, and nitric acid and water are formed in place of ammonia; the acid combines with the alkali, and nitrate of potash or soda results. These salts are frequently found in mixtures of decomposing organic manures; they are formed in the same way in the soil itself.

900. The effects produced on different plants by alkaline nitrates are very various; to some they are far more beneficial than others. Some plants, such as the sun-flower, tobacco, lettuce, and many others, always contain more or less of these salts. Others do not contain them, but when supplied with nitrates, are subsequently found to contain the base, without the acid. The soda, potash, or lime is combined with some organic acid, whilst the nitric acid has disappeared. It is probable that in these cases the nitrogen of the acid is assimilated by the plant, or that it assists in the formation of gluten and albumen.

901. Nitrates can have but very little value as manures on the soils which naturally contain salts of
nitric acid, or which, in consequence of the substances they contain, are constantly forming nitrates. On soils neither containing nitrates nor other alkaline salts, they appear to produce very beneficial results. It has been found that wheat manured with alkaline nitrates, contains more gluten and albumen than wheat grown in land not so manured.

902. These remarks apply equally to nitrate of soda and nitrate of potash; at least similar effects are produced by the two salts, as far as regards the increased formation of gluten and albumen. Nitrate of soda contains weight for weight more nitrogen than does the nitrate of potash. It is not known in how far soda is able to replace potash, or whether it is indifferent which of the two alkalies is supplied to plants.

903. Silicates of potash and soda have also been proposed and partially employed as manures. These salts are easily made by melting together sand and carbonate of potash or soda (262); it was supposed that they would be particularly valuable for corn and grass crops, as these plants contain a considerable quantity of silica; but the result of trials hitherto made with these compounds has been such as to render it probable that the silica which they contained produced much less effect than the alkali. It appears that almost all soils contain silica in a soluble form; and consequently that plants are always able to obtain as much silica as they require, from the soil, except in very rare cases.
CHAPTER IX.

COMPOSITION OF PARTICULAR CROPS AND MANURES.

904. Plants differ considerably from each other in chemical composition; not only do various plants consist of different proportions of the proximate elements which compose them, but also the same plant is found to contain variable proportions of these substances, depending on the peculiarities of soil, situation, mode of cultivation, manure, and the weather, or season when they were grown. In the following pages a brief account is given of the average mean composition of some of the principal plants which are cultivated as crops.

905. The first point to be ascertained is the relative proportion of water, organic matter, and inorganic matter in different plants. This is done by very carefully drying a weighed portion; the loss which it sustains indicates the quantity of water. The dry matter is then burnt, the weight of the ashes shows the proportion of inorganic matter. The chemical composition of the organic matter, as well as the inorganic constituents, requires a separate analysis.
906. The average composition of wheat, grain, and straw is—

<table>
<thead>
<tr>
<th></th>
<th>Grain</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter</td>
<td>866</td>
<td>835</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>17</td>
<td>45</td>
</tr>
<tr>
<td>Water</td>
<td>117</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

907. According to Boussingault, the ultimate composition of some wheat which he analyzed was—

<table>
<thead>
<tr>
<th></th>
<th>Grain</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>461</td>
<td>484</td>
</tr>
<tr>
<td>Oxygen</td>
<td>434</td>
<td>389</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>58</td>
<td>52</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>23</td>
<td>3</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>24</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

908. The proportion of bran and flour obtained from wheat varies very considerably. The flour, as it is obtained from the mill, is usually separated by sifting into four parts: namely, fine flour, which amounts to from 70 to 80 per cent.; boxings, from 6 to 10 per cent.; sharps, from 5 to 7 per cent.; and 6 to 8 per cent. of bran. There is usually a loss of rather more than 3 per cent. The bran always contains a considerable quantity of starch, soluble matters, &c.; in the following analysis the term bran is used to denote the insoluble fibrous husk alone.
### Wheat

<table>
<thead>
<tr>
<th></th>
<th>Hermbstädt.</th>
<th>Vauquelin.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>666</td>
<td>402</td>
</tr>
<tr>
<td>Gluten</td>
<td>95</td>
<td>351</td>
</tr>
<tr>
<td>Albümen</td>
<td>7</td>
<td>14</td>
</tr>
<tr>
<td>Sugar</td>
<td>19</td>
<td>14</td>
</tr>
<tr>
<td>Gum</td>
<td>18</td>
<td>16</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Soluble phosphates</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>Bran</td>
<td>140</td>
<td>142</td>
</tr>
<tr>
<td>Water</td>
<td>42</td>
<td>42</td>
</tr>
</tbody>
</table>

1000 1000 1000 1000

909. The inorganic substances contained in wheat are also found to vary greatly in composition and proportion from different sorts of wheat. According to the experiments of Sprengel, 100,000 parts of dry wheat contain 1777 parts of inorganic matter; the same quantity of wheat straw contains 3518 parts of earthy matters. These consist of the following substances:

<table>
<thead>
<tr>
<th></th>
<th>Grain.</th>
<th>Straw.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>225</td>
<td>20</td>
</tr>
<tr>
<td>Soda</td>
<td>240</td>
<td>29</td>
</tr>
<tr>
<td>Lime</td>
<td>96</td>
<td>240</td>
</tr>
<tr>
<td>Magnesia</td>
<td>690</td>
<td>32</td>
</tr>
<tr>
<td>Alumina</td>
<td>26</td>
<td>90</td>
</tr>
<tr>
<td>Silica</td>
<td>400</td>
<td>2870</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>50</td>
<td>37</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>40</td>
<td>170</td>
</tr>
<tr>
<td>Chlorine</td>
<td>10</td>
<td>30</td>
</tr>
</tbody>
</table>

1777 3518
910. According to a more recent analysis by Way, a sample of white wheat from Whitfield farm contained in 100,000 parts, grain 1730, and in the straw 4680 parts of inorganic matter. Hopeton wheat from the same place contained in the grain 1760, in the straw 4160, and in the chaff 10,360 parts of inorganic matter. 10,000 parts of these ashes respectively consisted of—

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Grain.</td>
<td>Grain.</td>
</tr>
<tr>
<td>Straw.</td>
<td>Straw.</td>
</tr>
<tr>
<td>Silica</td>
<td></td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td></td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td></td>
</tr>
<tr>
<td>Lime</td>
<td></td>
</tr>
<tr>
<td>Magnesia</td>
<td></td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td></td>
</tr>
<tr>
<td>Potash</td>
<td></td>
</tr>
<tr>
<td>Soda</td>
<td></td>
</tr>
</tbody>
</table>

911. With such data it is easy roughly to calculate about the quantity of each substance taken off from the soil with a crop of wheat. Suppose, for example, a crop of Hopeton wheat, consisting of 22 cwt. of grain, 30 of straw, and 15 of stubble and roots; the latter, of course, remaining in the soil need not be reckoned. The grain, then, would contain about 43 lbs., and the straw (including 4-cut chaff) nearly 180 lbs. of inorganic matter; the former containing nearly 20 lbs. of phosphoric acid, 14 lbs. of potash, 1 lb. of soda, and nearly 4½ lbs. of magnesia. The inorganic matters in the straw would include about 120 lbs. of silica, 12½ lbs. of phos-
phoric acid, 10 lbs. of sulphuric acid, 6 lbs. of magnesia, 18 lbs. of potash, and 1½ lbs. of soda, removed from each acre.

912. The composition of barley is about—

<table>
<thead>
<tr>
<th></th>
<th>Grain.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter</td>
<td></td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>25</td>
</tr>
<tr>
<td>Water</td>
<td>150</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1000</td>
</tr>
</tbody>
</table>

913. The proportion of the proximate elements of barley, according to the experiments of Hermbstädt, is—

<table>
<thead>
<tr>
<th></th>
<th>No Manure</th>
<th>Manure, Urine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>625</td>
<td>596</td>
</tr>
<tr>
<td>Gluten</td>
<td>29</td>
<td>59</td>
</tr>
<tr>
<td>Albumen</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Sugar</td>
<td>50</td>
<td>44</td>
</tr>
<tr>
<td>Gum</td>
<td>47</td>
<td>44</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Soluble phosphates</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>Husk</td>
<td>136</td>
<td>136</td>
</tr>
<tr>
<td>Water</td>
<td>110</td>
<td>105</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

914. In the process of malting barley (412, 690), a large quantity of starch, which the grain naturally contains, is converted into sugar and gum. The following analyses by Prout show the relative proportions of these substances which he found in the raw and in the malted grain. By Hordein, is meant a peculiar kind of starch, which exists together with
common starch in barley. Hordein is insoluble in hot water, whilst common starch is readily soluble (326).

<table>
<thead>
<tr>
<th></th>
<th>Barley</th>
<th>Malt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>320</td>
<td>560</td>
</tr>
<tr>
<td>Hordein</td>
<td>550</td>
<td>120</td>
</tr>
<tr>
<td>Gluten</td>
<td>30</td>
<td>10</td>
</tr>
<tr>
<td>Sugar</td>
<td>50</td>
<td>150</td>
</tr>
<tr>
<td>Gum</td>
<td>40</td>
<td>150</td>
</tr>
<tr>
<td>Resin</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

915. According to the experiments of Sprengel, 100,000 parts of dry barley contain 2349, and 100,000 parts of straw 5242 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Grain</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>278</td>
<td>180</td>
</tr>
<tr>
<td>Soda</td>
<td>290</td>
<td>48</td>
</tr>
<tr>
<td>Lime</td>
<td>106</td>
<td>554</td>
</tr>
<tr>
<td>Magnesia</td>
<td>180</td>
<td>76</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>trace</td>
<td>14</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>—</td>
<td>20</td>
</tr>
<tr>
<td>Alumina</td>
<td>25</td>
<td>146</td>
</tr>
<tr>
<td>Silica</td>
<td>1182</td>
<td>3856</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>59</td>
<td>118</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>210</td>
<td>160</td>
</tr>
<tr>
<td>Chlorine</td>
<td>19</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>2349</td>
<td>5242</td>
</tr>
</tbody>
</table>

916. According to two analyses by Way, Chevalier barley contains from 225 to 243 parts of inorganic
matter in 10,000 parts of the grain. 10,000 parts of the ash consisted of—

<table>
<thead>
<tr>
<th></th>
<th>I</th>
<th>II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>3273</td>
<td>2360</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>3169</td>
<td>2601</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>79</td>
<td>272</td>
</tr>
<tr>
<td>Lime</td>
<td>148</td>
<td>279</td>
</tr>
<tr>
<td>Magnesia</td>
<td>745</td>
<td>867</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>51</td>
<td>9</td>
</tr>
<tr>
<td>Potash</td>
<td>2077</td>
<td>2743</td>
</tr>
<tr>
<td>Soda</td>
<td>456</td>
<td>5</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>—</td>
<td>860</td>
</tr>
</tbody>
</table>

917. Oats consist on the average of—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter</td>
<td>872</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>28</td>
</tr>
<tr>
<td>Water</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

918. According to Boussingault, the grain of oats is composed of—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>503</td>
</tr>
<tr>
<td>Oxygen</td>
<td>372</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>63</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>22</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

919. The proportion of the proximate elements of oats are (Hermbstädt)—
According to Sprengel, 100,000 parts of dry oats contain 2580, and 100,000 parts of straw 5740 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Grain</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>150</td>
<td>870</td>
</tr>
<tr>
<td>Soda</td>
<td>132</td>
<td>2</td>
</tr>
<tr>
<td>Lime</td>
<td>86</td>
<td>152</td>
</tr>
<tr>
<td>Magnesia</td>
<td>67</td>
<td>22</td>
</tr>
<tr>
<td>Alumina</td>
<td>14</td>
<td>6</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>40</td>
<td>2</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td></td>
<td>2</td>
</tr>
<tr>
<td>Silica</td>
<td>1976</td>
<td>4588</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>35</td>
<td>79</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>70</td>
<td>12</td>
</tr>
<tr>
<td>Chlorine</td>
<td>10</td>
<td>5</td>
</tr>
</tbody>
</table>

2580 5740

According to Way, Hopeton oats contain 227, and potato oats 245 parts of inorganic matter in 10,000. The composition of the ash was in 10,000 parts—
922. Rye consists on the average of—

<table>
<thead>
<tr>
<th>Organic matter</th>
<th>Inorganic matter</th>
</tr>
</thead>
<tbody>
<tr>
<td>889</td>
<td>11</td>
</tr>
<tr>
<td>Water</td>
<td>100</td>
</tr>
<tr>
<td>100</td>
<td></td>
</tr>
</tbody>
</table>

923. According to Boussingault, the grain of rye consists of—

<table>
<thead>
<tr>
<th>Carbon</th>
<th>Oxygen</th>
<th>Hydrogen</th>
<th>Nitrogen</th>
<th>Inorganic matter</th>
</tr>
</thead>
<tbody>
<tr>
<td>463</td>
<td>442</td>
<td>54</td>
<td>17</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1000</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

924. The proportion of the proximate elements of rye is (Hermbstädt)—

<table>
<thead>
<tr>
<th>Starch</th>
<th>Gluten</th>
</tr>
</thead>
<tbody>
<tr>
<td>563</td>
<td>86</td>
</tr>
<tr>
<td>502</td>
<td>120</td>
</tr>
<tr>
<td>649</td>
<td>622</td>
</tr>
</tbody>
</table>

29*
925. According to the experiments of Sprengel, 100,000 parts of the grain of rye contain 1040, and 100,000 parts of straw contain 2793 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Grain</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>532</td>
<td>32</td>
</tr>
<tr>
<td>Soda</td>
<td></td>
<td>11</td>
</tr>
<tr>
<td>Lime</td>
<td>122</td>
<td>178</td>
</tr>
<tr>
<td>Magnesia</td>
<td>44</td>
<td>12</td>
</tr>
<tr>
<td>Alumina</td>
<td>24</td>
<td>25</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>34</td>
<td></td>
</tr>
<tr>
<td>Silica</td>
<td>164</td>
<td>2207</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>23</td>
<td>170</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>46</td>
<td>51</td>
</tr>
<tr>
<td>Chlorine</td>
<td>9</td>
<td>17</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1040</td>
<td>2793</td>
</tr>
</tbody>
</table>

926. According to Way, the grain of rye contains 136, and its straw 313 parts of inorganic matter in 10,000. The composition of the ash from the grain was—
Silica ................................................. 922
Phosphoric acid ...................................... 3992
Sulphuric acid ........................................ 17
Lime .................................................. 261
Magnesia .............................................. 1281
Peroxide of iron ...................................... 104
Potash .................................................. 3383
Soda ................................................... 39

927. The grain of maize, or Indian corn, consists of about—

<table>
<thead>
<tr>
<th>Organic matter</th>
<th>Inorganic matter</th>
</tr>
</thead>
<tbody>
<tr>
<td>857</td>
<td>13</td>
</tr>
</tbody>
</table>

Water .............................................. 130

1000

928. According to the experiments of Payen, dry maize contains—

| Starch          | 712 |
| Gluten          |     |
| Albumen         | 123 |
| Fixed oil       | 90  |
| Gum             | 4   |
| Woody matter    | 59  |
| Inorganic matter| 12  |

1000

929. Sprengel found in 100,000 parts of maize grain 1312 parts of inorganic matter, and in 100,000 parts of maize straw 3985 parts of inorganic matter, consisting of—
RICE.

<table>
<thead>
<tr>
<th></th>
<th>Grain.</th>
<th>Straw.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>200</td>
<td>189</td>
</tr>
<tr>
<td>Soda</td>
<td>250</td>
<td>4</td>
</tr>
<tr>
<td>Lime</td>
<td>35</td>
<td>652</td>
</tr>
<tr>
<td>Magnesia</td>
<td>128</td>
<td>236</td>
</tr>
<tr>
<td>Alumina</td>
<td>16</td>
<td>6</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>traces</td>
<td>4</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td></td>
<td>20</td>
</tr>
<tr>
<td>Silica</td>
<td>434</td>
<td>2708</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>17</td>
<td>106</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>224</td>
<td>54</td>
</tr>
<tr>
<td>Chlorine</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>1312</td>
<td>3985</td>
</tr>
</tbody>
</table>

930. The grain of rice consists of about—

Organic matter ... 856
Inorganic matter  ... 4
Water             ... 140

1000

931. According to the analyses of Braconnot, rice contains—

<table>
<thead>
<tr>
<th></th>
<th>Carolina</th>
<th>Piedmont</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>851</td>
<td>838</td>
</tr>
<tr>
<td>Gluten</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td>Albumen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sugar</td>
<td>3</td>
<td>0\frac{1}{2}</td>
</tr>
<tr>
<td>Gum</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>1</td>
<td>2\frac{1}{2}</td>
</tr>
<tr>
<td>Earthy phosphates</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Husk</td>
<td>48</td>
<td>48</td>
</tr>
<tr>
<td>Water</td>
<td>50</td>
<td>70</td>
</tr>
</tbody>
</table>

1000 1000
932. The more recent experiments of Payen and Boussingault, however, indicate a larger proportion of albumen and gluten as existing in rice; these chemists found 75 in place of 36 of those substances.

933. The dry grain of buckwheat consists of, according to Zenneck—

<table>
<thead>
<tr>
<th>Component</th>
<th>Seed</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>523</td>
<td></td>
</tr>
<tr>
<td>Gluten</td>
<td>107</td>
<td></td>
</tr>
<tr>
<td>Sugar and gum</td>
<td>83</td>
<td></td>
</tr>
<tr>
<td>Fixed oil</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Husk</td>
<td>269</td>
<td></td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td></td>
</tr>
</tbody>
</table>

934. From the experiments of Sprengel, it appears that 100,000 parts of the seed of buckwheat contain 1354 parts of inorganic matter. 100,000 parts of the dry straw contain 3203 parts, consisting of—

<table>
<thead>
<tr>
<th>Component</th>
<th>Seed</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>204</td>
<td>332</td>
</tr>
<tr>
<td>Soda</td>
<td>330</td>
<td>62</td>
</tr>
<tr>
<td>Lime</td>
<td>156</td>
<td>704</td>
</tr>
<tr>
<td>Magnesia</td>
<td>183</td>
<td>1292</td>
</tr>
<tr>
<td>Alumina</td>
<td>26</td>
<td>26</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>8</td>
<td>15</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>44</td>
<td>32</td>
</tr>
<tr>
<td>Silica</td>
<td>114</td>
<td>140</td>
</tr>
<tr>
<td>Chlorine</td>
<td>15</td>
<td>95</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>47</td>
<td>217</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>170</td>
<td>288</td>
</tr>
<tr>
<td></td>
<td>1354</td>
<td>3203</td>
</tr>
</tbody>
</table>
935. The proportion of starch, &c., in seeds which contain much oil is, of course, considerably less than in the grains hitherto described: The following is an analysis of linseed by L. Meyer:—

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>15</td>
</tr>
<tr>
<td>Gluten</td>
<td>29</td>
</tr>
<tr>
<td>Albumen</td>
<td>28</td>
</tr>
<tr>
<td>Azotized matter</td>
<td>151</td>
</tr>
<tr>
<td>Gum</td>
<td>61</td>
</tr>
<tr>
<td>Sugar, &amp;c.</td>
<td>109</td>
</tr>
<tr>
<td>Coloring matter</td>
<td>25</td>
</tr>
<tr>
<td>Resin</td>
<td>25</td>
</tr>
<tr>
<td>Wax</td>
<td>1</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>113</td>
</tr>
<tr>
<td>Husk</td>
<td>443</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>

936. In this analysis the quantity of oil is considerably below the usual average (938). The large quantity of azotized matter which exists in linseed explains the value of oil-cake, both in fattening cattle and as a manure (854). From the experiments of Sprengel, it appears that 100,000 parts of linseed contain 2340 parts of inorganic matter, and 100,000 parts of the stem contain 1456 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th>Component</th>
<th>Seed</th>
<th>Stalks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>438</td>
<td>510</td>
</tr>
<tr>
<td>Soda</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lime</td>
<td>630</td>
<td>230</td>
</tr>
<tr>
<td>Magnesia</td>
<td>234</td>
<td>480</td>
</tr>
<tr>
<td>Alumina</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td><strong>Carried forward</strong></td>
<td><strong>1304</strong></td>
<td><strong>1222</strong></td>
</tr>
</tbody>
</table>
HEMPSEED.

Brought forward 1304 1222
Oxide of iron trace 10
Oxide of mangenese — —
Silica 120 20
Chlorine 12 20
Sulphuric acid 24 66
Phosphoric acid 880 118

2340 1456

937. Hempseed contains, according to Bucholz—

Albumen 247
Gum 90
Sugar 16
Resin 16
Woody fibre 50
Fixed oil 191
Husk 383
Loss in analysis 7

1000

938. The proportion of oil yielded by different seeds varies greatly with the year, climate, soil, &c. The following table shows the average proportion of oil obtained from 100 parts of some of the most important oil seeds:

<table>
<thead>
<tr>
<th>Seed</th>
<th>Oil (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linseed</td>
<td>22</td>
</tr>
<tr>
<td>Poppy</td>
<td>47-50</td>
</tr>
<tr>
<td>Walnut</td>
<td>40-70</td>
</tr>
<tr>
<td>Hemp</td>
<td>25</td>
</tr>
<tr>
<td>Rape</td>
<td>33</td>
</tr>
<tr>
<td>Castor</td>
<td>50-60</td>
</tr>
<tr>
<td>Olives</td>
<td>10</td>
</tr>
<tr>
<td>Sunflower</td>
<td>15</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Seed</th>
<th>Oil (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Almond</td>
<td>30-40</td>
</tr>
<tr>
<td>Beech-nut</td>
<td>15-17</td>
</tr>
<tr>
<td>Colza</td>
<td>39</td>
</tr>
<tr>
<td>Rockcot</td>
<td>18</td>
</tr>
<tr>
<td>White mustard</td>
<td>36</td>
</tr>
<tr>
<td>Black</td>
<td>18</td>
</tr>
<tr>
<td>Earth-nut</td>
<td>45</td>
</tr>
<tr>
<td>Gold of Pleasure</td>
<td>27</td>
</tr>
</tbody>
</table>
939. Common field beans, according to an analysis by Einhof, consist of (347)—

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>501</td>
</tr>
<tr>
<td>Albumen and legumine</td>
<td>117</td>
</tr>
<tr>
<td>Sugar</td>
<td>82</td>
</tr>
<tr>
<td>Gum</td>
<td></td>
</tr>
<tr>
<td>Husk</td>
<td>100</td>
</tr>
<tr>
<td>Water</td>
<td>156</td>
</tr>
<tr>
<td>Salts and loss</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>

940. Kidney beans were found by Braconot to consist of—

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>430</td>
</tr>
<tr>
<td>Legumine</td>
<td>182</td>
</tr>
<tr>
<td>Albumen</td>
<td>54</td>
</tr>
<tr>
<td>Sugar</td>
<td>2</td>
</tr>
<tr>
<td>Gum</td>
<td>15</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>7</td>
</tr>
<tr>
<td>Husk</td>
<td>70</td>
</tr>
<tr>
<td>Water</td>
<td>230</td>
</tr>
<tr>
<td>Salts and loss</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>

941. According to the experiments of Sprengel, 100,000 parts of common beans contain 2136, and 100,000 parts of bean straw contain 3121 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th>Component</th>
<th>Seed</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>415</td>
<td>1656</td>
</tr>
<tr>
<td>Soda</td>
<td>816</td>
<td>50</td>
</tr>
<tr>
<td>Carried forward</td>
<td>1231</td>
<td>1706</td>
</tr>
</tbody>
</table>
942. Field beans, according to Way, contain 237 in the seed, and 497 parts of inorganic matter in 10,000 of the straw. These ashes, analyzed, were found to contain in 10,000 parts—

<table>
<thead>
<tr>
<th>Substance</th>
<th>Beans</th>
<th>Bean straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>42</td>
<td>261</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>2872</td>
<td>49</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>305</td>
<td>140</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>342</td>
<td>2532</td>
</tr>
<tr>
<td>Lime</td>
<td>520</td>
<td>1985</td>
</tr>
<tr>
<td>Magnesia</td>
<td>690</td>
<td>253</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>trace</td>
<td>61</td>
</tr>
<tr>
<td>Potash</td>
<td>5172</td>
<td>3285</td>
</tr>
<tr>
<td>Soda</td>
<td>54</td>
<td>277</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>trace</td>
<td>1154</td>
</tr>
</tbody>
</table>

943. The composition of ripe peas, according to Braconnot, is—
944. Peas consist of about 854 parts organic, 26 inorganic matter, and 120 parts water. According to Sprengel, 100,000 parts of peas contain 2464, and 100,000 parts of pea straw contain 4971 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Seed.</th>
<th>Straw.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>810</td>
<td>235</td>
</tr>
<tr>
<td>Soda</td>
<td>739</td>
<td>—</td>
</tr>
<tr>
<td>Lime</td>
<td>58</td>
<td>2730</td>
</tr>
<tr>
<td>Magnesia</td>
<td>136</td>
<td>342</td>
</tr>
<tr>
<td>Alumina</td>
<td>20</td>
<td>60</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>—</td>
<td>7</td>
</tr>
<tr>
<td>Silica</td>
<td>410</td>
<td>996</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>53</td>
<td>337</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>190</td>
<td>240</td>
</tr>
<tr>
<td>Chlorine</td>
<td>33</td>
<td>4</td>
</tr>
</tbody>
</table>

2464 4971

945. White peas, according to Way, contain 197, and the straw 752 parts of inorganic matter in 10,000. The composition of 10,000 parts of this ash was—
<table>
<thead>
<tr>
<th></th>
<th>Pea</th>
<th>Pea Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>176</td>
<td>253</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>2420</td>
<td>131</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>470</td>
<td>185</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>318</td>
<td>3033</td>
</tr>
<tr>
<td>Lime</td>
<td>697</td>
<td>4692</td>
</tr>
<tr>
<td>Magnesia</td>
<td>666</td>
<td>836</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>25</td>
<td>114</td>
</tr>
<tr>
<td>Potash</td>
<td>4402</td>
<td>387</td>
</tr>
<tr>
<td>Soda</td>
<td>—</td>
<td>186</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>823</td>
<td>176</td>
</tr>
</tbody>
</table>

946. Lentils, according to an analysis of Einhof, consist of—

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td></td>
<td>282</td>
</tr>
<tr>
<td>Albumen and legumine</td>
<td></td>
<td>331</td>
</tr>
<tr>
<td>Gum</td>
<td></td>
<td>51</td>
</tr>
<tr>
<td>Sugar</td>
<td></td>
<td>27</td>
</tr>
<tr>
<td>Husk</td>
<td></td>
<td>161</td>
</tr>
<tr>
<td>Water</td>
<td></td>
<td>140</td>
</tr>
<tr>
<td>Salts and loss</td>
<td></td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

947. In the preceding analysis of lentils, no mention is made of fixed oil. The following analysis, quoted by Boussingault, is therefore probably more correct:

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td></td>
<td>400</td>
</tr>
<tr>
<td>Legumine</td>
<td></td>
<td>220</td>
</tr>
<tr>
<td>Gum</td>
<td></td>
<td>70</td>
</tr>
<tr>
<td>Sugar</td>
<td></td>
<td>15</td>
</tr>
<tr>
<td>Fibrous matter</td>
<td></td>
<td>120</td>
</tr>
<tr>
<td>Fixed oil</td>
<td></td>
<td>25</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td></td>
<td>25</td>
</tr>
<tr>
<td>Water and loss</td>
<td></td>
<td>125</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>
According to Sprengel, 100,000 parts of lentils contain 1528, and 100,000 parts of lentil straw contain 3899 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Seed</th>
<th>Straw</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>736</td>
<td>420</td>
</tr>
<tr>
<td>Soda</td>
<td>118</td>
<td>33</td>
</tr>
<tr>
<td>Lime</td>
<td>57</td>
<td>2040</td>
</tr>
<tr>
<td>Magnesia</td>
<td>85</td>
<td>119</td>
</tr>
<tr>
<td>Alumina</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>52</td>
<td>34</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>traces</td>
<td>traces</td>
</tr>
<tr>
<td>Silica</td>
<td>180</td>
<td>686</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>94</td>
<td>38</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>140</td>
<td>480</td>
</tr>
<tr>
<td>Chlorine</td>
<td>54</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>1528</td>
<td>3899</td>
</tr>
</tbody>
</table>

Common vetch, according to an analysis by Crome, is stated to contain—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>26</td>
</tr>
<tr>
<td>Albumen</td>
<td>19</td>
</tr>
<tr>
<td>Gum, &amp;c.</td>
<td>76</td>
</tr>
<tr>
<td>Woody fibre</td>
<td>104</td>
</tr>
<tr>
<td>Water</td>
<td>775</td>
</tr>
<tr>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

According to Sprengel, 100,000 parts of vetch seed contain 5101, and 100,000 parts of vetch straw contain 2290 parts of inorganic matter consisting of—
A great number of different analyses of potatoes have been published, as the varieties of the tuber are found to contain very different proportions of starch, azotized matter, &c. The ultimate composition of dry potato is (Boussingault):

<table>
<thead>
<tr>
<th>Component</th>
<th>Grain.</th>
<th>Straw.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>897</td>
<td>1810</td>
</tr>
<tr>
<td>Soda</td>
<td>622</td>
<td>52</td>
</tr>
<tr>
<td>Lime</td>
<td>160</td>
<td>1955</td>
</tr>
<tr>
<td>Magnesia</td>
<td>142</td>
<td>324</td>
</tr>
<tr>
<td>Alumina</td>
<td>22</td>
<td>15</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td>Silica</td>
<td>200</td>
<td>442</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>50</td>
<td>122</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>140</td>
<td>280</td>
</tr>
<tr>
<td>Chlorine</td>
<td>43</td>
<td>84</td>
</tr>
</tbody>
</table>

952. According to the experiments of Einhof and Lampadius, potatoes contain:
953. The composition of good potatoes varies from about 10 to 25 starch, 3 to 8 fibre, 2 to 4 gum, 1 to 2 azotized matters (albumen, &c.) and 70 to 80 parts of water per cent.

954. According to Sprengel, 100,000 parts of dry potato tuber contain 2653 parts; and 100,000 of potato haulm, 4786 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Tubers</th>
<th>Haulm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>1291</td>
<td>138</td>
</tr>
<tr>
<td>Soda</td>
<td>748</td>
<td>0 (?)</td>
</tr>
<tr>
<td>Lime</td>
<td>106</td>
<td>2928</td>
</tr>
<tr>
<td>Magnesia</td>
<td>104</td>
<td>488</td>
</tr>
<tr>
<td>Alumina</td>
<td>16</td>
<td>52</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>9</td>
<td>58</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>trace</td>
<td>44</td>
</tr>
<tr>
<td>Silica</td>
<td>27</td>
<td>801</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>174</td>
<td>245</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>128</td>
<td>32</td>
</tr>
<tr>
<td>Chlorine</td>
<td>50</td>
<td>0 (?)</td>
</tr>
</tbody>
</table>

2653 4786

955. The batatas or sweet potato of the West India Islands, contains, according to O. Henry—
<table>
<thead>
<tr>
<th>Component</th>
<th>Tuber</th>
<th>Stem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>133</td>
<td>355</td>
</tr>
<tr>
<td>Albumen</td>
<td>9</td>
<td>10</td>
</tr>
<tr>
<td>Sugar</td>
<td>33</td>
<td>148</td>
</tr>
<tr>
<td>Cellular matter</td>
<td>68</td>
<td>4</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>11</td>
<td>28</td>
</tr>
<tr>
<td>Malic acid, and salts</td>
<td>14</td>
<td>4</td>
</tr>
<tr>
<td>Water</td>
<td>732</td>
<td>770</td>
</tr>
</tbody>
</table>

956. The ultimate composition of the tubers and stem of the Jerusalem artichoke, according to Boussingault, is—

957. The tubers, according to the analysis of Braconnot, contain a peculiar variety of starch, to which the name of Inulin is given. The tubers were found to contain—

<table>
<thead>
<tr>
<th>Component</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch (Inulin)</td>
<td>30</td>
</tr>
<tr>
<td>Albumen</td>
<td>10</td>
</tr>
<tr>
<td>Sugar, uncrystallizable</td>
<td>148</td>
</tr>
<tr>
<td>Gum</td>
<td>12</td>
</tr>
<tr>
<td>Fixed oil</td>
<td>1</td>
</tr>
<tr>
<td>Woody fibre</td>
<td>12</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>17</td>
</tr>
<tr>
<td>Water</td>
<td>770</td>
</tr>
</tbody>
</table>

1000
958. Jerusalem artichoke, according to Way, contains in the tubers 179 parts, in the stem 194, and in the leaves 1500 parts of inorganic matter per 10,000. These three ashes contained respectively in 10,000 parts—

<table>
<thead>
<tr>
<th></th>
<th>Tuber.</th>
<th>Stems.</th>
<th>Leaves.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>150</td>
<td>151</td>
<td>1725</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>1699</td>
<td>297</td>
<td>61</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>377</td>
<td>323</td>
<td>221</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>1180</td>
<td>2540</td>
<td>2431</td>
</tr>
<tr>
<td>Lime</td>
<td>334</td>
<td>2031</td>
<td>4015</td>
</tr>
<tr>
<td>Magnesia</td>
<td>130</td>
<td>191</td>
<td>195</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>45</td>
<td>88</td>
<td>114</td>
</tr>
<tr>
<td>Potash</td>
<td>5589</td>
<td>3840</td>
<td>681</td>
</tr>
<tr>
<td>Soda</td>
<td>—</td>
<td>69</td>
<td>372</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>—</td>
<td>468</td>
<td>182</td>
</tr>
<tr>
<td>Chloride of potassium</td>
<td>488</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

959. The bulbs of the Oxalis crenata were found by Payen to contain—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>25</td>
</tr>
<tr>
<td>Albumen</td>
<td>15</td>
</tr>
<tr>
<td>Gum, &amp;c.</td>
<td>55</td>
</tr>
<tr>
<td>Woody fibre</td>
<td>44</td>
</tr>
<tr>
<td>Water</td>
<td>861</td>
</tr>
</tbody>
</table>

1000

960. Common cabbage consists of about 62 parts of organic matter, 7 of inorganic matter, and 931 parts of water. According to Sprengel, 100,000 parts of dry cabbage contain 7546 parts of inorganic matter, consisting of—
961. The composition of turnips and Swedes is very similar; they consist of the same proximate elements united in nearly the same proportions. The following table shows the composition of three varieties according to Hermbstädt:

<table>
<thead>
<tr>
<th></th>
<th>Swede</th>
<th>White</th>
<th>Cabbage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch and fibre</td>
<td>53</td>
<td>72</td>
<td>60</td>
</tr>
<tr>
<td>Albumen</td>
<td>20</td>
<td>25</td>
<td>25</td>
</tr>
<tr>
<td>Sugar</td>
<td>90</td>
<td>80</td>
<td>90</td>
</tr>
<tr>
<td>Gum</td>
<td>30</td>
<td>25</td>
<td>35</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Water</td>
<td>800</td>
<td>790</td>
<td>780</td>
</tr>
<tr>
<td>Loss</td>
<td>2</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1000</td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

962. Dry turnip, according to Boussingault, consists of:

- Carbon: 429
- Oxygen: 423
- Hydrogen: 55
- Nitrogen: 17
- Inorganic matter: 76

**Total:** 1000
963. According to the same chemist, 100,000 parts of dry turnip contain 6226 parts of inorganic matter, consisting of—

Potash .......................... 2610
Soda ................................ 317
Lime ................................ 844
Magnesia .......................... 333
Alumina and oxide of iron .......... 93
Silica ................................ 496
Sulphuric Acid ................. 844
Phosphoric Acid .................. 465
Chlorine .......................... 224

6226

964. According to two more recent analyses of Way, Skirving's Swede contains in the bulb 76 and 88 parts inorganic matter, and 161 and 195 in the top. The composition of these ashes was per 10,000 parts—

<table>
<thead>
<tr>
<th></th>
<th>I. Top.</th>
<th>I. Bulb.</th>
<th>II. Top.</th>
<th>II. Bulb.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>411</td>
<td>163</td>
<td>114</td>
<td>173</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>654</td>
<td>1251</td>
<td>621</td>
<td>1017</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>650</td>
<td>1126</td>
<td>1220</td>
<td>1553</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>616</td>
<td>954</td>
<td>1297</td>
<td>1196</td>
</tr>
<tr>
<td>Lime</td>
<td>2399</td>
<td>1136</td>
<td>3038</td>
<td>1433</td>
</tr>
<tr>
<td>Magnesia</td>
<td>292</td>
<td>244</td>
<td>318</td>
<td>327</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>190</td>
<td>28</td>
<td>66</td>
<td>51</td>
</tr>
<tr>
<td>Potash</td>
<td>2063</td>
<td>3616</td>
<td>2079</td>
<td>2688</td>
</tr>
<tr>
<td>Soda</td>
<td>—</td>
<td>499</td>
<td>—</td>
<td>1331</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>1769</td>
<td>977</td>
<td>1031</td>
<td>219</td>
</tr>
<tr>
<td>Chloride of potassium</td>
<td>977</td>
<td>—</td>
<td>209</td>
<td>—</td>
</tr>
</tbody>
</table>

965. The same chemist found in the green-topped white turnip, in the tops 182, and in the bulb 50
parts of inorganic matter. The analyses of these ashes gave—

<table>
<thead>
<tr>
<th></th>
<th>Top</th>
<th>Bulb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>205</td>
<td>96</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>315</td>
<td>765</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>783</td>
<td>1286</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>1464</td>
<td>1482</td>
</tr>
<tr>
<td>Lime</td>
<td>2873</td>
<td>673</td>
</tr>
<tr>
<td>Magnesia</td>
<td>285</td>
<td>226</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>80</td>
<td>66</td>
</tr>
<tr>
<td>Potash</td>
<td>1268</td>
<td>4856</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>1067</td>
<td>544</td>
</tr>
<tr>
<td>Chloride of potassium</td>
<td>1656</td>
<td></td>
</tr>
</tbody>
</table>

966. Common beet and mangel-wurzel, also, are very similar in composition. The following table shows the composition of four varieties:

<table>
<thead>
<tr>
<th></th>
<th>Organic matter</th>
<th>Inorganic matter</th>
<th>Water</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red mangel-wurzel</td>
<td>88</td>
<td>11</td>
<td>901</td>
</tr>
<tr>
<td>Castelnaudary beet</td>
<td>139</td>
<td>11</td>
<td>850</td>
</tr>
<tr>
<td>White sugar beet</td>
<td>120</td>
<td>11</td>
<td>869</td>
</tr>
<tr>
<td>Bassano beet</td>
<td>115</td>
<td>12</td>
<td>873</td>
</tr>
</tbody>
</table>

967. According to Boussingault, dried field beet consists of—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>428</td>
</tr>
<tr>
<td>Oxygen</td>
<td>434</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>58</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>17</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>63</td>
</tr>
</tbody>
</table>

968. According to the experiments of Sprengel, 100,000 parts of dry beet root contain 5986 parts of
inorganic matter, and 100,000 parts of the dry leaves contain 15,439 parts, consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Root.</th>
<th>Leaves.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>1481</td>
<td>5600</td>
</tr>
<tr>
<td>Soda</td>
<td>3178</td>
<td>3290</td>
</tr>
<tr>
<td>Lime</td>
<td>285</td>
<td>2316</td>
</tr>
<tr>
<td>Magnesia</td>
<td>139</td>
<td>839</td>
</tr>
<tr>
<td>Alumina</td>
<td>20</td>
<td>130</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>58</td>
<td>50</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>50</td>
<td>60</td>
</tr>
<tr>
<td>Silica</td>
<td>105</td>
<td>425</td>
</tr>
<tr>
<td>Chlorine</td>
<td>380</td>
<td>1064</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>123</td>
<td>975</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>167</td>
<td>690</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5986</td>
<td>15,439</td>
</tr>
</tbody>
</table>

969. The yellow globe mangel-wurzel contains, according to Way, 102 parts of inorganic matter in the bulb, and 140 in its top. The long red contains in the bulb 64, and in the top 179 parts of inorganic matter in 10,000. These ashes analyzed were found to contain respectively, in 10,000 parts—

<table>
<thead>
<tr>
<th></th>
<th>Yellow globe.</th>
<th>Long red.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>235</td>
<td>222</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>589</td>
<td>449</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>654</td>
<td>368</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>692</td>
<td>1814</td>
</tr>
<tr>
<td>Lime</td>
<td>872</td>
<td>178</td>
</tr>
<tr>
<td>Magnesia</td>
<td>984</td>
<td>175</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>146</td>
<td>74</td>
</tr>
<tr>
<td>Potash</td>
<td>834</td>
<td>2354</td>
</tr>
<tr>
<td>Soda</td>
<td>1221</td>
<td>1908</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>3766</td>
<td>2454</td>
</tr>
</tbody>
</table>
970. The carrot, according to Hermbstädt and Einhof, contains—

<table>
<thead>
<tr>
<th></th>
<th>Einhof.</th>
<th>Hermbstädt.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>3</td>
<td>—</td>
</tr>
<tr>
<td>Fibre</td>
<td>46</td>
<td>90</td>
</tr>
<tr>
<td>Albumen</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Gum</td>
<td>—</td>
<td>18</td>
</tr>
<tr>
<td>Sugar</td>
<td>81</td>
<td>78</td>
</tr>
<tr>
<td>Volatile oil</td>
<td>—</td>
<td>4</td>
</tr>
<tr>
<td>Water</td>
<td>861</td>
<td>799</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

971. According to Sprengel, 100,000 parts of dry carrot contain 5090 parts of inorganic matter. 100,000 parts of the leaves contain 10,420 parts consisting of—

<table>
<thead>
<tr>
<th></th>
<th>Root.</th>
<th>Leaves.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>2718</td>
<td>3236</td>
</tr>
<tr>
<td>Soda</td>
<td>709</td>
<td>921</td>
</tr>
<tr>
<td>Lime</td>
<td>505</td>
<td>5050</td>
</tr>
<tr>
<td>Magnesia</td>
<td>205</td>
<td>398</td>
</tr>
<tr>
<td>Alumina</td>
<td>30</td>
<td>78</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>25</td>
<td>15</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>46</td>
<td>0</td>
</tr>
<tr>
<td>Silica</td>
<td>105</td>
<td>454</td>
</tr>
<tr>
<td>Chlorine</td>
<td>54</td>
<td>223</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>208</td>
<td>1082</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>395</td>
<td>963</td>
</tr>
<tr>
<td></td>
<td>5090</td>
<td>10,420</td>
</tr>
</tbody>
</table>

972. Two samples of white Belgian carrot, according to Way, contained respectively, in the root 77, and in the top 532 parts of inorganic matter; and 82 31
parts in the root, and 420 parts of inorganic matter in 10,000 parts of the top. These ashes contained—

<table>
<thead>
<tr>
<th></th>
<th>I. Top.</th>
<th>I. Root.</th>
<th>II. Top.</th>
<th>II. Root.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>739</td>
<td>76</td>
<td>183</td>
<td>110</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>255</td>
<td>837</td>
<td>112</td>
<td>786</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>263</td>
<td>634</td>
<td>457</td>
<td>695</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>1629</td>
<td>1515</td>
<td>2275</td>
<td>1772</td>
</tr>
<tr>
<td>Lime</td>
<td>3498</td>
<td>976</td>
<td>2950</td>
<td>826</td>
</tr>
<tr>
<td>Magnesia</td>
<td>250</td>
<td>378</td>
<td>303</td>
<td>320</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>406</td>
<td>74</td>
<td>90</td>
<td>166</td>
</tr>
<tr>
<td>Potash</td>
<td>728</td>
<td>3755</td>
<td>753</td>
<td>2800</td>
</tr>
<tr>
<td>Soda</td>
<td>946</td>
<td>1263</td>
<td>1069</td>
<td>1753</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>877</td>
<td>491</td>
<td>1714</td>
<td>765</td>
</tr>
</tbody>
</table>

973. The parsnip consists of—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter</td>
<td>195</td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>12</td>
</tr>
<tr>
<td>Water</td>
<td>793</td>
</tr>
</tbody>
</table>

974. According to Crome, the parsnip contains—

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>18</td>
</tr>
<tr>
<td>Albumen</td>
<td>21</td>
</tr>
<tr>
<td>Gum</td>
<td>61</td>
</tr>
<tr>
<td>Sugar</td>
<td>55</td>
</tr>
<tr>
<td>Fibre</td>
<td>51</td>
</tr>
<tr>
<td>Volatile oil</td>
<td>trace</td>
</tr>
<tr>
<td>Water</td>
<td>794</td>
</tr>
</tbody>
</table>

975. From the experiments of Sprengel, it appears that 100,000 parts of dry parsnip roots contain 4643 parts of inorganic matter. 100,000 parts of the dry leaves contain 15,661 parts consisting of—
CLOVER.

<table>
<thead>
<tr>
<th></th>
<th>Root.</th>
<th>Leaves.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>2310</td>
<td>3207</td>
</tr>
<tr>
<td>Soda</td>
<td>780</td>
<td>2448</td>
</tr>
<tr>
<td>Lime</td>
<td>520</td>
<td>4160</td>
</tr>
<tr>
<td>Magnesia</td>
<td>300</td>
<td>473</td>
</tr>
<tr>
<td>Alumina</td>
<td>26</td>
<td>132</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silica</td>
<td>180</td>
<td>1400</td>
</tr>
<tr>
<td>Chlorine</td>
<td>198</td>
<td>950</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>213</td>
<td>1198</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>111</td>
<td>1784</td>
</tr>
</tbody>
</table>

4643 15,661

976. Clover, according to the analysis of Boussingault, consists of—

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>474</td>
<td></td>
</tr>
<tr>
<td>Oxygen</td>
<td>378</td>
<td></td>
</tr>
<tr>
<td>Hydrogen</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>Nitrogen</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Inorganic matter</td>
<td>77</td>
<td>1000</td>
</tr>
</tbody>
</table>

977. The composition of white and red clover, as given by Crome, is—

<table>
<thead>
<tr>
<th></th>
<th>White.</th>
<th>Red.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>10</td>
<td>14</td>
</tr>
<tr>
<td>Albumen</td>
<td>15</td>
<td>20</td>
</tr>
<tr>
<td>Gum</td>
<td>34</td>
<td>35</td>
</tr>
<tr>
<td>Sugar</td>
<td>15</td>
<td>21</td>
</tr>
<tr>
<td>Woody fibre</td>
<td>115</td>
<td>139</td>
</tr>
<tr>
<td>Wax and resin</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Earthy matter</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>Water</td>
<td>800</td>
<td>760</td>
</tr>
</tbody>
</table>

1000 1000
978. According to Sprengel, 100,000 parts of white clover in the fresh state contain 1735 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th>Inorganic Matter</th>
<th>White Clover</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>590</td>
</tr>
<tr>
<td>Soda</td>
<td>110</td>
</tr>
<tr>
<td>Lime</td>
<td>446</td>
</tr>
<tr>
<td>Magnesia</td>
<td>58</td>
</tr>
<tr>
<td>Alumina</td>
<td>36</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>12</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td></td>
</tr>
<tr>
<td>Silica</td>
<td>280</td>
</tr>
<tr>
<td>Chlorine</td>
<td>40</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>67</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>96</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1735</strong></td>
</tr>
</tbody>
</table>

979. According to Way, 10,000 red clover contains 695, and white clover 765 parts of inorganic matter. These ashes analyzed were found to contain—

<table>
<thead>
<tr>
<th>Inorganic Matter</th>
<th>Red Clover</th>
<th>White Clover</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>334</td>
<td>368</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>635</td>
<td>1153</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>418</td>
<td>721</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>1693</td>
<td>1803</td>
</tr>
<tr>
<td>Lime</td>
<td>3539</td>
<td>2641</td>
</tr>
<tr>
<td>Magnesia</td>
<td>1122</td>
<td>815</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>97</td>
<td>196</td>
</tr>
<tr>
<td>Potash</td>
<td>1485</td>
<td>1433</td>
</tr>
<tr>
<td>Soda</td>
<td>140</td>
<td>372</td>
</tr>
<tr>
<td>Chloride of potassium</td>
<td>236</td>
<td>494</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>296</td>
<td></td>
</tr>
</tbody>
</table>
980. The composition of Lucern, according to Crome, is—

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>22</td>
</tr>
<tr>
<td>Albumen</td>
<td>19</td>
</tr>
<tr>
<td>Gum</td>
<td>44</td>
</tr>
<tr>
<td>Sugar</td>
<td>8</td>
</tr>
<tr>
<td>Woody fibre</td>
<td>143</td>
</tr>
<tr>
<td>Wax and resin</td>
<td>6</td>
</tr>
<tr>
<td>Earthy matter</td>
<td>8</td>
</tr>
<tr>
<td>Water</td>
<td>750</td>
</tr>
</tbody>
</table>

Total 1000

981. According to Sprengel, 100,000 parts of fresh Lucern contain 2580 parts of inorganic matter, consisting of—

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>362</td>
</tr>
<tr>
<td>Soda</td>
<td>166</td>
</tr>
<tr>
<td>Lime</td>
<td>1304</td>
</tr>
<tr>
<td>Magnesia</td>
<td>94</td>
</tr>
<tr>
<td>Alumina</td>
<td>8</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>7</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>—</td>
</tr>
<tr>
<td>Silica</td>
<td>90</td>
</tr>
<tr>
<td>Chlorine</td>
<td>86</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>109</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>353</td>
</tr>
</tbody>
</table>

Total 2580

982. According to Sprengel, 100,000 parts of fresh saintfoin or esparcette contain 1671 parts of inorganic matter consisting of—

31*
<table>
<thead>
<tr>
<th></th>
<th>In flower</th>
<th>In seed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>322</td>
<td>349</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>935</td>
<td>797</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>328</td>
<td>233</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>1520</td>
<td>1736</td>
</tr>
<tr>
<td>Lime</td>
<td>2430</td>
<td>2967</td>
</tr>
<tr>
<td>Magnesia</td>
<td>503</td>
<td>459</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>61</td>
<td>58</td>
</tr>
<tr>
<td>Potash</td>
<td>3190</td>
<td>2961</td>
</tr>
<tr>
<td>Soda</td>
<td>—</td>
<td>125</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>78</td>
<td>312</td>
</tr>
<tr>
<td>Chloride of potassium</td>
<td>624</td>
<td>—</td>
</tr>
</tbody>
</table>

983. Saintfoin in flower, according to Way, contains 565 parts, and, in seed, 570 parts, of inorganic matter. These ashes consisted of—

984. Hay. According to the experiments of Boussingault, 10,000 parts of good meadow hay contain 547 parts of inorganic matter, consisting of—
Potash 130
Soda 10
Lime 107
Magnesia 43
Oxide of iron 5
Silica 189
Sulphuric acid 16
Phosphoric acid 32
Chlorine 15

547

985. 10,000 parts of Italian rye-grass in flower, according to Way, contain 616 parts, and, in seed, 556 parts of inorganic matter. These ashes contained in 10,000 parts—

<table>
<thead>
<tr>
<th></th>
<th>In flower</th>
<th>In seed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>5918</td>
<td>6062</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>634</td>
<td>632</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>282</td>
<td>131</td>
</tr>
<tr>
<td>Lime</td>
<td>995</td>
<td>1229</td>
</tr>
<tr>
<td>Magnesia</td>
<td>223</td>
<td>264</td>
</tr>
<tr>
<td>Peroxide of iron</td>
<td>78</td>
<td>30</td>
</tr>
<tr>
<td>Potash</td>
<td>1245</td>
<td>1077</td>
</tr>
<tr>
<td>Soda</td>
<td>398</td>
<td>13</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>227</td>
<td>558</td>
</tr>
</tbody>
</table>

986. As has been several times stated, the exact relative proportions of these different constituents of plants vary very considerably, depending on soil, climate, manure, &c. Thus, for example, mangel-wurzel root contains from 9 to 11 parts of inorganic matter in every 1000 parts; or, abstracting the water, from 87 to 150 parts in the dry root; hence it is easy
to see that a crop of 25 tons may in truth take more inorganic matter out of the land than one of 30 tons; and that the mere weight of the crop is no proof of the extent to which it has exhausted the soil.

987. This effect is quite independent of the mere quantity of water in the crop, because the same result is shown when we ascertain the relation which exists between the inorganic matter and the dry organic matter which the plants contain; in different crops of mangel-wurzel it is found that the quantity of dry organic matter which is formed for every 100 parts of inorganic matter absorbed from the soil, varies from 568 to 1045.

988. It is consequently evident that, in order to raise the most profitable crops, we must endeavor to obtain the largest possible quantity of useful vegetable matter, with the smallest relative proportion of inorganic matter. This is a subject of the greatest practical importance, and one well worthy of the attention of scientific agriculturists.

989. Having now briefly stated the average composition of some of the most important plants cultivated as crops, it may be useful to enter shortly into the composition of the most common substances employed as manure, in order more fully to show the relation which exists between plants and the manures employed in their cultivation. Of the simple animal manures the most important are dung, urine, and bones.

990. The dung of sheep (according to Zierl) con-
sists of 291 parts dry organic matter, 30 parts inorganic matter, and 679 parts water. The earthy matters consist of about half silica, a quarter carbonate and phosphate of lime, and the remainder alkaline salts. The urine of the sheep consists of 28 parts organic matter, 12 parts inorganic matter, and 960 parts water, containing a portion of ammonia; because even in fresh urine a portion of the azotized matter which it contains, is generally beginning to decompose and form ammonia.

991. Fresh cow dung (according to Boussingault) consists of 124 parts dry organic matter, 17 parts inorganic matter, and 859 parts water. The inorganic matter contains two-thirds of silica, nearly one-third of earthy phosphates, and a small quantity of alkaline salts. Cows' urine consists of 53 parts dry organic matter, 21 parts inorganic matter, and 926 parts water (Sprengel).

992. The composition of cows' urine, and the nature of the changes which it undergoes when suffered to putrefy, either alone or diluted with water, are shown in the following table (Sprengel):

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Alone.</td>
<td>With water.</td>
</tr>
<tr>
<td>Urea</td>
<td>4000</td>
<td>1000</td>
<td>600</td>
</tr>
<tr>
<td>Albumen</td>
<td>10</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Mucus</td>
<td>190</td>
<td>40</td>
<td>30</td>
</tr>
<tr>
<td>Benzoic acid</td>
<td>90</td>
<td>250</td>
<td>120</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>516</td>
<td>500</td>
<td>500</td>
</tr>
<tr>
<td>Carried forward</td>
<td>4806</td>
<td>1790</td>
<td>1250</td>
</tr>
</tbody>
</table>
### Horse Dung—Pig Dung.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Alone.</td>
<td>With water.</td>
</tr>
<tr>
<td>Brought forward</td>
<td>4806</td>
<td>1790</td>
<td>1250</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>5256</td>
<td>165</td>
<td>1533</td>
</tr>
<tr>
<td>Ammonia</td>
<td>205</td>
<td>487</td>
<td>1622</td>
</tr>
<tr>
<td>Potash</td>
<td>664</td>
<td>664</td>
<td>664</td>
</tr>
<tr>
<td>Soda</td>
<td>554</td>
<td>554</td>
<td>554</td>
</tr>
<tr>
<td>Silica</td>
<td>36</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td>Alumina</td>
<td>2</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>4</td>
<td>1</td>
<td>—</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Lime</td>
<td>65</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Magnesia</td>
<td>36</td>
<td>22</td>
<td>30</td>
</tr>
<tr>
<td>Chlorine</td>
<td>272</td>
<td>272</td>
<td>272</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>405</td>
<td>388</td>
<td>332</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>70</td>
<td>26</td>
<td>46</td>
</tr>
<tr>
<td>Acetic acid, or vinegar</td>
<td>—</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>Sulphuretted hydrogen</td>
<td>—</td>
<td>1'</td>
<td>30</td>
</tr>
<tr>
<td>Insoluble earthy phosphates and carbonates</td>
<td>—</td>
<td>180</td>
<td>150</td>
</tr>
<tr>
<td>Water</td>
<td>92,624</td>
<td>95,442</td>
<td>93,481</td>
</tr>
</tbody>
</table>

100,000 100,000 100,000

993. Fresh horse dung consists of 284 parts dry organic matter, 18 parts inorganic matter, and 698 parts water. Of the inorganic matter about one-ninth is carbonate and phosphate of lime, one-twelfth alkaline salts, and the remainder silica (Zierl). Horses' urine consists of 27 parts dry organic matter, 33 parts inorganic matter, and 940 parts water.

994. Fresh pigs' dung, consisting of the excrement and urine together, contains 93 parts dry organic matter, 87 parts inorganic matter, and 820
parts water. Pigs' urine contains 56 parts dry organic matter, 18 parts inorganic matter, and 926 parts water (Sprengel). The inorganic matters consist chiefly of alkaline salts.

995. Human excrement (according to Berzelius) contains 227 parts dry organic matter, 100 parts inorganic matter, and 733 parts water. Its constituents are—

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Parts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albumen</td>
<td>9</td>
</tr>
<tr>
<td>Extractive</td>
<td>27</td>
</tr>
<tr>
<td>Mucus, fat, resin</td>
<td>140</td>
</tr>
<tr>
<td>Bile</td>
<td>9</td>
</tr>
<tr>
<td>Vegetable remains</td>
<td>70</td>
</tr>
<tr>
<td>Soluble salts</td>
<td>12</td>
</tr>
<tr>
<td>Water</td>
<td>733</td>
</tr>
<tr>
<td></td>
<td>1000</td>
</tr>
</tbody>
</table>

996. The inorganic matter contained in 1000 parts consequently weighs 150, and contains 100 parts earthy phosphates, 12 parts carbonate of soda, 8 parts sulphate and phosphate of soda, and sulphate of potash.

997. Human urine (according to Berzelius) consists of 49 parts dry organic matter, 7 parts salts of ammonia, 11 parts inorganic matter, and 933 parts water. Its composition in detail is—

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Parts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urea</td>
<td>3010</td>
</tr>
<tr>
<td>Mucus</td>
<td>32</td>
</tr>
<tr>
<td>Lactic acid, lactate of ammonia, animal</td>
<td>1714</td>
</tr>
<tr>
<td>matter insoluble in alcohol</td>
<td></td>
</tr>
<tr>
<td>Carried forward</td>
<td>4756</td>
</tr>
</tbody>
</table>
BONES.

<table>
<thead>
<tr>
<th>Substance</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brought forward</td>
<td>4756</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>445</td>
</tr>
<tr>
<td>Sulphate of potash</td>
<td>371</td>
</tr>
<tr>
<td>Sulphate of soda</td>
<td>316</td>
</tr>
<tr>
<td>Phosphate of soda</td>
<td>294</td>
</tr>
<tr>
<td>Phosphate of ammonia</td>
<td>165</td>
</tr>
<tr>
<td>Muriate of ammonia</td>
<td>150</td>
</tr>
<tr>
<td>Earthy phosphates</td>
<td>100</td>
</tr>
<tr>
<td>Uric acid</td>
<td>100</td>
</tr>
<tr>
<td>Silica</td>
<td>3</td>
</tr>
<tr>
<td>Water</td>
<td>93,300</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100,000</strong></td>
</tr>
</tbody>
</table>

998. The composition of bones varies considerably in the proportion of the constituents, though all bones consist chiefly of phosphate and carbonate of lime, together with gelatine and a portion of fat. Sheep's bones consist of about—

<table>
<thead>
<tr>
<th>Substance</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate of lime</td>
<td>560</td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>130</td>
</tr>
<tr>
<td>Organic matter</td>
<td>310</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>

999. The bones of oxen and cows (according to Berzelius) contain:—

<table>
<thead>
<tr>
<th>Substance</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate of lime</td>
<td>573</td>
</tr>
<tr>
<td>Phosphate of magnesia</td>
<td>20</td>
</tr>
<tr>
<td>Soda and chloride of sodium</td>
<td>35</td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>39</td>
</tr>
<tr>
<td>Fluoride of calcium</td>
<td>trace</td>
</tr>
<tr>
<td>Organic matter</td>
<td>333</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>

1000. The bones of horses contain nearly the
same proportion of phosphate of lime. They consist of:

<table>
<thead>
<tr>
<th>Component</th>
<th>Fresh.</th>
<th>Dry.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate of lime</td>
<td>546</td>
<td></td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>113</td>
<td></td>
</tr>
<tr>
<td>Salts</td>
<td>19</td>
<td></td>
</tr>
<tr>
<td>Organic matter</td>
<td>322</td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1000</strong></td>
<td></td>
</tr>
</tbody>
</table>

1001. The bones of pigs contain 520 parts of phosphate of lime, 10 parts carbonate of lime, and 470 parts organic matter and water; those of calves, 540 phosphate of lime, and 460 animal matter and water. The bones of fishes, generally speaking, contain less earthy matter than those of land animals. The bones of the cod-fish contain 480 phosphate of lime, 55 carbonate of lime, 20 phosphate of magnesia, 6 salts of soda, and 440 organic matter and water.

1002. The composition of farmyard dung, and all mixtures of animal and vegetable manures, is very variable. The following analysis of Boussingault may be considered as expressing very nearly the average composition of good yard dung:

<table>
<thead>
<tr>
<th>Component</th>
<th>Fresh.</th>
<th>Dry.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>74</td>
<td>358</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>9</td>
<td>42</td>
</tr>
<tr>
<td>Oxygen</td>
<td>53</td>
<td>258</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>4</td>
<td>20</td>
</tr>
<tr>
<td>Inorganic</td>
<td>67</td>
<td>322</td>
</tr>
<tr>
<td>Water</td>
<td>793</td>
<td>—</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>1000</strong></td>
<td><strong>1000</strong></td>
</tr>
</tbody>
</table>
1003. The composition of the inorganic part of such manure, like that of the organic, varies considerably. The following shows the composition of 10,000 parts of the inorganic matter in yard dung (Richardson):

<table>
<thead>
<tr>
<th>Substance</th>
<th>Parts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>322</td>
</tr>
<tr>
<td>Soda</td>
<td>273</td>
</tr>
<tr>
<td>Lime</td>
<td>34</td>
</tr>
<tr>
<td>Magnesia</td>
<td>26</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>327</td>
</tr>
<tr>
<td>Chlorine</td>
<td>315</td>
</tr>
<tr>
<td>Soluble silica</td>
<td>2705</td>
</tr>
<tr>
<td>Phosphate of lime</td>
<td>711</td>
</tr>
<tr>
<td>Phosphate of magnesia</td>
<td>226</td>
</tr>
<tr>
<td>Phosphate of iron</td>
<td>468</td>
</tr>
<tr>
<td>Phosphate of manganese</td>
<td>trace</td>
</tr>
<tr>
<td>Phosphate of alumina (?)</td>
<td>trace</td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>934</td>
</tr>
<tr>
<td>Carbonate of magnesia</td>
<td>163</td>
</tr>
<tr>
<td>Sand</td>
<td>3099</td>
</tr>
<tr>
<td>Alkali and loss</td>
<td>397</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>10,000</td>
</tr>
</tbody>
</table>

1004. The putrefied dung of birds has of late years been much employed as manure, under the name of guano. The composition of this substance varies greatly, according to the extent to which decomposition has proceeded, and the degree to which it has been exposed to the atmosphere. The following table shows three different analyses of guano:
### GUANO.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Urate of ammonia</td>
<td>32</td>
<td>122</td>
<td>147</td>
</tr>
<tr>
<td>Oxalate of ammonia</td>
<td>134</td>
<td>177</td>
<td>32</td>
</tr>
<tr>
<td>Oxalate of lime</td>
<td>164</td>
<td>13</td>
<td>10</td>
</tr>
<tr>
<td>Phosphate of ammonia</td>
<td>64</td>
<td>60</td>
<td>143</td>
</tr>
<tr>
<td>Phosphate of ammonia and magnesia</td>
<td>42</td>
<td>116</td>
<td>45</td>
</tr>
<tr>
<td>Phosphate of lime</td>
<td>100</td>
<td>202</td>
<td>220</td>
</tr>
<tr>
<td>Muriate of ammonia</td>
<td>65</td>
<td>22</td>
<td>30</td>
</tr>
<tr>
<td>Chloride of sodium</td>
<td>-1</td>
<td>4</td>
<td>-</td>
</tr>
<tr>
<td>Carbonate of ammonia</td>
<td>-</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>-</td>
<td>.16</td>
<td>-</td>
</tr>
<tr>
<td>Sulphate of potash</td>
<td>42</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>Sulphate of soda</td>
<td>11</td>
<td>49</td>
<td>-</td>
</tr>
<tr>
<td>Sulphate of ammonia</td>
<td>-</td>
<td>-</td>
<td>20</td>
</tr>
<tr>
<td>Phosphate of soda</td>
<td>53</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Humate of ammonia</td>
<td>-</td>
<td>11</td>
<td>-</td>
</tr>
<tr>
<td>Wax and resin</td>
<td>6</td>
<td>7</td>
<td>-</td>
</tr>
<tr>
<td>Sand. Insoluble residue</td>
<td>58</td>
<td>17</td>
<td>12</td>
</tr>
<tr>
<td>Alumina</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water</td>
<td>227</td>
<td>43</td>
<td>85</td>
</tr>
<tr>
<td>Organic matter</td>
<td>1000</td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

1005. The value of guano as manure depends on the ammonia which it contains, or the presence of matter capable of forming ammonia by its slow putrefaction, and on the presence of phosphoric acid. No manure is more subject to fraudulent adulteration than guano. One of the most common modes of doing this is by the addition of earth or brickdust. The presence of these substances may be roughly ascertained in guano by burning a portion. The
ashes left by pure guano are white, or nearly so; while, if soil or brickdust has been added, the oxide of iron, which they always contain, will give the ashes more or less a red-brown color.

1006. The composition of wood-ashes varies very considerably, according to the wood from which it is obtained. The composition of three kinds of wood-ashes, as given by Sprengel, is:

<table>
<thead>
<tr>
<th></th>
<th>Oak</th>
<th>Beech</th>
<th>Scotch fir.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>162</td>
<td>221</td>
<td>22</td>
</tr>
<tr>
<td>Soda</td>
<td>67</td>
<td>33</td>
<td>22</td>
</tr>
<tr>
<td>Lime</td>
<td>174</td>
<td>250</td>
<td>232</td>
</tr>
<tr>
<td>Magnesia</td>
<td>14</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Alumina</td>
<td></td>
<td></td>
<td>23</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>81</td>
<td>38</td>
<td>170</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>38</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silica</td>
<td>269</td>
<td>55</td>
<td>66</td>
</tr>
<tr>
<td>Chlorine</td>
<td>24</td>
<td>19</td>
<td>23</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>34</td>
<td>77</td>
<td>22</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>19</td>
<td>56</td>
<td>28</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>155</td>
<td>140</td>
<td>365</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1000</td>
<td>1000</td>
<td>1000</td>
</tr>
</tbody>
</table>

1007. In the process for making potash or pearl-ash (199), the greater part of the soluble salts contained in the wood-ashes is washed out; the remainder is a valuable manure, and often contains a considerable quantity of phosphoric acid. The refuse ashes from soap factories consist of the same matters. The following table exhibits the composition of three kinds of “lixiviated ashes,” according to Berthier:
### PEAT-ASHES.

<table>
<thead>
<tr>
<th></th>
<th>Oak</th>
<th>Beech</th>
<th>Scotch fir.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lime</td>
<td>548</td>
<td>426</td>
<td>423</td>
</tr>
<tr>
<td>Magnesia</td>
<td>6</td>
<td>70</td>
<td>105</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>—</td>
<td>15</td>
<td>1</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>—</td>
<td>45</td>
<td>4</td>
</tr>
<tr>
<td>Silica</td>
<td>38</td>
<td>58</td>
<td>46</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>8</td>
<td>57</td>
<td>10</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>396</td>
<td>329</td>
<td>360</td>
</tr>
<tr>
<td>Charcoal, &amp;c.</td>
<td>4</td>
<td>—</td>
<td>51</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Best</th>
<th>Middling</th>
<th>Inferior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potash</td>
<td>2</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Soda</td>
<td>10</td>
<td>39</td>
<td>4</td>
</tr>
<tr>
<td>Lime</td>
<td>136</td>
<td>86</td>
<td>61</td>
</tr>
<tr>
<td>Magnesia</td>
<td>49</td>
<td>16</td>
<td>39</td>
</tr>
<tr>
<td>Alumina</td>
<td>45</td>
<td>35</td>
<td>41</td>
</tr>
<tr>
<td>Oxide of iron</td>
<td>66</td>
<td>54</td>
<td>41</td>
</tr>
<tr>
<td>Oxide of manganese</td>
<td>10</td>
<td>43</td>
<td>2</td>
</tr>
<tr>
<td>Silica</td>
<td>471</td>
<td>559</td>
<td>704</td>
</tr>
<tr>
<td>Chlorine</td>
<td>12</td>
<td>30</td>
<td>5</td>
</tr>
<tr>
<td>Sulphuric acid</td>
<td>72</td>
<td>64</td>
<td>34</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>20</td>
<td>8</td>
<td>13</td>
</tr>
<tr>
<td>Carbonic acid</td>
<td>41</td>
<td>64</td>
<td>55</td>
</tr>
<tr>
<td>Charcoal</td>
<td>66</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

|               | 1000  | 1000    | 1000     |

1008. The ashes of peat, as might be expected, contain less alkaline salts than those of wood; they usually contain a considerable quantity of gypsum (237). The following table shows the composition of three sorts of Dutch ashes analyzed by Sprengel:—

1009. Kelp, the ashes of sea-weeds (218, 891) re-
KELP.

seems wood-ashes in containing a large quantity of alkaline salts; but it contains little or no phosphoric acid. The following analyses are by Dr. Ure:

<table>
<thead>
<tr>
<th></th>
<th>Rona.</th>
<th>Heisker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonate of soda</td>
<td>55</td>
<td>85</td>
</tr>
<tr>
<td>Sulphuret of sodium</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulphate of soda</td>
<td>190</td>
<td>80</td>
</tr>
<tr>
<td>Chlorides of sodium and potassium</td>
<td>375</td>
<td>365</td>
</tr>
<tr>
<td>Carbonate of lime</td>
<td>100</td>
<td>240</td>
</tr>
<tr>
<td>Sulphate of lime</td>
<td>95</td>
<td>-</td>
</tr>
<tr>
<td>Alumina and oxide of iron</td>
<td>100</td>
<td>90</td>
</tr>
<tr>
<td>Silica</td>
<td></td>
<td>80</td>
</tr>
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