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The practical methods of organic chemistry

Ludwig
Gattermann,
William Bush ...



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**THE PRACTICAL METHODS OF
ORGANIC CHEMISTRY**



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THE PRACTICAL METHODS
OF
ORGANIC CHEMISTRY

BY
LUDWIG GATTERMANN, Ph.D.
PROFESSOR IN THE UNIVERSITY OF FREIBURG

WITH NUMEROUS ILLUSTRATIONS

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AUTHORISED TRANSLATION
THE THIRD AMERICAN FROM THE ELEVENTH GERMAN EDITION

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**Tym czarownym rękom, których
dotknięcie pokój pozynosi.**

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TRANSLATOR'S PREFACE

THE success of Professor Gattermann's book in the original has warranted its reproduction in English. The translation is intended for those students of chemistry who have not yet become sufficiently familiar with scientific German to be able to read it accurately without constant reference to a dictionary. To such students this translation is offered, in the hope that it will increase their interest in the science without causing a corresponding decrease in their efforts to acquire a knowledge of German, which is indispensable to every well-trained chemist.

My grateful acknowledgments are due to my colleague, Dr. H. M. Ullmann, for many valuable suggestions, and to Professor Gattermann for his courtesy in pointing out several inaccuracies in the German edition.

WILLIAM B. SCHOBBER.

SOUTH BETHLEHEM, PENNSYLVANIA,
April, 1896.

PREFACE

THE present book has resulted primarily from the private needs of the author. If one is obliged to initiate a large number of students at the same time into organic laboratory work, it is frequently impossible, even with the best intentions, to direct the attention of each individual to the innumerable details of laboratory methods. In order that students, even in the absence of the instructor, can gain the assistance necessary for the carrying out of the common operations, a General Part, dealing with crystallisation, distillation, drying, analytical operations, etc., is given before the special directions for Preparations. In the composition of this General Part, it has been considered of more value to describe the most important operations in such a way that the beginner may be able to carry out the directions independently, rather than to give as fully as possible the numerous modifications of individual operations. In the Special Part, to each preparation are added general observations, which relate to the character and general significance of the reaction carried out in practice; and the result follows, that the student during the period given to laboratory work, becomes familiar with the most varied theoretical knowledge, which, acquired under these conditions adheres more firmly, as is well known, than if that knowledge were obtained exclusively from a purely theoretical

book. And so the author hopes that his book, along with the excellent "Introductions" of E. Fischer and Levy, may here and there win some friends.

For the assistance given by his colleagues, in pointing out deficiencies of his work, the author will always be grateful.

GATTERMANN.

HEIDELBERG, August, 1894.

PREFACE TO THE SECOND AMERICAN EDITION

IN the preparation of this new edition advantage has been taken of the opportunity offered to correct a number of errors in the first edition, and to make the text a reproduction of the fourth German edition of Professor Gattermann's book. In many cases the laboratory directions have been improved, a number of new illustrations have been added, and the Special Part now includes methods for the preparation of glycol, dimethylcyclohexenone, s-xilenol, phenylhydroxylamine, nitrosobenzene, p-tolyl aldehyde (Gattermann-Koch synthesis), salicylic aldehyde (Reimer and Tiemann's oxyaldehyde synthesis), cuprous chloride, the decomposition of inactive mandelic acid into its active constituents, and a zinc dust determination. The preparations of acetylene and acetylene tetrabromide have been omitted.

WILLIAM B. SCHOBBER.

SOUTH BETHLEHEM, PENNSYLVANIA,
May, 1901.

PREFACE TO THE THIRD AMERICAN EDITION

DURING my absence from the University on sick leave, my colleague, Professor Vahan S. Babasinian, has undertaken the entire labor and responsibility, — including the translation, editing, and reading the proofs, — involved in the preparation of the new edition of this book. To him, therefore, is due all the credit for the satisfactory results presented in the following pages.

WILLIAM B. SCHOBBER.

THIS new edition is a reproduction of the eleventh German edition of Professor Gattermann's book. A number of errors in the second edition have been corrected, four new illustrations have been added, improved methods for the preparation of ethylene, glycol and carbon monoxide have been outlined, Dennstedt's Method of Analysis and Grignard's Reaction have been fully described. This edition contains also discussions of the theoretical grounds upon which rest the processes of distillation with steam, salting out, separation by extraction and esterification.

VAHAN S. BABASINIAN.

SOUTH BETHLEHEM, PENNSYLVANIA,
September, 1913.

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THE PRACTICAL METHODS OF ORGANIC
CHEMISTRY

GENERAL PART



THE compounds directly obtained by means of chemical reactions are, only in rare cases, pure; they must therefore be subjected to a process of purification before they can be further utilised. For this purpose the operations most frequently employed are:

1. CRYSTALLISATION.
2. SUBLIMATION.
3. DISTILLATION.

CRYSTALLISATION

Methods of Crystallisation.—The crude solid product obtained directly as the result of a reaction is generally amorphous or not well crystallised. In order to obtain the compound in uniform, well-defined crystals, as well as to separate it from impurities like filter-fibres, inorganic substances, by-products, etc., it is dissolved, usually with the aid of heat, in a proper solvent, filtered from the impurities remaining undissolved, and allowed to cool gradually. The dissolved compound then separates out in a crystallised form, while the dissolved impurities are retained by the mother-liquor. (*Crystallisation by Cooling.*) Many compounds are so easily soluble in all solvents, even at the ordinary temperature, that they do not separate from their solutions on mere cooling. In this case, in order to obtain crystals, a portion of the solvent must be allowed to evaporate. (*Crystallisation by Evaporation.*)

Solvents.— As solvents for organic compounds, the following substances are principally used :

CLASS I. Water,
Alcohol,
Ether,
Ligroin (Petroleum Ether),
Glacial Acetic Acid,
Benzene.

Also mixtures of these :

CLASS II. Water + Alcohol,
Water + Glacial Acetic Acid,
Ether + Ligroin,
Benzene + Ligroin.

Less frequently used than these are : hydrochloric acid, carbon disulphide, acetone, chloroform, ethyl acetate, methyl alcohol, amyl alcohol, toluene, xylene, solvent naphtha, etc.

But rarely used are : pyridine, naphthalene, phenol, nitrobenzene, aniline, and others.

Choice of the Solvent.— The choice of a suitable solvent is often of great influence upon the success of an experiment, in that a solid compound does not assume a completely characteristic appearance until it is uniformly crystallised. In order to find the most appropriate solvent, preliminary experiments are made in the following manner : successive small portions of the finely pulverised substance (a few milligrammes will suffice) are treated in *small* test-tubes, with small quantities of the solvents of Class I. If solution takes place at the ordinary temperature, or on gentle heating, the solvent in question is, provisionally, left out of consideration. The remaining portions are heated to boiling, until, after the addition of more of the solvent if necessary, solution takes place. The tubes are now cooled by contact with cold water, and an observation will show in which tube crystals have separated in the largest quantity. At times crystallisation does not occur on mere cooling ; in this case the walls of the vessel

are rubbed with a sharp-angled glass rod, or the solution is "seeded," *i.e.* a small crystal of the crude product is placed in the solution; by this means, crystallisation is frequently induced. If the individual solvents of Class I. are shown to be unsuitable, experiments are made with the mixtures, — Class II. Compounds which are easily soluble in alcohol or glacial acetic acid, and which consequently do not separate out on cooling, are, as a rule, difficultly soluble in water. In order to determine whether a separation of crystals will take place on cooling, the hot solutions in the pure solvents are treated with more or less water, according to the conditions. Substances easily soluble in ether, benzene, toluene, etc., often dissolve in ligroin with difficulty. Hence mixtures of these solvents can be frequently utilised with advantage, in the manner just described. If these experiments have shown several solvents to be suitable, the portions under examination are again heated until solution takes place, and this time are allowed to cool slowly. That solvent from which the best crystals separate in the largest quantity is selected for the crystallisation of the entire quantity of the substance. If a substance is easily soluble in all solvents, recourse must be had to crystallisation by evaporation, *i.e.* by allowing the different solutions to stand some time in watch-glasses. That solvent from which crystals separate out first is the most suitable. Frequently a compound dissolves in a solvent only on heating and yet does not crystallise out again on cooling; compounds of this class are said to be "sluggish" (*träge*). In this case, the solution may be allowed to stand for some time, if necessary over night, in a cool place. If a compound is very difficultly soluble, solvents with high boiling-points are used, as toluene, xylene, nitrobenzene, aniline, phenol, and others. The crystals obtained in these preliminary experiments, especially if they are of easily soluble substances, are preserved, so that if from the main mass of the substance no crystals can be obtained, the solution may be seeded, thus inducing crystallisation. The crystallisation of substances which boil without decomposition may often be facilitated by first subjecting them to distillation.

Substances that are solids in the crude form, will sometimes

separate out as liquids from solvents. This may be due to the presence of small quantities of water. In this case, the solution in ether, ligroïn, benzene, etc., is heated with fused Glauber's salt ; crystals will often separate out when this is filtered.

To dissolve the Substance. — When water or glacial acetic acid, or a solvent which is not inflammable or not easily inflammable, is employed, the heating may be done in a beaker on a wire gauze over a free flame if the quantity is small ; if large, a flask is always used. In either case care must be taken to prevent the flask from breaking, by stirring up the crystals from the bottom with a glass rod, or by frequently shaking the vessel. This precaution is especially to be observed when, on heating, the substance to be dissolved melts at the bottom of the vessel. Alcohol and benzene may also be heated in like manner directly over a moderately large flame, if the student has already had a sufficient amount of experience in laboratory work and does not use too large quantities. If the liquid becomes ignited, no attempt to extinguish the flame by blowing on it should be made, but the burner is removed and the vessel covered with a watch-glass, a glass plate, or a wet cloth. In working with large quantities of alcohol, benzene, ether, ligroïn, carbon disulphide, or other substances with low boiling-points, they are heated on a water-bath in a flask provided with a vertical glass tube (air condenser) or a reflux condenser. A substance to be crystallised from a solvent which is not miscible with water must be dried, in case it is moist, before dissolving.

An error which even advanced students too often make in crystallising substances consists in this : an excessive quantity of the solvent is poured over the substance at once. When heat is applied, it is true, solution takes place easily, but on cooling nothing crystallises out. So much of the solvent has been taken that it holds the substance in solution even at ordinary temperatures. The result is that a portion of the solvent must be evaporated or distilled off, which involves a loss of time and substance, as well as decomposition of the substance. The following rule should, therefore, always be observed : *The quantity of solvent taken at first should be insufficient to dissolve the substance completely, even*

on heating; then more of the solvent is gradually added, until all of the substance is just dissolved. In this way only is it certain that on cooling an abundant crystallisation will take place. If a mixture of two solvents is used, one of which dissolves the substance easily and the other with difficulty, *e.g.* alcohol and water, the substance is first dissolved in the former with the aid of heat; the heating is continued while small amounts of the second are gradually added (if water is used it is better to add it hot) until the first turbidity appearing does not vanish on further heating. In order to remove this cloudiness, a small quantity of the first solvent is added. On the addition of the first portions of the second liquid (water or ligroin) resinous impurities separate out at times; in this case, these are filtered off before a further addition of the solvent is made.

At times it happens that the last portions of a compound will dissolve only with difficulty. The beginner often makes the mistake here of adding more and more of the solvent to dissolve this last residue, which for the most part generally consists of difficultly soluble impurities, like inorganic salts, etc. The result of this is that on cooling nothing crystallises out. In such cases the difficultly soluble portions may be allowed to remain undissolved, and on filtering the solution are retained by the filter.

Filtration of the Solution. — When a substance has been dissolved, the solution must next be filtered from the insoluble impurities like by-products, filter-fibres, inorganic compounds, etc. For filtration a funnel with a very short stem is generally used, *i.e.* an ordinary funnel the stem of which has been cut off close to the conical portion (Fig. 1). The funnels used in analytical operations have the disadvantage that when a hot solution of a compound flows through the stem, it becomes cooled to such an extent that crystals frequently separate out, thus causing an obstruction of the stem. The funnel with a shortened stem or no stem is prepared with a folded filter. In case the solution contains a substance that easily crystallises out, the filter



FIG. 1.

is made of rapid-filtering paper (Fig. 2). The solution to be filtered is not allowed to cool before filtering, but is poured on the filter immediately after removing it from the flame or water-bath. If inflammable solvents are used, care must be taken that the vapours are not ignited by a neighbouring flame. Under normal conditions, no crystals or only a few should separate out on the filter during filtration. If large quantities of crystals appear in a solution as soon as it is poured on the filter, it is



FIG. 2.

an indication that too small an amount of the solvent has been used. In a case of this kind, the point of the filter is pierced and the crystals are washed into the unfiltered portion of the solution with a fresh quantity of the solvent; the solution is further diluted with the solvent, heated, and filtered.

Very difficultly soluble compounds crystallise during the filtration in the space between the filter and funnel, in consequence of the contact of the solution with the cold walls of the funnel.

This may be prevented when a small quantity of liquid is to be filtered, by warming the funnel previously in an air-bath, or directly over a flame. If the quantity of the liquid is large, hot water or hot air funnels may be used (Figs. 3 and 4), or the funnel may be surrounded by a cone of lead tubing wound around it through which steam is passed (Fig. 5). Before filtering inflammable liquids, the flame with which the hot water or hot air funnel has been heated is extinguished. Substances which easily crystallise out again, may also be conveniently filtered with the aid of suction and a funnel having a large filtering surface (Büchner funnel, see Fig. 38, p. 58). After filtration the solution is poured into the proper crystallisation vessel. In order to prevent the thick-walled filter-flasks from being cracked by solvents of a high boiling-point, they are somewhat warmed before filtering by immersion in warm water.

Boiling nitrobenzene, aniline, phenol, and similar substances may be filtered in the usual way through ordinary filter-paper.

Choice of the Crystallisation Vessel.—The size and form of the

crystallisation vessel is not without influence upon the separation of the crystals. If a compound will crystallise out on simple cooling, without the necessity of evaporating a portion of the solvent, a beaker is used for the crystallisation. The shallow dishes known as "crystallising dishes" are not recommended for this purpose, since they cannot be heated over a free flame, and further, the solution easily "creeps" over the edge, involving a loss of the substance. Moreover, the crusts collecting on the edges are very impure, since, in consequence of the complete evaporation of the solvent, they contain all the impurities which should remain dis-



FIG. 3.

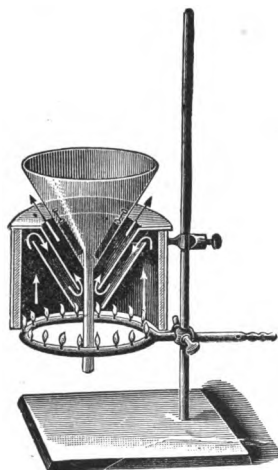


FIG. 4.

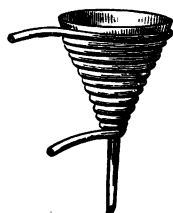


FIG. 5.

solved in the mother-liquor. The beaker is selected of such a size that the height of the solution placed in it is approximately equal to the diameter of the vessel, which is thus about one-half to two-thirds filled.

Heating after Filtration. — Many compounds crystallise out in the beaker during filtration. The crystals thus obtained are never well formed, in consequence of the rapid separation; therefore, after the entire solution has been filtered, it is heated again until the crystals have redissolved, and is then allowed to cool as slowly

as possible without being disturbed. In order to protect the solution from dust as well as to prevent it from cooling too rapidly, the vessel is covered first with a piece of filter-paper and then with a watch-glass or glass plate. The paper is used to prevent drops of the solvent formed by the vapours condensing on the cold cover-glass from falling into the solution, by which the crystallisation would be disturbed. The paper need not be used if the vessel is covered with a watch-glass, the convex surface of which is uppermost : the condensed vapours will thus flow down the walls of the beaker.

Crystallisation. — In order to obtain as good crystals as possible, the solution is allowed to cool slowly without being disturbed. In exceptional cases only is it placed in cold water to hasten the separation of crystals. The vessel must not be touched until the crystallisation is ended. If a substance, on slow cooling, separates out in very coarse crystals, it is expedient, in case a sample of the substance for analysis is desired, to accelerate the crystallisation by artificial cooling, so that smaller crystals will separate out. Very coarse crystals are commonly more impure than smaller ones, in that they enclose portions of the mother-liquor. If a deposit of crystals as abundant as possible is desired, the vessel is put in a cool place — in a cellar or ice-chest if practicable. Should a compound crystallise sluggishly, the directions given on page 2, under “Choice of the Solvent,” may be followed (rubbing the sides of the vessel with a glass rod ; seeding the solution ; allowing to stand over night). At times a compound separates out on cooling, not in crystals, but in a melted condition. This may be caused by the solution being so concentrated that crystallisation already takes place at a temperature above the fusing-point. In this case the solution is again heated until the oil which has separated out is dissolved, more of the solvent is then added, the quantity depending upon the conditions. In other cases this may be prevented by rubbing the walls of the vessel a short time with a sharp-edged glass rod, as soon as a slight turbidity shows itself, or by seeding the solution with a crystal of the same substance. This difficulty may also be avoided, in many cases, by allowing the solution to cool very slowly ; *e.g.* the beaker is placed in a larger vessel filled with hot water and allowed to cool in this.

At times the separation of crystals takes place suddenly, within a few seconds, throughout the entire solution. Since the crystals thus obtained are generally not well formed, the liquid, after some of the crystals have been removed, is heated until solution again takes place. After it has partially cooled, those crystals which were taken out are now added to it, by which a gradual crystallisation is caused.

Separation of Crystals from the Mother-Liquor. — When crystals have been deposited, they are then to be separated from the liquid (mother-liquor). This is always done with the aid of suction, and never by merely pouring off the liquid. The filter to be used is previously moistened with the same substance which was employed as the solvent. Crusts, formed on the sides and edges of the vessel by the complete evaporation of the solvent, are not filtered with the crystals; they are removed with a spatula before the filtering, and are worked up with the mother-liquor. In order to remove the last traces of the mother-liquor adhering to the crystals, they are washed several times with fresh portions of the solvent; obviously, if the substance is easily soluble, too large quantities of the solvent must not be used. If a solvent that will not evaporate easily in the air or on the water-bath has been used, *e.g.* glacial acetic acid, toluene, nitrobenzene, etc., it must be removed from the crystals by a more volatile substance, like alcohol or ether. This is done by first washing with a fresh quantity of the solvent, then with a mixture of the solvent and a small quantity of the more volatile liquid, the proportion of the latter in the washing mixture being gradually increased, until finally the volatile substance is used alone. Glacial acetic acid may, in this way, be displaced by water.

Drying of Crystals. — When crystals have been freed from the mother-liquor they must be dried. This may be effected (1) at the ordinary temperature by the gradual evaporation of the solvent in the air, and (2) at higher temperatures by heating on a water-bath or in an air-bath. In the first case the crystals are spread out in a thin layer upon several thicknesses of filter-paper and covered with a watch-glass, funnel, beaker, or similar vessel. In order that the vapours of the solvent may escape, the covering must be so placed that the air is not shut off completely from the crys-

tals ; this is conveniently done by supporting it on several corks. Crystals may also be dried in a desiccator which is partially exhausted, if necessary. In drying substances at higher temperatures the crystal form may be lost by the fusion of the substance or by the separation of the water of crystallisation. Since many substances will liquefy far below their melting-point if they contain even small quantities of the solvent, a preliminary experiment with a small portion is always made when the drying is to be effected at higher temperatures. Compounds, not easily soluble in ether, which crystallise from a solvent miscible with ether, can be very quickly dried by being washed several times with it. After a short exposure to the air they are dry.

Treatment of the Mother-Liquor. — The mother-liquor filtered off from crystals still contains more or less of the substance, in proportion to its solubility at the ordinary temperature ; in many cases it is advantageous to extract the last portions remaining in solution. A "second crystallisation" is obtained by distilling or evaporating off a portion of the solvent. The mother-liquor may also be diluted with a second liquid, in which the dissolved substance is difficultly soluble ; *e.g.* a solution in alcohol or glacial acetic acid may be diluted with water, or a solution in ether or benzene with ligroin.

Crystallisation by Evaporation. — If a compound is so easily soluble in all solvents that it will only crystallise out on partial evaporation, then, in order to get good crystals, a solution, not too dilute, is made, by the aid of heat if necessary, and filtered from the impurities remaining undissolved. In this case, as a crystallisation vessel, one of the various forms of shallow dishes — the so-called crystallising dishes — is used, in which the solution is allowed partially to evaporate. In order to protect the vessel from dust, it is covered with a funnel or watch-glass, in the manner indicated under "Drying of Crystals." In crystallising by this method, it sometimes happens that the solution, owing to capillary action, will "creep" over the edge of the dish. To avoid loss of the substance from this source, the dish is placed on a watch-glass or glass plate. Under these conditions, the vessel

is never covered with filter-paper, since, after standing some time, it may absorb the entire quantity of the substance. If, in order to obtain well-formed crystals, the solvent is to be evaporated as slowly as possible the solution is placed in a beaker or test-tube, which is then loosely covered with filter-paper. Evaporation may be hastened by placing the crystallisation vessel in a desiccator, charged, according to the nature of the solvent, with different substances; for the absorption of water or alcohol, calcium chloride or sulphuric acid is used; glacial acetic acid is absorbed by soda-lime, solid potassium hydroxide, or sodium hydroxide. The evaporation of all solvents may be hastened by exhausting the desiccator.

Since the purifying effect of crystallisation depends upon the fact that the impurities remain dissolved in the mother-liquor, and with this are filtered off, in no case must the solvent be allowed to evaporate completely, but the crystals must be filtered off while still covered with the mother-liquor. Before filtering, crusts deposited, generally on the edges of the vessel, are removed with the aid of a small piece of filter-paper or a spatula. Even though the substance is very soluble, the mother-liquor adhering to the crystals is washed away with small quantities of the solvent. If the quantity of crystals is very small, the adhering mother-liquor may be separated, in cases of necessity, by placing them on porous plates and moistening with a spray of the solvent.

Fractional Crystallisation.—Up to this point, it has been assumed that the substance to be crystallised possessed an essentially homogeneous nature, and the object of crystallisation was only to change it to a crystallised form. Crystallisation is often employed for another purpose—that of separating a mixture of different substances into its individual constituents,—a task that is generally far more difficult than the crystallisation of an individual substance. The simplest case is one in which two substances are to be separated. If the solubilities of the two substances are very different, as is generally the case when a mixture of two different highly substituted compounds is under examination, it is frequently not difficult to find a solvent which will dissolve a considerable portion of the more easily soluble substance, and but a

small portion of the less soluble. If, now, the mixture be treated with such a solvent, in not too large quantities, a solution will be obtained containing all of the easily soluble substance and a small

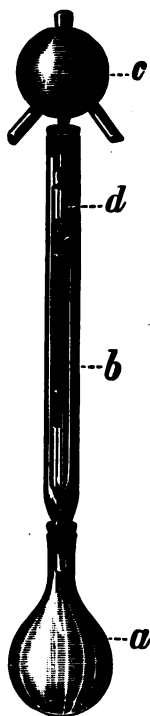


FIG. 6.



FIG. 7.

portion of the difficultly soluble substance. This is filtered from the residue remaining undissolved. The mixture has thus been divided into two fractions. By evaporating the solution to a certain point, the more insoluble compound will crystallise out, unaccompanied by any of the other compound; the crystals are filtered off, and the solution further evaporated. If the crystallisation of the two fractions be repeated a second time, a complete separation will be effected. For separating a mixture of this kind, specially constructed apparatus—the so-called extraction apparatus—may be employed, the use of which possesses the advantage over the method of simple heating, that much smaller quantities of the solvent are required. An apparatus of this kind is represented in Figs. 6 and 7. To a wide glass tube *d* is fused a narrow tube which acts as a siphon, bent as in Fig. 7. This portion of the apparatus is surrounded by a glass jacket *b*, narrowed at its lower end. This is connected with the flask that is to contain the solvent. A cork bearing a reflux condenser

—a ball condenser is convenient—is fitted in the opening at the upper end of the jacket. A shell of filter-paper is next prepared in the following manner: Three layers of filter-paper are rolled around a glass tube with half the diameter of the inner tube *d*. One end of the roll must extend somewhat beyond the edge of the glass tube; this is turned over and securely fastened with thread. To preserve the form of the roll, thread is loosely wound around its middle and upper portion. The length of the roll is such that it

extends 1 cm. above the highest point of the narrow siphon-tube. In the shell is placed the mixture of the easily soluble and difficultly soluble substance to be extracted; the upper end is closed by a loose plug of absorbent cotton. The flask *a*, containing the solvent, is now heated on a water-bath or over a free flame, according to the nature of the solvent. The condensed vapours drop from the condenser into the shell, dissolve the substance, filter through the paper, and fill the space between shell and inner glass tube. As soon as the liquid has reached the highest point of the siphon-tube, the solution siphons off and flows back into the flask *a*. This operation may be continued as long as necessary. The amount of solvent used should be one and a half or two times the volume of the inner tube up to the highest point of the siphon. The construction of a ball condenser is represented in Fig. 8. In order to distinguish the tube by which the water enters from the outlet-tube, the former is marked with an arrow. Comparatively easy also is the separation of two substances about equally soluble, if the one is present in larger quantity than the other. If a mixture of this kind is dissolved, then, on cooling, the substance which was present in larger quantity generally crystallises out. Occasionally, after standing some time, crystals of the second substance will appear; under these conditions the crystallisation must be carefully watched, and as soon as crystals differing from those first appearing are observed, the solution is filtered with suction at once, even though it is still warm.

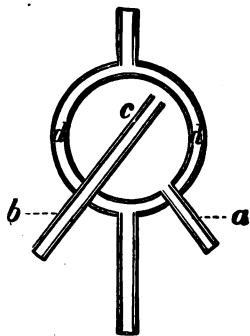


FIG. 8.

If two compounds crystallise simultaneously at the outset, as is the case when they possess approximately the same solubility and are present in almost equal quantities, they can be separated mechanically. If, *e.g.*, one of the compounds crystallises in coarse crystals, and the other in small ones, they may be separated by sifting through a suitable sieve or wire gauze. A compound crys-

tallising in leaflets can frequently be separated from one crystallising in needles by a sieve. If these methods fail, the separation may be effected by picking out the crystals with small pincers or a quill. In all these mechanical operations, the crystals must be as dry as possible.

In many cases, when one of the compounds is heavier than the other, it is possible to separate them by causing the lighter crystals to rise to the top of the liquid, by imparting to it a rotatory motion by rapid stirring with a glass rod. The heavier compound collects at the bottom of the vessel, and the liquid with the lighter compound floating in it can be poured off.

Double Compounds with the Solvent. — Many substances crystallise from certain solvents in the form of double compounds, composed of the substance and the solvent. It is well known that many substances, in crystallising from water, combine with a certain portion of water. Alcohol, acetone, chloroform, benzene, and others also have the power of uniting with other substances to form double compounds. As a familiar example, the combination of triphenylmethane with benzene may be mentioned in this connection. If double compounds of this kind are heated, the combined solvent is generally vaporised.

SUBLIMATION

Much less frequently than crystallisation, sublimation is used to purify a solid compound. The principle involved is this: A substance is converted by heat into the gaseous condition. The vapours do not assume the liquid phase when they are condensed on a cold surface, but deposit in the form of crystals.

The sublimation of a small quantity of a substance can be conveniently effected between two watch-glasses of the same size. The substance to be sublimed is placed on the lower one, which is then covered with a round filter perforated several times in its centre and projecting over the edges; the second watch-glass with its convex side uppermost is placed on it, and the two are held together by a watch-glass clamp. If the lower glass is now heated

very slowly on a sand-bath with a free flame, the vaporised substance condenses on the cold surface of the upper watch-glass in crystals; the filter-paper prevents the very small, light crystals from falling back on the hot surface of the lower glass. To keep the upper glass cool, it is covered with several layers of wet filter-paper or with a small piece of wet cloth. If large quantities of a substance are to be sublimed, the upper watch-glass in the apparatus just described is replaced by a funnel somewhat smaller than the lower glass (Fig. 9). To prevent the escape of vapours, the stem of the funnel is closed by a plug of cotton or is covered with a small cap of filter-paper. The apparatus for sublimation designed by Brühl is admirably adapted to the purpose for which it is intended (Fig. 10). It consists of a hollow metal plate through which water flows. In the conical opening is placed a crucible containing the substance to be sublimed. The plate is covered with a concave glass dish, the ground edges of which fit the plate tightly. The crucible is heated directly with a small flame, while cold water flows through the plate. The vapours condense in part on the glass cover, but



FIG. 9.

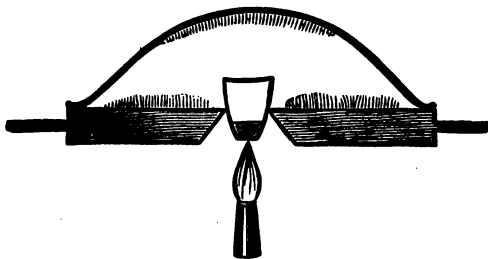


FIG. 10.

more abundantly on the upper cold surface of the plate in crystals. The glass cover is not removed until the apparatus is completely cold.

Sublimations can also be conducted in crucibles, flasks, beakers, retorts, tubes, etc. The heating may be done in an air- or oil-bath. In order to lead off the vapours rapidly, a current of an indifferent gas is sent through the apparatus.

Of late years substances of high purity have been obtained by causing them to sublime *in vacuo*. An apparatus devised for this purpose is described in the *Journal für praktische Chemie*, Vol. 78 (1908), page 201.

DISTILLATION

Kinds and Objects of Distillation. — By distillation is meant the conversion by heat of a solid or liquid substance into a vapour and the subsequent condensation of this. When a solid is distilled it does not condense directly in crystals, as is the case in sublimation. The distillate is a liquid which may solidify into a crystalline mass on standing. When distillation is conducted at the atmospheric pressure, it is called *ordinary distillation*; if in a partial vacuum, *vacuum distillation*. The object of distillation is either to test the purity of an individual substance by the *determination* of its *boiling-point*, or to separate a mixture of substances boiling at different temperatures into its constituents. (*Fractional Distillation.*)

Distillation Vessels. — The heating of the substance to be distilled is generally effected in a fractionating flask (Figs. 11, 12, 13). These flasks differ, not only in size, but in the diameter of the condensation-tube (side-tube), as well as in the distance of the latter from the bulb. In selecting a fractionating flask the following points are to be observed. For distillation at the atmospheric pressure a flask is selected having a bulb of such a size that when it contains the substance to be distilled it will be about two-thirds filled. There are two objections to distilling small quantities of a substance from a large flask: the vapours are easily overheated, thus giving a boiling-point that is too high; a loss of the substance follows, in that, after the distillation is finished, a larger volume of vapours which condense on cooling, remains behind in the bulb, than if a smaller flask had been used. In the distillation of low boiling

compounds, a flask is selected which has its condensation-tube as high as possible above the bulb, so that the entire thread of mercury of the thermometer employed is heated by the vapour of the liquid. By using a flask of this kind it is not necessary to correct the observed boiling-point, as is the case when the mercury column is not entirely surrounded by the vapour. The higher a substance boils, the nearer must the side-tube be to the bulb, in

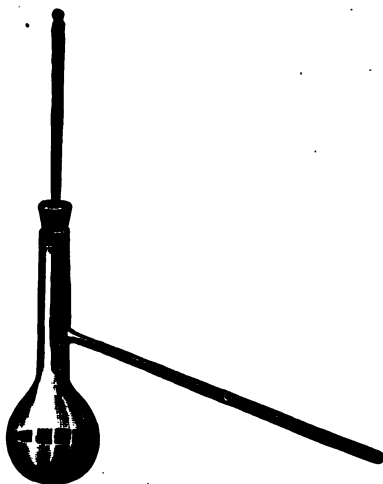


FIG. 11.

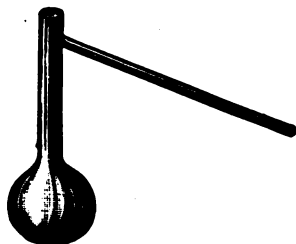


FIG. 12.



FIG. 13.

order that the vapours shall have as little opportunity as possible of condensing below the tube and flowing back into the bulb.

If large quantities of a substance are to be distilled, an ordinary flask is used. This can be converted into a fractionating flask with the aid of a cork bearing a T-tube, as illustrated in Fig. 14.

For the distillation of solid substances which solidify in the condensation-tube, a fractionating flask with a wide side-tube is used.

A fractional distillation can also be conducted in the fractionating flasks just described ; but the operation can be carried out more

rapidly and more completely by the use of apparatus especially adapted to fractionating (Fig. 15). These can be fused directly on the bulb or they can be attached to an ordinary flask by means of a cork (Fig. 14); the round, short-necked flasks such as represented in Fig. 16, are well adapted to this purpose. Flasks of

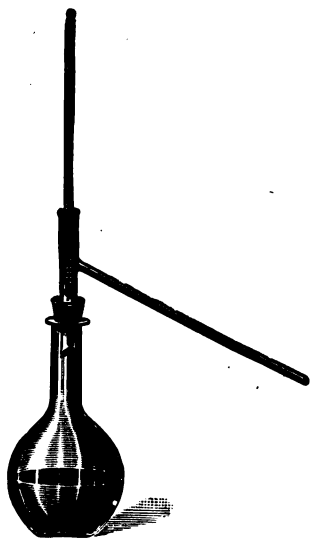


FIG. 14.

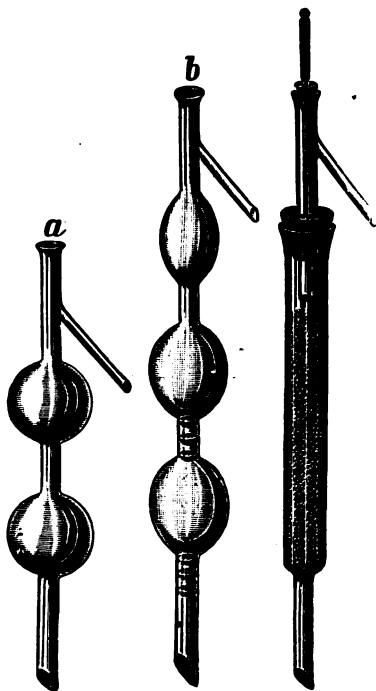


FIG. 15.

Fractionating Apparatus.

WURTZ

LINNEMANN

HEMPEL

this description can be obtained in different sizes but still possessing the same width of neck; this enables one to use the same cork with any flask. The value of these different forms of fractionating apparatus depends upon the fact that the higher boiling portions carried along with the vapours do not pass immediately

to the outlet tube, but before entering this they have an opportunity of condensing and flowing back into the flask. In the apparatus of Wurtz (*a*) the condensation takes place on the large upper surfaces of the bulbs. More complete condensation is obtained in Linnemann's apparatus (*b*), which differs from that of Wurtz in that the narrow tubes between the bulbs contain small platinum-wire sieves. Since the lower boiling portions condense to a liquid and collect in these, the ascending vapours are so far cooled by the passage through them that the accompanying portions of the higher boiling substances are likewise condensed. The apparatus of Hempel is filled with glass beads which act like the sieves in the Linnemann apparatus. For the distillation of large quantities of a liquid the Hempel apparatus is particularly well adapted; in working with it as well as the Linnemann form, the heating must be interrupted from time to time, in order that the liquid collecting in the beads or sieves may have an opportunity to flow back to the distillation flask. If the Le Bel-Henninger form is used, this precaution is unnecessary, since in this apparatus special tubes for conducting off the condensed liquid are joined to the sides of the bulb somewhat above the sieves.

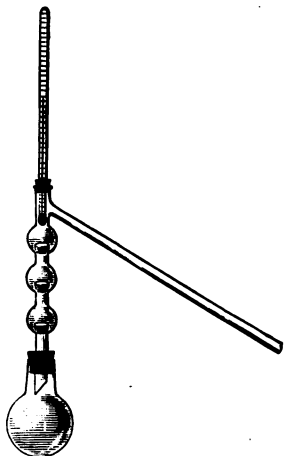


FIG. 16.

Experiments have shown that a single distillation with one of the forms of apparatus just described, effects a more complete separation than repeated fractionations in an ordinary fractionating flask.

Supporting the Fractionating Flask.—If it is necessary to support the fractionating flask with a clamp, it is placed as far above the outlet tube as possible, never below it; the glass expands by contact with the hot vapours, and since the expansion is impeded by the clamp, particularly if it is firmly attached, the flask frequently breaks.

Supporting the Thermometer.—The thermometer is passed through a cork (no rubber) which fits the neck of the flask. The most exact determinations of the boiling-point are obtained if the entire thread of mercury is surrounded by the vapour of the substance. With low boiling compounds this condition is easily obtained by the use of a fractionating flask having its outlet tube at a sufficient distance above the bulb. In this case the thermometer is so placed that the degree corresponding to the boiling-point of the liquid is opposite the outlet tube, but the bulb of the thermometer must not extend into the bulb of the flask and never into the liquid; if it does, another flask must be used, the outlet tube of which is still higher above the bulb. If in dealing with high boiling compounds such an arrangement is not possible, the thermometer is thrust so far into the neck of the flask that the thermometer-bulb is somewhat below the outlet tube. In this case, if an exact determination of the boiling-point is desired, the observed reading is corrected in the manner described below. In order to avoid making a correction a special form of thermometer is used, the graduation of the scale beginning at 100° , 200° , or at other convenient points. By employing an instrument of this kind the mercury column may be kept in the vapours at any temperature.

In making distillations, it occasionally happens that the mercury column ascends to that point in the scale which is hidden by the cork supporting the thermometer, thus preventing the temperature from being read. In a case of this kind the thermometer is either raised or lowered, so that the top of the mercury is visible, or if this is not possible, from that portion of the cork which projects above the flask, a section is cut which will enable the scale to be seen.

Condensation of Vapours.—The condensation of vapours is effected in various ways, depending upon the height of the boiling-point. If a compound boils at a relatively low temperature (up to 100°), the outlet tube of the fractionating flask is connected with a Liebig condenser by a cork (not a rubber stopper). For very low boiling compounds a long condenser is used, and for those of

high boiling-points a short one. If the boiling-point of a compound is very low, the flask in which the condensed liquid collects (the receiver) is connected with the condenser by means of a cork and a bent adapter (Fig. 63), and the receiver is cooled by ice or a freezing mixture. If the boiling-point is moderately high, between 100° and 200° , the receiver, connected to the condensing tube by a cork, is cooled by running water (Fig. 17). If the substance is to be distilled again, a fractionating flask is employed as a receiver; a tubulated suction-flask may also be used. It is often unnecessary to employ running water for cooling purposes if to the outlet tube of the flask a wide glass tube 50 cm. long (extension tube) is connected by a cork (Fig. 18). With still higher boiling substances even this is superfluous, since the condensation tube of the fractionating flask, provided it is not too short, will suffice for the condensation.

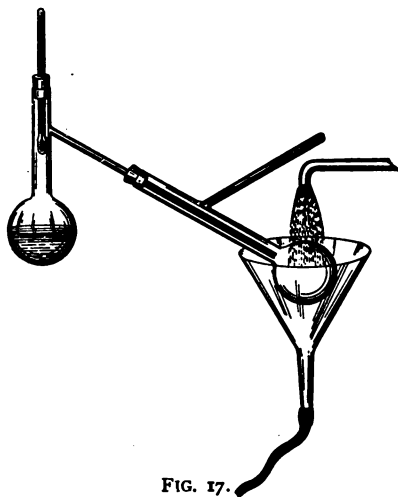


FIG. 17.

If a small quantity of a substance is to be distilled, and it is desired to avoid the loss of substance necessarily incident to the use of a condenser, the distillation even of low boiling compounds is conducted in a small distillation flask as slowly and carefully as possible, the source of heat being a minute flame (the so-called microburner).

If large quantities are to be distilled, a condenser is always used, since when other condensation apparatus is employed, the tube finally becomes so hot that the vapours are not completely condensed. If the vapours of a substance attack corks, the outlet tube is inserted far enough into the condenser or extension tube

so that the vapours do not come in contact with the cork. But generally a cork is not used ; the outlet tube being inserted sufficiently far into the condenser.

Heating.—Low boiling substances (those boiling up to about 80°) are not generally heated over the free flame, but on the water-bath gently or to full boiling. Frequently it is more convenient to immerse the bulb of the fractionating flask as far as the level of the liquid which it contains in a dish or beaker filled with water, which is heated gently or strongly as the case requires. Low boiling substances may also be heated by immersing the bulb

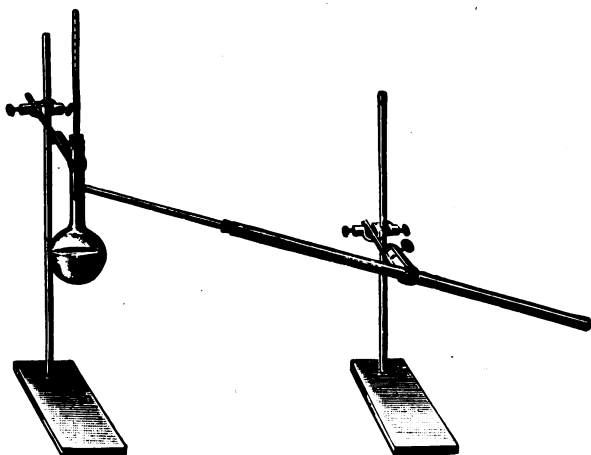


FIG. 18.

of the flask from time to time in a vessel filled with warm water. If a substance is not distilled over a free flame, in order to prevent "bumping" a few pieces of platinum wire or foil, or bits of glass, are thrown into the liquid (see below). When a substance to be distilled is heated on the water-bath, it may easily happen that the vapour inside the flask may be overheated by the steam escaping between the rings. For this reason, in the determination of exact boiling-points it is better to use a small free flame. The so-called microburner is well adapted to this purpose. High boil-

ing substances are always heated over the free flame. In this case the flask may be protected by heating it on a wire gauze ; still by working carefully the gauze need not be used. In heating, the flame is not placed under the flask at once, since the latter is likely to break easily on sudden heating ; it is better to pass the flame back and forth slowly and uniformly over the bottom of the flask until the liquid is brought to incipient ebullition. Substances which have been previously dissolved, after the evaporation of the solvent on the water-bath, often stubbornly refuse to give up the last portions of the solvent, particularly when ether has been used. If now a free flame be applied, it frequently happens that in consequence of a retarded boiling during which the solution becomes overheated, a sudden active ebullition and foaming will take place. In order to prevent this the flask is shaken repeatedly during the heating, since if the liquid is kept in motion, overheating cannot easily take place. It may also be prevented frequently by heating the flask on the side. During the actual distillation the heating may be continued by slowly passing the flame over the bottom of the flask as in the preliminary heating, but in this case care must be taken not to apply the flame to the flask at any point above the liquid inside, since an overheating of the vapours would result. In order to protect the hand in case the flask should break, the burner is held obliquely and not directly under the flask ; or during the distillation the burner may be placed under the flask and allowed to remain stationary. The size of the flame is so regulated that the condensed distillate flows into the receiver regularly in drops. If vapours escape from the receiver, it is an indication that the heating is too strong. Toward the end of the distillation the burner is turned down somewhat.

To collect the Fractions.—If a substance which is not quite pure is being treated, and it is desired to test the purity by a determination of its boiling-point, then on distillation a small portion will generally pass over below the true boiling-point (“first runnings”); this is collected separately in a small receiver. Then follows the principal fraction, passing over at the true boiling-point, the temperature remaining constant. If there is only a small

quantity of the liquid in the bulb of the flask, it is difficult, in spite of using a small flame, to prevent the vapours from being somewhat overheated; this will cause a rise of the mercury. The portion passing over a few degrees above the true boiling-point can, in preparation work, be collected with that portion which boils at the correct temperature, without evil results. High boiling portions collected separately are designated as "last runnings." The operation of fractional distillation is conducted in a wholly different manner. The preparation of benzoyl chloride (see page 317) will furnish a practical example of the method of procedure. This compound is obtained by treating benzoic acid with phosphorus pentachloride. The product of the reaction is a mixture of phosphorus oxychloride (b. p. 110°) and benzoyl chloride (b. p. 200°). If this mixture is subjected to distillation, the entire quantity of phosphorus oxychloride does not pass over at about 110° , and afterwards the benzoyl chloride at 200° ; but the distillation will begin below 110° , and a mixture consisting of a large quantity of the lower boiling substance and a small quantity of the higher boiling substance will pass over; the temperature then rises gradually; while the quantity of the former steadily decreases, that of the latter increases, until finally, at 200° , a mixture consisting essentially of the higher boiling substance passes over. A quantitative separation of the constituents of a mixture cannot be effected by the method of fractional distillation. However, in most cases, it is possible to obtain fractions which contain the largest part of the individual constituents, particularly when, as in the example selected, the boiling-points of the constituents lie far apart, by collecting the different fractions and repeating the distillation a number of times. It is almost impossible to give definite rules of general application for fractional distillation; the number of fractions to be collected depends upon the difference of the boiling-points, upon the number of compounds to be separated, upon the relative proportion of the compounds present, and upon other factors. If but two substances are to be separated, as is generally the case in preparation work, the procedure is, very commonly, as follows: as a basis for the fractions to be collected, the interval

between the boiling-points is divided into three equal parts; in the case of the example selected the temperatures would be 110° , 140° , 170° , 200° . The fraction passing over between the temperature at which the distillation first begins, up to 140° , is collected (fraction I.), then in another vessel the fraction passing over between 140° – 170° (fraction II.), and finally in another receiver that passing over between 170° and 200° (fraction III.). The quantities of the three fractions thus obtained are about equal. Fraction I. is now redistilled from a smaller flask, and the portion passing over up to 140° is collected as in the first distillation in the empty receiver I., which in the meantime has been washed and dried. When the temperature reaches 140° , the distillation is stopped, and to the residue remaining in the flask is added fraction II., and the distillation continued. The portion passing over up to 140° is collected in receiver I., that from 140° – 170° in the empty receiver II. When the temperature reaches 170° , the distillation is again interrupted, and to the residue in the flask is added fraction III., and the distillation is again continued: in this way the three fractions are collected. These are again distilled as in the first distillation, but now the lower and higher boiling fractions are much larger than the intermediate one; further, a larger portion of these end fractions boil nearer the true boiling-points than in the first distillation. If it is now desired to obtain the two substances in question in a still purer condition, the two end fractions are once more distilled separately, and the portion passing over a few degrees above and below the true boiling-point, for phosphorus oxychloride about 105° – 115° , for benzoyl chloride, 195° – 205° are collected.

Vacuum Distillation. — Many compounds, not volatile at the atmospheric pressure without decomposition, may be distilled undecomposed in a partial vacuum. The vacuum distillation is used advantageously for the fractionation of small quantities of a substance, since the separation of the individual constituents can be effected more rapidly and more completely than at the atmospheric pressure.

Vacuum Apparatus. — The simplest form of a vacuum apparatus

is represented in Fig. 19. Two fractionating flasks *a* and *b* are connected by a cork. The neck of *a* is closed by a tightly fitting cork bearing the glass tube *d*, reaching to the bottom of the flask, its lower end being drawn out to a fine point, the object of which will be explained below. A thermometer is placed in the tube. A thermometer is placed in the tube.

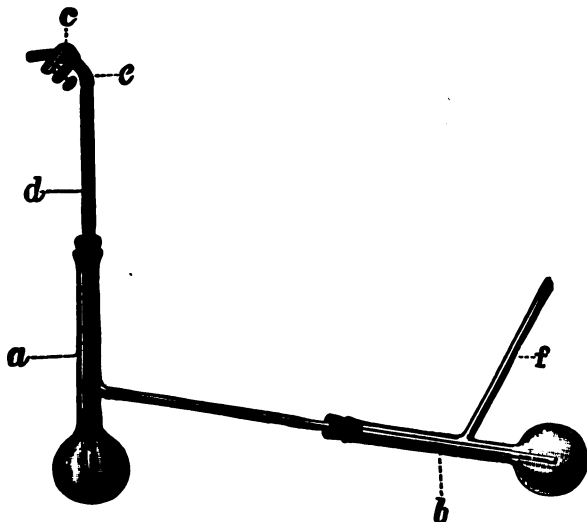


FIG. 19.

In place of the flask *b*, a suction-flask such as finds application in filtering under pressure, may be used (Fig. 20). But this kind of flask is used only in case low boiling substances are to be distilled, since the contact of too hot liquids with the thick walls causes them to crack easily: this is likely to prove very destructive in vacuum distillation. With low boiling substances, in order to get complete condensation of the vapours, the jacket of a Liebig condenser through which water is allowed to flow is fitted over the outlet tube of the fractionating flask. These simple forms of apparatus are used only when it is desired to collect a few fractions, since it is troublesome to be obliged to change the receiver, and thus destroy the vacuum, for each new fraction.

If it is desired to collect a larger number of fractions, an

apparatus is employed by means of which the receiver can be changed without destroying the vacuum.

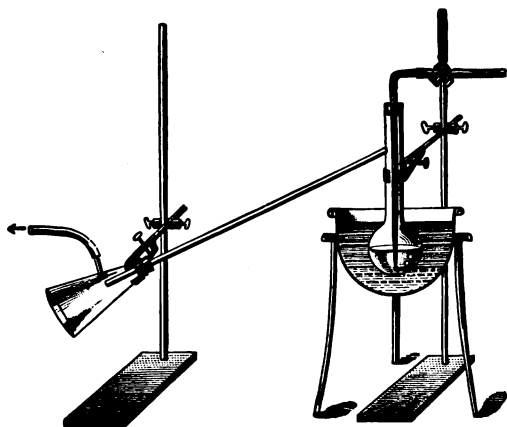


FIG. 20.

Brühl's apparatus is very well adapted to this purpose (Figs. 21 and 22). By turning the axis *b*, so arranged that it supports the receivers firmly, each receiver may in turn be brought under the end of the condenser tube *c*.

The receiver shown in Fig. 23 is also very convenient for fractional distillation in a vacuum. By grasping the cork *a* and the tube *c* firmly with the fingers and turning, the different portions of the receiver may be brought under the condensing tube.

Construction of a Vacuum Apparatus. — In vacuum distillations the evolution of bubbles of vapour occurs to a much greater extent than under ordinary conditions. In order to prevent the liquid from foaming up and passing over, a flask of such a size is selected, that when it contains the liquid it must in no case be more than half full; it is better to have it but one-third full. The individual parts of the apparatus are connected by rubber stoppers. Ordinary corks may also be used with almost equally good results, but only those are selected which are as free as possible from

pores; they are pressed in a cork-press, and then very carefully bored. If, after the apparatus is put together, the corks are coated

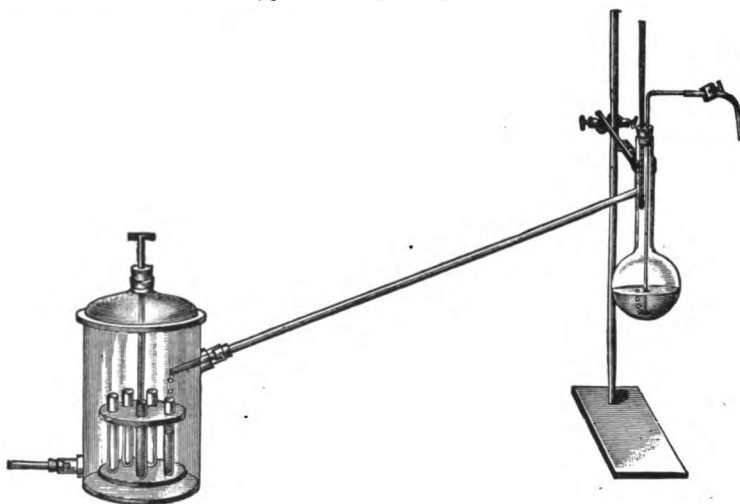


FIG. 21.

with a thin layer of collodion, there is no difficulty in obtaining a vacuum. The thermometer and capillary tube may be arranged as

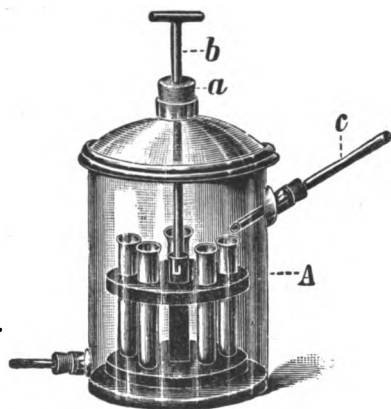


FIG. 22.

shown in Fig. 19. It is also a very excellent arrangement to use a two-hole cork, the thermometer passing through one, and the capillary tube through the other, as in Fig. 21. The capillary tube is made by drawing out a glass tube of 1-2 mm. diameter; the narrow hole in the cork through which this passes is made conveniently by a hot knitting-needle. Instead of using a capillary tube to prevent "bumping," other

means may be employed (see below), in which case the thermometer is supported in the fractionating flask as in ordinary distillations. When a tube drawn out to a capillary point is used, a short piece of thick-walled rubber tubing, which can be closed by a screw pinch-cock (Fig. 19, *e* and *c*), is attached to the upper end.

The flasks recommended by Claisen (Fig. 24) may be used advantageously in vacuum distillations in place of the common fractionating flasks. A tube drawn out to a capillary point is secured in the limb *a* by a piece of thick-walled rubber tubing or a cork. The thermometer is inserted in *b*. When a few large pieces of broken glass are placed in *b*, these flasks possess the advantage of preventing portions of the liquid (even in cases of violent boiling) from being carried over into the condenser. The space above the pieces of broken glass may be filled, partially or

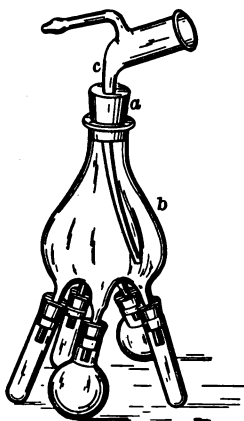


FIG. 23.

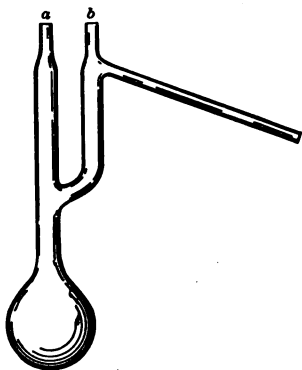


FIG. 24.

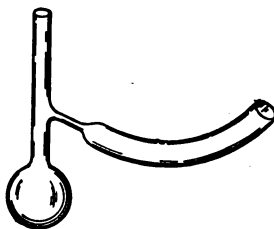


FIG. 25.

wholly, with glass beads — obviously these are only to be used in the distillation of liquids not having a too high boiling-point —

thus combining the advantages of a Hempel column with vacuum distillation.

For the distillation of solids a fractionating flask with a wide, bent sabre-shaped condensing tube is used (Fig. 25). In order to determine the efficiency of the vacuum, the lower tube of the Brühl apparatus is connected with a manometer (Fig. 26), by means of a thick-walled rubber tubing which will not collapse upon exhausting the apparatus. The other end of the manometer is connected with suction, by the same kind of rubber tubing.

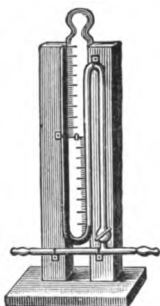


FIG. 26.

Since in consequence of the varying water pressure, it happens, at times, that the water from the suction pump may be forced into the manometer or receiver, it is advisable to insert a thick-walled suction flask between the suction pump and manometer.

In order that the apparatus may be perfectly tight, the corks, ends of the rubber tubing, as well as the ground surfaces of the Brühl receiver, are covered with a thin layer of grease or vaseline. If ordinary corks are used, these, as well as the ends of the tubing, are covered with collodion after the apparatus is set up. Before the distillation, the apparatus is tested to determine whether it will give the desired vacuum. For this purpose, the pinch-cock on the capillary tube is closed, the suction attached, and after some time the manometer is read: this will indicate whether the desired vacuum has been obtained. In case it is not, the corks are pressed more firmly into the tubes, greased again or covered with more collodion, and the rubber tubing is pushed farther over the ends of the glass. Frequently the suction pump will not work satisfactorily; it is then examined to see if it is stopped up, or a better pump is used. When the apparatus has been exhausted, the air must not be admitted suddenly, by removing a rubber joint, for the sudden rushing in of the air may easily destroy the apparatus. The rubber tube attached to the suction is closed by a screw pinch-cock which has been placed on it beforehand, and in case a

capillary tube has been used, the pinch-cock on this is gradually opened and the air allowed to enter through it, or after disconnecting the rubber tubing from the suction, the pinch-cock which has just been closed may be opened. The same object may be accomplished most rapidly by closing the tubing leading to the suction with the fingers, detaching it and opening the tube repeatedly for an instant at a time, until the rushing sound made by the inflowing air ceases. After a test has shown that the apparatus does not leak, the liquid to be distilled is poured in the flask and the distillation begun.

Heating. — In vacuum distillation the flask can be heated directly with a free flame, but the flame must be applied to the side, and not to the bottom of it, as in the ordinary way. Care must be taken to keep the flame constantly moving. It is much more satisfactory and safer to use an oil- or paraffin-bath, or better a metallic air-bath (iron crucible). The latter is covered with a thick asbestos plate containing a round opening in the centre, through which the neck of the fractionating flask may pass; from the opening to the edge of the plate there is a straight narrow slit. The air-bath must not be too large; the bottom is covered with a thin layer of asbestos, which will prevent the flask from coming in contact with the metal. The temperature of the oil- or air-bath should, in exact experiments, not be more than 20° – 30° higher than the boiling-point indicated by the thermometer. A thermometer is immersed in the bath and the flame so regulated that the difference between the two thermometers is not greater than that mentioned. The heating is not begun until the apparatus is exhausted.

To prevent Bumping. — In vacuum distillations a troublesome bumping (a sudden, violent ebullition) frequently occurs. To prevent this a slow continuous current of air is drawn through the liquid, thus keeping it in constant motion. The air current, controlled by a pinch-cock, must not be allowed to enter too rapidly, otherwise it will be difficult to maintain a high vacuum. The same effect may be obtained by placing certain substances in the liquid — splinters of wood the size of a match, capillary tubes, bits of glass, pieces of porcelain, powdered talc, scraps of platinum wire or foil. Small pieces of pumice-stone bound with platinum

wire also act satisfactorily. For further details concerning vacuum distillation consult "Die Destillation unter vermindertem Druck im Laboratorium," R. Anschütz.

Lowering of the Boiling-Point. — In order that some idea may be obtained as to the approximate lowering of the boiling-point, by diminishing the pressure, the following table is given :

Substance.	Boiling-point at 12 mm.	Boiling-point at Ordinary Pressure.	Difference.
Acetic acid	19°	118°	99°
Monochloroacetic acid . .	84°	186°	102°
Chlorobenzene	27°	132°	105°
p-Nitrotoluene	108°	236°	128°
Acetanilide	167°	295°	128°

Corrections of the Boiling-Point. — If it is not possible in making an exact determination of the boiling-point to have the mercurial column entirely surrounded by the vapour of the liquid, — a condition usually obtained by employing a flask, the side-tube of which is at a sufficient distance from the bulb, or a sectional thermometer, or both, — then a correction may be applied to the observed boiling-point in one of two ways. The portion of the mercurial column not heated by the vapours — that portion above the side-tube — is read in degrees (L). Another thermometer is brought as near as possible to the middle point of this column, the temperature of which is also read (t). If T is the observed boiling temperature, then the following correction is added: $L(T - t) \cdot 0.000154^\circ$. The so-called "corrected" boiling-point may also be obtained as follows: The boiling-point is determined in the usual way; after the distillation, another substance, the corrected boiling-point of which is known, and which lies near the one in question, is placed in the same flask and distilled under the same conditions. The difference between the corrected and observed boiling-points is applied to the boiling-point of the first substance.

Distilling off a Solvent. — An operation frequently employed in organic work is distilling off a solvent from the substance dis-

solved in it. When the boiling-point of the solvent is sufficiently far away from that of the dissolved substance, a complete separation can be effected by a single distillation. The methods used depend upon the quantity of the solution, that of the dissolved substance and the boiling-point of the solvent. The methods which can be used for distilling off low boiling solvents, like ether, ligroïn, carbon disulphide, alcohol, and others, will be described first. If a small quantity of a solvent is to be evaporated, and it is not worth the trouble to recover it by condensation, then, in case the solvent is ether, ligroïn, or carbon disulphide, the solution is poured into a small flask, and this is immersed in a larger vessel filled with warm water. The vaporisation is considerably accelerated by shaking the flask. The operation is more rapidly performed by heating the flask on a water-bath. To prevent a sudden foaming, due to retarded ebullition, some small pieces of platinum wire or capillary tubes are placed in the liquid; the evaporation is also facilitated by frequent shaking. Should the vapours become ignited from the flame of the water-bath, no attempt to blow out the burning vapours should be made; but the burner is extinguished, the flask removed from the bath with a cloth, and the mouth covered with a watch-glass. Carbon disulphide, on account of its great inflammability, is never vaporised in this way, but always without a flame.

Large quantities of solvents may also be evaporated by these two methods, but the entire quantity is not treated at once. A portion is placed in a small flask, and when this has been evaporated, a second portion is added, and so on. The danger of ignition of the solvent may be avoided by inserting in the flask a glass tube, extending to within a few centimetres of the level of the liquid, supported firmly by a clamp, and attached by rubber tubing to the suction. The tube must at no time touch the liquid.

For rapid evaporation of small quantities of ether, the following method of procedure is recommended: A few cubic centimetres of the solution are placed in a sufficiently wide test-tube; this is warmed, with continuous shaking, over a small, luminous flame.

After the first portion is evaporated, the second is added, and so on. Since the vapours of the ether almost regularly become ignited, this event should always be expected, and should occasion no alarm. When it happens, the heating is interrupted for a moment, and the flame is easily extinguished by blowing on it or covering the mouth of the test-tube. If the tube is held as nearly horizontal as possible during the heating, the danger of ignition is lessened.

If it is desired to distil off a larger quantity of ether, ligroïn, or carbon disulphide, and to recover it by condensation, the receiver is attached to the condenser tube by a cork, and the flask is heated by immersing it in a water-bath containing hot water. To prevent the liquid from being superheated, a silk thread as frayed as possible at the end reaching to the bottom of the flask is suspended from the neck and the flask is shaken frequently during the distillation (Fig. 28). The entire quantity of the liquid is not placed in the flask at once, but only a portion: after the solvent has been distilled off from this, another portion is added, and so on.

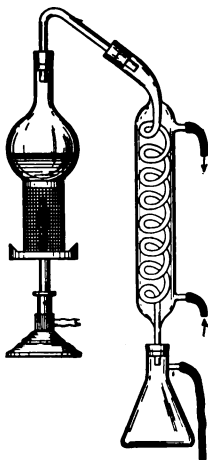


FIG. 27.

By the use of the so-called safety water-bath, *i.e.* one in which the flame is surrounded by a wire gauze as in Davy's Safety Lamp, ether and ligroïn can be distilled by continuous heating with a flame. It is not safe to distil off carbon disulphide even from this apparatus, since, when it becomes sufficiently hot, it will ignite spontaneously without the intervention of a flame.

By the use of a coil condenser (Fig. 27) the distillation of solvents is greatly facilitated. The free flame, if it be surrounded by a cylinder of wire gauze, may be employed in place of a water-bath.

A piece of rubber tubing attached to the side tube of the receiver carries the vapours to a hood or below the surface of the table.

The apparatus best adapted to distilling off any desired quantity of ether is represented in Fig. 29. A fractionating flask, into the neck of which a dropping-funnel is inserted, is connected with an ordinary condenser or an upright coil condenser. During the heating by means of hot water, or in special cases, the water-bath may be heated with a flame, or the flask may be heated directly by a flame protected by a safety gauze, the ethereal solution is allowed to flow gradually from the dropping-funnel into the flask in the bottom of which are a few scraps of platinum, or pieces of unglazed porcelain.

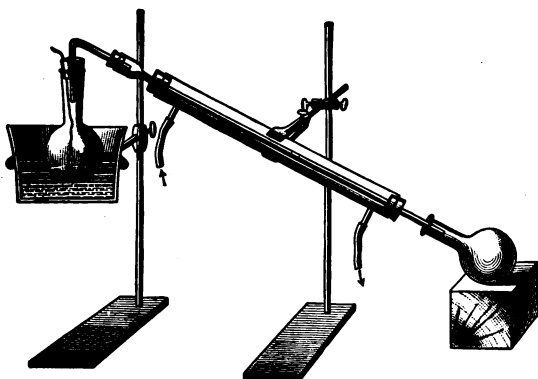


FIG. 28.

If the flow of the solution is regulated so that the same quantity of liquid is added as that distilled, the operation may be carried on continuously for hours. The quantity of ether collected in the receiver is prevented from becoming too large, by pouring it into a larger vessel from time to time. To protect the ether from ignition, the mouth of the receiver is closed by a loose plug of cotton, or the receiver, united to the condensing tube by a cork, is connected with the hood by rubber tubing. Besides its convenient manipulation, this method possesses the further advantage that after the completion of the distillation the dropping-funnel may be replaced by a thermometer, and the residue can be distilled directly from the fractionating flask. This is an especially eco-

nomical procedure when the quantity of the dissolved substance is small. In a case of this kind the size of the flask is selected with reference to the residue that may be expected. In distilling off alcohol, it is necessary to heat the water-bath continuously, and to always use threads. The distillation may be hastened by placing the flask not upon, but *in*, the water-bath. If a solution of common salt is employed in the bath, the temperature is raised, and the distillation proceeds still more rapidly.

Cylinders that fit in the water-bath, and are perforated at the bottom and on the sides, may be used with great advantage in place of the simple ring covers. This device permits the heating of the distillation flask with steam, not only at the bottom, but also on the sides.

If one has had sufficient experience in laboratory work, alcohol

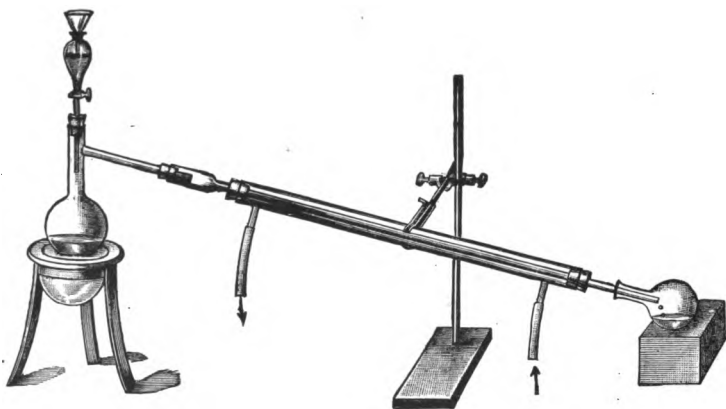


FIG. 29

may be distilled off by heating the flask on a wire gauze or sand-bath over a flame. In this case especial care must be taken not to use too large quantities at one time. Benzene can be distilled off under the same conditions as alcohol. The methods applicable to high boiling liquids have been given under "Distillation." (See page 16.)

In comparatively few cases the difference between the boiling-points of the solvent and the dissolved substance is a slight one; under these conditions the separation must be effected by a systematic fractional distillation with the aid of fractionating apparatus.

DISTILLATION WITH STEAM

A particular kind of distillation, very frequently employed in organic work for the purification or separation of a mixture, is distillation with steam. Many substances, even those distilling far above 100° , or those not volatile without decomposition, possess the property, when heated with water, or when steam is passed over or through them, of volatilising with the steam. This phenomenon is explained as follows: Suppose we take a mixture of two liquids that are absolutely insoluble in one another. Each liquid will exert its own vapour pressure as it would if it were alone, and will not in any way be influenced by the other. A practical example of this, to be taken up later, is a mixture of water (B. P. 100°) and brombenzene (B. P. 155°). When this is gradually heated, the vapour pressure of both substances will increase, and boiling will begin when the sum of the vapour pressures is equal to the barometric pressure, which we shall assume to be 760 mm. The mixture will boil at 95.25° , as will be seen from the following table:

t	Vapour pressure ¹ of C_6H_5Br	Vapour pressure of H_2O	Total
95°	120 mm.	634 mm.	754 mm.
95.25°	121 mm.	639 mm.	760 mm.
96°	124 mm.	657 mm.	781 mm.

Thus at this temperature a mixture of water and brombenzene distils over. The proportion of the two substances will be seen from the following considerations: According to Avogadro's Law,

¹ The vapour pressure of brombenzene is calculated by interpolation from the values found by Young at 90° and 100° . (Journ. Chem. Soc. 55, p. 486.)

under the same conditions of temperature and pressure equal volumes of all ideal gases contain the same number of molecules. If temperature is the same and pressure different, the number of molecules in the same volume will be proportional to the pressure. Let us now consider the mixture consisting of the vapours of water and brombenzene at 95.25° . Since at this temperature the former exerts a vapour pressure of 639 mm., and the latter a vapour pressure of 121 mm., the molecular quantities must be in the same ratio, *i.e.* for every 639 molecules of water 121 molecules of brombenzene will be present. In order to calculate the weights of the substances that distil over, we must multiply the number of molecules with the molecular weight of each substance. In our example, for 639×18 parts by weight of water, 121×157 parts by weight of brombenzene will distil over, which corresponds approximately to 3 parts by weight of water and 5 parts by weight of brombenzene. This proportion remains constant until one of the two has completely distilled over.

It must be stated that such an ideal condition of distillation with steam is never realised. There are no substances that are absolutely insoluble in one another. Furthermore, there is the disturbing influence of vapour pressure, which is quite small in the example given above. And again, since vapours do not strictly obey Avogadro's Law, the ratio of the two substances in the distillate is somewhat irregular. Finally, since the heating is carried out by steam, and the temperature of the substance is never raised to the exact boiling point, the mean temperature is somewhat different from that given above, and the proportions are therefore altered.

Apparatus.—The apparatus used for distillation with steam is represented in Fig. 30. A round flask inclined at an angle is closed by a two-hole cork; through one hole passes a not too narrow glass tube reaching to the bottom and serving to lead in the steam; the other hole bears a short glass tube the end of which is just below the cork, the other end is connected with a long condenser. The distillation flask selected is of such a size that the liquid fills it not more than half full.

In order that the steam may act on an oil at the bottom of the flask, the inlet tube is bent so that it may reach the lowest point of the flask. Steam is generated in a tin vessel about half-filled with water, the neck being closed by a two-hole stopper; into one hole is inserted a safety-tube partially filled with mercury; the lower end of this tube does not touch the water; through the other hole passes the outlet tube bent at a right angle.

Method of Procedure.— The experiment is begun by heating the steam generator and the flask simultaneously, the former conveniently by means of a low burner (Fletcher burner). The flask may be heated on a wire gauze over a free flame; but since at times a very troublesome “bumping” will occur, it is better, if this happens, to heat on a briskly boiling water-bath. As soon as the water in the generator boils and the liquid in the flask has been heated to the proper point, the tubes of the two vessels are connected with rubber tubing. The distillation is then continued until the condensed steam passes over unaccompanied by any of the substance. Should the steam escape from the safety-tube, the generator is being heated too strongly, and the flame should be lowered. To prevent the partial condensation of vapours in the upper cool part of the flask this should be covered with several layers of thick cloth to lessen the radiation of heat. If the quantity of substance to be distilled is small, so that only a small flask need be used, the preliminary and continued heating of the latter is superfluous: the steam can be passed at once into the cold liquid. If a compound is very easily volatile with steam, the introduction of the latter may be omitted; in this case it is only necessary to mix the compound with several times its volume of water and distil directly from the flask. If the substance to be distilled is solid, and its vapour forms crystals in the condenser, these may be removed provided the substance melts below 100° , by drawing off the water in the condenser for a short time. The substance is melted by the hot steam and flows into the receiver. If after this operation the water is to be turned into the condenser again, it must be done slowly at first, otherwise the cold water

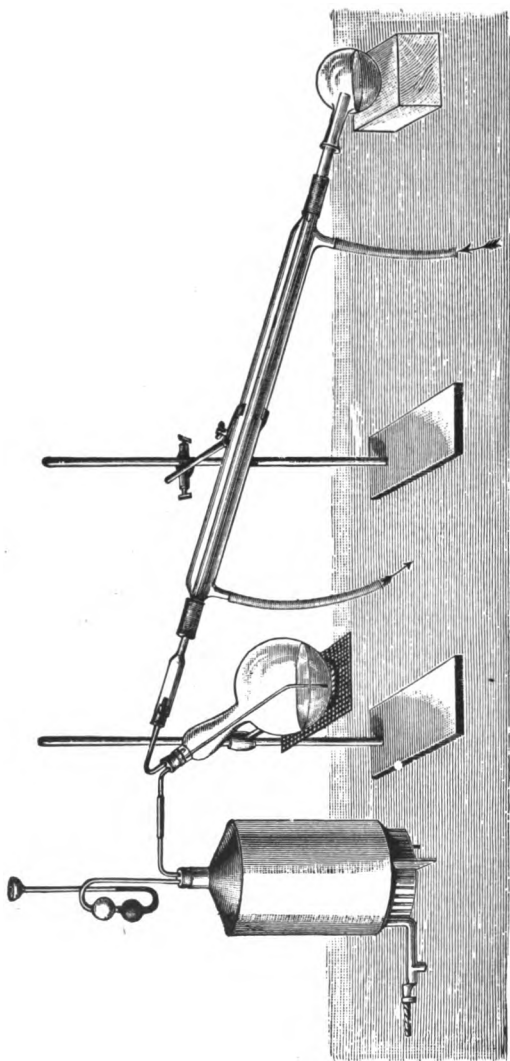


FIG. 30.

coming in contact with the hot condenser may easily crack it. When the melting-point of the substance is above 100° , in order to keep the condenser free from crystals, the distillation is interrupted for a short time, and the crystals are pushed out of the tube by a long glass rod.

The end of the operation is indicated, if the substance is difficultly soluble in water, by the fact that the water passing over carries no drops of oil or crystals with it. But when the substance is soluble, even though the condensed water is apparently pure, it may still contain considerable quantities of the dissolved substance. In this case, to determine when the end of the operation has been

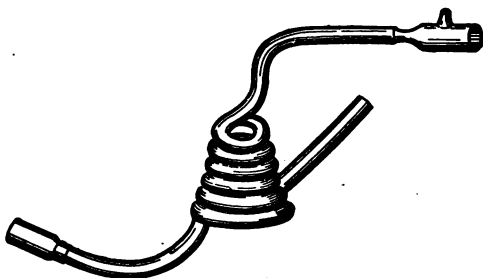


FIG. 31.

reached, a small quantity, about 10 c.c., is collected in a test-tube, shaken up with ether, the ether decanted and evaporated. If no residue remains, the distillation is finished. When a substance shows a colour reaction, *e.g.* aniline with bleaching powder, advantage is taken of this to decide the question. After the distillation is ended the rubber tubing is first removed from the distilling flask, and then, *after this has been done*, the flame under the generator is extinguished. This point is also carefully observed when the distillation is interrupted; otherwise it may happen that the contents of the flask will be drawn back into the generator.

Superheated Steam.—In dealing with very difficultly volatile compounds, it is frequently necessary to conduct the distillation

with the aid of superheated steam. A conically wound copper tube is interposed between the steam generator and the distilling flask (Fig. 31). In order to superheat the steam, a large burner is placed under the spiral in such a position that the flame comes in contact with the interior of the spiral. Since for many purposes it is necessary to use superheated steam at a definite temperature, a small opening is made in the steam exit and a piece of metal tubing affixed to it. A thermometer is inserted into this tubulure and held in position by means of asbestos twine. The distillation is still further facilitated by heating the flask to a high temperature in an oil- or water-bath. Under these conditions the substance to be distilled is not covered with a layer of water.

The apparatus represented in Fig. 31 A was proposed by V. Haehl and Co. of Strassburg. It is made in different sizes,

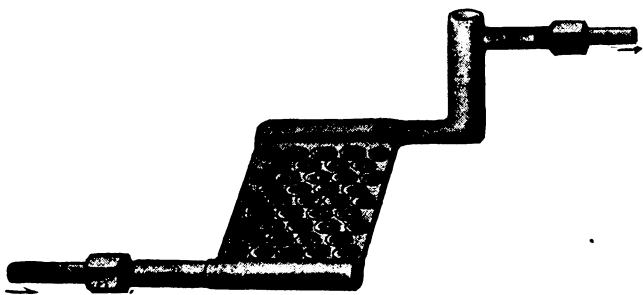


FIG. 31 A.

and may be used with great advantage in the laboratory and for work on the large scale. It consists of a hollow filter-plate, surrounded with a double layer of thick asbestos board. The latter is provided with a large opening in the lower half for the heat of the flame. A number of smaller openings are found in the upper half, which permit the escape of the hot gases. The superheating of the steam is rendered very efficient by the passage of the hot gases through the numerous canals.

SEPARATION OF LIQUID MIXTURES. SEPARATION BY EXTRACTION. SALTING OUT

Separation of Liquids. — If the separation of large quantities of two non-miscible liquids, one of which is for the most part water or a water solution, is to be effected, it can be done with a separating funnel (Fig. 32). If the liquid desired is of a greater specific gravity than that of water, it is allowed to flow off through the stem by opening the cock. If, however, it floats upon the water, the latter is first allowed to flow off, and the liquid remaining is poured out of the *top of the funnel*. By this manipulation the liquid is prevented from coming in contact with the portion of water remaining in the cock, and adhering to the sides of the stem. For the separation of small quantities of liquids a small separating funnel, the so-called dropping funnel, is employed. If the quantity of liquid is so small that even a dropping funnel is too large, a capillary pipette is used (Figs. 33 and 34). The mixture to be separated is placed in a narrow test-tube, the pipette is immersed in the mixture almost to the surface of contact of the two liquids in case the upper layer is to be removed, the test-tube is brought to the level of the eyes, and the upper layer drawn off. The tubing is then closed by pressure with the teeth or fingers and the pipette removed from the tube. If the lower layer is the one desired, the pipette is immersed to the bottom of the test-tube, and the operation conducted as before. Pipettes of this kind are very easily made from glass tubing; and if one is accustomed to work with them, are indispensable.

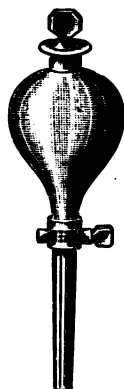


FIG. 32.

Separation by Extraction. — When a substance is held in suspension or dissolved in a liquid, generally water, the removal of the dissolved substance may be effected by agitating the solution with another solvent which will more readily dissolve the substance, but which is not miscible with the first liquid, drawing off this and

distilling it. For extraction, ether is generally used ; in special cases carbon disulphide, ligroïn, chloroform, benzene, amyl alcohol, etc., may be used. In the discussion following it will be assumed that the extraction is made with ether.

If the liquid to be separated is insoluble in water and is present in such a small quantity that a direct separation would cause a loss owing to the adhesion of the liquid to the walls of the vessel, or if it is held in suspension by the water in the form of individual

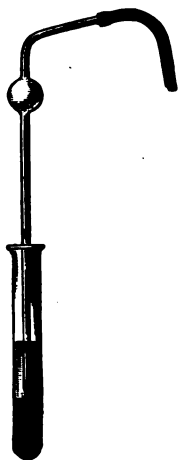


FIG. 33.



FIG. 34.

drops, ether is added to the mixture ; it is then shaken, allowed to stand, the two layers separated, and the ether evaporated.

A similar method is followed if the substance is completely soluble in water, or if a portion of it is soluble and the remainder held in suspension. If from an aqueous solution a considerable portion of a solid or liquid separates out, it is not immediately extracted with ether, but if the substance separating out is a solid, it is first filtered off ; if a liquid, the oily layer is separated in a dropping funnel, and then the filtrate is

extracted with ether. In many cases, *e.g.* when the layer of oil is very turbid, or the solution strongly acid or alkaline, thus rendering filtration difficult, the entire solution may be treated at once with ether. With substances soluble in water the extraction must be repeated one or more times in proportion to the greater or less solubility of the substance. If it is desirable to avoid using large quantities of ether, after the first extraction it is distilled off and used for the second extraction, and so on. The extraction is repeated until a test portion of the ethereal solution, evaporated on a watch-glass, leaves no residue. If the test portion is evaporated by blowing the breath on it, the beginner will probably often be deceived by the appearance of an abundant quantity of crystals of *ice*, formed by the condensation of the moisture of the breath, due to the cold produced by the evaporation of the ether. Another error into which the beginner often falls when extracting with ether is this: in most chemical processes small quantities of coloured impurities are formed; on extraction, these impart a colour to the ether. The tendency is to continue the extraction until the ether is no longer coloured; but the proper method of procedure is just the reverse of this. On extracting colourless compounds, the colouring of the ether does not show that it still contains any of the substance dissolved; the above-mentioned test gives the only safe indication of the presence of the substance.

In an extraction the following points are to be observed: Hot liquids are allowed to stand until they are *cold* before they are extracted with ether, carbon disulphide, or other low boiling solvents. It frequently happens that, after long standing, two layers of liquids will not separate, in consequence of the formation of a flocculent precipitate floating in the liquids at the surface of contact. This difficulty may be obviated either by stirring the liquids with a glass rod, or by giving the separating funnel a circular motion in a horizontal plane, in such a way as not to cause the two layers to be mixed by the shaking. Under certain conditions the neck of the funnel may be closed with a cork bearing a glass tube attached to the suction; by this means the space over the liquid is exhausted. The bubbles of gas which now rise through the liquid frequently

destroy the troublesome emulsion. The same object may also be attained by adding a few drops of alcohol to the ether. If the separation of the layers is very imperfect, the cause is often due to the fact that an insufficient quantity of ether has been used ; in this case more ether is added. If all of these methods prove ineffectual, then a complete separation may be obtained by first filtering the mixture, best with the aid of a Büchner funnel, which will retain the precipitate causing the emulsion, and then allowing it to stand.

Occasionally when extracting an aqueous solution containing inorganic salts with ether, they will separate out as solids. In this case water is added until the salts are redissolved, or the solution is filtered, with suction.

If the specific gravity of an ethereal solution is approximately the same as that of the water solution, the separation often takes place only with difficulty. Some common salt is then added, by which the specific gravity of the aqueous solution is increased.

Under certain conditions both the ethereal and aqueous solutions are so coloured that they cannot be distinguished. In this case the separating funnel is held toward the light ; in the evening a luminous flame is placed behind it, and the eye is directed to the liquid at a point just above the cock. On opening the cock, the eye will readily detect the layer separating the two liquids, as it approaches the opening.

Theory of Separation by Extraction. — Suppose we take a mixture of two immiscible solvents 1. and 2., and add to this a substance that has the same molecular weight in both liquids, and is furthermore twice as soluble in 1. as in an equal volume of 2. If we use a quantity of the substance that will saturate both solvents, then in a unit volume of 1. we shall have twice as much in solution as we have in a unit volume of 2., and the concentrations will be in the ratio of 2 : 1. If to both solvents in a vessel we add a quantity of the substance that will only form an unsaturated solution, or if we dissolve the substance in one solvent and shake this with the second solvent, we shall find that, in either case, the substance distributes itself in such a way that the ratio of the concentrations (the distribution coefficient k) is again 2 : 1 (Berthelot and

Jungfleisch). Suppose we represent the maximum concentrations by K_1 and K_2 , and let k_1 and k_2 represent the concentrations of the unsaturated solutions. We have the equation :

$$k = \frac{K_1}{K_2} = \frac{k_1}{k_2}.$$

If, for example, the amount of the dissolved substance is a , and if equal volumes of solvents 1. and 2. are taken, after the first extraction solvent 1. will contain $\frac{2}{3} a$, while solvent 2. will only contain $\frac{1}{3} a$. The greater the difference in solubility the more complete will the extraction be. When two volumes of 1. and one volume of 2. are used for extraction, the amount of a in two volumes of 1. may be calculated in the following manner :

Solvent.	Volumes.	Dissolved Substance.	Concentration, <i>i.e.</i> quotient of dissolved substance and volumes of solvents.
1.	2	x	$\frac{x}{2}$
2.	1	$a - x$	$(a - x)$

The ratio of concentrations must now be equal to the distribution coefficient, *i.e.* :

$$\frac{\frac{x}{2}}{(a - x)} = \frac{2}{1}, \text{ and hence } x = \frac{4}{5} a.$$

The principle of distribution may be generally expressed as follows :

Solvent.	Volumes.	Dissolved substance.	Concentration.	Coefficient of distribution of 1. according to 2.
1.	v_1	x	$\frac{x}{v_1}$	} k
2.	v_2	$a - x$	$\frac{a - x}{v_2}$	

$$\frac{\frac{x}{v_1}}{\frac{a-x}{v_2}} = k, \quad x = \frac{kav_1}{v_2 + kv_1}.$$

In practical work we have this question to consider : Is it expedient to carry on the extraction in one operation by a given volume of the solvent, or do this in several operations by the use of small portions of the solvent ? The question may be answered as follows : In the example under consideration, a single extraction by an equal volume of 1. dissolved $\frac{2}{3}a$. If in a second experiment we only use a half volume of 1. then x parts of a will go into solution, and x will be smaller than $\frac{2}{3}a$. The portion that remains in solution in 2. will be $(a-x)$. We therefore have the following ratio :

Solvent.	Volumes.	Dissolved substance.	Concentration, <i>i.e.</i> quotient of dissolved substance and vol- umes of solvents.
1.	$\frac{1}{2}$	x	$\frac{x}{\frac{1}{2}} = 2x$
2.	1	$a-x$	$a-x$

But the ratio of concentrations must be equal to the coefficient of distribution ; consequently :

$$\frac{2x}{a-x} = \frac{2}{1}; \text{ or } x = \frac{a}{2}.$$

One half of a is therefore in solvent 1. and the other half remains in solvent 2. If we now shake 2. with the second half of solvent 1., a portion equal to one half of $\frac{a}{2}$, namely $\frac{a}{4}$, will be extracted by the latter. The total amount extracted by using solvent 1. in two portions is $\frac{a}{2} + \frac{a}{4} = \frac{3}{4}a$, whereas the amount extracted when the solvent is used in one operation is $\frac{2}{3}a$. Thus a given quantity

of the solvent is much more efficient when it is used in small portions than it is when all of it is used in one operation. The first procedure has the disadvantage of requiring more time than the second. But since in laboratory work time is of greater consequence than the price of the solvent, large quantities of the solvent are used in one operation. — It is clear from these observations that a quantitative extraction, in the strict sense of the word, is impossible, even by a repetition of the process. Thus

$$\frac{a}{2} + \frac{a}{4} + \frac{a}{8} + \frac{a}{16} \cdots \text{cannot possibly be equal to } a.$$

The above discussions apply to solvents only which are not themselves miscible. This is not strictly true for mixtures of water and ether, since ether is soluble in water, as water is in ether.

The same formulas may be used to express the solubilities of unit weights, as well as of unit volumes of solvents. In each case the distribution coefficient will naturally have a different value.

Salting Out. — A very valuable method to induce substances dissolved in water to separate out is known as “salting out.” Many substances soluble in pure water are insoluble or difficultly soluble in an aqueous solution of certain salts; if, therefore, sodium chloride, potassium chloride, potash, calcium chloride, ammonium chloride, Glauber’s salt, sodium acetate, ammonium sulphate, or other salt is added to the solution, this is dissolved, and the substance previously in solution separates out. By this method many compounds like alcohol, acetone, etc., which are so easily soluble in water that they cannot be removed from it by extraction with ether, can be separated out with ease. The method of procedure is this: One of the above-mentioned salts, usually solid potash, is added to the solution until no more will dissolve. The substance thus forced out of solution collects above the heavier salt-solution and is removed by decantation or suction.

A combination of extraction and salting out also presents many advantages. If to the solution of a compound in water one of the salts mentioned is added — it is best to use finely pulverized sodium chloride — before the extraction with ether, this latter is greatly facilitated for several reasons. In the first place, a portion

of the dissolved substance will separate out, due to the "salting out" action; furthermore, the solubility of the substance in the the new solvent—sodium chloride solution—will be diminished so that on extracting, a larger portion is dissolved by the ether than on treating the solution directly with it, and finally, ether does not dissolve so readily in a sodium chloride solution as in water, so that the volume of the ethereal solution is larger. The amount of sodium chloride to be added is about 25–30 grammes of the finely pulverised salt to 100 c.c. of the aqueous solution. Unfortunately the method of "salting out" has not been so generally adopted in scientific laboratories as it deserves, while in the laboratories of technical chemists it has long been in daily use. Among the reagents constantly used, a bottle of solid sodium chloride should not be wanting. In many cases, instead of the salt, a concentrated aqueous solution may also be used. Concerning the salting out of electrolytes see theoretical considerations under "Benzenesulphonic Acid."

DECOLOURISING. REMOVAL OF TARRY MATTER

As is well known, animal charcoal possesses the property of being able to remove the colour from certain solutions; for this reason it is frequently employed in the laboratory to free a colourless substance from coloured impurities. If it is to be used to remove the colour of a solid substance, the latter is first dissolved in a suitable solvent, then boiled with the animal charcoal and filtered. Before treating a hot solution, it is allowed to cool somewhat, since when animal charcoal comes in contact with liquids heated nearly to the boiling-point, a violent ebullition is frequently caused, and an overflowing of the liquid may easily take place. When a solvent not miscible with water is used, the animal charcoal, which is generally moist, is previously dried on the water-bath. The solvent selected is such that upon cooling the decolourised solution, the substance will crystallise out. In carrying out this operation, the general rule that no animal charcoal is added until the substance to be decolourised has completely dis-

solved, should be followed. Under these conditions only, is it certain that a portion of the substance does not remain undissolved mixed with the charcoal. The quantity of animal charcoal to be added to a solution depends upon the intensity of the colour of the latter. To a solution very slightly coloured, a small quantity is added ; to a deeply coloured solution, a larger quantity. Very finely divided precipitates in water which pass through the filter may also be removed by the use of animal charcoal. When, *e.g.* tin is precipitated with hydrogen sulphide, the tin sulphide is often so finely divided that it runs through the filter. If the liquid is boiled with animal charcoal, the filtration presents no difficulty.

The use of animal charcoal, especially when it is in a very finely divided condition, has the disadvantage that at times it passes through the filter and contaminates the filtrate. This may be prevented frequently, by filtering again, or by boiling the filtrate a few minutes before the second filtration. When substances to be analysed have been decolourised with animal charcoal, care must always be taken to prevent the contamination of the substance. In such cases it is again crystallised without the use of animal charcoal. This difficulty may also be prevented or essentially lessened, by washing the charcoal with water several times before using ; the portion suspended in the water is decanted, and only the coarser residue which easily settles at the bottom is used.

Recently the use of animal charcoal in the sugar industry has been replaced in part by a mixture of fine wood meal and floated infusorial earth (*kieselguhr*). This mixture ought to be of great advantage in the laboratory, for decolourising purposes, if used in the same way on a small scale. To the mixture is ascribed very superior purifying properties, so that by using much smaller quantities the same effect is obtained as with far larger quantities of animal charcoal. In order to prevent an easily oxidisable liquid from decomposing when it is heated in the air — this action being generally attended with more or less colouration — a gaseous reducing or protecting agent is passed through it ; *e.g.* sulphur dioxide, hydrogen sulphide, or carbon dioxide. Very easily oxidisable sub-

stances are not evaporated in a dish, but in a flask, since in this the liquid is better protected from the action of the air.

Not only coloured impurities, but those of a tarry character, may also be removed by boiling with animal charcoal as above described. A mixture of wood meal and infusorial earth with which the solution may likewise be boiled is said to be of great value.

For the absorption of tarry impurities, in so far as they are liquid or oily, unglazed, porous plates (drying plates) may be used with advantage, the substances being firmly pressed out with a spatula in a thin layer. If one pressing out is insufficient, the substance is spread out again upon a fresh, unused portion of the plate. The absorption of an oil may often be facilitated by moistening the substance on the plate with alcohol, ether, or ligroïn, which at times will dissolve the impurities without causing a solution of the substance. Oily by-products may also be removed by pressing the substance between a number of layers of filter-paper. For this purpose either a screw-press is used, or the substance is placed in layers of filter-paper between two wooden blocks, the upper one of which bears a heavy object.

DRYING

Drying Solid Compounds.—Under the chapter on “Crystallisation,” page 9, the method of drying moist crystals has already been given. This method is naturally applicable to all solids, even if they are not crystallised, or only imperfectly crystallised, so that it will be unnecessary to repeat the directions already given. But a few methods, not so refined, and generally employed in dealing with crude products will be referred to here. Before a substance is dried by allowing it to lie in the air or in a desiccator, or by heating, the greatest portion of the moisture is removed by pressure, as follows: The substance lying between a number of layers of filter-paper is placed in a screw press, and pressure applied. The operation is repeated, and the paper renewed, until it is no longer moistened. If a solid is not contaminated by water or other

solvent, but by a liquid by-product, which one desires to obtain, the paper, after it has absorbed this liquid substance, can be extracted with a solvent, like ether. Large masses of a compound not too finely granulated can be tied up in a piece of filter-cloth of fine texture, placed in the screw press, and pressure applied. Smaller quantities of a substance may be pressed out between two wooden blocks, the upper of which bears a heavy object.

Very often solids may be dried by making use of the power of unglazed porcelain to absorb liquids with avidity. The substance to be dried is pressed out in a thin layer upon a suitable piece of an unglazed porcelain plate, with a spatula, and is allowed to stand for some time, longer or shorter, as may be necessary. If one pressing out is not sufficient, the operation is repeated, using a fresh plate. Oily and tarry impurities may also be removed in this way, as mentioned above.

Compounds which fuse without decomposition may be dried either upon the water-bath or in an air-bath, or by heating over a free flame until they melt, allowing them to solidify, and then pouring off the water.

In order to dry a substance at a high temperature in a vacuum, two glass hemispheres, the edges of which are ground and fitted together, are used. The upper vessel is supplied with a tubulure, the opening of which is closed by a cork bearing a glass tube bent at a right angle connected with suction. The sphere may be heated by immersion in a large quantity of hot water or on a boiling water-bath. The upper hemisphere is enveloped in a cloth to prevent the condensation of the vapours.

Drying Agents for Liquids. — Liquids are dried (deprived of water) either by placing in them, or in a solution of them, drying agents. The most frequently employed drying agents are :

Calcium chloride,
 (a) granulated,
 (b) fused,
 Potassium hydroxide,
 Sodium hydroxide,

Ignited potash,
Fused sodium sulphate,
Dehydrated magnesium sulphate.

Less frequently used are : lime, barium oxide, anhydrous sodium carbonate, anhydrous copper sulphate, phosphorus pentoxide, sodium, and others.

In the choice of a drying agent care must be taken to select one which will not react with the substance to be dried. For example, calcium chloride unites to form double compounds with alcohols as well as with bases. Consequently, for drying these two classes of compounds, calcium chloride is never used. Caustic potash and caustic soda, as is well known, react with acids and phenols to form salts, upon alcohols to form alcoholates, and upon esters, saponifying them. These drying agents are never used with these substances. Further, acids are never dried with carbonates, owing to the salt formation taking place.

Calcium chloride is employed in two forms, — granulated and fused. The former acts more energetically, since it possesses a larger acting surface. Still, it has the disadvantage of being more porous than the fused variety and in consequence of this porosity the loss of the substance being dried is greater. For drying small quantities of a substance, or liquids containing very little moisture, it is better, therefore, to use fused calcium chloride.

On drying bases with caustic potash or caustic soda, it must be borne in mind that these drying agents may be contaminated, at times, with potassium nitrite or sodium nitrite. Since these latter act upon bases, decomposing them, it is necessary to use the pure alkalis, or in place of them potassium carbonate or Glauber's salt.

Methods of Drying. — As already mentioned, liquids may be dried either in the undiluted form or in solution. The first method is followed when the quantity of the liquid is considerable, so that the loss of the substance necessarily incident to the adhesion of the liquid to the drying agent cannot amount to a large percentage of the whole. Low boiling liquids are always dried directly, without the use of a solvent. If a solution of a higher boiling

compound is to be dried, it is done before the solvent is distilled off. A small quantity of a substance or a viscous substance is designedly treated with a diluting agent, generally ether, and is then dried.

The drying is accomplished by placing the drying agent in the liquid and allowing the two substances to remain in contact for a longer or shorter time, according to circumstances. So long as a liquid appears turbid, it has not been deprived of its moisture. A liquid about to be dried must never contain drops of water which are visible; in case it does, it must be treated in a separating funnel or the water drawn off with a capillary pipette: it is then dried. If only a few small drops of water are present, the liquid is first filtered through a small folded filter, or it is poured carefully into another vessel, and the water drops will remain in the first vessel, adhering to the walls. When a separation of an ethereal from an aqueous solution is to be made, to prevent a portion of the water from being carried along with the ethereal solution, the former is not drawn off through the cock of the vessel, but is poured out of the mouth, as has already been mentioned.

If a liquid contains very much moisture, and this is the case especially in turbid, milky liquids, it frequently happens that the drying agent will absorb enough water to dissolve itself and thus form an aqueous solution. In this case a fresh quantity of the drying agent is not added at once, but the separation of the two layers is effected by a separating funnel, pipette, or by decanting one of the layers.

A similar rule obtains for undiluted liquids. The drying of high boiling substances, if they are not volatile at the temperature of the water-bath, may be greatly facilitated by heating them with the drying agent on the water-bath.

If the boiling point of a liquid is above 200° , it may be freed from water, without the aid of a drying agent, by heating for some time on a water-bath under diminished pressure. The water will thus distil over. The apparatus represented in Fig. 20, page 27 is employed for this purpose. The use of a manometer will be unnecessary if this is connected with a good suction pump.

Before distilling a liquid which has been dried, or before distilling off the solvent from such a liquid, it is poured off from the drying agent. To obtain the small portions which adhere to the latter it may be washed with a small quantity of the dried solvent. Low boiling individual liquids (boiling on water-bath) can, under certain conditions, be distilled without a previous separation from the drying agent. If the liquid to be dried is of such a specific gravity that the drying agent will float in it, then, in order to effect the separation, it is poured through a funnel containing a small quantity of glass wool, or asbestos. In some cases, which, however, are rare, a liquid not easily volatile may be dried by exposing it in a dish as shallow as possible in a partially exhausted desiccator.

FILTRATION

While in analytical operations it is much more desirable to conduct filtrations without employing pressure, the precipitates obtained in organic preparation work are filtered with pressure whenever it is possible. The method presents a number of advantages: the filtration may be made in a much shorter time; the liquid may be much more completely separated from the precipitate, in consequence of which the latter will dry more rapidly, etc.

The student has already learned the methods of filtering without pressure in the operations of analytical chemistry, but he is advised to reread the chapter on Crystallisation (see page 1).

Filtration with Suction. — For filtering under pressure (suction), a filtering flask *a* (suction flask) with a side-tube *b* (Fig. 35) is used. An ordinary flask may be converted into a suction flask by fitting to it a two-hole rubber stopper; through one hole is passed the stem of a funnel, through the other a glass tube, bent at a right angle, one end of which passes just through the cork,

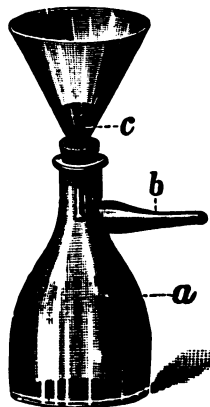


FIG. 35.

while the other is attached to the suction. For this purpose flasks with thick walls are selected, in order that they may not be crushed by the atmospheric pressure on exhaustion; if a thin-walled flask is used, it must be exhausted but slightly.

The funnel used is, in many cases, the ordinary conical glass form, in which is placed the filter. If the funnel is imperfect in construction, and does not possess the correct angle (60°), the filter is made narrower or wider, as the case may be, to accommodate it to the angle of the funnel. In order that the point of the filter not supported by the glass walls of the funnel, may not tear on exhaustion, a platinum cone *c* is previously placed in the funnel.

If a platinum cone is not at hand, it may be replaced by a conically folded piece of parchment paper or filter-cloth. The filter is moistened with the same liquid which is to be filtered, otherwise it may happen that the filtration is prevented, or, at least, rendered difficult; *e.g.* if the filter has been moistened with water, and an alcoholic solution is to be filtered through it, the substance dissolved in the alcohol may be precipitated in the pores of the filter by the water. If a liquid foams excessively on filtering, as happens at times with alkaline liquids, the rubber tubing is removed suddenly from time to time from the filter-flask. The pressure of the in-rushing air destroys the bubbles. The foaming may also be prevented at times by treating the filtrate with a few drops of alcohol or ether. This is one of the common methods of preventing foaming in general. When filtering very small quantities of a liquid, a test-tube is placed in the filter-flask, as is represented in Fig. 36.

The suction surface may be increased by placing a so-called filter-plate of glass or porcelain in the funnel (Fig. 37). If a filter-plate is used, the filter-paper should be of two thicknesses. Upon the plate is first placed a round filter of exactly the same size as the plate, and upon this another round filter, the edge of which projects about 2–3 mm. beyond that of the plate.

The Büchner funnel is indispensable in working with organic substances. In consequence of its large suction surface, a very rapid filtration is possible. In the filtration of large quantities of substance it should always be used (Fig. 38).

A double filter (described above) may be used in this, but in most cases a single filter is sufficient. Since the Büchner funnels are made of porcelain, and consequently are opaque, they must be carefully cleaned immediately after using.



FIG. 36.

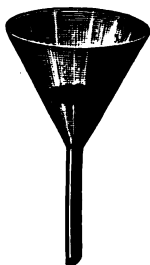


FIG. 37.



FIG. 38.

Similar to the Büchner funnel in its construction and action is the so-called "Nutsch" filter. This consists of a shallow dish with a perforated bottom, which is fitted to the cover of a tubulated cylinder by means of a rubber ring, the joint being air-tight (Fig. 39).

If the solution to be filtered acts on the filter-paper, filter-cloth may be used in its place. A fine or coarse meshed cloth is selected, according to the nature of the precipitate; it is moistened before the filtration.

If this is also attacked, nitrocellulose cloth may be used; it is made by treating a cloth woven from plant fibres with a mixture of nitric and sulphuric acids. Concentrated sulphuric acid may be filtered through

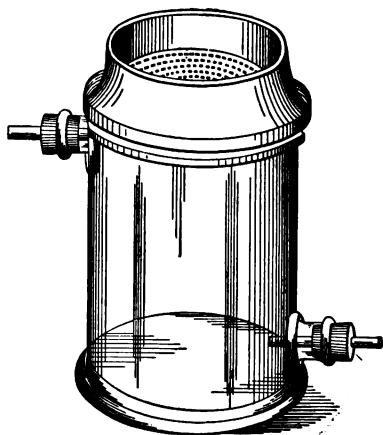


FIG. 39.

it. Such cloth and other substances containing nitrocellulose must always be preserved under water, on account of their explosiveness.

In cases of this kind the precipitate is retained by using glass wool, or better, long fibrous asbestos, with which the bottom of the funnel, containing in this case a platinum cone, is filled, or it is spread out in thin layers over a filtering plate, or on the surface of a Büchner funnel. Under these conditions, the suction is applied gently at the beginning of the filtration; as soon as a large quantity of the precipitate has accumulated, the suction is increased. Recently acid- and alkali-proof filter disks have also been used in the arts. By the aid of asbestos linings these may be placed in conical or Büchner funnels, and made to render excellent service. Very coarse-grained precipitates can be filtered without the use of a filter by placing in the point of an ordinary glass funnel a sphere of glass (a marble); this is surrounded by glass wool, or asbestos, if necessary.

Pukall Cells. — For the filtration of precipitates, like calcium sulphate, barium sulphate, of strongly acid liquids, etc., Pukall's

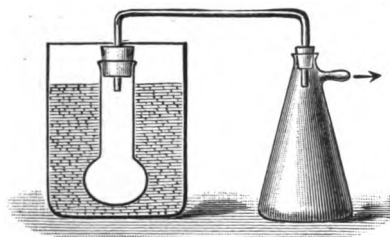


FIG. 40.

cells (porous cells), made of unglazed clay, are very useful. They may be procured in different sizes in the market, and possess either the form of a cylinder or a mortar pestle. The operation is performed as follows: In the mouth of the cell is placed a closely fitting stopper bearing

a glass tube bent twice at right angles, connected with a filter-flask. The tube at each end projects slightly below the stopper (Fig. 40). The cell is now immersed in the liquid to be filtered, contained in a beaker, not too wide, until it almost touches the bottom. When the suction is applied, the liquid filters through the porous walls until the cell is filled, and is then drawn into the flask; the precipitate remains behind in the vessel, and for the most part is deposited on the exterior walls of the cell.

Filter-Press. — For the filtration of large quantities of substances which filter with difficulty, especially dye-stuffs, barium sulphate,

calcium sulphate, etc., filter-presses are often used, of which the Hempel form will be described (Fig. 41). The separation of the liquid from the precipitate is effected in the cell *c*, which consists of two perforated porcelain plates between which is a rubber ring. The first operation in working with the press is the preparation of the cells. Two circular pieces of filter-cloth and two of filter-paper the same size as the plates are cut; after the cloth (linen or muslin) has been thoroughly moistened with water, the cells are made as follows: At the bottom comes the perforated plate upon which is placed one layer of the filter-paper, and upon this the cloth. After a wide glass tube *g*, which extends almost to the opposite side of the cell, has been inserted into the opening of the rubber ring, this is placed upon the cloth, then follows the other piece of cloth, filter-paper, and finally the second plate. The cell is now secured by three clamps, one of which is attached near the glass

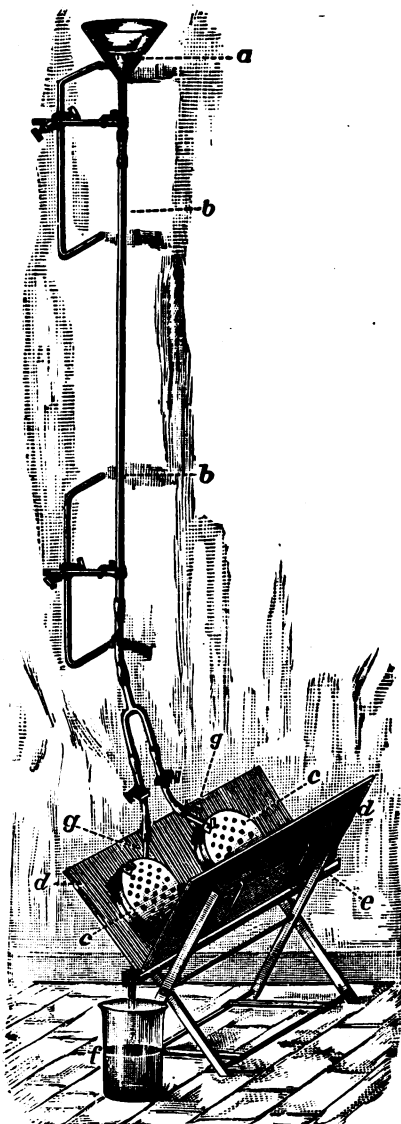


FIG. 41.

tube, and the others equally distant from this. The cell is now ready for the filtration, and is placed between the two corrugated glass plates *d*. Before it is connected with the vertical tube *b*, the pinch-cock on this is closed, water is poured into the funnel *a*, and the cock is now opened until the vertical tube is filled. The cock is again closed and the tube is connected with the cells, the liquid to be filtered poured into the funnel and the cock opened. During the resulting filtration care is taken to keep the funnel partially filled so that the vertical tube is constantly full. If the first portions run through turbid, they are returned to the funnel. In order to wash the precipitate collecting in the cell, the glass tube passing through the rubber ring is partly withdrawn, so that it projects into the cell but a few centimetres. This causes a canal to be formed in the cell from which the wash water can permeate the precipitate in all directions. If the precipitate is large enough to completely fill the interior space of the cell, it forms a solid cake that can be removed without difficulty. But if the precipitate is small, and it is desired to obtain it, the glass tube is withdrawn from the rubber ring, the contents of the cell,

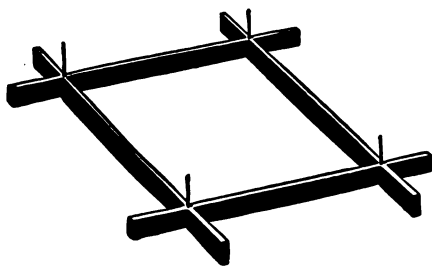


FIG. 42.

generally half-fluid, are poured into a beaker, the cell taken apart, and the precipitate adhering to the sides scraped off with a spatula. By filtering with suction a complete separation of the liquid and precipitate is effected. If it is

desired to filter larger quantities of a precipitate than can conveniently be done in a single cell, two cells connected by a Y-tube may be used.

Filtering through Muslin.—Precipitates which are not too finely divided may be filtered off through a filter-cloth (muslin) stretched over a wooden frame (filter-frame) (Fig. 42). A square piece of muslin or linen, after being thoroughly moistened, is

fastened on the four nails of the frame in such a way as to cause a shallow bag in the middle. The frame is placed over a dish of the proper size and the liquid to be filtered is poured on the cloth and generally filters rapidly through it. If it is desired after washing the precipitate to press it out, the cloth is taken from the four corners, folded together, and squeezed with the hands. The precipitate may be further dried, by tying up the opening of the bag with twine, and then pressing it out carefully under a screw-press.

HEATING UNDER PRESSURE

Sealed Tubes. Method of Filling. — If it is desired to induce a reaction between two substances at a temperature above their boiling-points, they are generally heated in sealed tubes. If a quantitative determination is not to be made, if the substances to be heated do not attack the glass or generate no gases, and if the heating is not to be high, soft glass tubes may be used. But generally, and in quantitative determinations always, difficultly fusible tubes of potash glass are used, since they are not so easily acted upon and do not crack so readily as the former. In filling the tubes the following points are observed. The tube is dried before placing the substance in it. Never put solid or liquid substances directly in the tube, but with the aid of a funnel-tube which should be as wide as possible when the substance is a solid. In proportion to the temperature of the heating a greater or less pressure is developed; therefore more or less of the substance is placed in the tube, depending on the conditions. The tube is never more than half-filled. Easily volatile substances as well as those giving off vapours, like hydrochloric and hydriodic acids, which render the sealing of the tube difficult, are transferred to the tube just before the sealing is to be done. In withdrawing the funnel-tube care is taken to avoid bringing it in contact with the walls of the tube.

Sealing. — To seal the open end of a tube charged with the substance, it is warmed by holding it at an angle of about 45° , with constant turning, in the small luminous flame of a blast-lamp, and then heated strongly in a larger non-luminous flame; when the glass becomes soft, a previously somewhat warmed glass rod is fused to it (Fig. 43, I). The flame is then applied to the tube at a short distance from the opening, and as soon as the glass has become soft the tube is narrowed by drawing it out suddenly (II). After breaking off or cutting off the end of the capillary tube at a , to allow the air to escape on further heating, it is heated at b , when the tube is softened at this point it is drawn out slightly,

the heat is applied just below *b*, it is drawn out again, and so on; the result is that the form of the end of the tube gradually changes from a cylinder to a sharp-pointed cone. The narrowest part of the latter is then heated with a not too large flame without drawing it further. The soft glass melts together, and there is thus obtained a thick-walled capillary tube which is melted off at the proper place (III). Figure 44 shows the sealed portion of a tube in its natural size. In the formation of

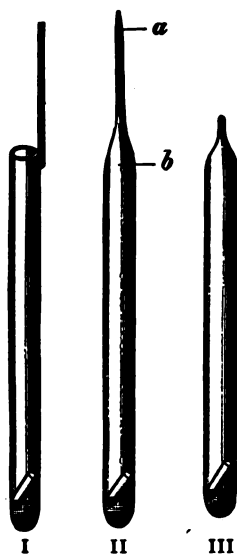


FIG. 43.

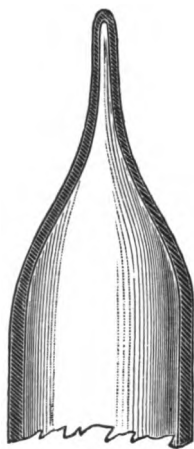


FIG. 44.

the capillary portion, it is desirable not to turn the tube in the manner previously directed, but to give it a few turns in one direction and then to reverse the motion, otherwise a spiral would be formed owing to the smallness of the glass at that point. After sealing, the heated portion is cooled gradually by holding it in the luminous flame until it is blackened. The sealing of hard glass tubes may be facilitated by placing a brick or tile near the flame in such a position that the heat will be reflected. If one

is in possession of a cylinder of oxygen, it may be attached to the blast-lamp in place of the blast. At the high temperature of the illuminating gas-oxygen flame, the sealing may be effected with great ease.

In many cases the operation is rendered difficult by the vapours of the substance attacking the glass, or by the decomposition of the substance with the evolution of troublesome products like carbon, iodine, etc. Under these conditions, the tube is not drawn out first to a narrow tube, as above, but the glass rod fused on is allowed to remain, and this is used to draw out the tube. The sealing is rendered less difficult by allowing the air to have free access to the tube, in order that the evolved vapours may pass out unimpeded. The separation of carbon may be avoided by having an assistant direct a continuous current of air, during the heating, through a narrow tube into the upper part of the tube being sealed; this will cause the oxidation of the carbon. When dealing with very volatile substances, during the sealing the lower part of the tube is cooled by water, ice, or a freezing mixture. In this case, the services of an assistant will be needed to give to the vessel containing the cooling agent a circular motion corresponding to that of the tube. Under these conditions, it is often advisable to narrow the tube before charging it with the substance, so that it will just admit a funnel-tube as narrow as possible.

Heating. — The heating of sealed tubes (bombs) is conducted in the so-called "bomb-furnace," of which a convenient form is represented in Fig. 45. To be able to carry out the operation of heating at a definite temperature, a cork, covered with asbestos paper, bearing a thermometer, is fitted into the opening at the top of the furnace. The bulb of the thermometer must be about 1 cm. above the bottom of the iron tube. The sealed tube is not heated directly, but in a thick-walled protecting case of iron closed at one end, in which the glass tube is so placed that the capillary portion is at the open end. In transferring the glass tube to the iron casing the latter is not held vertically, but is slightly inclined from the horizontal, so that the glass tube may

not be broken by suddenly striking the bottom. The iron case is pushed into the furnace open end first, so that in case of an explosion the fragments of glass are not thrown out of the forward end but from the rear of the furnace, directed toward a wall. A "fragment cage" renders the flying pieces of glass harmless. After the tube is in position the front opening is closed by a "drop-slide." The tubes are not heated at once up to the desired temperature, but are warmed gradually. If it is desired to heat a furnace similar to the one represented to a low temperature, the gas tubes are raised and small flames used, rather than a lowering of the gas-tube and the corresponding increase in size

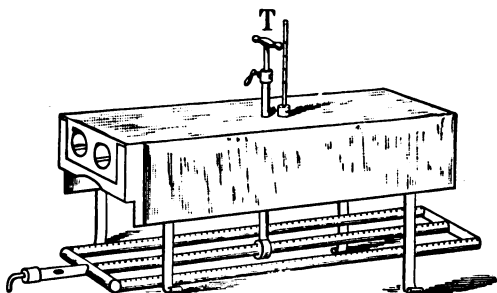


FIG. 45.

of flame. The danger of the bursting of the glass tubes may be diminished in many cases, particularly in those in which a very high pressure is developed, by interrupting the heating after a certain length of time, opening the capillary after the tube has completely cooled, and allowing the gases which have been generated to escape. The tube is then resealed and heated again.

If tubes are to be heated not higher than 100° , the convenient so-called "water-bath cannon" is used, in which the case enclosing the tube is heated by steam at ordinary pressure; in this case overheating is impossible.

Opening the Tubes.—*Sealed tubes must not be opened until after they are completely cold. The protecting case of iron, containing the tube, is removed from the furnace and held in a slightly*

inclined position, the end of the capillary being higher than the rear end. By means of a slight jerk the capillary end of the glass tube is caused to project from the iron case. The extreme end of the capillary is now held in the flame of a Bunsen burner. In case there is an internal pressure in the tube, the glass on becoming soft will be blown out and the gases will escape from the opening thus made, often with such force as to extinguish the flame. If on the softening of the glass the capillary is not blown out, it may be due to the absence of internal pressure or the tube may be stopped up by some of the substance. In the latter case the substance is removed by heating. To show that there is an internal pressure the capillary is held after it has been opened near a small luminous flame; if the latter is blown out in a long thin flame sidewise, obviously there is pressure. If great pressure exists in a tube to be opened, before blowing the capillary the hand holding the iron casing is protected by a thick glove or a cloth is wrapped around the casing several times at the point where it is held, so that if the tube bursts, in consequence of the sudden diminution of pressure, and the seam of the case should be torn open, the hand is protected from injury. *In handling an unopened tube the greatest care possible must be observed. It is never removed from the iron casing to look at it or for any other purpose. On opening, it is held in such a position that neither the operator nor any one else can be injured in case of bursting.*

On heating substances with hydriodic acid and phosphorus, it sometimes happens, that the tube on being opened by a flame, explodes. In this case the explosion is due to the fact that the phosphine as well as the hydrogen evolved in the reaction have formed an explosive mixture with the oxygen of the air present in the tube. Under these conditions, the capillary is opened by snipping off the end with pincers or tongs, but in doing so the greatest care must be observed. To remove the end of the cone, it is not necessary to proceed as described below, but the end of the tube is broken directly with a blow of a hammer.

In order to break off the end of a tube after it has been opened, so that the contents may be emptied out, the procedure is as

follows: At that point of the tube where the cone begins, a well-defined file mark is made, not extending completely around the tube; this is touched lightly with the hot end of a glass rod, previously heated to fusion in the blast-flame. If the crack caused by this does not extend entirely around the tube, the extreme end of it is again touched with a hot glass rod, by which it is extended, so that the conical end may be lifted off. Instead of a glass rod, a thick iron wire, the end of which has been bent around the iron casing to a semicircle, may be used. If this is heated to redness, the file mark touched with it, and the wire turned, the end of the tube breaks off smoothly. To prevent the fragments of glass from falling into the tube (when a quantitative determination is being made), the method of procedure is this: As before, a deep file mark is made, and on each side of it, at a distance of $\frac{1}{2}$ cm., a strip of moistened filter-paper 1 cm. wide is wrapped around the tube several times. That portion of the tube between the strips is heated by a small flame, the tube being constantly turned, this causes the end to split off smoothly without splintering. If the glass does not crack at once, the heated portion is moistened with a few drops of water, and the breaking off will follow with certainty.

Volhard Tubes.—The tube described by Volhard (Fig. 46) may be used to great advantage when it is desired to heat large quantities of substances in a single tube. It consists of a wide tube to the end of which a narrower one is fused. A tube of this kind, 35 mm. in diameter and 45 cm. in length, contains about $\frac{1}{4}$ of a litre, and possesses the further advantage of being easy to seal. If on opening the tube care be taken to cut off as small a portion of the narrow end as possible, it may be used repeatedly. If, finally, the narrowed portion becomes too short, another piece of the same kind of tubing is sealed on.



FIG. 46.

Pressure Flasks. Autoclaves.—In order to heat substances under pressure at a moderate temperature which on reacting with each other evolve no gaseous products, so that no pressure due

to the reaction is developed, they are sometimes enclosed in strong-walled flasks (pressure flasks), wrapped up in a cloth and heated in a water-bath.



FIG. 47.

Very well adapted to this purpose are the common soda-water or beer bottles, of the kind represented in Fig. 47. In using them they are not immersed in water already heated, but are slowly heated with the water. The water-bath is closed by a loosely fitting cover, so that in case the bottle bursts, one may not be burned by the hot water. The flasks are not opened until after they are completely cold.

Large quantities of substances which do not act on metals may be heated under pressure in closed vessels, generally made of iron, bronze or copper (autoclaves). Such vessels are not suited for heating acid substances, but may be used for neutral or alkaline substances. In this laboratory Mannesmann tubes (without seams) are in use, one end being welded together, and the other is supplied with a screw-thread and cap. The open end is cone-shaped. The tube is closed by a threaded cap, which in section shows a cone. The cap is partially filled with lead. After the substance has been put in, the cover is screwed on as far as possible with the hand, the tube is then clamped in a vise, and the cap made fast with a wrench. The conical end of the tube is pressed into the soft lead, thus giving an excellent joint. The heating may be conducted in an oil-bath, or directly in the bomb-furnace. If the heating is to be carried beyond the point at which lead softens, a

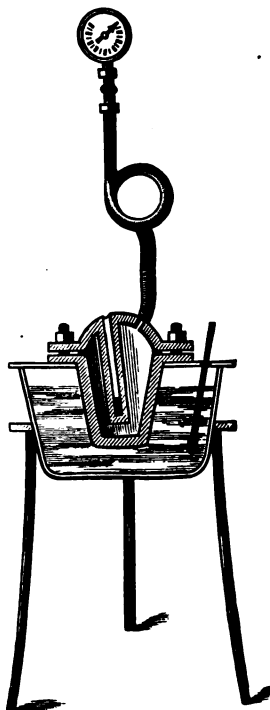


FIG. 48.

short metallic condenser about 10 cm. in length may be screwed on the threaded portion of the tube. A slow current of water is passed through the condenser.

Another form of autoclave is represented in Fig. 48. For the packing a ring of lead or asbestos is used. The tube leading to the interior is designed for a thermometer. The lower portion contains oil in which the thermometer is placed.

MELTING-POINT

In organic work the most common method of testing the purity, of characterising and of recognising a solid compound, is the determination of its melting-point. The apparatus most generally used for this purpose is represented in Figs. 49 and 50. A long-necked flask is closed by a cork provided with several canals cut in the sides, through which the heated air and vapours

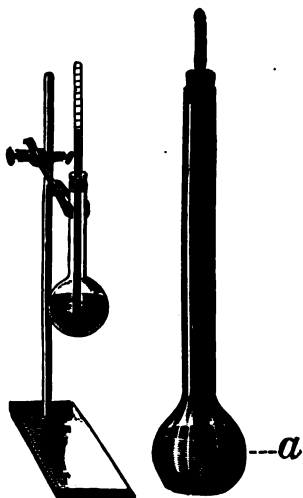


FIG. 49.

FIG. 50.

may escape, bearing a thermometer. The bulb of the flask is two-thirds filled with pure concentrated sulphuric acid, into which is dropped a crystal of potassium nitrate the size of a pin-head, to prevent it from becoming dark in colour. The substance is placed in a small narrow tube (melting-point tube), made in the following way: A glass tube 4-5 mm. wide is heated at one point while constantly turned, in a small, blast-lamp flame, until it becomes soft, and is then drawn out from both ends to a tube 1 mm. wide. The narrow tube thus produced is then fused off at its middle part; the portion lying next to that part

of the glass tube which has not been drawn out is heated as before and is again drawn out, and so on. There is thus produced a tube having the form represented in Fig. 51 *a*. In order to prepare the melting-point tube from this a file-mark is made at the points indicated, the tube broken off and fused at the narrow end by holding it nearly vertical in a Bunsen flame. Fig. 51 *b* represents the melting-point tube in its natural size. A supply of several dozen of these is made and preserved in a closed bottle. To transfer to the tube the substance the melt-

ing-point of which is to be determined, a small portion of it is pulverised, the end of the tube dipped into it; by gentle tapping the substance is caused to fall from the upper end to the bottom of the tube. In order that it may not form a too loose layer, it is packed by a thin glass rod or platinum wire. The height of the layer should be 1 mm. and in no case more than 2 mm. To attach the tube different methods may be used. The upper end of the tube may be touched with a drop of sulphuric acid; this, when brought in contact with the thermometer, will cause it to adhere. It is safer to fasten the tube, just below the mouth, to the thermometer with a thin platinum wire or a rubber ring 1 mm. wide. The substance is placed at the middle point of the thermometer bulb. The thermometer is now immersed in the sulphuric acid until the bulb is at about the centre of the liquid; the flask is heated with a free flame which is given a continuous, uniform motion as in distillation. The burner is inclined at a convenient angle, so that, if the flask should break, the hand would not be directly under it. When the melting temperature is reached, it is observed that the previously opaque, unfused substance suddenly becomes transparent and a meniscus is formed on its upper surface. If it is known at about what point the substance will melt, it may be heated rapidly to within 10° of this point, and then slowly with a small flame so that the behaviour of the substance from degree to degree can be easily observed. If the melting-point is not known, it can be readily ascertained on heating it rapidly to a high temperature. In this case the determination is repeated, heating rapidly until the temperature approaches the melting-point, and then slowly. In many cases when the temperature nears the melting-point this is shown by a softening of the substance before melt-

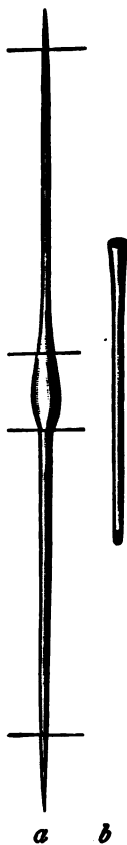


FIG. 51.

ing; it loosens from the walls of the tube and collects in the middle. If this phenomenon occurs the heating is conducted very slowly from degree to degree. At times proximity to the melting-point may also be recognised by the fact that the particles of the substance which adhered to the upper portion of the tube during the filling, melt before the mass of the substance; since the hotter and therefore lighter layers of the acid rise to the top, the upper layers of the bath are heated somewhat higher than the lower.

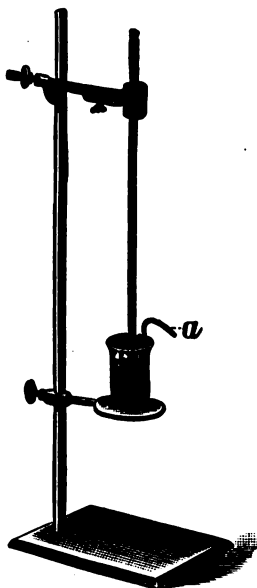


FIG. 52.

Instead of the apparatus just described the one represented in Fig. 52 serves very well for the same purpose. The liquid used may be water or sulphuric acid, depending on the melting-point of the substance to be examined, or in case of a substance with a high melting-point paraffin is placed in a beaker supported on a wire gauze. In order to keep the liquid at a uniform temperature, it is stirred by an up-and-down motion of the glass stirrer *a*.

A substance is regarded as pure in most cases, if it melts sharply within one-half or a whole degree, and if after repeated crystallisation the melting-point does not change. In determining the melting-point of a newly discovered substance, one determination is not sufficient even if it is very sharp; a small portion is recrystallised and the melting-point again determined. Many substances decompose on fusing, if this takes place suddenly at a definite temperature, this may also be regarded as a characteristic of the substance.

Since many compounds on heating decompose explosively, and since in the last few years it has happened that the explosion of minute quantities of a compound has shattered the melting-point

apparatus, and serious wounds have been caused by the hot sulphuric acid, it is safer before the melting-point of a hitherto unknown substance is determined in the apparatus described above, to take the slight trouble of making a preliminary test by heating a small tube containing the substance directly in a small flame to the melting temperature, and by this means ascertaining if the substance will explode.

Testing the Thermometer.—At this point a few observations concerning the testing and correcting of the thermometer will be added. Since the ordinary thermometers, at least the cheaper varieties, are never exact, they must be corrected before using. If a normal thermometer is at hand, the correction to be applied may be determined by slowly heating the thermometer to be tested by the side of the normal instrument in a bath of sulphuric acid, glycerol, or vaseline, and noticing the reading of both thermometers for every 10° . There is thus obtained a table from which the corrections may be read directly. For many purposes it is sufficient to determine the deviation at only a few points; the corrections for the degrees lying between these may be calculated by interpolation. Thus, *e.g.*, the point to be considered as the true zero point may be determined as follows: A thick-walled test-tube of about $2\frac{1}{2}$ cm. in diameter and 12 cm. in length is one-third filled with distilled water. The mouth is closed by a cork bearing a thermometer dipping into the water. Through an opening cut out of the side of the cork is introduced a thick copper wire, the end of which is bent into a circle at a right angle to its length. The test-tube is surrounded by a freezing mixture of ice and salt. The water is frequently agitated with the stirrer; the temperature at which crystals first begin to form is carefully noted.

The true 100° point is found by placing distilled water in a not too small fractionating flask and determining the boiling-point of it, the entire column of mercury being in the vapour. In an analogous manner, the boiling-point of naphthalene (218° at 760 mm. pressure) and of benzophenone (306° at 760 mm. pressure) may serve for the correction of the higher degrees. Since the boiling-point is influenced by the pressure, the barometer must be read at

the same time with the thermometer and a correction, taken from the table given below, applied.

Pressure.	Water.	Naphthalene.	Benzophenone.
720 mm.	98.5°	215.7°	303.5°
725	98.7	216.0	303.8
730	98.9	216.3	304.2
735	99.1	216.6	304.5
740	99.3	216.9	304.8
745	99.4	217.2	305.2
750	99.6	217.5	305.5
755	99.8	217.8	305.8
760	100.0	218.1	306.1
765	100.2	218.4	306.4
770	100.4	218.7	306.7

DRYING AND CLEANING OF VESSELS

While in analytical operations, since one generally deals with aqueous solutions, the cleaned vessels may be used even if wet, it frequently happens in organic work, in experimenting with liquids not miscible with water, that dry vessels must be employed. In order to dry small pieces of apparatus rapidly, they should be rinsed first with alcohol and then with ether. To remove the last portions of the easily volatile ether, air from a blast is blown through the vessel for a short time, or the ether vapours are removed by suction. The alcohol and ether used for rinsing can frequently be used again; it is convenient to keep two separate bottles for the wash alcohol and wash ether, into which the substances, after being used, may be poured.

For rapid drying of large vessels this method is costly. In this case the procedure is as follows: The wet vessel is first drained as thoroughly as possible, and then heated with constant turning in a large luminous blast-flame, while, by means of a blast of air from bellows or other source, the water vapour is driven out. It may also be removed by careful heating and simultaneous suction.

Thick-walled vessels like suction flasks must not be heated over a flame, but are dried by the first method.

Vessels may be cleaned in part by rinsing them out with water with the use of a feather or flask-cleaner. If the last portions of the solution of a solid, *e.g.* in alcohol, are to be removed from a flask, it is not washed out at once with water, but first with a small quantity of the solvent, and then afterwards with water. If the vessel contained a liquid not miscible with water, it is first washed with alcohol and then with water. Resinous or tarry impurities adhering firmly to the walls can be removed by crude concentrated sulphuric acid. The action of this latter may be strengthened by adding a little water to it, by which heat is generated; also by the addition of some crystals of potassium dichromate. At times the impurities adhere so firmly that the vessel must be allowed to stand in contact with sulphuric acid for a long time. Crude concentrated nitric acid, or a mixture of this with sulphuric acid, is also used at times for cleaning purposes. Impurities of an acid character can, under certain conditions, be removed by caustic soda or caustic potash.

Finally, a method for cleaning the hands may be mentioned if they are discoloured by dyes which cannot be removed by water. If the dye, *e.g.* fuchsine, contains an amido (NH_2) group, the hands are dipped into a dilute, weakly acid solution of sodium nitrite. The dye is diazotised, and may be removed by washing in water. The two methods following are applicable to all dyes; the hands are immersed into a dilute solution of potassium permanganate to which some sulphuric acid has been added, and are allowed to remain for some time; the dye is oxidised, and thereby destroyed. After the permanganate has been washed off with water, the hands, especially the nails, are coloured brown by manganese dioxide. This is removed by washing the hands with a little sulphurous acid, or oxalic acid (ammonium oxalate plus hydrochloric or sulphuric acid). The second method is this: A thick paste of bleaching powder and a sodium carbonate solution is rubbed on the hands. This causes the oxidation and destruction of the dye as above. In order to take away the unpleasant odour of the bleaching powder, the hands are scrubbed with a brush, care being taken to remove the particles adhering to the upper and under surface of the nails, and are then washed, as just described, with sulphurous acid, or oxalic acid.

ORGANIC ANALYTICAL METHODS

DETECTION OF CARBON, HYDROGEN, NITROGEN, SULPHUR, AND THE HALOGENS

Tests for Carbon and Hydrogen. — If on heating a substance on platinum foil, it burns with a flame (exceptions, *e.g.* S), or decomposes with charring, it is an organic substance. Carbon and hydrogen may be detected in one operation, by mixing a small portion of the dried substance with several times its volume of ignited fine cupric oxide, placing the mixture in a small test-tube, adding more cupric oxide to the top of the mixture, and heating strongly, the tube being closed by a cork bearing a delivery tube bent twice at right angles. If the gas evolved (carbon dioxide) will cause a clear solution of barium hydroxide to become turbid, the original substance contained carbon; if it also contained hydrogen, small drops of water will collect in the upper cold part of the tube.

Test for Nitrogen. — To test an organic substance for nitrogen, it is heated in a small test-tube of difficultly fusible glass, about 5 mm. wide and 6 cm. long, with a piece of bright potassium the size of a lentil, which has been pressed between layers of filter-paper, in a Bunsen flame until decomposition, generally accompanied by slight detonations and dark colouration, takes place. The tube is finally heated to redness; while still hot it is dipped into a small beaker containing 10 c.c. of water; by this the tube is shattered, and any potassium unacted upon becomes ignited. The aqueous solution containing potassium cyanide, if nitrogen was present in the substance, is filtered from the carbon and glass fragments, the filtrate treated with a few drops of caustic potash or caustic soda until it shows an alkaline reaction; to this solution is then added a small quantity of ferrous sulphate solution and ferric chloride solution; it is boiled 1–2 minutes, and if potassium cyanide was present, potassium ferrocyanide will be formed. After cooling, the alkaline liquid is acidified with hydrochloric acid, the

precipitated ferric and ferrous hydroxides will be dissolved, and being acted upon by the potassium ferrocyanide, will form Berlin blue. Accordingly, if nitrogen was present, a blue precipitate is obtained, otherwise only a yellow solution will be formed. If the substance contains only a small proportion of nitrogen, at times no precipitate is obtained at first, but only a bluish-green solution. If this is allowed to stand some time, under certain conditions, over night, the precipitate will separate out. In testing easily volatile substances for nitrogen, a longer tube is used and the portions of substance condensing in the upper cold part of the tube flow back a number of times on the potassium. In place of potassium, sodium may also be used in most cases, but the former acts more certainly. In testing for nitrogen, in a substance containing sulphur, a larger quantity of potassium or sodium than that given above is used (for 0.02 grm. of substance, about 0.2 grm. potassium, in order to prevent the formation of an alkali sulphocyanate). Substances which evolve nitrogen at moderate temperatures, *e.g.* diazo-compounds, cannot be tested in the manner described. In dealing with a substance of this kind it must be determined whether on heating the substance with cupric oxide in a tube filled with carbon dioxide, a gas is given off which is not absorbed by a solution of caustic potash. (See quantitative determination of nitrogen.)

In a limited number of substances containing nitrogen, the presence of the latter may be proved by heating the substance with an excess of pulverised soda-lime in a test-tube with a Bunsen flame; this causes decomposition with evolution of ammonia, which is detected by its odour or by means of a black colour imparted to a piece of filter-paper moistened with a solution of mercurous nitrate. Nitro-compounds, *e.g.*, do not give this reaction.

Test for Sulphur. — The qualitative test for sulphur is made in the same manner as that for nitrogen. The substance is heated in a small tube with sodium. After the mass has cooled it is treated with water, and to one-half of the solution is added a small quantity (a few drops) of a solution of sodium nitroprussiate, just prepared by shaking a few crystals with water at the ordinary temperature. A violet colouration indicates the presence

of sulphur. Since the nitroprussiate reaction is very delicate, no conclusion as to the amount of sulphur can be drawn from the test, therefore the second half of the solution is treated with a lead acetate solution and acidified with acetic acid. In proportion to the amount of lead sulphide formed, the liquid will assume a dark colour, or a more or less heavy precipitate will appear, in this way indicating the original quantity of sulphur.

Easily volatile substances cannot usually be tested by this method. They are heated with fuming nitric acid in a bomb-tube to about 200 or 300°. After diluting with water the solution is tested with barium chloride for sulphuric acid. (See method for the quantitative determination of sulphur.)

Test for the Halogens. — The presence of chlorine, bromine, and iodine in organic compounds can only in rare cases be shown by precipitation with silver nitrate. This is explained by the fact that most organic compounds are non-electrolytes; *i.e.* that the solutions of the same do not contain free halogen ions, as is the case in solutions of the inorganic salts of the halogen hydric acids.

In order to detect the halogens, the substance to be tested is heated in a not too narrow test-tube with a Bunsen flame with an excess of chemically pure lime, the tube while still hot is dipped into a little water, chemically pure nitric acid is added to acid reaction, the solution is then filtered and treated with silver nitrate.

In compounds containing no nitrogen, a test for the halogens may be made by the same method given for nitrogen — heating with sodium. In this case the solution, filtered from the decomposition products and fragments of glass, is acidified with nitric acid and silver nitrate added. Substances containing nitrogen cannot be tested in this way for the halogens, since, as shown above, these on fusion with sodium give sodium cyanide, which, like the sodium halides, reacts with silver nitrate.

The presence of halogens may be recognised very quickly and conveniently by Beilstein's test. A piece of cupric oxide the size of a lentil, or a small rod of the oxide $\frac{1}{2}$ cm. long, is wrapped around with a thin platinum wire, the other end of which is fused to a glass handle, and heated in the Bunsen flame until it becomes

colourless. If, after cooling, a minute particle of the substance containing a halogen is placed on this and then heated in the outer part of the flame, the carbon burns first and a luminous flame is noticed. This soon vanishes, and there appears a green or bluish-green colour due to the vaporisation of the copper halide. From the length of time the colour is visible, conclusions may be drawn concerning the presence of a trace or more of the halogen in the original substance.

QUANTITATIVE DETERMINATION OF THE HALOGENS CARIUS' METHOD

The method consists in heating a weighed amount of the substance to be analysed in a sealed glass tube with silver nitrate and fuming nitric acid, by which it is completely decomposed (oxidised), and weighing the quantity of the silver halide thus formed.

Requisites for the analysis :

1. A tube of difficultly fusible glass sealed at one end, length about 50 cm. ; outside diameter, 18–20 mm. ; thickness of walls, about 2 mm. (Sealing-tubes, bomb-tubes.)
2. A funnel-tube about 40 cm. long, for transferring the silver nitrate and nitric acid to the glass tube.
3. A weighing-tube of hard glass (length, 7 cm. ; outside diameter, about 6–8 mm.).
4. Solid silver nitrate and pure fuming nitric acid. The purity of the latter is tested by diluting 2 c.c. of it with 50 c.c. of distilled water, and adding a few drops of a silver nitrate solution. Neither an opalescence nor a precipitate should appear.

Filling and Sealing the Tube. — After the bomb-tube, weighing-tube, and funnel-tube have been cleaned with distilled water, they are dried, not with alcohol and ether, but by heating over a flame. (See page 75, Drying.) The exact weight of the weighing-tube is next determined. Into this, with the help of a spatula, is placed 0.15 to 0.2 gramme of the substance to be analysed, finely powdered. The open end of the tube is wiped off with a cloth, and the exact

weight of the tube plus the substance is found. With the aid of the funnel-tube, about 0.5 gramme of finely powdered silver nitrate is transferred to the bomb-tube (a correspondingly larger amount up to 1 gramme is used for substances containing a high percentage of halogen) and 2 c.c. of fuming nitric acid. If a number of halogen determinations are to be made, it is advisable to measure off 2 c.c. of water in a narrow test-tube, mark the volume with a file on the outside, and then use this to measure the acid for the different determinations. After removing the funnel-tube, care being taken not to touch the walls of the bomb-tube with it, the weighing-tube is inserted in the bomb held at a slight angle, and is allowed to slide down to the bottom, but the substance must not come in contact with the acid. The tube is now sealed in the manner described on page 63. During the sealing, the substance must be prevented from coming in contact with the acid. Even after the tube is closed, this is not brought about purposely, as by violently shaking the tube.

If the substance to be analysed is liquid, it is placed in the weighing-tube with a capillary pipette, otherwise the procedure is just as described. In dealing with easily volatile substances, the weighing-tube is closed by a glass stopper, made by heating a piece of glass rod in the blast-flame until it softens, and then pressing it on a metal surface until a head is formed (Fig. 53).



FIG. 53.

Heating the Tube. — After cooling, the tube is transferred to the iron protecting case, and heated in the bomb-furnace, in accordance with the directions on page 65. The temperature and time of heating depend upon the greater or less ease with which the compound is decomposed. In many cases, it is necessary to heat aliphatic compounds, or aromatic compounds that oxidise easily, 2–4 hours at a temperature of 150–200°, while substances that do not easily oxidise, especially those that contain sulphur, must be heated 8–10 hours, and finally up to 250–300°. In this case it is convenient to so plan the analysis that the bombs may be sealed in the evening, so that the heating may be begun the first thing the next

day. The sealed tube is kept under the hood in the bomb-room, in the iron case, over night, which is clamped with its open end directed vertically upwards. The tube is never allowed to remain at the working table. If the furnace is not loaded, naturally it is most convenient to place the bomb at once in that. Since in many cases the oxidation begins even at the ordinary temperature, pressure is developed in the tube; therefore, after it has been standing over night, it must not be removed from the iron case to be examined. The heating is done gradually; at first, with a small flame, the gas-tubes being lowered from the furnace. Gradually these are raised, and the flames increased in size. The following table will show how the heating of a moderately refractory substance should be regulated.

The heating is begun at 9 o'clock A.M.

From 9-10 the temperature is raised to about 100°,

10-11	"	"	"	150°,
11-12	"	"	"	200°,
12-3	"	"	"	250°,
3-6	"	"	"	300°.

If an especially high pressure is generated by the decomposition of a substance, the danger of the bursting of the tube may be lessened by turning off the gas before leaving the laboratory at noon, and then in the afternoon opening the capillary, sealing and heating again to a higher temperature. The same method is followed in working with a substance so refractory that several days' heating is required; in this case at the beginning of the second day the pressure is reduced by opening the tube.

If two bombs are heated in the furnace at the same time, an entry is made in the note-book to show which tube lies to the right and which to the left. If this has been neglected, and the identity of the two tubes is in doubt, the neglect may be corrected by again weighing the two weighing tubes.

To Open and Empty the Tube.—The perfectly cooled tube is opened according to the directions given on page 66. Especial care must be taken before heating the capillary to softening in a

large flame, to drive back into the tube by gentle heating over a small flame, any of the liquid which may have collected in the capillary. Before the conical end is broken off the tube is examined to see whether it still contains crystals or oily drops of the undecomposed substance. In case it does, the capillary is again sealed and the tube reheated; but if it does not, the conical end is removed according to the directions given on page 66. The part broken off is first washed free from any liquid or any of the precipitate which may have adhered to it, with distilled water, into a beaker; the portion in the tube is diluted with distilled water, upon which there is generally obtained a bluish-green solution, coloured by nitrous acid; this is poured, together with the weighing-tube into a beaker by inverting the tube, care being taken that the sudden falling of the weighing-tube does not break the bottom of the beaker. In pouring out the tube-contents, the attention should be directed to the open end of the tube and not to the liquid in the rear end, otherwise some of the liquid may easily be spilled. After the outer open portion of the tube has been washed with distilled water the tube is revolved and the precipitate in the interior is washed out; this is repeated as often as may be necessary. If a portion of the silver halide adheres firmly to the glass, it may be removed by loosening it with a long glass rod over the end of which has been drawn a piece of rubber tubing (such as is used in the quantitative analysis of inorganic substances) and then washing it out with distilled water. The weighing-tube is removed from the bottom of the beaker with a glass rod or thick platinum wire held against the walls above the liquid, washed thoroughly inside and out with distilled water, and then raised with the fingers and washed several times again. At times the weighing-tube becomes yellow to deep brown in colour due to the formation of silver silicate. This is not detrimental to the results of the analysis.

To filter off and weigh the Silver Halide. — The beaker is now heated on a wire gauze until the silver halide has settled to the bottom and the supernatant liquid is clear. Since the excess of silver nitrate at times packs together with the silver halide to form thick, solid lumps, the precipitate is from time to time crushed

with a glass rod, the end of which has been flattened out to a broad head. After cooling, the silver halide is collected on a filter, the weight of the ash of which is known, and washed with hot water until, on testing the filtrate with hydrochloric acid, no turbidity follows; the filter, together with the funnel, is then dried in an air-bath at $100-110^{\circ}$, the funnel being covered with a piece of filter-paper. In order to weigh the dry halide, as large an amount as possible is separated from the paper carefully, and transferred to a watch-glass placed on a piece of black glazed paper. The portions which fall on the paper are swept into the watch-glass with a small feather. The filter is rolled up tightly, wrapped with a platinum wire, and ignited in the usual way over a weighed porcelain crucible; the heating is done only with the outer part of the flame, and not with the inner, reducing part. The folded filter may also be incinerated directly in the crucible, which is first heated over a small flame, and the temperature increased later; the heating is continued until the filter ash appears uniformly light. In order to convert the silver which has been reduced in the incineration back to the silver halide, the fused residue is moistened, by the aid of a glass rod, with a few drops of nitric acid,—if the latter method of incineration has been employed, only after complete cooling of the crucible. It is now evaporated to dryness on the water-bath. It is then treated with a few drops of the corresponding halogen acid, and again evaporated to dryness on the water-bath. The principal mass of the silver halide on the watch-glass is transferred to the crucible with the aid of a feather, and heated directly over a small flame until it just begins to fuse: the crucible is then placed in a desiccator, and allowed to cool. If the analysis is intended to be very exact, the principal mass of the silver halide may be moistened before fusion, with a few drops of nitric acid, and then evaporated on the water-bath with the halogen acid.

The silver halide can also be conveniently weighed, either in an asbestos tube, or in a Gooch crucible. Concerning this compare Jannasch, "*Praktischer Leitfaden der Gewichtsanalyse*," Second Ed.,

pages 10 and 145; also *Chemical News*, 37, 181; and *Zeitschrift für analytische Chemie*, 19, 333.

Even after taking the usual precautions, it sometimes happens that the silver halide is mixed with fragments of glass, which will, of course, cause the percentage of halogen to be too high. If the substance under examination is silver chloride, and the presence of glass is noticed in the beaker or on filtering, an error may be avoided, by pouring over the completely washed, moist silver chloride on the filter, slightly warmed dilute ammonium hydroxide several times, then washing the filter with water, and precipitating the pure silver chloride in the filtrate by acidifying with hydrochloric acid. If the compound under examination is silver bromide or iodide, and glass fragments have been noticed, the analysis is carried out to the end in the usual way. To determine the amount of glass present, the silver halide in the crucible is treated with very dilute pure sulphuric acid, and a small piece of chemically pure zinc is added. In the course of several hours, the silver halide is reduced to spongy, metallic silver. By careful decantation, the liquid is separated from the silver, water is added and decanted; this is repeated several times. It is then treated with dilute nitric acid, and heated on the water-bath until all the silver is dissolved. After dilution with water, it is filtered through a quantitative filter, the undissolved glass fragments are also well washed on the filter, and the latter incinerated. The weight of the glass is to be subtracted from the weight of the halide obtained. It is obvious that the purity of the fused silver chloride may also be tested in this way.

In conclusion, the atomic weights of the halogens, the molecular weights of the corresponding silver compounds, and the logarithms of the analytical constants are here given :

$$\text{Cl} = 35.46; \text{AgCl} = 143.34; \log \frac{\text{Cl}}{\text{AgCl}} = 0.39337 - 1$$

$$\text{Br} = 79.92; \text{AgBr} = 187.80; \log \frac{\text{Br}}{\text{AgBr}} = 0.62896 - 1$$

$$\text{I} = 126.92; \text{AgI} = 234.80; \log \frac{\text{I}}{\text{AgI}} = 0.73283 - 1$$

Küster's Modification.—The determination of the halogens may be materially facilitated by using 16–20 drops of fuming nitric acid instead of $1\frac{1}{2}$ –2 c.c. The tube so charged is heated directly up to 320° – 340° ,—it is unnecessary to heat it gradually. When an ordinary thermometer is employed to register temperatures above 300° , the column of mercury frequently parts. A bomb-thermometer made by C. Desaga, Heidelberg, Germany, at the suggestion of the author, corrects this defect. It contains nitrogen over the mercury column, and possesses but two degree marks, corresponding to 320° and 340° .

QUANTITATIVE DETERMINATION OF SULPHUR CARIUS' METHOD

This method, like the preceding one, depends upon the complete oxidation of the weighed substance, by heating it with fuming nitric acid in a sealed tube. The sulphuric acid thus formed is weighed as barium sulphate. The charging, sealing, heating, opening, and emptying of the tube are performed in exactly the same way as in the halogen determinations; but in this case it is evident that the use of silver nitrate is superfluous. Before breaking off the conical end, the tube is examined to see that no undecomposed portions of the substance are present; if there should be, the capillary is again sealed and the tube reheated. Before the sulphuric acid is precipitated with barium chloride, the bottom of the beaker must be examined for any fragments of glass which may be present; if there are any, they are filtered off through a small filter.

Precipitation of the Barium Sulphate.—The liquid from the bomb, diluted with water up to 400 c.c., is heated almost to boiling on a wire gauze and acidified with hydrochloric acid; a solution of barium chloride heated to boiling in a test-tube is gradually added until a precipitate is no longer formed. This can be easily observed by allowing the precipitate to settle somewhat before adding more of the solution. The liquid is then heated over a

small flame until the barium sulphate settles at the bottom of the beaker and the supernatant liquid is perfectly clear: at times from one to two hours' heating may be necessary. After cooling, the liquid is filtered, without disturbing the precipitate at the bottom, through a small filter the weight of the ash of which is known; the precipitate remaining in the beaker is boiled several minutes with 100 c.c. water and filtered through the same filter. The precipitate occasionally at first goes through the paper; in case it does, another beaker is placed under the funnel so that the entire quantity of liquid need not be refiltered. The precipitate is washed with hot water until a portion of the filtrate tested with dilute sulphuric acid shows no turbidity. Before throwing away the filtrate, barium chloride is added in order to be sure that a sufficient quantity was used in the first instance. If a precipitate is formed, the above process is repeated and the second precipitate collected on the filter containing the first.

The method just described has the disadvantage that if a smaller quantity of water be used for diluting the contents of the tube than that given above, the barium sulphate may easily carry along with it some barium nitrate, which is only removed with difficulty on washing with water. Since, in consequence of this, the percentage of sulphur is too high, it is for many reasons preferable to wash the contents of the bomb into a porcelain dish instead of a beaker, and to evaporate the liquid on the water-bath until the acid vapours vanish, before adding the barium chloride; by this operation the nitric acid is removed. After evaporating, the residue is diluted with water, filtered if necessary, from any glass fragments, and the operation just described above repeated. Under these conditions, too much of an excess of barium chloride is to be avoided.

Ignition and Weighing of the Barium Sulphate.—In order to prepare the barium sulphate for weighing, it is not necessary to dry it before incineration; if Bunsen's method is followed, it may be incinerated while still moist. With the aid of a small spatula or knife the moist filter is removed from the funnel and folded in the form of a quadrant. Should any barium sulphate adhere to

the funnel, it is removed with a small piece of filter-paper, which is incinerated with the main mass. After the filter has been carefully folded toward the centre, it is pressed into the bottom of a weighed platinum crucible, placed on a platinum triangle in such a position that its axis is inclined $20-30^\circ$ from a vertical position. The cover, also inclined at an angle of $20-30^\circ$, in the opposite direction, however, is supported before the crucible, so that the upper half of the opening of the latter is uncovered. The burner under the crucible is placed in such a position that the flame, which must not be too large at first, is directly under the angle formed by the crucible and cover. This will allow the ignition of the filter to take place at so low a temperature that reduction of the barium sulphate need not be feared. It sometimes happens that on heating the filter, the gases formed take fire at the mouth of the crucible, which, however, does no harm. After some time the burner is placed under the bottom of the crucible, the flame increased, and the heating continued until the residue has become white. The crucible is now placed in an upright position, heated a short time with the full flame, and then allowed to cool in a desiccator. It is entirely superfluous to treat the barium sulphate with sulphuric acid and then evaporate it off. The barium sulphate may also be weighed in a Gooch crucible. (See page 84.) If the percentage of sulphur found is too high, this may have been caused, under certain conditions, by the fact that in the precipitation too great an excess of barium chloride has been used, and that the barium sulphate has carried along some of it. This source of error may be rectified by treating the ignited barium sulphate with water until the crucible is half full, then adding a few drops of concentrated hydrochloric acid, and heating on the water-bath for fifteen minutes. The liquid is filtered from the precipitate through a quantitative filter; the greatest portion of the precipitate remaining in the crucible is again treated with water and hydrochloric acid, and the contents of the crucible poured on the filter already used; after washing repeatedly with water, the filter and precipitate are again ignited as before. This process is obviously only employed when the barium sulphate has

not been evaporated down with sulphuric acid. For the calculation of the analysis the atomic and the molecular weights are given :

$$S = 32.07; \text{BaSO}_4 = 233.44; \log \frac{S}{\text{BaSO}_4} = 0.13792 - 1$$

Simultaneous Determination of the Halogens and Sulphur. —

If a substance contains both a halogen and sulphur, they may be determined in a single operation by the following method : As in the determination of the halogens, the bomb is charged with silver nitrate and nitric acid, and the silver halide filtered off after the heating, as above described. The filtrate thus obtained contains, besides the excess of silver nitrate, the sulphuric acid formed by oxidation. This latter cannot be precipitated as before with barium chloride, since the silver as silver chloride would also be thrown down. In its place is used a solution of barium nitrate, the purity of which has been tested by adding silver nitrate to it. The precipitation is made hot as above directed, the solution used being as dilute as possible — the volume of which must be at least 500 c.c. A large excess of barium nitrate is particularly to be avoided. If the barium nitrate solution contains halogen salts as impurities, it is heated, and silver nitrate added so long as a precipitate is formed, the precipitate filtered off, and the solution, which is now free from halogens, is used for the precipitation.

**QUANTITATIVE DETERMINATION OF NITROGEN
DUMAS' METHOD**

In scientific laboratories, the method almost exclusively used for determining nitrogen quantitatively is that of Dumas. The principle involved is that the substance is completely burned by cupric oxide in a tube filled with carbon dioxide, the nitrogen is evolved as such, and its volume measured, while the carbon and hydrogen are completely oxidised to carbon dioxide and water.

Requisites for the analysis :

1. A combustion tube of difficultly fusible glass, 80–85 cm. long ; outside diameter, about 15 mm.
2. A glass funnel-tube with wide stem (at least 10 mm. in diameter).
3. 400 grammes of coarse and 100 grammes of fine cupric oxide. The former is kept in a large flask, the latter in a small one, both of which are closed by a cork covered with tin-foil.
4. 500 grammes of magnesite, in pieces the size of a pea. The fine powder, which cannot be used, is sifted out in a wire sieve. The dark grains which have become discoloured by impurities are thrown out.
5. A small flask of pure methyl alcohol (50 grammes) for reducing the copper spiral.
6. A copper spiral, 10–12 cm. long. This is made by winding an oblong piece of copper wire gauze spirally around a thin glass rod. It is made of such a width that when in position it will touch the walls of the combustion tube ; a space between the walls and spiral is disadvantageous. Also a short copper spiral from 1–2 cm. long.
7. A solution of 150 grammes of potassium hydroxide in 150 grammes of water. It is prepared in a porcelain dish, and not in a glass beaker or flask, since these are frequently broken by the heat generated by the solution. After cooling, it is preserved in a well-closed bottle.

8. A nickel crucible 6 cm. high ; diameter of top 7 cm., for the ignition of the coarse cupric oxide.
9. A moderately large porcelain crucible for the ignition of the fine cupric oxide.
10. A small mortar with a glazed bottom.

Besides these, a weighing-tube, a one-hole rubber stopper for closing one end of the combustion tube, a sieve to sift the copper oxide, a small feather, thermometer, absorption apparatus, and a eudiometer.

Preparations for the Analysis. — The analysis is conveniently begun by heating the entire quantity of coarse copper oxide in the nickel crucible over a large flame (Fletcher burner), and the fine copper oxide in the porcelain crucible over a Bunsen flame for a long time, the crucibles being supported on wire triangles. The covers are placed on the crucibles loosely, and the copper oxide occasionally stirred with a thick wire. While the copper is being heated, one end of the combustion tube is sealed to a solid head, the narrower end being selected for this purpose, if the tube is not perfectly cylindrical. The sealing is done as follows : The end of the tube is first warmed in a luminous flame, with constant turning ; it is then heated to softening, in the blast-flame, a glass rod fused on it, and the heated portion suddenly drawn out to a narrow tube. The glass rod is now fused off, and the conical part of the tube just produced is heated and drawn out. The cone is then heated in the hottest flame until it falls together ; it is finally allowed to cool gradually over a small luminous flame. When this operation is finished, the open end of the tube is warmed in a luminous flame, and, with constant turning, the sharp edges are rounded by the blast-flame : it is then allowed to cool in the luminous flame again. After complete cooling, the soot is removed, the tube rinsed out several times with water, the water allowed to drain off as completely as possible, and the tube finally dried in one of the two following ways : The tube, with constant turning, is repeatedly passed through the large luminous flame of a blast-lamp, while a current of air is blown

from a blast into the bottom of it by means of a narrower, longer (10 cm.) tube inserted in the larger tube ; this operation is continued until all moisture is removed. Or the combustion tube is clamped in a horizontal position, a narrower tube extending to the sealed end, attached to suction, is inserted, and the combustion tube equally heated with a Bunsen burner throughout its entire length ; the water vapour is drawn off by the suction. To reduce the long copper spiral which is to be used for the reduction of oxides of nitrogen which may be formed, the method of procedure is as follows : Into a test-tube large enough to admit the spiral, 1 c.c. of methyl alcohol is placed ; the spiral, held by crucible tongs, is then heated to glowing in a large, somewhat roaring blast-flame, and dropped as quickly as possible into the test-tube ; since this becomes strongly heated at its upper end, it is clamped in a test-tube holder, or wrapped in a cloth or strips of paper. The dark spiral soon assumes a bright metallic lustre, while vapours, having a sharp, pungent odour (oxidation products of methyl alcohol like formic aldehyde and formic acid), which frequently become ignited, are formed ; after a few minutes, the tube may be loosely corked, and the spiral allowed to cool. When this operation is ended, the copper oxide will have been sufficiently heated, and the flames may be removed. During the cooling, the substance to be analysed is weighed. A convenient method is this : The weight of the weighing-flask is determined with exactness to centigrammes, this weight is entered in the note-book at a convenient place for future use. The substance to be analysed is now placed in the weighing-tube, and the weight of the tube, plus substance, is determined exactly to the tenth of a milligramme. In the meantime, the copper oxide has cooled sufficiently to be transferred to the appropriate flask. The combustion tube is next filled.

Filling the Tube. — At the edge of the working table is placed a stand ; fastened firmly near the bottom of this is a clamp projecting over the edge of the table supporting the combustion tube in a vertical position, the mouth being at about the level of the table. The tube is now directly filled with the magnesite until the layer has a height of 10–12 cm. (Fig. 54). A small roll of

copper gauze 1–2 cm. long, held with pincers or tongs, is heated for a short time in a Bunsen flame (it need not be reduced) and dropped on the magnesite. The funnel-tube is then placed in the tube, and from the flask coarse copper oxide is poured in until the layer measures 8 cm., and upon this is poured a layer of 2 cm. of the fine oxide. To the operation following — the mixing of the

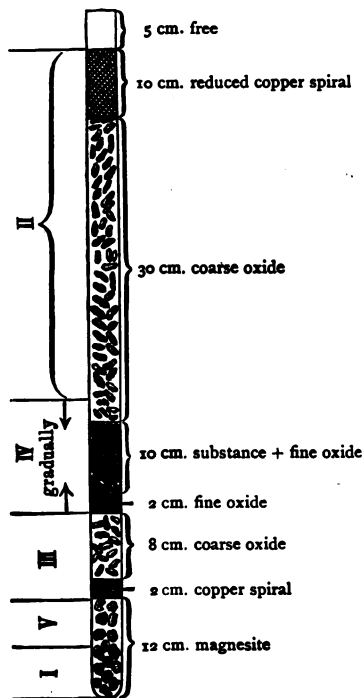


FIG. 54.

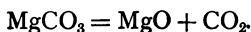
substance with copper oxide and the transference of the mixture to the tube — especial care must be given. In the bottom of a small mortar, standing on black, glazed paper a $\frac{1}{2}$ cm. layer of the fine, perfectly cooled copper oxide is placed; to this is added from the weighing-tube the substance to be analysed, of which 0.15–0.20 gramme is taken, unless the substance contains a small proportion of nitrogen, when more is taken. Since the weight of the empty tube is known as well as that of the substance contained therein, one can easily decide, by measuring with the eye, how much of the substance to take. Fine copper oxide is now added until the substance is completely covered, and the two are carefully mixed by

stirring with the pestle, without pressure; during the mixing care must be taken not to stir so rapidly as to cause dust-like particles of the mixture to leave the mortar. With the aid of a clean, clipped feather, such as is used in quantitative operations, or a small brush, the contents of the mortar are transferred through

the funnel-tube into the combustion tube. The operation must be done cautiously to prevent the light, dusty particles from being blown away. The mortar, as well as the pestle, is now rinsed with a fresh portion of the fine copper oxide, and this is likewise transferred to the tube with the aid of the feather. The layer of substance plus copper oxide should be about 10 cm. long. Then follows a layer of 30 cm. of coarse copper oxide, and finally the reduced copper spiral.

The length of the tube, as well as that of the single layers, is regulated in accordance with the size of the combustion furnace; the figures given above refer to a furnace possessing a flame surface of 75 cm. Generally the tubes are 5 cm. longer than the furnace; the tube contents are of the same length as the flame surface.

Heating the Tube. — After the tube is filled it is held in a horizontal position and tapped gently on the table in order that a canal may be formed in the upper portion of the fine copper oxide; it is then connected with a rubber stopper to the absorption apparatus which has been charged with caustic potash solution, and placed in the combustion furnace, the rear end of which (that under the magnesite) has been raised on a block (Fig. 55). The following points are to be observed: In the lower part of the absorption apparatus there must be a sufficient amount of mercury to extend almost to the side-tube; if this is not the case, more mercury is added: the end of the glass tube passing through the rubber stopper must be flush with the end of the stopper. In order to protect the latter from the heat, there is placed over the portion of the tube projecting beyond the furnace, an asbestos plate having a circular opening in the centre. After opening the pinch-cock of the absorption apparatus, the burners under the last half of the magnesite are lighted; the flames, being small at first, are increased in size, as soon as the tube becomes warmed, but not sufficiently to cause them to meet above the tube. In order to raise the temperature higher when it becomes necessary, the tube is covered from both sides with the tiles. After about ten minutes a rapid current of carbon dioxide is evolved, the magnesite being decomposed by heat as represented in the following equation:



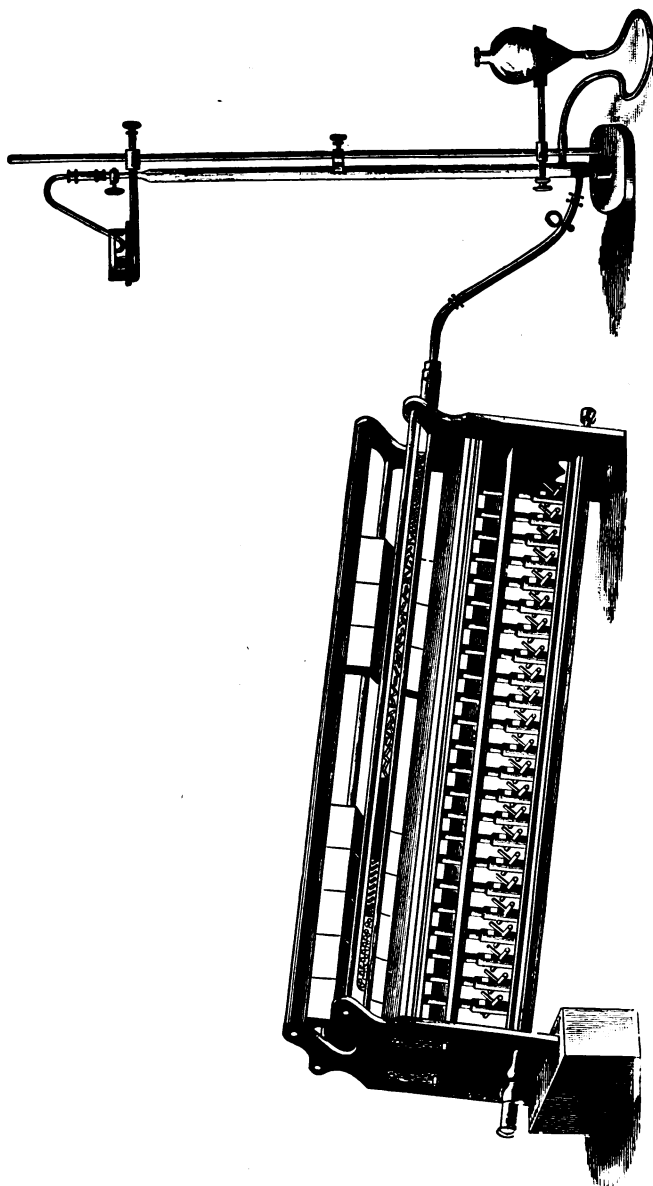


FIG. 55.

During this operation the glass stop-cock of the absorption apparatus is opened, and the pear-shaped vessel placed as low as possible, so that it contains the principal portion of the caustic potash. After a rapid current of carbon dioxide has been evolved for about fifteen minutes, the burners under the copper spiral are lighted in order to drive out any occluded gas (hydrogen), the pear-shaped vessel is raised high enough to cause the caustic potash to ascend somewhat above the tubulure in the glass cock, the latter is closed, and the pear-shaped vessel again lowered as far as possible. When the air in the tube has been completely replaced by carbon dioxide, only a minimum quantity of light foam should collect over the potash in the course of two minutes. If this is not the case, and a large air volume collects, the glass cock is opened, upon which the potash flows in to the lowered pear vessel, and carbon dioxide is caused to pass through the tube for five minutes longer. The pear vessel is then raised as high as at first, the glass cock closed, and the former lowered. An observation will show whether the air has been displaced, which should be the case under normal conditions. If now after two minutes only a trace of foam has collected, the end of the delivery tube is dipped under the water in a dish as shown in Fig. 55, the pear raised to the highest point of the delivery tube, and the glass cock opened in order that the potash may drive out the air in the delivery tube: when this has been done, the cock is closed again and the pear lowered to the bottom. All the flames but one under the magnesite are now extinguished or lowered, and those under the long copper spiral as well as those under four-fifths of the adjacent layer of coarse copper oxide are lighted at the same time; the flames, small at first, are increased in size, after the tube has become somewhat heated, until the copper oxide is heated to dull redness. Concerning the steps taken in heating the tube, refer to Fig. 54 — the numbers on the left indicate the portions of the tube to be heated successively. At this point care is taken that the flames are not so large as to meet above the tube. As before in heating the magnesite, after the first warming the heated portions of the tube are covered on both sides with the tiles. As soon as the forward layer of coarse copper oxide becomes dark red, the burners under the rear layer of coarse oxide adjacent to

the magnesite are lighted — small flames at first, which are increased after a time, the tube being covered simultaneously with the tiles. Care must be taken that the flames nearest the layer of substance plus fine copper oxide are not too large, in order that the substance may not yet be burned. Upon the operation which now follows — the gradual heating of the fine oxide containing the substance — virtually depends the success of the analysis. For the proper manipulation of this operation especial care must be taken. It is a rule that the heating had better be somewhat too slow than too rapid. A small flame is now lighted at the point adjacent to the short layer of coarse oxide : an observation of the absorption apparatus will show whether after some time any unabsorbed gas collects. If this is the case, no other burners are lighted until the evolution of gas ceases. When the gas no longer collects, another burner on the opposite side of the fine oxide is lighted. In this way the burners are gradually lighted from both sides, toward the middle of the fine oxide, and after, in each case, the cessation of the evolution of the gas, the flames are gradually made larger until finally the tube covered with tiles is heated with full flames ; thus the substance is regularly and quietly burned. The combustion must be so conducted that the bubbles of gas ascend in the absorption apparatus with a slow regularity. If the single bubbles cannot be counted, or if they are so large as to occupy almost the entire cross-section of the absorption tube, the heating is too strong, and the last burners lighted must be extinguished or lowered, the tiles being also laid back at the same time if necessary, until the generation of the gas is lessened. When this is ended, small flames are again lighted under the entire layer of magnesite, and increased in size after some time. As soon as the evolution of carbon dioxide has become active, the flames under the rear half of the magnesite lighted at the beginning of the analysis are extinguished. After a rapid current of carbon dioxide has passed through the tube for ten minutes, all the nitrogen is carried over to the absorption apparatus. This is shown by the complete absorption of the gas bubbles by the potash, as at the beginning of the analysis, except for a minimum foam-like residue. The absorption apparatus is

then closed by the pinch-cock and the rubber stopper bearing the connecting tube is withdrawn from the combustion tube. The gas is not immediately transferred to the eudiometer, but the pear is raised until the surfaces of the liquid in the pear and that in the tube are at the same level: the apparatus is then allowed to stand for at least half an hour. The flames under the combustion tube are not turned out simultaneously, but first one is extinguished, and then after a short time another, and so on. During the cooling of the tube the weighing-flask is weighed again.

Transferring the Nitrogen.—After the nitrogen has stood in contact for at least half an hour with the caustic potash, in order that the last portions of carbon dioxide may be absorbed, the end of the delivery tube is dipped under the surface of the water contained in a wide-mouth cylinder, as represented in Fig. 56, care being taken that in the lower bent portion of the delivery tube no air bubbles are present; if there are, they must be removed with a capillary pipette. The eudiometer is now filled with water, the end closed with the thumb, inverted and dipped below the surface of the water, the thumb removed and the tube clamped to the cylinder, at an oblique angle, so that the end of the delivery tube may be passed under it. The pear supported by the clamped ring is raised as high as possible above the delivery tube, and the glass cock gradually opened. The nitrogen is thus transferred to the eudiometer, the cock being left open until the delivery tube is completely filled with the caustic potash. The absorption apparatus is then removed, the eudiometer wholly immersed in the water. To obtain the temperature, a thermometer, held in the clamp which supported the

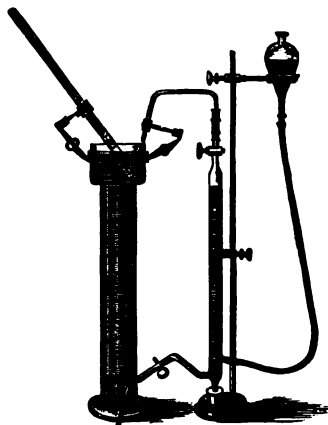


FIG. 56.

delivery tube, is immersed in the water as far as possible. After about ten minutes, the nitrogen has come to the same temperature as the water, the eudiometer is then seized with a clamp especially adapted to this purpose, or crucible tongs,—never with the hands,—and is raised so far out of the water that the level of water inside and outside the tube is the same. The volume of gas thus read off, is under the same pressure as that indicated by a barometer.

Calculations of the Analysis.—If s is the amount of substance in grammes, v the volume of nitrogen read at the temperature t° , and b the height of the barometer in millimetres, w the tension of the water vapour in millimetres at t° , then the percentage of nitrogen is p :

$$p = \frac{v \cdot (b - w) \cdot 0.12505}{760 \cdot (1 + 0.00367 \cdot t) \cdot s}.$$

The calculation of the analysis is rendered easier by referring to the table in which the weight of one cubic centimetre of moist nitrogen is given in milligrammes at different temperatures and pressures. If this, under the observed conditions, is g , then the percentage of nitrogen is:

$$p = \frac{100 \times v \times g}{s}.$$

In this formula, s is the weight of the substance in milligrammes.

A table is given at the end of the book for these calculations. The values for the height of the barometer not found in the table may be obtained by interpolation. For ordinary work it is unnecessary to read off fractions of degrees on a thermometer, or of millimetres on a barometer, since the slight differences in values found in this manner lie within the limits of experimental error.

The upper figure in each space in the table is the weight of moist nitrogen in milligrammes, and the lower figure is the mantissa of the logarithm of the weight of nitrogen.

Length of Time for an Analysis.—The following abstract will give an approximate idea of the length of time that the single

operations of a well-conducted combustion ought to occupy. From the beginning of the heating of the magnesite to the appearance of a rapid current of carbon dioxide requires about 10 minutes, the first test as to whether air is still present in the tube follows after a further 15 minutes ; length of time for various tests, 5 minutes. From the warming of the forward layer of the copper oxide with the spiral to the heating of the rear layer of oxide to a dark red heat, 15 minutes. The actual combustion of the substance requires 30 minutes. The displacement of the last portions of nitrogen by heating the magnesite requires 10 minutes. Total, 1 hour and 25 minutes.

These time figures are, of course, only to be considered as approximate, since they depend upon the efficiency of the furnace, upon the nature of the substance burned, upon the skill of the experimenter, and upon other factors.

Subsequent Operations.—After the tube has cooled and the copper spiral taken out, all the copper oxide is sifted to separate the coarse from the fine, and may be used again for further analyses as often as desired, provided that it is reheated each time in the nickel crucible to oxidise it. The tube may also be used again if it has not been distorted by high heating. The magnesite is useless for further analyses.

The caustic potash in the absorption apparatus, which can be used a second time, is poured into a bottle, which is then well closed. The absorption apparatus, including the rubber tubing, is washed out repeatedly with water, so that the latter may not be corroded by the caustic potash.

General Remarks.—The above-described method of Dumas for nitrogen is used in variously modified forms, but the principle is the same in all. It is preferred in many places to generate the carbon dioxide from acid sodium carbonate or manganese carbonate. A combustion tube open at both ends may be used, if a number of nitrogen determinations are to be made. The tube is charged as represented in Fig. 60 (page 107). The substance is placed in a porcelain or copper boat. In order to replace the air by carbon dioxide, the rear end of the tube is connected with

another tube of difficultly fusible glass (closed at one end), 25–30 cm. long and 15–20 mm. wide, which is three-fourths filled (in cross section) with sodium bicarbonate. In order to absorb the water generated from this on heating, a small sulphuric acid wash bottle is interposed between the two tubes. The layer of bicarbonate is heated with a single Bunsen burner, beginning at the fused end. In order to protect the bicarbonate tube from the direct flame, it is surrounded by a cylinder of coarse iron gauze (Fig. 57). The bicarbonate may be replaced by a Kipp generator.

Further, the mixing of the substance with the fine copper oxide may be done in the tube. Instead of the absorption apparatus

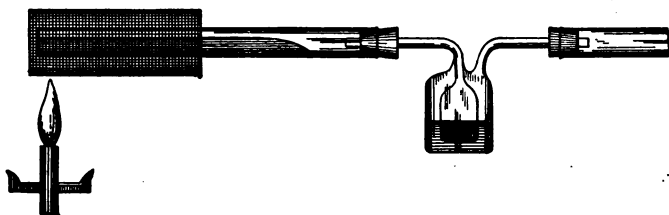


FIG. 57.

of Schiff, described above, a graduated tube from which the volume of the gas may be read directly may be used, thus obviating the necessity of transferring it to a eudiometer. This modification carries with it, however, the disadvantage that the tension of caustic potash is not exactly known, and therefore a somewhat arbitrary correction must be applied. But as mentioned these modifications do not differ essentially.

QUANTITATIVE DETERMINATION OF CARBON AND HYDROGEN LIEBIG'S METHOD

The essential part of the method consists in completely burning with copper oxide a weighed amount of the substance, and then weighing the combustion products, carbon dioxide and water.

The requisites for analysis are :

1. A hard glass tube open at both ends ; outside diameter 12–15 mm. It should be about 10 cm. longer than the furnace.
2. Four hundred grammes of coarse and 50 grammes of fine copper oxide, preserved in bottles closed with tin-foil-covered corks as in the nitrogen determination. But the copper oxide used for the latter purpose and that for the carbon and hydrogen determinations are always kept in separate bottles.
3. A U-shaped and a straight calcium chloride tube.
4. A caustic potash apparatus. The Geissler form is the most convenient.
5. A drying apparatus for air or oxygen.
6. Two one-hole rubber stoppers fitting the ends of the combustion tube.
7. A glass tube provided with a cock.
8. Two copper spirals of 10 and 12–15 cm. length, respectively ; two short spirals 1–2 cm. long.
9. A piece of good rubber tubing 20 cm. long ; six pieces rubber tubing 2 cm. long (thick-walled and seamless).
10. A porcelain and a copper boat.
11. A screw pinch-cock.
12. Two asbestos plates for the protection of the rubber stoppers.

Preparations for the Analysis.—The sharp edges of the combustion tube are rounded by careful heating in a blast-flame. After cooling the tube is rinsed out with water several times ; this is allowed to drain off, and the tube dried by one of the methods given on pages 91 and 92.

The coarse copper oxide is not previously heated in the nickel crucible as in the determination of nitrogen, but this is done later in the tube itself. If the nature of the substance to be analysed is such that it is necessary to mix it with fine copper oxide, the latter is ignited for a quarter hour in the porcelain crucible and allowed to cool in a desiccator.

The U-tube for the absorption of the water (Fig. 61) is filled with granulated, not fused, calcium chloride, which must be freed

from any powder by sifting. In order to prevent the calcium chloride from falling out, both ends of the tube are provided with loose plugs of cotton. The open leg is closed by a rubber stopper or a good cork bearing a glass tube bent at a right angle. The cork stopper is covered with a thin layer of sealing-wax. Calcium chloride tubes, in which the open leg is longer than the other, are very convenient. After the tube is filled the open end may be sealed in a blast-flame. In this case the plug in this end is not cotton, but asbestos or glass-wool. In order that the tube may be suspended from the arm of the balance in weighing, a platinum wire with a loop in the centre is attached to both legs. Calcium chloride often contains basic chlorides, which not only absorb water, but also carbon dioxide, thus causing an error in the results of the analysis; before the filled tube is used a stream of dry carbon dioxide is passed through it for about two hours, dried air is then drawn through for half an hour to displace the carbon dioxide. The two side tubes of the calcium chloride tube are closed by pieces of rubber tubing 2 cm. long, in which is inserted a glass rod rounded at both ends, $1\frac{1}{2}$ cm. long. The tube may be used repeatedly until the calcium chloride begins to liquefy. The straight calcium chloride tube is filled in like manner, but it is unnecessary to pass carbon dioxide through this before using.

The three bulbs of the potash apparatus similar to the one represented in Fig. 58 are three-fourths filled with a solution of caustic potash (2 parts potassium hydroxide, 3 parts water) as follows: the horizontal tube which is to be charged with solid caustic potash is removed, and to the free end of the bulb tube rubber tubing is attached. The inlet tube represented in Fig. 58 at the left is now dipped into the caustic potash solution, contained in a shallow dish, and this is sucked up with the rubber tubing until the three bulbs are three-fourths filled. Care must be taken not to suck too strongly, otherwise some of the caustic potash solution may be drawn into the mouth. This may be prevented by inserting an empty wash bottle between the potash apparatus and the mouth, or the suction-pump may be used, in which case the water-cock must be opened to a very slight extent. After filling

the bulbs that part of the tube immersed in the potash solution is cleaned with pieces of rolled-up filter-paper. The horizontal potash tube, removed before filling the bulbs, is now filled with coarse-grained soda-lime and solid caustic potash in pea-size pieces as follows: In the bulb is placed a plug of glass-wool or asbestos, then follows a layer of the soda-lime, a layer of caustic potash, and finally another plug of glass-wool or asbestos. When this is done, it is closed in the same way as the calcium chloride tube. In handling the Geissler tubes it is always to be remembered that they are very fragile, and in all cases the lever-

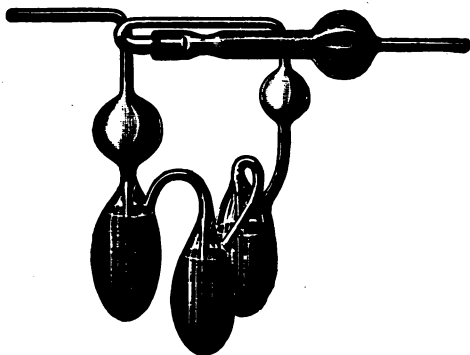


FIG. 58.

arm formed in lifting them should be as short as possible. When the apparatus is to be closed by rubber tubing, *e.g.*, it is not grasped by the bulbs, but immediately behind the place over which the tubing is to be drawn or pushed. When the potash apparatus has been used twice, it must be refilled. The longer of the two so-called copper oxide spirals need not be reduced before the combustion; on the contrary, it is oxidised in the combustion tube, as will be pointed out below. In order to be able to remove it from the tube conveniently a loop of copper wire is fastened in the meshes of the gauze near the end, or a not too thin copper wire is passed through the centre of the spiral and bent at one end to a right angle and at the other in the form of a loop. The

shorter spiral, which, as in the nitrogen determination, serves to reduce any oxides of nitrogen, is next reduced according to the directions given on page 92. To remove any adhering organic substances like methyl alcohol or its oxidation products the spiral is placed, after cooling, in a glass tube 20 cm. long, one end of which is narrowed; carbon dioxide is passed through it, and as soon as the air has been displaced, it is heated for a few minutes with a Bunsen flame and then allowed to cool in a current of carbon dioxide. To remove the mechanically adhering gas the spiral is placed in a vacuum desiccator. If this is not at hand, an ordinary desiccator containing a small dish of solid caustic potash or unslaked lime is used. (It may also be heated in an air-bath at 100–110°.)

For drying the oxygen or air an apparatus consisting of two wash cylinders and two U-shaped glass tubes mounted on a wooden stand is employed. The gas passes first through a wash cylinder containing a solution of potassium hydroxide (1 : 1), then a tube filled with soda-lime, then one filled with granulated calcium chloride, and finally a wash cylinder containing sulphuric acid (Fig. 59).

The legs of the glass tube containing the stop-cock are fused off and slightly narrowed at the ends, so that on either side of the cock the length is 5 cm.

Filling the Tube.—The simplest case of combustion with which one can deal is that involving the analysis of a substance containing no nitrogen. In a case of this kind, assuming that the furnace has a flame surface of 75 cm., the tube is filled in the following manner: A short copper gauze roll, 1–2 cm. long, of sufficient diameter to fit the tube tightly, and somewhat elastic, is pushed into the tube 5 cm., and then the opposite side of the tube is partially filled with a layer of coarse copper oxide 45 cm. held in position by another

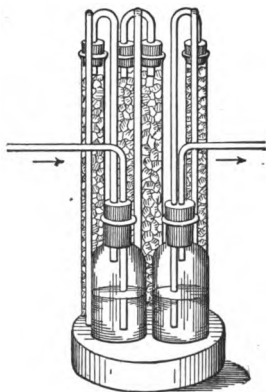


FIG. 59.

small elastic copper spiral at its upper end. Into the tube lying in a horizontal position the copper oxide spiral is pushed so far that its loop is 5 cm. from the mouth of the tube (Fig. 60).

Igniting the Copper Oxide. — The charged tube is placed in the furnace, the end nearest the copper oxide spiral is closed by a rubber stopper bearing the glass stop-cock tube, and the latter is connected with the drying apparatus by means of rubber tubing provided with a screw pinch-cock. The other end of the tube is allowed to remain open at first; while a current of oxygen is passed through the tube, slow enough to enable one to count the bubbles (the glass stop-cock is opened wide and the current regulated with the pinch-cock), the entire length of the tube is heated, at first with flames as small as possible; these are gradually increased until finally, the tiles being in position, the copper oxide begins to appear dark red. The water deposited at the beginning of the heating, in the forward cool end of the tube, is now removed with filter paper wrapped around a glass rod. When no more water collects, the front end of the tube is closed by a rubber stopper bearing the straight calcium chloride tube. After about 20 or 30 minutes' heating the burners under the copper oxide spiral, the adjacent empty space, and those under about 5 cm. of the copper oxide layer lying next, are extinguished, and at the same time the current of oxygen is cut off.

Weighing the Absorption Apparatus and the Substance. — While the rear part of the tube is cooling, the calcium chloride tube, the potash bulbs, and the substance are weighed. Before the absorption apparatus is weighed, it is wiped off with a clean cloth, free from lint, and the rubber tubing and glass rods removed; after the weighing, these are replaced. The substance, if solid, is weighed in a porcelain boat which has previously been heated strongly, and cooled in a desiccator. The boat is first weighed empty, 0.15 to 0.20 gramme of the substance placed in it and weighed again; it is then placed on a tin-foil-covered cork, in which a suitable groove has been cut, and transferred to a desiccator.

The Combustion. — When the rear end of the tube is cold, the copper oxide spiral is withdrawn with a hooked glass rod or wire,

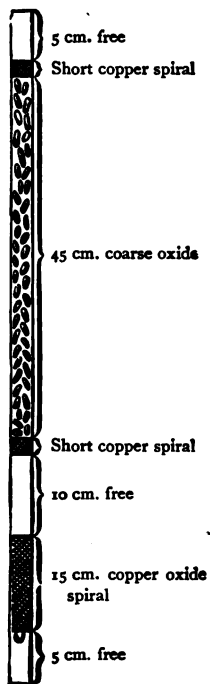
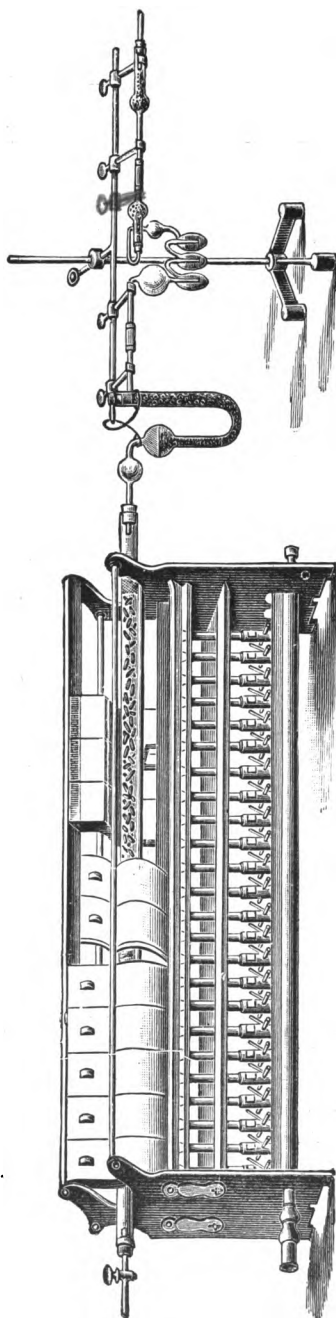


FIG. 60.



the porcelain boat is inserted as far as the coarse copper oxide, care being taken not to upset the boat, and finally the spiral is replaced. The stop-cock tube, with the cork closed, is then put in position. The straight calcium chloride tube is replaced by the weighed U-tube, with its empty bulb, which will condense the greater portion of the water, nearest the furnace. To the U-tube is connected, by a rubber joint, the potash apparatus, and the soda-lime tube of the latter with the straight calcium chloride tube in the same way (Fig. 61). The connecting of the different parts of the apparatus may be facilitated by blowing air from the lungs through each rubber joint before pushing it on the glass tubes. Especial care is taken to have a good joint between the U-calcium chloride tube and the potash bulbs, since at this point very commonly lies the source of error in analyses not concordant. A thick-walled seamless rubber tubing is employed; it is drawn over the two ends of the glass tubes until they touch. In order to provide against any possible leak, two ligatures of thin copper wire or "wax ends" are bound around the joints. A test as to whether the apparatus is perfectly tight is not always convincing when the combustion is conducted in an open tube; since, on the one hand, the heating is not constant, and on the other, in consequence of the friction of the solution in the narrow tubes, a leak, at times, may not be detected. The rubber stoppers closing the tube may be protected from the heat by placing on the tube, close to the furnace, an asbestos plate with a circular hole in the centre. After closing the screw pinch-cock, the glass-cock is opened, and a slow current of oxygen (two bubbles per second) is admitted to the tube by carefully opening the pinch-cock. Small flames are now lighted under the copper oxide spiral, which are increased after some time, until, finally, the spiral is brought to a dark red glow. When this is done, the flames under the unheated copper oxide are gradually lighted, care being taken not to allow any flame near the porcelain boat to be too large. Now follows the most difficult operation of the analysis, upon which the success of it virtually depends, viz. the gradual heating of the substance. This is conducted in exactly the same

way as that given under the nitrogen determination. The heating is begun, at first, with a single small flame; this is gradually increased in size, or several others may be lighted, then the tube is covered on one side with the tiles, and after a short time, on the other; and finally the full flames are used. With easily volatile substances, the heating at the beginning is not done with the flame, but by covering that portion of the tube containing the boat with hot tiles, taken from the forward highly heated portion of the furnace. Numerous modifications have been applied to this most difficult part of the analysis, concerning which no satisfactory general directions can be given. A valuable rule is to conduct the heating in such a way that the gas bubbles passing through the potash apparatus follow one another with as slow a regularity as possible. If the passage of bubbles becomes too rapid, the heating is moderated. If, during the combustion, water should condense in the glass-cock, or in the rear, cold portion of the tube, as it always does in the front end, it is removed by holding a hot tile under it, or by heating with a small flame. When the boat has been heated some time with the full flames, the combustion is considered to be ended. In order to drive the last portions of carbon dioxide and water from the tube into the absorption apparatus, a somewhat more rapid current of oxygen is passed through the tube, until a glowing splinter held before the opening of the straight calcium chloride tube is ignited. During this operation, the water, condensed for the most part in the front end of the tube, is also driven over into the calcium chloride tube, as above described. When this has been done, the rubber stopper is withdrawn from the front end of the combustion tube, care being taken to prevent the water in the calcium chloride tube from running out. To remove the oxygen in the absorption apparatus, a slow current of air which need not be dried is drawn through it for 1-2 minutes, with the mouth or suction. The apparatus is taken apart, closed up as above described, allowed to stand in the weighing-room for half an hour, and is then weighed. From the difference in the weights of the absorption apparatus before and after the combustion, the percentage of carbon and hydrogen is found from the following equations:

$$\text{Percentage of Carbon} = \frac{\text{Wt. CO}_2 \times 300}{\text{Wt. Substance} \times 11},$$

$$\log \frac{C}{\text{CO}_2} = 0.43573 - 1$$

$$\text{Percentage of Hydrogen} = \frac{\text{Wt. H}_2\text{O} \times 201.6}{\text{Wt. Substance} \times 18.016},$$

$$\log \frac{\text{H}_2}{\text{H}_2\text{O}} = 0.04884 - 1$$

Modifications of the Method. — In many cases instead of using oxygen for the ignition of the copper oxide, the same result may be obtained by using a current of air. The combustion may also be conducted in a current of air; but when the substance is difficult to burn, it is still necessary toward the end of the operation to pass oxygen through the tube for some time. As soon as a glowing splinter held at the end of the straight calcium chloride tube is ignited, the combustion is ended. The combustion may also be conducted without passing a current of air or oxygen into the tube at the beginning, in which case the glass stop-cock is closed. Under these conditions, as soon as the substance has been heated for some time with the full flames, toward the end of the operation the glass stop-cock is opened and a current of air or oxygen passed through the tube. Substances which burn with great difficulty can also be mixed with fine copper oxide in a copper boat (see below), and then burned in the same way in oxygen.

Combustion of Substances containing Nitrogen. — Since in the combustion of nitrogenous compounds, the reduced copper spiral serving for the reduction of the oxides of nitrogen must be used, the combustion tube is charged somewhat differently in this case. The first copper roll is inserted in the tube, not 5 cm., but 15 cm., the space in front of it being reserved for the reduced spiral. Consequently the layer of coarse copper oxide is but 35 cm., and not 45 cm., in length. No change is made in the disposition of the copper oxide spiral. The ignition of the copper oxide is conducted exactly as above, except that a current of air is used.

If, however, the ignition should be conducted throughout with oxygen, at the end of the operation this is displaced by air. The further operations are the same as those described above, except that the reduced copper spiral is put in position last — just before connecting the combustion tube with the absorption apparatus. In order to prevent the oxidation of the copper, the combustion proper is performed with the glass-cock closed, and oxygen is not admitted to the tube until at the end. As soon as the oxygen is admitted, the flames under the reduced copper spiral are extinguished. The gas is passed through until it can be detected at the end of the apparatus as above described. In the combustion of substances which leave a charred, difficultly combustible, nitrogenous residue, it is necessary to burn them by mixing with fine copper oxide. Since the porcelain boats are generally too small to contain a sufficient quantity of this, a boat made of sheet copper, 8 cm. long and of a width sufficient to enable it to be just passed into the tube, is used. It is filled as follows: After it has been previously ignited, it is placed upon a sheet of black glazed paper, and half filled with fine copper oxide also previously ignited and afterwards cooled in a desiccator. Upon this is carefully spread the weighed substance from a weighing-tube as in the nitrogen determination, then a layer of fine copper oxide is added until the boat is three-fourths full: the substances are now well mixed by careful stirring with a thick platinum wire. If some of the mixture should fall upon the glazed paper, it is returned to the boat with the aid of a feather or brush. The combustion is made with the glass-cock closed. Oxygen is not admitted until at the end of the operation.

Combustion of Substances containing Sulphur or a Halogen.—Sulphur compounds cannot be burned with copper oxide in the manner described, since at a red heat the copper sulphate formed gives off sulphurous acid, which is absorbed by the potash apparatus along with the carbon dioxide, giving a result in which the percentage of carbon is too high. In this case the oxidation is accomplished with granulated lead chromate.

The filling of the tube, open at both ends, is done just as described above: copper oxide spiral, empty space for boat, long layer of lead chromate. The ignition in oxygen, etc., is also the same. But two points are here to be observed: (1) the lead chromate is not heated as strongly as the copper oxide, otherwise it fuses in the glass; and (2) the most forward portion of the lead chromate layer, nearest the calcium chloride tube (that above about three burners), is heated very slightly, since lead sulphate is not completely stable at a red heat. The substance is mixed in the copper boat with powdered, ignited lead chromate.

Halogen compounds can be burned in the usual way with copper oxide; but since the copper halides are partially volatile and give up the halogen on being heated to redness, a silver spiral must be inserted in the tube to retain the halogen. The tube is filled in the same way as for the combustion of a nitrogen compound, only in place of the reduced copper spiral, one of silver is used. But it is better to perform the combustion with lead chromate, in which case it will not be necessary to use a silver spiral. Since the lead halides are also somewhat volatile at a red heat, so, as above, the front part of the tube containing the lead chromate is heated but slightly.

Combustion of Liquids. — If the compound to be analysed is a liquid, it can be weighed directly in the porcelain boat, provided it is very difficultly volatile. Moderately volatile substances are weighed in a small glass tube which is loosely closed with a glass stopper (see Fig. 53, page 81). In order to introduce this into the tube, it is placed in the porcelain boat in such a position that the mouth of the tube is directed upwards. A preliminary trial will show whether the boat containing the empty tube will pass into the combustion tube. Very easily volatile substances are weighed in small bulb-tubes which are sealed after weighing (Fig. 62). The filling is done as follows: The empty tube is weighed, heated gently, and the open end dipped under the liquid to be analysed. On cooling, the liquid will be drawn up into the bulb.



FIG. 62.

If a sufficient quantity is not obtained the first time, the operation is repeated; before it is sealed care must be taken that the capillary contains none of the liquid; if it does, it must be removed by heating. It is now sealed, and the tube *plus* substance weighed. Care must again be taken to prevent any of the liquid from finding its way into the capillary, due to sudden movements or other causes. To prepare the tube for the combustion, the extreme end is filed and broken off, during which operation the tube is not held by the bulb. It is placed in the boat with its open end elevated and directed toward the front end of the furnace. The precaution to ascertain beforehand whether the boat loaded with the tube will pass into the combustion tube, should always be taken. If necessary, the capillary is shortened.

ELEMENTARY ANALYSIS

DENNSTEDT'S METHOD

Dennstedt's method consists in burning the substance with free oxygen¹ exclusively, in the presence of platinum as contact substance (catalyser).

In addition to the frame of a combustion furnace, and an oxygen gasometer,—Dennstedt recommends the use of two flasks provided with tubulures at the bottom and having a capacity of 5 litres,—the requisites for analysis are:

1. A hard glass tube open at both ends; length 86 cm.; diameter 18–20 mm.; also an inner tube and a small wash bottle.
2. A drying tower for the oxygen.
3. A U-shaped calcium chloride tube with two ground stop-cocks and a ground-in stopper (next to the bulb).
4. A soda-lime tower with two ground stoppers.
5. A U-shaped soda-lime-calcium chloride tube with two ground stop-cocks ("Testing-tube").

¹ Not electrolytically prepared.

6. A wash bottle for palladious chloride ("Palladium bottle").
7. A straight calcium chloride tube.
8. A strip of star-shaped platinum, and a roll of thin platinum foil.
9. A rod of hard glass for the inner tube, with a loop and platinum thread or platinum wire.
10. A porcelain boat (for the substance) divided into three parts, length 8 cm. ; also several ordinary porcelain boats (Absorption boats).
11. Rubber stoppers and seamless rubber tubing, to be used for the analysis only.

12. Chemicals: calcium chloride, soda-lime, palladious chloride, lead peroxide, minium, sodium carbonate, molecular silver.

1. **The Combustion Furnace** consists of two stands supporting three loose iron troughs. The middle one is covered with thin asbestos paper, and serves as a support for the combustion tube, while the other two troughs support the covers, the inner surfaces of which are lined with asbestos. Of these five are needed ; a large cover 25 cm. in length, and four smaller covers each one having a length of 12 cm. Of the latter, one is provided with a

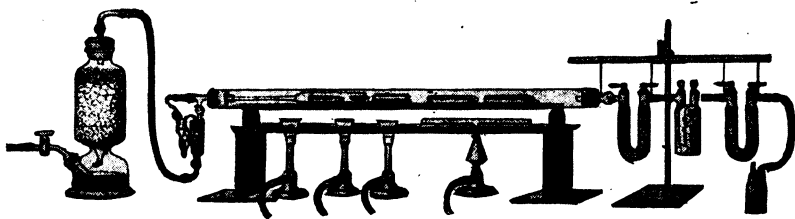


FIG. 62 A.

sliding window of mica (10 cm. : 2 cm.), through which the catalyst may be observed. The sheet of mica is renewed when it becomes opaque. The iron covers are provided with projections on the upper edges, and may be moved back and forth with crucible tongs. They should not touch the combustion tube. The combustion tube is heated with three, or preferably four, burners. The

two burners shown at the left of Fig. 62 A are used to vaporise the substance. They are Bunsen or Teclue burners to which wing burners may be attached. One burner is sometimes used for the purpose. Then follows a stronger Bunsen or Teclue burner, provided with a wing burner and a regulation device. This is placed under the contact star and supplies the combustion flame. The rear of the tube (the part preceding the absorption apparatus) is heated with a movable flame tube consisting of 20 non-luminous flames. During the combustion a temperature of $300-320^{\circ}$ is necessary in this part of the tube. The height of the flames that will produce this temperature is determined in a preliminary experiment by inserting a thermometer in the combustion tube.

2. Drying Tower for the Oxygen.—Pure concentrated sulphuric acid is poured into the lower part of the tower (Fig 62 A, left) to a height of 2 cm., so that the end of the conducting tube, which is 2 mm. wide, dips into this to a depth of $\frac{1}{2}$ –1 cm. Ligatures (string or wire) are bound around the joints. A small funnel, with its tube bent sidewise, is introduced into the lower narrow part of the tower. The funnel is covered with a little glass-wool or cotton, and the lower half of the cylindrical tower (about 10 cm.) is filled with coarse, sifted soda-lime; the upper half is filled with calcium chloride. Then follows a layer of glass-wool or cotton. Concerning the quality of the soda-lime and the calcium chloride, see below (Filling of absorption apparatus).

3. The Small Wash Bottle.—A plug of cotton is placed at the bottom of the wide tube attached to the small wash bottle (Fig. 62 B); it is then filled with calcium chloride and covered with a layer of cotton (see 4 a). The lower narrow part of the small wash bottle is filled with concentrated sulphuric acid. A capillary tube is used for this purpose. The bottle is held in an inverted position; the tip of the capillary is now inserted as far into the U-shaped curve as possible, and by proper manipulation the acid is allowed to flow through the inner bulb into the lower narrow end. The limb nearest the calcium chloride tube is dried with blotting-paper. The small wash bottle must be constructed so that there is a space of at least 1 mm. between the inner bulb and

the outer tube. The connecting parts of the apparatus may be made tight by moistening the two holes of the stopper, as well as its lower surface, with a saturated solution of calcium chloride. In order to regulate the stream of oxygen as perfectly as possible by the aid of the three stop-cocks of the small wash bottle, the perforations of the cocks are filed on both sides with a sharp triangular file, as seen in Fig. 62 c. The calcium chloride is renewed from time to time, and the stoppers are kept closed when the apparatus is not in use.

4. Absorption Apparatus.—(a) *Calcium Chloride Tube.* It contains two ground stop-cocks, a small ground glass stopper,

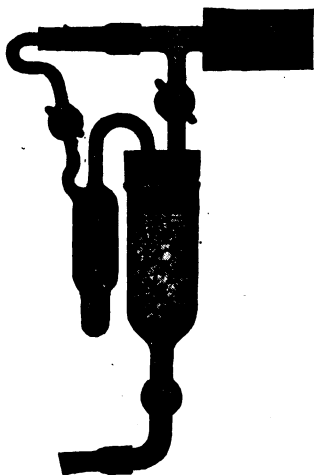


FIG. 62 B.



FIG. 62 C.

and the bulb in which a greater part of the water condenses. The previously sifted calcium chloride is poured into a wide test-tube, clamped in a slanting position, and carefully heated with a free flame until no more water deposits in the cooler part of the tube. The heating must be done cautiously, and only for a short time. In filling the U-tube care must be taken that no dust particles or calcium chloride settle in the side-tubes; to prevent this the side-

tubes are plugged with cotton during the filling. The cotton is removed with forceps after filling the tube. A short roll of paper may also serve the same purpose, protecting not only the side-tubes but also the ground part of the U-tube. A roll of cotton is finally introduced into the calcium chloride tube. The ground parts of the tube are then cleaned and slightly oiled. Before the newly filled tube is used, dry carbon dioxide is passed through it (see page 103). The carbon dioxide is then displaced by oxygen.

(b) *Soda-lime Tower.* Commercial soda-lime is often too dry and does not therefore readily absorb carbon dioxide. Before filling the absorption apparatus a few grammes are carefully heated in a test-tube over a free flame. Much water should condense in the cooler parts of the tube. If this is not the case, the entire quantity is moistened with the necessary amount of water (spray). The soda-lime must be freed from small particles by sifting, and care must be taken that no small particles settle in the side-tubes. The precautions outlined for calcium chloride are followed here. As soda-lime expands by the absorption of carbon dioxide, in order to avoid the danger of breaking the apparatus care is taken not to use too much. The perforations of the glass stoppers are then filled with dry cotton; the ground surfaces are very carefully cleaned, and slightly greased. When it is desired to refill a tower that has been used, the stoppers are removed, the apparatus is freed from grease, and dipped for several hours in water that has been slightly acidified with hydrochloric acid. During the combustion the tower is always kept in the same position; in order to prevent mistakes the side-tubes are marked (arrow, or coloured glass-button).

(c) *Soda-lime-calcium Chloride Tube ("Testing-Tube").* One limb of this tube is filled with soda-lime and the other with calcium chloride. The precautions mentioned in (a) and (b) are observed. The testing-tube is inserted in such a manner that the soda-lime limb is next to the soda-lime tower.

(d) *The Weighing of the Absorption Apparatus.* The three absorption tubes are always filled with oxygen before weighing. When used for the first time, dry oxygen is conducted into the

apparatus until all the air has been driven out. (Test : vigorous flaming of a glowing splinter.) One of the two stop-cocks in each apparatus is opened for a moment before weighing, in order to equalise the differences in pressure. In weighing after a combustion, the temperature should be as nearly the same as it was before combustion. In most cases it is sufficient to carry out the final weighing when the apparatus has remained in the balance room for one or two hours. It is safer to check the weighing on the following morning. In winter-time the temperature of the balance room should be kept uniform over night. Should a long period elapse between two analyses, the apparatus is again filled with oxygen.

(e) *The Palladious Chloride Wash Bottle (" Palladium Bottle")* is half filled with a clear dilute water solution of palladious chloride of a straw-yellow colour. The solution is filtered when it becomes cloudy. The separation of palladium (by the action of carbon monoxide) indicates an incomplete combustion. In addition to this the solution is also of service in showing the rate of flow of the gas stream.

5. The Combustion Tube and Accessories.—All combustions are carried out by the use of a duplex supply of oxygen. For this purpose a T-tube is inserted in the stopper in the rear end of the combustion tube. The vertical limb of the T-tube is provided with a stop-cock, and is connected with the calcium chloride tube of the small wash bottle as shown in Fig. 62 B. Through the horizontal part of the T-tube passes the capillary of the inner tube ; these two are connected with a short rubber tubing (ligature). The wider part of the inner tube fills the space inside the combustion tube almost completely, and serves as a receiver for the boat containing the substance taken for analysis. In order to prevent the fusion of the inner tube into the combustion tube, a coil of fairly thick platinum wire is placed around the wide, open end of the former. Furthermore, in order to prevent the formation of an explosive mixture by the combustible vapours and the oxygen, a rod of hard glass is introduced into the inner tube. The rod is of a size that nearly fills the empty space in the inner

tube. It is provided with a loop in the fore-part; to this is attached a roll of platinum wire which reaches the open end of the inner tube. Or, instead of the platinum roll, a thin platinum wire is wound around the rod. This arrangement will also prevent the rod from fusing. The glass rod is unnecessary in most instances. Its place is taken by a roll of thin platinum foil. This is placed at the opening of the inner tube in such a way that one half rests in the inner tube, the other half in the combustion tube; then follows the contact star. Vapours of high boiling substances will sometimes condense and run into the inner tube. In order to protect the tube, it is covered to within 1 cm. of the opening and over one-third of the inner surface with a strip of thin asbestos paper. This must be heated with the combustion tube in a stream of oxygen during the drying of the apparatus. (In the simultaneous estimation of sulphur the asbestos is not used.) Before the analysis, the previously washed and dried combustion tube, the inner tube (including the glass rod), the platinum coil, and the platinum star are heated for a while in a moderate stream of oxygen at 250–300°. During the heating the burners are evenly distributed, and the covers are placed in position. The opening at the forward end is finally closed with a straight calcium chloride tube, and the combustion tube is allowed to cool in a stream of oxygen. As the bulky Dennstedt tubes break more readily than those used in Liebig's method, special care is exercised in the heating as well as the cooling of the former, and in exposing them to sudden changes of temperature. It is also necessary, at all times, to prevent the covers from coming in contact with the hot tube.

6. The Combustion Proper.—In combustions by Liebig's method it has not been found possible to prescribe details that will cover all cases. This is still less possible in Dennstedt's method, in which the number of modifications and combinations is much greater. A general outline of the method will be given for substances containing carbon, hydrogen and oxygen, and Dennstedt's own words will be used in part.

While the combustion tube is being heated and cooled in a

stream of oxygen, the absorption apparatus is weighed (a stop-cock is opened for an instant) ; the boat having the three divisions is also weighed, first empty, and then with the substance. The supply of oxygen is now cut off, and the inner tube (with small wash bottle, etc.) is withdrawn by removing the stopper at the rear end of the combustion tube. The long rubber connection between the drying tower and the small wash bottle is not removed. The boat containing the substance is introduced into the inner tube, and pushed in, as far as possible. Then follows the glass rod with its platinum wire, or the platinum roll alone, as described above, and the apparatus is securely attached to the combustion tube. The different parts of the absorption apparatus are finally connected by the use of short, thick-walled, seamless rubber joints. The rubber should be drawn over the ends of the glass tubes until they touch. The limb of the U-tube containing soda-lime should be kept next to the soda-lime tower. The drying apparatus is most conveniently connected with the combustion tube in the following manner : A rubber stopper is first placed on the calcium chloride tube (at the bulb end) ; the tube is then connected with the soda-lime tower, which is in turn attached to the testing-tube ; and finally, the end of the combustion tube is securely closed by the rubber bearing the calcium chloride tube. Then follows the palladium bottle.

Before the combustion proper, a test as to whether the apparatus is perfectly tight is conducted in the following manner : All the stop-cocks, from the stopper of the gasometer to the last stopper of the U-tube (testing-tube) are closed. The stop-cock of the gasometer and the lower stop-cock of the drying tower are now opened, so that a few gas bubbles pass through the sulphuric acid. If the joints are perfectly tight, the bubbling will stop as soon as the pressure between gasometer and drying tower is equalised. If the bubbling continues, then there is a leak which must be prevented. The upper stop-cock of the drying tower is now opened, and left open until the bubbling stops. This is continued to the last stop-cock of the testing-tube. During this test, the stop-cock of the small wash bottle containing sulphuric acid is opened

in such a way that not more than two bubbles per second escape. The combustion is commenced when the apparatus is found to be perfectly tight. "For its success it is absolutely necessary to thoroughly mix the vapour of the burning compound when it reaches the contact substance (platinum) with as much oxygen at every instant as is required for its complete combustion, *i.e.* there should always be present an excess of oxygen. In the combustion of volatile substances, in order to mix the vapour with the proper quantity of oxygen, the oxygen stream is introduced in two parts; one part is known as the slow *vaporising stream*, which sweeps over the burning substance, and carries away its vapour in a regular manner; the other part is known as the *combustion stream*, which meets the vaporising stream near the glowing platinum." The former is regulated through the stop-cock of the small wash bottle (Fig. 62 B, upper left hand), the latter through the stop-cock of the calcium chloride tube (Fig. 62 B, upper right hand). The lower stop-cock of the calcium chloride tube permits the passage of both streams. The small wash bottle shows the velocity of the vaporising stream, and the palladium bottle that of the combustion stream. Whenever the velocity of these streams is specified, *e.g.* by the number of bubbles for every 10 seconds, it is understood that the bubbles in the small wash bottle are much smaller than those in the palladium bottle. The outer stream is regulated in such a manner that in 10 seconds 10-15 bubbles pass through the palladium bottle. "When the vaporisation of the substance becomes too rapid, the velocity of this stream may be temporarily doubled, or increased even more, without hesitation." The inner stream is so regulated that if the substance is easily volatile, 5-10 bubbles pass through the small wash bottle every 10 seconds, and if the substance is not easily volatile 10-30 bubbles pass through the wash bottle every 10 seconds. The velocity of the inner stream should not be altered in the early part of the combustion. A small flame is first lighted under the contact star. This is gradually increased, until finally the star appears to glow brightly through the mica window of the small cover. The flame, the small cover, and the end of the inner tube lie in the same vertical plane. The

forward empty part of the combustion tube is heated at the same time to about 300° with the burner tube, and a large cover is placed next to the small cover with the mica window. "The burner used to vaporise the substance is turned on almost entirely, and burns with a moderately high flame. It is now placed at the rear and kept so far away from the boats that even very volatile substances are just vaporised without undergoing much change. When the contact substance begins to glow brightly, the vaporising flame is quickly moved forward, about 1 cm. every 3-4 minutes, depending upon the volatility of the substance, until the latter melts or begins to decompose. With substances that do not vaporise or decompose readily, the flame is brought near the boat. It is then allowed to remain undisturbed. An observation will now show whether the contact substance is glowing, whether there is condensation of water at the forward end of the tube, in short, whether combustion proper has started. If this is the case, everything is left unchanged for 15 minutes. If combustion does not progress smoothly at the end of this period, the combustion flame and the cover are moved 1-2 mm. to the rear. This is repeated after a short time, until it is found that, depending upon the nature of the substance, it begins to burn completely, *i.e.* it first melts, and then chars, etc., even in the first division of the boat. Should the decomposition slow down, the combustion flame and the cover are slowly, millimetre by millimetre, moved backwards, until finally there is complete vaporisation, decomposition, or charring. *The combustion flame should never be moved far enough to diminish the glow of the platinum at the open end of the inner tube.* The cover, however, may be moved back and kept directly above the boat. The position of the burners need not be changed when the contact mass is seen to glow brightly, since this is only an indication that the combustion is progressing briskly. But if the glowing is very sudden and too active, or, if flames appear at the open end of the inner tube, both burners are pushed backwards in order to prevent rapid decomposition, and the vaporising flame is again brought near the boat in order to secure normal conditions once more. The combustion flame is also moved back gradually to the extreme

limit. Should this stop the progress of the combustion, the vaporising flame is moved forward, and the cover at the rear of the tube is put in position. When the contents of the boat have either entirely disappeared, or are completely carbonised, the inner stream of oxygen is increased, and the tube heated to redness." The part of the tube adjacent to the absorption apparatus is finally covered with a small cover.

When the combustion is ended, the flames are first lowered and then turned off entirely, and the apparatus is cooled in a stream of oxygen. The palladium bottle is now removed. *The lower stop-cock of the drying tower is first closed*, and the absorption apparatus is taken apart after turning off the stop-cocks. The small stopper near the bulb of the calcium chloride tube must not be forgotten. Concerning the weighing of the apparatus see directions given above. Under normal conditions, the soda-lime-calcium chloride tube should not increase in weight more than a few milligrammes. An increase in weight amounting to more than 10 mg. indicates that the soda-lime tower is exhausted and needs refilling. In this case its original position should not be reversed by oversight.

Very often in dealing with substances that are not very volatile, it is more convenient to use two vaporising flames in place of one. Even in this case the combustion flame under the platinum star is first lighted. The flame is not placed under the substance, but is allowed to remain under the contact star. The vaporising flame at the rear is also kept, as before, as far away from the boat as possible; its position is changed only towards the end of the combustion. In this case the substance is vaporised by the forward vaporising flame, which is kept very low at first and placed near the combustion flame, and is slowly brought near the boat with a gradual increase of the flame, the time required for this being the same as that given above for the combustion flame. As soon as the forward flame is brought under the boat, the rear flame is also pushed towards the substance.

It is to be noted once more that the rate of the combustion depends upon the number and the height of flames, the use of

burners without or with wing burners, their distance from the boat, the use of covers, the velocity of the oxygen stream, etc. These are conditions that may be readily controlled.

Should the palladious chloride become very black during the combustion, the analysis may be considered unsuccessful. Under these conditions either the absorption apparatus is removed, and the tube is heated to glowing in the presence of a stream of oxygen, as described above (the absorption apparatus being re-filled with oxygen for a new analysis), or the combustion is brought to completion as under normal conditions. The weight of the absorption apparatus is, of course, worthless in this case. A combustion is also considered unsuccessful when the substance or its decomposition products sublime, or distil beyond the contact star. If this is the case, the absorption apparatus is immediately removed, and the tube is heated to glowing in the presence of a stream of oxygen. If necessary, the absorption apparatus is freshly prepared. Finally, weak or strong explosions may occur inside the tube during the combustion. If these do not produce a sublimate, or a distillate, and cause no blackening of the palladious chloride solution, then the analysis, including the final weighing, is brought to a normal end. If analyses give persistently inaccurate results, a blank combustion is carried on (without substance) to ascertain the defect in the apparatus.

The Combustion of Substances containing **Nitrogen** or **Sulphur**, or both at the same time, is in principle exactly the same as described above, with the only modification that, in order to absorb the nitric oxide, the sulphurous acid, and sulphuric acid, a mixture of minium and lead peroxide is placed in the combustion tube and heated to 320–350° with the flame-tube.

The quality of lead peroxide. Lead peroxide especially prepared for the Dennstedt analysis is placed on the market by the firms of Merck and Kahlbaum under the label of "Lead Peroxide according to Dennstedt." But Dennstedt's investigations show that even these pure preparations almost always contain small quantities of organic substances, such as fibres, dust particles, etc., which must be destroyed before the substance is used. This

is done by drying a large quantity on a watch crystal in an oven, at a temperature of $120-140^{\circ}$, and heating this for at least half an hour, or longer, in a hard glass tube in a stream of oxygen at $320-350^{\circ}$. For the preparation of minium the required amount of pure lead peroxide is heated in a stream of air at $400-450^{\circ}$, or in an open porcelain dish, until the dark-brown colour is changed to a dull red; it is then allowed to cool in a desiccator. Equal quantities of lead peroxide and minium are intimately mixed and used for the analysis. The quantity used for each analysis is 7-8 grammes (not over 10 grammes) of the mixture, weighed out to within $\frac{1}{2}$ gramme, by the use of ordinary weights. The substance is introduced into the tube in three small boats, the boat in the rear containing the main portion. The boats should be in the tube during the heating of the latter in a stream of oxygen.

When a substance contains sulphur, the boat in the rear is pushed to within 6-7 cm. of the contact star, but when sulphur is not present, the boat is kept at a distance of 8-10 cm. from the contact star. Under these conditions the heating in a stream of oxygen ($320-350^{\circ}$) is done very cautiously in the course of at least a half hour, or better still, a whole hour, or more. The course of the combustion is otherwise the same as that described above. A rapid current of oxygen is avoided as much as possible, otherwise complete absorption by lead peroxide will be prevented.

Should the platinum (star, roll, etc.) come in contact with lead peroxide or minium at any time, the particles of the latter are removed first mechanically, and then by the use of hot dilute hydrochloric acid.

If a substance explodes on heating, it is mixed in the boat with quartz-powder or quartz-sand (previously treated with hot hydrochloric acid and heated to glowing), or with lead chromate which has been heated to redness.

Simultaneous Determination of Sulphur with Carbon and Hydrogen.—In this case the lead peroxide-minium mixture should naturally be free from sulphates. In order to test the substance for sulphuric acid 20-25 grammes of the mixture is digested on a

water-bath for one hour with a 20 % solution of sodium carbonate, free from sulphuric acid. The mixture is crushed and stirred frequently. It is then filtered, washed with 50 c.c. of water, and acidified with hydrochloric acid. The hot solution is now treated with barium chloride. The liquid should remain perfectly clear, even after standing overnight.

At the end of the combustion, and after the removal of the absorption apparatus, the boats are carefully drawn out by the use of a strong copper wire with a little hook at one end. The surface of the substance in the first two boats will be covered with a white layer of lead sulphate. The contents of the boats are transferred into a beaker; the empty boats are placed into a wide test-tube and heated on a water-bath with a 5 % solution of pure sodium carbonate; the liquid is added to the main portion of lead peroxide and the boats are once more washed with a dilute solution of sodium carbonate. Traces of sulphuric acid may also adhere to the contact substance, the roll, the combustion tube and the inner tube. These are, therefore, rinsed freely with water. The mouth of the inner tube is also dipped in a few cubic centimetres of water, but the small wash bottle is not removed. If the salt of a sulphonic acid was analysed the combustion boat is also washed with water, or better still, it is treated, with the boats, with a solution of sodium carbonate. In this case the wider portion of the inner tube is also rinsed with water. All the washings containing sulphuric acid are poured into the beaker, which holds the contents of the boats. The mixture should give an alkaline reaction. It is heated over an actively boiling water-bath for at least one hour. It is frequently stirred, and the precipitate is crushed with a glass rod which has been flattened out at one end.

The sulphuric acid is determined quantitatively as barium sulphate by one of the following two methods: The entire liquid is separated from the precipitate by filtration. The latter is washed with water. Should the filtrate show turbidity by this treatment, it is once more poured through the same filter. The clear filtrate is then acidified with dilute hydrochloric acid, and is finally precipitated at the boiling point with barium chloride by the usual method.

The second method is that proposed by Dennstedt. The contents of the beaker (solution and precipitate) are transferred into a measuring cylinder. (The beaker must be rinsed.) The volume is now increased to a convenient round number by the addition of water. Suppose the volume is 100 c.c. Since the specific gravity of the precipitate is about 7 (if 7 grammes of lead peroxide were used), then in 100 c.c. of the mixture we have 1 c.c. of the precipitate and 99 c.c. of the liquid. To simplify the calculation another cubic centimetre of water is added so as to make the liquid volume just 100 c.c. An aliquot part is then taken, *i.e.* 95 c.c., and filtered through a dry filter as described above. The washings from the precipitate are also saved. After acidifying with hydrochloric acid, the sulphuric acid is precipitated as barium sulphate. From the quantity of barium sulphate in 95 c.c. of the liquid the amount in 100 c.c. is calculated; in other words, from the quantity in the partial volume that in the total volume is calculated.

Determination of Sulphur alone. — It is evident that the method described above may also be used for the determination of sulphur alone, by leaving out the absorption apparatus. It is, however, simpler and more convenient to use anhydrous sodium carbonate as an absorption medium, in place of lead peroxide. The boats are filled with pure, calcined sodium carbonate (free from sulphuric acid). The substance is gently pressed down with a spatula. The combustion tube containing the boats is heated in a stream of oxygen, the sodium carbonate being heated to about 400° with the flame-tube. The absorption apparatus is replaced by a straight calcium chloride tube which is connected with the palladium bottle. At the end of the combustion the cooled boats are drawn out of the tube by means of a bent wire, and their contents are transferred into a beaker. The boats are placed in a wide test-tube and treated with hot water. The liquid is added to the dry sodium carbonate. The combustion tube, the platinum star, the inner tube, and the combustion boat are rinsed with water. In short, the process described above is exactly followed. In order to change traces of sulphites in the alkaline solution into

sulphates, it is heated and then treated with a few cubic centimetres of a saturated solution of bromine water. The beaker is now covered with a watch crystal, and the solution is *carefully* acidified (evolution of carbon dioxide) with moderately strong hydrochloric acid. The solution is heated until the yellow colour, due to excess of bromine, disappears. It is then treated hot with barium chloride.

This method is so easy and convenient after a few trials, that it seems to be destined to finally replace the Carius method. The analysis, including the weighing of the barium sulphate, may be usually completed without difficulty in half a day. The method has the further advantage over Carius' method that whereas in the latter oxidation is sometimes incomplete with substances like sulphonic acids or sulphones, the difficulty is not experienced in this method. If a sufficient quantity of the substance is on hand, it is much more advantageous to carry on a separate determination for sulphur by the sodium carbonate method, and ignore the results from lead peroxide in the *simultaneous* determination of sulphur, carbon and hydrogen.

If the substance contains a small percentage of sulphur, the addition of barium chloride may not precipitate barium sulphate, even after long heating, due to the solubility of the latter in excess of barium chloride. But the precipitate will form when the solution is allowed to stand overnight. If a larger quantity of sulphur is present, and a precipitate forms as soon as barium chloride is added, the precipitate is filtered in the usual way, after the solution has become clear. The filtrate is then allowed to stand for a day so as to recover any precipitate that may form. The same procedure is also followed in the lead peroxide method.

Analysis of Substances containing a Halogen. — If an organic substance contains a halogen in addition to carbon, hydrogen and oxygen (but no nitrogen or sulphur), a simultaneous determination of carbon, hydrogen and halogen may be carried out by the use of "molecular silver" (Kahlbaum). A long porcelain boat is filled with a few grammes of molecular silver. During the heating of the combustion tube this is also heated by the flame-tube, a

somewhat higher temperature being used than that prescribed for lead peroxide.

The warm boat is then removed and transferred into a weighing tube (preferably a tube with two short legs near the open end), and is weighed after cooling. During the combustion the boat is kept at a distance of about 6 cm. from the contact star, and is heated to 300–350° with the flame-tube. At the end of the analysis the weight of the silver boat gives directly the amount of halogen in the substance.

Halogen compounds, and especially compounds very rich in halogen, burn much more slowly than halogen-free substances. On this account they easily distil over the contact star without decomposition, and the typical glow of the platinum, as well as the blackening of the tube, are not noticed. When this is the case the combustion is carried out very carefully and slowly, and the oxygen is not allowed to stream too rapidly.

The method described above cannot be used directly if the halogen compound contains nitrogen or sulphur, due to the formation of silver nitrite and silver nitrate or silver sulphate. For the determination of halogen in this case, as well as the determination of sulphur and halogen (by the use of lead peroxide) see Prof. Dr. M. Dennstedt's "Anleitung zur vereinfachten Elementaranalyse" (Hamburg, Otto Meissners Verlag, 2. Aufl., 1906).

Calculation of the Atomic Formula. — In order to calculate the simplest formula of a substance from the figures giving its percentage composition, the method is as follows: If a substance contains, *e.g.*:

Carbon = 48.98 %

Hydrogen = 2.72 %

Chlorine = 48.30 %

the percentage figures are divided by the corresponding atomic weights. There is thus obtained:

$$48.98 \div 12 = 4.08 \text{ C}$$

$$2.72 \div 1 = 2.72 \text{ H}$$

$$48.30 \div 35.5 = 1.36 \text{ Cl}$$

These figures are divided by the smallest — in this case 1.36 :

$$4.08 \div 1.36 = 3 \text{ C}$$

$$2.72 \div 1.36 = 2 \text{ H}$$

$$1.36 \div 1.36 = 1 \text{ Cl}$$

The simplest atomic formula, therefore, is $\text{C}_3\text{H}_2\text{Cl}$. If the numbers obtained in the last division are not integers, they are multiplied by the smallest integer which will convert the fractions into whole numbers. If, *e.g.*, the following numbers have been found, 1.25, 1.75, and 0.5, they are multiplied by 4, the results being 5, 7, and 2.

The simplest formula thus obtained from the analytical data does not always correspond with the true molecular weight. This must be determined by one of the usual methods, unless it may be inferred from the nature of the reaction by which the substance analysed was produced.

The exact atomic weights of the elements used in analytical calculations are :

$$\text{H} = 1.008$$

$$\text{C} = 12.00$$

$$\text{N} = 14.01$$

$$\text{S} = 32.07$$

$$\text{Cl} = 35.46$$

$$\text{Br} = 79.92$$

$$\text{I} = 126.92$$

$$\text{O} = 16.00$$

These figures are taken from Kuster's "Logarithmischen Rechen-tafeln für Chemiker."

SPECIAL PART

I. ALIPHATIC SERIES

1. REACTION: THE REPLACEMENT OF AN ALCOHOLIC HYDROXYL GROUP BY A HALOGEN

1. EXAMPLE: Ethyl Bromide from Ethyl Alcohol¹

To 200 grammes (110 c.c.) of concentrated sulphuric acid contained in a round litre-flask, add quickly with constant shaking, without cooling, 90 grammes of alcohol (about 95%). After cooling the mixture to the room temperature, add carefully 75 grammes of ice-water, the cooling being continued, and then 100

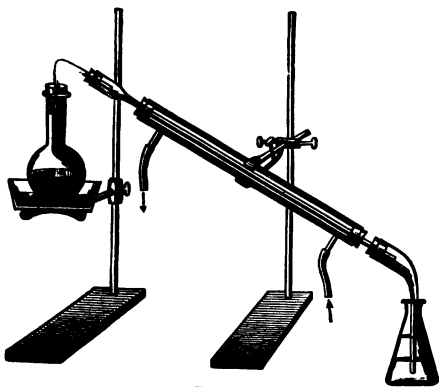


FIG. 63.

grammes of finely pulverised potassium bromide (see Hydrobromic Acid, page 379). The mixture is subjected to distillation, which must not be too slow, the flask being heated on a small sand-bath with a large flame (Fig. 63). Since the boiling-point of ethyl bromide is low (38°); a long condenser, with a

quite rapid current of water passing through it, is used. An upright coil condenser (see Fig. 27, page 34) may be employed advantageously. At the beginning of the operation, the receiver is filled with a sufficient amount of water containing a few pieces of ice to allow the end of the adapter to dip under the surface. The reaction is ended as soon as the oily drops which sink to the bottom of the receiver cease passing over. If, during the distillation, the contents of the receiver should be drawn up into the con-

¹ J. 1857. 441.

denser, this difficulty may be overcome by placing the receiver in such a position that the end of the adapter reaches just below the surface of the water. The same result may be attained by turning the adapter to one side, so that air may enter it. The lower layer of the distillate consisting of ethyl bromide is washed in the receiver several times with water, and finally with a dilute solution of sodium carbonate, during which the flask must not be closed. The lower layer is then run out of a separating funnel, dried with calcium chloride, and finally distilled, the same precautions as to cooling, mentioned above, being observed. In this case a very small flame of a microburner is used, or the burner tube of a Bunsen burner is removed and a luminous flame, a few millimetres high, is lighted at the nozzle. The ethyl bromide distils between $35-40^{\circ}$, the main portion at $38-39^{\circ}$. In consequence of the low boiling-point of ethyl bromide, it is never allowed to stand in open vessels for any length of time; during the drying over calcium chloride, the flask must be closed by a tight-fitting cork. The finished preparation, particularly at summer temperature, must not be preserved in thin-walled vessels of any kind, but always in thick-walled, so-called specimen bottles. Yield, 70–80 grammes.

At the conclusion of this experiment, as well as in all the following preparations, the amount of actual yield is compared with the percentage required by theory. The following points are therefore noted: According to the chemical equation, one molecular weight of potassium bromide (119) requires one molecular weight of alcohol (46). In actual work, especially in a large number of organic reactions which do not take place quantitatively, and where economy is to be considered, one of the reacting substances is taken in excess, in accordance with the Law of Mass Action. Since the price of one kg. of potassium bromide is about 4 marks, and that of one kg. of alcohol (including duty) is about 1.80 marks, — duty-free alcohol 0.90 mark, — the price of one molecule of potassium bromide (119×4) compared with that of one molecule of alcohol (including duty) (46×1.80), or duty-free alcohol (46×0.90), is in the ratio of 6 : 1, or 12 : 1. From the standpoint of economy it is more rational to use an excess

of the cheaper alcohol and convert the more expensive potassium bromide, as completely as possible, into ethyl bromide. The quantities used in the above experiment are in accordance with these considerations. 100 grammes of potassium bromide require theoretically 39 grammes of alcohol, while the quantity of alcohol used is 86 grammes (90 grammes of 95 %), *i.e.* more than twice the theoretical amount. In order to calculate the yield theoretically possible, the quantity of potassium bromide must be taken as the basis, since the conversion of the total amount of alcohol into ethyl bromide is impossible. When the alcohol used for the preparation of its corresponding bromide is more expensive than potassium bromide, then the substance to be taken in excess is naturally potassium bromide. (Compare the discussion of the Law of Mass Action under Acetic Ester.)

The ethyl bromide thus obtained is contaminated with a small amount of ether. If it be desired to remove this, the well-cooled crude ethyl bromide, contained in a flask surrounded by a freezing mixture of ice and salt, is treated, before drying with calcium chloride, with concentrated sulphuric acid, added drop by drop and with frequent shaking, until it separates out in a layer under the ethyl bromide. The acid containing the dissolved ether is then run off (in a separating funnel) from the ethyl bromide, which is shaken up several times with ice water, dried with calcium chloride, and redistilled as before. For the preparation of ethyl benzene (which see) the ethyl bromide need not be thus purified.

2. EXAMPLE: Ethyl Iodide from Ethyl Alcohol ¹

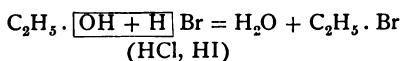
To a mixture of 5 grammes of red phosphorus and 40 grammes of absolute alcohol, contained in a small flask of about 200 c.c. capacity, 50 grammes of finely pulverised iodine are added gradually in the course of a quarter hour; the flask is frequently shaken during the addition, and cooled from time to time by immersion in cold water. An air condenser — a straight vertical glass tube is the common form — is connected with the flask, and the reac-

¹ A. 126, 250.

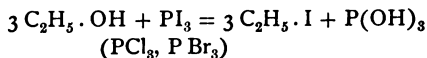
tion mixture is allowed to stand at least four hours. (In case the experiment was begun late in the afternoon, it may stand until the next day.) In order to complete the reaction, the mixture is heated for two hours on the water-bath, a reflux condenser being attached to the flask. The ethyl iodide is then distilled off conveniently by immersing the flask in a rapidly boiling water-bath. The distillation is facilitated by the use of a frayed thread (see page 34). Should the last portions go over with difficulty, the water-bath is removed, the flask is dried and heated for a short time with a luminous flame kept in constant motion. The distillate, coloured brown by iodine, is washed several times with water to free it from alcohol, and then with water to which a few drops of caustic soda have been added, to remove the iodine; the colourless oil is now separated in a dropping funnel, dried with a small quantity of granular calcium chloride, and distilled. If the calcium chloride should float on the oil, it is poured through a funnel containing some asbestos or glass-wool into the fractionating flask. The boiling-point of ethyl iodide is 72° . Yield, about 50 grammes.

The following questions are to be answered at this point : Which one of the reacting substances was taken in excess ? Why ? What percentage of the theoretical yield of ethyl iodide is obtained ?

Both of these reactions are special cases of a reaction of general application, viz. : the replacement of an alcoholic hydroxyl group with a halogen atom. This may be effected in two ways : (1) by causing the alcohol to react with the halogen hydracid as in the preparation of ethyl bromide, *e.g.* :

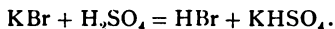


(2) by treating the alcohol with a phosphorus halide, as in the preparation of ethyl iodide, *e.g.* :



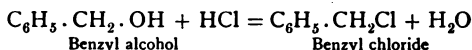
1. The first reaction takes place most easily with hydriodic acid ; in many cases saturation with the gaseous acid being sufficient to induce the reaction. Hydrobromic acid reacts with greater difficulty ; it is

frequently necessary to heat the alcohol saturated with the acid in a sealed tube. The above method for the preparation of ethyl bromide is a simple case of the general reaction. In place of using hydrobromic acid directly, it can in some cases, like the one given, be generated by warming a mixture of potassium bromide and sulphuric acid:



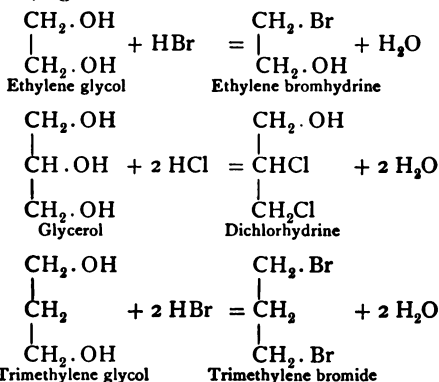
Hydrochloric acid reacts with most difficulty, and it is, *e.g.*, in the preparation of methyl chloride and ethyl chloride, necessary to employ a dehydrating agent — zinc chloride is the best — or with the alcohols of high molecular weights, to heat in a closed vessel under pressure.

This reaction is not only applicable to the aliphatic, but also to the aromatic alcohols; *e.g.*:

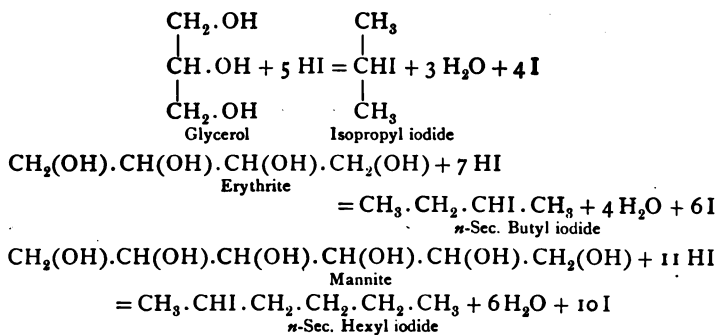


But phenol hydroxyl groups cannot be replaced by the action of a halogen hydricid.

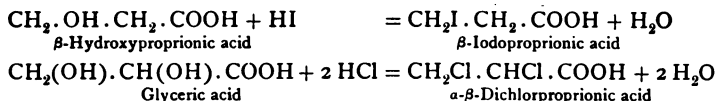
With di-acid and poly-acid alcohols the reaction takes place, at least with hydrochloric and hydrobromic acids; but, in this case, the number of hydroxyl groups which are replaced by the halogen depends upon the conditions of the experiment, the quantity of the halogen acid, the temperature, etc., *e.g.*:



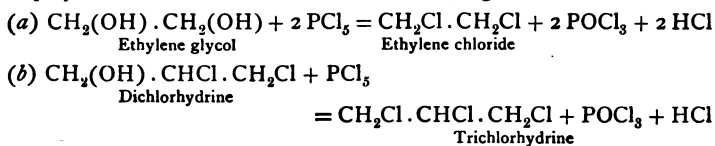
Hydriodic acid, in consequence of its reducing properties, acts upon the poly-acid alcohols in a different manner. A *single* hydroxyl group, and that particular one in combination with a carbon atom which is in turn in combination with other carbon atoms, is replaced by iodine, while at times other hydroxyl groups are replaced by the hydrogen of the acid, *e.g.*:



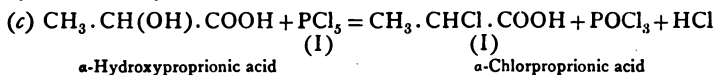
With derivatives of alcohols, *e.g.* alcohol-acids (hydroxy acids) the first reaction takes place:



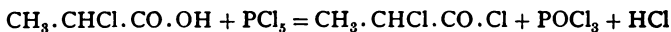
2. The second reaction takes place much more energetically, especially when a phosphorus halide which has been previously made is used. This is not always necessary, at least in introducing bromine and iodine; in many cases it is better to generate the phosphorus halides in the course of the reaction, by adding to the mixture of alcohol and red phosphorus either bromine from a dropping funnel, or as above, finely pulverised iodine. This reaction, as well as the first, is applicable to poly-acid alcohols and substituted alcohols, *e.g.*:



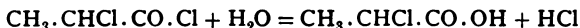
This example illustrates the more energetic action of the phosphorus halide as compared with the corresponding hydrogen halide; it is impossible to replace the third hydroxyl group of glycerol with chlorine by the use of hydrochloric acid.



In cases of this kind a complication arises, due to the fact that the phosphorus halide also acts upon the hydroxyl of the carboxyl group, replacing it with the halogen, giving rise to an acid-chloride:

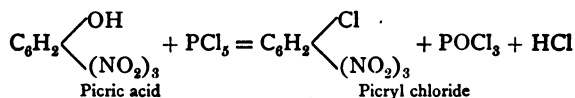
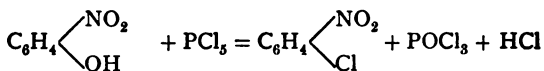
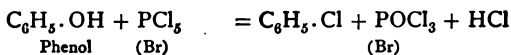


The acid may be regenerated by treating the acid-chloride with water:

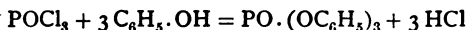


The action of phosphorus iodide on poly-acid alcohols is similar to the action of hydriodic acid referred to under Reaction 1.

The more energetic action of the phosphorus halides may also be perceived in the fact that phenol hydroxyl groups can be replaced by a halogen, by the use of the phosphorus compounds, which, as mentioned above, is impossible with the halogen hydracids, *e.g.*:



But the quantities obtained are much less satisfactory, the reason being that the phosphorus oxychloride attacks the unacted-upon phenol, forming phosphoric acid esters, *e.g.*:

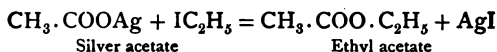
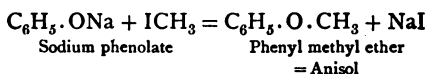
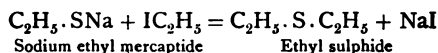
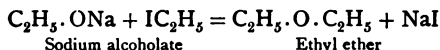


In this way a large portion of the phenol is withdrawn from the main reaction.

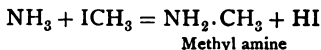
The monohalogen alkyls $\text{C}_n\text{H}_{(2n+1)}\text{Cl}(\text{Br}, \text{I})$ are in most cases colourless liquids, the exceptions being methyl and ethyl chlorides and methyl bromide, which are gaseous at the ordinary temperature, and the members of the series having high molecular weights, like cetyl iodide, $\text{C}_{16}\text{H}_{33}\text{I}$, which are semi-solid, salve-like substances. The iodides are only colourless when freshly prepared; on long standing, especially under the influence of light, a slight decomposition, resulting in the separation of iodine, takes place, imparting to them a faint pink colour at first, which becomes brownish red after a long time. This decomposition can be prevented if some finely divided, so-called molecular silver is added to the liquid. A coloured iodide can be made colourless by shaking it with some caustic soda solution. The halogen alkyls mix readily with organic solvents, as alcohol, ether, carbon disulphide,

benzene, etc., but *not* with water. The chlorides are lighter, the bromides and iodides heavier than water; the latter have the highest specific gravities. This property decreases in all three classes with the decrease in halogen percentage from the lower up to the higher members of the series; *i.e.* the higher molecular weight compounds have a smaller specific gravity than the lower members. The chlorides have the lowest boiling-points; the corresponding bromides boil about 25°, and the iodides 50° higher.

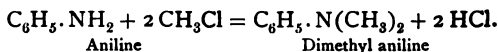
The ease with which the monohalogen alkyls react with other compounds gives them great importance; they are used primarily as a means of introducing alkyl groups into other molecules, *i.e.* by replacing an hydrogen atom with an alkyl group. If it is desired, *e.g.*, to replace in an alcohol, mercaptan, phenol, or acid the hydrogen of the (OH)-, (SH)-, or (COOH)-group by an alkyl radical, *i.e.* to prepare an ether, or ester of these substances, the corresponding sodium compound, or in case of acids, better the silver salt, is treated with the halogen alkyl, *e.g.* :



The alkyl groups may be introduced into the ammonia molecule and into organic amine molecules by means of the halogen alkyls; *e.g.* :

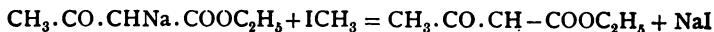


Di- and tri-methyl amine are also formed at the same time.



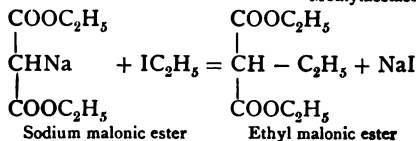
Hydrogen atoms in combination with carbon may also be replaced by alkyl radicals, by means of the halogen alkyls. Since under these conditions another radical is introduced into the molecule, it presents a method of preparing the higher members of a series from the lower, simpler ones. Various examples of this will be taken up later in labor-

atory practice. Here it will be sufficient to refer to several equations showing this kind of reaction.



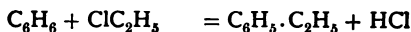
Sodium acetacetic ester

Methylacetacetic ester



Sodium malonic ester

Ethyl malonic ester

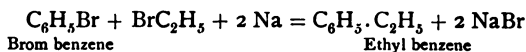


Benzene

Ethyl benzene

(In presence of AlCl_3)

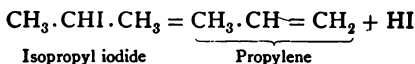
Fittig's Synthesis, to be taken up later, is a case of this kind, by which a halogen atom is replaced by an alkyl group; *e.g.* :



Brom benzene

Ethyl benzene

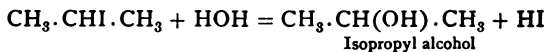
Further, the halogen alkyls serve for the preparation of the unsaturated hydrocarbons of the ethylene series.



Isopropyl iodide

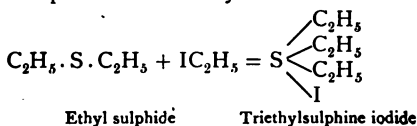
Propylene

In many cases the alcohols may also be prepared from the halogen alkyls; *e.g.* :



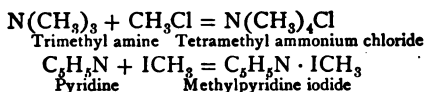
Isopropyl alcohol

This reaction is obviously only of importance when the halogen alkyl is not obtained from the corresponding alcohol, as is the case in the example given. As above mentioned, the isopropyl iodide is most simply obtained from glycerol and hydriodic acid, so that by this reaction the glycerol can be converted into the isopropyl alcohol (compare Reaction 13). Halogen alkyls also unite directly with other compounds, like sulphides and tertiary amines :



Ethyl sulphide

Triethylsulphine iodide



With these examples the list of the many-sided reactions of the halogen alkyls is not exhausted; they are also used for the preparation of the metallic alkyls, *e.g.* zinc alkyls; for the preparation of the phosphines, and for many other compounds. Through the Grignard reaction, with its many-sided applications, the halogen alkyls have recently become extremely important. Finally, attention is called to the characteristic difference between the organic and inorganic halides. While, *e.g.*, potassium chloride, bromide, or iodide in solution act *instantly* with a silver nitrate solution to form a *quantitative* precipitate of silver chloride, bromide, or iodide respectively, silver nitrate in a water solution does not act on most organic halides, so that this reagent does not serve in the usual way to show the presence of a halogen.

EXPERIMENT: Treat a solution of silver nitrate with a few drops of ethyl bromide. Not the least trace of silver bromide is formed.

It has been customary to explain this by saying that the affinity between the halogen atom and carbon is greater than that between the halogen and the metallic atom. According to our later views the difference between the two classes of compounds is explained thus: The metallic halides belong to the class of so-called electrolytes, *i.e.* substances which are dissociated in water solution, *e.g.* the molecule KCl is dissociated into its ions, K^+ and Cl^- . The organic halides are non-electrolytes, *i.e.* the solutions of these contain the undissociated molecules. According to this conception, the potassium chloride reacts with silver nitrate, because no further separation of the potassium from the chlorine (other than that effected by solution) is necessary, while in case of the brom alkyls, the brom-carbon union must first be severed. Only halogen ions react, at once, quantitatively with the silver ions of silver nitrate.

Ethyl iodide is an exception; when shaken with a solution of silver nitrate, it gives an abundant precipitate of silver iodide. Other iodides, especially tertiary iodides, do not react with a water solution of silver nitrate, as readily as ethyl iodide. Methyl and ethyl iodides react quantitatively when an *alcoholic* solution of silver nitrate is used. The reaction is employed, according to Zeisel, for the determination of methoxyl groups. Ethyl bromide reacts with alcoholic silver nitrate on heating; ethyl chloride reacts more slowly.

2. REACTION: PREPARATION OF AN ACID-CHLORIDE FROM THE ACID

EXAMPLE: Acetyl Chloride from Acetic Acid¹

To 100 grammes of glacial acetic acid contained in a fractionating flask connected with a condenser (coil condenser), 80 grammes of phosphorus trichloride are added through a dropping funnel, the flask being cooled by water. The bulb is then im-

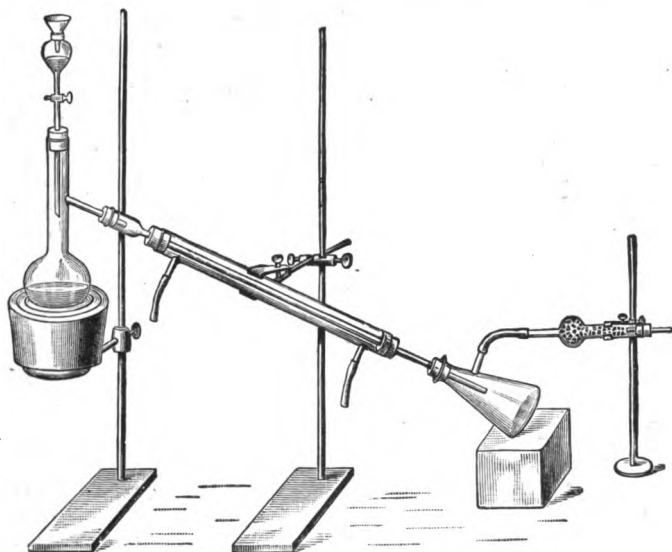


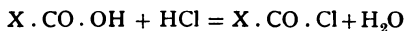
FIG. 64.

mersed in a porcelain dish filled with water at a temperature of 40–50°, and the heating continued until the active evolution of hydrochloric acid gas slackens, and the liquid which was homogeneous before heating has separated into two layers. To separate the acetyl chloride which forms the upper, lighter layer, from the heavier layer of phosphorous acid, the mixture is heated on a

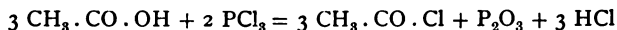
¹ A. 87, 63.

rapidly boiling water-bath until nothing more passes over. Since acetyl chloride is very easily decomposed by moisture, the distillate must not be collected in an open receiver, but the condenser-tube must be tightly connected with a tubulated flask (suction flask), protected from the air by a calcium chloride tube, as represented in Fig. 64. For complete purification, the distillate is distilled in a similar apparatus, except that the dropping funnel is replaced by a thermometer. The apparatus represented in Fig. 17, page 21, may be used for the redistillation. The portion distilling from $50-56^{\circ}$ is collected in a separate vessel. Boiling-point of pure acetyl chloride, 51° . Yield, 80–90 grammes.

In order to replace the hydroxyl of a carboxyl group ($\text{CO} \cdot \text{OH}$) by chlorine, a reaction, similar to the one employed above for the substitution of an alcoholic hydroxyl group by a halogen, may be used. If, *e.g.*, a mixture of an acid and phosphoric anhydride is treated with gaseous hydrochloric acid (heating if necessary), there is formed an acid-chloride, the reaction being analogous to that by which ethyl bromide was prepared.



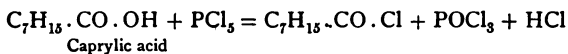
This reaction is without practical importance, since the reactions involved in the methods described under (2), page 136, take place much more smoothly and easily, and, therefore, are exclusively used. In practice, the acid-chlorides are almost always prepared by the action of phosphorus tri- or penta-chloride on the acid directly, or in many cases, on the sodium or potassium salt. Phosphorus oxychloride is employed in rare cases. The selection of the chloride of phosphorus depends upon (1) the ease with which the acid under examination reacts, and (2) upon the boiling-point of the acid-chloride. If, as in the case of acetic acid and its homologues, the trichloride of phosphorus reacts easily with the formation of the acid-chloride, this is selected in preference to the more energetic pentachloride. The reaction probably takes place in accordance with the following equation :



The quantity of acetic acid used is made the basis for the calculation of the yield required by theory.

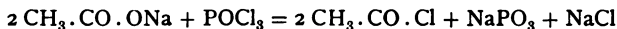
In cases in which the boiling-point of the acid-chloride desired does

not lie far from that of phosphorus oxychloride (110°), thus rendering a fractional distillation for the separation of the products difficult, the trichloride is always used. If an acid does not react too energetically, as is the case with the higher members of the acetic acid series with the pentachloride, this is used. With the aromatic acids, the latter is used exclusively, since the trichloride and oxychloride react with great difficulty:

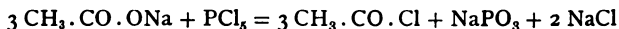


Attention is called to the fact that for one molecule of phosphorus pentachloride, but one molecule of the acid-chloride is obtained.

The phosphorus oxychloride is used generally only when dealing with the salts of carbonic acids, upon which it acts as indicated by the equation:

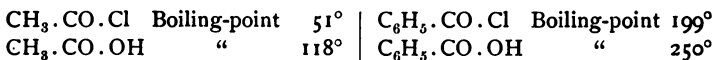


This reaction may be used with advantage in order to utilise more of chlorine of the phosphorus pentachloride than is the case when the latter acts upon the free acids. If the pentachloride is allowed to act on a sodium salt, as above, there is formed, for an instant, phosphorus oxychloride, and while this no longer acts upon the free acid, it can convert two other molecules of the salt into the chloride:



In this way, with the use of one molecule of phosphorus pentachloride, three molecules of the acid-chloride are obtained.

The lower members of the series of acid-chlorides are colourless liquids; the higher, colourless crystalline substances. They boil, generally, at ordinary pressure without decomposition, but the higher members are more conveniently distilled in a vacuum. The boiling-points of the acid-chlorides are lower than those of the acids; the replacement of hydroxyl by chlorine usually causes a lowering of the boiling-point.

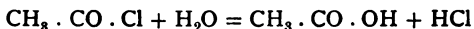


The acid-chlorides possess pungent odours. They fume in the air, since they unite with the moisture and decompose, thus forming the corre-

sponding acid and hydrochloric acid. They are heavier than water, and do not mix with it, but are easily soluble in indifferent organic solvents like ether, carbon disulphide, benzene.

To separate the chlorides from the by-products formed by the phosphorus chloride, one can proceed as in the case of acetyl chloride, by distilling off the volatile chloride from the non-volatile phosphorus acid, either on a water-bath or over a free flame. In order to separate a chloride, in case it distils without decomposition, from the volatile phosphorus oxychloride formed when phosphorus pentachloride is used, a fractional distillation is made. In other cases, the mixture is heated in a vacuum apparatus on an actively boiling water-bath, upon which the phosphorus oxychloride passes over. The non-volatile residue can be used, in many cases, without further purification; it may be obtained perfectly pure by distilling it in a vacuum.

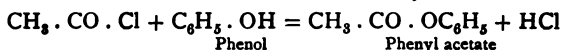
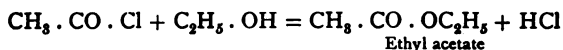
Chemical Reactions. — The acid-chlorides are decomposed by water with the formation of the corresponding acid and hydrochloric acid.



This decomposition takes place often with extreme ease; the chlorine atom is united to the acid radical much less firmly than it is in the case of an alkyl radical. While it is generally necessary, in order to convert a halogen alkyl into an alcohol, to boil it a long time with water, often with the addition of sodium hydroxide, or potassium hydroxide, a carbonate or acetate, the analogous transformation of an acid-chloride takes place with far greater ease. With the lower members, *e.g.* acetyl chloride, the reaction begins almost instantly at the ordinary temperature, and continues with violent energy; but it is necessary to heat the higher members to induce the transformation, *e.g.* benzoyl chloride, which will be prepared later.

EXPERIMENT: To 5 c.c. of water in a test-tube is gradually added $\frac{1}{2}$ c.c. of acetyl chloride. If the water is very cold, the oily drops, sinking to the bottom, do not mix with it, and may be observed for a short time. On shaking the tube, an energetic reaction sets in with evolution of heat, and the chloride passes into solution, which happens immediately if the water is not cold.

The acid-chlorides react with alcohols and phenols to form esters:



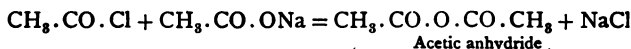
EXPERIMENT: To 1 c.c. of alcohol in a test-tube, cooled with water, add an equal volume of acetyl chloride, drop by drop; this mixture is then treated with an equal volume of water, the tube being cooled as before; the liquid is then carefully made weakly alkaline with sodium hydroxide. If the pleasant-smelling ethyl acetate does not separate out on the water solution in a mobile layer, finely pulverised salt is added until no more will dissolve. This will cause the ethyl acetate to separate out.

The acid-chlorides are also used to determine whether a substance under examination contains an alcoholic or a phenol hydroxyl group or not. If the compound reacts readily with an acid-chloride, it contains either alcoholic or phenol hydroxyl, since compounds containing oxygen in some other form of combination, *e.g.* as in ethers, do not react with acid-chlorides.

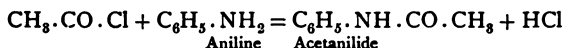
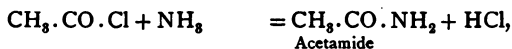
The addition of anhydrous sodium acetate often materially assists the reaction.

Finally, the action of the acid-chlorides upon alcohols and phenols is made use of to separate the latter from solution, or in order to detect them. However, benzoyl chloride is most generally used for this purpose, concerning the importance of which more will be said later.

Acid-chlorides act upon the salts of carbonic acids, forming anhydrides:



The next preparation will deal with this reaction. The acid-chlorides react with ammonia as well as with primary and secondary organic bases with great ease:

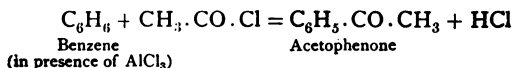


EXPERIMENT: To 1 c.c. of aniline, acetyl chloride is added in drops; an energetic reaction takes place with a hissing sound,

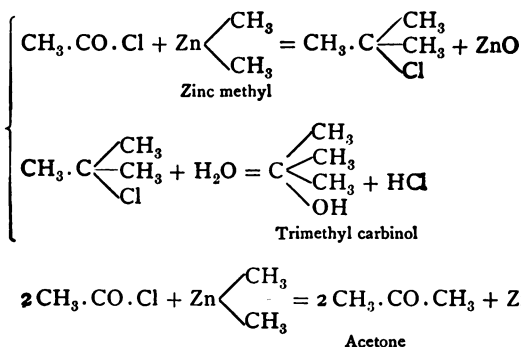
which ceases when about the same volume of the chloride is added. The mixture is cooled with water, and five times its volume of water is added, upon which an abundant precipitate of acetanilide separates out, the quantity of which is increased by rubbing the walls of the test-tube with a glass rod. The precipitate is filtered off, and recrystallised from hot water. Melting-point, 115° .

This reaction is also used to characterise organic bases by converting them into their best crystallised acid derivatives, and in order to detect small quantities, especially of liquid bases, by a melting-point determination. Since the tertiary bases do not react with acid-chlorides, because they no longer contain any ammonia hydrogen, this reaction may be employed to decide whether a given base is, on the one hand, primary or secondary, or, on the other, tertiary.

The fact that hydrogen in combination with carbon can be replaced by an acid radical by the use of an acid-chloride is of special importance. In this connection, the Friedel-Crafts ketone synthesis is particularly mentioned. This will be taken up later, in practice. The reaction is indicated by the following equation :



The acid-chlorides are also of service for the synthesis of tertiary alcohols (Butlerow's Synthesis), as well as for that of ketones. The final reactions will only be indicated here; in regard to the details, reference must be made to larger works.



3. REACTION: PREPARATION OF AN ANHYDRIDE FROM THE ACID-CHLORIDE AND THE SODIUM SALT OF THE ACID

EXAMPLE: Acetic Anhydride from Acetyl Chloride and Sodium Acetate¹

For the preparation of acetic anhydride an apparatus similar to that used in the preparation of acetyl chloride is employed, except that the fractionating flask is replaced by a tubulated retort (Fig. 65, page 148).²

To 70 grammes of finely pulverised, anhydrous sodium acetate (for the preparation of this, see below) contained in the retort, 50 grammes of acetyl chloride are added drop by drop from a separating funnel. As soon as the first half of the chloride is added, the reaction is interrupted for a short time, in order that the pasty mass may be stirred up with a glass rod. The second half is then allowed to run in. If in consequence of a too rapid addition of acetyl chloride some of it should distil over into the receiver undecomposed, this is poured back into the funnel and again allowed to act on the sodium acetate. The separating funnel is then removed, the tubulure closed with a cork, and the anhydride distilled off from the salt residue by means of a *luminous* flame which is kept in constant motion. The distillate is finally purified by distilling in an apparatus similar to the one used in the rectification of the acetyl chloride (see also Fig. 17) with the addition of 3 grammes of finely pulverised anhydrous sodium acetate, which serves to convert the last portions of acetyl chloride into the anhydride. Boiling-point of acetic anhydride, 138°. Yield, about 50 grammes.

Preparation of Anhydrous Sodium Acetate: Crystallised sodium acetate contains three molecules of water of crystallisation. In order to dehydrate it is placed in a shallow iron or nickel dish and heated over a free flame (120 grammes for this experiment). The salt first melts in the water of crystallisation, on further heating steam is copiously evolved, and the mass solidifies as soon as

¹ A. 87, 149.

² An upright coil condenser may be used, in which case an adapter is unnecessary.

the main portion of the water has been driven off, provided the flame is not too large. In order to remove the last portions of the water, the mass is now heated with a large flame, the burner being constantly moved, until the solidified mass melts for the second time. Care must be taken not to overheat; in case this should happen,

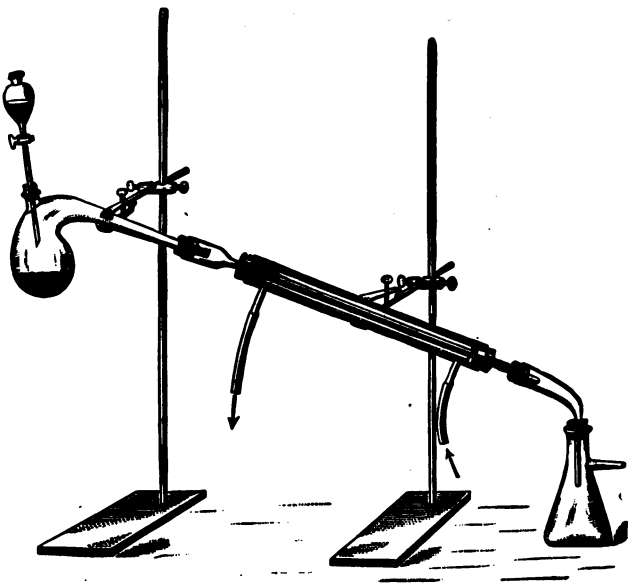
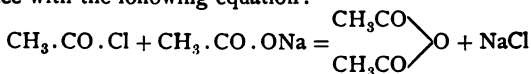


FIG. 65.

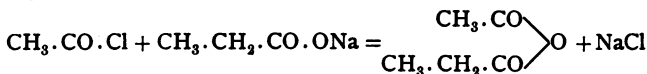
the fact will be recognised by the evolution of combustible gases and the charring of the substance. After cooling, the salt is removed from the dish with a knife. If commercial anhydrous sodium acetate is at hand, it is recommended that this also be melted once, since when it is kept for a long time it always takes up water.

The reaction of acetyl chloride with sodium acetate takes place in accordance with the following equation :

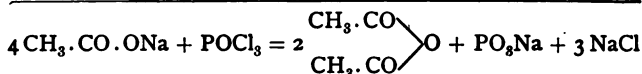
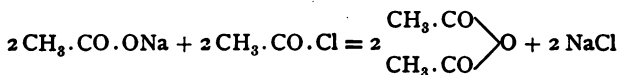
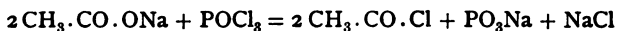


This reaction is capable of general application, and the anhydride of the acid may be made by treating its chloride with the corresponding sodium salt. The so-called mixed anhydrides, containing two different

acid radicals, can also be prepared by this reaction, by using the chloride and sodium salt of two different acids :



Since, as stated, an acid-chloride may be obtained from an alkali salt of the acid and phosphorus oxychloride, it is not necessary for the preparation of an anhydride to first isolate the chloride; it is better to allow the same to act directly on an excess of the salt, so that from the oxychloride and salt an anhydride is directly obtained :



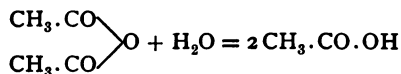
The lower members of the acid-anhydride series are colourless liquids; the higher members, crystallisable solids. They possess a sharp odour, are insoluble in water, but soluble in indifferent organic solvents. Their specific gravities are greater than that of water. The boiling-points are higher than those of the corresponding acids :

Acetic acid, 118° ,

Acetic anhydride, 138° .

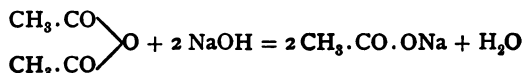
The lower members can be distilled without decomposition at ordinary pressure; but the higher members must be distilled in a vacuum.

The chemical conduct of anhydrides toward water, alcohols, and phenols, as well as bases, is wholly analogous to that of the chlorides; but the anhydrides react with more difficulty than the chlorides. Thus with water, the anhydrides yield the corresponding acids :



EXPERIMENT : 5 c.c. of water are treated with $\frac{1}{2}$ c.c. of acetic anhydride. The latter sinks to the bottom and does not dissolve even on long shaking. It will be recalled that the corresponding chloride reacts instantly with water very energetically. If the mixture be warmed, solution takes place.

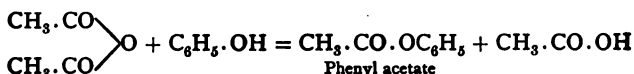
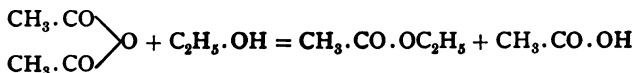
In the presence of alkalis, solution takes place much more readily with the formation of the alkali salts :



EXPERIMENT : Mix 5 c.c. of water with $\frac{1}{2}$ c.c. of acetic anhydride, and add a little caustic soda solution. On shaking, without warming, solution takes place.

Anhydrides of high molecular weight react with water with still greater difficulty, and require a longer heating to convert them into the corresponding acid.

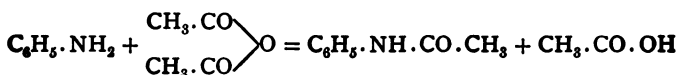
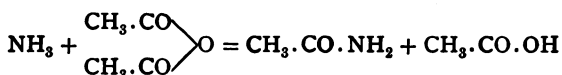
With alcohols and phenols, the anhydrides form acid-esters on heating, while the acid-chlorides react at the ordinary temperature :



It is to be noted that one of the two acid radicals in the anhydride is not available for the purpose of introducing the acetyl group into other compounds, — acetylating, — since it passes over into the acid.

EXPERIMENT : 2 c.c. of alcohol are added to 1 c.c. of acetic anhydride in a test-tube, and heated gently for several minutes. It is then treated with water and carefully made slightly alkaline. The acetic ester can be recognised by its characteristic pleasant odour. If it does not separate from the liquid, it may be treated with common salt, as in the experiment on page 145.

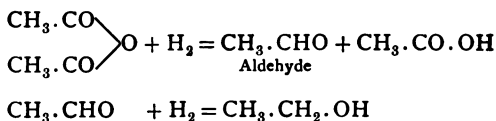
With ammonia and primary or secondary organic bases, the anhydrides react like the chlorides :



EXPERIMENT : Add 1 c.c. of aniline to 1 c.c. of acetic anhydride, heat to incipient ebullition, and then, after cooling, add twice the volume of water. The crystals of acetanilide separate out easily if the walls of the vessel be rubbed with a glass rod ; these are filtered off, and may be recrystallised from a little hot water.

The acid-anhydrides can, therefore, be used, like the chlorides, for the recognition, separation, characterisation, and detection of alcohols, phenols, and amines.

In order to complete the enumeration of the reactions of the acid-anhydrides, it may be mentioned briefly that they yield alcohols, and the intermediate aldehydes when treated with sodium amalgam :



It is, therefore, possible to pass from the anhydride of an acid to its aldehyde or alcohol.

4. REACTION: PREPARATION OF AN ACID-AMIDE FROM THE AMMONIUM SALT OF THE ACID

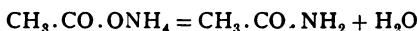
EXAMPLE : Acetamide from Ammonium Acetate¹

To 75 grammes of glacial acetic acid heated to 40–50° in a porcelain dish on a water-bath, finely pulverised ammonium carbonate is added (100 grammes will be necessary) until a test-portion diluted in a watch-glass with water just shows an alkaline reaction. The viscous mass is warmed on an actively boiling water-bath to 80–90°, until a few drops of it diluted with water just show an acid reaction ; it is then poured (without the use of a funnel-tube) directly into two wide bomb-tubes of hard glass, which have been previously warmed in a flame. A single Volhard tube (see page 68) is much more convenient. After the portions of substance adhering to the upper end of the tube have been removed by melting down carefully with a flame, the last traces are removed with filter-paper, the tube sealed and heated

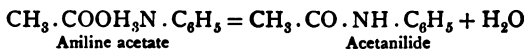
¹ B. 15, 979.

for five hours in a bomb-furnace at $220-230^{\circ}$.¹ The liquid reaction product is fractionated under the hood in a distilling-flask provided with a condenser. There is first obtained a fraction boiling between $100-130^{\circ}$, consisting essentially of acetic acid and water. The temperature then rises rapidly to 180° (an extension tube is substituted for the condenser, see page 22), at which point the acetamide begins to distil. The fraction passing over between $180-230^{\circ}$ is collected in a beaker, cooled by ice water at the end of the distillation, and the walls are rubbed with a sharp-edged glass rod; the crystals separating out are pressed on a drying plate to remove the liquid impurities. By another distillation of the pressed-out crystals, the almost pure acetamide boiling at 223° passes over. Yield, about 40 grammes. The product thus obtained possesses an odour very characteristic of mouse excrement; this is not the odour of pure acetamide, but of an impurity accompanying it. In order to remove the impurity, a portion of the distilled amide is again pressed out on a drying plate, and then crystallised from ether. There are thus obtained colourless, odourless crystals, melting at 82° .

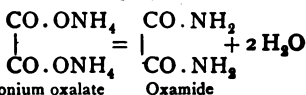
The reaction involved in the preparation of an amide from the ammonium salt of the acid is capable of general application. The latter is subjected to dry distillation, or more conveniently, heated in a sealed tube at $220-230^{\circ}$ for five hours:



In order to purify the amide thus obtained, the reaction-mixture may be fractionated, as in the case of acetamide, or if the amide separates out in a solid condition, it may be purified by filtering off the impurities and crystallising. Substituted acid-amides, and especially easily substituted aromatic amides, *e.g.* acetanilide, can also be readily obtained by this method, by heating a mixture of the acid and amine a long time in an open vessel:

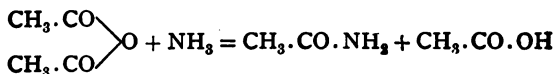
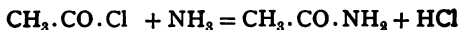


The ammonium salts of di- and poly-basic acids react in a similar way, *e.g.*:

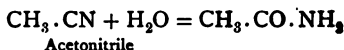
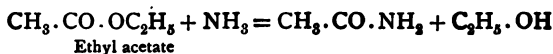


¹ Above this temperature the tube is liable to explode.

Concerning further methods of preparation, it may be stated that acid-chlorides or anhydrides when treated with ammonia, primary or secondary bases form acid-amides very easily:



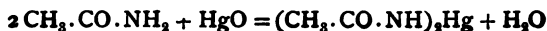
The acid-amides may be furthermore obtained by two methods of general application: (1), by treating an ethereal salt with ammonia, and (2), by treating a nitrile with water:



The acid-amides are, with the exception of the lowest member, formamide, $\text{H}\cdot\text{CO}\cdot\text{NH}_2$ (a liquid), colourless, crystallisable compounds, the lower members being very easily soluble in water, *e.g.*, acetamide; the solubility decreases with the increase of molecular weight, until finally they become insoluble. The boiling-points of the amides are much higher than those of the acids:

Acetic acid, Boiling-point,	118°	Propionic acid, Boiling-point,	141°
Acetamide, ,,	223°	Propionamide, ,,	213°

While the entrance of an alkyl residue into the ammonia molecule does not change the basic character of the compound, as will be discussed more fully under methylamine, the entrance of a negative acid radical enfeebles the basic properties of the ammonia residue, so that the acid-amides possess only a very slight basic character. It is true that a salt corresponding to ammonium chloride — $\text{CH}_3\cdot\text{CO}\cdot\text{NH}_2\cdot\text{HCl}$ — can be prepared from acetamide by the action of hydrochloric acid; but this shows a strong acid reaction, is unstable, and decomposes easily into its components. If it is desired to assign to the acid-amides a definite character, they must be regarded as acids rather than bases. One of the amido-hydrogen atoms possesses acid properties in that it may be replaced by metals. The mercury salts of the acid-amides may be prepared with especial ease, by boiling the solution of the amide with mercuric oxide:

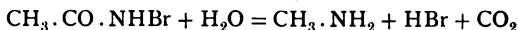


EXPERIMENT : Some acetamide is dissolved in water, treated with a little yellow mercuric oxide, and warmed. The latter goes into solution, and the salt of the formula given above is formed.

The amido-hydrogen atoms can also be replaced by the negative chlorine and bromine atoms, as well as by the positive metallic atoms. These substitution compounds are obtained by treating the amide with chlorine or bromine, in the presence of an alkali :

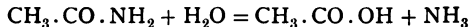


The monohalogen substituted amides are of especial interest, since, on being warmed with alkalis, they yield primary alkylamines :



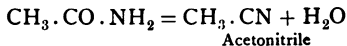
This important reaction will be taken up later, under the preparation of methyl amine from acetamide.

In the acid-amides, the acid radical is not firmly united with the ammonia residue ; this is shown by the fact that they are saponified, *i.e.* decomposed into the acid and ammonia, on boiling with water, more rapidly by warming with alkalis :

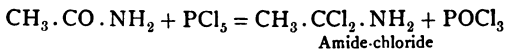


EXPERIMENT : Heat some acetamide in a test-tube with caustic soda solution. A strong ammoniacal odour is given off, while the solution contains sodium acetate.

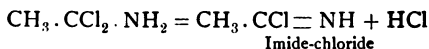
If an acid-amide is treated with a dehydrating agent, *e.g.*, phosphorus pentoxide, it is converted into a nitrile :



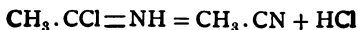
The same result is obtained by treating it with phosphorus pentachloride ; but in this case the intermediate products, the amide-chlorides or imide-chlorides are formed :



The very unstable amide-chloride then passes over, with the loss of one molecule of hydrochloric acid, into the more stable imide-chloride :



And this finally into the nitrile,

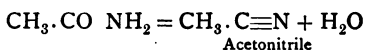


5. REACTION: PREPARATION OF AN ACID-NITRILE FROM AN ACID-AMIDE

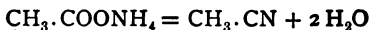
EXAMPLE: Acetonitrile from Acetamide¹

To 15 grammes of phosphoric anhydride, contained in a small, dry flask, 10 grammes of dry acetamide are added. After the two substances are shaken well together, the flask is connected with a short condenser, and then heated *carefully*, with a not too large *luminous* flame kept in *constant motion*. The reaction proceeds with much foaming. After the mixture has been heated a few minutes, the acetonitrile is then distilled over with a *large* luminous flame, kept in *constant motion*, into the receiver (test-tube). The distillate is treated with half its volume of water, and then solid potash is added until it is no longer dissolved by the lower layer of liquid. The upper layer is removed with a capillary pipette and distilled, a small amount of phosphoric anhydride being placed in the fractionating flask for the complete dehydration of the nitrile. Boiling-point, 82°. Yield, about 5 grammes.

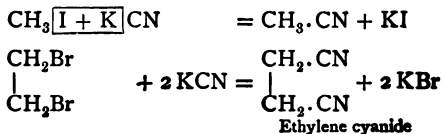
If an acid-amide is heated with a dehydrating agent (phosphorus pentoxide, pentasulphide, or pentachloride), it loses water, and passes over into the nitrile, *e.g.*:



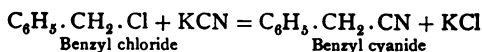
Since, as has just been done, the acid-amide may be made by dehydrating the ammonium salt of an acid, thus, in a single operation the nitrile may be obtained directly from the ammonium salt, if it is treated with a powerful dehydrating agent, *e.g.* ammonium acetate heated with phosphoric anhydride:



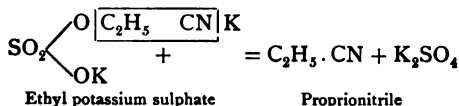
The acid-nitriles may also be obtained by heating alkyl iodides (or bromides, chlorides) with alcoholic potassium cyanide:



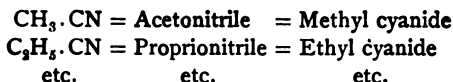
¹ A. 64, 332.



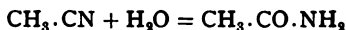
or by the dry distillation of alkyl alkali sulphates with potassium cyanide:



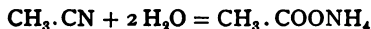
These two reactions differ from those above in that the introduction of a new atom of carbon is brought about. The nitriles thus appear to be cyanides of the alkyls, and, therefore, may be equally well designated as cyanides, *e.g.*:



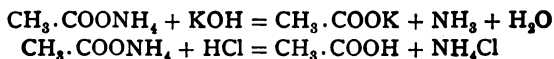
The lower members of the nitrile series are colourless liquids, the higher members, crystallisable solids; the solubility in water decreases with the increase in molecular weights. If they are heated with water up to 180° under pressure, they take up one molecule of water and are converted into the acid-amides:



On heating with acids or alkalis, they take up two molecules of water, and pass over into the ammonium salt as an intermediate product:

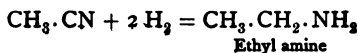


which immediately reacts with the alkali or acid, in accordance with the following equations:



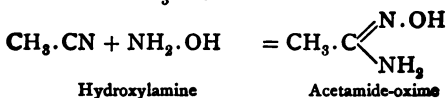
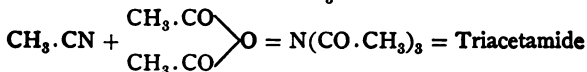
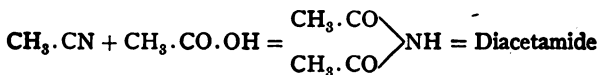
This process is called "saponification."

If nascent hydrogen (*e.g.* from zinc and sulphuric acid) be allowed to act on nitriles, primary amines are formed (Mendius' reaction):¹



¹ A. 121, 129.

Further, but of less importance, general reactions may be indicated by the following equations:



6. REACTION: PREPARATION OF AN ACID-ESTER FROM THE ACID AND ALCOHOL

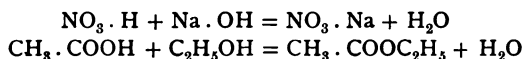
EXAMPLE: Acetic Ester from Acetic Acid and Ethyl Alcohol¹

A $\frac{1}{2}$ -litre flask, containing a mixture of 50 c.c. of alcohol and 50 c.c. of concentrated sulphuric acid, is closed by a two-hole cork; through one hole passes a dropping funnel, through the other a glass delivery tube connected with a long condenser or coil condenser. The mixture is heated in an oil-bath to 140° (thermometer in oil); when this temperature is reached, a mixture of 400 c.c. of alcohol and 400 c.c. of glacial acetic acid is gradually added through the funnel, at the same rate at which the ethyl acetate (acetic ester), formed in the reaction, distils over. In order to remove the acetic acid carried over, the distillate is treated in an *open* vessel with a dilute solution of sodium carbonate until the upper layer will not redden blue litmus paper. The layers are now separated with a dropping funnel; the upper layer is filtered through a dry folded filter, and shaken up with a solution of 100

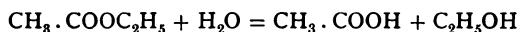
¹ Bl. 33, 350.

grammes of calcium chloride in 100 grammes of water, in order to remove the alcohol.¹ The two layers are again separated with the funnel, the upper one dried with granular calcium chloride and then distilled on the water-bath (see page 16). Boiling point, 78°. Yield, about 80-90 % of the theory.

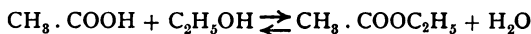
The formation of an ester from acid and alcohol is analogous to the formation of a salt from an acid and a metallic hydroxide:



The two reactions take place quantitatively, but not in a similar manner. A strong acid reacts almost quantitatively with an equivalent weight of a strong base, and the product of this neutralisation is a salt. Upon this depend the processes of acidimetry and alkalimetry. But equimolecular quantities of an acid and an alcohol do not yield the theoretical quantity of the ester. A maximum quantity of ester is formed, but this falls short of the quantity required by theory, and it is impossible, even when the reacting substances are kept in contact, to convert the unchanged acid and alcohol into ester beyond a certain limit. If, for example, equimolecular quantities of acetic acid and alcohol are allowed to interact, only two-thirds of these enter into the reaction, the maximum yield of ester being 66.7 % of the theory. It is impossible to cause a union between the remaining one-third of acetic acid and alcohol, even when the reaction is continued for a long time. The difference in the quantitative course of the reaction in the formation of an ester is due to the "reversibility of the reaction"; *i.e.* the reaction products on the right-hand side of the equation (ester and water) will interact in such a manner as to reverse the reaction:



In reactions of this order the two sides of the equation are united, not by an equation sign, but, as proposed by van't Hoff, by two arrows pointing in opposite directions:

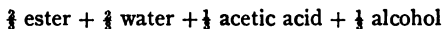


The reaction in the neutralisation of a strong acid with a strong base is, on the other hand, unlike esterification, since it is "an irreversible or a complete reaction"; the water that is liberated does not react with the salt to reverse the reaction and regenerate the acid and the base. In reality this difference does not exist. All reactions are

¹ Calcium chloride forms a compound with alcohol. (Compare page 54.)

reversible. But when a reaction product is extremely insoluble, or when it is a gas, or when for other reasons the final products have little tendency to react and bring about the reverse change (and this is the case in the above example of the neutralisation of a strong acid with a strong base), then one of the two opposing reactions is said to be complete "within measurable limits," and it is called an "irreversible reaction in the ordinary sense," although not in the strictest sense. — While in irreversible reactions chemical equations enable us to calculate the amount of the products from given quantities of reacting substances, in reversible reactions strictly quantitative stoichiometric methods do not give the desired information. But by the aid of the highly important **Law of Mass Action** (Guldberg and Waage, 1867), it is possible to determine to what extent a reversible reaction may be complete.

As has already been mentioned, when equimolecular quantities of acetic acid and ethyl alcohol are allowed to react for some time, only two-thirds of these substances will be transformed into acetic ester and water. A "state of equilibrium" will finally be established, and the reaction mixture will have the following constant composition :



The same equilibrium is established, if, instead of a mixture of acid and alcohol, a similar mixture of ester and water is taken. In this case the ester will be partially saponified into acetic acid and alcohol, but the reaction will proceed only until $\frac{1}{3}$ of the ester is saponified. An equilibrium will once more be established as above, so that $\frac{2}{3}$ of ester and water, and $\frac{1}{3}$ of acetic acid and alcohol will be obtained.

It must not be assumed that in an equilibrium of this kind the molecules of the four substances remain unaltered (static equilibrium). On the contrary, while acetic acid and alcohol are forming ester and water, the molecules of ester and water are simultaneously reacting to bring about the reverse change (dynamic equilibrium). In spite of this continuous reaction an equilibrium will exist, *i.e.* the composition of the system will remain unaltered, when the velocities of the two opposing reactions are the same, *i.e.* when in unit time an equal number of ester molecules is formed and saponified.

The formation of ethyl acetate from acetic acid and alcohol may be expressed by the following equation of mass action :

$$\frac{C_s \cdot C_a}{C_e \cdot C_w} = K$$

where C_s , C_a , C_e , C_w show the "concentration" of acetic acid, alcohol, ester, and water respectively, and K is a constant. By concentration,

or "active mass" (Guldberg and Waage) is not meant the weight of each substance in the total volume or in unit volume, but the relative number of molecules, *i.e.* the weight of each substance divided by its molecular weight (the number of gramme-molecules or moles). The equation shows that when the four substances are in equilibrium with one another, the product of the concentrations of acetic acid and alcohol divided by the product of the concentrations of ester and water will be equal to a constant. How can this constant be calculated? We simply ascertain by analytical methods the weights of the four substances that are in equilibrium with one another in a concrete example, calculate the concentrations in accordance with deductions mentioned above, and introduce these values into the equation given. This may be readily done in our example, since three of the substances (alcohol, ester, and water) are neutral, and the quantity of the fourth, the acetic acid, may be easily ascertained by titration. If we take, *e.g.*, equimolecular quantities of acetic acid and alcohol, namely, 60 grammes of acetic acid and 46 grammes of alcohol (and this will contain an equal number of molecules), we can determine the amount of acid in one c.c. of this mixture, at the first minute, by titration. If we now allow the two substances to react for some time, and titrate one c.c. of the mixture at stated intervals, the titre of the acid will be found to diminish gradually, until finally it remains constant, due to equilibrium. If now we compare the maximum first titre with the final minimum titre, we shall find that the latter is exactly one-third of the former; *i.e.* at equilibrium only one-third of the original molecules of acetic acid remain uncombined, the other two-thirds having been changed into the ester. Since one molecule of acid yields one molecule of ester, the number of ester molecules is exactly two-thirds of acetic acid molecules used originally for the experiment. Since further, with the formation of every ester molecule a water molecule is also formed, the number of water molecules is also exactly two-thirds of acetic acid molecules taken originally. And finally, since with every molecule of acid one molecule of alcohol reacts to form the ester, two-thirds of the alcohol molecules are used up, and only one-third remain unchanged at equilibrium. We have thus determined the amount of all four substances at equilibrium by a mere titration. Consequently, we have the following values in our equation:

$$C_S = \frac{1}{3}; C_A = \frac{1}{3}; C_E = \frac{2}{3}; C_W = \frac{2}{3}.$$

If we carry these values into the above equation, we obtain:

$$K = \frac{\frac{1}{3} \cdot \frac{1}{3}}{\frac{2}{3} \cdot \frac{2}{3}} = \frac{1}{4}.$$

Thus when we have accurately determined the value of K in a single experiment, we shall be in a position to calculate the quantitative yield of acetic ester at equilibrium for all proportions of acetic acid and alcohol. Suppose we use, *e.g.*, one gramme-molecule of acetic acid with two gramme-molecules of alcohol, and let x be the number of gramme-molecules of ester at equilibrium; the gramme-molecular quantity of water will also be represented by x . The gramme-molecular quantity of unchanged acetic acid will then be $(1 - x)$, and that of unchanged alcohol will be $(2 - x)$. Making these substitutions in our equation, we obtain :

$$\frac{(1 - x) \cdot (2 - x)}{x \cdot x} = \frac{1}{4}$$

$$x = 2 \pm 2\sqrt{\frac{1}{3}}$$

Since the quantity of acetic acid used is only one gramme-molecule, x cannot be greater than one, and we are thus concerned only with the negative sign. Thus x is equal to $2 - 2\sqrt{\frac{1}{3}} = 0.85$, *i.e.* 0.85 gramme-molecules of ester are obtained at equilibrium, *i.e.* 85% of the acetic acid is transformed into ester. If, therefore, instead of using equimolecular quantities of acetic acid and alcohol, we double the theoretical quantity of the latter, 85% of the acetic acid will be transformed into ester instead of 66.7%.

The following problems may be solved in this connection :

How much ester will be formed when one gramme-molecule of acetic acid is treated with three gramme-molecules of alcohol? How much ester will be formed when 30 grammes of acetic acid and 50 grammes of alcohol are used? What proportions of acetic acid and alcohol must be used in order to transform 75% of the former into ester?

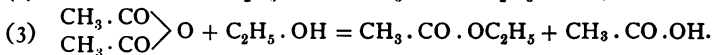
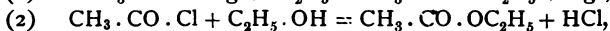
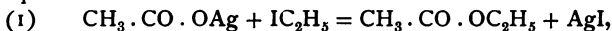
As the above example shows, the yield of ester from the same quantity of acetic acid is greater, the larger the amount of alcohol used. This also follows directly from the equation of mass action given above. As we have seen, K must have the constant value of $\frac{1}{4}$ for all proportions of reacting substances. Thus when C_A , the concentration of the alcohol, is increased, the remaining three quantities will also be changed in such a manner that the quotient will have a constant value. This can only happen when the quantities in both denominators increase, *i.e.* when the concentrations of the ester and the water formed at the same time become greater. The equalisation of the quotient, however, takes place not only through the denominator, but also through the numerator; because when more ester is formed, more acetic acid will also be used up. This is shown by the decrease in the concentration of acetic acid. Thus a large excess of alcohol is taken when it is desired

to convert an acid, as completely as possible, into an ester. If, on the other hand, it is desired to convert an alcohol into an ester quantitatively, a large excess of acid is used.

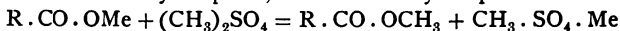
The practical application of these principles in preparation work has already been pointed out under ethyl bromide.

In the preparation of ethyl acetate described above, sulphuric acid is also used in addition to acetic acid and alcohol. This reacts in two ways. It is a well-known fact that concentrated sulphuric acid combines with water chemically. Consequently, the water liberated during the formation of ester is either completely, or in part, removed, and the reverse reaction from right to left, *i.e.* the saponification of the freshly formed ester, is either rendered impossible or difficult. Thus the yield of ester is much larger in the presence of sulphuric acid. But sulphuric acid also acts as a catalytic agent in the reaction, *i.e.* it accelerates the formation of ester, as well as its saponification, and indeed in the same ratio, so that the state of equilibrium is not changed. Other acids may also be employed as catalytic agents in this sense. Thus esters may be formed by simply conducting hydrochloric acid gas into a mixture of acid and alcohol, or by heating the acid with alcoholic hydrochloric acid (containing a small percentage of HCl). (Compare B. 28, 3252.) The speed of the catalytic action is proportional to the strength of the acid used as a catalytic agent. The stronger the dissociation of the acid, *i.e.* the greater the concentration of hydrogen ions in the reaction mixture, the more rapid the formation of the ester. By this method, and others, is the strength of many acids determined.

In many cases, where the salts of organic acids are more readily obtained than the free acids, they may be used for the preparation of the esters by heating them directly with alcohol and sulphuric acid. Other methods for the preparation of acid-esters have been referred to, and, in part, carried out practically on the small scale in the foregoing preparations, so that at this point it is only necessary to recall the equations:



(4) Acid-esters may also be readily obtained by treating the alkali salts of acids with alkyl sulphate,¹ at the ordinary temperature:



(Compare B. 37, 3658.) It must be remembered that methyl sulphate is poisonous.

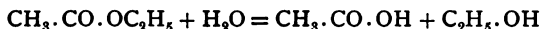
¹ The hydrogen atom in a hydroxyl group in phenols, as well as a hydrogen atom in combination with nitrogen, may readily be exchanged for a methyl group by the use of dimethyl sulphate. See A. 327, 104.

Concerning the purification of the acid-esters, it may be mentioned, that the crude reaction-product is shaken with a sodium carbonate solution until the ester no longer shows an acid reaction. The alcohol may be removed from esters difficultly soluble in water by repeatedly washing with water; in case an ester is moderately soluble in water, as ethyl acetate, it is better to use a solution of calcium chloride.

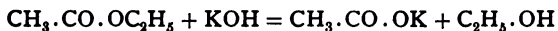
The lower members of the series of acid-esters are colourless liquids with pleasant, fruit-like odours; the higher members, as well as those of the aromatic acids, are crystallisable compounds. The boiling-points of esters containing alkyl residues of small molecular weights (CH_3 , C_2H_5 , C_3H_7) are lower than those of the corresponding acids; the entrance of more complex alkyl residues raises the boiling-points:

$\text{CH}_3 \cdot \text{CO} \cdot \text{OCH}_3$	Boiling-point,	57°
$\text{CH}_3 \cdot \text{CO} \cdot \text{OC}_2\text{H}_5$	“ “	78°
$\text{CH}_3 \cdot \text{CO} \cdot \text{OH}$	“ “	118°
$\text{CH}_3 \cdot \text{CO} \cdot \text{OC}_6\text{H}_{13}$	“ “	169°
Hexyl acetate		

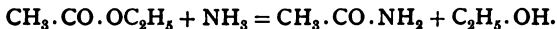
As already mentioned, the esters are saponified by heating with water:



The saponification is effected more readily by heating with alkalis:



Other methods of saponification will be further discussed when a practical example is taken up. The action of ammonia upon acid-esters, forming acid-amides, has already been referred to under acetamide:



7. REACTION: SUBSTITUTION OF HYDROGEN BY CHLORINE

EXAMPLE: Monochloroacetic Acid from Acetic Acid and Chlorine¹

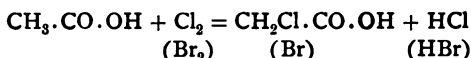
A current of dry chlorine is passed into a mixture of 150 grammes of glacial acetic acid and 12 grammes of red phosphorus, contained in a flask provided with a delivery tube and an inverted

¹ R. 23, 222; A. 102, 1.

condenser; the flask is heated on a rapidly boiling water-bath, and must be placed in such a position as to receive as much light as possible. The best result is obtained by performing the reaction in the direct sunlight, since the success of the chlorination depends essentially on the action of the sun's rays. The reaction is ended when a small test-portion cooled with ice-water solidifies on rubbing the walls of the vessel (test-tube) with a glass rod. In summer the chlorination may require a single day, while during the cloudy days of winter, two days may be necessary. For the separation of the monochloroacetic acid the reaction-product is fractionated from a distilling flask connected with a long air condenser. The fraction passing over from 150–200° is collected in a separate beaker; this is cooled in ice-water and the walls rubbed with a glass rod. The portion solidifying, consisting of pure monochloroacetic acid is rapidly filtered with suction, the loose crystals being pressed together with a spatula or mortar-pestle. The suction must not be continued too long, because the monochloroacetic acid gradually becomes liquid in warm air. The filtrate is again distilled, and the portion passing over between 170–200° is collected in a separate vessel. This is treated as before (cooling and filtering), and there is obtained a second portion of monochloroacetic acid; this is united with the main quantity, which is again distilled. The product thus obtained is perfectly pure. Boiling-point, 186°. Yield varying, 80–125 grammes.

Since monochloroacetic acid, especially when warm, attacks the skin with great violence, care must be taken in handling it.

Chlorine or bromine substitution products of aliphatic carbonic acids can be obtained by the direct action of the halogen on the acids:

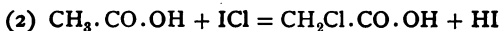


If the reaction is allowed to continue for a long time, other substitution products can also be obtained. But the action of chlorine or bromine on acids is very sluggish. It may be substantially facilitated by certain conditions. If, *e.g.*, the operation is conducted in direct sunlight, the reaction proceeds much more rapidly than in a dark place. The reaction is assisted more effectively by adding a so-called "carrier." Iodine

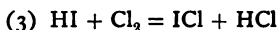
may be used as such for the introduction of chlorine or bromine. When added in small quantities to the substance to be substituted, it causes the substitution to take place more rapidly and completely. The continuous action of this carrier depends upon the following facts: In the first phase of the reaction, chlorine iodide is formed:



This acts, then, as a chlorinating agent in the second phase, according to the following reaction:

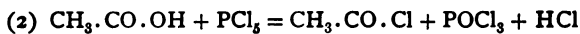
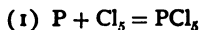


The chlorine acts upon the hydriodic acid as follows:

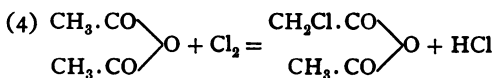


The molecule of chlorine iodide is thus formed anew (equation 1) and can chlorinate another molecule of acetic acid, and so on. The action of the iodine in the last case depends upon the fact that the molecule of chlorine iodide (ICl) is more easily decomposed into its atoms than the molecule of chlorine (Cl_2). The disadvantage necessarily incident to the use of iodine as a carrier is, that the reaction-product is easily contaminated with iodine derivatives,—in small quantities, it is true.

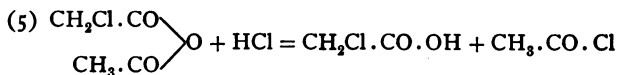
In an entirely different way the chlorination is facilitated by the addition of red phosphorus. In this case, phosphorus pentachloride is first formed from the phosphorus and chlorine; this, acting on the acetic acid, generates acetyl chloride, and this latter, with an excess of the acid, forms the anhydride. Direct experiments have shown that acid-chlorides, as well as anhydrides, are substituted by chlorine with much greater ease than the corresponding acids; in this fact the action of red phosphorus finds its explanation. Since a small amount of phosphorus is sufficient for the chlorination of a large amount of acetic acid, the question as to how this is continuously effected remains to be answered. In accordance with the above statements, the following reactions take place:



If the chlorine now acts on the anhydride, monochloroacetic anhydride is formed :



But this reacts directly with the hydrochloric acid, in accordance with this equation :



There is thus obtained, besides the molecule of chloroacetic acid, the molecule of acetyl chloride, first formed in reaction 2, which is utilised repeatedly by its regeneration in accordance with reactions 3, 4, and 5.

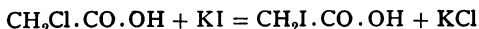
As a substitute for red phosphorus, sulphur is also recommended for the chlorination of aliphatic acids. This acts in a wholly similar manner, since it first forms sulphur chloride, which, reacting on the acid, like phosphorus chloride, converts it into an acid-chloride. The other phases of the reaction are similar to those given above.

The bromination of aliphatic carbonic acids, which is not only of great importance in preparation work, but also as a means for determining constitution, is also conducted with the addition of red phosphorus (Hell-Volhard-Zelinsky Method).¹

The halogen atoms always enter the α -position to the carboxyl group. Thus, *e.g.*, when propionic and butyric acids are brominated, they yield :



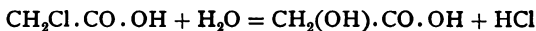
If no α -hydrogen atom is present, *e.g.*, in trimethyl-acetic acid $(\text{CH}_3)_3\text{C} \cdot \text{CO} \cdot \text{OH}$, bromination will not take place. The ability of an acid to form a bromine substitution product can, therefore, be used as a test for the presence of an α -hydrogen atom. Iodine cannot be introduced directly into aliphatic acids like chlorine and bromine. To obtain iodine substitution products, it is necessary to treat the corresponding chlorine or bromine compound with potassium iodide :



The halogen derivatives of the fatty acids are in part liquids, in part

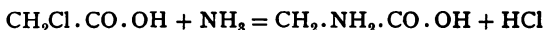
¹ B. 14, 891; 21, 1726; A. 242, 141; B. 21 1904; B. 20, 2026; B. 24, 2216.

solids. In their reactions they resemble the acids, on the one hand, since they form salts, chlorides, anhydrides, esters, etc.; on the other hand, the halogen alkyls. They are of great value in the preparation of oxy- and amido-acids, of unsaturated acids, for the synthesis of polybasic acids, etc. Below are given a few equations capable of general application :



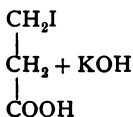
Oxyacetic acid =

Glycolic acid

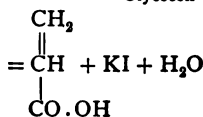


Amidoacetic acid =

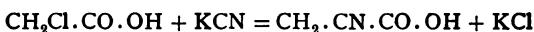
Glycocoll



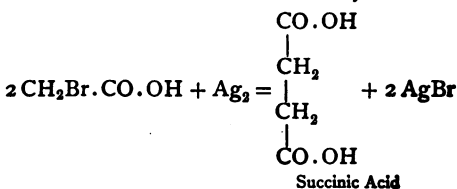
(From glyceric acid + PI_3)



Acrylic acid



Cyanacetic acid



Succinic Acid

8. REACTION: OXIDATION OF A PRIMARY ALCOHOL TO AN ALDEHYDE

EXAMPLE : Acetaldehyde from Ethyl Alcohol¹

A 1½-litre flask containing 110 grammes of concentrated sulphuric acid and 200 grammes of water is closed by a two-hole cork; through one hole passes a dropping funnel, through the other a glass delivery tube connected with a long condenser. To the lower end of the condenser is attached an adapter bent downwards, the narrower portion of which passes through a cork in

¹ A. 14, 133; J. 1853, 329.

the neck of a thick-walled suction flask of about $\frac{1}{2}$ -litre capacity. (See Fig. 65, page 148.) By using an upright coil condenser connected directly with the suction flask, an adapter is unnecessary. The suction flask is placed in a water-bath filled with a freezing mixture of ice and salt. The larger flask is heated over a wire gauze until the water just begins to boil; a solution of 200 grammes of sodium dichromate in 200 grammes of water which has been treated with 100 grammes of alcohol is then added in a small stream through the dropping funnel, the lower end of which is about 3 cm. above the surface of the liquid in the flask. During

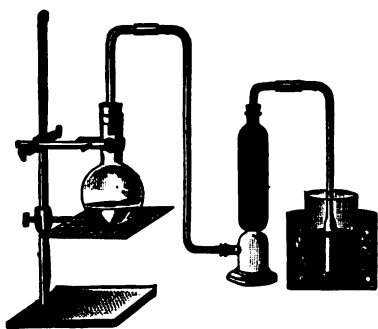


FIG. 66.

the addition of the mixture, it will be unnecessary to heat the flask, since the heat produced by the reaction is sufficient to cause ebullition. The aldehyde thus formed distils into the receiver, besides some alcohol, water, and acetal. If uncondensed vapours of the aldehyde escape from the receiver, the mixture is admitted to the flask more slowly. On

the other hand, if boiling is not caused by the flowing in of the mixture, the reaction is assisted by heating with a small flame. After all of the mixture has been added, the flask is heated for a short time by a flame, until boiling begins.

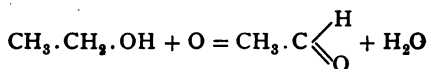
Since the aldehyde cannot be obtained easily from the reaction-products by fractional distillation, it is first converted into aldehyde-ammonia, which, on proper treatment, readily yields the pure aldehyde.

The apparatus for this purpose is arranged as follows: A small flask to contain the aldehyde, placed on a wire gauze, is connected with a moderately large reflux condenser. Into the upper end of the condenser is placed a cork bearing a $\sqcap$ -shaped glass tube, which is connected with two wash-bottles, each containing 50 c.c. of dried ether. After the condenser has been filled with water at

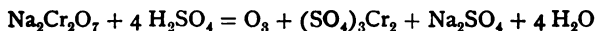
30° (the lower side-tube of the condenser is closed with rubber tubing and a pinch-cock), the crude aldehyde is heated for 5-10 minutes to gentle boiling, and the aldehyde that is not condensed passes over, and is absorbed by the ether. Should the ether begin to ascend in the connecting tube, the flame must be somewhat increased immediately. To obtain aldehyde-ammonia, a current of dry ammonia (see page 382) is conducted, with the aid of a wide adapter or funnel (Fig. 66), into the ethereal solution contained in a beaker surrounded by a freezing mixture of ice and salt, until the liquid smells strongly of it. After an hour, the aldehyde-ammonia which has separated out is scraped from the sides of the vessel and adapter with a spatula or knife, filtered with suction, washed with a little ether, and then allowed to dry on filter-paper in a desiccator. Yield, about 30 grammes.

In order to obtain pure aldehyde, 10 grammes of aldehyde-ammonia are dissolved in 10 grammes of water, treated with a cooled mixture of 15 grammes of concentrated sulphuric acid and 20 grammes of water, and heated on the water-bath. Since aldehyde has a low boiling-point (21°), the receiver is connected with the condenser by a cork, and well cooled with ice and salt.

Aldehydes can be obtained by the use of the general reaction, which in many cases serves as a method of preparation, of extracting two hydrogen atoms from a primary alcohol by oxidation.



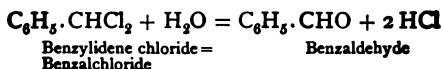
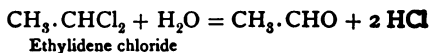
The name of the class of compounds is derived from this action: Aldehyde = Al(cohol)dehyd (rogenatus). As an oxidising agent in the above case, chromic acid is the most suitable in the form of potassium, or sodium dichromate in the presence of sulphuric acid:



The rather difficultly soluble potassium dichromate (1 part dissolves in 10 parts water) was formerly generally used, but at present the more soluble and cheaper sodium salt (1 part dissolves in 3 parts of water) is employed wherever it is possible. But in the preparation of the simplest aldehyde (formaldehyde) from an alcohol a different oxidising agent is

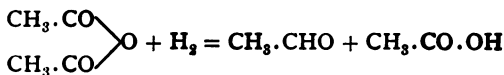
used, viz. the oxygen of the air. On passing a mixture of the vapour of methyl alcohol and air over a heated copper spiral, formaldehyde is produced.

While by the first reaction one proceeds from substances which in comparison with the aldehydes are oxidation products of a lower order, the aldehydes may also be obtained by a second method involving the use of compounds of the same substitution series, viz. the dihalogen derivatives of the hydrocarbons containing the group CHCl_2 or CHBr_2 . If these are boiled with water, or, better, water containing sodium carbonate, potash, lead oxide, or calcium carbonate, etc., the two halogen atoms are replaced by one oxygen atom :

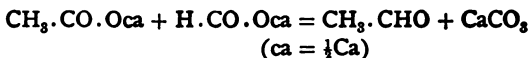


This method is used on the large scale for the manufacture of the commercially important benzaldehyde. It will be referred to under benzaldehyde.

Finally, aldehydes can be prepared from their oxidation products, the carbonic acids, by two methods, one of which has been already mentioned under acetic anhydride. If sodium amalgam is allowed to act on acid-anhydrides, an aldehyde is first formed :



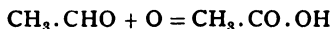
But this reaction is of little practical value for the preparation of aldehydes. The second method, which is the real preparation method, consists in the dry distillation of a mixture of the calcium or barium salt of the acid with calcium or barium formate :



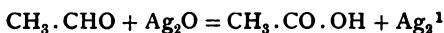
The lower members of the aldehyde series are colourless liquids, soluble in water, possessing pungent odours. The intermediate members are also liquids, but insoluble in water ; the higher members are solid, crystallisable substances. The boiling-points of the aldehydes are lower than those of the corresponding alcohols.

{	CH ₃ .CHO	Boiling-point, 21°
{	CH ₃ .CH ₂ .OH	" " 78°
{	CH ₃ .CH ₂ .CHO	" " 50°
{	CH ₃ .CH ₂ .CH ₂ .OH	" " 97°

Aldehydes are oxidised to acids by free as well as combined oxygen (compare benzaldehyde) :



Upon this action depends the fact that aldehydes cause metals to separate from certain salts, *e.g.* silver nitrate :



EXPERIMENT : Treat a few cubic centimetres of a diluted silver nitrate solution with a few drops of ammonium hydroxide and 5 drops of aldehyde. Silver will be deposited in the form of a brilliant mirror on the walls of the vessel ; the deposition is especially beautiful if the vessel has previously been treated with a solution of caustic soda to remove any fatty matter. The reaction frequently takes place at the ordinary temperature, but in many cases only on gentle warming. The reaction is used for the detection of aldehydes.

Another reaction which can also be employed for the recognition of aldehydes, depends upon the fact that they give a red colour to fuchsine-sulphurous acid. (Caro's reaction.)

EXPERIMENT : Fuchsine-sulphurous acid is prepared by dissolving fuchsine in water ; a sufficient quantity of the latter is taken to prevent the solution from being too intense in colour. Sulphur dioxide is conducted into this until a complete decolouration takes place. To a few cubic centimetres of this solution add several drops of aldehyde. On shaking, a violet-red colour will be produced.

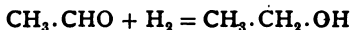
Finally aldehydes may also be detected by a method depending upon the fact that when treated with diazobenzene sulphonic acid and sodium amalgam, they give a violet colour.

EXPERIMENT : To as much diazobenzene sulphonic acid as can be held on the point of a knife, add 5 c.c. of water, a few drops

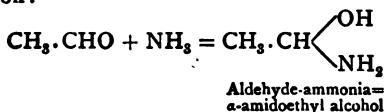
¹ B. 15, 1635 and 1828.

of caustic soda, and then a few drops of aldehyde, and finally a piece of solid sodium amalgam as large as a pea. After some time the red-violet colour appears.

That aldehydes upon reduction pass over to primary alcohols has been mentioned under acetic anhydride:

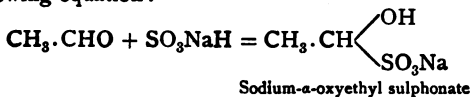


It is wholly characteristic of the aldehydes that they unite directly (1) with ammonia, (2) sodium hydrogen sulphite, and (3) hydrocyanic acid. The union with ammonia takes place in accordance with the following equation:



This reaction is not so common as the second and third. Thus, *e.g.*, formic aldehyde and most of the aromatic aldehydes behave differently toward ammonia. Whenever this reaction does take place it can also be used with advantage for the purification of the aldehyde, as in case of acetaldehyde; by allowing the well-crystallised double compound to separate out, on treating it with dilute sulphuric acid, the free aldehyde is obtained.

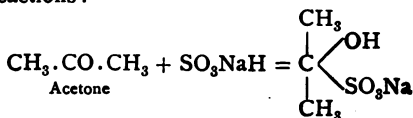
The union with sodium hydrogen sulphite takes place in accordance with the following equation:



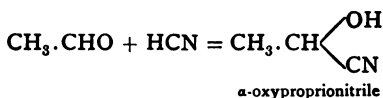
This reaction may also be used for the purification of aldehydes, since when a concentrated solution of the sulphite is employed, the double compound generally separates out in a crystallised condition. The free aldehydes can be obtained from the sulphite compounds by heating with dilute acids or alkali carbonates. (Compare benzaldehyde.)

EXPERIMENT: Treat 5 c.c. of a cooled concentrated solution of sodium hydrogen sulphite with 1 c.c. of aldehyde and shake the mixture. The double compound separates out in a crystallised condition.

As distinguished from the union of aldehyde with ammonia, this reaction is wholly general, and is frequently of great value in dealing with the aldehydes of the aromatic series. It should be noticed in this connection, that the ketones, which are closely related to the aldehydes, show similar reactions:



The addition of hydrocyanic acid to ketones as well as to aldehydes is also general:

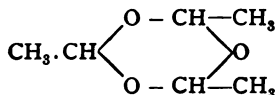


This reaction is of especial interest, since the addition of a new carbon atom is brought about. Concerning the value of this reaction for the synthesis of α -oxyacids, see Mandelic Nitrile, page 307.

Aldehydes possess further a marked tendency to combine with themselves (polymerise).

EXPERIMENT: Treat 1 c.c. of aldehyde with one drop of concentrated sulphuric acid. The aldehyde boils, and polymerisation (condensation) takes place.

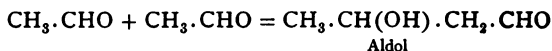
The compound thus obtained is called paraldehyde; it boils much higher (124°) than ordinary aldehyde; the determination of its vapour density shows that one molecule is composed of three molecules of ordinary aldehyde. Paraldehyde does not show the characteristic aldehyde reactions; on distillation with dilute sulphuric acid it is converted back into the ordinary variety. For this reason it is believed that no new union of carbon atoms takes place in the molecule, but that three molecules are united by means of the oxygen atoms:



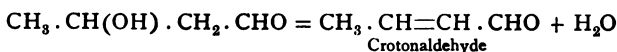
If aldehyde is *cooled* and treated with sulphuric acid, or if at the ordinary temperature gaseous hydrochloric acid, sulphur dioxide, or other compounds are passed into it, a solid polymerisation product,

metaldehyde, is formed; this can also be converted back into the ordinary variety.

The aldehydes undergo a wholly different kind of polymerisation under certain conditions, concerning which reference must be made to the chemical literature and treatises. For example, two molecules of acetaldehyde can unite with the formation of a compound in which a new carbon union is present.



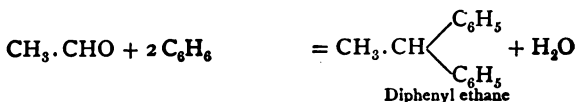
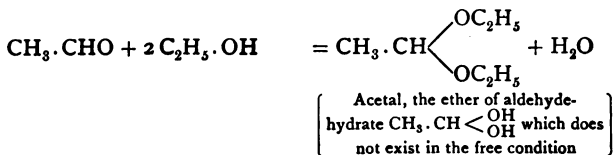
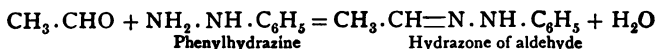
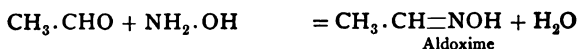
This compound, as distinguished from paraldehyde and metaldehyde, is a true aldehyde, in that it cannot be converted back to acetaldehyde. Aldol loses water easily, and is converted into an unsaturated aldehyde:



In connection with these condensations, it may be pointed out that many aldehydes, when heated with alkalis, polymerise to resinous products of high molecular weight (aldehyde resins).

EXPERIMENT: Treat a few cubic centimetres of caustic potash solution with several drops of aldehyde, and warm. A yellow colouration takes place with the separation of a resinous mass.

In order, finally, to represent the great activity of the aldehydes, the following equations are given:

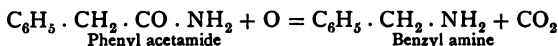


9. REACTION: PREPARATION OF A PRIMARY AMINE FROM AN ACID-AMIDE OF THE NEXT HIGHER SERIES**EXAMPLE: Methyl Amine from Acetamide¹**

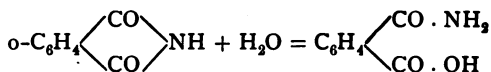
To a mixture of 25 grammes of acetamide, which has been previously well pressed out on a porous plate, and 70 grammes (23 c.c.) of bromine contained in a $\frac{1}{2}$ -litre flask, add a solution of 40 grammes of caustic potash in 350 c.c. of water (the flask is well cooled with water), until the brownish red colour formed at first is changed to a bright yellow, for which the greater portion of the potash solution will be required. This reaction-mixture is then, in the course of a few minutes, allowed to flow from a dropping funnel in a continuous stream into a litre flask containing a solution of 80 grammes of caustic potash in 150 c.c. of water heated to 70–75°. In case the temperature rises higher than 75°, the flask must be cooled by immersion for a short time in cold water. The liquid is maintained at this temperature until it becomes colourless, which usually requires a quarter to half hour. The methyl amine is then distilled off with steam, and collected in a receiver containing a mixture of 60 grammes of concentrated hydrochloric acid and 40 grammes of water. In order that the methyl amine may be completely absorbed by the acid, the end of the condenser is connected with an adapter which dips 1 cm. below the surface of the liquid in the receiver. If a coil condenser is employed, the end of it dips directly into the acid. As soon as the liquid in the condenser no longer shows an alkaline reaction, the distillation is discontinued. The methyl amine hydrochloride is partially evaporated in a porcelain dish over a free flame, then to dryness on the water-bath, and is finally heated for a short time in an air-bath at 100° to dusty dryness. In order to separate the methyl amine salt from the ammonium chloride mixed with it, the finely pulverised substance is crystallised from absolute alcohol, and the crystals separating out dried in a desiccator. Yield, varying.

¹ B. 15, 762; B. 17, 1406 and 1920.

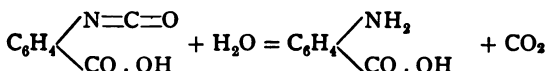
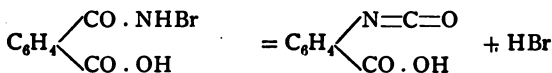
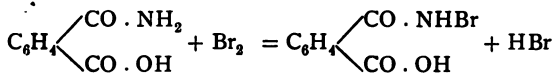
may be obtained from the easily accessible nitro-compounds; and since, on the other hand, if the above reaction is employed, bromine substitution products are easily formed. But in those cases in which the nitro-compound corresponding to the amine is not known, or can be prepared only with difficulty, the reaction is also of importance in the aromatic series. Two cases of this kind may be mentioned in this place. If phenyl acetamide is treated in accordance with Hofmann's reaction, there is formed in the usual way benzyl amine:



Further, the reaction is of practical value in the preparation of o-amido-benzoic acid, used in the manufacture of artificial indigo. If, as above, bromine and caustic potash are allowed to act on phthalimide (technically bleaching powder is used), there is first formed, by the addition of water, an acid-amide:



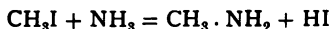
which in accordance with the following reactions gives the amido-acid.



Since the nitro-acid corresponding to the o-amido benzoic acid is difficult to obtain, and phthalimide is easily prepared (naphthalene is changed to phthalic anhydride by oxidation, and when this is treated with ammonia, phthalimide is formed with the separation of water), the Hofmann reaction in this case gives a very convenient method of preparation for the amido-acid.

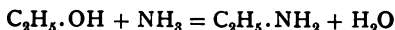
Primary aliphatic amines can also be prepared according to the following equations:

(1) By the action of alcoholic ammonia on halogen alkyls:

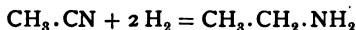


In this case, secondary and tertiary bases, or the corresponding ammonium compounds, are also formed.

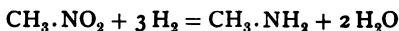
(2) From alcohols and zinc chloride-ammonia :



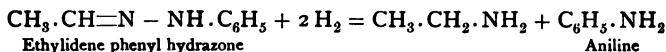
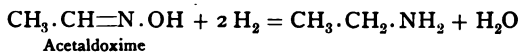
(3) By the reduction of nitriles (Mendius' reaction) :



(4) By the reduction of nitro-compounds :



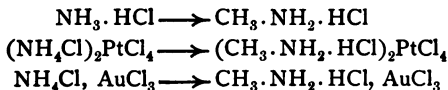
(5) By the reduction of oximes and hydrazones :



The lowest members of the amines in the free condition are gaseous compounds soluble in water, possessing odours suggestive of ammonia : they differ from ammonia in being inflammable.

EXPERIMENT : Treat some solid methyl amine hydrochloride in a small test-tube with a concentrated solution of caustic potash, or caustic soda, and warm gently. A gas, smelling like ammonia, is evolved, which burns with a pale flame.

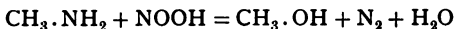
The higher members are liquids or insoluble solids. Since they are derivatives of ammonia, they possess basic properties, and, like ammonia, unite with acids to form salts, the composition of which is analogous to that of the ammonium compounds :



The hydrochlorides of organic bases are distinguished from ammonium chloride by their solubility in absolute alcohol. Use was made of this property above.

The numerous reactions of the primary amines need not be mentioned here, since, under the aromatic amines, frequent reference will be made to them. At this place, one difference between the aromatic and aliphatic amines will be pointed out. If nitrous acid is allowed to

act on an aliphatic primary amine, an alcohol is formed with evolution of nitrogen:



while, under these conditions, an aromatic amine is converted into a diazo-compound (see Diazo-compounds).

10. REACTION: SYNTHESSES OF KETONE ACID-ESTERS OR POLYKETONES WITH SODIUM OR SODIUM ALCOHOLATE

EXAMPLE: Acetacetic Ester from Acetic Ester and Sodium¹

For the successful preparation of acetacetic ester, the character of the acetic ester used is of great importance, since, if it is completely free from alcohol, it will be attacked very slowly by sodium, even on heating; if, on the other hand, it contains too much alcohol, the sodium acts easily, but the yield of the product is varying and usually small. According to the experiments of the author, the following method of procedure gives a good yield and is one that does not fail.

Purification of Acetic Ester: The acetic ester prepared according to Reaction 6, even after it has been freed from acetic acid and alcohol by shaking with sodium carbonate and calcium chloride respectively, dried over calcium chloride, and finally rectified, is not suitable for this preparation, since it reacts too violently with sodium. But if it is allowed to stand, after distilling, for some hours, over night at least, in a well-closed flask, over about $\frac{1}{2}$ its volume of granulated calcium chloride, and is then filtered, it may be used for the successful preparation of acetacetic ester.

If commercial acetic ester is to be used, it must be shaken with a sodium carbonate solution, as described on page 157, treated with calcium chloride solution, etc.; in short, it is treated as the crude product obtained in the preparation of acetic ester. Obviously, it is also necessary to allow it to stand over night in contact with calcium chloride, after the distillation.

The yield of acetacetic ester may be further increased if the acetic ester, after being filtered from the calcium chloride, is again distilled, care being taken to prevent the absorption of moisture. All parts of the apparatus must be perfectly dried, and the end of

¹ A. 186, 214.

the condenser tube must be connected to the receiver (suction-flask) by a good cork.

Preparation of Acetacetic Ester: 25 grammes of sodium from which the outside layers have been removed are cut with the aid of a sodium knife (Fig. 89) into pieces as thin as possible, and placed in a dry litre-flask. After this is connected with a long reflux condenser, inclined at an oblique angle, 250 grammes of dried acetic ester is poured into the top of the condenser, by a funnel which must not be attached to the condenser, but is held in the hand, so that the air may escape. If the acetic ester is added properly, no violent ebullition will occur, but at first a gradual, gentle boiling. After 10 minutes, the flask is placed on a previously heated water-bath, the temperature of which is so regulated that the acetic ester boils but gently; the reaction-mixture is heated until all the sodium is dissolved, which will require from 3-4 hours. To the warm liquid is added a mixture of 80 grammes of glacial acetic acid and 80 grammes of water, until it just shows an acid reaction. If a thick, porridge-like mass should separate out, this is again dissolved by vigorous shaking, or carefully breaking up the small lumps with a glass rod. To the liquid is then added an equal volume of a cold saturated solution of sodium chloride, and the lower aqueous layer is separated from the upper one, consisting of acetic ester and acetacetic ester, by allowing it to run off from a dropping funnel. Should a precipitate settle out on the addition of the salt solution, it is dissolved by adding some water. To separate the acetacetic ester from the main portion of the excess of the acetic ester used, the mixture is distilled from a flask, heated by a free flame over a wire gauze, or, more conveniently, without the wire gauze, with a luminous flame. As soon as the thermometer indicates 95° , the heating is discontinued, and the residue is subjected to vacuum-distillation, as described on page 25. In place of the usual condenser, the outside jacket of a Liebig condenser is pushed over the long side-tube of the distillation flask, and water is allowed to circulate through it. The heating is done in an air-bath. After small quantities of acetic ester, water, and acetic acid have

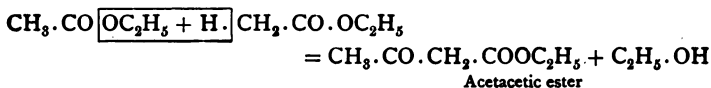
passed over, the temperature becomes constant, and the main portion of the acetacetic ester distils over within one degree. The following table gives the boiling-points at various pressures. A reference to this will show at what approximate points the collection of the preparation should begin :

Boiling-point	71° at 12.5 mm. pressure.			
"	74°	"	14	"
"	79°	"	18	"
"	88°	"	29	"
"	94°	"	45	"
"	97°	"	59	"
"	100°	"	80	"

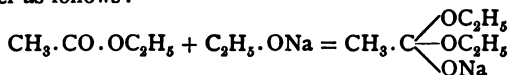
The yield of acetacetic ester amounts to 55–60 grammes.

In the preparation of this substance, it must be borne in mind that the experiment must be completed in one day. The operation should be begun in the morning, the acetic ester heated with sodium at midday, and the experiment completed in the afternoon. If the reaction is discontinued at any point, and the unfinished preparation allowed to stand over night, the yield is materially diminished.

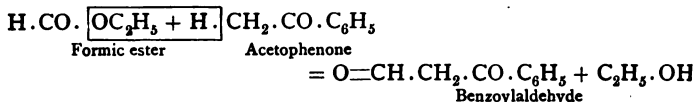
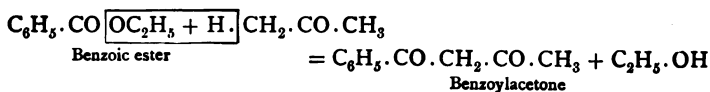
The formation of acetacetic ester from acetic ester, discovered by Geuther in 1863, takes place in accordance with the following equation :



But the mechanism of the reaction is much more complicated than here indicated. According to the views of Claisen, the sodium first acts on the alcohol, which, as above mentioned, must be present in small quantities, forming sodium alcoholate, and this unites with the acetic ester as follows :

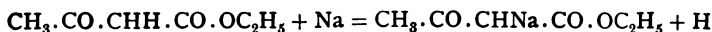


Reaction then takes place between this addition product and a second molecule of acetic ester, with the elimination of two molecules of alcohol, and the formation of the sodium salt of the acetacetic ester :



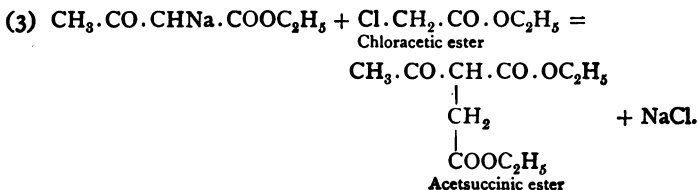
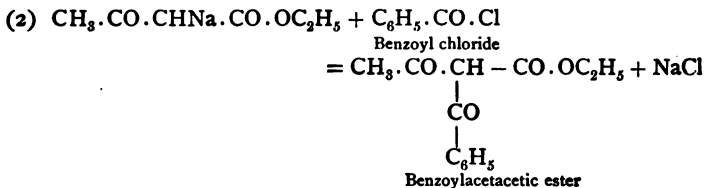
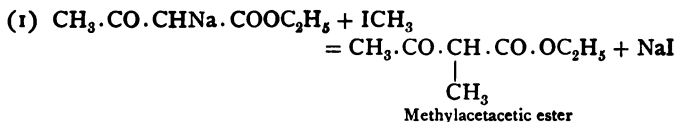
These few examples are sufficient to show the many-sided applications of the above reaction.

The most remarkable characteristic of acetacetic ester is that a portion of its hydrogen may be substituted by metals. If sodium is allowed to act on it, the sodium salt is formed with the elimination of hydrogen:

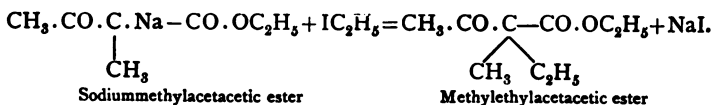


The same salt is also formed by shaking the ester with a solution of sodium hydroxide. The reason for this phenomenon is to be sought in the acidifying influence of the two neighbouring carbonyl (CO) groups.

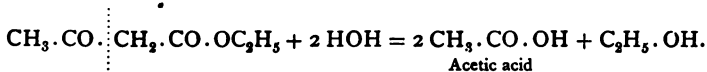
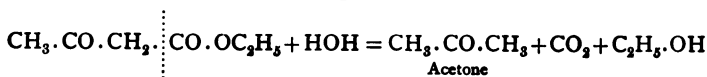
The synthetical importance of acetacetic ester depends on the fact that the most various organic halogen substitution products react with sodium acetacetic ester, the halogen uniting with the sodium, with the condensation of the two remaining residues. Thus a large number of compounds may be built up from their constituents. A few typical examples may elucidate these statements:



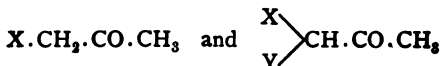
In the compounds thus obtained the second methylene hydrogen atom is also replaceable by sodium, and this salt is likewise capable of entering into similar reactions, by which the number of derivatives is largely increased, *e.g.* :



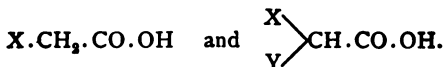
From all these compounds simpler ones may be obtained on saponification. The acetacetic ester breaks up in one of two ways, depending upon the conditions of the saponification :



The first kind of decomposition is called "ketone decomposition," the second, "acid decomposition." Since, as shown above, either one or both of the methylene hydrogen atoms in acetacetic ester can be replaced by different radicals, X or Y, these substances yield either mono- or di- substituted acetones :



as well as mono- and di- substituted acetic acids,



The variety of the acetacetic ester syntheses is still further increased by the fact that *two molecules* of the ester, by reaction with aldehydes or alkylene bromides, may be united with one another by the most various bivalent radicals.

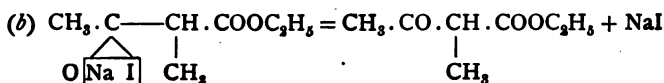
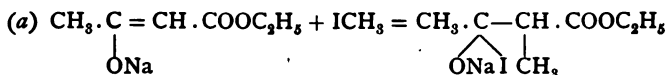
According to recent researches of K. H. Meyer¹ and Knorr² especially, the optical behaviour of ordinary acetacetic ester indicates that it consists of a mixture of the ketoform mainly, with a small quantity of the enolform (β -oxycrotonic ester), these two being in a state of

¹ A. 280, 212.

² B. 44, 1138.

equilibrium. When the mixture is cooled with solid carbon dioxide and ether to -78° , the pure ketoform crystallises out. When the dry sodium salt of acetacetic ester is decomposed at a low temperature with dry hydrogen chloride, pure enolform is obtained which does not solidify at -78° . It *readily* reacts with ferric chloride and is coloured red. The ketoform does not behave in this manner. It is true that the latter may also give the red colouration on standing for a short or a longer period; this, however, is due to the enolform produced in the reaction mixture. The two forms are converted one into the other at ordinary temperatures, the final product being the "equilibrium acetic ester" mentioned above.

The sodium salt of acetacetic ester is also derived, according to its optical behaviour, from the enolform. The transpositions mentioned above are brought about by the action of methyl iodide. A double compound is first formed, with a subsequent decomposition into sodium halide.



11. REACTION: SYNTHESIS OF THE HOMOLOGUES OF ACETIC ACID BY MEANS OF MALONIC ESTER

EXAMPLE: Butyric Acid from Acetic Acid

(a) *Preparation of Malonic Ester*¹

Dissolve 50 grammes of chloracetic acid in 100 grammes of water; warm gently (about 50°), and neutralise with solid, dry, potassium carbonate, for which 30–40 grammes will be required. The solution is then heated gradually, with thorough stirring, while 40 grammes of pure, finely pulverised potassium cyanide are added. (Use a sand-bath or asbestos plate, *under the hood*.) The forma-

¹ A. 204, 121; *Journ. Amer. Chem. Soc.* 18, 1105.

tion of cyanacetic acid takes place, with vigorous ebullition. When the reaction is complete, the mixture is evaporated as quickly as possible on the sand-bath until a thermometer placed in the viscous brownish salt indicates 135° . Since the substance "bumps" and spatters during the evaporation, it is constantly stirred with the thermometer, the hand being protected by a glove or cloth. It is allowed to cool, the stirring being continued during the cooling, otherwise the product bakes into a hard, scarcely pulverisable mass. It is then quickly powdered as finely as possible, placed in a $\frac{1}{2}$ -litre flask provided with a reflux condenser, and treated with 20 c.c. of absolute alcohol. A cooled mixture of 80 c.c. of absolute alcohol and 80 c.c. of concentrated sulphuric acid is then added gradually, with good shaking, through the condenser tube. The pasty mass is now heated, with frequent shaking, two hours in a water-bath (hood); it is then well cooled, and treated with 150 c.c. of water (shaking). After the undissolved salt has been filtered off with suction, it is washed on the filter several times with ether. The filtrate, consisting of the water solution and the ether washings, is then carefully extracted with a sufficiently large quantity of ether. The ethereal extract is shaken up in a separating funnel with a concentrated solution of sodium carbonate until it no longer shows an acid reaction. (*Owing to the copious evolution of gas the funnel is not closed at first.*) It is then dried over fused Glauber's salt, and after the evaporation of the ether, distilled. Boiling-point, 195° . Yield, 45–50 grammes.

The compound may also be advantageously dried, without the use of Glauber's salt, by evaporating off the ether, and heating the residue about a quarter hour in a vacuum on a water-bath. (Compare page 55.)

(b) *The Introduction of an Ethyl Group*

Dissolve 2.3 grammes of sodium in 25 grammes of absolute alcohol in a small flask connected with a reflux condenser; treat the cooled solution gradually with 16 grammes malonic ester, — the transparent crystals of sodium ethylate separating out at first pass over into a voluminous pasty mass of sodium malonic ester, —

and then, with shaking, add through the condenser 20 grammes of ethyl iodide in small portions. The mixture is then heated on the water-bath until the liquid no longer shows an alkaline reaction, for which one or two hours may be necessary. The alcohol is then distilled off on an actively boiling water-bath, a thread being placed in the flask to facilitate the boiling; the residue is taken up with water and extracted with ether, the ether evaporated, and the residue distilled. Boiling-point, 206–208°. Yield, about 15 grammes.

(c) *Saponification of Ethyl Malonic Ester*

For the saponification of the ester, a concentrated solution of caustic potash is prepared; for every gramme of the ester, a solution of 1.25 grammes potassium hydroxide in 1 gramme of water is used. The cooled solution is placed in a flask provided with a reflux condenser; through this the ester is gradually added; an emulsion is first formed, which soon solidifies to a white solid mass, probably potassium ethylmalonic ester. On heating the mixture on a water-bath a sudden energetic boiling-up sets in, especially if the flask be shaken. The heat generated in the reaction causes the alcohol, liberated in the saponification, to boil. After about an hour's heating on the water-bath, the oily layer disappears, showing that the saponification is complete.

The free ethyl malonic acid may be obtained by following either of the methods given below:

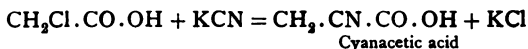
(1) The solution is diluted with $1\frac{1}{2}$ times the volume of water previously used to dissolve the caustic potash. To this is added, gradually, with cooling, a quantity of concentrated hydrochloric acid equivalent to the total amount of caustic potash used (the strength of the acid is determined by a hydrometer). The ethylmalonic acid liberated is taken up with not too little ether, the ethereal solution dried over anhydrous Glauber's salt, the ether evaporated, and the residue heated, with stirring, on a water-bath, in a large watch crystal or dish, until it begins to solidify. After cooling, it is pressed out on a drying plate and crystallised from benzene. Melting-point, 111.5°. Yield, about 7 grammes.

(2) The solution is diluted with the same volume of water previously used to dissolve the caustic potash. To this is added carefully and with cooling concentrated hydrochloric acid, until an acid reaction may just be detected. The ethyl malonic acid is precipitated out in the form of its difficultly soluble calcium salt, by the addition of a cold solution of calcium chloride, *as concentrated as possible*. This is filtered off, well pressed on a porous plate, and the ethyl malonic acid liberated by treating carefully with concentrated hydrochloric acid is obtained pure as in (1).

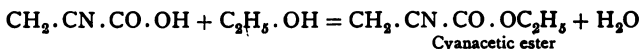
(d) *Elimination of Carbon Dioxide from Ethyl Malonic Acid*

The ethyl malonic acid is placed in a small fractionating flask provided with a *long* condensing tube supported in an oil-bath at an oblique angle, so that its outlet tube is inclined upward. The mouth is closed by a cork bearing a thermometer. The acid is heated at 180° , until carbon dioxide is no longer evolved, which will require about a half-hour. The residue is distilled from the same flask in the usual way; the butyric acid passes over between $162-163^{\circ}$. Yield, about 80-90% of the theory.

(a) In the first phase of the reaction which gives ethylmalonic ester, the potassium cyanide acts on the chloracetic acid, or on its potassium salt, with the formation of cyanacetic acid:

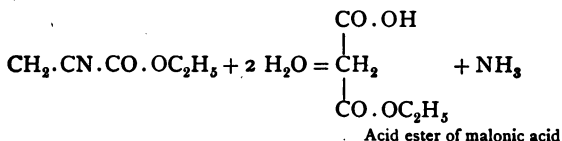


As already mentioned in the preparation of acetonitrile, a halogen united with aliphatic residues may generally be replaced by the cyanogen group, on heating with potassium or silver cyanide. If alcohol and sulphuric acid or ethylsulphuric acid are now allowed to act on the cyanacetic acid, three reactions take place. At first there is an esterification in accordance with the equation:

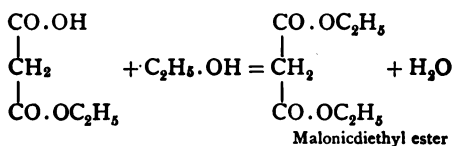


Under the discussion of acetic ester, it has already been brought forward that, in general, acid esters can be obtained by treating a mixture of the alcohol and acid with sulphuric acid. In the second place,

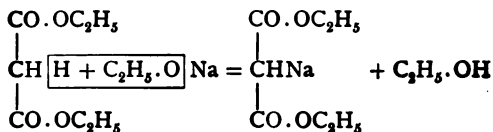
the sulphuric acid has a saponifying action on the cyanacetic ester, *i.e.*, the cyanogen group is converted into carboxyl (COOH).



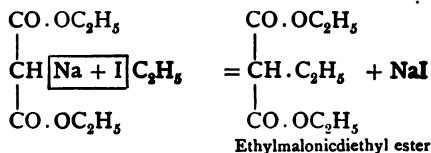
The carboxyl group thus formed is then acted on in the same way as the carboxyl group of cyanacetic acid above, with the formation of an ester:



(b) The ester of malonic acid, like acetacetic ester, possesses the property in virtue of which one of the two methylene hydrogen atoms can be replaced by sodium, in consequence of the acid properties imparted by the two neighbouring carbonyl (CO) groups. When the sodium compound is treated with organic halides, like alkyl halides, halogen derivatives of acid-esters, acid-chlorides, etc., the sodium is replaced by alkyl residues, acid residues, etc., just as in the case of the closely related acetacetic ester. In the above-mentioned examples, the sodium salt of the malonic ester is first formed from sodium alcoholate and the ester:



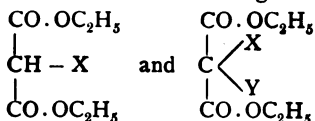
Ethyl iodide reacts on this as follows:



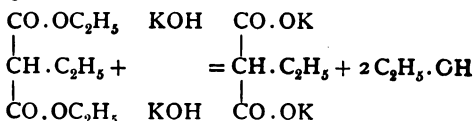
As in acetacetic ester, the second hydrogen of the malonic ester can also be replaced by sodium; consequently the malonic ester is capable

of reacting a second time with organic halides, so that disubstituted malonic esters can also be prepared.

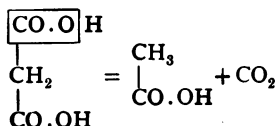
(c) The compounds thus obtained of the general formulæ:



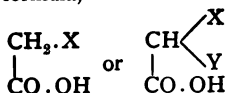
are distinguished from the corresponding derivatives of acetacetic ester in that on saponification they do not decompose, but yield the free substituted malonic acids. Thus, the ethylmalonicdiethyl ester reacts with caustic potash as follows:



(d) From the substituted malonic acid thus obtained, derivatives of acetic acid may be prepared by heating it to a high temperature. It is a general law that one carbon atom cannot hold two carboxyl groups in combination at high temperatures, since carbon dioxide will be eliminated from one. By this means, a dicarbonic acid is converted into a monocarbonic acid, *e.g.*:



From the mono- or di- substituted malonic acid a substituted acetic acid is obtained of the formula,



Thus from ethylmalonic acid, there is formed ethylacetic acid = butyric acid. If, instead of ethyl iodide, methyl or propyl iodide is used, propionic acid or valerianic acid respectively is obtained. If two methyl groups are introduced into malonic ester, then, on decomposition, a dimethylacetic or isobutyric acid will be formed.

As shown above, similar acids may be prepared from the acetacetic ester. Since the decomposition of the acetacetic ester derivatives may take place in two different ways (acid- and ketone-decomposition);

and since these decompositions frequently take place side by side, while the malonic acid derivatives decompose in only one way, so in most cases it is more advantageous to use the malonic ester for the synthesis of the homologous fatty acids.

12. REACTION: PREPARATION OF A HYDROCARBON OF THE ETHYLENE SERIES BY THE ELIMINATION OF WATER FROM AN ALCOHOL. COMBINATION OF THE HYDROCARBON WITH BROMINE

EXAMPLE: Ethylene from Ethyl Alcohol. Ethylene Bromide¹

(a) From Ethyl Alcohol and Sulphuric Acid

A mixture of 25 grammes of alcohol, 150 grammes of concentrated sulphuric acid, and 30 grammes of coarse-grained sea-sand (freed from fine particles) is heated, *not too strongly*, in a litre round flask on a sand-bath or a wire gauze covered with thin asbestos paper. As soon as an active evolution of ethylene takes place, add, through a dropping funnel, a mixture of 1 part alcohol and 2 parts concentrated sulphuric acid (made by pouring 150 grammes of alcohol into 300 grammes of sulphuric acid, with constant stirring), slowly, so that a regular stream of gas is evolved. If the mixture in the flask foams badly with a separation of carbon, it has been too strongly heated, and it is advisable to empty the flask and begin the operation anew with a smaller flame. In order to free the ethylene from alcohol, ether, and sulphur dioxide, it is passed through a wash-bottle containing sulphuric acid, and a second one, provided with three tubulures, the central one supplied with a safety-tube, containing a dilute solution of caustic soda. It then enters two wash-bottles, each containing 25 c.c. of bromine, covered with a layer of water 1 cm. high. Since the combination of ethylene with bromine causes the evolution of heat, the bromine bottles are placed in thick-walled vessels filled with cold water. In order to get rid of the bromine vapours which escape from the last bottle, it is connected with the hood or with a flask containing a solution of caustic soda; to prevent the

¹ A. 168, 64; A. 192, 244.

caustic soda from being drawn back into the bromine bottle, the delivery tube must not dip under the surface of the caustic soda, and the stopper must be provided with canals cut in the sides. As soon as the bromine is decolourised, which requires 4-5 hours under normal conditions, the operation is discontinued, care being taken to disconnect all of the vessels immediately;

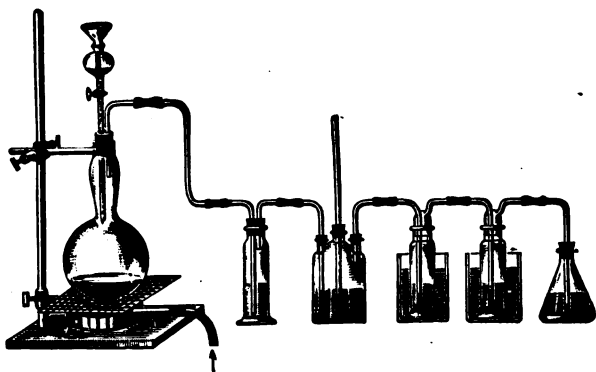


FIG. 67.

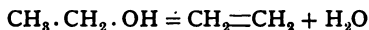
otherwise, in consequence of the cooling of the large flask, the contents of the bottles will be drawn back. The ethylene bromide is then washed repeatedly with water in a dropping funnel, and, finally, with caustic soda solution. It is dried over calcium chloride, and, on distillation, is obtained perfectly pure. Boiling-point, 130° . Yield, 125-150 grammes.

The addition of the alcohol-sulphuric acid mixture is often attended with difficulty, in that as soon as the cock is opened, the gas passes out through the funnel, thus preventing the entrance of the mixture. This difficulty may be obviated by taking the precaution of always keeping the stem of the funnel filled with the mixture. Before the heating is begun, a portion of the mixture is placed in a porcelain dish, the end of the stem of the funnel immersed in it and filled by suction. The cock is then closed, the funnel placed in the cork of the generating flask, and the heating begun.

(b) From Ethyl Alcohol and Phosphoric Acid

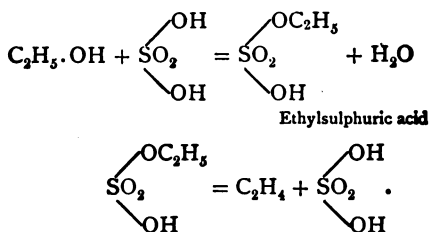
Separation of carbon and foaming will be prevented in the preparation of ethylene when syrupy phosphoric acid is used in place of sulphuric acid. A 200 c.c. round flask, with a wide mouth, is provided with a three-hole cork. Through one hole passes a thermometer extending to the bottom of the flask, through the second hole is inserted a dropping funnel; the stem of the latter is at least 25 cm. long and is drawn to a fine point. The end of the stem is kept only a few millimetres below the stopper. Through the third hole passes a delivery tube, not too narrow, bent at right angles. The tube is attached with rubber to a Wolfe flask, provided with two tubulures flush with the stoppers. The flask is completely surrounded with broken ice. Then follow two wash-bottles surrounded with ice for bromine, and finally a flask containing a solution of sodium hydroxide, as in method (a). For the preparation of ethylene 120 grammes of syrupy phosphoric acid, sp. gr. 1.7–1.75, are placed in an open vessel and gradually heated. The acid is stirred with a thermometer. At 160° water begins to be given off. The heating is continued, until at 220° the evolution of vapour is very slight. The acid is somewhat cooled and poured into the flask. It is heated to 210–220° over an asbestos gauze; at this temperature ordinary alcohol is allowed to run into the flask, drop by drop, from the funnel, the steam of which should have been previously filled with alcohol, as in method (a). A steady current of ethylene is thus evolved. The decolourisation of bromine requires 3½–4½ hours. The crude product is purified as in method (a).

The hydrocarbons of the ethylene series may be prepared, in general, by abstracting water from the corresponding alcohol, *e.g.*:

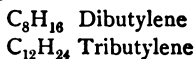


If sulphuric acid or phosphoric acid is used as the dehydrating agent, the reaction does not follow the above equation, but ethylsulphuric acid or ethylphosphoric acid is first formed, and this, on heating, yields sulphuric acid or phosphoric acid.

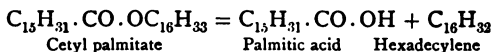
Q



In many cases the elimination of water takes place so easily that the use of concentrated sulphuric acid is unnecessary, since the diluted acid answers the purpose. With the higher members of the series the reaction is complicated by the fact that the simple alkylenes polymerise under the influence of sulphuric acid. Thus there is formed, besides butylene, C_4H_8 , hydrocarbons having respectively twice and three times its molecular weight, *e.g.* :

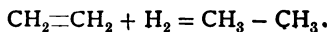


In these cases it is much more convenient to prepare an ester from the alcohol by the action of the chloride of a higher fatty acid, and subjecting this to distillation by which it is decomposed into an hydrocarbon of the ethylene series and the free fatty acid, *e.g.* :

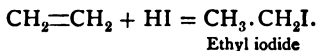


The first four members of the alkylene series are gases at ordinary temperatures, which burn with strongly luminous, smoky flames. The intermediate members are colourless liquids, not miscible with water, which can be distilled at ordinary pressures without decomposition; the higher members are solids, and can only be distilled without decomposition in a vacuum. Chemically these compounds are characterised primarily by the property of uniting with two univalent atoms, or a univalent atom and a univalent radical, upon which the double union is changed to single union.

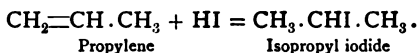
They take up, especially in the presence of platinum-black, two atoms of hydrogen, thus passing over to the hydrocarbons of the saturated series (paraffins) :



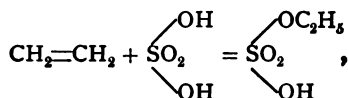
Hydrogen halides may also be added to them; hydriodic acid with the greatest ease, hydrobromic acid with less, and hydrochloric acid only with difficulty :



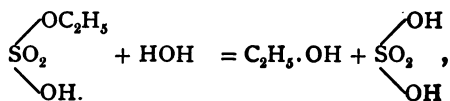
The homologues of ethylene also form addition products; the halogen atom seeks that carbon atom which is combined with the smallest number of hydrogen atoms :



The constituents of water (H and OH) may also be added indirectly to the alkynes. If concentrated sulphuric acid be allowed to act on one of them, it dissolves, forming a sulphuric acid ester :



If this is boiled with water, the ester is decomposed into alcohol and sulphuric acid :

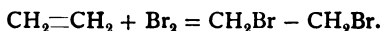
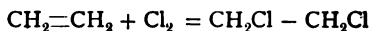


so that finally H and OH have been added to ethylene :



Analogous to the halogen atoms, the hydroxyl (OH) group unites with that carbon atom holding in combination the smallest number of hydrogen atoms.

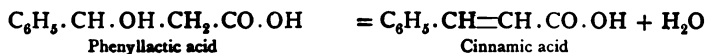
The alkynes take up two atoms of chlorine or bromine with great ease :



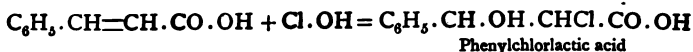
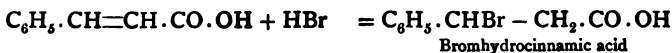
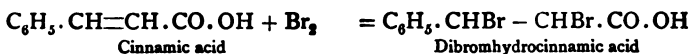
Finally they combine directly with hypochlorous acid to form glycol-chlorhydrines.

The reactions taking place in the formation of the alkynes as well as those in the formation of addition products are not only applicable

to the hydrocarbons but also to their substitution products. Thus, *e.g.*, unsaturated acids are commonly obtained from oxyacids by the elimination of water:



All compounds in which the ethylene condition is present show the addition phenomena, in accordance with the following equations:



13. REACTION: REPLACEMENT OF HALÖGEN ATOMS BY ALCOHOLIC HYDROXYL GROUPS

EXAMPLE : Ethylene Alcohol (Glycol) from Ethylene Bromide

(a) *Conversion of Ethylene Bromide into Glycoldiacetate*

A mixture of 60 grammes ethylene bromide, 20 grammes glacial acetic acid, and 60 grammes of freshly fused, finely pulverised

potassium acetate,¹ placed in a $\frac{1}{2}$ -litre, short-necked, round flask, provided with a reflux condenser, is heated to active boiling for two hours on a sand-bath over a large flame. The reaction product is then distilled (with a condenser) over a large luminous flame kept in continuous motion. Toward the end of the distillation the flame is gradually made non-luminous. The distillate is then further treated with 60 grammes ethylene bromide and 80 grammes potassium acetate, and the mixture, as above, heated to active boiling for two to three hours on a sand-bath. The reaction product is then again distilled over (with a condenser) by a luminous flame. The distillate is fractioned — using a 10 cm. long Hempel tube. The fractions are collected as follows: 1. up to 140° ; 2. from 140 – 175° ; 3. above 175° . Fractions 2 and 3 are then again distilled separately. The pure glycoldiacetate goes over between 180 – 190° , the main portion at 186° . Yield, about 70 grammes.

If it is desired to increase the yield, the portion going over under 180° is heated for three hours longer with potassium acetate. The product is then treated as above described. This causes an increase of about 15 grammes.

(b) *Saponification of Glycoldiacetate*²

Glycoldiacetate is saponified by heating with a solution of hydrochloric acid in methyl alcohol. For this purpose a mixture of 100 grammes ordinary methyl alcohol and a quantity of compact slaked lime (about one-third of the alcoholic volume) is heated for several hours on a water-bath to active boiling, in a flask at-

¹ Potassium acetate (Kalium aceticum pur. Ph. G. III.), differing from sodium acetate (compare page 147), crystallises without water of crystallisation. Nevertheless, for this experiment it must be heated to fusion over a free flame in an iron or nickel dish. The melted salt is poured into a shallow, flat iron or copper dish, in a thin layer. While still warm it is pulverised as finely as possible, and must be at once transferred to a bottle which is to be kept tightly closed. For this experiment 200 grammes of the salt are fused.

² A private method; by courtesy of Prof. Henry (Löwen).

tached to a reflux condenser. The dehydrated methyl alcohol is now distilled. It is then subjected to fractional distillation, and the portion distilling at $66-67^{\circ}$ is collected separately. 44 grammes of pure methyl alcohol are weighed in a small flask tared with its delivery tube. Gaseous hydrochloric acid is now passed into the alcohol with cooling under water, until a gain of 1.1 grammes is obtained. Should the increase in weight be more than this, a calculated quantity of pure methyl alcohol is added in order to obtain a $2\frac{1}{2}\%$ alcoholic solution of hydrochloric acid, which is the necessary strength.

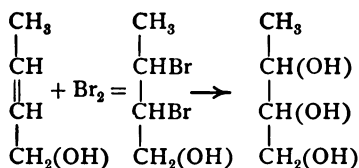
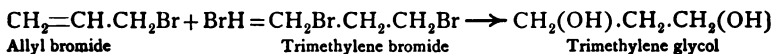
In a flask, provided with a reflux condenser, 45.1 grammes of the alcoholic solution of hydrochloric acid, and 50 grammes of glycoldiacetate are heated on an actively boiling water-bath for half an hour. The reaction mixture is then *quickly* distilled from a water-bath with frequent shaking; methyl alcohol and methyl acetate will thus distil over, while glycol will remain in the flask with a small quantity of unsaponified ester. These cannot be separated from one another by distillation, as their boiling points are close together. But the thick liquid in the flask is shaken twice with an equal volume of dry ether, whereby glycoldiacetate is taken up, while glycol remains undissolved. The ethereal layer is removed, either by decantation, or by the use of a pipette. The glycol is then poured into a small fractionating flask connected with a long condenser. During the distillation (heat slowly at first) a low-boiling fraction (to 100°) is first obtained, when the thermometer rises rapidly to 190° . The main portion of glycol distils over at 195° . Yield about 80-90 % of the theory (17-19 grammes).

This preparation shows a method for replacing a halogen atom by an alcoholic hydroxyl group. In Reaction 1 the reverse replacement was brought about, — the substitution of a hydroxyl group by a halogen. This method is obviously only of importance in those cases in which it is more convenient to obtain the halogen derivative than the alcohol. Among the monacid alcohols it is of value for preparing isopropyl alcohol, normal secondary butyl, and normal secondary hexyl alcohols.

As stated on page 135, the action of hydriodic acid on polyacid alcohols does not yield, as might be expected, the poly-iodine derivatives, but mono-iodine derivatives. Thus from glycerol, isopropyl iodide is obtained; from erythrite, normal secondary butyl iodide; from mannite, the normal secondary hexyl iodide. These iodides, as pointed out, may be converted into the corresponding alcohols. The method is of practical value in the preparation of tertiary alcohols from acid-chlorides and zinc alkyls. — Butlerow's synthesis. Compare page 146. In this reaction the tertiary chloride is formed as an intermediate product. The method is of importance for the preparation of di-acid alcohols (glycols), especially for the α -glycols, in which the hydroxyl groups are combined with the two adjacent carbon atoms. The dibromides corresponding to these alcohols are easily obtained by the addition of bromine to the hydrocarbons of the ethylene series. In this way glycol was first prepared by Wurtz.¹

Other glycols may be obtained in a similar manner, *e.g.*, if allyl bromide be treated with hydrobromic acid, trimethylene bromide is formed, from which a β -glycol — trimethylene glycol — may be obtained by the above reaction.

If, to unsaturated mono-acid alcohols containing a double union, two bromine atoms be added, dibrom-alcohols are obtained which, by replacing the bromine with hydroxyl, yield tri-acid alcohols:



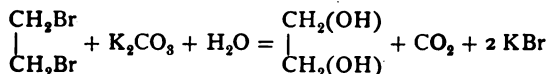
From these examples the value of this method for obtaining alcohols is evident.

Oxyaldehydes, oxyketones, and oxyacids may also be obtained by this reaction, from the corresponding halogen compounds. Finally, it may be employed to replace the halogen, in side-chains of aromatic compounds, by hydroxyl.

The substitution of halogen atoms by hydroxyl groups may be done

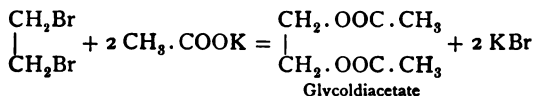
¹ A. ch. (3), 55, 400.

by two methods: (1) Directly in a single operation. (2) In two reactions; (a) by preparing an acid-ester of the desired alcohol, and (b) subjecting this to saponification. If method (1) be used, the halogen derivative is heated with water at the ordinary pressure, or if necessary, at an increased pressure. The same object is attained more quickly, and in many cases with a better yield, by the addition to the reaction-mixture of certain oxides, hydroxides, or carbonates. Silver oxide, lead hydroxide, barium hydroxide, potassium or sodium carbonate, and others may be used for this purpose. It appears to be true that a tertiary halogen atom reacts more easily than a secondary or primary, and that a secondary, more easily than a primary. By following this method, glycol may be obtained directly from ethylene bromide, if the latter be heated with water and potassium carbonate:

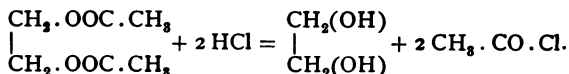


The separation of the glycol from the large excess of water used is troublesome.

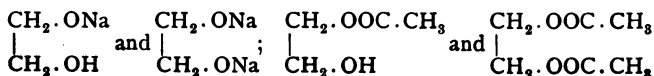
In accordance with method (2) certain salts, as silver, potassium, or sodium acetate are allowed to act on the halogen substitution product. This results in the formation of an ester of the desired alcohol:



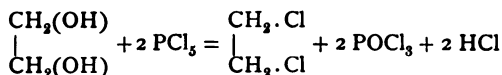
The ester is then saponified under the proper conditions, upon which the free alcohol is obtained:



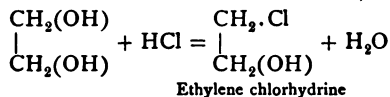
The acetyl chloride reacts with methyl alcohol forming methyl acetate with the liberation of a fresh quantity of hydrochloric acid. Glycol is a thick, colourless, odourless liquid, boiling at 195° ; it melts at 11.5° after having been solidified by low temperature. Like all poly-acid alcohols, it has a sweet taste. It is easily soluble in water and in alcohol, but not in ether. Chemically it differs only from the mono-acid alcohols in its ability to form mono- or di-derivatives according to the conditions:



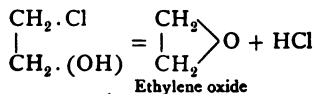
Both hydroxyl groups are replaced by the action of phosphorus pentachloride :



But if glycol be heated with hydrochloric acid, only one hydroxyl group is replaced :



From these so-called halogen hydrines, by the action of alkalies the inner anhydrides of the glycols are obtained :



TRANSITION FROM THE ALIPHATIC TO THE AROMATIC SERIES

Dimethylcyclohexenone and *s*-Xylenol from Ethylidenebisacetacetic Ester. (Ring Closing in a 1.5 Diketone. Knoevenagel Reaction.¹)

1. ETHYLIDENEBISACETACETIC ESTER

In a thick-walled flask closed by a cork bearing a thermometer reaching almost to the bottom, treat 50 grammes of *pure* (in vacuum distilled), cooled acetacetic ester with 8.5 grammes of *pure* aldehyde distilled just before the experiment. The flask is cooled to $-10-15^{\circ}$ in a freezing mixture of ice and salt. To the reaction-mixture is then added a few drops of diethyl amine from a small medicine "dropper." In most cases no elevation of temperature takes place at first. Since it is very difficult to obtain acetacetic ester and aldehyde absolutely free from acids, the first portions of the amine are neutralised by the acids present, and are thus not available for the main reaction. The addition of the amine is continued slowly until at a certain point an elevation of a few degrees in the temperature is observed. Normally this should occur on the addition of the first ten drops. When this takes place, the liquid, clear at first, becomes turbid. From this point, during the gradual addition of a further portion of ten drops of the base, the temperature is slowly allowed to rise to 0° . The addition of the base in drops is continued, gradually and with frequent shaking, until, collectively, 60 drops = 1.5 grammes have been used. The length of the operation is about an hour. After the reaction-mixture has stood a further quarter hour, it is removed from the freezing mixture and allowed to come to the room temperature. If, in consequence of a secondary reaction, the temper-

¹ A. 281, 25.

ature should go up to 20° , the flask is cooled off a short time in ice water. The reaction-product is a viscous, bright yellow liquid in which numerous drops of water are suspended. It is allowed to stand undisturbed until it solidifies to a crystalline mass, which generally requires from two to three days.

A small specimen is pressed out on a porous plate and recrystallised from diluted alcohol. Colourless needles are thus obtained which melt at $79-80^{\circ}$. Concerning the constitution of this substance see the remarks on page 207.

The solidification of the crude product may be hastened by seeding it, after one day's standing, with crystals obtained in a previous preparation. This is best done on the upper portion of the flask, which is only moistened by the liquid.

2. DIMETHYLCYCLOHEXENONE

The crude product liquefied by heating in a water-bath is poured into a mixture of 400 grammes of water and 100 grammes of concentrated sulphuric acid contained in a round litre flask provided with a long reflux condenser. The reaction-mixture is heated to lively boiling on a wire gauze; a few pieces of unglazed porcelain are placed in the flask to insure a regular ebullition.

After about seven hours' heating (the experiment should be commenced in the morning of a working-day), the reflux condenser is replaced by an ordinary condenser, and steam is passed into the mixture until the distillate measures about 100 c.c. The flask is heated by a free flame up to the boiling-point of its contents. The distillate is preserved in a well-closed vessel.

On the second day the mixture is again heated for seven hours (with a reflux condenser, new porcelain scraps in the flask), and then 100 c.c. are again distilled off with steam. This is repeated on the third day, and finally steam is passed into the mixture until from a test portion of the distillate saturated with solid potash no oil, or only a minute quantity, separates out. The three distillates in which the reaction-product is for the most part dissolved are now united. Solid potash is added until it is no longer dissolved.

For the success of the salting out, one must use anhydrous potash

as pure as possible. From the potash solution a brownish red oily layer separates out: it consists of dimethylcyclohexenone and alcohol. It is separated from the water solution in a dropping funnel, and the alcohol is distilled off by the aid of a Hempel tube 10 cm. long filled with glass beads. The residue is dried over fused Glauber's salt and distilled from an ordinary fractionating flask. The portion going over between $200-215^{\circ}$ is collected separately. Boiling-point of the pure compound, 211° . Yield, 15-20 grammes.

3. s-XYLENOL

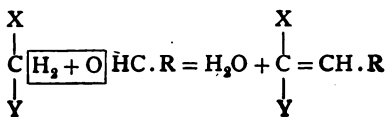
A mixture, cooled by ice water, of 10 grammes of the ketone dissolved in 20 grammes of glacial acetic acid (the acid must not be allowed to solidify) is treated gradually with a mixture of 13 grammes of bromine and 10 grammes of glacial acetic acid from a dropping funnel. The reaction-mixture is then allowed to stand, under the hood, at least half a day, or better over night, at the room temperature. Hydrobromic acid is evolved copiously. The mixture is heated, with frequent shaking, about an hour on a water-bath to about 50° , the temperature is then increased until the water boils, and the heating is continued until there is only a slight evolution of hydrobromic acid. It is heated finally, using an air condenser, on a wire gauze, to incipient ebullition of the acetic acid, until the evolution of hydrobromic acid almost entirely ceases. After cooling, it is poured carefully into a cooled solution of 75 grammes of caustic potash in 150 grammes of water, upon which only a small quantity of an oil should separate out. The by-products insoluble in the alkaline solution are extracted with a sufficient quantity of ether, the alkaline solution is saturated with carbon dioxide, and the s-xyleneol liberated is distilled over in the presence of carbon dioxide with steam (use a three-hole cork). The end of the distillation may be readily determined. So long as the xyleneol is coming over, a test of the distillate by adding a few drops of bromine will show a precipitate of tribromxyleneol.

If the distillate be allowed to stand in a cool place over night,

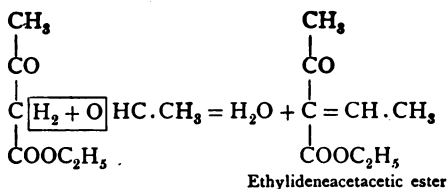
the larger portion of the xylenol will crystallise out. In order to obtain the portion remaining dissolved, the crystals are filtered off, and the filtrate saturated with solid salt and extracted with ether. Melting-point of *s*-xylenol, 64°. Boiling-point, 220–221°. Yield, 5–6 grammes.

A better characterisation of this phenol is obtained by covering a few drops of it in a test-tube with 5 c.c. of water and then adding bromine drop by drop until the reddish brown colour of the latter does not disappear. The excess of bromine is removed by the addition of a solution of sulphur dioxide. The precipitate is recrystallised from alcohol. There are thus obtained colourless needles of tribromxylenol, which melt at 165°.

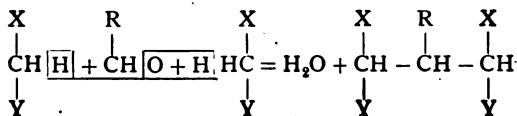
1. Aldehydes unite, with the elimination of water, with compounds containing the group CH_2 between two negative radicals (acetacetic ester, malonic ester, acetylacetone, and others), in two ways. 1. Equal molecules unite in accordance with the following equation:



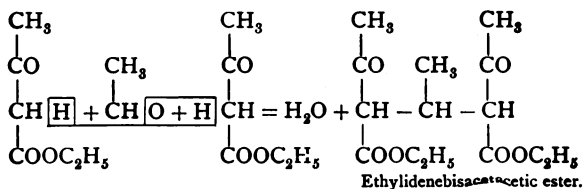
Example:



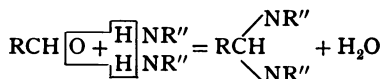
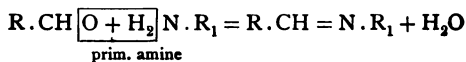
2. The reaction may take place between *one* molecule of the aldehyde and *two* molecules of the other compound:



Example, carried out in practice :

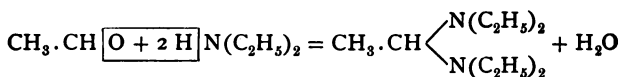


For bringing about the first reaction the following-named substances may be used as condensation agents: hydrochloric acid, acetic anhydride, as well as primary and secondary amines (ethyl amine, diethyl amine, piperidine, and others). For the second reaction the bases mentioned may be used. A small quantity of one of these may produce large quantities of the condensation product: this is a case of a so-called continuous reaction. It is probable that the amine reacts first with the aldehyde, water being eliminated: ¹

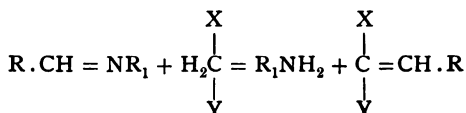


(R'' = a bivalent radical or two univalent radicals.)

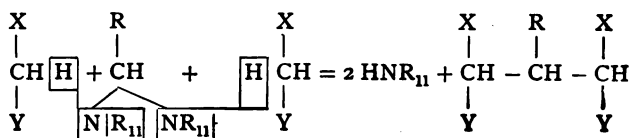
In the example above :



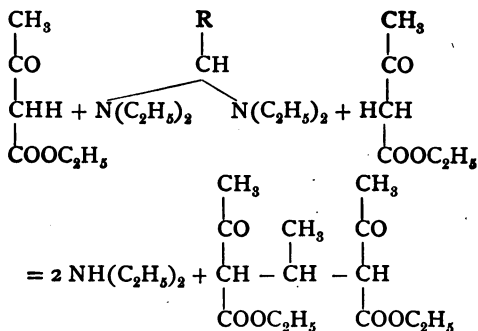
The aldehyde derivative thus formed then acts upon the second compound with the regeneration of the amine :



¹ B. 31, 738.

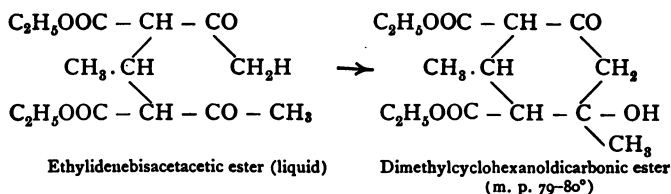


In the above example :



The amine thus regenerated carries over anew the aldehyde residue to the acetacetic ester, and so on.

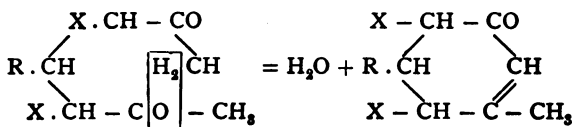
According to Rabe (A. 323, 83 and 332, 1), the substance melting at 79–80° does not possess the constitution of an ethylidenebisacetacetic ester, but it shows a desmotropic modification, forming a cyclic compound as follows :



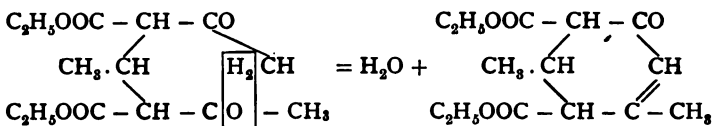
But compare the objections of Knoevenagel (B. 36, 2118).

2. Of the compounds which can be obtained by reaction (2), of especial interest are those which, like the ethylidenebisacetacetic ester prepared above, contain two carbonyl groups (1.5-diketones) and in addition a methyl group. If such compounds are treated with those

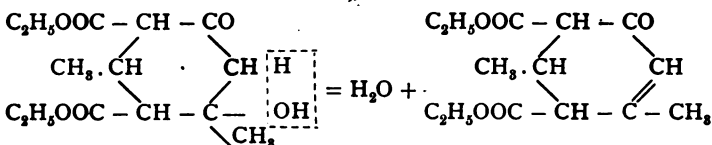
substances which have the power to eliminate water (alkalies or acids), six-membered carbon rings are formed as follows :



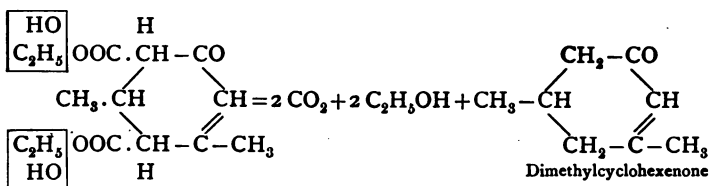
In the above example :



According to Rabe, ring formation takes place as indicated above, and the change at this stage is as follows :

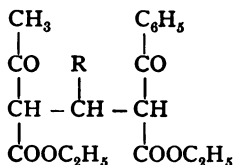


Beside this ring closing, a second reaction takes place in the experiment made above. The sulphuric acid saponifies the primarily formed acid-ester, with the elimination of carbonic acid :

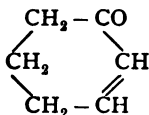


This ring closing with 1.5 diketones is capable of many modifications. By using formaldehyde, acetaldehyde, propionaldehyde, benzaldehyde, etc., R = H, CH₃, C₂H₅, C₆H₅, etc. By using acetacetic ester, acetylacetone, benzoylacetone, etc., X = COOC₂H₅, CH₃CO, C₆H₅CO, etc.

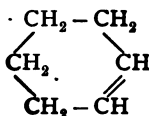
The many-sidedness of the reaction is materially increased by starting with the unsymmetrical 1.5 diketone, *e.g.* :



The nature of the reaction requires, however, that one of the two carbonyl groups must be connected with a methyl group, otherwise the elimination of water cannot take place. The compounds so obtained are all derivatives of the mother substance :

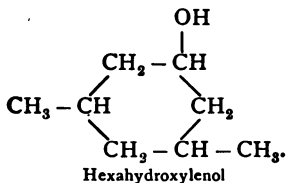


which is called cyclohexenone, and which may be considered as the keto-derivative of tetrahydrobenzene

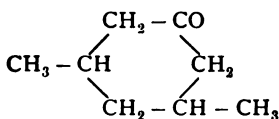


This, therefore, is a transition from the aliphatic to the (hydro) aromatic series.

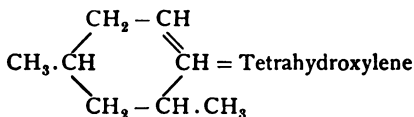
This primarily obtained compound may by different reactions be converted into other hydroaromatic and aromatic substances. If, *e.g.*, the dimethylcyclohexenone be reduced, the ketone group is converted into the secondary alcohol group, at the same time the double union is severed, and two hydrogen atoms are added on, so that there is obtained an alcohol derivative of hexahydro-benzene or -xylene.



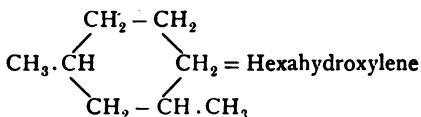
If this compound be oxidised, the secondary alcohol group is changed to a ketone group, and a keto-derivative of a hexahydroxylene is formed :



If the hexahydrogen addition alcohols be treated with substances having the power to eliminate water, there is obtained a tetrahydrogen addition product of the hydrocarbon, *e.g.*:

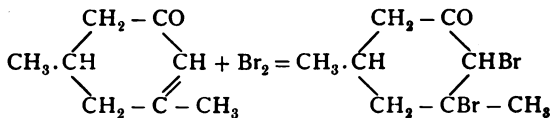


If in compounds like hexahydroxyleneol, the hydroxyl be replaced by iodine, and the resulting iodide reduced, there is obtained a hexahydrogen addition product of the hydrocarbon, *e.g.*:

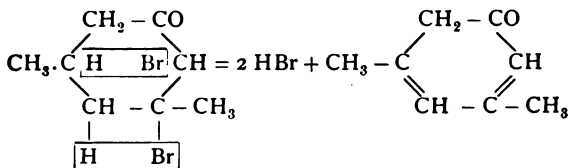


It is thus evident that the syntheses of a great variety of hydroaromatic compounds are possible.

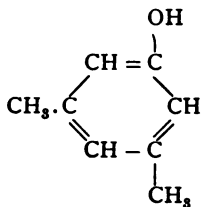
3. By these reactions the compounds of the pure aromatic series may be reached. If bromine be allowed to act on the primarily arising ring compound, as has been done practically, the double union is broken up by the addition of two atoms of bromine:



These dibromides are very unstable, and even in the cold give off two molecules of hydrobromic acid:



Finally, this unstable keto-form ($\text{CH}_2\text{-CO}$) changes itself into the stable enol-form ($\text{CH} = \text{C} \cdot \text{OH}$), and the *s*-xlenol is obtained.



For further information concerning the transition from aliphatic to aromatic or hydroaromatic compounds, compare Bernthsen, XI Ed., p. 382; Richter, X Ed., Vol. II, pp. 4-6, and 35; Krafft, IV Ed., p. 446; Meyer-Jacobson, II Vol., p. 79.

II. AROMATIC SERIES

1. REACTION: NITRATION OF A HYDROCARBON

EXAMPLES: Nitrobenzene and Dinitrobenzene¹*Nitrobenzene*

To 150 grammes of concentrated sulphuric acid contained in a $\frac{1}{2}$ -litre flask, add gradually, and with frequent shaking, 100 grammes of concentrated nitric acid (sp. gr. 1.4). After cooling the mixture to the room temperature, by immersion in water, gradually add 50 grammes of benzene, with frequent shaking. If the temperature should rise above 50–60°, the operation is interrupted, and the flask immersed in water for a short time. When all of the benzene has been added, a vertical air condenser is attached to the flask; it is then heated in a water-bath for an hour at 60° (thermometer in the water); during the heating the flask is frequently shaken. After cooling, the lower layer, consisting of sulphuric and nitric acids, is separated from the upper layer of nitrobenzene in a separating funnel. The nitrobenzene is then agitated in the funnel several times with water: it must be borne in mind that the nitrobenzene now forms the *lower* layer. After being washed, it is placed in a dry flask, and warmed on a water-bath with calcium chloride until the liquid, milky at first, becomes clear.² It is finally purified by distillation from a fractionating flask provided with a long air condenser. Boiling-point, 206–207°. Yield, 60–70 grammes.

Dinitrobenzene

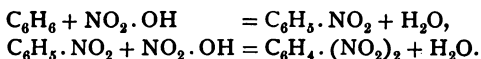
To a mixture of 25 grammes of concentrated sulphuric acid and 15 grammes of fuming nitric acid, 10 grammes of nitrobenzene are

¹ A. 9, 47; 12, 305. Ostwald's Klassiker der exacten Wissenschaften Nr. 98.

² The crude product, separated from the acid and treated with water, may also be distilled with steam. The higher nitro derivatives are not volatile with steam. The distillate is now treated as above.

gradually added (hood) ; the reaction-mixture is then heated for half an hour on a water-bath, with frequent shaking ; after cooling somewhat, it is poured, with stirring, into cold water. The dinitrobenzene which solidifies is filtered off, washed with water, pressed out on a porous plate, and recrystallised from alcohol. Melting-point, 90° . Yield, 10–12 grammes.

The property of yielding nitro-derivatives, when treated with nitric acid, is a characteristic of aromatic compounds. According to the conditions under which the nitration is carried out, one or more nitro-groups can be introduced at the same time. The above reactions take place in accordance with the following equations :



If a saturated aliphatic residue is present in an aromatic compound, the nitration under the above conditions always affects the benzene ring, and not the side-chain. Since the benzene carbon atoms are in combination with only one hydrogen atom, the nitro-compounds obtained on nitration are tertiary ; they therefore do not have the power to form salts, nitrolic acids, or pseudo-nitroles, like the primary and secondary nitro-compounds.

Recently the nitro-group has been introduced directly into the side-chain.¹ If, *e.g.*, toluene or ethyl benzene be heated with weak nitric acid (sp. gr. 1.076) in a bomb up to about 100° , phenylnitromethane, $\text{C}_6\text{H}_5 \cdot \text{CH}_2 \cdot \text{NO}_2$, or phenylnitroethane, $\text{C}_6\text{H}_5 \cdot \text{CH} \cdot \text{NO}_2 \cdot \text{CH}_3$ is obtained.

Not only can the mother substances, the aromatic hydrocarbons, but all their derivatives, as phenols, amines, aldehydes, acids, etc., undergo similar reactions. But the nitration does not take place in every case with the same ease. In each case, therefore, the most favourable conditions for the experiment must be determined. If a compound is very easily nitrated, the nitration may be effected, according to the conditions, by nitric acid diluted with water, or the substance may be dissolved in a solvent which is not attacked by nitric acid ; glacial acetic acid is frequently used for this purpose, and then treated with nitric acid. The reverse process may also be employed, *i.e.* the substance is added to a mixture of nitric acid and water, or nitric acid and glacial acetic acid. If a substance is moderately difficult to nitrate, it is added

¹ B. 27. Ref. 194 and 468.

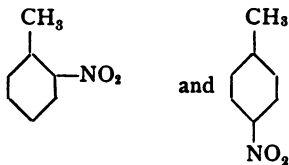
to concentrated or fuming nitric acid. If the nitration is difficult, the elimination of water is facilitated by the addition of concentrated sulphuric acid to ordinary or fuming nitric acid. In the nitration, the substance may either be added to the mixture of nitric acid and sulphuric acid, or the nitric acid is added to the substance dissolved in concentrated sulphuric acid. In working with sulphuric acid solutions, at times either potassium nitrate or sodium nitrate may be used instead of nitric acid. The three nitration methods just described may be still further modified in two ways: (1) the temperature may be varied; (2) the quantity of nitric acid may be varied. The nitration can be effected in a freezing mixture, in ice, or in water, by gentle heating, or finally, at the boiling temperature. Further, the theoretical amount of nitric acid, or an excess, may be used. In order to determine which of these numerous modifications will give the best results, preliminary experiments on a small scale must be made. Since the nitro-compounds are generally insoluble in water, or difficultly soluble, they can be separated from the nitrating mixture by diluting it with water, or in many cases better, with a solution of common salt.

The chemical character of a substance is not changed in kind, but in degree, by the introduction of a nitro-group. Thus, the nitro-derivatives of the hydrocarbons are indifferent compounds like the hydrocarbons. If a nitro-group is introduced into a compound of an acid nature like phenol, it becomes more strongly acid, *e.g.* the nitro-phenols are more strongly acid than phenol. When a nitro-group is introduced in a basic compound, the resulting substance is less basic; *e.g.* nitro-aniline is less basic than aniline.

The great importance of the nitro-compounds is due to their behaviour on reduction; this will be considered under the next preparations.

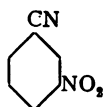
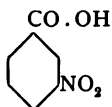
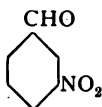
Concerning the introduction of the nitro-group,¹ the following laws are of general application.

The introduction of one nitro-group in the benzene molecule can, obviously, only result in the formation of one mononitrobenzene. If an alkyl radical is present in the benzene molecule, the nitro-groups enter the ortho- and para-, but only to a slight extent the meta-position to the radical. On nitrating toluene, *e.g.*, there are formed almost exclusively:

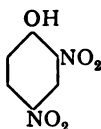
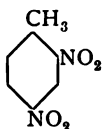


¹ The same laws apply also to the introduction of halogen and sulphonic acid groups.

The nitro-groups seek the same position when a benzene-hydrogen atom has been substituted by hydroxyl. Thus, *e.g.*, phenol gives on nitration a mixture of *o*- and *p*-nitrophenol. On the other hand, if a compound contains an aldehyde-, carboxyl-, or cyanogen-group, on nitration the nitro-group goes in the meta-position to this. Benzaldehyde, benzoic acid, and benzonitrile give on nitration respectively :

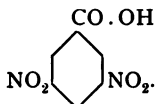


If a compound already contains a nitro-group, a second one will take the meta-position to this. Thus, on nitrating nitrobenzene, *m*-dinitrobenzene is formed. *O*-nitrotoluene or *o*-nitrophenol yield on nitrating :



respectively.

From *m*-nitrobenzoic acid the following dinitrobenzoic acid is formed :



The nitro-compounds are in part liquids, in part solids; in case these latter distil without decomposition, they possess a higher boiling-point than the mother substance.

2. REACTION: REDUCTION OF A NITRO-COMPOUND TO AN AMINE

EXAMPLES: (1) Aniline from Nitrobenzene¹

(2) Nitroaniline from Dinitrobenzene

A mixture of 90 grammes of granulated tin² and 50 grammes of nitrobenzene is placed in a 1½-liter round flask. To this are gradually added 200 grammes of concentrated hydrochloric acid

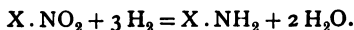
¹ A. 44, 283.

² If granulated tin is not at hand, it may be prepared by melting block-tin in an iron spoon over a blast flame. The molten mass is allowed to fall in drops into a pail of water from a height of ½-1 metre.

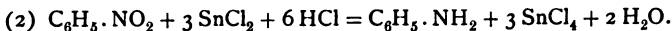
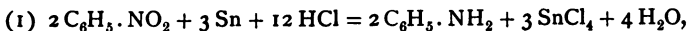
in the following manner: At first only about one-tenth of the acid is added; an air condenser, not too narrow, is then attached to the flask and the mixture well shaken. After a short time it becomes warm, and finally an active ebullition takes place. As soon as this happens, the flask is immersed in cold water until the reaction has moderated. The second tenth of the acid is then added, and the above operation repeated. After one half of the acid has been used, the reaction becomes less violent, and the second half may be added in larger portions. In order to effect the reduction of the nitrobenzene completely, the mixture is finally heated one hour on the water-bath. To separate the free aniline, the warm solution is treated with 100 c.c. of water, then a solution of 150 grammes of caustic soda in 200 grammes of water is gradually added. The mixture should finally give a strongly alkaline reaction. If the action of the caustic soda causes the liquid to boil, the flask is cooled by water for a short time, before a further addition of caustic soda. When all of the solution has been added, a long condenser is attached to the flask, and steam is passed into the hot liquid, upon which aniline, as a colourless oil, and water pass over, the aniline collecting under the water. As soon as the distillate no longer appears milky, and becomes clear, the receiver is changed and about 300 c.c. more of the liquid distilled over. The distillates are mixed, treated with 25 grammes of finely powdered sodium chloride for every 100 c.c. of the liquid, shaken until all the salt is dissolved, and the aniline extracted with ether. After the ethereal solution has been dried by treating it with a few pieces of solid potassium hydroxide, the ether is evaporated and the aniline subjected to distillation. Boiling-point, 182° . Yield, 90–100% of the theory. If the circumstances are such as not to permit the experiment to be completed without interruption, it is so arranged that the neutralisation with sodium hydroxide, and the distillation with steam immediately following, may take place within a short time, so that the heat of neutralisation may be utilised.

To the nitro-compounds of the aromatic series, as well as those of the aliphatic series, belongs the property of being converted into primary amines on energetic reduction. For the reduction of every nitro-group,

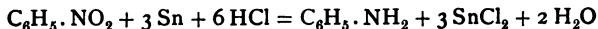
six atoms of hydrogen are necessary, and the following equation is the general expression of the reaction :



For the reduction of nitro-compounds on the small scale in the laboratory, it is most convenient to use, as the reducing agent, granulated tin and hydrochloric acid, or stannous chloride and hydrochloric acid :



To 1 molecule of a mononitro-compound, $1\frac{1}{2}$ atoms of tin, or 3 molecules of stannous chloride, are therefore used. In calculating the amount of the latter necessary for a reaction, it is to be remembered that the salt crystallises with two molecules of water ($SnCl_2 + 2 H_2O$). If the reduction is to be effected by metallic tin, double the above quantity is frequently used, *i.e.* to 1 nitro-group, 3 atoms of tin. In this case, the tin is not converted into stannic chloride, but into stannous chloride :

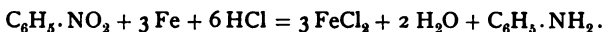


Since, in the cases mentioned, hydrochloric acid is always present in excess, and the amines unite with it to form soluble salts, the end of the operation occurs when no more of the insoluble nitro-compound is present, and the reaction-mixture dissolves clear in water. In order to get the free amine from the acid mixture, various methods may be employed. If, as in the above example, the amine is volatile with steam, and insoluble in alkali, then the acid solution is treated with caustic potash, or caustic soda, until the oxide of tin which separates out at first is redissolved in the excess of alkali; the liberated amine is driven over with steam. Further, volatile or non-volatile amines can be extracted from an alkaline solution by a proper solvent, like ether. But this process is often troublesome, since the alkaline tin solution forms an emulsion with ether, which subsides with great difficulty. If the free amine is solid, it may be obtained by filtering off the alkaline liquid. In many cases, where a non-volatile amine is under examination, it is advisable to precipitate the tin before liberating the amine. This is done by diluting the acid solution with much water, heating on the water-bath, and as soon as the liquid has reached the temperature of the bath, hydrogen sulphide is passed into it. The tin is precipitated as stannous or stannic sulphide; this is separated from

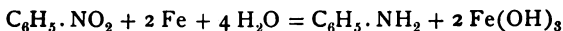
the amine hydrochloride by filtering. Since tin, in the presence of a large excess of hydrochloric acid, is precipitated only with difficulty by hydrogen sulphide, it is frequently necessary to drive off the excess of the acid before treating with hydrogen sulphide. This is done by evaporating to dryness on the water-bath.

After the tin sulphide has been filtered off, a portion of the filtrate is tested with hydrogen sulphide for tin; if it should be present, the whole filtrate is evaporated on the water-bath, as completely as possible, to remove the hydrochloric acid, then diluted with water, and hydrogen sulphide is again passed into it. At times, the amine forms with hydrochloric acid, a difficultly soluble salt, or the amine hydrochloride combines with the tin chloride to form a difficultly soluble double salt. In this case, the isolation of the amine may be facilitated by filtering it off, washing with hydrochloric acid, and pressing out on a porous plate, if necessary. If one is dealing with amines, which, like amido-acids, possess an acid character, obviously, these cannot be separated by the use of an alkali, as in the above example. In a case of this kind, the tin is always removed first, the acid solution evaporated to dryness, and the amido-compound is now liberated by the addition of sodium acetate. With amido-phenols, sodium hydrogen carbonate, sodium carbonate, or sodium sulphite may be used to decompose the hydrochloric acid salt.

In the laboratory, other metals, like iron, zinc, etc., in connection with an acid, are only rarely used in the place of tin or stannous chloride, for the reduction of nitro-compounds. On the large scale, iron, owing to its cheapness, is used in the preparation of bases like aniline, toluidine, α -naphthyl amine, etc., from the corresponding nitro-compounds. By the use of iron and hydrochloric acid, the reduction should theoretically take place in accordance with the following equation:

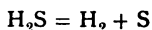


As a matter of fact, on the large scale, much less hydrochloric acid (only $\frac{1}{4}$) is used than that required by the above equation. In the presence of ferrous chloride, the nitro-compound is reduced by the iron without the action of hydrochloric acid, according to the equation:



For the neutralisation of the hydrochloric acid, a small quantity of which is always used on the large scale, slaked lime is employed in preference to the more costly alkalis.

The complete reduction of nitro-compounds containing several nitro-groups is conducted in the same way as for mononitro-compounds. If it is desired to reduce but one or two of several nitro-groups, it cannot be done by adding just the calculated amount of the reducing agent; for cases of this kind, special methods are necessary. For this purpose, hydrogen sulphide in the presence of ammonia or ammonium sulphide is often used for the reduction:

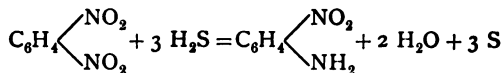


The compound to be reduced is dissolved in water or alcohol, according to circumstances, treated with ammonia, heated, and hydrogen sulphide passed into it; or it is heated in a water or alcohol solution with a previously prepared water or alcohol solution of ammonium sulphide. In this way, *e.g.*, dinitrohydrocarbons may be converted into nitro-amines. A second method, which may be generally used for the reduction, step by step, of compounds containing several nitro groups, is this: An alcoholic solution of the theoretical amount of stannous chloride saturated with hydrochloric acid is gradually allowed to flow into an alcoholic solution of the substance to be reduced, which is well cooled, and constantly shaken. (B. 19, 2161.)

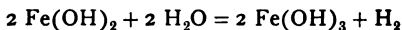
EXPERIMENT:¹ The recrystallised dinitrobenzene is dissolved in alcohol (4 grammes alcohol to 1 gramme dinitrobenzene), in a flask, the solution is quickly cooled down, upon which a portion of the dinitrobenzene separates out; it is then treated with 0.8 gramme of concentrated ammonia for 1 gramme dinitrobenzene (the ordinary dilute solution of ammonia employed as a reagent must not be used). After the flask and its contents have been tared, the mixture is saturated with hydrogen sulphide at the ordinary temperature; the current of hydrogen sulphide is then shut off, and the flask, provided with a reflux condenser, is heated for about half an hour on a water-bath. It is then allowed to cool to the ordinary temperature, and hydrogen sulphide again passed into it to saturation, etc. This operation is repeated until there is an increase of 0.6 gramme in weight for every gramme of dinitrobenzene used. If in consequence of insufficient cooling the required increase in weight does not take place, hydrogen sulphide

¹ A. 176 14.

is again passed into the mixture. It is then diluted with water, the precipitate filtered off, washed with water, and extracted several times by warming with dilute hydrochloric acid. From the acid filtrate, the nitro-aniline is set free by neutralising with ammonium hydroxide; it is recrystallised from water. Melting-point, 114° . Yield, 70–80% of the theory.

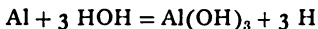


Special methods are necessary for the reduction of nitro-compounds containing groups capable of being acted upon by hydrogen, *e.g.*, an aldehyde-group, an unsaturated side-chain, etc. In cases of this kind, ferrous hydroxide is frequently used:



The reduction is effected by adding to the substance to be reduced, in the presence of an alkali (potassium-, sodium-, or barium-hydroxide), a weighed quantity of ferrous sulphate. By this reaction, o-nitrobenzaldehyde is reduced to o-amidobenzaldehyde; o-nitrocinnamic acid to o-amidocinnamic acid.

As a perfectly neutral reducing agent, which appears to be well adapted for a great variety of reduction reactions, aluminium amalgam¹ is recommended. It is made by treating aluminium filings or shavings, which have been slightly acted on by caustic soda, with a solution of mercuric chloride. It reacts with water in accordance with this equation:



Besides the reducing agents mentioned, there is still a large number of others which find only an occasional application in reducing nitro-compounds to amines. They will be referred to under the different preparations.

The primary mon-amines are in part colourless liquids, *e.g.*, aniline, o-toluidine, xylidine; or colourless solids like p-toluidine, pseudocuminidine, the naphthyl amines, etc. They can be distilled without decomposition, are volatile with steam, and difficultly soluble in water. The di- and poly-amines are for the most part solids, non-volatile with steam, and much more readily soluble in water than the mon-amines.

¹ B. 28, 1323.

The amines possess a basic character, but the basicity is weaker than that of the aliphatic amines, in consequence of the negative nature of the phenyl group.

Salts: $\text{C}_6\text{H}_5\cdot\text{NH}_2\cdot\text{HCl}$ Aniline hydrochloride
 $\text{C}_6\text{H}_5\cdot\text{NH}_2\cdot\text{HNO}_3$ Aniline nitrate
 $(\text{C}_6\text{H}_5\cdot\text{NH}_2)_2\cdot\text{H}_2\text{SO}_4$ Aniline sulphate

Like ammonia, the amines unite with calcium chloride to form double compounds; for this reason they must not be dried with this substance (see page 54).

The primary mon-amines find numerous applications in the laboratory, as well as on the large scale, in consequence of their great activity. Frequent reference will be made to the subject in the following pages.

With the aniline prepared above, the following experiments are made:

(1) Add 3 drops of aniline to 10 c.c. of water in a test-tube, and shake the mixture. The aniline dissolves. At moderate temperatures, 1 part of aniline dissolves in about 30 parts of water.

(2) Dilute 1 c.c. of this aniline solution with 10 c.c. of water, and add a small quantity of a filtered water solution of bleaching powder. A violet colouration takes place; by this reaction (Runge's), the most minute quantity of free aniline may be detected. If in this experiment the solution should not remain clear, but a dirty violet precipitate separate out, a too concentrated solution has been used; the aniline water is diluted further, and the experiment repeated. If a salt of aniline is to be tested, it is dissolved in water, treated with alkali, the free aniline extracted with ether, this latter evaporated, and the residue dissolved in water. Then proceed exactly as just directed.

This reaction may also be used to detect small quantities of benzene or nitrobenzene. In a test-tube mix 5 drops of concentrated sulphuric acid with 5 drops of concentrated nitric acid, then add 1 drop of benzene, shake, and warm gently by passing the tube through a flame several times. Then add 5 c.c. of water, and extract the nitrobenzene with a little ether; the ether layer is removed with a capillary pipette, and the ether evaporated. The residue is treated with 1 c.c. of concentrated hydro-

chloric acid, and to this is added a piece of zinc the size of a lentil, to effect the reduction. When the zinc is dissolved, the mixture is diluted with water, and made strongly alkaline, until the hydroxide of zinc precipitated at first is redissolved; the aniline is then extracted with a little ether. Then proceed as just described.

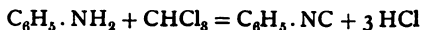
If it is desired to determine whether a given compound is nitrobenzene, it is at once reduced with zinc and hydrochloric acid.

(3) In a small porcelain dish place 5 drops of concentrated sulphuric acid, and with a glass rod add 1 drop of aniline. The aniline sulphate thus formed solidifies for the most part on the rod; remove it by rubbing it against the walls of the dish. Then add 4 drops of an aqueous solution of potassium dichromate, and mix the liquid by revolving the dish. After a short time the liquid assumes a beautiful blue colour. If the reaction does not take place, add 2 more drops of the dichromate, or heat a moment over a small flame.

(4) *Isonitrile Reaction*: Heat a piece of caustic potash the size of a bean with 5 c.c. of alcohol, pour off the solution from the undissolved residue into another test-tube; the warm solution is treated with 1 drop of aniline and 4 drops of chloroform. A reaction takes place immediately, or on gentle warming; this is recognised not only by the separation of potassium chloride, but by a most highly characteristic, disagreeable odour. The odour becomes more pronounced on pouring off the liquid and adding some cold water to the tube. If the vapours of the isonitrile are inhaled through the mouth, a peculiar sweet taste is noticed in the throat.

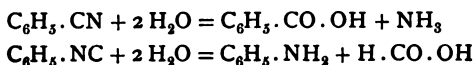
The reaction must be carried out under a hood with a good draught.

While the two colour reactions with bleaching powder and chromic acid are used especially for the recognition of aniline, the isonitrile reaction will show the presence of any primary amine of the aliphatic or aromatic series. The reaction takes place in accordance with the following equation:



For the elimination of hydrochloric acid, caustic potash is added. Since all isonitriles or carbylamines possess a characteristic odour, on the one hand the smallest quantity of a primary base may be detected by this reaction, and on the other a base may be shown to be primary. Secondary and tertiary bases do not give the reaction.

In the isonitriles it is very probable that the carbon atom combined with the nitrogen atom is only bivalent: $C_6H_5 \cdot N=C$. The isonitriles are isomeric with the acid-nitriles, *e.g.*, $C_6H_5 \cdot C \equiv N$, benzonitrile. While the nitriles on saponification give acids, the isonitriles decompose into a primary amine and formic acid:



3. REACTION: (a) REDUCTION OF A NITRO-COMPOUND TO A HYDROXYLAMINE DERIVATIVE. (b) OXIDATION OF A HYDROXYLAMINE DERIVATIVE TO A NITROSO-COMPOUND

EXAMPLES: (a) Phenylhydroxylamine from Nitrobenzene
(b) Nitrosobenzene from Phenylhydroxylamine

(a) *Phenylhydroxylamine*: In a thick-walled $\frac{1}{2}$ -litre battery jar treat a solution of 5 grammes of ammonium chloride in 160 c.c. of water with 10 grammes freshly distilled nitrobenzene. In the course of an hour add, with constant stirring, 15 grammes of zinc dust. The jar containing the liquid is surrounded with water and kept at a temperature of 13° (thermometer in water; small pieces of ice are used if necessary). In order to secure an intimate mixture, the zinc dust is divided into four equal portions, and each portion is added in the course of a quarter of an hour. After the addition of the last portion the stirring is continued for 10 minutes; it is then filtered, using suction and a Büchner funnel, from the zinc oxide; the filtrate (solution I) is poured into a beaker, and the zinc oxide deposit on the funnel is washed with 200 c.c. of water at 45° ; before the water is poured on the residue, the suction is disconnected from the funnel, and is only attached later to draw the liquid through gently, drop by drop. The residue is then pressed together and filtered with strong suction (solution II).

The two water solutions are separately saturated (with stirring) with finely pulverised salt; for solution I, about 45 grammes, and for solution II, about 60 grammes of salt will be required. They are cooled in ice for 15 minutes to 0° . The colourless crystals separating out are filtered off with suction, and, without washing, are pressed out on a porous plate. Yield, almost quantitative.

A small test-portion of the crude product is recrystallised from benzene. Melting-point, 81° . The remainder, without further purification, is worked up into nitrosobenzene.

The success of the reaction depends essentially upon the quality of the zinc dust used. It is therefore necessary to make a zinc dust determination (see page 390), and then use about 10% more than is required by the theory. Zinc dust of 75% is referred to above.

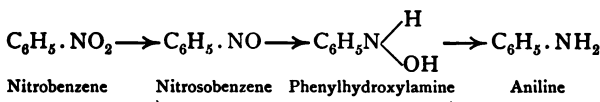
Precautions: In the preparation of phenylhydroxylamine, care is taken to prevent it, and particularly a warm solution of it, from coming in contact with the skin, since it causes very painful and annoying inflammation. Even in pressing it out or pulverising it under the hood, care must be taken not to breathe in any of the dust, since it causes extraordinarily violent attacks of sneezing.

(b) *Nitrosobenzene:* To a solution of 30 grammes of concentrated sulphuric acid in 270 c.c. of water, well cooled by ice water, add 4 grammes of freshly prepared and finely pulverised phenylhydroxylamine; the solution is then quickly treated with an ice-cold solution of 4.6 grammes of potassium dichromate in 200 c.c. of water; the pure nitrosobenzene separates out immediately in crystals.

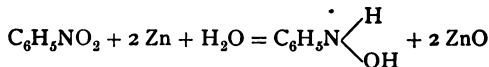
"On account of the splendid phenomena, steam should be passed into the liquid containing the oxidation product; the total quantity of nitrosobenzene is carried over in 4–5 minutes. At the beginning of the heating the walls and neck of the flask take on a deep green colour, and soon the nitrosobenzene sublimes in white, lustrous plates in the bent tube entering the condenser; a few moments later beautiful emerald-green oil drops appear which solidify so completely, in the lower part of the condenser, to snow-white crystals, that the distillate presents the appearance of a faintly green liquid containing only a few minute crystals. The

crystals of nitrosobenzene are pushed out of the condenser with a glass rod, the end being covered with a cotton plug, spread out on a porous plate and washed upon the plate with ligroin (boiling-point 40–70°). Melting-point 67.5–68°."

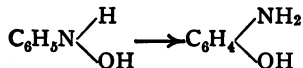
(a) The primary amines discussed in the preceding reaction are the lowest reduction products of nitro-compounds. Recently two classes of compounds have been discovered which appear to be intermediate products between the nitro-compounds and amines. In order to distinguish them from the compounds referred to in the next preparation, they may be called "monomolecular intermediate reduction products."



Phenylhydroxylamine¹ was obtained simultaneously by Bamberger and Wohl by the reduction of nitrobenzene with zinc dust in a neutral solution:



The presence of certain salts, *e.g.*, calcium or ammonium chloride, promotes the reaction. Nitro-compounds may also be reduced into hydroxylamine derivatives by the action of ammonium sulphide.² Phenylhydroxylamine acts like a base towards acids. If it be warmed with mineral acids, it undergoes a noteworthy transformation into paraamidophenol:



This behaviour explains the electrolytic reduction of aromatic nitro-compounds.³

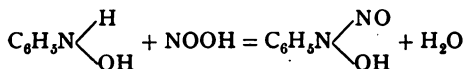
If a nitro-compound dissolved in concentrated sulphuric acid is subjected to electrolytic reduction, not only is the nitro-group reduced to the amido-group, but a hydroxyl group enters the para position (to the amido-group) if it is vacant. Thus from nitrobenzene p-amidophenol is obtained. In accordance with our present knowledge the reaction is

¹ B. 27, 1347, 1432, 1548; 28, 245, 1218. ³ B. 26, 1844, 2810; 27, 1927; 29, 3040

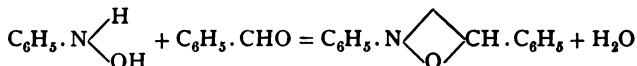
² B. 41, 1936.

no longer considered remarkable. Phenylhydroxylamine is first formed, which immediately undergoes a molecular transformation into the amidophenol.

Phenylhydroxylamine is a strong reducing agent, which reduces Fehling's solution and an ammoniacal solution of silver nitrate even in the cold. With nitrous acid it forms a nitroso-derivative:

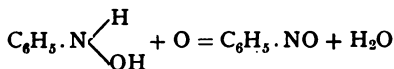


With aldehydes it reacts thus:



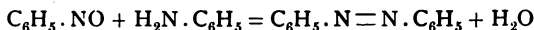
By the oxygen of the air it is oxidised to azoxybenzene; more energetic oxidising agents convert it into nitrosobenzene.

(b) Nitrosohydrocarbons may best be obtained by the oxidation of hydroxylamine derivatives:

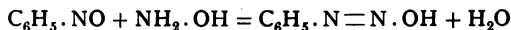


The nitrosohydrocarbons in the solid state form colourless crystals, but when fused or in solution an emerald-green liquid. They possess a characteristic piercing odour which suggests quinone and the mustard oils; they are extremely volatile. On reduction they go over into amines.

With primary amines they combine to form an azo-compound, *e.g.*,



Combined with hydroxylamine they form isodiazoo-compounds, *e.g.*,



4. REACTION: REDUCTION OF A NITRO-COMPOUND TO AN AZOXY-, AZO-, AND HYDRAZO-COMPOUND

EXAMPLES: Azoxybenzene, Azobenzene, Hydrazobenzene

(1) *Azoxybenzene*.¹ To 200 grammes of methyl alcohol contained in a 2-litre flask provided with a wide reflux condenser, 20 grammes of sodium in pieces the size of a bean are gradually

¹ J. pr. 36, 93; B. 15, 865.

added ; the flask is not cooled (heat being generated by the reaction). Since methyl alcohol frequently contains much water, the first portions of the sodium must not be added too rapidly. When the metal is dissolved, 30 grammes of nitrobenzene are added, and the mixture heated for 3 hours on an actively boiling water-bath (reflux condenser). Crystals of sodium formate soon begin to separate out ; this often causes a troublesome bumping. The greater portion of the methyl alcohol is then distilled off (the flask being placed *in* the water-bath ; silk thread). The residue is treated with water, and the reaction-mixture poured into a beaker. After long standing, especially in a cool place, the oil at the bottom solidifies to a bright yellow crystalline mass, which is separated from the liquid by decanting the latter ; it is washed several times with water and finally pressed out on a porous plate. If the azoxybenzene does not solidify, the main quantity of the water solution is poured off and the oil treated with small pieces of ice. If solidification does not take place now, it is due to the presence of nitrobenzene ; this is distilled off with steam, and the difficultly volatile residue, after it has cooled, is further cooled with ice. From methyl alcohol (use 3 c.c. of the alcohol for every gramme of the substance) the azoxybenzene crystallises in bright yellow needles, melting at 36°. Yield, 20–22 grammes.

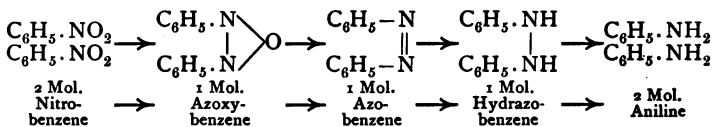
(2) *Azobenzene*.¹ Five grammes of crystallised azoxybenzene, dried completely by heating on a water-bath for an hour, are finely pulverised and intimately mixed in a mortar with 15 grammes of *coarse* iron filings, which must also be completely dry ; the mixture is distilled from a small retort, not tubulated. It is first warmed with a small luminous flame kept in constant motion ; the size of the flame is increased after some time ; finally the last portions are distilled over with a non-luminous flame. If, on heating, a sudden but harmless explosion should occur, it is due to the fact that the substances were not dry ; the experiment must be repeated. The reddish distillate is collected in a small beaker, and, after it has solidified, is washed with hydrochloric acid to remove

¹ A. 12, 311 ; 207, 329.

the aniline, then with water, and pressed out on a porous plate. The experiment is repeated a second time with a fresh quantity of azoxybenzene; by working carefully the same retort can be used again. The two pressed-out crude products are united. Azobenzene crystallises from ligroïn, after a partial evaporation of the solvent, in the form of coarse red crystals melting at 68° .

(3) *Hydrazobenzene*:¹ Dissolve 5 grammes of azobenzene in 50 grammes of alcohol (about 95%) in a flask provided with a reflux condenser, and treat with a solution of 2 grammes of caustic soda in 4 grammes of water. To the boiling solution gradually add zinc dust in small portions (best by occasionally removing the cork) until the orange-coloured solution becomes colourless: about 8 grammes of zinc dust will be necessary. The hot solution is then filtered with suction (Büchner funnel) from the excess of zinc; 20 c.c. of a water solution of sulphur dioxide and 100 c.c. of water are previously placed in the filter-flask. The hydrazobenzene precipitating out of the alcoholic solution is quickly filtered, washed with water containing sulphur dioxide, and pressed out on a porous plate. By crystallising from ligroïn it is obtained pure. Melting-point, 126° . Yield, 80–90% of the theory.

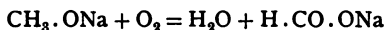
Under the influence of suitable reducing agents, nitro-compounds undergo a partial reduction in such a way that two molecules enter into combination. There are thus obtained first the azoxy-, then the azo-, and finally the hydrazo-compound, which in order to distinguish them from the compounds obtained in Reaction 3 may be called "dimolecular intermediate reduction products."



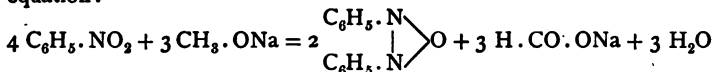
In order to reduce a nitro-compound to an azoxy-compound, either sodium amalgam or alcoholic-caustic potash or caustic soda is used. With nitrobenzene, particularly, the reaction takes place most surely by dissolving sodium in methyl alcohol as above. The reducing action of sodium methylate depends upon the fact that it is oxidised to

¹ Z. 1868, 437.

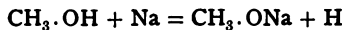
sodium formate, two hydrogen atoms of the methyl group being replaced by one atom of oxygen :



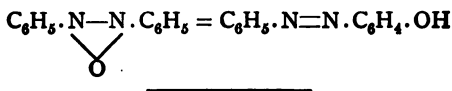
In the operation carried out above the reaction is expressed by the equation :



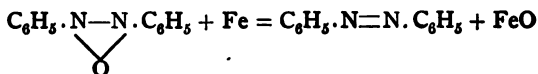
A few words may be said here concerning the relatively weak reducing power of *previously prepared* alcoholates, in comparison with the extremely energetic action of a mixture of undissolved sodium and alcohol. While the previously prepared alcoholates can generally only abstract oxygen, the mixture just referred to belongs to the class of very strong reducing agents. With the aid of this, it is possible to break up the double or centric union of the benzene ring, and thus prepare hydrogen derivatives of benzene. In this case the alcoholate does not act as a reducing agent as above, but the hydrogen effects the reduction :



The azoxy-compounds are yellow- to orange-red crystallisable substances, which, like the nitro-compounds, are of an indifferent character; but they are not volatile with steam, and cannot be distilled without undergoing decomposition. On reduction they yield first the azo-compounds, then the hydrazo-compounds, and finally two molecules of a primary amine. By heating with sulphuric acid, azoxybenzene is converted into its isomer oxyazobenzene :

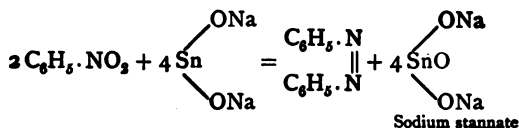


If an azoxy-compound is distilled carefully over iron filings, its oxygen atom is removed, and an azo-compound is formed :

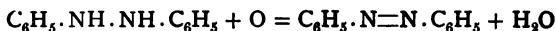


Azo-compounds may also be obtained directly from nitro-compounds, since they are reduced by sodium amalgam, or, in an alkaline solution, by

zinc dust or stannous chloride (sodium stannous oxide). The latter reducing agent acts in accordance with this equation :



Azo-compounds may also be obtained by the oxidation of hydrazo-compounds :



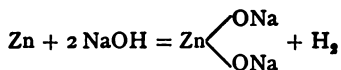
The azo-hydrocarbons are orange-red to red crystalline substances which can be distilled without decomposition, differing in this respect from the azoxy-compounds.

EXPERIMENT : A few crystals of azobenzene are heated in a test-tube to boiling, over a free flame. A red vapour is evolved, which again condenses to crystals on cooling.

The azo-compounds thus differ in their stability from the very easily decomposable diazo-compounds, which also contain the group $\text{N}=\text{N}$, but it is in a different combination.

By the reduction of an azo-compound, a hydrazo-compound is first formed and finally an amine.

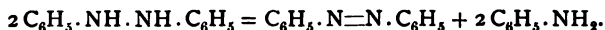
The hydrazo-compounds are formed by the reduction of azo-compounds with ammonium sulphide or zinc dust and an alkali. Zinc dust with caustic soda acts as follows :



They may also be formed on the direct reduction of nitro-compounds in alcoholic solution by zinc dust and an alkali; this method is used practically on the large scale.

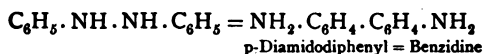
The hydrazo-compounds, in contrast with the azoxy-, and especially with the intensely coloured azo-compounds, are colourless. They are derived from hydrazine, NH_2-NH_2 , in which one hydrogen atom of the two amido-groups has been replaced by a hydrocarbon radical. The

basic character of hydrazine is so weakened by the presence of the negative hydrocarbon residues, that the hydrazo-compounds no longer possess a basic character. On oxidation hydrazo-compounds pass over to azo-compounds, a reaction which takes place slowly but completely, under the influence of the oxygen of the air. The hydrazo-compounds decompose, on heating, into azo-compounds and primary amines.



EXPERIMENT: A few crystals of hydrazobenzene are heated in a small test-tube to boiling; the colourless compound becomes red, azobenzene being formed. In order to show the presence of aniline, after cooling, the substance is shaken with water and the bleaching-powder test applied.

If the hydrazo-compounds are treated with concentrated acids like hydrochloric or sulphuric acids, they are converted into derivatives of diphenyl:¹

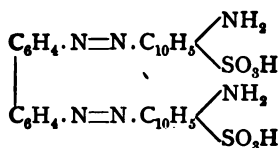


The molecular transformation takes place essentially in para position to the imide (NH) groups.

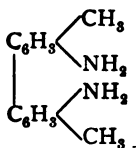
EXPERIMENT: Hydrazobenzene is covered with concentrated hydrochloric acid, and allowed to stand for about 5 minutes. It is then treated with water, and half the solution is made alkaline with caustic soda: the free benzidine is extracted several times with ether, the ether evaporated, and the substance crystallised from hot water. Leaflets of a silvery lustre are obtained. Melting-point, 122°. The other half of the solution is treated with dilute sulphuric acid, upon which the difficultly soluble benzidine sulphate separates out.

Benzidine differs from hydrazobenzene, in that it is a strong, di-acid primary base. It is prepared technically, since the azo dyes derived from it possess the important property of colouring unmordanted cotton fibre directly; for most azo dyes the cotton must first be mordanted. The first representative of these dyes made was Congo Red, prepared from the bisdiazo-compound of benzidine and naphthionic acid. In consequence, the entire class of these dyes is called the "Congo Dyes."

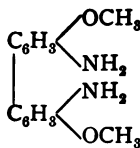
¹ J. pr. 36, 93; J. 1863, 424.



In a wholly analogous manner, from *o*-nitrotoluene and *o*-nitroanisol are prepared *o*-tolidine and dianisidine, respectively.



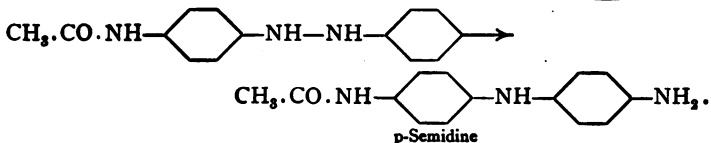
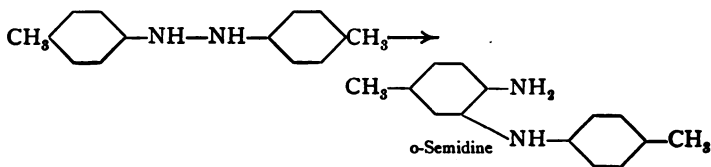
Tolidine



Dianisidine

If, in hydrazo-compounds, the *para* position to the imido (NH) group is occupied as, *e.g.*, in *p*-hydrazotoluene, then the benzidine transformation cannot occur.

In such cases, derivatives of *o*- and *p*-amidodiphenyl amine are formed through the so-called "Semidine transformation."¹



5. REACTION: PREPARATION OF A THIOUREA AND A MUSTARD OIL FROM CARBON DISULPHIDE AND A PRIMARY AMINE

EXAMPLE: Thiocarbanilide and Phenyl Mustard Oil from Carbon Disulphide and Aniline

Thiocarbanilide: A mixture of 40 grammes of aniline, 50 grammes of carbon disulphide, 50 grammes of alcohol, and

¹ B. 26, 681, 688, 699; A. 287, 97.

10 grammes of finely pulverised potassium hydroxide is gently boiled for 3 hours on a water-bath in a flask provided with a long reflux condenser. The excess of carbon disulphide and alcohol is then distilled off, the residue treated with water, the crystals separating out are filtered off, and washed first with water, then with dilute hydrochloric acid, and finally with water. For the preparation of phenyl mustard oil, the crude product is used directly, after it has been dried on the water-bath. In order to obtain pure thiocarbanilide, 2 grammes of the dried crude product are recrystallised from alcohol. Large, colourless tablets are thus obtained, which melt at 154° . Yield, 30–35 grammes. If a mixture of equal parts, by weight, of aniline, carbon disulphide, and alcohol (40 grammes of each) placed in a flask provided with a reflux condenser is heated, with the addition of 0.3 gramme of crystallised sulphur,¹ to gentle boiling on the water-bath for 6 hours, a better, almost quantitative yield of thiocarbanilide is obtained, although a longer time is required. After the heating, proceed as above.

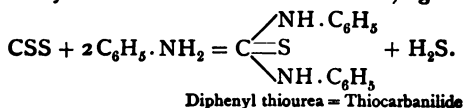
Phenyl Mustard Oil:² In a flask of about 400 c.c. capacity place 30 grammes of the crude thiocarbanilide, and treat with 120 grammes of concentrated hydrochloric acid; the mixture is distilled by heating to the boiling-point of the acid, on a sand-bath, with a large flame under a hood. When only about 20 c.c. of the liquid remain in the flask, the distillation is discontinued. The distillate is treated with an equal volume of water, the mustard oil separated in a dropping funnel, dried with a little calcium chloride, and distilled. Boiling-point, 222° . Yield, almost quantitative.

Triphenyl Guanidine: The residue remaining in the flask after the distillation with hydrochloric acid is treated with 100 c.c. of water, and then allowed to stand for several hours, when colourless crystals of triphenylguanidine hydrochloride separate out. These are filtered off, and warmed with some dilute caustic soda solution. The salt is decomposed, and the free base obtained, which on recrystallising from alcohol forms colourless crystals. Melting-point, 143° .

¹ B. 32, 2245.

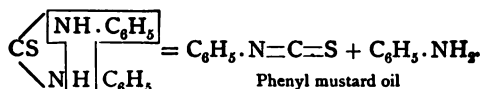
² B. 15, 986. Z. 1869, 589.

Carbon disulphide acts upon primary aromatic amines (NH_2 in the nucleus) to form symmetrical disubstituted thioureas, *e.g.*

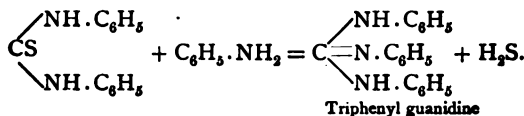


By the addition of caustic potash the elimination of hydrogen sulphide is facilitated, so that the reaction takes place in a shorter time than without the use of the alkali.

From the thioureas thus obtained the mustard oils may be prepared by heating with acids, as hydrochloric acid, sulphuric acid, phosphoric acid. The reaction takes place in accordance with the following equation:



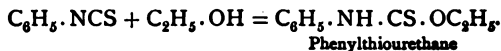
The primary amine formed in addition to the mustard oil combines with the acid. Besides this reaction a second one takes place, *viz.* : the amine formed acts upon some still undecomposed thiourea, resulting in the formation of a guanidine derivative:



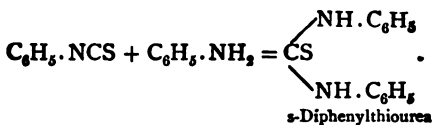
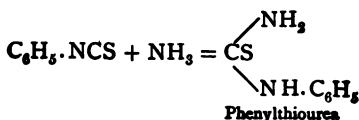
Since guanidine $\begin{array}{c} \text{NH}_2 \\ \diagup \\ \text{C} = \text{NH} \\ \diagdown \\ \text{NH}_2 \end{array}$ is an extremely strong base, which, like

caustic potash and caustic soda, absorbs carbon dioxide from the air, the introduction of the three negative phenyl groups in the above compound has not neutralised the basic properties entirely, and it still has the power to form salts.

The aromatic mustard oils are in part colourless liquids, in part crystallisable solids, the lower members are easily volatile with steam, and possess a characteristic odour. In chemical behaviour they are very active. If they are warmed for a long time with an alcohol, they combine with the alcohol, addition taking place, and a thiourethane is formed:

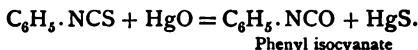


In the same way, ammonia and primary bases are added with the formation of a thiourea:



EXPERIMENT: Treat 2 drops of phenyl mustard oil on a watch-glass with 2 drops of aniline, and warm gently over a small flame. On stirring the reaction-product after cooling, with a glass rod, the thiocarbonyl will solidify in crystals, from which in the above reverse reaction the mustard oil itself was prepared.

By heating with yellow mercuric oxide, the sulphur is replaced by oxygen, and an isocyanate is formed, which may be easily recognised by its extremely disagreeable odour:



EXPERIMENT: Heat $\frac{1}{2}$ c.c. of phenyl mustard oil in a test-tube with the same volume of yellow mercuric oxide for some time, until the oil boils. The yellow oxide is changed to the black sulphide, at the same time the extremely disagreeable odour of the phenyl isocyanate arises; the vapour of the compound attacks the eyes, causing tears.

6. REACTION: THE SULPHONATION OF AN AMINE

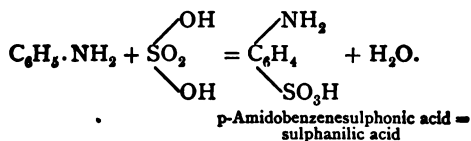
EXAMPLE: Sulphanilic Acid from Aniline and Sulphuric Acid¹

To 100 grammes of pure concentrated sulphuric acid in a dry flask, 30 grammes of freshly distilled aniline are added gradually, with shaking; the mixture is heated in an oil-bath up to 180–190°, until, from a test-portion diluted with water and treated

¹ A. 60, 312; 100, 163; 120, 132.

with caustic soda, no aniline separates out: about 4-5 hours' heating will be necessary. The cooled reaction-mixture is poured, with stirring, into cold water, upon which the sulphanilic acid separates out in crystals. It is filtered off, washed with water, and recrystallised from water, with the addition of animal charcoal, if necessary. Yield, 30-35 grammes.

When an aromatic compound is treated with sulphuric acid, a portion of the benzene-hydrogen is replaced by a sulphonic acid group, the reaction taking place in accordance with the equation below. The aliphatic compounds do not react in a similar manner. Under the preparation of benzene sulphonic acid, the details of the reaction will be discussed.



In the above example, it happens, as in many cases, that the sulphonic acid group enters in the para-position to the amido-(NH₂) group. The amido sulphonic acids are colourless crystallisable compounds melting with decomposition; they possess acid properties, *i.e.* in dissolving in alkalis. The basic character of the amine is so greatly weakened by the introduction of the negative sulphonic acid group that the amido sulphonic acids cannot form salts with acids. They differ

in this from the analogous carbonic acids $\left(\text{e.g., } \text{C}_6\text{H}_4 \begin{array}{c} \text{NH}_2 \\ \diagup \quad \diagdown \\ \text{CO}\cdot\text{OH} \end{array} \right)$, which

Amidobenzoic acid

dissolve in both acids and alkalis.

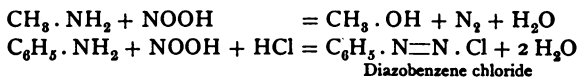
The amido-sulphonic acids, since they are derivatives of a primary amine, may like them be diazotised by the action of nitrous acid; upon this fact depends their great technical importance. If the diazo-compounds thus obtained are combined with amines or phenols, azo dyes are formed which contain the sulphonic acid group, and in the form of their alkali salts are soluble in water. Sulphanilic acid particularly, and its isomer, metanilic acid, obtained by the reduction of m-nitrobenzenesulphonic acid, as well as the numerous mono- and poly-sulphonic acids derived from α and β naphthyl amines, find extensive technical application in the manufacture of azo dyes.

7. REACTION: REPLACEMENT OF THE AMIDO- AND DIAZ-GROUPS BY HYDROGEN

EXAMPLE: Benzene from Aniline

Dissolve 5 grammes of aniline in a mixture of 15 grammes of concentrated hydrochloric acid and 30 c.c. of water; cool with ice, and treat with a solution of 5 grammes of sodium nitrite in 15 c.c. of water, until free nitrous acid may be recognised with starch-potassium-iodide paper. The diazobenzenechloride solution thus obtained is allowed to flow carefully into a solution of 10 grammes caustic soda in 30 c.c. of water contained in a 400 c.c. flask which is *well cooled* with ice. Further, dissolve 20 grammes of stannous chloride in 50 c.c. of water, and treat this solution with a concentrated solution of sodium hydroxide (2 parts to 3 of water), until the precipitate at first formed (stannous oxide) is redissolved in the excess of the alkali. Treat the alkaline diazobenzene solution, *well cooled with ice-water*, gradually with small portions of the sodium-stannous oxide solution, previously well cooled, waiting after each addition until the lively evolution of nitrogen has ceased before adding more. When all the reducing liquid has been added, the flask is connected with a condenser, and the liquid heated to boiling. The benzene formed passes over first, and is collected in a test-tube. By a careful distillation from a small fractionating flask (without condenser), it is obtained perfectly pure. Boiling-point, 81°. Yield, 3-4 grammes.

As already mentioned, under the preparation of methyl amine, the behaviour of the aliphatic primary amines toward nitrous acid is very different from that of the aromatic compounds. While the former yield alcohols with the elimination of nitrogen, the latter, in a mineral acid solution, under the influence of nitrous acid, yield diazo-compounds, discovered by Peter Griess,¹ in the form of their mineral acid salts.

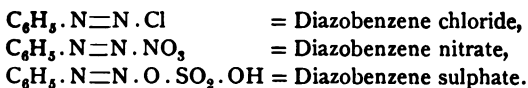


It has been held that the mother substance of the diazo-compounds, — free diazobenzene — possessed the constitution:

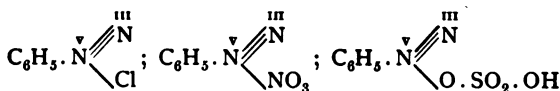


¹ A. 137, 39.

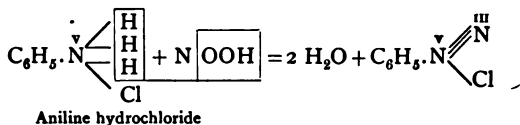
In accordance with which the diazo salts were expressed by :



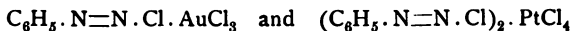
More recently this view has been abandoned, and the one proposed earlier by Blomstrand taken up. It is, however, not accepted generally. According to this conception the above salts are represented thus :



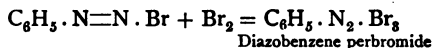
In accordance with this view the diazotising process consists in replacing the three hydrogen atoms combined with a nitrogen atom having a valence of 5, by a trivalent nitrogen atom :



To emphasise the similarity to ammonium compounds, in which the valence of the nitrogen atom is probably 5, the diazo salts are called diazonium salts. The diazo-compounds can also form double salts, *e.g.* :

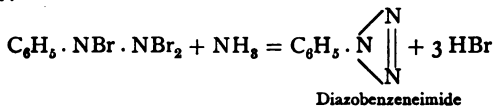


Diazobromides have the power of taking up two atoms of bromine to form perbromides :



EXPERIMENT : Dissolve 1 c.c. of aniline in an excess of hydrochloric acid, diazotise as above, and add 1 c.c. of bromine dissolved in a water solution of hydrobromic acid, or in a concentrated solution of potassium bromide. A dark oil separates out, from which the solution is decanted. It is washed several times with water ; on cooling, it solidifies to crystals.

If ammonia is allowed to act on the perbromide, diazobenzeneimide is obtained :



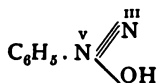
EXPERIMENT: The perbromide just obtained is covered with water, and concentrated ammonium hydroxide added to it. A vigorous reaction takes place with the formation of an oil possessing a strong odour (diazobenzeneimide).

Under the influence of alkalis the diazonium compounds yield salts, thus acting like acids.

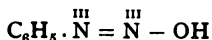


These can exist in *two isomeric* modifications. The one primarily obtained is characterised by the fact that in alkaline solution it unites with phenols to form azo dyes; while the second modification obtained by a longer action of the alkali, at higher temperature if necessary, does not possess this property at all, or only in a slight degree. If they (the latter) be treated with acids, they are converted back into the diazonium salts, and now have the property of combining with phenols in alkaline solutions.

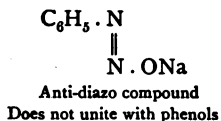
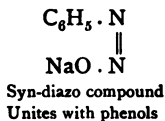
The view of Hantzsch is that the salts in which the diazonium compound behaves as an *acid*, are not derived from diazonium hydroxide, *e.g.*:



but they are derived from a compound having the following constitution:

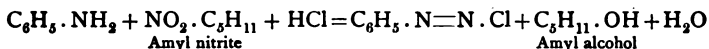


Consequently, for *metallic salts*, the first formula must be modified. The differences underlying the constitution of the two metallic salts are due to stereoisomerism, *e.g.*:



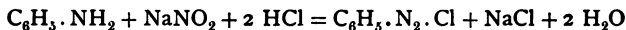
The principles underlying the space arrangement of the three valences of nitrogen, as illustrated in these examples, will be discussed under Benzophenoxim. Compare Hantzsch: "Die Diazoverbindungen" (Stuttgart, 1902), also Hantzsch: "Stereochemie" (2 Ed.), p. 142. The salts of the diazo-compounds formed with acids are, in most cases, colourless, crystallisable substances, easily soluble in water, insoluble in ether.

In order to prepare them in the solid condition, various methods may be used. Thus, *e.g.*, the very explosive diazobenzene-nitrate may be obtained in colourless needles by conducting gaseous nitrous acid into a well-cooled pasty mass of aniline nitrate and water, and treating the diazo-solution with alcohol and ether. In general, the solid diazo-salts may be prepared by adding to an alcoholic solution of the amine that acid the salt of which is desired, and then treating the well-cooled mixture with amyl nitrite: ¹



If the solid diazo-compound does not separate out at once, ether is added. On heating, the dry diazo-salts decompose either, as in the case of diazobenzene nitrate, with explosion, or a sudden evolution of gas takes place without detonation. A few diazo-compounds are so stable that they may be recrystallised from water.

In rare cases only, in working with diazo-compounds, is it necessary to isolate them in a pure condition; generally, the very easily prepared water solutions are used. These compounds were formerly obtained by passing gaseous nitrous acid into a salt of the amine until it was diazotised. But at present this method is employed only in rare cases; the free nitrous acid obtained from sodium nitrite is used. In order to diazotise an amine, a solution of it in a dilute acid—most frequently hydrochloric acid or sulphuric acid—is first prepared. Theoretically, two molecules of a monobasic acid are required to diazotise one molecule of a monamine:



but an excess is always taken, — not less than three molecules of hydrochloric acid or two of sulphuric acid to one molecule of a monamine. In many cases, the hydrochloride or sulphate of the amine is difficultly soluble in water. Under these conditions, it is not necessary to add water until the salt is entirely dissolved, but the solution of the nitrite may be poured into the pasty mass of crystals; when the undissolved salt is diazotised, it passes into solution. For the diazotisation of one molecule of a monamine, one molecule of sodium nitrite is necessary, theoretically; but since the commercial salt is never perfectly pure, it is advisable to weigh off from 5–10 % more than the calculated amount, and to determine by the method given below when a sufficient quantity

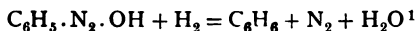
¹ B. 23, 2994.

of this has been added. The nitrite is dissolved in water, generally 5-10 parts of water to 1 part of salt. The nitrite solution must be added gradually to the amine solution, and the liquid must not be allowed to become warm. In many cases, the experimenter is often too careful, in that he cools the amine solution with a freezing mixture, and adds the nitrite solution drop by drop from a separating funnel. Frequently it is sufficient to place the solution in a water-bath filled with cold water, or ice is thrown into the water, or the amine solution is cooled by ice. It is very convenient to cool the solution, not from without, but by throwing into it from time to time small pieces of ice. The nitrite solution may be poured directly from a flask. If the addition causes evolution of gas bubbles or vapours of nitrous acid, the temperature of the solution must be lowered and the nitrite added more slowly. In order to be cognisant of the course of the reaction, as well as to be able to determine when it is completed, starch-potassium-iodide paper, prepared as follows, is used :

A piece of starch the size of a pea is finely pulverised, and added to 200 c.c. of boiling water; it is boiled a short time, with stirring. After cooling, a solution of a crystal of potassium iodide the size of a lentil, in a little water, is added to it. With this mixture, saturate long strips of filter-paper 3 cm. wide; the strips are dried by suspending them from a string in a place free from acids. After drying, the strips are cut up and preserved in a closed vessel.

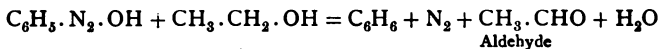
In order now to diazotise an amine, the cooled solution is first treated with a small portion of the nitrite solution; it is well stirred, and a drop of it transferred with a clean glass rod to the starch-potassium-iodide paper. If the nitrous acid is already used up in the diazotisation, no dark spots appear, and further portions of the sodium nitrite may be added, the test is again repeated, and so on. But if a dark spot is formed at once, the nitrous acid is still present; and in this case, before more of the nitrite is added, one waits until the reaction has been completed, and so on. After the addition of three-fourths of the nitrite solution, larger quantities may be added at one time, but toward the end of the reaction small quantities must again be employed. The diazotisation is ended when, after standing some time, the mixture shows the presence of nitrous acid. Since the diazotisation of the last portions of the amine often requires some time, the addition of the nitrite is not discontinued at once, even if after one minute the test for nitrous acid is obtained, but the solution is allowed to stand 5-10 minutes, and is then tested again. At times, it happens that the

weighed-off quantity of sodium nitrite is apparently not sufficient to complete the diazotisation, and that even after the addition of a fresh quantity, the test will not show the presence of nitrous acid. This phenomenon has its cause generally in the fact that the acid (hydrochloric or sulphuric) has been used up, and consequently the nitrite cannot enter into the reaction. Thus, in case the weighed-off amount of sodium nitrite is not sufficient, some acid is first added to a small portion of the liquid, and this is then tested to determine whether the desired reaction has taken place. Further, often the diazo-solution becomes cloudy toward the end of the reaction, or a precipitate separates out. This is the diazoamido-compound; its formation is also caused by the lack of free acid. On the addition of acid and solution of the nitrite, the precipitate disappears. The replacement of the diazo-group by hydrogen in the above reaction takes place in accordance with the following equation :



In this way it is possible in many cases to replace a primary amido-group by hydrogen. Obviously, such a reaction is superfluous, if, as in the above case, the amine is obtained by the nitration of the hydrocarbon and the reduction of the nitro-compound. But there are cases in which an amine is not obtained in this way, and where it is of importance to prepare the amido-free compound (see below).

The replacement of a diazo-group by hydrogen may be effected by other reducing agents. If, *e.g.*, a diazo-compound is boiled with alcohol, the latter is converted into aldehyde, thus liberating two hydrogen atoms, by which the diazo-compound is reduced :



The reaction is effected either by conducting gaseous nitrous acid into the boiling alcohol solution of the amine, or by heating the amine with alcohol saturated with ethyl nitrite ; or the boiling alcohol solution of the amine, acidified with sulphuric acid, may be treated with sodium nitrite.

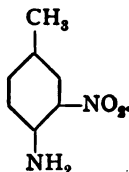
At this place, two examples may be mentioned which illustrate the theoretical as well as the practical value of the reaction : by the oxidation of a mixture of aniline and p-toluidine, there is formed a complex dye, para-fuchsine, the constitution of which was unknown for a long time. This was first explained by E. and O. Fischer. They heated the diazo-compound of the leuco-base of the dye, paraleucaniline with

¹ B. 22, 587.

alcohol, which gave the mother substance — the hydrocarbon triphenyl methane (A. 194, 270).

As an example of the preparation value of the reaction, the following case is cited :

No method is known by which m-nitrotoluene can be prepared on a large scale by the nitration of toluene; this results in the formation of the o- and p-compounds mainly. In order to obtain the m-nitrotoluene, the starting-point is p-toluidine. This is nitrated, upon which a nitro-toluidine of the following constitution is obtained :



If the amido-group is replaced by hydrogen, using the method last described, the desired m-nitrotoluene is obtained.

By boiling a diazo-compound with alcohol the reaction may take place in a different way; at times the diazo-group is not replaced by hydrogen, but by the ethoxy ($-\text{OC}_2\text{H}_5$) group, thus giving rise to a phenol ether.



In conclusion, special attention is called to the fact that not only aniline and its homologues can be diazotised, but all the derivatives of these, as the nitro-amines, halogen-substituted amines, amino-aldehydes, amino-carbonic acids, etc.

8. REACTION: REPLACEMENT OF THE DIAZO-GROUP BY HYDROXYL

EXAMPLE : Phenol from Aniline

Pour 20 grammes of concentrated sulphuric acid as rapidly as possible, with stirring, into 50 grammes of water; to the hot solution add 10 grammes of freshly distilled aniline, with stirring, by allowing it to flow down the side of the beaker, then add 100 c.c. of water. After the hot liquid has been cooled by immersion in cold water, it is treated with a solution of 8.5 grammes of sodium nitrite in 40 c.c. of water, until it shows a blue spot on starch-potassium-iodide paper. The diazobenzene sulphate solu-

tion thus obtained is gently heated (40–50°) for half an hour on the water-bath, the phenol is then distilled over with steam, and the distillate, after being saturated with salt, is extracted several times with ether. The ethereal solution is allowed to stand for some time over fused sodium sulphate. The ether is then evaporated, and the residue of phenol is subjected to distillation in a small flask. Boiling-point, 183°. Yield, 7–8 grammes.

The liquid remaining back in the flask after the steam distillation is filtered hot. On cooling, a small quantity of oxydiphenyl crystallises out.

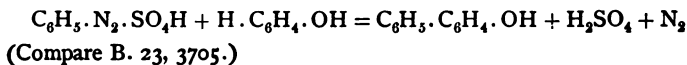
If a diazo-compound is heated with water, it will pass over to a phenol with the evolution of nitrogen, *e.g.* :



For this reaction the diazosulphate is most advantageously used. Under certain circumstances, the diazo-chloride may also be employed. But the use of the diazonitrate is avoided, since, in this case, the nitric acid liberated, acting upon the phenol, readily forms nitro-compounds. In many cases, it is more convenient not to isolate the diazo-compound, but to add a water solution of the calculated amount of sodium nitrite to a boiling solution of the amine in dilute sulphuric acid. The diazotisation of the substance and the immediate decomposition of the diazo-compound take place in one operation.

The same reactions are also applicable to substituted amines, like amido-carbonic acids, amido-sulphonic acids, halogen substituted amines, etc.

The oxydiphenyl obtained as a by-product is formed in consequence of the action of some of the undecomposed diazo-compound on phenol :



9. REACTION: REPLACEMENT OF A DIAZO-GROUP BY IODINE

EXAMPLE: Phenyl Iodide from Aniline

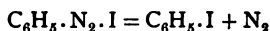
(*Phenyliodidechloride, Iodoso-benzene, Phenyl iodite, and Diphenyliodonium iodide*)

A solution of 10 grammes of aniline ¹ in a mixture of 50 grammes of concentrated hydrochloric acid and 150 grammes of water

¹ Twenty grammes of aniline should be used if the Grignard reaction is to be carried out later on.

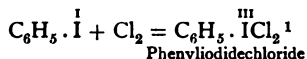
cooled with ice-water is gradually treated with a solution of 8.5 grammes of sodium nitrite in 40 c.c. of water, until a test will give a blue colour to the starch-potassium-iodide paper. The diazo-solution is then treated in a flask, not too small, with a solution of 25 grammes of potassium iodide in 50 c.c. of water, the mixture is allowed to stand several hours, being cooled by water, finally it is gently heated on the water-bath until the evolution of nitrogen ceases. The liquid is made strongly alkaline with caustic soda or caustic potash, and the iodobenzene distilled over with steam; the steam delivery tube should reach almost to the bottom of the flask. The iodobenzene is separated from the water in a dropping funnel, dried with calcium chloride and redistilled. Boiling-point, 189–190°. Yield, about 20 grammes.

If a diaziodide is heated, the diazo-group is replaced by iodine, the reaction taking place smoothly in most cases.



The reaction is effected by diazotising the amine in a hydrochloric acid or sulphuric acid solution, and then treating it with potassium iodide. From the diazochloride or diazosulphate there is formed a diaziodide, the reaction, in many cases, taking place at the ordinary temperature; in others, on heating, as above. Since the reaction occurs without difficulty, it is used as the method of preparation of many iodides.

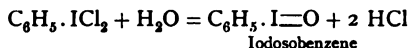
The aromatic iodides (iodine in the nucleus) possess the noteworthy property of combining with two atoms of chlorine, the iodine previously univalent becoming trivalent :



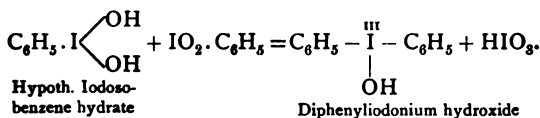
EXPERIMENT : A portion of the phenyl iodide obtained is dissolved in five times its volume of chloroform, the solution is cooled by ice water, and a current of dry chlorine is passed into it from a very wide delivery tube, until no more is absorbed. The crystals separating out are filtered off, washed with a fresh quantity of chloroform, spread out in a thin layer on a pad of filter-paper, and allowed to dry in the air.

¹ J. pr. 33, 154. B. 25, 3494; 26, 357; 25, 2632. Concerning aliphatic iodide-chlorides see A. 369, 119.

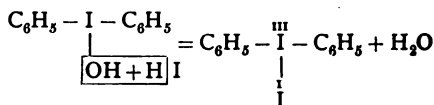
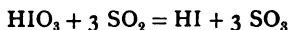
If caustic soda is allowed to act on an iodochloride, the two chlorine atoms are replaced by one oxygen atom, and an iodoso-compound is obtained :



Besides this reaction another takes place, resulting in the formation of an iodonium base. This formation is probably due to the fact that a small part of the iodosobenzene is oxidised to phenyl iodite, and this condenses with iodosobenzene, iodic acid being eliminated :



This base is present in the alkaline solution filtered off from the iodosobenzene. If the filtrate be treated with sulphur dioxide, this reduces the iodic acid to hydriodic acid, which, combining with the iodonium base, forms an iodide insoluble in cold water :



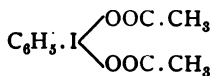
EXPERIMENT : The iodochloride is carefully triturated with dilute caustic soda in a mortar (for 1 gramme of the iodochloride, use a solution of 0.5 gramme sodium hydroxide in 4 grammes of water), and allowed to stand over night. The iodosobenzene is then filtered off, washed with water, and pressed out on a porous plate.

The alkaline filtrate is treated with a solution of sulphur dioxide until it smells strongly of it. The precipitate formed is filtered off and dissolved in hot water. On cooling, colourless needles of diphenyliodonium iodide are obtained.

The iodoso-compounds have the power of uniting with acids to form salts, in which they act like a di-acid base, *e.g.*, $\text{C}_6\text{H}_5 \cdot \text{I} \begin{array}{l} \text{OH} \\ \diagup \\ \diagdown \\ \text{OH} \end{array}$

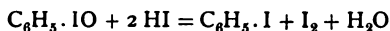
EXPERIMENT : Several grammes of iodosobenzene are dissolved with heat in as small a quantity of glacial acetic acid as possible ;

the solution is evaporated on the water-bath to dryness, in a watch-glass, or shallow dish. The solid residue is pulverised and recrystallised from a little benzene. Iodosobenzene acetate is thus obtained,



in the form of colourless prisms, melting at 157° .

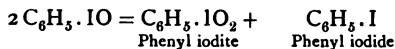
The iodoso-compounds, on treatment with hydriodic acid, are reduced to iodides, with a separation of iodine.



This reaction is used for the quantitative determination of iodoso-oxygen.

EXPERIMENT: Some potassium iodide is dissolved in water, acidified with dilute sulphuric acid, or acetic acid; and a few grains of iodosobenzene are added. The iodine separates out as a brown precipitate.

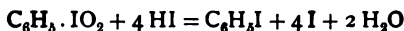
If an iodoso-compound is heated carefully to 100° , it passes over to an iodite (*Iodo-Verbindung*):



The same compound may also be obtained by treating an iodoso-compound with steam.

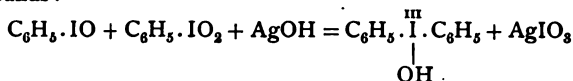
EXPERIMENT: Iodosobenzene is treated in a flask with enough water to form a thin paste. Into this steam is conducted (apparatus for distillation with steam), until no more phenyl iodide passes over with the steam and all the iodosobenzene has been dissolved. If the phenyliodite formed does not dissolve completely, water is added until solution takes place. The solution is then evaporated on the water-bath until a test-portion cooled off yields an abundant crystallisation of phenyl iodite.

The iodites, like the iodoso-compounds, puff up and suddenly decompose on heating. (Try it.) They also abstract iodine from hydriodic acid, and in double the quantity as compared to the similar action of the iodoso-compounds.

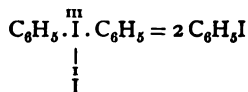


They do not form salts with acids.

The iodonium bases¹ are prepared most conveniently by the action of silver oxide on a mixture of equal molecules of iodoso and iodite compounds:



They are soluble in water, show a strong alkaline reaction like the sulphonium and ammonium compounds, and give with halogen hydrides precipitates of the corresponding salts. If the dried salts be heated, they decompose into two molecules of the hydrocarbon substitution product, *e.g.*:



EXPERIMENT: The diphenyliodonium iodide obtained as a by-product in the preparation of iodosobenzene is heated carefully in a test-tube over a small flame. Suddenly the substance begins to melt in one place; the fusion increases, without the need of further heating, and the whole mass boils up. Iodobenzene, easily recognised by its odour, is obtained.

10. REACTION: REPLACEMENT OF A DIAZO-GROUP BY CHLORINE, BROMINE, OR CYANOGEN

EXAMPLE: *p*-Tolyl Nitrile from *p*-Toluidine

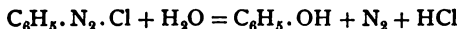
Dissolve 50 grammes of copper sulphate in 200 grammes of water in a 2-litre flask by heating on the water-bath; then add gradually, with continuous heating, a solution of 55 grammes of potassium cyanide in 100 c.c. of water. Since cyanogen is evolved, the reaction must be conducted under a hood, with a good draught, and the greatest care taken not to breathe the vapours.

While the cuprous cyanide solution is further gently heated up to about 60–70° on the water-bath, the diazotoluenechloride solu-

¹ B. 27, 426, 502, and 1592.

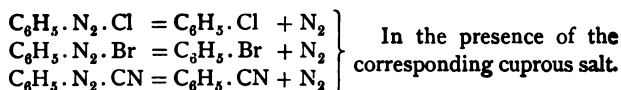
tion is prepared in the following way: 20 grammes of p-toluidine are heated with a mixture of 50 grammes of concentrated hydrochloric acid and 150 c.c. of water until solution takes place; the liquid is then quickly immersed in cold water and vigorously stirred with a glass rod, in order that the toluidine hydrochloride may separate out in as small crystals as possible. A solution of 16 grammes of sodium nitrite in 80 c.c. of water is then added to the amine hydrochloride, cooled by ice-water, until a permanent reaction of nitrous acid upon the starch-potassium-iodide paper is obtained. The diazotoluene chloride thus formed is poured from a flask into the cuprous cyanide solution, with frequent shaking. After the addition of the diazo-solution, which should require about 10 minutes, the reaction mixture is heated on the water-bath for about a quarter-hour. The tolyl nitrile is then distilled over with the steam. This operation must also be done under a hood with a *good draught*, since hydrocyanic acid passes over. The nitrile distils as a yellow oil, which after some time solidifies in the receiver. It is separated by decanting the water, pressed upon a porous plate, and purified by distillation. If the oil will not solidify, the entire distillate may be taken up with ether, the ethereal solution shaken with caustic soda solution to remove the cresol, and then, after evaporating the ether, the residue remaining is distilled directly, or, in case it is solid, it is pressed out on a porous plate, as above, and then distilled. Boiling-point, 218°. Yield, about 15 grammes.

The diazo-group *cannot* be replaced in the same way by iodine as by chlorine, bromine, or cyanogen. If a water solution of a diazo-chloride, -bromide, or -cyanide is heated, a phenol is formed, as is also the case on heating a diazo-sulphate:



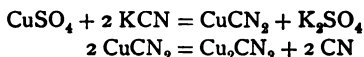
To Sandmeyer¹ we are indebted for the important discovery that, if the heating be done in the presence of cuprous chloride, bromide, or cyanide, the reaction taking place is analogous to the one by which phenyl iodide is formed:

¹ B. 17, 1633 and 2650; 18, 1492 and 1496.



The manner in which the cuprous salts act is not known; in any case they unite first with a diazo-compound to form a double salt, which plays a part in the reaction.

The reaction in the above preparation of cuprous cyanide takes place in accordance with the following equation:



In order to replace the diazo-group by chlorine or bromine, the above method is followed exactly. A diazo-solution is first prepared, and gradually added to a heated solution of cuprous chloride or bromide. With easily volatile chlorine or bromine compounds, it is desirable to use a reflux condenser, and to allow the diazo-solution to flow in from a dropping funnel. In some cases it is more advantageous not to use a previously prepared diazo-solution, but to proceed as follows: The amine is dissolved in an acid solution of a copper salt; this is heated, and to the hot solution, the solution of nitrite is added from a dropping funnel. The diazotisation and replacement of the diazo-group then takes place in one reaction. If the reaction-product is not volatile with steam, it may be obtained from the reaction-mixture by filtering, or extracting with ether.

The Sandmeyer reaction is capable of general application. Since the yield of the product is generally very good, for many substances it is used as a method of preparation. It should be finally pointed out that by the replacement of the diazo-group by cyanogen a new carbon union takes place.

11. REACTION: (a) REDUCTION OF A DIAZO-COMPOUND TO A HYDRAZINE. (b) REPLACEMENT OF THE HYDRAZINE-RADICAL BY HYDROGEN

EXAMPLES: (a) Phenyl Hydrazine from Aniline
(b) Benzene from Phenyl Hydrazine.

(a) Add 10 grammes of freshly distilled aniline to 100 c.c. of concentrated hydrochloric acid in a beaker, with stirring; aniline hydrochloride partially separates out in crystals. To the mixture,

cooled with ice, add slowly from a dropping funnel, a solution of 10 grammes of sodium nitrite in 50 c.c. of water, until a test with starch-potassium-iodide paper shows free nitrous acid. In this case the strong acid solution must not be brought directly upon the test-paper, but a test-portion is diluted with water in a watch-glass and then the test applied. To the diazo-solution add, with stirring, a solution of 60 grammes of stannous chloride in 50 c.c. of concentrated hydrochloric acid cooled with ice; a thick paste of crystals of phenyl hydrazine hydrochloride separates out. After standing several hours this is filtered off with suction (Büchner funnel and filter-cloth), the precipitate is pressed firmly together on the filter with a pestle; it is then transferred to a small flask and treated with an excess of caustic soda solution. Free phenyl hydrazine separates out as an oil, it is taken up with ether, the ethereal solution dried with ignited potash, and the ether evaporated. For the later experiments the phenyl hydrazine thus obtained can be used directly. If it is desired to purify the substance, the best method is to distil it in a vacuum, or it can be cooled by a freezing-mixture, and the portions remaining liquid are poured off. Yield, about 10 grammes.

Since the diazotisation in a strong hydrochloric acid solution as well as the filtration of strongly acid liquids is liable to miscarry, it is better to perform the experiment as follows: Dissolve 10 grammes of freshly distilled aniline in a mixture of 30 grammes of concentrated hydrochloric acid and 75 c.c. of water and diazotise it with a solution of 8 grammes of sodium nitrite in 30 c.c. of water, the beaker being cooled with ice-water. The diazo-solution is saturated with finely pulverised salt (about 30 grammes) with shaking; the solution is poured off from any undissolved salt, and being cooled with ice is treated with a cold solution of 60 grammes of stannous chloride in 25 grammes of concentrated hydrochloric acid. After several hours' standing, the phenylhydrazine hydrochloride separates out, this is filtered off with suction, washed with a little saturated salt solution, pressed out on a porous plate, and treated as above.

(b) In a 1-litre flask provided with a dropping funnel and con-

denser (Fig. 68) 150 grammes of water and 50 grammes of copper sulphate are heated to boiling, then from the funnel add gradually a solution of 10 grammes of free phenyl hydrazine in a mixture of 8 grammes of glacial acetic acid and 75 grammes of water. The oxidation of the phenyl hydrazine proceeds with an energetic evolution of nitrogen; the benzene is immediately distilled over with steam and collected in a test-tube. By another careful rectification from a small fractionating flask (without con-

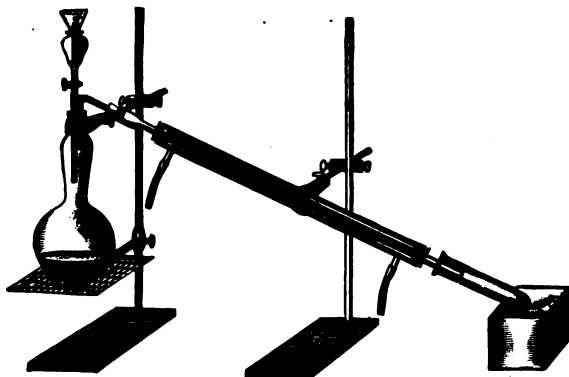
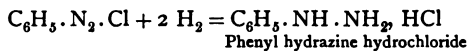


FIG. 68.

denser), pure benzene, boiling at 81° , is obtained. Yield, about 5 grammes.

Monosubstituted hydrazines of the type of phenyl hydrazine may be obtained according to the method of V. Meyer and Lecco,¹ by reducing the diazo-compounds with stannous chloride and hydrochloric acid:

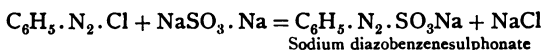


The reaction is always conducted as above: The amine is diazotised in a strong hydrochloric acid solution, and then a solution of stannous chloride in strong hydrochloric acid is added to it. Since the hydrochlorides of the hydrazines are difficultly soluble in concentrated hydrochloric acid, these separate out directly on the addition of the

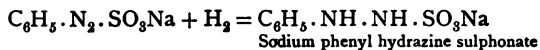
¹ B. 16, 2976.

stannous chloride, and can easily be obtained pure by filtration, as above.

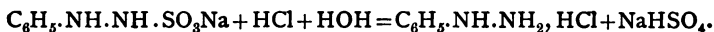
The reduction of the diazo-compounds to hydrazines may be accomplished by the method of Emil Fischer¹ which led to the discovery of this class of compounds, and also by another method. If neutral sodium sulphite is allowed to act on a diazo-salt, the acid radical of the diazo-compound is replaced by a residue of sulphurous acid, *e.g.* :



If this salt is now reduced with sulphurous acid, or with zinc dust and acetic acid, it takes up two atoms of hydrogen and is converted into a hydrazine sulphonate :

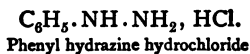


If this is heated with hydrochloric acid, the sulphonic acid group is split off, and phenyl hydrazine hydrochloride is formed, which, on evaporation, crystallises out :

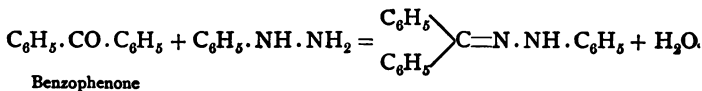
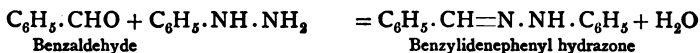


According to this method, which is slower, but cheaper than the former, phenyl hydrazine is prepared on the large scale.

The monosubstituted hydrazines possess a basic character ; in spite of the fact that they contain two ammonia residues, they combine with only one molecule of a monobasic acid, *e.g.* :



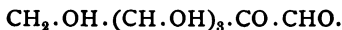
Phenyl hydrazine reacts with aldehydes and ketones, the two hydrogen atoms of the amido-groups unite with the oxygen atom of the CHO- or CO-groups, and are eliminated as water :²



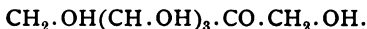
¹ A. 190, 67.

² B. 17, 572.

If this compound is heated with hydrochloric acid, it acts in the same way as all hydrazones, and phenyl hydrazine is eliminated; the original unchanged sugar is not formed again, but an oxidation product of it is obtained, a so-called "Osone." In the example selected, the osone is:



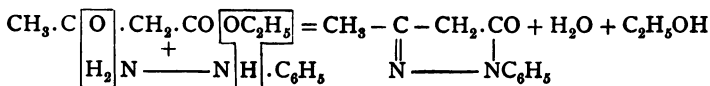
If this compound is treated with a reducing agent, the aldehyde group and not the ketone group is reduced, and the original sugar is not obtained:



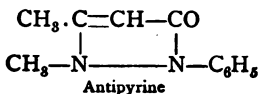
The aldose is converted into a ketose, the grape sugar into fruit sugar. The general importance of the reaction as applied to the sugars may be inferred from these brief statements.

EXPERIMENT : A cold solution of 2 grammes of phenyl hydrazine hydrochloride and 3 grammes of crystallised sodium acetate in 15 c.c. of water is treated with a solution of 1 gramme of pure grape sugar in 5 c.c. of water, and warmed on the water-bath. After about 10 minutes, the fine, yellow needles of the osazone begin to separate out; the quantity is increased by a longer heating. After heating an hour, the crystals are filtered off, washed with water, and allowed to dry in the air. Melting-point, 205°.

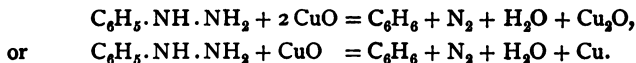
Phenyl hydrazine undergoes condensation with β -diketones and β -ketone-acid-esters with the formation of ring compounds containing nitrogen — the so-called pyrazoles and pyrazolones. The phenyl methyl pyrazolone formed from acetacetic ester and phenyl hydrazine is of importance:



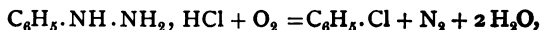
from which, by the action of methyl iodide, the important febrifuge—“Antipyrine”—dimethyl phenyl pyrazolone is obtained:



If the primary hydrazines are boiled with copper sulphate,¹ or ferric chloride,² the hydrazine radical is replaced by hydrogen, and there is obtained, *e.g.*, from phenyl hydrazine, benzene :



The statements made above concerning the replacement of a diazo-group by hydrogen are also applicable to this reaction. If it is desired to prepare an amido-compound from an amido-free compound, and if the direct reduction of the diazo-compound by sodium stannous oxide or alcohol (see page 237) has been shown to be impracticable, then, as above, the hydrochloric acid salt of the corresponding hydrazine is prepared, the free hydrazine is liberated, and oxidised with caustic soda. The amido-free substance is not always easily volatile, as in the example cited. In a case of this kind, the oxidation may be effected in an open vessel; the reaction product is obtained either by filtering or by extracting with ether. It may be pointed out here that it is more convenient to separate the hydrazine from the hydrochloric acid salt, and subject this to oxidation. If a hydrochloric acid salt of a hydrazine is oxidised, it may happen that the hydrazine radical will be replaced by chlorine :



which may give rise to complications.

12. REACTION: (a) PREPARATION OF AN AZO DYE FROM A DIAZO-COMPOUND AND AN AMINE. (b) REDUCTION OF THE AZO-COMPOUND

EXAMPLES: (a) Helianthine from Diazotised Sulphanilic Acid and Dimethyl Aniline

(b) Reduction of Helianthine

(a) Dissolve 10 grammes of sulphanilic acid, dried on the water-bath, in a solution of 3.5 grammes of dehydrated sodium carbonate in 150 c.c. of water, and treat with a solution of 4.2 grammes of pure sodium nitrite in 20 c.c. of water. To this mixture, after being cooled by water, is added a quantity of hydrochloric acid solution corresponding to 12.5 grammes of an-

¹ B. 18, 90.

² B. 18, 786.

hydrous hydrochloric acid. For this purpose, concentrated hydrochloric acid is diluted with an equal volume of water, and the specific gravity of the dilute acid is determined by a hydrometer. Consult a table, to find the amount of anhydrous hydrochloric acid corresponding to the reading of the hydrometer. (See Graham-Otto, Vol. II 1, p. 318.)¹

Before diazotising the sulphanilic acid, a solution of 7 grammes of dimethyl aniline in the theoretical amount of hydrochloric acid is prepared. Aromatic bases cannot be neutralised with hydrochloric acid in the same way as caustic potash, caustic soda, and ammonium hydroxide, by gradually treating with the acid and testing with blue litmus-paper until the liquid is just acid. In consequence of the weak basic character of the amines, their hydrochlorides still give an acid reaction with blue litmus-paper, therefore an acid reaction can be obtained even at the beginning of the neutralisation. Red fuchsine-paper possesses the property of becoming decolourised by free hydrochloric acid, which converts the red monoacid fuchsine into a colourless polyacid salt. The hydrochloric acid salts of bases, on the contrary, do not produce this decolourisation. In order to neutralise the dimethyl aniline (7 grammes), it is treated with 25 c.c. of water, and, with stirring, small quantities of concentrated hydrochloric acid are added; after each addition a test is made to show whether or not the fuchsine-paper is decolourised.

The dimethyl aniline hydrochloride thus obtained is added to the diazo-solution, and the mixture is made distinctly alkaline by the addition of not too much caustic soda solution. The dye separates out directly; the quantity can be increased if 25 grammes of finely pulverised sodium chloride are added to the solution. After filtering off and pressing out on a porous plate, the dye is recrystallised from a little water.

¹ If the specific gravity of the hydrochloric acid has been determined, the percentage of free anhydrous acid may be found without a table, by the following calculation: The decimal number is multiplied by 2, and a decimal point placed after the first two figures thus obtained, *e.g.*, sp. gr. = 1.134; $2 \times 134 = 268$. Percentage contents = 26.8. If the sp. gr. is greater than 1.18, a table must be consulted.

Preparation of Fuchsine-Paper: A crystal of fuchsine, the size of a lentil, is pulverised, dissolved by heating in 100 c.c. of water, and the solution filtered.

Into this immerse strips of filter-paper 2 cm. wide; they are dried either by suspending from a string in an acid-free place, or on the water-bath. The paper must not be an intense red, but only a faint rose colour. If the colour is too intense, the fuchsine solution must be correspondingly diluted with water.

Instead of the fuchsine-paper, the commercial Congo-paper will serve, the red colour of which is changed to blue by free acid.

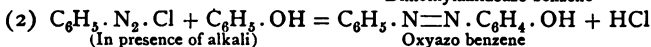
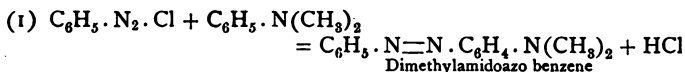
(b) Dissolve 2 grammes of the dye in the least possible amount of water by heating; while the solution is still hot, treat with a solution of 8 grammes of stannous chloride in 20 grammes of hydrochloric acid until decolourisation takes place. The colourless solution is then well cooled, upon which, especially if the sides of the vessel are rubbed with a glass rod, sulphanilic acid separates out: it is filtered off through asbestos or glass-wool. The filtrate is diluted with water, and caustic soda solution is added until the oxyhydrate of tin separating out at first is again dissolved. It is then extracted with ether several times, the ethereal solution dried with potash, and the ether evaporated, upon which the p-amidodimethyl aniline remains as an oil: on cooling and rubbing with a glass rod, it solidifies.

*Reactions of p-Amidodimethyl Aniline:*¹ The amidodimethyl aniline is treated gradually with small quantities of dilute sulphuric acid until it is just dissolved. Add a few drops of this solution to a dilute solution of hydrogen sulphide in a beaker which has been treated with $\frac{1}{50}$ of its volume of concentrated hydrochloric acid. To this mixture now add several drops of a dilute solution of ferric chloride. An intensely blue colouration, due to the formation of methylene blue, takes place.

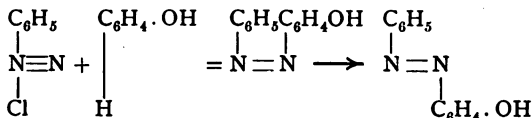
Diazo-compounds react with amines, as well as phenols, to form the Azo dyes:²

¹ B. 16, 2235.

² A. 137, 60; B. 3, 233.



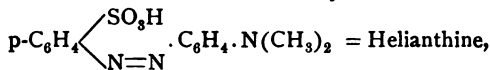
According to Hantzsch the reaction takes place in the following manner:



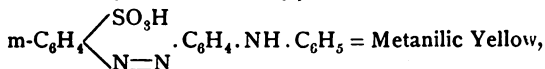
An unstable syn-azo compound is formed first, which rapidly changes into the more stable anti-azo compound.

In accordance with these two typical reactions, the vast number of monoazo dyes are prepared. The great number of possible combinations can be inferred from the following considerations: In reaction (1), instead of diazotised aniline, other bases, like o-toluidine, p-toluidine, xyloidine, cumindine, α -naphthyl amine, β -naphthyl amine, etc., may be used. In addition, the most varied derivatives of these bases, especially their sulphonic acids, like sulphanilic acid, metanilic acid, the large number of α - and β -naphthyl amines mono- to poly-sulphonic acids, may also be employed. Instead of dimethyl aniline, the diazo-compound can be combined or "coupled" with other tertiary, and in part also with secondary and primary amines, like diphenyl amine, or m-diamines, etc. In the second reaction, the diazo-compounds of the just mentioned bases can be employed as the starting-point, and these can be combined with mon-acid phenols, like cresol, naphthols, or di-acid phenols like resorcinol, or the sulphonic acids of these phenols, especially the numerous sulphonic acids of both naphthols. Since a dye must be soluble in water, and the alkali salts of the sulphonic acids of the dyes are more easily soluble than the mother substance containing no sulphonic acid groups, therefore, in the preparation of the azo dyes, the starting-point is usually a sulphonic acid. A few examples will explain these statements:

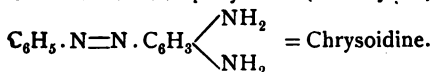
I. Amidoazo Dyes



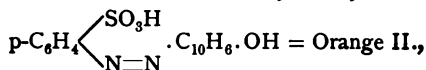
Diazotised sulphanilic acid + dimethyl aniline



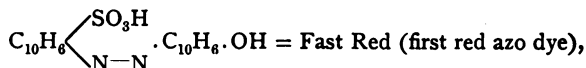
Diazot. Metanilic acid + diphenylamine (secondary base)



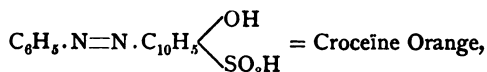
Diazot. Aniline + m-phenylenediamine (primary base)

II. *Oxyazo Dyes*

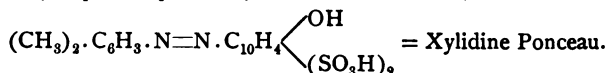
Diazot. Sulphanilic acid + β -naphthol



Diazot. α -Naphthionic acid + β -naphthol

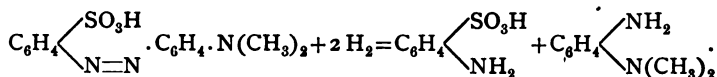


Diazot. Aniline + croceïne acid
(β -Naphthol sulphonic acid)



Diazot. Xylidine + β -naphthol disulphonic acid

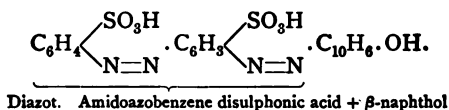
Concerning the constitution of the azo dyes, provided the components are known, the only question to solve is: which hydrogen atom of the undiazotised component combines with the acid radical of the diazo-compound (the acid thus formed being eliminated). The question may be answered by investigating the reduction products of the azo dyes. By energetic reduction, best in acid solution with stannous chloride, the double $\text{N}=\text{N}$ union is broken up, thus forming, with the addition of 4 atoms of hydrogen, two molecules of a primary amine, *e.g.* :



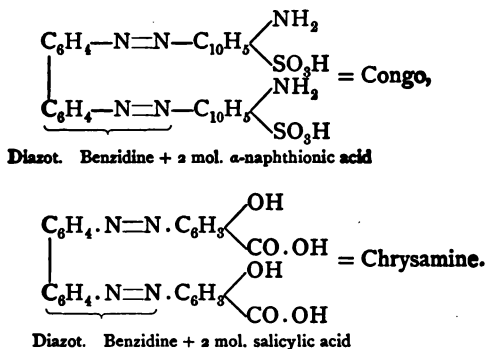
From this equation it is evident that by reduction, the amine which was diazotised — in the above case sulphanilic acid — may be obtained again on the one hand, on the other an amido-group is introduced into the second component. If the constitution of this second product can be determined, then the constitution of the azo dyes is also determined. It may be stated as a general rule that, when a diazo-compound combines with an amine or phenol, the hydrogen atom in the para position to the amido- or hydroxyl-group is always substituted. In accordance with this, in the above case, *p*-amidodimethyl aniline ought to be obtained on reduction. If the para position is already occupied, then the *o*-hydrogen atom unites with the acid radical.

In some cases, the formation and consequent reduction of an azo dye with the introduction of an amido-group into a phenol or amine is of practical value.

Azo dyes which contain two "chromophore groups," $\text{N}=\text{N}$, and which are called dis- or tetr-azo dyes, can be prepared; two methods may be employed: (1) The starting-point is an amido-azo-compound which already contains one azo group; this is diazotised, and then united with an amine or phenol. "Biebrich scarlet" is obtained in this way, by diazotising the disulphonic acid of amidoazobenzene, and combining it with β -naphthol:



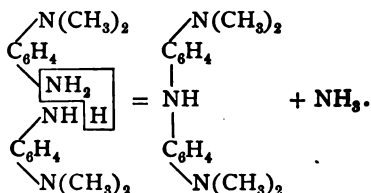
(2) A diamine is the starting-point; this is diazotised, and the bisdiazocompound is combined with two molecules of an amine or phenol. To this class belong the important dyes of the Congo group, prepared from the benzidine bases (see page 231), *e.g.*:



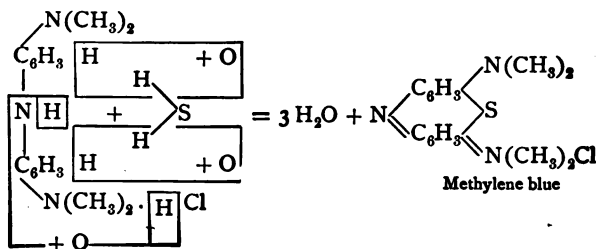
These Congo dyes possess the noteworthy property of colouring vegetable fibres (cotton) directly, whereas, with almost all other azo dyes, the cotton must be mordanted before dyeing.

In conclusion, the above dye-stuff reaction (which may be used for detecting the smallest amount of hydrogen sulphide) is technically carried out on the large scale, for the manufacture of the important methylene blue. The reaction takes place as follows: From two molecules of the diamine there is split off on oxidation with ferric chloride,

one molecule of ammonia, while a derivative of diphenyl amine is formed :



In the presence of hydrogen sulphide and hydrochloric acid, there is formed from this, by the oxidising action of ferric chloride, a derivative of thiodiphenylamine, as follows :



13. REACTION: PREPARATION OF A DIAZOAMIDO-COMPOUND

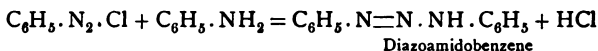
EXAMPLES : Diazoamidobenzene from Diazobenzenesulphide and Aniline¹

Dissolve 10 grammes of freshly distilled aniline in a mixture of 100 c.c. of water, and that quantity of concentrated hydrochloric acid corresponding to 12 grammes anhydrous hydrochloric acid (determine the sp. gr. by a hydrometer). The solution is cooled with ice-water, and diazotised with a solution of 8 grammes of sodium nitrite in 50 c.c. of water, in the manner already described. A solution containing 10 grammes of aniline, 50 grammes of water, and the exact theoretical amount of hydrochloric acid is previously prepared according to the directions given on pages

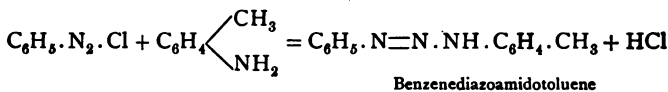
¹ A. 121, 257.

256 and 257; this is well cooled with ice water, and added to the diazo-solution, with stirring. Further, 50 grammes of crystallised sodium acetate are dissolved in the least possible amount of water, the cooled solution is added, with stirring, to the mixture of the diazo-compound with aniline hydrochloride. After standing half an hour, the diazoamidobenzene separates out, and is filtered off with suction, washed several times with water, well pressed out on a porous plate, and recrystallised from ligroin. Melting-point, 98°. Yield, almost theoretical.

If one molecule of a diazo-compound is allowed to act on one molecule of a primary amine, the acid radical of the former unites with the hydrogen atom of the latter, upon which the organic residues combine, as in the formation of the azo dyes. In this case, an amido-hydrogen atom is eliminated, so that a compound containing a chain of three nitrogen atoms is formed; in the formation of an azo dye, one of the benzene-hydrogen atoms of the amine is eliminated:

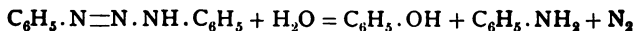


Mixed diazo-compounds may also be prepared by causing the diazo-derivative of an amine to combine with another amine:

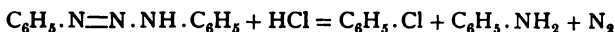


Diazo-compounds combine only with the *free* amines to form diazo-amido compounds; the object of the addition of sodium acetate at the end of the reaction (see above) is to set free the base from aniline hydrochloride.

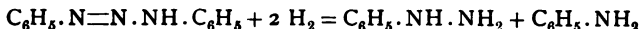
The diazoamido-compounds are yellow substances which do not dissolve in acids. They are far more stable than the diazo-compounds, and may be recrystallised without decomposition. Still, if they are heated rapidly they puff up suddenly and decompose. In their reactions they behave like a mixture of a diazo-compound and an amine. If, *e.g.*, they are boiled with hydrochloric acid, they decompose with evolution of nitrogen, and form a phenol and an amine:



On heating with cuprous chloride and hydrochloric acid, the Sandmeyer reaction takes place:

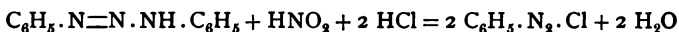


By reduction with acetic acid and zinc dust, they form a hydrazine:

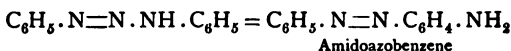


But, in addition to the reaction-product of the diazo-radical, there is always formed one molecule of an amine.

Under the influence of nitrous acid, they decompose, the amine residue being diazotised, into two molecules of a diazo-compound:

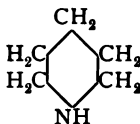


If a diazoamido-compound is warmed with an amine in the presence of some amine hydrochloride, transformation to the isomeric amidoazo-compound takes place:

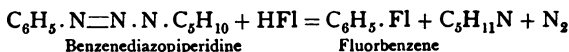


The next preparation deals with this reaction.

The diazo-compounds also have the power of combining with secondary amines to form diazoamido-compounds, — the combinations with an alkaloid base, piperidine $\text{C}_5\text{H}_{11}\text{N}$:



are of especial value for preparations. If they are gradually warmed with hydrofluoric acid, they are decomposed with the evolution of nitrogen into piperidine and a fluoride:¹



In this way it has been possible to prepare the aromatic fluorides.

In accordance with the older views it was believed that the aromatic fluorides could not be obtained from the diazo-compounds in the same way in which the analogous chlorides, bromides, and iodides are prepared; recently however they have been obtained by the direct decomposition of the diazo-fluorides.

¹ A. 243, 239.

14. REACTION: THE MOLECULAR TRANSFORMATION OF A DIAZO-AMIDO-COMPOUND INTO AN AMIDOAZO-COMPOUND

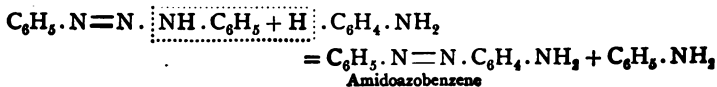
EXAMPLE: Amidoazobenzene from Diazoamidobenzene

To a mixture of 10 grammes of crystallised diazoamidobenzene, finely pulverised, and 5 grammes of pulverised aniline hydrochloride, contained in a small beaker, add 25 grammes of freshly distilled aniline; the mixture is then heated, with frequent stirring, one hour, on the water-bath, at 45°. It is then transferred to a larger vessel, and treated with water; dilute acetic acid is added, until all the aniline has passed into solution, and the undissolved precipitate remaining is completely solid. This is filtered off, washed with water, heated in a large dish with a large quantity of water (about a litre), and gradually treated with hydrochloric acid until the greatest portion of the precipitate is dissolved. From the filtered solution, steel-blue crystals of amidoazobenzene hydrochloride separate out, on long standing; these are filtered off, and washed with dilute hydrochloric acid, *not* with water.

If aniline hydrochloride is not at hand, prepare it by adding aniline to concentrated hydrochloric acid, with stirring. After cooling, the pasty mass of crystals separating out is filtered on glass-wool, pressed firmly together on the filter with a pestle, and then spread in thin layers on a porous plate.

In order to obtain the free amidoazobenzene, the hydrochloride is warmed with dilute ammonia, the free base filtered off, dissolved in alcohol by heating, and hot water is added until the liquid begins to be turbid. Melting-point, 127–128°. Yield, 6–8 grammes.

If a diazoamido-compound is heated with an amine and some amine hydrochloride, it goes over to an amidoazo-compound. The most probable cause of the reaction is that the amine residue of the diazo-amido-compound unites with a benzene-hydrogen atom of the amine hydrochloride, upon which the diazo-residue unites with the residue of the amine salt to form amidoazobenzene:



While the amidoazobenzene does not unite with hydrochloric acid, the new molecule of the amine formed in the reaction does, and thus there is a molecule of the amine hydrochloride present, which again causes the transformation, so that a small amount of the hydrochloride may transform an indefinitely large amount of the diazoamido-compound.

If amidoazobenzene is reduced, p-phenylene diamine and aniline are obtained. The transformation accordingly results in the formation of a compound in which the amido-groups are in the para position, which always happens when the para position is unoccupied. The amidoazo-compounds possess weakly basic properties; but if their salts are treated with much water, they partially dissociate.

The amidoazobenzene hydrochloride came into the market formerly, as a yellow dye, under the name of "Aniline Yellow." At present, it is scarcely used, but there is prepared from it, by heating with sulphuric acid, a mono- or di-sulphonic acid, which in the form of its alkali salts finds application as a dye under the name of "Acid Yellow," or "Fast Yellow." As already mentioned under the dis-azo dyes, from the diazo-compound of this dye, "Biebrich Scarlet" may be made by combination with β -naphthol. Finally, the amidoazobenzene is still used for the preparation of the Induline dyes.

15. REACTION: OXIDATION OF AN AMINE TO A QUINONE

EXAMPLE: Quinone from Aniline¹

To a solution of 25 grammes of aniline in a mixture of 200 grammes of concentrated pure sulphuric acid and 600 c.c. of water contained in a thick-walled beaker (a small battery jar), cooled to 5° by being surrounded with ice, add gradually, with constant stirring (use a small motor), from a dropping funnel, a solution of 25 grammes of sodium dichromate in 100 c.c. of water (Fig. 69). Should the temperature rise above 10°, the addition of the dichromate must be discontinued for a short time and a few pieces of ice thrown into the beaker. The reaction-mixture is then allowed to stand over night in a cool place, and the next morning it is again cooled and stirred, while a solution of 50 grammes of sodium dichromate in 200 c.c. of water is added. After the mixture has been allowed to stand until midday, it is divided into two equal parts, one of which is worked up into quinone as follows: In a large separating

¹ A. 27, 268; 45, 354; 215, 125; B. 19, 1467; 20, 2283.

funnel one half is treated with $\frac{2}{3}$ its volume of ether, and the two layers thus formed are *carefully* shaken together. If the shaking is too vigorous, the layers will not readily separate. After allowing it to stand for half an hour, the lower layer is run off (see page 44, Separation of coloured liquids), the ethereal solution is filtered through a folded filter, and the ether distilled off (water-bath with warm water). The water solution is again extracted with the condensed ether, and the ether again distilled from the same flask as before. In order to obtain perfectly pure quinone, a rapid current of steam is passed over the crude product — it is not treated with

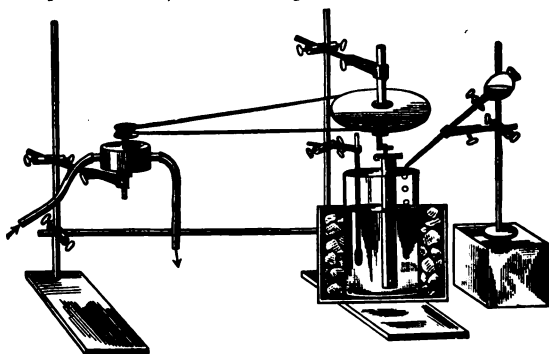


FIG. 69.

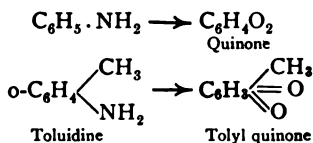
water; the pure quinone is carried over with the steam to the condenser and receiver, where it crystallises in the form of golden-yellow needles; they are filtered off and dried in a desiccator. Melting-point, 116° . Yield, 10–12 grammes.

If sodium dichromate is not at hand, the potassium salt may be used for the oxidation. In this case, 25 grammes of aniline are dissolved in a mixture of 200 grammes of sulphuric acid and 800 c.c. of water; then add, as above, with stirring and good cooling, 25 grammes of potassium dichromate, powdered extremely fine. On the next day add 50 grammes of this salt. In other respects, proceed as above.

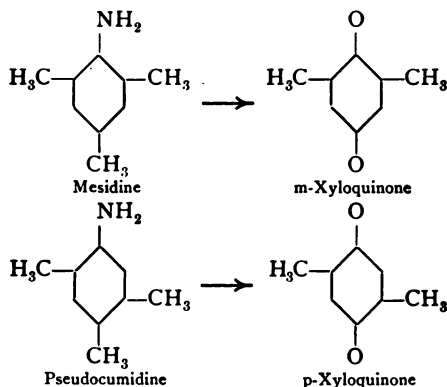
For the preparation of hydroquinone from quinone see p. 270.

Many primary aromatic amines yield quinones on oxidation with chromic acid. But the reaction cannot be expressed in a simple equation; still, it is always true that the amido-group and the hydrogen

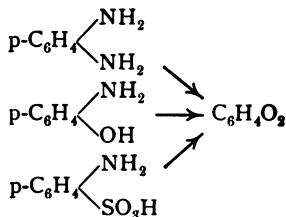
atom in the para position to this are each replaced by an oxygen atom, *e.g.*:



The tendency to form quinones is so great that even in cases where the para position to the amido-group is occupied by an alkyl (methyl) radical, the latter is split off and a quinone (poorer in carbon contents) is formed. Indeed, in the simplest cases, like p-toluidine and asym. m-xylidine, the reaction is very incomplete; however, mesidine, as well as pseudocumidine, give satisfactory yields of quinones belonging to the next lower series:

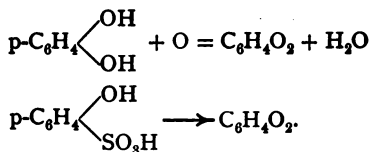


But if the para position is occupied by an amido-, oxy-, or sulphonic acid-group, this is eliminated and a quinone formed:

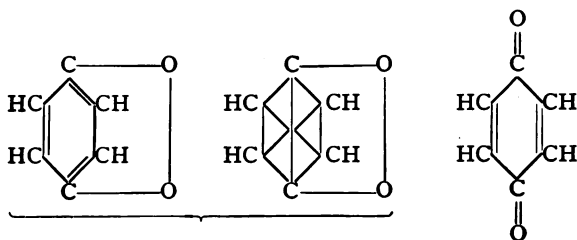


From these methods of formation it follows that the two quinone-oxygen atoms are in the para position to each other. The quinone reaction can be used in doubtful cases to decide whether a compound

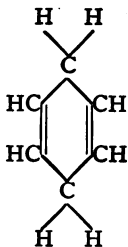
belongs to the para series. The quinones can also be obtained very easily from p-dioxy-compounds as well as from the p-sulphonic acids of mon-acid phenols:



Two formulæ for the quinones have been proposed:—



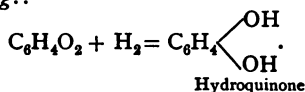
According to the former the quinones still contain the true benzene ring with either three double or six centric bonds. The two oxygen atoms are only singly united with the benzene-carbon atoms, and are united to each other. According to the second formula, the quinones do not contain the true benzene ring, but they are derived from a dihydrobenzene,



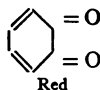
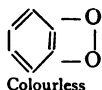
and are regarded as the di-ketone derivative of this. According to this conception, the oxygen atoms are connected by two bonds, as in the ketones, with the carbon atoms of the benzene nucleus. The facts in favour of the first formula are these: In many reactions both of the

oxygen atoms are replaced by two univalent atoms or radicals. Thus, *e.g.*, by the action of phosphorus pentachloride on quinone, p-dichlorobenzene is formed, while the second formula would lead one to expect a tetra-chloride. In support of the second formula is the fact that hydroxylamine acts directly on quinones, as on ketones, with the formation of a mono- or di-oxime.

The p-quinones are coloured compounds, possessing a characteristic odour; they are easily volatile with steam, but with a slight decomposition. They are somewhat volatile even with the vapour of ether, as one observes in the preparation of quinone. On reduction they take up two hydrogen atoms and pass over to hydroquinones. (See the next preparation) *e.g.* :



Of late years ortho-benzoquinone has also been prepared by the oxidation of pyrocatechin (o-dihydroxy benzene) with silver oxide.¹ Under certain conditions a *colourless*² o-quinone is first formed, but being very unstable it passes over into *red* quinone. For the colourless quinone the so-called Peroxide formula has been proposed, and for the red quinone the Ketone formula :



The o-quinones are less stable than the p-quinones. They are odourless, and non-volatile with steam.

16. REACTION : REDUCTION OF A QUINONE TO A HYDROQUINONE

EXAMPLE : Hydroquinone from Quinone

Conduct sulphur dioxide into the second half of the quinone solution obtained above, until the liquid smells intensely of the gas, then allow it to stand for 1-2 hours. Should the odour of sulphur dioxide vanish, it is passed in again and the mixture allowed to stand for some time as before. It is then extracted with the ether distilled from the quinone in the preceding experi-

¹ B. 37, 4744.

² B. 41, 2580.

ment, several times; the ether is evaporated or distilled, and the hydroquinone, well pressed out on a porous plate, is crystallised with the use of animal charcoal from a little water. Melting-point, 169° . Yield, 8–10 grammes.

Since the hydroquinone solution may be extracted with ether with much greater ease than the quinone solution, and since the hydroquinone is smoothly oxidised to quinone, the preparation of quinone may be done as follows: The *entire quantity* of the oxidation product is saturated with sulphur dioxide, and as just described the hydroquinone may be obtained by repeated extraction with ether. In order to convert it into quinone it is dissolved in the least possible amount of water, to which is added 2 parts of concentrated sulphuric acid to 1 part of hydroquinone; the well-cooled liquid is treated with a water solution of sodium dichromate until the green crystals of quinhydrone (an intermediate product between quinone and hydroquinone) separating out in the beginning have changed into pure yellow quinone.

The equation for the formation of hydroquinone from quinone has been given above. All homologous quinones react in the same way. The hydroquinones are di-acid phenols, which dissolve in alkalis and show all the properties of phenols. They are not volatile with steam.

17. REACTION: BROMINATION OF AN AROMATIC COMPOUND

EXAMPLE: Mono- and Di-brombenzene from Bromine and Benzene

A wide-neck 250 c.c. flask is connected with a vertical tube 50 cm. long and $1\frac{1}{2}$ cm. wide, the upper end of which is closed by a cork bearing a glass tube, not too narrow, bent twice at right angles. The other end is connected with a flask containing 250 c.c. of water, by a cork having a *small canal* in the side (Fig. 70). The tube does not touch the liquid, but the end is about 1 cm. above the surface. After 50 grammes of benzene¹ and 1 gramme of coarse iron filings (the bromine carrier) have been placed in the flask, it is cooled in a large vessel (battery jar) filled

¹ One hundred grammes of benzene should be used if the Grignard reaction (see p. 348) is to be carried out later on.

with ice-water ; through the vertical tube there is added 40 c.c. = 120 grammes of bromine : the narrow tube is immediately

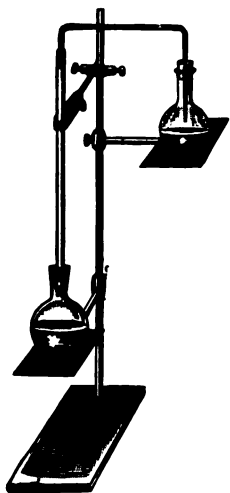


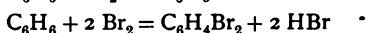
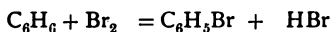
FIG. 70.

connected with the vertical tube. After some time an extremely energetic reaction will begin, generally spontaneously, with the evolution of hydrobromic acid, which is completely absorbed by the water. Should the reaction not begin at once, the ice-water is removed for a short time, and if necessary the flask is immersed for a moment in slightly warm water. But as soon as even a weak gas evolution begins, the flask is at once cooled again, since otherwise the reaction easily becomes too violent. Should this happen in spite of the cooling, the cause is found in the fact that the iron filings used were too fine. In other experiments use coarser filings or small iron nails. When the main reaction is over, the ice-water is removed, the flask dried and heated over a

small flame until the red bromine vapours are no longer visible above the dark-coloured liquid. The reaction-product is washed several times with water and then distilled with steam. As soon as crystals of dibrombenzene separate out in the condenser, the receiver is changed and the distillation continued until all the dibrombenzene has passed over. The liquid monobrombenzene is separated from the water, dried with calcium chloride, and subjected to a fractional distillation ; the portion passing over between $140-170^{\circ}$ is collected separately. This is again distilled, and the portion going over between $150-160^{\circ}$ collected. The boiling-point of the pure monobrombenzene is 155° . Yield, 60-70 grammes. The residue boiling above 170° remaining in the flask after the two distillations, is poured, while still warm, on a watch-glass, and after cooling is pressed out, together with the separately collected dibrombenzene, on a porous plate. On crystallising from alcohol, coarse colourless crystals of p-dibrombenzene are obtained, which melt at 89° .

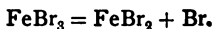
The by-product, hydrobromic acid, is purified as described in the Inorganic Part. (See page 379.)

A portion of the hydrogen of the aromatic hydrocarbons is very easily replaced by bromine, especially in the presence of a carrier, even at low temperatures; while in the aliphatic series the direct substitution of bromine is not used as a preparation method for alkyl bromides, the aromatic bromides are readily prepared in this way. According to the amount of bromine used one or more hydrogen atoms may be substituted; it may happen, *e.g.*, particularly with benzene under the influence of an energetic bromination, that all the hydrogen atoms may be replaced by bromine. A single bromide, even on using only the theoretical amount of bromine, is never formed; but rather a portion of the hydrocarbon is brominated short of the theoretical action, and another portion is always acted upon farther, with the formation of a higher bromine substitution product. Thus in the example above cited, besides the principal product, monobrombenzene, a small quantity of dibrombenzene is formed:



In most cases, however, the principal product may be separated from the by-product without difficulty by distillation or crystallisation. Since the hydrogen atoms substituted by bromine combine with bromine to form hydrobromic acid, therefore, for the introduction of each bromine atom, a molecule (two atoms of bromine) must be used.

The introduction of bromine can be essentially facilitated by the use of a so-called bromine carrier. As such, the bromides of metalloids, or metals, are used; (1) either in the already prepared condition, or (2) they can be generated from their elements in the reaction. To the first class belong ferric bromide and aluminium bromide. The action of ferric bromide depends on the fact that on being reduced to ferrous bromide, it yields bromine in *statu nascendi*:

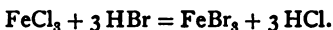


Ferric bromide . Ferrous bromide

Since the ferrous bromide unites with bromine again, to form ferric bromide, a small quantity of this has the power to transfer an indefinitely large quantity of bromine:

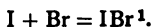


Instead of ferric bromide, ferrous bromide or anhydrous ferric chloride may be used. The latter decomposes with hydrobromic acid to ferric bromide and hydrochloric acid:



The activity of aluminium bromide is explained by the fact that it unites with the hydrocarbon to form a double compound which is more capable of reacting with other substances than the hydrocarbon itself.

To the second class belong iodine, sulphur, phosphorus, iron, aluminium, etc. If these elements are added to the brominating mixture, the corresponding bromides are formed, *e.g.*:



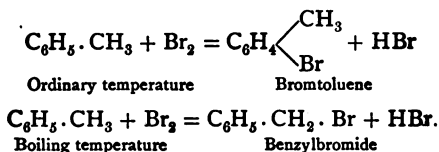
While these give up all their bromine, or a portion of it, as is the case with ferric bromide, in the atomic condition, the residue again unites with bromine, and as above, a small quantity of the carrier may transfer large quantities of atomic bromine.

Bromine can also act on aromatic hydrocarbons to form addition products, since it may be *added* in one, two, or three molecules, and thus break up the double or centric union. Thus, *e.g.*, the hexabrom-addition product, $\text{C}_6\text{H}_6\text{Br}_6$, is obtained from the action of bromine on benzene in the sunlight. Since the addition products render difficult the purification of substitution products, especially on distillation (they decompose when distilled), it is often necessary to remove them before the purification, by long boiling with alcoholic caustic potash, or alcoholic caustic soda. Under these conditions, one-half of the bromine atoms added in common with the same number of hydrogen atoms are abstracted as hydrobromic acid; the residue of the molecule is converted into a substitution derivative, which is not troublesome in the purification:



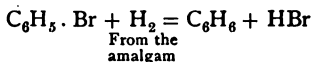
On brominating benzene, the same products will be formed, whether the temperature is high or low, but when its homologues are treated with bromine, the nature of the products depends upon the temperature. As will be pointed out more fully, under the chlorination of toluene, the law holds here, that at low temperatures the halogen enters the ring; at high temperatures, the side-chain, *e.g.*:

¹ Compare page 165.

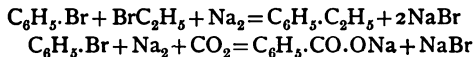


The aromatic bromides which contain bromine in the benzene nucleus are either colourless liquids or crystals, which in contrast with the side-chain substituted isomers in part possess an aromatic odour, and their vapours do *not* attack the eyes and nostrils. The bromine is held very firmly in them, more firmly than in the aliphatic bromides, and cannot be detected by silver nitrate. While the aliphatic bromides, as mentioned under bromethyl, decompose with ammonia, alcoholates, alkalis, etc., to form amines, ethers, alcohols, etc., respectively, these reagents do not act on the aromatic bromides. The bromides containing the bromine in the side-chain, behave like their aliphatic analogues.

By the action of sodium amalgam, the bromine may be replaced by hydrogen, *e.g.* :



The aromatic bromides are of synthetical importance,¹ especially for the building up of homologous hydrocarbons and the preparation of carbonic acids :



The next preparation will take up the first of these reactions in detail.

The hydrocarbons and most of their derivatives, like nitro-, amido-compounds, aldehydes, acids, etc., may be brominated with greater or less ease. At this place, the various modifications by which the bromination may be effected will be mentioned. If a substance is very easily brominated, the bromine may be used in a diluted condition. For this purpose, either bromine water or a mixture of bromine with carbon

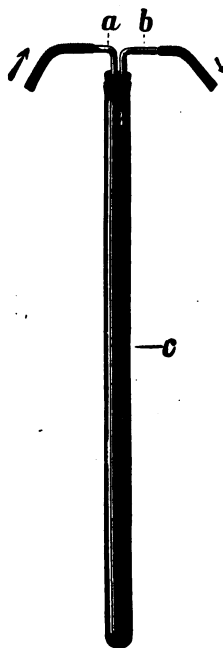


FIG. 71.

¹ Compare also the Grignard reaction (p. 348).

disulphide or glacial acetic acid may be employed. In many cases a bromination may be very well effected by using gaseous bromine. The method of procedure is as follows: The substance is spread out in thin layers on a watch-glass and placed under a glass bell-jar, under which is also a small dish containing bromine. If it is desired to cause bromine to act gradually, it is allowed to drop from a separating funnel, in concentrated form or in solution, on the compound to be brominated. If an extremely slow and very careful bromination is desired, the bromine may be allowed to flow drop by drop from a siphon-shaped capillary tube. If bromination takes places with difficulty, the brominating mixture is heated, either in an open vessel or in a sealed tube. In the first case the condensing apparatus cannot, as usual, be connected to the flask with a cork or rubber stopper, since this is soon attacked and destroyed by the bromine. Instead, the condenser is well wrapped with asbestos twine and then pushed into the conical part of the neck of the flask, the asbestos being pressed in with a knife. A condenser of the kind represented in Fig. 71 can also be used. A long tube *c*, sealed at one end, is closed by a two-hole cork, through one of which passes a long glass tube reaching almost to the bottom *a*; the other bears a short tube just passing through the cork. Water is caused to flow through *a*; it flows out of *b*. This cooling apparatus is suspended in the heating-flask, which is selected with as long a neck as possible.

18. REACTION: FITTIG'S SYNTHESIS OF A HYDROCARBON

EXAMPLE: Ethyl Benzene from Brombenzene and Bromethyl¹

In a dry, round, $\frac{1}{2}$ -litre flask, provided with a long reflux condenser (the flask is supported on a straw ring in an empty water-bath), place 27 grammes of sodium, cut in scales as thin as possible with a sodium knife, and add 100 c.c. of alcohol-free, dry ether prepared as described below. As soon as this has been completely dried by the sodium, which may be recognised by the fact that the upper surface is no longer disturbed by wave-like motions (after several hours' standing), pour through the condenser a mixture of 60 grammes of brombenzene and 60 grammes of bromethane, and allow to stand until the next day. Should the

¹ A. 131, 303.

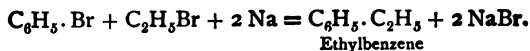
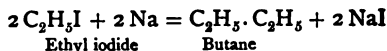
liquid begin to boil gently, which may easily happen at a summer temperature, cold water is poured into the water-bath. Water is not allowed to run through the condenser over night. During the reaction, the bright sodium will be changed to a blue powder, and an ethereal solution of ethylbenzene is formed. The ether is then distilled off on a water-bath, and the condenser is replaced with an air condenser 40–50 cm. long and 1 cm. wide, containing a short bend. After the flask has been placed in an oblique position, the extreme end of its neck is clamped *loosely*, and the ethylbenzene is distilled from the sodium bromide and sodium *by a large, luminous flame, which is kept in constant motion*. With the use of a Linnemann apparatus, the crude product is finally subjected to two distillations. The boiling-point of pure ethylbenzene is 135°. Yield, about 25 grammes.

The residue of sodium bromide and sodium remaining in the flask must be handled with extreme caution. Water must not be added to it, nor must it be thrown into the sink or waste-jars, nor allowed to stand a long time; it is better to throw the flask, which cannot be used again, and its contents into some open place. The sodium residue may be rendered harmless by throwing water on it from a great distance.

Preparation of Anhydrous, Alcohol-free Ether

Shake 200 grammes of commercial ether in a separating funnel with half its volume of water; the latter is allowed to run off, and the operation repeated a second time with a fresh quantity of water, by which the alcohol is removed. The ether is dried by standing over calcium chloride, not too little, two hours. It is then filtered through a folded filter, and can now be used for the above reaction.

Fittig's synthesis of the aromatic hydrocarbons is the analogue of Wurtz' synthesis of the aliphatic hydrocarbons, *e.g.*:



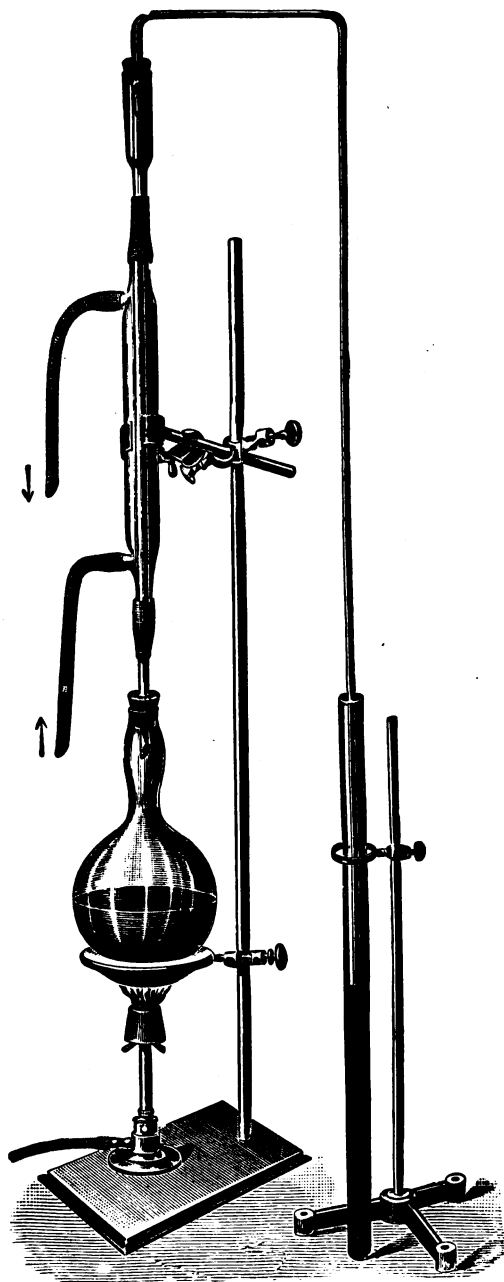


FIG. 72.

or benzene is employed. These substances are not alike in their activity, since ligroin and benzene generally prolong the reaction, and on this account find application in a very energetic reaction; ether does not retard the reaction, but causes it to be more regular. At times, the reaction-mixture will not act, even on long standing. In this case, the reaction can frequently be started by a short heating, or the addition of a few drops of ethyl acetate. Since the use of this compound, at times, causes a very stormy action, it is more advantageous to wait for the reaction to begin spontaneously, even if a long time is necessary. In syntheses which are moderately difficult, the reaction-mixture, treated with a diluent, can be heated on the water-bath or in an oil-bath; while, if the reaction takes place with great difficulty, the mixture, generally without dilution, must be heated in an oil-bath. In the latter case, the reaction may be still further facilitated by heating under pressure of a mercury column. By this means, it is possible to heat the reacting substances in an open vessel above their boiling-points. (Fig. 72.)

19. REACTION: SULPHONATION OF AN AROMATIC HYDRO-CARBON (I)

EXAMPLE: (a) Benzenemonosulphonic Acid from Benzene and Sulphuric Acid ¹

(b) Sulphobenzide. Benzenesulphonchloride. Benzenesulphonamide

(a) To 150 grammes of liquid fuming sulphuric acid, containing from 5–8 % of anhydride, placed in a 200 c.c. flask provided with an air condenser, gradually add, with good shaking and cooling with water, 40 grammes of benzene; before the addition of a new portion, always wait until the last portion, which at first floats on the surface of the acid, dissolves on shaking. The sulphonation requires about 10–15 minutes. The reaction-mixture is then added, with stirring, drop by drop, from a separating funnel, to three to four times its volume of a cold, saturated solution of sodium chloride contained in a beaker. In order that the solution may not be heated above the room temperature, the beaker is

¹P. 31, 283 and 631; A. 140, 284; B. 24, 2121.

placed in a large water-bath filled with ice-water. After some time, but with especial ease when the walls of the vessel are rubbed with a sharp-edged glass rod, the sodium salt of benzenesulphonic acid separates out in the form of leaflets of a fatty lustre; the quantity is increased, on long standing, to such an extent that the contents of the beaker are converted into a thick pasty mass of crystals. If the separation of crystals does not begin, 10 c.c. of the liquid is shaken in a corked test-tube, and cooled by immersion in water. The solidified content of the tube is then added to the main quantity in the beaker. In summer, at times, it may require a several hours' standing before the separation of crystals is ended. The pasty mass of crystals is then filtered off with suction on a Büchner funnel, firmly pressed together with a pestle, and washed with a little saturated sodium chloride solution.

To obtain the salt perfectly dry it is transferred to a linen bag and well squeezed under a screw-press. After being pulverised it is heated to dusty dryness in an air-bath at 110° . Yield, about 100 grammes.

If even after long standing an abundant separation of crystals does not take place, the cause is probably due to the large percentage of anhydride in the fuming sulphuric acid. Under these conditions it is diluted with concentrated acid, and the experiment repeated. If on the other hand the acid is too weak, the benzene will not dissolve in it. In this case, during the sulphonation the mixture is not cooled, and the reaction is allowed to take place at $40-50^{\circ}$.

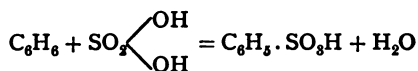
In order to obtain pure sodium benzenesulphonate, 5 grammes of the crude product is crystallised from absolute alcohol, upon which it is noticed that the sodium chloride mixed with it is insoluble in alcohol.

(b) In order to obtain the by-product, sulphobenzide, 30 grammes of the pulverised salt is warmed with 50 c.c. of ether, filtered with suction while hot, and washed with ether. After evaporating the ether, a small amount of a crystalline residue is obtained; this is recrystallised in a test-tube from ligroïn. Melting-point, 129° .

To prepare benzenesulphonchloride from sodium benzenesulphonate, the extracted salt and unextracted salt are treated in a dry flask (under the hood) with finely powdered phosphorus pentachloride (for 3 parts dry sodium benzenesulphonate, use 4 parts phosphorus pentachloride). Mix by thorough shaking. The mixture is warmed $\frac{1}{4}$ to $\frac{1}{2}$ hour on an actively boiling water-bath. The cold reaction-product is then poured gradually into ice-water in a flask (use ten times the weight of the sodium salt); it is shaken up from time to time, and, after standing for two to three hours, the sulphonchloride is taken up with ether and the generally turbid ethereal solution filtered; the ether is then evaporated off. Yield, 40–50 grammes.

In a porcelain dish 10 grammes of finely powdered ammonium carbonate are treated with about 1 c.c. of benzenesulphonchloride, and rubbed together intimately; the mixture is heated, with good stirring, over a small flame, until the odour of the sulphonchloride has vanished. After cooling, it is treated with water, filtered with suction, washed several times with water, and the benzenesulphonamide crystallised from alcohol to which hot water is added until turbidity begins. Melting-point, 156° .

Under the sulphonation of aniline it was mentioned that the aromatic compounds differ from the aliphatic compounds in that they can be sulphonated by the action of sulphuric acid; *i.e.* the benzene-hydrogen atoms are replaced by the sulphonic acid group, SO_3H . Thus the above reaction takes place in accordance with the following equation:

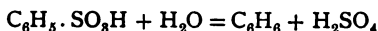


Since, in the sulphonation, an excess of sulphuric acid is always used, after the reaction is complete it is necessary to separate the sulphonic acid from the excess of sulphuric acid. Many sulphonic acids, especially those of the hydrocarbons, are very easily soluble in water, so that the pure acid cannot be separated out on mere dilution with water, as is the case with sulphanilic acid. There are three methods in common use for the isolation of sulphonic acids soluble in water. The sulphonic acids obtained most easily are those difficultly soluble in cold sulphuric acid. In this case it is only necessary to cool the sulphon-

ating mixture, and filter off the sulphonic acid separating out, with suction over asbestos or glass-wool. A second method consists in allowing the sulphuric acid solution to flow into a saturated solution of common salt; in many cases the difficultly soluble (in sodium chloride solution) sodium salt of the sulphonic acid separates out. Frequently it is more advantageous to use sodium acetate, potassium chloride, ammonium chloride, or other salts, instead of sodium chloride. Almost all soluble sulphonic acids, in the form of their alkali salts, can be separated by these two methods in the shortest time. In dealing with a new substance, preliminary experiments with small quantities of the substance are made to determine which salt is best adapted for the separation. The Theory of Salting out is discussed at the end of this chapter. The third method, which is the one generally applicable, depends upon the property of sulphonic acids, of forming soluble salts of calcium, barium, and lead in contradistinction to sulphuric acid. If the sulphuric acid solution, diluted with water, is neutralised with the carbonate of one of these metals and then filtered, the filtrate contains only the corresponding salt of the sulphonic acid, while the sulphuric acid in the form of calcium, barium, or lead sulphate remains on the filter. If the alkali salts of the sulphonic acids are desired, the water solution of one of the above salts is treated with the alkali carbonate until a precipitate is no longer formed. The precipitate is filtered off, and the pure alkali salt of the sulphonic acid is obtained in solution, which, on evaporation to dryness, yields the salt in the solid condition.

In order to obtain the free sulphonic acid, the lead salt is prepared and then decomposed with sulphuretted hydrogen.

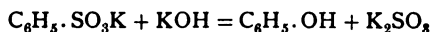
The sulphonic acids of the hydrocarbons are generally colourless, crystallisable compounds, very easily soluble in water, insoluble in ether, behaving like strong acids. By heating with hydrochloric acid, under pressure if necessary, or by the action of steam, the sulphonic acid group may be split off, *e.g.* :



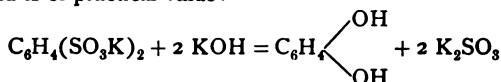
This reaction is of importance in many cases for the separation of hydrocarbon mixtures. If under certain conditions one hydrocarbon is sulphonated, and another is not, the latter can be separated from the former by removing the sulphuric acid solution of the sulphonic acid of the first, and from this the original hydrocarbon may be regenerated by one of the methods mentioned.

Of particular importance is the behaviour of sulphonic acids when

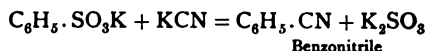
fused with caustic potash or caustic soda, by which the sulphonic acid group is eliminated and a phenol formed:



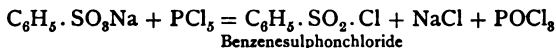
With benzenesulphonic acid this important reaction does not take place smoothly; for this reason the directions for carrying it out practically will be given later in another place (see β -naphthol). Poly-acid phenols may also be obtained from poly-basic sulphonic acids. The formation of m-dioxybenzene or resorcinol from benzenedisulphonic acid is of practical value:



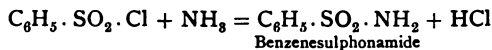
If an alkali salt of a sulphonic acid mixed with potassium cyanide or potassium ferrocyanide is subjected to dry distillation, the sulphonic acid group is replaced by cyanogen and an acid-nitrile is obtained, *e.g.*:



The sulphonic acids behave toward phosphorus pentachloride like the carbonic acids, with the formation of acid-chlorides:

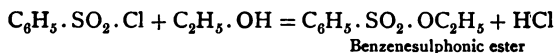


The sulphonch chlorides are not decomposed, or only slightly decomposed by cold water. In order to separate them from the phosphorus oxychloride, the mixture is generally poured into cold water; after long standing the oxychloride is converted into phosphoric acid, and the acid-chloride insoluble in water is obtained by decanting the water or extracting with ether; or in case it is solid, by filtering. The sulphonch chlorides are generally distinguished by a very characteristic odour. They can be distilled in a vacuum only, without decomposition. Treated with ammonia they form sulphonamides, which crystallise well and are used for the characterisation of the sulphonic acids:

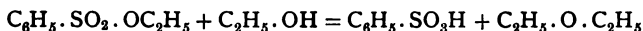


In the sulphonamides, in consequence of the strongly negative character of the $\text{X} \cdot \text{SO}_2$ -group, the hydrogen of the amido-group is

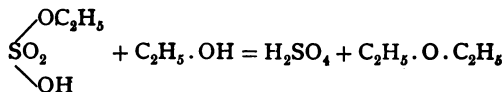
so easily replaced by metals, that they dissolve in water solutions or the alkalies to form salts of the amide. (Try it.) If a sulphon-chloride is allowed to stand a long time with an aliphatic alcohol, a sulphonic acid ester is formed, *e.g.*:



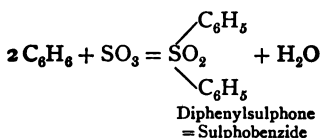
If this is now warmed with an alcohol, an aliphatic ether is formed, with the generation of the sulphonic acid, *e.g.*:



The formation of ether in this case is analogous to the formation of ethyl ether on heating ethyl sulphuric acid with alcohol:



Since this reaction is continuous, and since the benzene sulphonic acid formed in the reaction is a weaker acid than sulphuric acid, and consequently does not carbonise the alcohol like sulphuric acid, the operation may be continued for a long time uninterruptedly. For these reasons recently attempts have been made to employ the aromatic sulphonic acids for the technical preparation of ether. If the sulphonation is effected as above with fuming sulphuric acid, in many cases, besides the sulphonic acid a small quantity of sulphone is formed, *e.g.*:

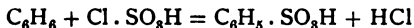


For sulphonating purposes, either ordinary concentrated sulphuric acid or the so-called monohydrate or fuming sulphuric acid of various grades is used, according to the conditions. The reaction is conducted with cooling, at the room temperature, or with heating.

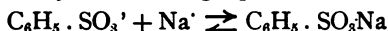
To facilitate the elimination of water, phosphorus pentoxide or potassium sulphate may be added to the sulphonating mixture.

In some cases it is of advantage to use chlorsulphuric acid instead

of sulphuric acid; the reaction takes place in accordance with the following equation :



Theory of Salting out. — The process of salting out depends upon the following conditions : According to the theory of electrolytic dissociation, the water solution of an electrolyte like sodium benzenesulphonate, contains the ions of $\text{C}_6\text{H}_5\text{SO}_3'$ — and Na' —, and the electrically neutral, undissociated molecules of $\text{C}_6\text{H}_5 \cdot \text{SO}_3\text{Na}$. Now it must not be assumed that in such a solution, at any given time, ions and molecules remain in this condition ; it is more likely that as the ions of $\text{C}_6\text{H}_5 \cdot \text{SO}_3'$ — and Na' — come into contact with each other under favourable conditions, they recombine forming new molecules, while at the same time undissociated molecules dissociate. We therefore have a dynamic equilibrium of a reversible reaction (similar to that mentioned under ethyl acetate), which may be expressed by the following equation :



If we indicate the concentrations of the ions and molecules by $\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3'}$, $\text{C}_{\text{Na}'}$ and $\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3\text{Na}}$, the mass action equation for this equilibrium will be :

$$(I) \frac{\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3'} \times \text{C}_{\text{Na}'}}{\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3\text{Na}}} = K$$

$$\text{or,} \quad (II) \text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3'} \times \text{C}_{\text{Na}'} = K \times \text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3\text{Na}}$$

where K is the dissociation constant. Suppose we add solid common salt to such a solution and take care that no solid salt separates out ; then since common salt is a strong electrolyte, it will largely dissociate into ions in the solution, and the concentration of Na' -ions will be increased. But since the constant K has the same value for all proportions, the increase of $\text{C}_{\text{Na}'}$ will decrease $\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3'}$ and will consequently increase $\text{C}_{\text{C}_6\text{H}_5 \cdot \text{SO}_3\text{Na}}$. In this way only can K have a constant value. Hence, the addition of common salt will cause a decrease in the number of $\text{C}_6\text{H}_5 \cdot \text{SO}_3'$ -ions, and an increase in the number of undissociated molecules. This is known as "diminution in dissociation by the addition of an electrolyte containing one ion in common." If we now continue to add more common salt, the concentration of $\text{C}_6\text{H}_5 \cdot \text{SO}_3'$ -ions will constantly diminish, while that of the undissociated molecules will constantly increase, and a point will finally be reached when the solution will become saturated with undissociated molecules ; i.e. a maximum number of molecules will be held in solution. If more common salt is now added, the newly formed undissociated molecules will separate out in the solid form.

At this stage $C_6H_5 \cdot SO_3Na$ becomes a constant, and the product of the concentration of ions is called the "*solubility product*." When the solid begins to separate out, the amount of the soluble undissociated portion becomes independent of the presence of the electrolyte having a common ion, although the concentration of the ions is variable, and is determined by the solubility product. Thus, to mention an extreme example, in one case $C_6H_5 \cdot SO_3$ may be large and C_{Na} small, while in another case C_{Na} may be large and $C_6H_5 \cdot SO_3$ small. Thus it becomes impossible, even by the addition of a large quantity of an electrolyte having a common ion, to salt out the undissociated molecules that are in solution. This may be termed the solution-tension of the undissociated molecules, and corresponds with Dalton's law of the vapour-tension of a substance, which is independent of external pressure. The vapour-tension of liquid water whether it be confined in a vacuum space, or kept at a definite pressure in physical equilibrium with other gases, will always be the same at any given temperature.

In the practical example described above, the conditions are somewhat complicated. Neglecting the excess of sulphuric acid used, we have a water solution of benzenesulphonic acid, treated with common salt. But according to the ionic dissociation theory, such a solution will contain ions of $C_6H_5 \cdot SO_3'$, Cl' , Na' and H' in addition to undissociated molecules of $C_6H_5 \cdot SO_3Na$, $NaCl$, $C_6H_5 \cdot SO_3H$ and HCl .

But the main fact to be taken into consideration is the equilibrium existing between $C_6H_5 \cdot SO_3'$, Na' and $C_6H_5 \cdot SO_3Na$, which is not influenced by the presence of other substances. When the solubility limit for this equilibrium is exceeded by increasing the number of Na -ions, by the addition of common salt, then the undissociated sodium benzenesulphonate will separate out in the solid state.

It is hardly necessary to state that with certain solubility proportions, not only solid salt, but also its solution, can be used for salting out.

20. REACTION: REDUCTION OF A SULPHONCHLORIDE TO A SULPHINIC ACID OR TO A THIOPHENOL

EXAMPLES: (a) Benzenesulphinic Acid.¹ (b) Thiophenol²

(a) Heat 40 grammes of water to boiling in a 300 c.c. flask provided with a short reflux condenser and a dropping funnel; add 10 grammes of zinc dust, and without further heating by the flame, gradually allow to flow in from the funnel, with thorough

¹ A. 9, 1585.

² A. 119, 142.

shaking, 10 grammes of benzenesulphonchloride in small portions. After each addition wait until the vigorous reaction accompanied by a hissing sound has moderated. The mixture is then heated a few minutes over a small flame, filtered after cooling from the precipitate of zinc dust and the zinc salt of benzenesulphinic acid, and the precipitate washed several times with water. The insignificant-looking gray precipitate is the reaction-product, and not the filtrate, which can be thrown away. The precipitate is then heated for about ten minutes, not quite to boiling, with a solution of 10 grammes of dehydrated sodium carbonate in 50 c.c. of water, and then filtered with suction. The precipitate remaining on the filter is worthless, while the filtrate contains the sodium benzenesulphinate in solution. This is evaporated to about one-half its original volume, and, after cooling, acidified with dilute sulphuric acid, upon which the free benzenesulphinic acid separates out in colourless crystals; the separation is facilitated by rubbing the sides of the vessel with a glass rod. After filtering, the substance is recrystallised from a little water. Melting-point, $83-84^{\circ}$.

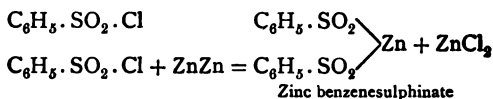
Should the free acid not separate on acidifying the sodium salt, it is extracted several times with ether; this is evaporated, and the residue, in case it does not solidify of itself, is rubbed with a glass rod and then recrystallised.

(b) In order to convert the residue of benzenesulphonchloride obtained in Reaction 19 into thiophenol, it is heated on a water-bath with granulated tin and concentrated hydrochloric acid, in a large flask provided with a long reflux condenser and dropping funnel; the sulphonchloride is allowed to flow in gradually from the dropping funnel. (To 1 part of the chloride use $2\frac{1}{2}$ parts of tin and 5 parts of concentrated acid.) The heating is continued until most of the tin is dissolved. The thiophenol formed is distilled over with steam, extracted with ether, dried over anhydrous Glauber's salt, and, after the evaporation of the ether, rectified. Boiling-point, 173° .

In the preparation of thiophenol, care is taken that there are no flames in the neighbourhood of the flask in which the reaction is conducted, otherwise there may be an explosion of the mixture

of oxygen and hydrogen. Since the thiophenol possesses an extremely unpleasant odour, and the vapours attack the eyes, causing tears, the experiment must not be carried out in the laboratory, but in a side room (hydrogen sulphide room), or in the open air, in the basement, or at least under a hood with a good draught. Further, care must be taken not to allow the substance to come in contact with the skin, since it produces a violent burning.

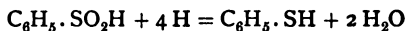
If zinc dust is allowed to act on a sulphonchloride, the zinc salt of the sulphinic acid is formed :



The zinc salts thus formed are insoluble in water, and can be easily obtained by filtering off. In order to prepare the free sulphinic acid from a zinc salt, it is first converted into the easily soluble sodium salt by boiling with a sodium carbonate solution ; the solution of the sodium salt is concentrated, and the free acid is precipitated with dilute sulphuric acid. The sulphinic acids differ from the sulphonic acids in that they are difficultly soluble in cold water, and can, therefore, be recrystallised from water. They are also soluble in ether, in which sulphonic acids do not dissolve. On fusing with potassium hydroxide, the sulphinic acids pass over to the hydrocarbons :



If they are reduced, a thiophenol is finally obtained, as above :

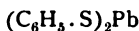


The thiophenols may also be prepared by the direct reduction of the sulphonchlorides :

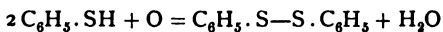


The thiophenols are liquids of unpleasant odours ; the higher members of the series are solids. Like the mercaptans of the aliphatic series, they form difficultly soluble salts with lead and mercury.

EXPERIMENT : Dissolve mercuric chloride or lead acetate in a test-tube with alcohol by heating ; then cool, and filter. If the alcoholic solution is treated with a few drops of thiophenol, a precipitate of the difficultly soluble salt is obtained. The lead salt is yellow, and possesses the composition represented by the formula :

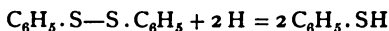


In the air, and on treatment with oxidising agents like nitric acid, chromic acid, iodine, etc., the thiophenols are oxidised to disulphides :

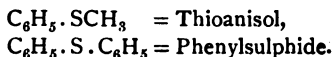


EXPERIMENT : A few drops of phenyl mercaptan are dissolved in alcohol, treated with some ammonia, and evaporated to dryness on the water-bath in a watch-glass. (Under the hood.) Colourless needles of the disulphide remain. Melting-point, 61° .

By reduction the disulphides are easily converted back to the thiophenols :



Like the phenols the thiophenols have the power of forming ethers, *e.g.* :



21. REACTION : SULPHONATION OF AN AROMATIC HYDRO-CARBON (II)

EXAMPLE : β -Naphthalenesulphonic Acid

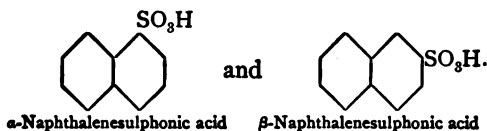
A mixture of 50 grammes of finely pulverised naphthalene and 60 grammes of pure concentrated sulphuric acid is heated in an open flask in an oil-bath for 4 hours to $170-180^\circ$. After cooling, the solution is carefully poured, with stirring, into 1 litre of water, and the naphthalene not attacked is filtered off ; in case the filtration takes place very slowly, only the turbid liquid is poured off from the coarse pieces of naphthalene ; the mixture is

neutralised at the boiling temperature in a large dish with a paste of lime, not too thin, prepared by triturating about 70 grammes of dry slaked lime with water. The mixture is filtered as hot as possible through a filter-cloth, which has been previously thoroughly moistened (see page 61) and the precipitate washed with hot water. The filter-cloth is then folded together and thoroughly squeezed out in another dish; the expressed, generally, turbid liquid, after filtering, is united with the main quantity. The solution is then evaporated in a dish over a free flame until a test-portion will solidify to a crystalline paste on rubbing with a glass rod. After the solution has been allowed to stand over night the calcium β -naphthalenesulphonate is filtered off with suction, washed once with a little water, pressed firmly together with a pestle, and spread out on a porous plate. In order to obtain the sodium salt, it is dissolved in hot water, and the solution gradually treated with a concentrated solution of 50 grammes of crystallised sodium carbonate until a test-portion filtered off no longer gives a precipitate with sodium carbonate. After cooling, the precipitate of calcium carbonate is filtered off with suction, washed with water, and the filtrate evaporated over a free flame until crystals begin to separate from the hot solution. After standing several hours at the ordinary temperature, the crystals are filtered off, and the mother-liquor further concentrated; after long standing, the second crystallisation is filtered off, and the mixture of the two lots of crystals dried on the wafer-bath. Yield, 60–70 grammes.

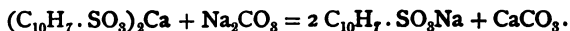
Naphthalene is sulphonated on heating with sulphuric acid, in accordance with the following equation:



There is formed not as in the case of benzene, in which the six hydrogen atoms are equivalent, a single sulphonic acid, but a mixture of two isomeric sulphonic acids:



According to the temperature at which the sulphonation takes place, more of one than of the other acid is formed; at lower temperatures an excess of the α -acid is obtained, at higher an excess of the β -acid. If the mixture is heated to 100° , a mixture of 4 parts of the α -acid and 1 part of the β -acid is formed, while at 170° a mixture of 3 parts of the β -acid and 1 part of the α -acid is obtained. In order to separate the sulphonic acids from the excess of sulphuric acid, advantage is taken of the fact that sulphonic acids differ from sulphuric acid in that they form soluble salts of calcium, barium, and lead, as mentioned under benzenesulphonic acid. For the separation of the sulphonic acid from sulphuric acid, the calcium salt is prepared by neutralising the acid mixture with chalk or lime, since it is cheaper than lead carbonate or barium carbonate. This method is followed technically on the large scale as well as in laboratory preparations. Since the calcium salts of the two isomeric sulphonic acids possess a very different solubility in water, — at 10° 1 part of the α -salt dissolves in 16.5 parts of water, and 1 part of the β -salt dissolves in 76 parts of water, — the β -salt, which is more difficultly soluble, and consequently crystallises out first, can be separated by fractional crystallisation from the α -salt which remains in solution. For the conversion into naphthol the calcium salt cannot be used directly; it must first be changed into the sodium salt by treatment with sodium carbonate:



In order to remove the last portions of the α -salt, it is advisable not to evaporate the solution of sodium salt directly to dryness, but to allow the more difficultly soluble β -salt to crystallise out, upon which the α -salt remains dissolved in the mother-liquor.

The reactions of the naphthalenesulphonic acids are similar to those given above under benzenesulphonic acid. It is still to be mentioned that the α -acid is converted into the β -acid by heating with concentrated sulphuric acid to almost 200° ; a reaction which is explained by the fact that the sulphonic acid decomposes in the small amount of water always present, into naphthalene and sulphuric acid, and that the former is then sulphonated to the β -acid at the higher temperature (200°). The sulphonation of naphthalene to the α - and β -acids is carried out on the large scale in technical operations, since when fused with sodium hydroxide these acids yield naphthols of great importance for the manufacture of dyes. The next preparation deals with this reaction.

22. REACTION: CONVERSION OF A SULPHONIC ACID INTO A PHENOL**EXAMPLE: β -Naphthol from Sodium- β -Naphthalene Sulphonate and Sodium Hydroxide¹**

In order to convert sodium- β -naphthalene sulphonate into β -naphthol the proportions of the necessary reagents used are :

10 parts sodium- β -naphthalene sulphonate ;
30 parts sodium hydroxide, as pure as possible ;
1 part water.

The sodium hydroxide is broken in pieces about a centimetre in length, or the size of a bean, treated with the water, and heated in a nickel crucible (a crucible 11 cm. high and 8 cm. in diameter is a convenient size), with stirring, to 280° (Fig. 73). The stirring is done with a thermometer, the lower end of which is protected by a case of copper or nickel, about 16 cm. long and 8 mm. wide. This is supported by a cork, containing a narrow canal at the side, fitting the case. In order to be able to determine the temperature as exactly as possible, a layer of oil 1 cm. high is placed in the case, in which the bulb of the thermometer is immersed. If the stirring is done with the case, the upper portion is covered with several layers of asbestos board, secured with wire, or a cork is pushed over the case (Fig. 73). Since, on fusion of the sodium hydroxide, a troublesome spattering takes place, the hand is protected by a glove, and the eyes by glasses. As soon as the temperature reaches 280°, the heating is continued with a somewhat smaller flame, and the sodium naphthalene sulphonate is gradually added, with stirring. After each new addition, the temperature falls somewhat ; no more of the salt is added, until the temperature again reaches 280°. After all the salt is added, the flame is made somewhat larger, upon which the fusion becomes viscid with evolution of steam and frothing, until finally, at about 310°, the real reaction takes place. After the temperature is held

¹ E. Fischer, Prep. of Organic Compounds, 7 Ed. page 55. Z. 1867, 299.

at 310–320° for about 5 minutes, the fusion becomes liquid, and the reaction is complete. The melted mass is then poured in a thin layer on a strong copper plate the edges of which have been turned up. The portions of dark sodium naphtholate may be easily distinguished from the brighter caustic soda. After cooling, the solid mass is broken up and dissolved in water. The naphthol is precipitated at the boiling temperature with concentrated hydrochloric acid (under the hood), and after cooling is extracted with ether. The ethereal solution is dried over anhydrous Glauber's salt, and then the ether is evaporated in an apparatus similar to the one described on page 35; a fractionating flask with a very wide condensing tube is used. After the removal of the ether, the naphthol remaining back is distilled over without the use of a condenser. Melting-point, 123°. Boiling-point, 286°. Yield, half the weight of the sulphonate used.

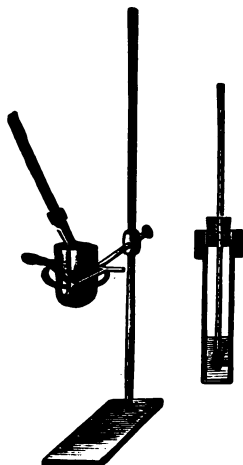
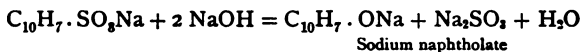


FIG. 73.

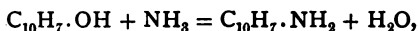
As above indicated, in a sodium hydroxide, or potassium hydroxide fusion of a sulphonic acid, besides the phenol, the alkali sulphite is formed, *e.g.* :



The free phenol is, therefore, not directly obtained on fusion, but the alkali salt of it, from which, after the solution of the fusion in water, the phenol is liberated on acidifying with hydrochloric acid.

The reaction just effected is in practice carried out on the largest scale in iron kettles to which stirring apparatus is attached. β -naphthol as well as its numerous mono- and poly-sulphonic acid derivatives obtained by treatment with sulphuric acid find extensive application for

the manufacture of azo dyes. Further, from the β -naphthol, β -naphthylamine is prepared by the action of ammonia under pressure :

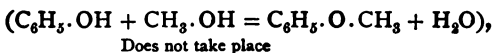


which also finds technical use for the manufacture of azo dyes, as such and in the form of its sulphonic acids. α -Naphthol is also prepared in the same way by fusion of α -sodiumnaphthalene sulphonate with sodium hydroxide, although not in so large quantities as the β -naphthol.

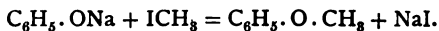
The phenols, in consequence of the negative character of the aromatic hydrocarbon residue, are weak acids which dissolve in water solutions of the alkalis to form salts. Still the acid nature is so weak that the salts can be decomposed by carbon dioxide; use is frequently made of this property for the purification and separation of phenols.

EXPERIMENT: A mixture of β -naphthol and benzoic acid is dissolved in a diluted caustic soda solution, and carbon dioxide passed into it for a long time. β -Naphthol only separates out; this is filtered off. The filtrate is acidified with concentrated hydrochloric acid upon which the benzoic acid is precipitated.

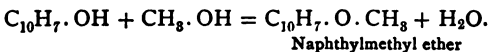
The naphthols differ from the phenols of the benzene series, in that their hydroxyl groups are more capable of reaction than those of the phenols, cresols, etc. While, for example, the ether of phenol cannot be prepared from the phenol and corresponding alcohol by abstracting water :



but can only be obtained by the action of halogen alkyls, or salts of alkyl sulphuric acid on phenol salts :



By heating the naphthols with an aliphatic alcohol and sulphuric acid, the ethers are easily prepared :



23. REACTION: NITRATION OF A PHENOL**EXAMPLE: o- and p-Nitrophenol**

Dissolve 80 grammes of sodium nitrate in 200 grammes of water by heating; after cooling, the solution is treated, with stirring, with 100 grammes of concentrated sulphuric acid. To the mixture cooled to 25° contained in a beaker, add drop by drop, from a separating funnel, with frequent stirring, a mixture of 50 grammes of crystallised phenol and 5 grammes of alcohol, melted by warming. During this addition the temperature is kept between $25-30^{\circ}$ by immersing the beaker in water. Should the phenol solidify in the separating funnel, it is again melted by a short warming in a large flame. After the reaction-mixture has been allowed to stand for two hours with frequent stirring, it is treated with double its volume of water; the reaction-product collects as a dark oil at the bottom of the vessel. The principal portion of the water solution is then decanted from the oil, this is washed again with water, and after the addition of $\frac{1}{2}$ litre of water, is distilled with steam until no more o-nitrophenol passes over. Concerning the removal of the o-nitrophenol solidifying in the condenser, see page 39 (temporary removal of the condenser-water).

After cooling, the distillate is filtered, the o-nitrophenol washed with water, pressed out on a porous plate, and dried in a desiccator. Since it is obtained completely pure, it is unnecessary to subject it to any further process of purification. In order to obtain the non-volatile p-nitrophenol remaining in the flask, the mixture is cooled by immersion in cold water, the water solution is filtered from the undissolved portions, and the filtrate boiled for a quarter-hour with 20 grammes of animal charcoal, the water evaporating being replaced by a fresh quantity. The charcoal is then filtered off and the filtrate allowed to stand in a cool place over night, upon which the p-nitrophenol separates out in long, almost colourless needles. The oil still present in the distillation flask is boiled with a mixture of 1 part by volume of concentrated

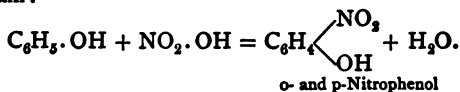
hydrochloric acid and 2 parts by volume of water, with the addition of animal charcoal, filtered after partial cooling and the filtrate allowed to stand over night. There is thus obtained a second crystallisation. If the crystals which have separated out are still contaminated by the oil, they are recrystallised from dilute hydrochloric acid with the use of animal charcoal.

Melting-point of o-Nitrophenol, 45° ;

Melting-point of p-Nitrophenol, 114° .

Yield, 30 grammes and 5–10 grammes respectively.

The mon-acid phenols of the benzene series are, in contrast to the corresponding hydrocarbons, very easily nitrated. In the nitration of benzene, in order to facilitate the elimination of the water, concentrated sulphuric acid must be added; whereas the action of concentrated nitric acid alone upon phenol is so energetic, that in this case it must be diluted with water. Upon nitrating phenol, the o- and p-nitrophenols are formed simultaneously, the former of which is volatile with steam:



On nitrating the homologues of phenol, the nitro-groups always enter the o- and p-positions to the hydroxyl group. In order to prepare m-nitrophenol, it is necessary to start from m-nitroaniline; this is diazotised and its diazo-solution boiled with water.

The nitrophenols behave in all respects like the phenols. But by the entrance of the negative nitro-group, the negative character of the phenol is so strengthened that the nitrophenols not only dissolve in alkalis, but also in the alkali carbonates.

EXPERIMENT: Dissolve some o-nitrophenol in a solution of sodium carbonate by warming; the scarlet red sodium salt is formed.

In consequence of this action, the nitrophenols cannot be precipitated from their alkaline solutions by carbon dioxide.

In addition the nitrophenols show the characteristics of the nitro-compounds in general, since they, for example, pass over to amido-phenols on energetic reduction, etc.

24. REACTION: (a) CHLORINATION OF A SIDE-CHAIN OF A HYDRO-CARBON. (b) CONVERSION OF A DICHLORIDE INTO AN ALDEHYDE

EXAMPLES: (a) Benzalchloride from Toluene

(b) Benzaldehyde from Benzalchloride

(a) A 100 c.c. round flask with a wide neck (Fig. 74) containing 50 grammes of toluene is placed in a well-lighted position, best in the sunlight. The toluene is heated to boiling and a current of dry chlorine conducted into it until its weight

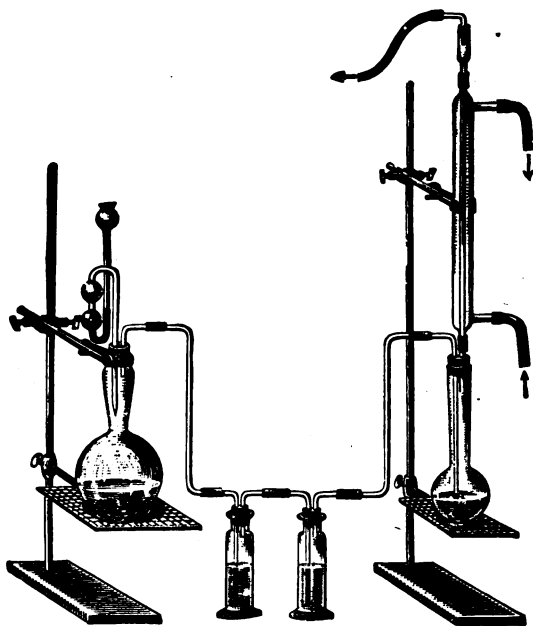


FIG. 74.

is increased by 40 grammes. In order to be able to judge of the course of the reaction, the flask, with the toluene, is weighed before the experiment. By interrupting the passage of the chlorine from time to time, cooling and weighing the flask, the increase in weight will indicate how far the chlorinating action

has proceeded. The length of the operation varies greatly. In summer the reaction is complete in a few hours; during the cloudy days of winter a half or a whole day may be necessary. The reaction may be materially assisted by adding 4 grammes of phosphorus pentachloride to the toluene.

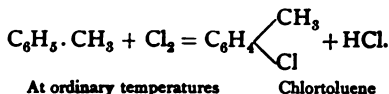
(b) In order to convert the benzalchloride into benzaldehyde, the crude product thus obtained is treated in a round flask provided with an effective reflux condenser, with 500 c.c. of water and 150 grammes of precipitated calcium carbonate (or floated chalk or finely pulverised marble) and the mixture heated four hours in a hemispherical oil-bath to 130° (thermometer in the oil). Without further heating, steam is passed through the hot contents of the flask until no more oil distils over. For this purpose the apparatus necessary (cork with a glass tube) has been prepared before the heating in the oil-bath.

Before the crude benzaldehyde is subjected to purification, the liquid remaining in the distilling flask is filtered while hot through a folded filter, and the filtrate acidified with much concentrated hydrochloric acid. On cooling, the benzoic acid obtained as a by-product in the preparation of benzaldehyde separates out in lustrous leaves. It is filtered off and recrystallised from hot water, during which it must not be heated too long, since it is volatile with steam. Melting-point, 121° .

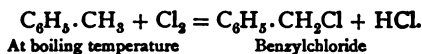
The oil passing over with the steam is treated, together with all of the liquid, with a concentrated solution of sodium hydrogen sulphite, until after long shaking the greater part of the oil has passed into solution. Should crystals of the double compound of benzaldehyde and sodium hydrogen sulphite separate out, water is added until they are dissolved. The water solution is then filtered through a folded filter from the oil remaining undissolved, and the filtrate treated with anhydrous sodium carbonate until it shows a strong alkaline reaction. This alkaline liquid is now subjected to distillation with steam, when perfectly pure benzaldehyde passes over; it is taken up with ether, and, after the evaporation of the ether, is distilled. Boiling-point, 179° .

Under the preparation of brombenzene it has already been mentioned that by the action of chlorine or bromine on aromatic hydrocarbons containing aliphatic side-chains, different products are formed,

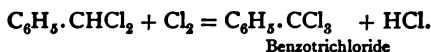
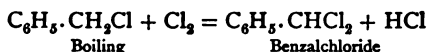
depending on the temperature at which the action takes place. If, *e.g.*, chlorine acts at lower temperatures on toluene, chlortoluene is formed, the chlorine entering the benzene ring :



If, on the other hand, chlorine is conducted into boiling toluene, the chlorine atom enters the side-chain :



If chlorine is conducted into toluene at the boiling temperature for a long time, a second, and finally a third, hydrogen atom of the methyl group is substituted :



The formation of benzotrichloride is the final result of the action of chlorine under these conditions, since the benzotrichloride is not changed even by passing in the chlorine for a longer time.

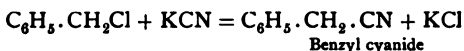
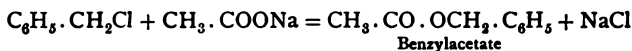
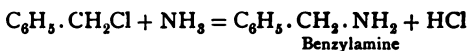
The replacement of hydrogen by chlorine directly presents the difficulty, unlike the liquid bromine, that weighed quantities cannot be used, and the exact point to which the introduction of chlorine should be continued in order to get a certain definite compound must be determined. This is accomplished if, from time to time, the increase in weight of the substance being chlorinated is determined. Since the conversion of one molecule of toluene to benzylchloride requires an increase of weight equal to the atomic weight of chlorine minus the atomic weight of hydrogen ($\text{Cl} - \text{H} = 34.5$), therefore in the preparation of benzylchloride, 100 parts by weight of toluene must take up an additional weight of chlorine = 37.5 parts by weight, and correspondingly in the preparation of benzalchloride or benzotrichloride, the increase in parts by weight must be respectively $2 \times 37.5 = 75$ and $3 \times 37.5 = 112.5$.

In most organic chlorinating reactions, besides the main reaction, a side reaction also takes place which, in the above example, results in the conversion of a portion of the toluene to benzalchloride and

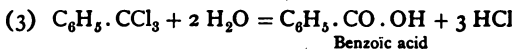
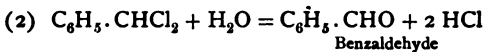
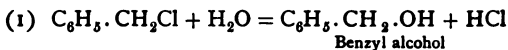
benzotrichloride, while another portion is only chlorinated to benzyl chloride. Accordingly the reaction-product obtained above consists essentially of benzalchloride mixed with a small quantity of benzyl chloride and benzotrichloride.

The halogen derivatives of the aromatic hydrocarbons containing the halogen in the side-chain are in part liquids, in part colourless crystallisable solids, which are distinguished from their isomers containing the halogen in the benzene ring, in that their vapours violently attack the mucous membrane of the eyes and nose. Care is taken therefore in the above preparation not to expose the face to the vapours, and further to prevent the chlorination products from coming in contact with the hands.

Concerning their chemical properties, these two isomeric series differ in that the compounds containing the halogen in the side-chain are far more active than those in which the halogen occurs in the benzene ring, as is apparent from the following equations :



The chlorides obtained by substituting a side-chain are of importance for making certain preparations,— the aromatic alcohols, aldehydes, and acids. On boiling with water, they decompose, in accordance with the following equations :



If the halogen atom is in the ring, none of these transformations will take place. But since, in cases 1 and 2, the hydrochloric acid formed in the reaction acts in the opposite way and may regenerate the original chloride, it must be neutralised. This is usually accomplished by the addition of a carbonate, upon which the acid acts with the liberation of carbon dioxide.

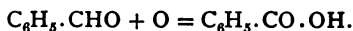
In practice, the cheap calcium carbonate (marble dust) is used, and the above method for the preparation of benzaldehyde is, as far as possible, an imitation of the technical process used for obtaining the substance. From benzylchloride, benzaldehyde may also be prepared directly by boiling it with water in the presence of lead nitrate or copper nitrate. From the benzylchloride, benzyl alcohol is first formed, which is oxidised by the nitrate to benzaldehyde.

As above mentioned, the chlorination product obtained consists essentially of benzalchloride, mixed with small amounts of benzylchloride and benzo-trichloride. If the mixture is boiled with water, with the addition of calcium carbonate, a mixture consisting mainly of benzaldehyde, besides benzylchloride and benzoic acid—the latter being converted into the calcium salt by the carbonate—is obtained. If the reaction-mixture is distilled with steam, benzaldehyde, benzyl alcohol, and a small amount of chlorides which have not taken part in the reaction, pass over with the steam, while the calcium benzoate remains in the distillation flask. By acidifying the residue, the free benzoic acid may be obtained as above. A large proportion of the so-called "benzoic acid from toluene" is obtained in this way as a by-product in the technical preparation of benzaldehyde. In order to separate the benzaldehyde from benzyl alcohol, the chlorides, and other impurities, advantage is taken of the general property of aldehydes of uniting with acid sodium sulphite to form soluble double compounds. If the distillate is shaken with a solution of sodium hydrogen sulphite, the aldehyde dissolves, while the impurities remain undissolved. These are filtered off, and the sulphite compound of the aldehyde decomposed with sodium carbonate; on a second distillation with steam, the pure aldehyde passes over.

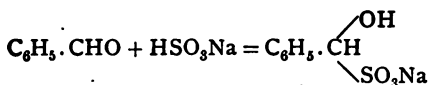
The aromatic aldehydes are in part liquids, in part solids, possessing a pleasant aromatic odour.

They show the characteristic aldehyde reactions, yielding primary alcohols on reduction and carbonic acids on oxidation.

EXPERIMENT: A few drops of benzaldehyde are allowed to stand in a watch-glass in the air. After a long time, crystals of benzoic acid appear.

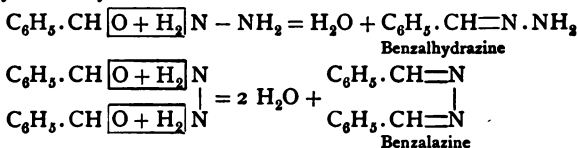


That they unite with acid sulphites to form crystalline compounds has been mentioned:

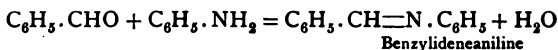


EXPERIMENT: To $\frac{1}{2}$ c.c. of benzaldehyde add a concentrated solution of sodium hydrogen sulphite, and shake. The mixture solidifies after a short time to a crystalline mass.

Further, the aldehydes react, as mentioned under acetaldehyde, with hydroxylamine and phenyl hydrazine, to form oximes and hydrazones. With hydrazine they yield, according to the experiment conditions, the not very stable hydrazines or the more stable and characteristic azines:



Aldehydes also condense readily with primary aromatic bases with the elimination of water:



EXPERIMENT: In a test-tube make a mixture of 1 c.c. of benzaldehyde and 1 c.c. of pure aniline, and warm gently. On cooling, drops of water separate out, and the mixture solidifies, crystals of benzylideneaniline being formed.

An additional number of characteristic aldehyde reactions will be taken up later in practice. Benzaldehyde is prepared technically on the large scale. Its most important application is for the manufacture of the dyes of the Malachite Green series, and of cinnamic acid (see these preparations).

25. REACTION: SIMULTANEOUS OXIDATION AND REDUCTION OF AN ALDEHYDE UNDER THE INFLUENCE OF CONCENTRATED POTASSIUM HYDROXIDE

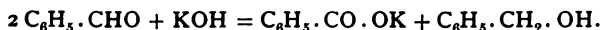
EXAMPLE: Benzoinic Acid and Benzyl Alcohol from Benzaldehyde¹

Treat 20 grammes of benzaldehyde in a stoppered cylinder or a thick-walled vessel with a cold solution of 18 grammes of potassium hydroxide in 12 grammes of water, and shake until a permanent emulsion is formed; the mixture is then allowed to stand over night. The vessel is closed by a cork, and not a glass stopper, since at times a glass stopper becomes so firmly fastened that it can be removed only with great difficulty. To the crys-

¹ B. 14, 2394.

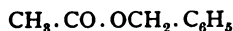
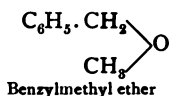
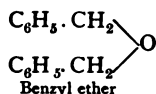
talline paste (potassium benzoate) separating out, water is added until a clear solution is obtained from which the benzyl alcohol is extracted by repeatedly shaking with ether. After the evaporation of the ether the residue is subjected to distillation; benzyl alcohol passes over at 206° . Yield, about 8 grammes. The benzoic acid is precipitated from the alkaline solution on acidifying with hydrochloric acid.

While many aliphatic aldehydes (see acetaldehyde) are converted by alkalis into more complex compounds, with higher molecular weights, the so-called aldehyde resins, the aromatic aldehydes under similar conditions react smoothly; two molecules are decomposed by one molecule of potassium hydroxide, one aldehyde molecule being oxidised to the corresponding acid, and the other being reduced to a primary alcohol:



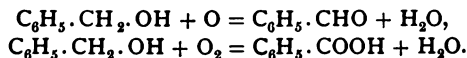
Since the aldehydes are in part easily obtained, the different primary alcohols may be prepared advantageously by this reaction.

The primary aromatic alcohols behave in all respects like the corresponding aliphatic alcohols in forming ethers and esters, *e.g.*:



Aceticbenzyl ester

On oxidation they are converted first into aldehydes and finally into acids:



26. REACTION: CONDENSATION OF AN ALDEHYDE BY POTASSIUM CYANIDE TO A BENZOIN

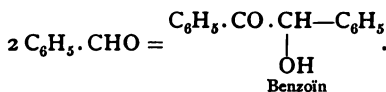
EXAMPLE: Benzoïn from Benzaldehyde¹

Mix 10 grammes of benzaldehyde with 20 grammes of alcohol and treat the mixture with a solution of 2 grammes of potassium cyanide and 5 c.c. of water. Boil on the water-bath for one hour (reflux condenser). The hot solution is poured into a beaker and allowed to cool slowly; the crystals separating out are filtered

¹ A. 198, 150.

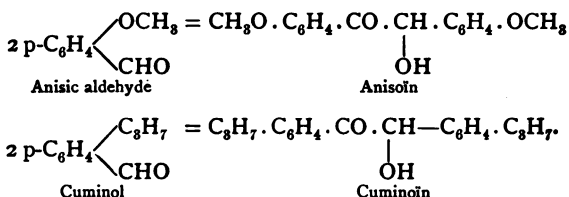
off, washed with alcohol, and dried on the water-bath. For conversion into benzil (see next preparation), they need not be recrystallised. In order to obtain perfectly pure benzoïn, a small portion of the crude product is recrystallised from a little alcohol in a test-tube. Melting-point, 134° . Yield, about 90% of the theory.

If an aromatic aldehyde of the type of benzaldehyde is warmed in alcohol solution with a small quantity of potassium cyanide, substances are obtained which possess the same composition, but with double the molecular weight of the aldehyde:

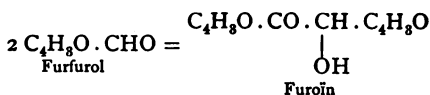


This is unlike aldol formation, since here condensation takes place between the two aldehyde groups (p. 174).

In the same way from anisic aldehyde and cuminol, there are obtained anisoïn and cuminoïn, respectively:

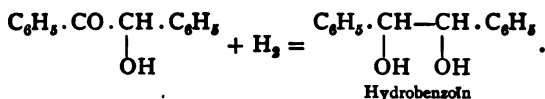


With potassium cyanide furfural yields furoïn:

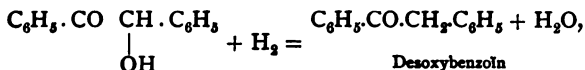


Benzoïn and its analogues are derivatives of the hydrocarbon dibenzyl, $\text{C}_6\text{H}_5 \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{C}_6\text{H}_5$, and in fact benzoïn on reduction with hydriodic acid is converted into this hydrocarbon.

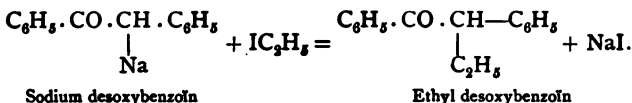
The benzoïns act, on the one hand, like ketones if the carbonyl group (CO) takes part in the reaction, and, on the other hand, like secondary alcohols if the group CH.OH (the secondary alcohol group) reacts. Thus they have the power to form oximes and hydrazones with hydroxylamine and phenyl hydrazine respectively. If benzoïn is reduced with sodium amalgam, the ketone group is converted into the secondary alcohol group.



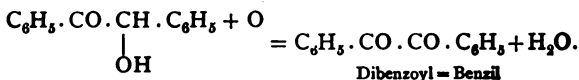
If the reduction is effected by zinc and hydrochloric acid or glacial acetic acid, the carbonyl group is not attacked, but the alcohol group is reduced and desoxybenzoïn is obtained :



a compound of especial interest, because in it, as in acetacetic ester, one of the two hydrogen atoms of the methylene group (CH_2), in consequence of the acidifying influence of the adjoining negative carbonyl and phenyl groups, may be replaced by sodium ; with the sodium compound the same kind of syntheses may be effected as with acetacetic ester :



Benzoïn, further, acts as an alcohol, the hydroxyl group being capable of reacting with alkyl- and acid-radicals to form ethers and esters. If oxidizing agents act on benzoïn, the alcohol group is oxidized to a ketone group, as is the case with all secondary alcohols :



The next preparation will deal with this reaction.

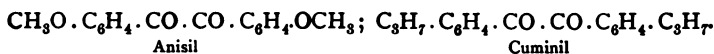
27. REACTION: OXIDATION OF A BENZOÏN TO A BENZIL

EXAMPLE: Benzil from Benzoïn

The crude benzoïn obtained in the preceding preparation is finely pulverised after drying, and heated in an open flask, with frequent shaking, with twice its weight of pure concentrated nitric acid, for $1\frac{1}{2}$ –2 hours on a rapidly boiling water-bath. When the oxidation is ended, the reaction-mixture is poured into cold water ;

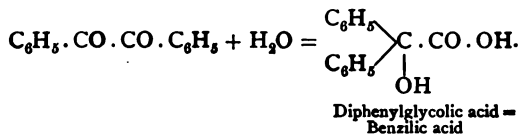
after the mass solidifies the nitric acid is poured off; it is then washed several times with water, pressed out on a porous plate, and crystallised from alcohol. After filtering off the crystals separating out, they are dried in the air on several layers of filter-paper. Melting-point, 95° . Yield, about 90% of the theory.

The equation representing the oxidation of benzoïn to benzil has been given under the preceding preparation. The analogues of benzoïn also give, on oxidation, compounds of the benzil series. Thus from anisoïn and cuminoïn, anisil and cuminil respectively are obtained:



Benzil acts like a ketone in that it forms oximes with hydroxylamine. The oximes are of exceptional interest, since our knowledge of the stereochemistry of nitrogen proceeds from them. Benzil forms *two* monoximes and *three* dioximes. The constitution of these compounds will be discussed later, under the preparation of benzophenone-oxime.

On fusion with potassium hydroxide or by long heating with a water solution of potassium hydroxide, benzil undergoes a remarkable change, in that by taking up water it passes over to the so-called benzilic acid:



Anisil and cuminil also yield, in a similar way, anisilic and cuminilic acids.

28. REACTION: THE ADDITION OF HYDROCYANIC ACID TO AN ALDEHYDE

EXAMPLE: Mandelic Acid from Benzaldehyde¹

(a) *Mandelic Nitrile*

In a flask containing 13 grammes of finely pulverised 100% potassium cyanide, or an equivalent amount of the purest salt

¹ B. 14, 235

available, pour 20 grammes of freshly distilled benzaldehyde, and add to this from a separating funnel, the flask being cooled with ice, a quantity of the most concentrated hydrochloric acid, corresponding to 7 grammes of anhydrous hydrochloric acid (about 20 grammes concentrated acid), drop by drop, carefully, under a hood, with frequent shaking. The reaction-mixture is then allowed to stand, with frequent shaking, for one hour, then poured into about 5 volumes of water, the oil washed with water several times, and finally separated in a dropping funnel. Owing to the ease with which the nitriles decompose, a further purification is not possible. Yield, almost quantitative.

Much better results are obtained by preparing mandelic nitrile¹ thus: Pour 50 c.c. of a concentrated solution of sodium bisulphite over 15 grammes of benzaldehyde in a beaker; stir the mixture

with a glass rod until the addition product $\text{C}_6\text{H}_5\cdot\text{C}\begin{array}{l} \text{H} \\ \text{—} \\ \text{SO}_3\text{Na} \end{array}$

solidifies to a pasty mass. It is then filtered with suction, pressed firmly together, and washed once with a *little* water. The double compound is stirred up with water to a thick paste, and treated with a cold solution of 12 grammes potassium cyanide in 25 grammes of water. After a short time, very easily on stirring, the crystals go into solution, and the nitrile appears as an oil, which is separated from the solution in a dropping funnel.

(b) *Saponification of the Nitrile*

The nitrile is mixed with four times its volume of concentrated hydrochloric acid in a porcelain dish, and evaporated on the water-bath until crystals begin to separate out on the upper surface of the liquid.

The reaction-mixture is then allowed to stand over night in a cool place; the crystals separating out, are triturated with a little water, filtered off with suction, and then washed with not too much water. A further quantity of the acid may be obtained by

¹ Ch. Z. 1896, 90.

extracting the filtrate with ether; after evaporating off the ether, the residue is heated in a watch-crystal some time on a water-bath. The crude mandelic acid is pressed out on a porous plate, and is obtained pure by recrystallising it from benzene. Melting-point, 118° . Yield, about 10–15 grammes.

(c) *Separation of the Inactive Mandelic Acid into its Active Components*¹

A mixture of 20 grammes of crystallised cinchonine, 10 grammes of crystallised mandelic acid, and 500 c.c. of water is heated with quite frequent shaking in an open flask for an hour on an actively boiling water-bath. After cooling, the portions undissolved are filtered off, and are *not* washed. To this clear solution (a) add a few crystals of d-cinchonine mandelate (see below), and allow it to stand, according to the conditions, one or more days in a cool place ($6-8^{\circ}$; in summer in a refrigerator, in winter in a cellar if necessary). In order to purify the crude d-cinchonine mandelate separating out, it is filtered off (filtrate A), pressed out on a porous plate, and recrystallised from water, using for each gramme of the dried salt 25 c.c. of water (heating as above described for an hour, with quite frequent shaking, in an open flask upon a water-bath). On filtering the cold solution the undissolved portions remaining are *not* washed. If the solution be seeded with a few crystals of d-cinchonine mandelate and allowed to stand under the same conditions referred to above, a purer salt will crystallise out. To obtain the free dextro-mandelic acid the purified salt is dissolved in not too much water, and then treated with a slight excess of ammonia, which precipitates the cinchonine; this is filtered off, and, after recrystallisation from diluted alcohol, may be used for other experiments. The filtrate, which contains dextro-ammonium mandelate, is acidified with hydrochloric acid and extracted with ether. If the residue obtained on evaporating the ether be heated in a watch-glass some time on a water-bath, then, on cooling, crystals of dextro-mandelic acid

¹ B. 16, 1773; 32, 2385.

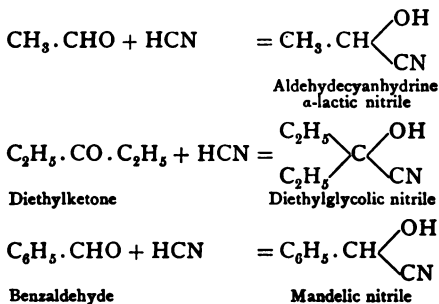
separate out ; they are pressed out on a porous plate and recrystallised from benzene. Melting-point, $133-134^{\circ}$.

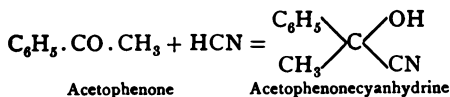
The pure *lævo*-mandelic cannot be obtained readily from small quantities of mandelic acid ; but a preparation showing to a slight extent *lævo*-rotatory power may be obtained in the following way : The filtrate A is worked up for the free acid exactly as in the method described for pure dextro-cinchonine mandelate ; since a portion of the d-modification has been removed from the solution, it should be *lævo*-rotatory.

From the three preparations thus obtained, viz. inactive mandelic acid, the pure d-acid, and the impure l-acid, water solutions of the proper concentration are prepared, and their properties investigated by a polariscope (consult text-books on Physics).

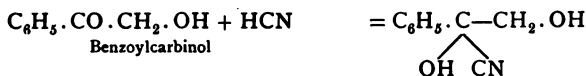
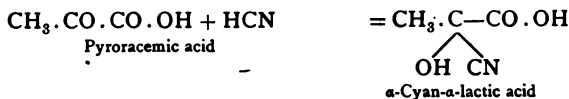
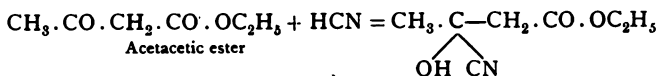
If one is not in possession of d-cinchonine mandelate for the first experiment, a proper seeding material is prepared as follows : To a few cubic centimetres of solution (a) obtained above is added, drop by drop, a saturated solution of salt until a slight precipitation takes place. The solution is now heated until the precipitate redissolves, and is allowed to stand until crystals separate out, which may require several days. The crystals thus obtained are those of cinchonine hydrochloride upon which small quantities of d-cinchonine mandelate have been deposited ; the latter are in sufficient quantity, however, to cause a further separating out of the d-salt.

Hydrocyanic acid unites with aromatic as well as aliphatic aldehydes and ketones with the formation of α -oxyacid nitriles :



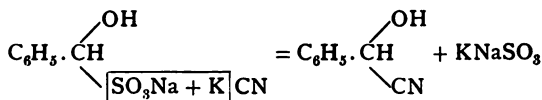


This reaction also takes place with more complex compounds containing the carbonyl group :

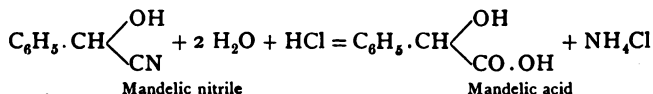


The reaction may be effected by digestion with already prepared hydrocyanic acid at ordinary or higher temperatures, but in most cases it is more advantageous to employ nascent hydrocyanic acid as above.

If the second method be followed, — treating the aldehyde with sodium bisulphite, — the reaction takes place in accordance with the following equation :



If the oxynitriles are subjected to saponification, for example, by boiling with hydrochloric acid, the free oxyacid is obtained, *e.g.* :

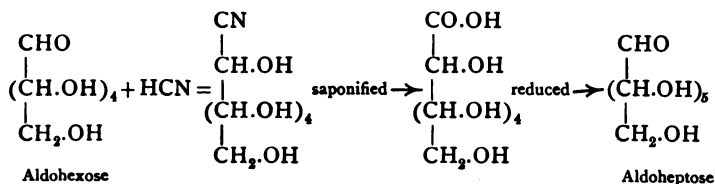


Since the cyanhydrine reaction takes place smoothly in most cases, it is frequently used for the preparation of α-oxyacids.

Thus in the sugar group the cyanhydrine reaction is of extreme importance, not only for its value in determining constitution, but also for the syntheses of sugars or sugar-like substances containing long chains of carbon atoms.

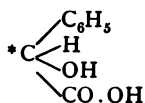
In reference to the latter, one example may be mentioned. If hydro-

cyanic acid is united with grape sugar, which is an aldehyde, there is first obtained an oxynitrile, which on saponification yields an oxyacid. If this, or rather the inner anhydride (lactone) into which it easily passes, is reduced, the carboxyl group is reduced to an aldehyde group, and there is thus obtained a sugar containing one more secondary alcohol (CHOH) group than the original grape sugar :



With the substance thus obtained a similar reaction may be carried out, and so on.

Mandelic acid belongs to the class of substances containing an asymmetric carbon atom, *i.e.*, one which is in combination with four different substituents :



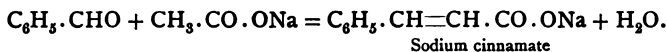
Like all compounds of this class, it exists in two different space modifications, which bear the same relation to each other as does an object and its image, and owing to their power of revolving the plane of polarisation, are called dextro- and lævo-mandelic acids. The acid obtained in the above synthesis is optically inactive; since, in the synthesis of compounds with an asymmetric carbon atom from inactive substances, an equal number of molecules of the dextro- and lævo-varieties are always obtained, which, in the above case, unite to form the inactive, so-called, para-mandelic acid. But, by different methods, the active acids can be obtained from the inactive modifications. If, *e.g.*, the cinchonine salt of para-mandelic acid is allowed to crystallise, the more difficultly soluble salt of the dextro-acid separates out first, and then, later, the lævo-salt crystallises.

With the aid of certain micro-organisms, the inactive compounds may be decomposed into their active constituents. If, *e.g.*, the well-known *Penicillium glaucum* is allowed to grow in a solution of ammonium para-mandelate, it destroys the lævo-modification; while another organism, *Saccharomyces ellipsoideus*, consumes the dextro-modification, and leaves the other.

29. REACTION: PERKIN'S SYNTHESIS OF CINNAMIC ACID¹**EXAMPLE : Cinnamic Acid from Benzaldehyde and Acetic Acid**

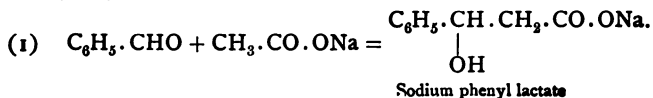
A mixture of 20 grammes of benzaldehyde, 30 grammes of acetic anhydride, both freshly distilled, and 10 grammes of anhydrous pulverised sodium acetate (for the preparation, see page 147), is heated in a flask provided with a wide vertical air-condenser about 60 cm. long, for 8 hours, in an oil-bath at 180°. If the experiment cannot be completed in one day, a calcium chloride tube is placed in the upper end of the condenser over night. After the reaction is complete, the hot reaction-product is poured into a large flask; add water, and then distil with steam, until no more benzaldehyde passes over. The quantity of water used here is large enough so that all of the cinnamic acid dissolves except a small portion of an oily impurity. The solution is then boiled a short time, with some animal charcoal, and filtered; on cooling, the cinnamic acid separates out in lustrous leaves. Should it not possess the correct melting-point, it is recrystallised from hot water. Melting-point, 133°. Yield, about 15 grammes.

The reaction involved in the Perkin synthesis takes place in accordance with this equation :



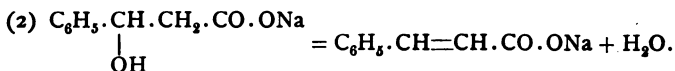
The reaction, however, does not take place, as appears from the equation, by the direct union of the aldehyde-oxygen atom with the hydrogen atoms of the methyl group and a combination of the resulting residues, but it proceeds in two phases.

In the first, the sodium acetate unites with the aldehyde, forming sodium phenyl lactate :

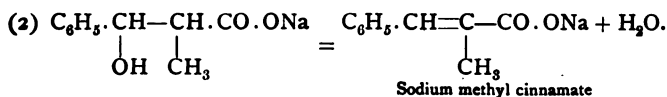


¹ J. 1877, 789; B. 10, 68; 16, 1436; A. 227, 48.

In the second phase, this salt, under the influence of acetic anhydride, loses water, upon which the sodium cinnamate is formed :

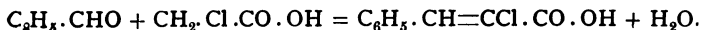


That sodium acetate, and not the acetic anhydride, condenses with the benzaldehyde, is proved by the following facts : If, instead of sodium acetate, sodium propionate is used, and this is heated with benzaldehyde and acetic anhydride, cinnamic acid is not obtained, but methyl cinnamic acid :

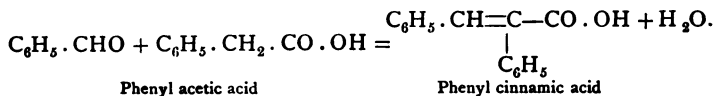


It follows from this that the sodium salt used always takes part in the reaction. In the experiment it is of course necessary that the fusion is not carried out at so high a temperature as in the above example, but only at the heat of the water-bath ; at higher temperatures the sodium salt of propionic acid and acetic anhydride decompose into sodium acetate and propionic anhydride, so that cinnamic acid is obtained, and therefore, *apparently*, the anhydride reacts with the aldehyde.

The Perkin reaction is capable of numerous modifications, since in place of benzaldehyde, its homologues, its nitro- and oxy-derivatives, etc., may be used. On the other hand, the homologues of sodium acetate may be used as has been pointed out. The condensation in these cases always takes place at the carbon atom adjoining the carboxyl group. Halogen substituted aliphatic acids will also react ; thus from benzaldehyde and chloracetic acid, chlorcinnamic acid is obtained :

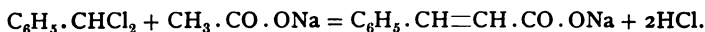


In place of the aliphatic homologues of acetic acid the aromatic substituted acetic acids can also be used, *e.g.* :

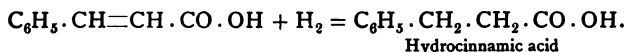


These examples are sufficient to show the wide application of the Perkin reaction.

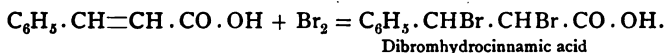
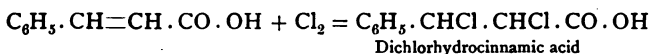
A very similar reaction takes place on heating sodium acetate with the cheaper benzalchloride, instead of benzaldehyde:



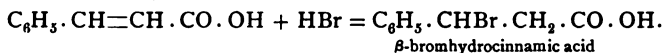
Cinnamic acid, its homologues and analogues, behave on the one hand like acids, since they form salts, esters, chlorides, amides, etc. Further, they show the properties of the ethylene series in that they take up by addition the most various kinds of atoms and groups. By the action of nascent hydrogen two atoms of hydrogen are added to the molecule of cinnamic acid with a change from double to single union:



It also combines with chlorine and bromine:

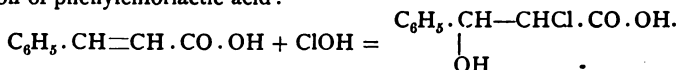


Further, it unites with hydrochloric, hydrobromic, and hydriodic acids, *e.g.*:

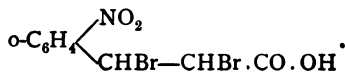


The halogen atom in these cases always unites with the carbon atom not adjoining the carboxyl group.

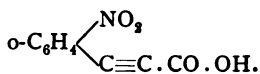
Hypochlorous acid also unites with cinnamic acid with the formation of phenylchlorlactic acid:



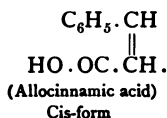
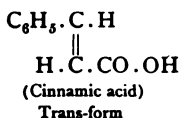
The *o*-nitrocinnamic acid from which indigo is synthetically prepared is of technical importance. If cinnamic acid, or better, an ester of it, is nitrated, a mixture of the *o*- and *p*-nitroderivatives is obtained which can be separated into its constituents. If bromine is allowed to act on the *o*-nitrocinnamic acid, there is obtained:



If this acid is boiled with alcoholic potash, two molecules of hydrobromic acid are split off as in the preparation of acetylene from ethylene bromide, and o-nitrophenylpropionic acid is formed, which, with alkaline reducing agents, yields indigo, and is used in indigo printing :



Cinnamic acid is known in two stereoisomeric forms :



30. REACTION: ADDITION OF HYDROGEN TO AN ETHYLENE DERIVATIVE

EXAMPLE: Hydrocinnamic Acid from Cinnamic Acid

In a glass-stoppered cylinder, or a thick-walled preparation glass, treat 10 grammes of cinnamic acid with 75 c.c. of water ; add a *very dilute solution* of caustic soda until the acid passes into solution and the liquid is *just alkaline*. If a precipitate of sodium cinnamate separates out at this point, too much caustic soda has been used. It is then treated gradually with about 200 grammes of 2 % sodium amalgam, and heated gently, as soon as this has become liquid, on the water-bath for a short time. The liquid is then decanted from the mercury and acidified with hydrochloric acid, upon which the hydrocinnamic acid separates out as an oil ; when cooled with ice-water and rubbed with a glass rod, it solidifies to a crystalline mass. After pressing it out on a porous plate, the acid is recrystallised from water. Since it possesses a low melting-point, it may separate out as an oil on cooling, in which case proceed according to the directions given on page 8. Melting-point, 47°. Yield, 8–9 grammes.

The equation for the reaction has been given under cinnamic acid. The same reaction also takes place on heating with hydriodic acid and red phosphorus.

31. REACTION: PREPARATION OF AN AROMATIC ACID-CHLORIDE FROM THE ACID AND PHOSPHORUS PENTACHLORIDE**EXAMPLE: Benzoyl Chloride from Benzoic Acid¹**

Treat 50 grammes of benzoic acid in a dry $\frac{1}{2}$ -litre flask, with 90 grammes of finely pulverised phosphorus pentachloride under the hood; the two are shaken well together, upon which, after a short time, reaction takes place with energetic evolution of hydrochloric acid, and the mass becomes liquid. In order to prevent the vessel, which has become strongly heated by the reaction, from cracking, it is not placed on the cold stone floor of the hood, but on a wooden block or straw ring. After standing a short time, it is gently heated on a water-bath, and the completely liquid mixture is twice fractionated (under the hood) with the use of an air condenser, observing the directions given on pages 24 and 25. Boiling-point of benzoyl chloride, 200° . Yield, 90 % of the theory.

The formation of benzoyl chloride takes place in accordance with the following reaction:



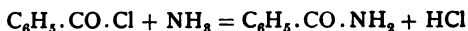
It has been mentioned under acetyl chloride that, for the preparation of the aromatic acid-chlorides, phosphorus pentachloride is generally used. Benzoyl chloride differs from acetyl chloride in that it is more difficultly decomposed by water.

EXPERIMENT: Treat $\frac{1}{2}$ c.c. of benzoyl chloride with 5 c.c. of water and shake. While acetyl chloride, under these conditions, decomposes violently, the benzoyl chloride is scarcely changed. It is then warmed somewhat. It must be subjected to a longer heating before all the oil has been decomposed.

In other respects, benzoyl chloride is a wholly normal acid-chloride, and what was said under acetyl chloride is applicable to this chloride; only it is possible to prepare aromatic amides by a different method from that used for the preparation of acetamide.

¹ A. 3, 262. Ostwald's Klassiker der exakten Wissenschaften, Nr. 22. (Investigations concerning the Radical of Benzoic Acid, by Wöhler and Liebig.)

EXPERIMENT: In a porcelain dish, 10 grammes of finely pulverised ammonium carbonate are treated with 5 grammes of benzoyl chloride; they are intimately mixed with a glass rod and heated on the water-bath until the odour of the acid-chloride has vanished. The mixture is then diluted with water, filtered with suction, washed with water, and crystallised from water. Melting-point of benzamide, 128°.

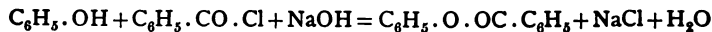


32. REACTION: THE SCHOTTEN-BAUMANN REACTION FOR THE RECOGNITION OF COMPOUNDS CONTAINING THE AMIDO-, IMIDO-, OR HYDROXYL-GROUP

EXAMPLE: Benzoicphenyl Ester from Phenol and Benzoylchloride¹

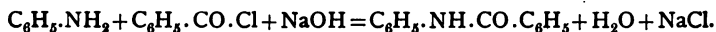
Dissolve a small quantity of crystallised phenol (about $\frac{1}{2}$ gramme) in 5 c.c. of water in a test-tube and add $\frac{1}{2}$ c.c. of benzoyl chloride; make the solution alkaline with a solution of caustic soda and, with shaking, heat gently a short time over a free flame. If the reaction-mixture is cooled by water and then shaken and the sides of the tube rubbed with a glass rod, the oil separating out solidifies to colourless crystals, which are filtered off with suction, washed with water, pressed out on a porous plate, and recrystallised in a small test-tube from a little alcohol. Melting-point, 68–69°.

As already mentioned under acetyl chloride, acid-chlorides react with alcohols, phenols, primary and secondary amines, the chlorine atom uniting with the hydrogen of the hydroxyl-, amido-, or imido-group, with the elimination of hydrochloric acid, while the residues combine to form an ester or a substituted amide. The value of the Schotten-Baumann reaction depends on the fact that this reaction is so essentially facilitated by the presence of sodium hydroxide or potassium hydroxide, that even in the presence of water the decomposition takes place.

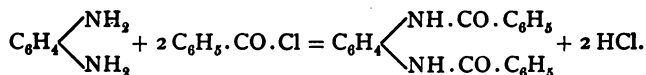
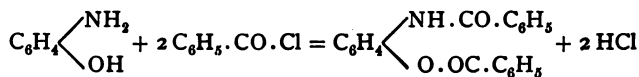


¹ B. 19, 3218; 21, 2744; 23, 2962; 17, 2545.

The reaction is of great importance, especially for the recognition and characterisation of soluble compounds containing the groups mentioned above. It is obvious that if it is desired to test even small quantities of those compounds, the most difficultly soluble acid derivatives of them must be prepared. The benzoyl derivatives are particularly well adapted to this purpose. A few examples may render this statement clearer: If a water solution of a poly-acid aliphatic alcohol, *e.g.*, glycerol, or of the various sugars, from which the dissolved substance will only separate with difficulty, is treated with benzoyl chloride and alkali, a benzoate is formed, which is generally insoluble in water, and which can be recognised by its melting-point. For the recognition of primary and secondary amines the method of procedure is the same. Thus, *e.g.*, it is not difficult to convert aniline (one drop dissolved in water) by the above method to benzanilide, which can be recognised by its melting-point, 163° . (Try the experiment.)



The soluble amido-phenols, di- and poly-amines are also converted into difficultly soluble benzoyl derivatives:



In place of benzoyl chloride, other chlorides, *e.g.*, phenylacetyl chloride, or benzenesulphon chloride, can be used, which act in a similar way. Acetyl derivatives may also be prepared in the presence of alkalies in water solution, only in this case acetic anhydride and not the easily decomposed acetyl chloride is used. At times the reaction takes place better by using potassium hydroxide, or pyridine, in place of sodium hydroxide.

- 33. REACTION:** (a) **FRIEDEL AND CRAFTS' KETONE SYNTHESIS**¹
(b) **PREPARATION OF AN OXIME**
(c) **BECKMANN'S TRANSFORMATION OF AN OXIME**

EXAMPLE: **Benzophenone from Benzoylchloride, Benzene and Aluminium Chloride**

(a) To a mixture of 30 grammes of benzene, 30 grammes of benzoyl chloride, and 100 c.c. (= 130 grammes) of carbon disulphide in a dry flask, add, in the course of about 10 minutes, with frequent shaking, 30 grammes of freshly prepared and finely pulverised aluminium chloride, which is weighed in a dry test-tube closed by a cork. The flask is then connected with a long reflux condenser, and heated on a gently boiling water-bath (a water-bath heated to 50° is better) until only small amounts of hydrochloric acid are evolved: this will require about 2-3 hours. The carbon disulphide is then distilled off, and the residue, while still warm, is carefully poured into a large flask containing 300 c.c. of water and small pieces of ice. The residue adhering to the walls of the first flask is treated with water, and the water added to the main quantity. After the reaction-mixture has been treated with 10 c.c. of concentrated hydrochloric acid, steam is passed into it for about a quarter-hour. The residue remaining in the flask is, after cooling, extracted with ether, the ethereal solution washed several times with water, filtered, and shaken up with dilute caustic soda solution. After drying with calcium chloride, the ether is evaporated, and the residue distilled from a fractionating flask, the side-tube of which is as near as possible to the bulb. Boiling-point, 297°. Melting-point, 48°. Yield, about 30 grammes.

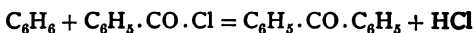
(b) A solution of 2 grammes of benzophenone in 15 c.c. of alcohol is, with cooling, treated with a cold solution of 1.5 grammes of hydroxylamine hydrochloride in 5 c.c. of water, and 3.5 grammes of caustic potash in 6 grammes of water; the mixture is heated two hours on the water-bath, with a reflux condenser. Then add 50 c.c. of water, and filter off, if necessary, any un-

¹ A. ch. [6] 1, 518.

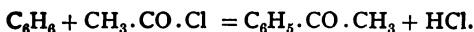
changed ketone which balls together very easily on shaking; acidify the filtrate slightly with dilute sulphuric acid, and recrystallise the free oxime from alcohol. Melting-point, 140° .

(c) A weighed amount of the oxime is dissolved in some anhydrous, alcohol-free ether, at the ordinary temperature, and gradually treated with $1\frac{1}{2}$ times its weight of finely pulverised phosphorus pentachloride. The ether is then distilled off, the residue, with cooling, is treated with water, and the precipitate separating out is recrystallised from alcohol. Melting-point, 163° .

(a) If an aromatic or an aliphatic acid-chloride is allowed to act on an aromatic hydrocarbon in the presence of an anhydrous aluminium chloride, one of the benzene-hydrogen atoms will be replaced by an acid radical, a *ketone* being formed:

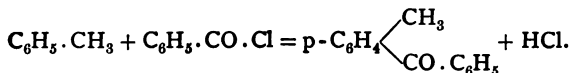


Diphenyl ketone
=Benzophenone



Phenylmethyl ketone
=Acetophenone

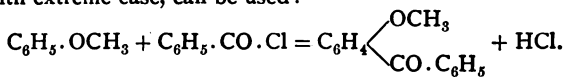
The reaction may be varied if (1) in place of benzene a homologue is used:



Toluene

Phenyltolyl ketone

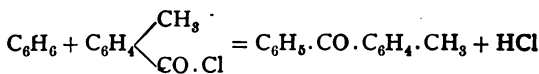
In cases of this kind, the acid-radical always enters the para position to the alkyl radical. If this is already occupied, it then goes to the ortho position. (2) In place of hydrocarbons, phenol-ethers, which react with extreme ease, can be used:



Anisol

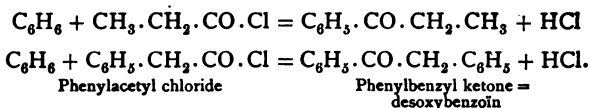
Anisylphenyl ketone

Concerning the entrance of the acid-radical, the statements made above are also true for this case. (3) In place of benzoyl chloride or acetyl chloride, their homologues can be used:

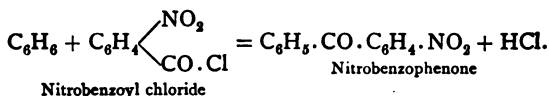
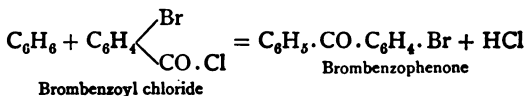


Toluyyl chloride

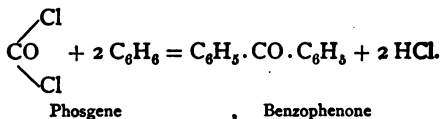
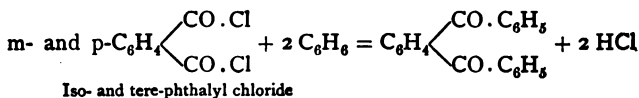
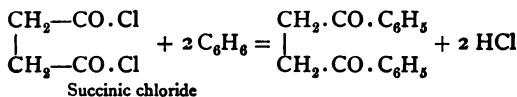
γ



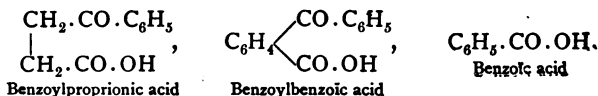
In this way, starting from o- or m-toluic acid, the o- or m-tolyl-phenyl ketone can be prepared; it cannot be obtained by the action of benzoyl chloride on toluene. (4) Substituted acid-chlorides like brombenzoyl chloride, nitrobenzoyl chloride, etc., can be used, and thus halogen or nitroketones are obtained:



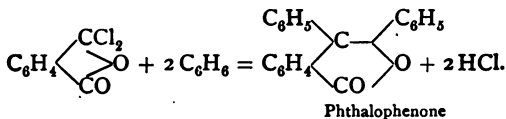
(5) Finally, the chlorides of dibasic acids react with the formation of diketones or ketonic acids:



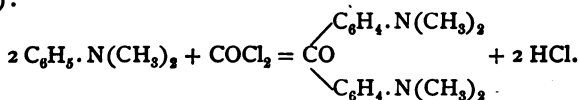
In these reactions if but one chlorine atom should react, the chlorides of the three following acids would be obtained:



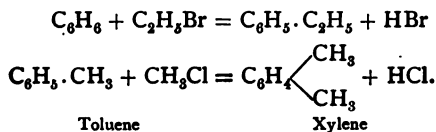
From the chloride of phthalic acid phthalophenone is formed, important on account of its relation to the fluorescein dyes:



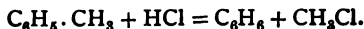
Michler's ketone, tetramethyldiamidobenzophenone is of technical importance; it is obtained from dimethyl aniline and phosgene, and is used in the preparation of dyes of the fuchsine series (see Crystal Violet):



The Friedel-Crafts reaction can also be used for the preparation of the homologous aromatic hydrocarbons, since in place of the acid-chloride, halogen alkyls may be caused to act on the hydrocarbons:¹

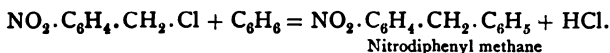
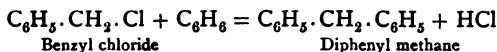


But in this connection the reaction is in many cases, and indeed in the simplest case, not of equal importance with its application for the ketone syntheses, for three reasons: First, the product of the reaction is a hydrocarbon which can again react; thus it is often difficult to limit the reaction to the desired point. For example, in the action of methyl chloride on toluene, not only is one hydrogen atom substituted, with the formation of dimethyl benzene, but varying quantities of tri-, tetra-, penta-, and hexa-methyl benzene are also formed. A second disadvantage is this: In the different series a mixture of isomers is obtained; in the above case, *e.g.*, not only one of the three dimethyl benzenes, but a mixture of the *o*-, *m*-, and *p*-varieties is formed, which cannot be separated like the homologues by fractional distillation. The reaction is still further complicated in that the aluminium chloride partially splits off the alkyl groups:

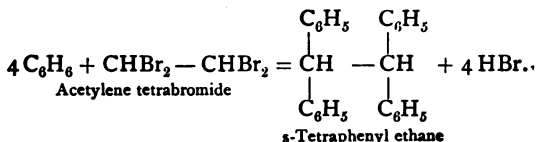
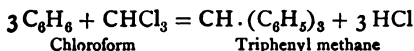
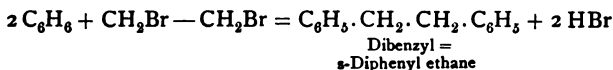


¹ B. 14, 2627.

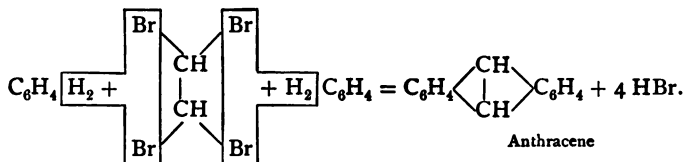
Since the lower homologues thus formed again react synthetically with the halogen alkyls, and the halogen alkyls on elimination also take part in the reaction, mixtures often difficult to separate are formed. In some favourable cases the reaction is of use in the preparation of the homologues of benzene. The reaction is also applicable to aromatic chlorides which contain the halogen in the side-chain :



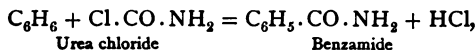
As the chlorides of dibasic acids yield diketones, the alkylene chlorides or bromides, as well as tri- and tetra-halogen substituted hydrocarbons, can react with several hydrocarbon molecules, *e.g.* :

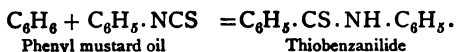
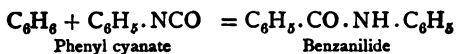


In the latter reaction, anthracene is also formed, according to the equation :

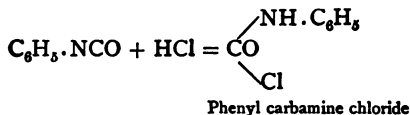


For the synthesis of aromatic acids the Friedel-Crafts reaction is also of value, although the acids themselves are not directly obtained, but derivatives of them, which upon saponification yield the free acid, *e.g.* :





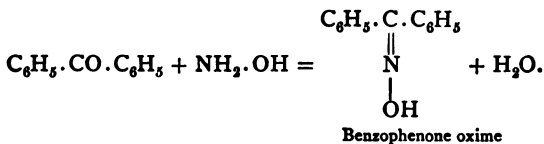
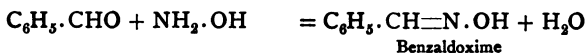
The last two reactions are to be considered as cases of the normal Friedel-Crafts reaction, since the cyanate and mustard oil unite in the first phase with hydrochloric acid, forming an acid-chloride, which then reacts with the hydrocarbon with elimination of hydrochloric acid, *e.g.* :



If one considers that in the modifications, in place of the hydrocarbons, ethers, mono- and poly-acid phenols, naphthalene, thiophene, diphenyl, naphthol-ethers, halogen substitution products of hydrocarbons, and many other compounds can be used, the great value of the Friedel-Crafts reaction will be readily understood.

Concerning the role which aluminium chloride plays in the reaction, it is still not perfectly clear; certain it is that hydrocarbons as well as phenol-ethers unite with it to form double compounds which are of assistance in causing the reaction to take place.

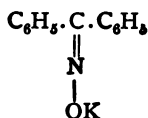
(b) By the action of hydroxylamine on aldehydes and ketones, oximes¹ (aldoximes, ketoximes) are formed in accordance with the following typical reactions :



Oximes may be obtained by three methods: (1) The alcoholic solution of the aldehyde or ketone may be treated, generally, with a concentrated water solution of hydroxylamine hydrochloride and the mixture allowed to stand at the ordinary temperature, or it may be heated in a flask provided with a reflux condenser, or in a bomb-tube. An addition of a few drops of concentrated hydrochloric acid often

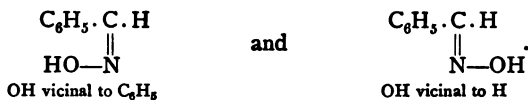
¹ B. 15, 1324.

expedites the reaction. (2) The formation of oximes may be brought about by the use of free hydroxylamine obtained by treating its hydrochloride with the theoretical amount of a solution of sodium carbonate (3) Oximes may in many cases be very easily obtained, if, as above, for one carbonyl group three molecules of hydroxylamine hydrochloride and nine molecules of potassium hydroxide are used; in the presence of a large excess of hydroxylamine in a strongly alkaline solution, generally a very smooth decomposition takes place. Since the oximes possess a weak acid character, under these conditions the alkali salt of the oxime is first obtained, *e.g.* :

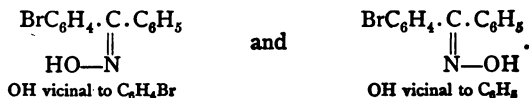


from which the free oxime is liberated by treating it with an acid.

Of especial significance for the stereo-chemistry of nitrogen are the oximes of aldehydes as well as those of the unsymmetrical ketones. By the action of hydroxylamine on benzaldehyde, *e.g.*, there is formed not only a single oxime, but a mixture of *two* stereo-isomers. This is also true when oximes are formed from many unsymmetrical ketones. The existence of these isomers is explained by the assumption that the three valencies of nitrogen do not lie in a plane, but that they extend into space, proceeding from a point like the three edges of a regular triangular pyramid.¹ Since, *e.g.*, in the formation of benzaldehyde-oxime, the hydroxyl-group of the hydroxylamine is vicinal to either the phenyl-group or hydrogen atom, the two following stereo-isomers are possible :



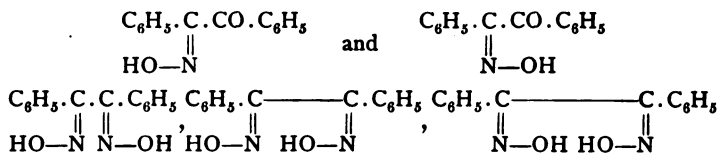
The stereoisomeric forms of an unsymmetrical ketone are, according to this conception, to be expressed by the following formulæ, *e.g.* :



¹ B. 23, 11, 1243.

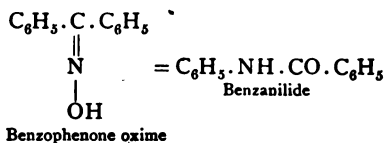
With symmetrical ketones it is obviously immaterial upon which of the two similar sides the hydroxyl-group finds itself, so that here only one oxime is possible.

In this place the two mono-oximes and three dioximes of benzil may be referred to again. These compounds gave the impetus to the investigations¹ of this class of compounds. They are explained by the following space-formulæ:

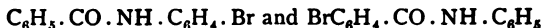


Not all aldehydes and unsymmetrical ketones yield two oximes. In many cases one form is so unstable (labile) that only the other (stable) modification exists.

(c) If phosphorus pentachloride is allowed to act on an oxime, it is transformed into an anilide,² *e.g.*:



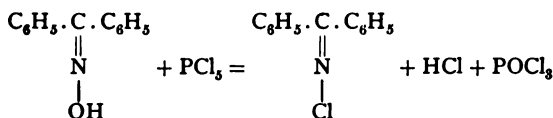
This so-called Beckmann transformation has been of great significance for the explanation of the constitution of the isomeric oximes. If, *e.g.*, phosphorus pentachloride is allowed to act on both of the above formulated stereoisomeric oximes of the brombenzophenone, the same compounds are not obtained from both, but two different ones, which, as follows from their saponification products, correspond on the one hand to the benzoyl derivative of bromaniline, and, on the other, to the brombenzoyl derivative of aniline:



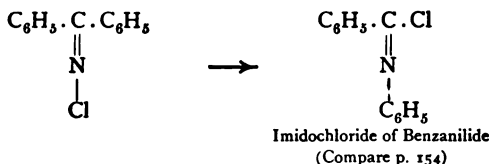
The transformation takes place, probably, in the following way: If phosphorus pentachloride is allowed to act on an oxime, the hydroxyl-group is replaced by chlorine:

¹ B. 16, 503; 21, 784, 1304, 3510; 22, 537, 564, 1985, 1996.

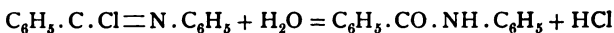
² B. 19, 988; 20, 1507 and 2580; A. 252, 1.



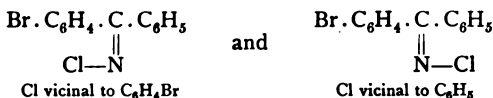
But a compound of this kind, in which chlorine is united with nitrogen, is unstable, and it is immediately transformed into a more stable imido-chloride, the chlorine atom being replaced by a phenyl-group :



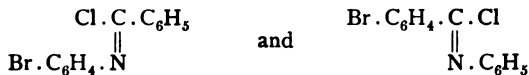
If this is now treated with water, benzanilide is formed, in accordance with the following equation :



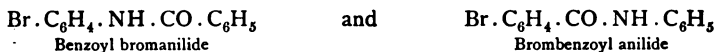
If the oxime of brombenzophenone, formulated above, is subjected to a similar reaction, the unstable chlorides are first obtained :



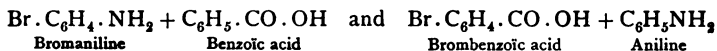
If the most probable assumption is now made, that the chlorine atom gives up its position to the vicinal hydrocarbon radical, there are formed :



from which, by treatment with water, there are obtained :

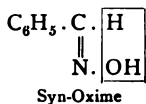


Upon saponification, these yield :

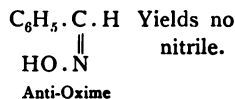


That hydrocarbon radical which in the oxime was vicinal to the hydroxyl-group, is, therefore, on saponification of the polymerised product, obtained in the form of a primary amine. In this way, the constitution of the stereoisomeric oximes of the unsymmetrical ketones is determined.

Whatever may be the space configuration of the stereoisomeric aldoximes, the derivatives containing an acid radical (acetyl derivative) of the one form easily yield an acid-nitrile on being treated with soda, while the second form does not. The hypothesis proposed, which seems very probable, suggests that in the first case the acid radical (or in the corresponding oxime) the hydroxyl-group is vicinal to the aldehyde hydrogen atom, while in the second case it is vicinal to the hydrocarbon residue :



Owing to the nearness of the oxygen and hydroxyl, it loses water easily, and yields a nitrile, $\text{C}_6\text{H}_5\text{C}\equiv\text{N}$.

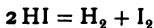


34. REACTION: REDUCTION OF A KETONE TO A HYDROCARBON

EXAMPLE: Diphenyl Methane from Benzophenone¹

A mixture of 10 grammes of benzophenone, 12 grammes of hydriodic acid (boiling-point, 127°), and 2 grammes of red phosphorus is heated in a sealed tube² for 6 hours at 160° . The reaction-mixture is then treated with ether, poured into a small separating funnel, and shaken up with water several times. The ethereal solution is filtered through a small folded filter, and dried, the ether evaporated, and the residue distilled. Boiling-point, 263° . On cooling, the diphenyl methane solidifies to crystals which melt at 27° . Yield, almost quantitative.

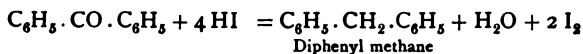
Hydriodic acid, especially at high temperatures, is an extremely energetic reducing agent, which can be used to effect reduction when, as in the above case, another reducing agent, *e.g.*, a metal and acid, could not be employed. The reducing action depends on the following decomposition :



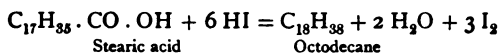
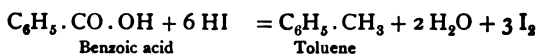
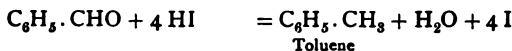
¹ B. 7, 1624.

² Before opening the tube, see page 67.

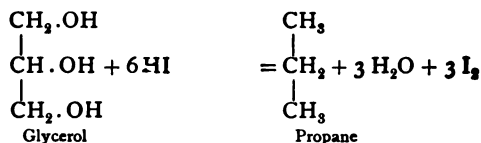
The above reaction takes place in accordance with the following equation :



With the aid of hydriodic acid, not only ketones but also aldehydes and acids may be reduced to the hydrocarbon from which they are derived, *e.g.* :



Alcohols, iodides, etc., can also be reduced to their final reduction products, the hydrocarbons, *e.g.* :



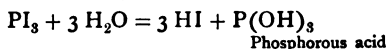
By heating with hydriodic acid, the unsaturated compounds take up hydrogen, *e.g.* :



The effect of hydriodic acid is increased by the addition of red phosphorus. Under these conditions, during the course of the reaction, the liberated iodine unites with the phosphorus to form phosphorus tri-iodide :



which with the water present again decomposes to form hydriodic acid :



A definite amount of hydriodic acid can thus, provided a sufficient quantity of phosphorus is present, act as a continuous reducing agent.

35. REACTION : ALDEHYDE SYNTHESIS — GATTERMANN-KOCH**EXAMPLE : p-Tolylaldehyde from Toluene and Carbon Monoxide¹**

To 30 grammes of freshly distilled toluene (boiling-point 110°) contained in a wide-necked vessel (an "extract of beef" jar is convenient) cooled with water, add, not too quickly, 45 grammes of pulverised, freshly prepared aluminium chloride and 5 grammes pure cuprous chloride (see page 389). The vessel is closed by a three-hole cork ; in the middle hole is inserted a glass tube which carries a stirrer (paddle wheel of glass) ; the other holes are used for the inlet and outlet tubes (Fig. 75).

After the apparatus has been fastened firmly in a clamp it is immersed into

a casserole filled with water at 20° . A current, not too rapid, of carbon monoxide and hydro-

chloric acid gas is led in through the prong-shaped tube while the stirrer is set in motion (a small motor is convenient).

The carbon monoxide, contained in a gasometer of about 10 litres, is passed first through a solution of caustic potash² (1 : 1)

and then through a wash-bottle containing concentrated sulphuric acid. The hydrochloric acid is generated in a Kipp apparatus from fused ammonium chloride and concentrated sulphuric acid. It is passed

through a wash-bottle containing concentrated sulphuric acid.

The gas currents are so regulated that the volume of the carbon monoxide is about twice as large as that of the hydrochloric acid.

The escaping gas is led directly to the hood opening: In the course of an hour when about 1-2 litres of carbon monoxide have been passed into the mixture, the temperature rises to $25-30^{\circ}$; the remainder of the gas is passed in during four to five hours.

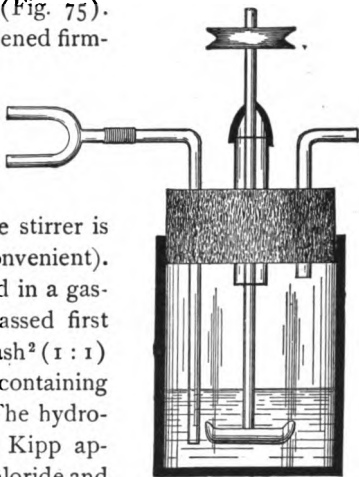


FIG. 75.

¹ B. 30, 1622; 31, 1149. A. 347, 347.

² If carbon monoxide is prepared from formic acid, the caustic potash wash-bottle is unnecessary.

If the reaction-mixture should become so viscous before the lapse of this time that the stirrer revolves only with difficulty, the reaction may be stopped. The viscid product is then poured into a large flask containing crushed ice; the aldehyde formed and any unattacked toluene is distilled over with steam. The distillate—oil and water—is then shaken up with a sodium bisulphite solution for a long time. The toluene, remaining undissolved, is separated in a dropping funnel. If the aldehyde-bisulphite compound should crystallise out, water is added until it dissolves. The filtered water solution is then treated with anhydrous soda until it shows a decided alkaline reaction; the aldehyde is then again distilled over with steam. It is extracted from the distillate with ether. Upon evaporating the ether, from 20–22 grammes of perfectly pure tolylaldehyde remains. Boiling-point, 204° .

Preparation of Carbon Monoxide

1. From Oxalic Acid.

In a round litre flask heat 100 grammes of crystallised oxalic acid with 600 grammes of concentrated sulphuric acid.¹ The gases evolved are passed into two large wash-cylinders (Fig. 74) filled with a solution of caustic potash (1 part caustic potash to 2 parts of water), and then into a gasometer (Fig. 76). At first the sulphuric acid is heated somewhat strongly. As soon as the oxalic acid has dissolved and a regular current of gas comes off, the flame is lowered. Before filling the gasometer the gas is tested by collecting a test-tube full over water and applying a match. So long as air remains in the apparatus, a slight explosion will occur. But as soon as pure gas is evolved, it burns quietly in the tube. It is then admitted into the gasometer. When the evolution of gas ceases, the apparatus is taken

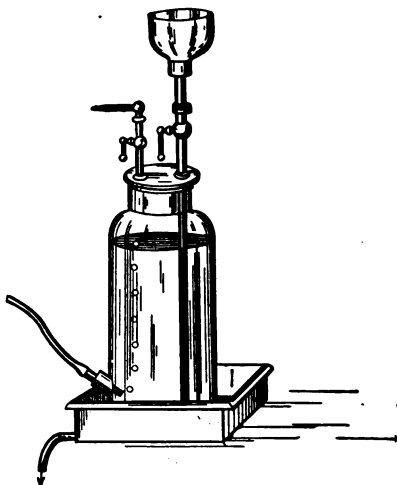


FIG. 76.

¹ The outlet tube through which the gaseous mixture passes before coming in contact with the caustic potash solution should be as wide as practicable, in order that it may not be clogged by the sublimed oxalic acid.

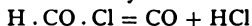
apart. Since carbon monoxide is poisonous, the experiment is carried out under the hood, and care is taken not to breathe the gas.

2. From Formic Acid.

Carbon monoxide is more readily liberated from formic acid. The latter may now be obtained at a low cost. In a half-litre flask, provided with a dropping funnel, a thermometer and an outlet tube, place 100 c.c. (110 grammes) of 98–100 % formic acid. To this add, gradually, concentrated sulphuric acid. Upon the addition of about 50 c.c. of acid the temperature will rise to 60–70°. The acid is now added more slowly, drop by drop, in order to keep the temperature at 50–60°. If necessary, the mixture is gently heated on a water-bath for a short time. When the air has been completely replaced, the gas, consisting of pure carbon monoxide, is admitted into the gasometer.

If it is desired to avoid the use of a gasometer, a regular and continuous current of carbon monoxide may be generated as follows: A litre flask, provided with a safety-tube, containing 200 grammes of oxalic acid and 200 grammes of sulphuric acid (conc.), is heated in an oil-bath to 120°; the temperature is gradually increased according to the conditions. If the gas be passed first into two caustic potash cylinders, and then into two sulphuric acid (conc.) cylinders, it may be used directly for the synthesis. Or, the carbon monoxide obtained from formic acid may be used directly, after it has been dried by passing through a wash-bottle containing concentrated sulphuric acid. But the yield of aldehyde is not as good as when a gasometer is used.

A direct synthesis of the aromatic aldehydes by means of the Friedel-Crafts reaction could not be brought about until recently, because of the instability of formyl chloride, which, if formed, decomposes immediately into carbon monoxide and hydrochloric acid:

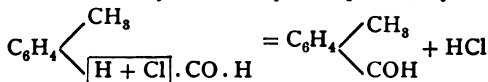


If it were stable, it should form aldehydes, in accordance with this equation:



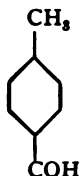
But now it is known that a mixture of carbon monoxide and hydrochloric acid in the *presence of cuprous chloride*, which combines with the former, behaves like formyl chloride.

The Gattermann-Koch synthesis may be expressed by the following equation:

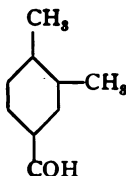


From other hydrocarbons like o- and m-xylene, mesitylene, ethylbenzene, diphenyl, etc., in an analogous way, aldehydes may be obtained. As has been shown in the ketone syntheses, the acid radical goes into the para position to the alkyl residue, so also in the aldehyde syntheses

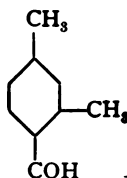
the aldehyde group always enters the para position to the alkyl residue
Thus from toluene there is obtained :



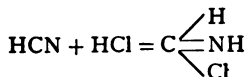
From o-xylene →



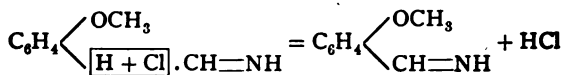
From m-xylene →



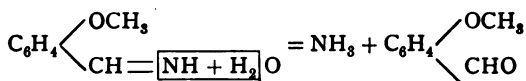
Since the Friedel-Crafts reaction, when applied to phenol ethers, yields the corresponding aldehydes far more easily than the same reaction applied to the hydrocarbons, it is remarkable that the Gattermann-Koch method cannot be used with phenol ethers. If it be desired to obtain aldehydes from them, hydrocyanic acid is used in place of carbon monoxide; in these cases the presence of cuprous chloride is unnecessary. The action takes place, due to the union of hydrocyanic and hydrochloric acids to form the chloride of imidoformic acid :



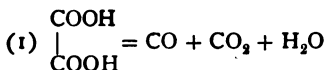
which, under the influence of aluminium chloride, reacts with the phenol ether, liberating hydrochloric acid :



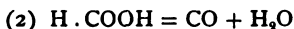
There is thus obtained first the aldehyde-imide which, through the action of acids, passes over into the aldehyde with great ease :



In this way it is possible to introduce the aldehyde group into phenol ethers as well as into the phenols. The latter always enters the para position to the oxalyl or hydroxyl groups. The carbon monoxide is obtained in accordance with the following equations :



The mixture of gases is separated by passing it through a solution of caustic soda or caustic potash ; the carbon dioxide is absorbed, and the carbon monoxide emerges in a pure condition.



36. REACTION: SAPONIFICATION OF AN ACID-NITRILE

EXAMPLE : Toluic Acid from Toly Nitrile¹

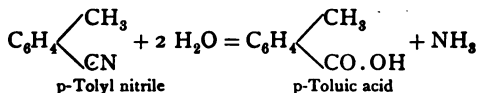
The p-tolyl nitrile obtained in Reaction 10 is heated with slightly diluted sulphuric acid on the sand-bath in a round flask with reflux condenser until crystals of toluic acid appear in the condenser. For each gramme of the nitrile a mixture of 6 grammes of concentrated sulphuric acid with 2 grammes of water is used. After cooling it is diluted with water, the acid separating out is filtered off and washed several times with water. A small portion is dissolved in a little alcohol, and hot water added until the solution just becomes turbid ; it is then boiled some time with animal charcoal. On cooling, the pure acid is obtained. Melting-point, 177°. Yield, 80–90 % of the theory.

By saponification in a narrow sense is understood the splitting up of an acid-ester into an alcohol and acid. It is, however, used in a wider sense to indicate the conversion of acid-derivatives, like nitriles, amides, substituted amides, *e.g.*, anilides, into acids of the same name. Saponification may be conducted either in an alkaline or an acid solution. Thus, for instance, acetamide reacts on heating with a solution

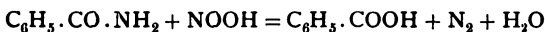
¹ A. 258, 10.

of caustic potash or caustic soda with the formation of the alkali salt of acetic acid and the evolution of ammonia. Nitriles and esters may frequently be saponified by water solutions of the alkalis. Further, alcoholic caustic potash or caustic soda can be used for a similar purpose. Finally, saponification may be effected by heating with a sodium carbonate solution under pressure; this method is especially well adapted for difficultly saponifiable amides or anilides.

In order to effect saponification in acid solution, the substance to be saponified is heated with either hydrochloric acid or sulphuric acid in varying degrees of dilution, *e.g.*:



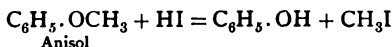
Acid amides may be easily saponified by dissolving in concentrated sulphuric acid, cooling, adding sodium nitrite, and then gradually heating,¹ *e.g.*:



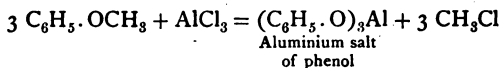
In order to saponify a nitrile by this method it is first converted by heating with 85% sulphuric acid into the amide, and this is treated as directed above.

Frequently it is better to allow the nitrite to act directly on the warm dilute sulphuric acid solution of the amide.

The decomposition of the ethers of phenols is also designated as saponification. Such decomposition cannot be effected by the methods hitherto given. Hydriodic acid is used which, when heated with phenol-ethers, decomposes them into the phenol and the alkyl iodide:



Anhydrous aluminium chloride may be used here with great advantage; upon heating, it acts on the phenol-ether in the manner indicated by the following equation:



If a phenol salt is treated with an acid, the free phenol will separate out. This method presents the advantage that it may be applied to substances containing, in addition to the phenol-ether radical, a reducible carbonyl group, which, if treated with hydriodic acid, would be changed.

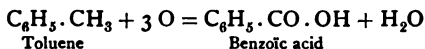
¹ B. 26, Ref. 773; 28, Ref. 917; 32, 1118.

37. REACTION: OXIDATION OF THE SIDE-CHAIN OF AN AROMATIC COMPOUND

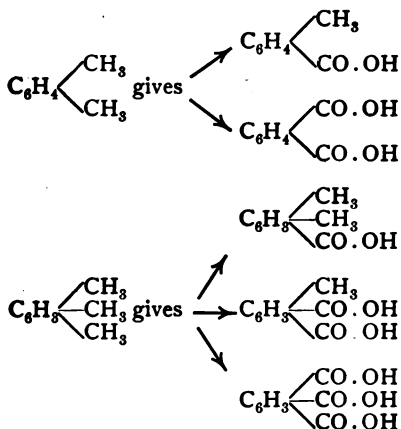
EXAMPLE: Terephthalic Acid from p-Toluic Acid

Dissolve 5 grammes of the crude toluic acid obtained in Reaction 36 in a solution of 3 grammes of sodium hydroxide in 250 c.c. of water; heat in a porcelain dish on the water-bath, and gradually treat with a solution of 12 grammes of finely powdered potassium permanganate in 250 c.c. of water until, after long boiling, the red colour of the permanganate no longer vanishes. Alcohol is then added until the liquid is colourless, and, after cooling, the manganese dioxide separating out is filtered off; this is washed with hot water, and the filtrate, heated to boiling, is acidified with concentrated hydrochloric acid. After cooling, the terephthalic acid is filtered off, washed with water, and dried on the water-bath. Yield, 90% of the theory. Terephthalic acid is insoluble in water. On heating, it sublimes without melting.

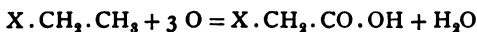
It is a common property of aliphatic side-chains, united with the benzene nucleus, to pass over to carboxyl groups on oxidation. A methyl group requires 3 atoms of oxygen for oxidation:



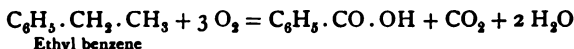
If several side-chains are present in a compound, either all or a portion of them may be converted into carboxyl groups:



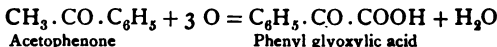
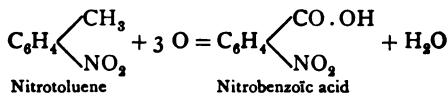
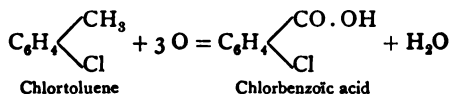
If a side-chain contains several carbon atoms, in many cases only the methyl group at the end of the chain can be oxidised, *e.g.* :



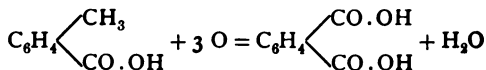
But by an energetic oxidation all the carbon atoms, with the exception of the last, are split off, *e.g.* :



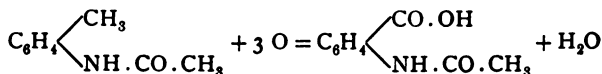
The basicity of the acid derived from the oxidation of a hydrocarbon accordingly gives an indication concerning the number of side-chains of the hydrocarbon. Derivatives of hydrocarbons are also capable of similar reaction, *e.g.* :



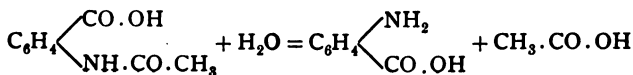
The reaction carried out above takes place in accordance with this equation :



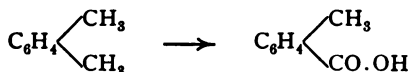
Amines and phenols cannot be directly oxidised in most cases, but an indirect method must be employed, by which the former are converted into an acid derivative, and the latter into an ester. If, *e.g.*, it is desired to convert p-toluidine into p-amidobenzoic acid, the base is first acetylated, and the acetatoluide is then oxidised :



The acid thus obtained is then saponified, and the desired amido-benzoic acid is formed :

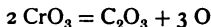


If it is desired, on the other hand, to oxidise a phenol, *e.g.*, cresol, $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH}_3 \\ \text{OH} \end{smallmatrix}$, the sulphuric acid- or phosphoric acid-ester of it is first prepared and oxidised; the reaction-product is then saponified. As oxidising agent, dilute nitric acid (1 vol. conc. nitric acid to 3 vol. water¹), chromic acid or potassium permanganate is used. The mildest effect is obtained with the nitric acid, which is therefore used when all the side-chains are not to be oxidised, but only a portion of them, *e.g.*:



Nitric acid is also used in other cases, where, as frequently happens with ortho derivatives, other oxidising agents totally destroy the substance.

Chromic acid, in the form of its anhydride generally, dissolved in glacial acetic acid, or as a water solution of potassium dichromate or sodium dichromate acidified with dilute sulphuric acid, can also be used as an oxidising agent, not only in the case in hand, but also for the oxidation of alcohols, ketones, etc. In oxidation reactions, two molecules of chromic anhydride (CrO_3) give three atoms of oxygen:



For the oxidation of aromatic hydrocarbons,² experience has shown that a good oxidising mixture is 40 parts of potassium dichromate, 55 parts of concentrated sulphuric acid, diluted with twice its volume of water.

With potassium permanganate³ oxidation can be effected either in alkaline or in acid solution. In the first case, manganese dioxide is deposited:



Two molecules of potassium permanganate yield, therefore, in alkaline solution, three atoms of oxygen.

In acid solution (sulphuric acid), no manganese dioxide separates out, since it is dissolved by the sulphuric acid, with evolution of oxygen, to form manganous sulphate:



Two molecules of the permanganate in acid solution, therefore, yield 5 atoms of available oxygen.

In oxidising with potassium permanganate, a 2-5 % solution is generally used. An excess of the permanganate can be removed by the addition of alcohol or sulphurous acid. The alcohol is oxidised into aldehyde or acetic acid, and the sulphurous acid into sulphuric acid.

¹ A. 137, 302.

² A. 133, 41.

³ B. 7, 1057.

38. REACTION: SYNTHESIS OF OXYALDEHYDES. REIMER AND TIEMANN¹**EXAMPLE: Salicylic Aldehyde from Phenol and Chloroform**

In a round litre flask dissolve 80 grammes of caustic soda in 80 c.c. of water by heating. Add 25 grammes of phenol; cool the solution without shaking to 60–65° by immersion in cold water. By means of a two-hole cork attach to the flask an effective reflux condenser, and insert a thermometer the bulb of which dips into the liquid. Add 60 grammes of chloroform gradually, as follows: At first add one-third through the condenser; on gentle shaking the liquid becomes a fuchsine-red in colour. After a short time the colour changes to orange, and the temperature rises. When it reaches 70° the entire flask is immersed in cold water until the thermometer indicates 65°. In this way during the entire reaction the temperature is always kept between 65° and 70°. Should it fall below 60°, the mixture is warmed by immersing it a short time in hot water until the mercury rises to 65°. After 10–15 minutes the second third of the chloroform is added, observing the precautions just given. Finally, after about 20 minutes, the remainder of the chloroform is added. Since toward the end the reaction takes place very quietly, the flask is frequently immersed in hot water in order to maintain the temperature between the prescribed limits. The synthesis requires in all from 1½ to 2 hours. Frequent shaking of the mixture, especially during the last phase, increases the yield materially. When the reaction is complete the chloroform is distilled off with steam. The orange-coloured alkaline liquid is allowed to cool somewhat, and is acidified carefully with dilute sulphuric acid, upon which it becomes almost colourless; finally, steam is passed into it until drops of oil no longer go over.

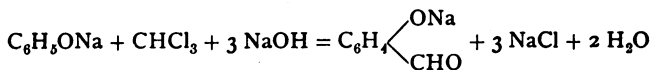
The distillate is then extracted with ether, the ethereal solution separated from the water, and the ether evaporated. The residue, consisting of unchanged phenol and salicylic aldehyde, is treated

¹ B. 9, 423, 824; 10, 1562; 15, 2685, etc.

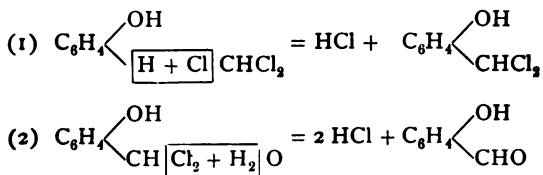
with twice its volume of a concentrated solution of commercial sodium bisulphite. Upon stirring well for a long time with a glass rod, a solid paste of the double compound of the aldehyde and bisulphite should separate out. After standing from $\frac{1}{2}$ –1 hour the crystals are filtered with suction, pressed firmly together, and washed several times with alcohol to remove completely the adhering phenol, and finally with ether. The pearly, lustrous leaflets are then well pressed out on a porous plate and the aldehyde set free by a gentle warming with dilute sulphuric acid on the water-bath. On cooling, the mixture is extracted with ether, the ethereal solution dried over anhydrous Glauber's salt, the ether is evaporated, and the residue of the pure aldehyde is distilled. Boiling-point, 196° . Yield, 10–12 grammes.

The small amount of the p-oxybenzaldehyde formed with the salicylic aldehyde is not volatile with steam, and remains back in the flask after the distillation with steam. In order to obtain it, the residue remaining in the flask after cooling is filtered through a folded filter, and the clear filtrate saturated with solid salt, upon which the p-oxybenzaldehyde separates out at once or on standing. If this be filtered off, and the filtrate extracted with ether, a further quantity is obtained, which, together with the first, is purified by recrystallisation from water with the addition of a solution of sulphur dioxide. Melting-point, 116° . Yield, 2–3 grammes.

The synthesis takes place in accordance with this equation :



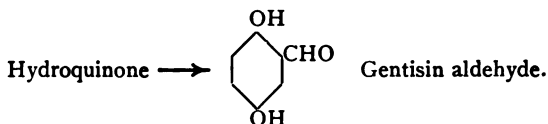
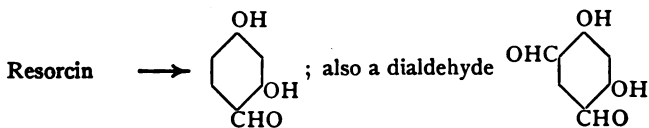
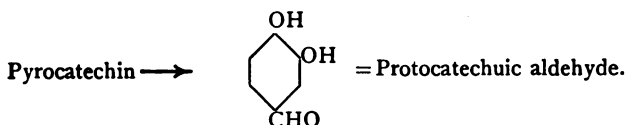
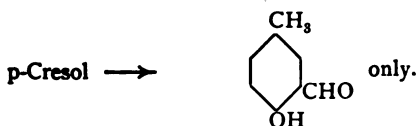
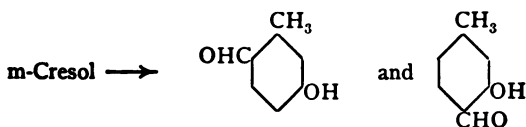
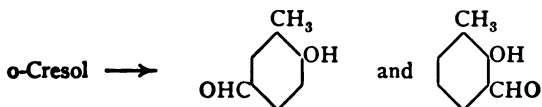
Probably the reaction takes place in the two phases :



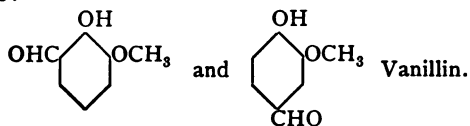
By this method the aldehyde group may be introduced into mono- and poly-acid phenols; it enters the ortho and para positions to a hydroxyl group:

Phenol gives

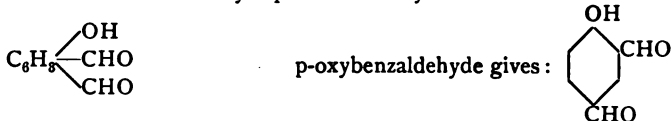
o- and p-oxybenzaldehyde,



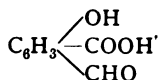
The reaction also takes place with the ethers of poly-acid phenols
Thus guaiacol gives:



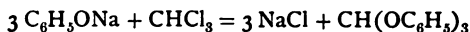
From resorcinmonomethyl ether there are formed two monoaldehydes and two dialdehydes. The reaction is also applicable to oxyaldehydes as well as oxycarbonic acids. Thus from salicylic aldehyde there is formed a mixture of two oxyisophthalic aldehydes :



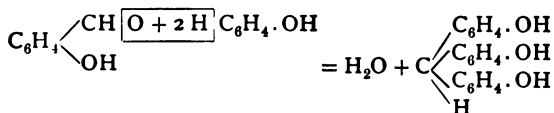
That resorcin, in addition to a monoaldehyde, yields a dialdehyde, has already been mentioned. From the three oxybenzoic acids are formed two oxyaldehyde acids,



This synthesis is capable of very wide application. But it has some defects. Thus the yield of aldehyde obtained in accordance with the original directions leaves much to be desired. The yield is very much decreased by the fact that a portion of the phenol does not enter into the reaction, and another portion reacts with the chloroform to produce an ester of ortho formic acid :

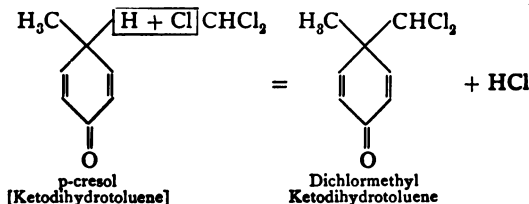


A portion of the aldehyde first formed is lost by condensation with some unattacked phenol, forming a derivative of triphenylmethane :



Further, a portion of the oxyaldehydes is converted into resins by the alkali.

And again, many phenols react as keto-compounds, and give rise to varying quantities of by-products, *e.g.* :



In some cases, the chief product of the reaction consists of chlorinated ketones of this order. (B. 35, 4209; A. 352, 288.)

In addition, the separation of the mixture of the mono- and dialdehydes is often attended with serious difficulty.

As already mentioned on page 334, the aldehyde group may be introduced into phenols by the use of condensation agents like aluminium chloride, zinc chloride, hydrocyanic and hydrochloric acids. These reactions possess many advantages over those discussed above. They take place more smoothly, yield only the p-oxyaldehydes, and introduce only one aldehyde group; further, only small amounts of resins are formed, and finally they may be applied to phenols like pyrogallol, phloroglucin, the two naphthols, poly-acid phenols of naphthalene, etc. With these substances the other reaction is useless. (See B. 31, 1765; 32, 278, etc.; A. 357, 313.)

39. REACTION: KOLBE'S SYNTHESIS OF OXYACIDS

EXAMPLE: Salicylic Acid from Sodium Phenolate and Carbon Dioxide¹

Dissolve $12\frac{1}{2}$ grammes of chemically pure sodium hydroxide in 20 c.c. of water in a porcelain dish, or better a nickel dish, and

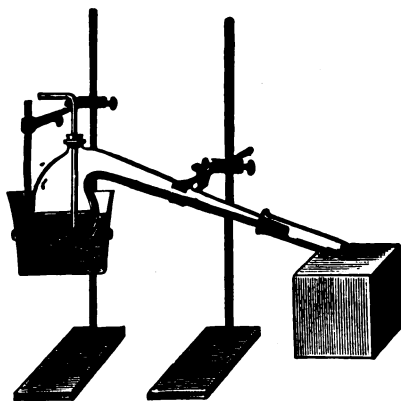


FIG. 77.

with stirring, treat gradually with 30 grammes of crystallised phenol. The greatest portion of the water is then evaporated by heating over a free flame, the mass being continually stirred. As soon as a crystalline film forms on the surface of the liquid, the heating is continued with a luminous flame, which is not placed directly under the dish, but is kept in constant motion. In order to fasten the dish, a pair of crucible tongs is clamped in a vertical position,

and the dish supported between its jaws. There is first obtained a caked, bright-coloured mass, which is crushed from time to time with a mortar-pestle. As soon as the particles no longer

¹ J. pr. [2] 10, 89; 27, 39; 31, 397.

bake together, the mass is pulverised quickly in a dry mortar, the dry mass is then heated with thorough stirring in a nickel dish to dusty dryness. It is then placed in a tubulated retort of 200 c.c. capacity. The retort is then immersed as far as possible in an oil-bath (Fig. 77). This is heated to 110° , and at this temperature a current of dry carbon dioxide is passed over the sodium phenolate (the end of the delivery tube is 1 cm. above the upper surface of the sodium phenolate); this is passed into the retort for an hour. The temperature is then gradually raised (20° per hour) during the course of four hours, while a not too rapid current is passed in, to 190° . The mixture is finally heated 1–2 hours at 200° . During the operation the mass is stirred several times with a glass rod. After cooling, the phenol in the neck of the retort is melted by the application of a flame to the outside, the dusty, fine powder is poured into a large beaker, the retort is washed out several times with water, and the salicylic acid precipitated with much concentrated hydrochloric acid. After the reaction-mixture has been cooled with ice-water a long time, and the sides of the vessel rubbed with a glass rod, the crude salicylic acid is filtered off, washed with a little water, and pressed out on a porous plate.

The purification of the crude salicylic acid is accomplished best with superheated steam. For this purpose the acid, in a *dry condition*, is placed in a short-necked flask, and heated in an oil-bath to 170° , a not too rapid current of steam at a temperature of 170 – 180° (see page 41) is passed over it. The connection between the flask and steam generator must not be made until the oil-bath and the steam have the same temperature. Since the acid distilling over very soon stops up a condenser tube of the usual width, one should use for this experiment a tube of 2.5 cm. width (width of mantel 5 cm., length of same 75 cm.). The connecting tube between the flask and condenser must be 2 cm. wide, and as short as possible. If the acid removed from the condenser be dissolved in the watery distillate in the receiver by heating, long colourless needles separate out on cooling. Melting-point, 156° . Yield, 5–10 grammes.

The preparation of salicylic acid does not always take place

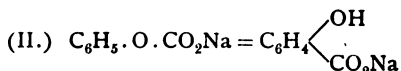
successfully the first time. The success of the experiment depends particularly on the condition of the sodium phenolate, which must be *perfectly dry*.¹ If it "cakes" on heating the retort, there is great probability that the experiment will be unsuccessful.

The operation should be so arranged that the sodium phenolate is prepared toward evening, so that it may be allowed to stand in a sulphuric acid desiccator over night. The drying in the current of carbon dioxide is begun immediately next morning.

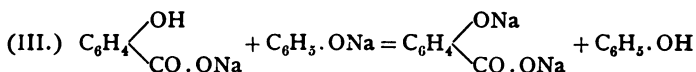
The synthesis is named after its discoverer, Kolbe. It takes place in three phases. In the first, the carbon dioxide is added to the sodium phenolate, which forms sodium phenyl carbonate:



In the above experiment this reaction is completed during the heating up to 110° for one hour. In the second phase, the sodium phenyl carbonate is transformed into the so-called neutral sodium salicylate:



while in the last phase a molecule of this salt reacts with a molecule of unchanged sodium phenolate in the following way:



These two latter reactions take place during the gradual heating up to 200° . Only one-half of the phenol, therefore, is converted into salicylic acid, the second half being obtained unchanged.

A modification of the Kolbe synthesis which permits the immediate conversion of all the phenol into salicylic acid is known as Schmitt's synthesis. According to this method, as in the other, the sodium phenyl carbonate is first prepared; this is then further heated in an

¹ The experiment is more certain of success if the sodium phenolate is heated a half hour in a current of dry hydrogen at 140° (retort in the oil-bath) before the introduction of the carbon dioxide; the mass must be cooled to 110° before the latter is led in.

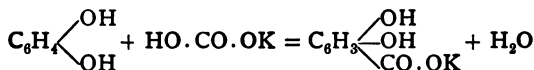
autoclave under pressure to 140° , upon which it is completely transformed into sodium salicylate according to Equation II. Instead of preparing the sodium phenyl carbonate with gaseous carbon dioxide, the sodium phenolate may be mixed directly with liquid or solid carbon dioxide in the autoclave.

The Kolbe synthesis is capable of very common application, since from each mon-acid phenol, a carbonic acid may be obtained in the same way as that used above. The carboxyl group under these conditions primarily seeks the ortho position to the hydroxyl group. The derivatives of phenols, *e.g.* the three chlorphenols, yield chlorinated salicylic acids. With acid-ethers of poly-acid phenols which still con-

tain a free hydroxyl group, as, *e.g.*, guaiacol, $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{OCH}_3 \\ \text{OH} \end{smallmatrix}$, this reaction likewise takes place. From the two naphthols $\text{C}_{10}\text{H}_7\text{.OH}$ the oxynaphthoic acids $\text{C}_{10}\text{H}_6 \begin{smallmatrix} \text{OH} \\ \text{CO.OH} \end{smallmatrix}$, can be obtained.

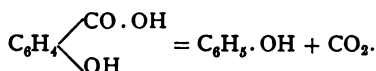
If in the Kolbe reaction instead of sodium phenolate, potassium phenolate is used, the para-oxybenzoic acid is obtained, and not the ortho-acid. The potassium phenolate, like the sodium phenolate, first absorbs carbon dioxide, and the potassium phenyl carbonate thus formed, heated in carbon dioxide up to 150° , also yields salicylic acid; but if the temperature is increased, an increasingly larger quantity of the para-acid is obtained, until finally at 220° the potassium para-oxybenzoate is the only product.

The addition of carbon dioxide is effected with greater ease in poly-acid phenols. With these compounds the reaction begins if the phenol is boiled in a water-solution of ammonium carbonate or potassium hydrogen carbonate, *e.g.* :



Salicylic acid is prepared technically on the large scale. Since it is an excellent antiseptic, it finds extensive application in preventing fermentation, for the preservation of meat, for the disinfection of wounds. It may easily be recognised, since its water solution gives a violet colour with ferric chloride; in this action it differs from the para- and meta-modifications. It is volatile with steam; for this reason it must not be boiled too long in an open vessel when it is to be recrystallised. All ortho-oxy-carbonic acids show this property; the

meta- and para-isomers are not volatile with steam. If salicylic acid is heated strongly, it decomposes into carbon dioxide and phenol :



The para-oxycarbonic acids show the same property while the meta-acids are stable.

40. REACTION: GRIGNARD'S REACTION

(a) Benzoic Acid from Iodobenzene. (b) Benzhydrol from Iodo- or Brombenzene and Benzaldehyde

(a) 2.4 grammes of magnesium shavings (or thin magnesium ribbon, about 2 mm. in width, rubbed with fine emery-cloth, cleaned with filter paper, and cut in pieces 1-2 cm. long) are placed in a flask provided with a reflux condenser, and treated with a mixture of 20.4 grammes of **thoroughly dried** iodobenzene, 40 c.c. of **absolute ether**¹ and a granule of iodine. The flask is dipped in hot water, or heated in a water-bath at the boiling point of the ether. In $\frac{1}{4}$ - $\frac{1}{2}$ hour a white, flocculent precipitate will begin to form, and the heat of the reaction will cause the ether to boil actively when the water-bath is removed. If, now, the bottom of the flask is wrapped up in dry cloth, so as to utilise the heat of the reaction, the greater part of the magnesium will go into solution in the course of two hours. At the end of this period

¹ For experiments (a) and (b) absolute ether is prepared as follows: 250 c.c. of commercial ether are shaken several times with 100 c.c. of water, and allowed to remain over granular calcium chloride over night. The ether is separated from the calcium chloride and treated with thin, bright scales of sodium. The vessel is stoppered with a cork to which is attached a calcium chloride tube, open at both ends. As soon as the evolution of gas ceases, the ether is distilled off from the sodium in the same vessel. It is collected in a suction flask attached to the lower end of the condenser with a cork, and having a calcium chloride tube connected to its side tube to prevent moisture from entering the receiver. For the success of the experiments the use of **thoroughly dry** materials and vessels is absolutely necessary. Should the progress of the experiment (the dissolving of magnesium) require more time than is prescribed above, then the reagents were not dry enough. The experiment is repeated. The ether must again be kept in contact with sodium over night, and must be freshly distilled from the sodium just before it is to be used.

the boiling of the ether will either slacken or entirely cease. It is once more heated for half an hour on a water-bath, then cooled by surrounding the flask with ice water. The condenser is removed, and in the course of 2-3 hours a slow current of carbon dioxide (dried by passing through two wash-bottles containing concentrated sulphuric acid) is run into the ethereal solution of phenyl magnesium iodide, which may still contain a small quantity of undissolved magnesium. The flask is kept cold during this operation. The reaction mixture consists of two layers, a light upper layer of ether, and a heavy, viscous layer of the reaction product. When too rapid a current of carbon dioxide is passed, a single layer of viscous mass will be obtained. This will not impair the success of the experiment if the mixture was cooled thoroughly. It is now treated with finely broken ice, and with a cooled mixture of 15 c.c. of concentrated hydrochloric acid and an equal volume of water. The benzoic acid is extracted with commercial ether. The ether is evaporated, and the residue is gently warmed with caustic soda or caustic potash. After filtering the undissolved portion,¹ the alkaline solution is treated with hydrochloric acid. The precipitated benzoic acid is filtered. More acid is obtained by extracting the mother liquor with ether. Yield of the crude product 10-12 grammes. It crystallises from water in colourless lustrous leaves. M.P. 121°.

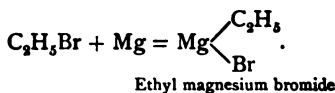
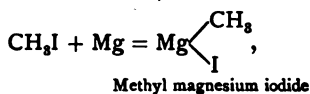
(b) 2.4 grammes of magnesium, 20.4 grammes of iodobenzene and 40 c.c. of absolute ether are converted into phenyl magnesium iodide as in (a). The product is cooled and carefully added, drop by drop, and with constant shaking, to a mixture of 10.6 grammes of freshly distilled benzaldehyde, and 30 c.c. of absolute ether. During this treatment the mixture is kept cold with ice water. A violent reaction takes place, with the formation of a yellow, and finally a white, precipitate. The reaction mixture is treated with ice as in (a), and then gradually with a cooled mixture of 15 c.c. concentrated hydrochloric acid in an equal volume of water. It is extracted with commercial ether. In order to remove benzaldehyde, the ethereal layer is shaken

¹ B. 40, 1584.

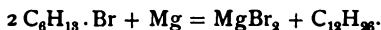
thoroughly with a dilute water solution of sodium bisulphite. Upon evaporating the ether an oily reaction product is obtained, which solidifies completely when it is cooled and rubbed with a glass rod. It is pressed on a porous plate and crystallised from ligroin. Yield of the crude product 10–14 grammes. Benzhydrol forms colourless needles which melt at 68°.

With brombenzene and magnesium the experiment is carried out in the following manner : 2.4 grammes of magnesium are covered, in the presence of a granule of iodine, with a mixture of 15.7 grammes of brombenzene that must be **perfectly dry** and must possess a **constant boiling point**, and 50 c.c. of **absolute ether**. If the materials used were **pure** and **perfectly dry**, the reaction will commence in a short time when the mixture is heated on a water-bath as in (a), *i.e.* the ether will continue to boil when the water-bath is removed. Almost all the magnesium goes into solution after $\frac{1}{4}$ – $\frac{1}{2}$ hour, and the boiling of the ether ceases. It is now heated moderately on the water-bath for a quarter of an hour. The treatment with benzaldehyde is exactly the same as with phenyl magnesium iodide.

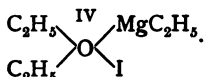
We are indebted to Grignard for the important observation that monobrom or monoiodo derivatives of different hydrocarbons (1 molecular weight) unite with magnesium (1 atomic weight) in the presence of absolute ether, *e.g.* :



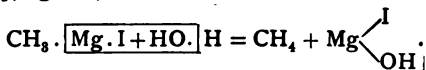
With long carbon chains in the aliphatic series, saturated hydrocarbons are also formed, and the longer the chain the more readily will this reaction take place. With molecules containing six or more carbon atoms, the main reaction consists in the removal of halogen, and the formation of the hydrocarbon :



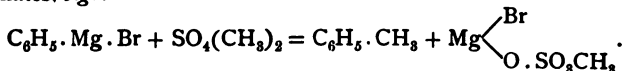
In an analogous way are formed $\text{C}_6\text{H}_5 \cdot \text{Mg} \cdot \text{Br}$ and $\text{C}_6\text{H}_5 \cdot \text{Mg} \cdot \text{I}$ = phenyl magnesium bromide and iodide ; $\text{C}_6\text{H}_5 \cdot \text{CH}_2 \cdot \text{Mg} \cdot \text{I}$ = benzyl magnesium iodide ; $\text{C}_6\text{H}_{11} \cdot \text{Mg} \cdot \text{I}$ = hexahydrophenyl magnesium iodide (from iodocyclohexamethylene), etc. Since these compounds are formed only in the presence of ether as solvent, it is probable that the ether not only serves as a solvent, but also plays an essential part in the reaction. As a matter of fact a compound $\text{C}_2\text{H}_5 \cdot \text{Mg} \cdot \text{I} + (\text{C}_2\text{H}_5)_2\text{O}$ has been isolated which may be regarded as an oxonium derivative :



The alkyl magnesium haloids or their ether derivatives are soluble in ether. On account of their unusual reactivity they may be used generally for the synthetic preparation of a large number of compounds. Thus *hydrocarbons* may be obtained when organo-magnesium compounds are treated with water or other substances containing the hydroxyl group, *e.g.* :

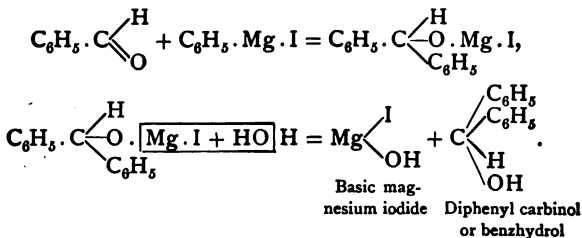


Hydrocarbons are likewise obtained synthetically by the action of alkyl sulphates, *e.g.* :



In the synthesis of alcohols (mentioned below) *hydrocarbons of the ethylene series* are formed, either as by-products, or as main products.

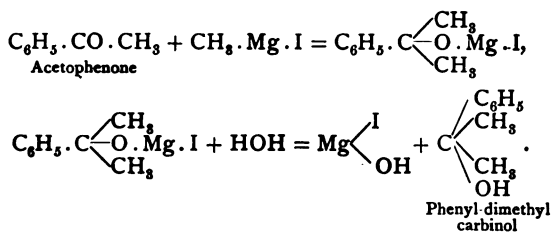
They unite with aldehydes to form double compounds, which are decomposed with water, yielding *secondary alcohols*, *e.g.* :



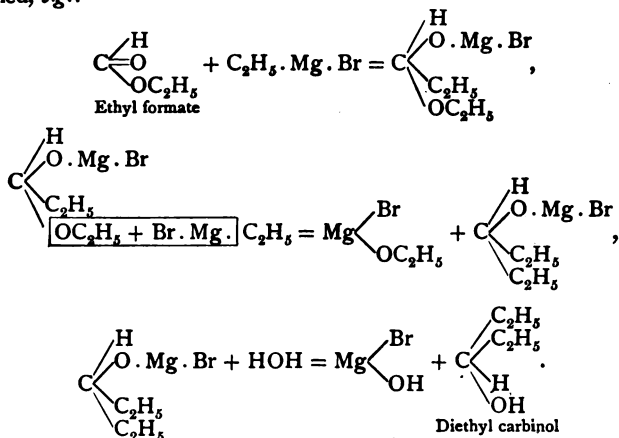
This reaction was carried out above in (b).

When the simplest aldehyde (formaldehyde) is used in the form of its polymer, trioxymethylene, *primary alcohols* are formed.

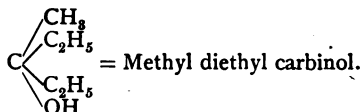
In an analogous manner, *tertiary alcohols* are formed when ketones are used, *e.g.* :



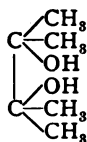
The esters of mono- and poly-basic carbonic acids also react readily with alkyl magnesium haloids with the formation of alcohols. In this case, two molecules of the magnesium compound react with one molecule of the ester. Thus from the esters of formic acid *secondary alcohols* are formed, e.g.:



In an analogous manner the esters of all the other monobasic acids form *tertiary alcohols*. Thus ethyl acetate forms with ethyl magnesium bromide:



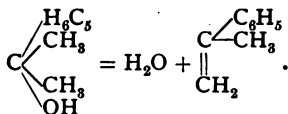
The reaction amounts essentially to the replacement of the carbonyl oxygen atom of the corresponding free acid by two univalent hydrocarbon residues. In this way, the esters of dibasic acids yield *diacid alcohols*. Thus from oxalic ester is formed:



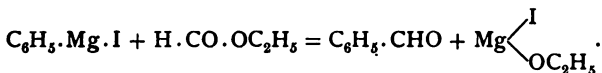
Tetramethyl glycol, or Pinacone.

In place of acid esters the corresponding chlorides or anhydrides may also be used.

As has already been mentioned, in the synthesis of alcohols, *unsaturated hydrocarbons* are also formed in addition to alcohols, either as by-products, or as main products. These are formed by the removal of water from alcohols, *e.g.* :

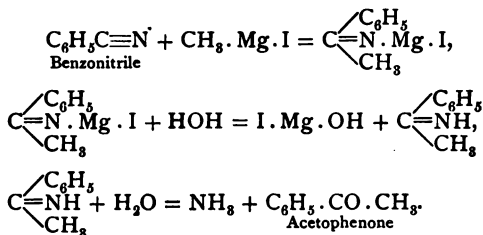


When organo-magnesium compounds are allowed to react with *excess* of formic ester, *aldehydes* are formed, and not secondary alcohols, *e.g.* :

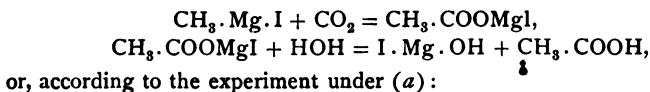


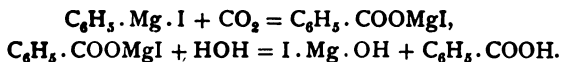
The same reaction will take place when, instead of formic ester, orthoformic ester, disubstituted formamide, and other derivatives of formic acid are used.

Nitriles unite with alkyl magnesium haloids to form compounds which yield *ketones* when they are decomposed with water :



When alkyl magnesium haloids are treated with dry carbon dioxide, *carbonic acids* are formed, *e.g.* :





Thus from the bromide or iodide of a hydrocarbon the monocarbonic acid¹ of the next higher series is formed.

These simple examples are sufficient to show the great importance of the Grignard reaction. Further details may be obtained from Jul. Schmidt's work on "Die organischen Magnesium-verbindungen und ihre Anwendung zu Synthesen I u. II" (Stuttgart, Enke; 1905 u. 1908).

41. REACTION: PREPARATION OF A DYE OF THE MALACHITE GREEN SERIES

EXAMPLE: Malachite Green from Benzaldehyde and Dimethylaniline²

(a) *Preparation of the Leuco-Base.* — A mixture of 50 grammes of dimethylaniline and 20 grammes of benzaldehyde (both freshly distilled) is heated in a porcelain dish, with frequent stirring, on the water-bath, for 4 hours, with 20 grammes of zinc chloride, which has been previously fused in a porcelain dish, and pulverised, after cooling. (See p. 358.) This viscous mass, which cannot be poured directly out of the dish, is melted by covering it with hot water, and heating it at the same time on the water-bath; while hot, it is transferred to a $\frac{1}{2}$ -litre flask. Steam is conducted into it, until no drops of it pass over. There is thus obtained the non-volatile leuco-base of the dye, in the form of a viscous mass, which adheres firmly to the walls of the distilling flask. After the liquid is cold, the water is poured off, the base adhering to the sides of the flask is washed with water several times, and then dissolved in the flask with alcohol, on the water-bath. After filtering, the solution is allowed to stand over night in a cool place, upon which the base separates out in colourless crystals; these are filtered off, washed with alcohol, and dried in the air on several layers of filter-paper. By concentrating the mother-liquor, a second crystallisation may be obtained. Should the base not crystallise, but separate out in an oily condition, which frequently

¹ Concerning the other course of the reaction see B. 40, 1584.

² A. 206, 83; 217, 250.

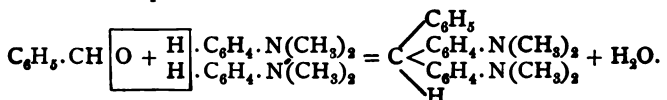
happens after a short standing of the filtered solution, this is due to the fact that an insufficient amount of alcohol has been used. In this case, more alcohol is added, and the mixture heated until the oil is dissolved.

(b) *Oxidation of the Leuco-base.*—Dissolve 10 parts, by weight, of the completely dry leuco-base by heating in a quantity of dilute hydrochloric acid corresponding to 2.7 parts, by weight, of anhydrous hydrochloric acid. For the purpose, dilute pure concentrated hydrochloric acid with double its volume of water, determine the specific gravity of the diluted acid, and refer to a table to find out how much anhydrous acid the solution contains, and from this calculate how much of the solution must be taken in order to get the required amount of the anhydrous acid (2.7 grammes). The colourless solution of the leuco-base is then diluted in a large flask with 800 parts, by weight, of water, and treated with 10 parts of 40 % acetic acid (sp. gr. 1.0523), prepared by gradually diluting glacial acetic acid with water; the mixture is *well cooled by throwing in pieces of ice*; then, with frequent stirring, gradually add (during 5 min.) a quantity of freshly prepared lead peroxide paste¹ corresponding to 7.5 grammes of pure lead peroxide. The peroxide is weighed off in a beaker, and treated with a quantity of water sufficient to form a very thin paste. The residue remaining in the beaker after the first emptying is washed out with water. After the addition of the peroxide, the reaction-mixture is allowed to stand five minutes, with frequent shaking; then add a solution of 10 parts of Glauber's salt and 50 parts of water; the solution is then filtered off through a folded filter from the precipitated lead sulphate and chloride. The filtrate is treated with a filtered solution of 8 parts of zinc chloride dissolved in as small a quantity of water as possible; then a saturated sodium chloride solution is added, until all the dye is precipitated. This is easily recognised by bringing a drop of the solution, with a glass rod on a piece of filter-paper; a bluish green precipitate surrounded by a circle of a still fainter bright green colour will be formed. The precipitated dye is filtered off with suction, washed with a little saturated sodium chloride solution, and pressed out

¹ See page 388.

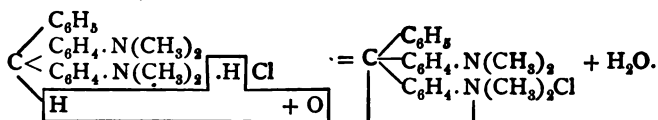
on a porous plate. In order to purify it further, it may be dissolved again in water; and from the filtered solution, after cooling, it is again thrown out by sodium chloride.

The reaction just carried out, discovered by Otto Fischer in 1877, is also used in the large scale for the preparation of Malachite Green, or Bitter Almond Green. In the formation of the leuco-base, the following reaction takes place:

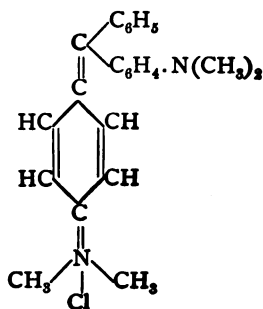


Tetramethyldiamidotriphenylmethane
= Leuco-base of Malachite Green

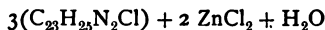
The substance thus obtained is not a dye, but the reduction product of the real dye, which, on oxidation, passes over to the dye. Formerly, the dye formation was believed to take place in accordance with the following equation:



The union, effected by the oxidation between the pentavalent nitrogen atom of the dimethyl aniline residue and the common methane-carbon atom, was considered to be the condition which determined the nature of the dye. At present, the view that the latter is determined by the presence of the quinone-like secondary benzene residue is generally accepted, and the formula of the dye-salt is written thus:

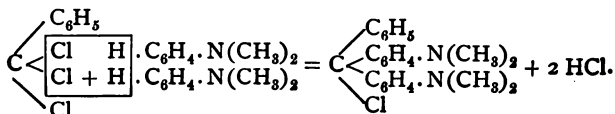


Further, it may be pointed out in this place that, in the formation of the leuco base, the hydrogen atom in the para position to the dimethyl-amido groups $N(CH_3)_2$ unites with the aldehyde oxygen atom to form water. The salt of the formula given above is difficult to separate from its solution. But if zinc chloride is added, a double salt of the same colour is formed:



which may be separated from its water solution by common salt.

Malachite Green may also be made by a second method, which was discovered by Döbner: it consists in heating benzotrichloride with dimethyl aniline in the presence of zinc chloride:



Since the chlorine atom remaining over migrates toward a nitrogen atom, the dyestuff salt is directly formed by the transformation. Still, since the preparation of pure benzotrichloride on the large scale is difficult, this method, which was formerly used, has been abandoned, and the dye is now prepared exclusively by Fischer's method.

Malachite Green is a representative of a whole series of dyes, — the Malachite Green Series. If, instead of dimethyl aniline, diethyl aniline is used, an analogous substance, which bears the name of Brilliant Green, is formed. In place of benzaldehyde, substituted benzaldehydes, etc., can be used. The dyes of the Bitter Almond Series colour only the animal fibres, silk and wool, directly. Vegetable fibre (cotton) is not coloured unless it has been previously mordanted.

42. REACTION: CONDENSATION OF PHTHALIC ANHYDRIDE WITH A PHENOL TO FORM A PHTHALEIN

EXAMPLE: (a) Fluorescein.¹ (b) Bromination of Fluorescein with the Formation of Eosin

(a) In a mortar grind up and intimately mix 15 grammes of phthalic anhydride with 22 grammes of resorcinol, and heat in an

¹ A. 183, 1.

oil-bath to 180° (Fig. 78). As a vessel for heating the mixture, an "extract of beef" jar, glazed inside, is well adapted to the purpose; it can be obtained readily at a small cost, and may be used several times for the same fusion. It is suspended by its projecting edge from a triangle into the oil-bath. To the fused

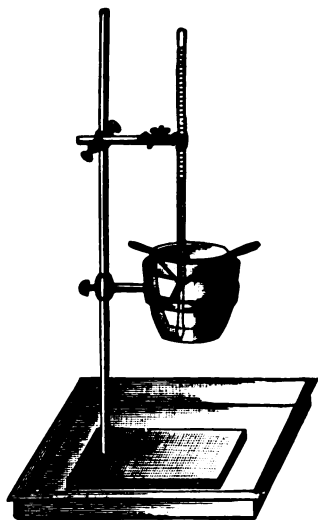


FIG. 78.

mass add, with stirring (glass rod), in the course of 10 minutes, 7 grammes of pulverised zinc chloride. This is prepared in the following way: 10 grammes of the commercial salt, which always contains water, is carefully heated to fusion over a free flame in a porcelain dish. After the mass has been kept in a fused condition for a few minutes, it is allowed to cool, and the solidified substance is removed from the dish with a knife and pulverised. After adding 7 grammes of the anhydrous salt thus obtained, the temperature is increased to 210° , and the heating continued until the liquid, which gradually thickens, becomes solid,

for which about 1-2 hours is required. The cold friable melt is removed from the crucible with a sharp instrument (it is best to use a chisel), finely pulverised, and boiled 10 minutes in a porcelain dish with 200 c.c. of water and 10 c.c. of concentrated hydrochloric acid. This causes the solution of the substance which did not enter into the reaction; the addition of hydrochloric acid is necessary to dissolve the zinc oxide and basic zinc chloride. The fluorescein is filtered from the solution, washed with water until the filtrate no longer gives an acid reaction; it is then dried on the water-bath. Yield, almost quantitative.

(b) Over 15 grammes of fluorescein in a flask, pour 60 grammes

of alcohol (about 95 %), add, with frequent shaking, 33 grammes of bromine, drop by drop, from a separating funnel. This should require about a quarter-hour. In place of a separating funnel, it is advisable, as in all cases of bromination, to use a burette, by which the troublesome weighing of bromine is obviated. Since the specific gravity of bromine at moderate temperatures is very nearly 3, it is only necessary to divide the required weight by 3, in order to find the number of cubic centimetres corresponding to the weight. Of the numerous kinds of burettes, the one best adapted to this

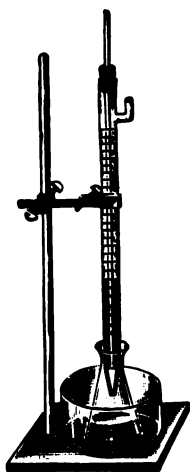


FIG. 79.

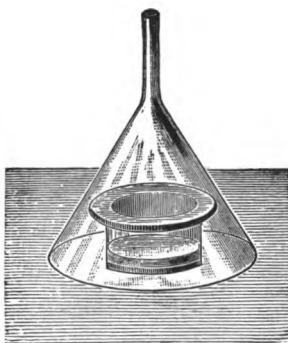


FIG. 80.

purpose is the Winckler form ; since it possesses no cock, it can be inserted into the body of a flask with a not too narrow neck, and by this manipulation the disagreeable bromine vapours may be avoided (Fig. 79). In the above case, 11 c.c. of bromine are necessary. On the addition of bromine, it is observed that the quantity of fluorescein insoluble in alcohol steadily decreases, and that when about one-half of the bromine has been added, a clear, dark, reddish-brown solution is formed. This is due to the fact that the dibromide is first formed, which is easily soluble in

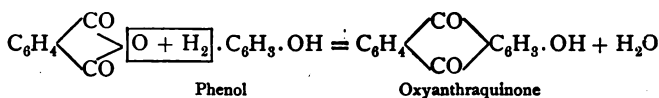
alcohol. On the further addition of bromine, the tetra-bromide is formed, which, since it is difficultly soluble in alcohol, separates out in the form of brick-red leaflets. After all the bromine has been added, the reaction-mixture is allowed to stand for 2 hours, the precipitate is filtered off, washed several times with alcohol, and dried on the water-bath. The product thus obtained is a compound of 1 molecule of eosin and 1 molecule of alcohol. In order to obtain pure eosin from it, the substance is heated a half-hour in an air-bath at 110° : during the heating, its colour becomes brighter. Since eosin is insoluble in water, the soluble potassium-, sodium-, or ammonium-salt is prepared on the large scale for dyeing.

Ammonium Eosin.—Over a flat-bottom crystallising dish, $\frac{1}{8}$ filled with a concentrated ammonia solution, place a filter, of paper as strong as possible. Upon this is spread the eosin acid, in a layer about $\frac{1}{2}$ cm. thick, and the whole is covered with a funnel (Fig. 80). The bright-red crystals of the free eosin acid very soon assume a darker colour, and, after about three hours, it is completely converted into the ammonium salt, which forms dark-red crystals with a greenish lustre. The end of the reaction is easily recognized, by testing a small portion with water. If it dissolves, the conversion is complete.

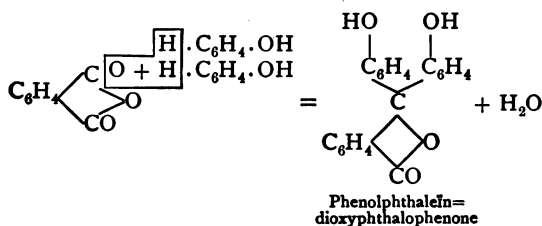
On the large scale, this reaction is carried out in wooden chests containing a number of frames covered with coarse linen, arranged like drawers. After the eosin is spread out on the linen in thin layers, dry ammonia evolved from ammonium chloride and lime is passed into the chest, until a test-portion of the substance will completely dissolve.

Sodium Eosin.—Grind 6 grammes of eosin with 1 gramme of dehydrated sodium carbonate, and in a not too small beaker moisten it with a little alcohol; after the addition of 5 c.c. of water, heat on the water-bath until the evolution of carbon dioxide ceases. To the water solution of sodium eosin thus obtained, add 20 grammes of alcohol, heat to boiling, and filter the hot solution. On cooling, the soluble sodium salt separates out in the form of splendid, brownish-red needles of a metallic lustre. As is the case with many dyes, the crystallisation requires a long time; one day, at least, is necessary.

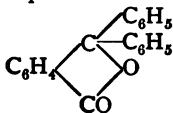
Phthalic anhydride and phenols can react with each other in two different ways. (1) An *equal* number of molecules of each can condense, the oxygen atom of the anhydride, which unites the carbonyl groups, can combine with two ring-hydrogen atoms of the phenol to form one molecule of water; this action results in the formation of an anthraquinone derivative:



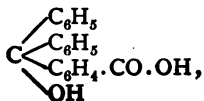
Or (2) *one* molecule of the anhydride can react with *two* molecules of the phenol in such a way that one of the two carbonyl-oxygen atoms of the former combines with one ring-hydrogen of the two phenol molecules to form a so-called phthalein:



For the knowledge concerning this class of compounds, to which belong numerous important dyestuffs, we are indebted to the investigations of A. Baeyer (1871). Phthalophenone is considered to be the mother-substance of the group:



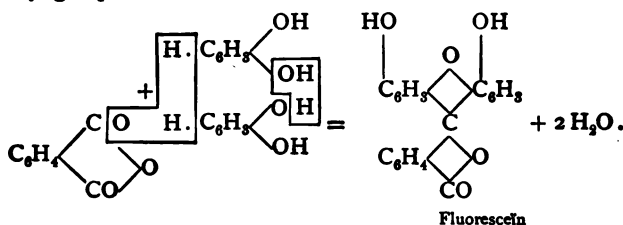
which, as already stated, is obtained from phthalyl chloride and benzene in the presence of aluminium chloride. If one conceives that the mother-substance can take up one molecule of water, a hypothetical mono-carbonic acid of triphenyl carbinol would result:



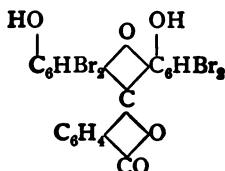
the formula of which shows very clearly the connection between the phthaleins and the triphenyl methane derivatives.

If, as expressed by the above equation, phthalic anhydride is allowed to act on phenol, phenolphthalein is obtained, a substance of acid properties, colourless in the free condition; its salts are red. It is used as an indicator in volumetric analysis.

By the action of phthalic anhydride on resorcinol, the formation of a *tetraoxyphthalophenone* would naturally be expected; but fluorescein, containing the constituents of one molecule of water less than this, is obtained, an inner anhydride formation taking place between the two hydroxyl groups:



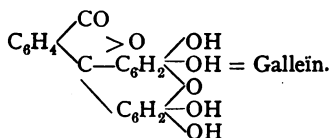
Fluorescein is technically prepared on the large scale by the method given above. While phenolphthalein, in spite of the intense colour of its salts, is not a dye, in that it does not colour fibres, fluorescein is a true dye which colours animal fibres a fast yellow. But it is not manufactured as a dye, since it has been replaced by other dyes that give as beautiful colours and are cheaper. A number of its halogen- and nitro-substitution products have valuable colouring properties, and are prepared from it. The simplest dye of this kind is eosin or tetra-brom-fluorescein, discovered in 1874 by Caro. The four bromine atoms are equally divided between the two resorcinol residues:



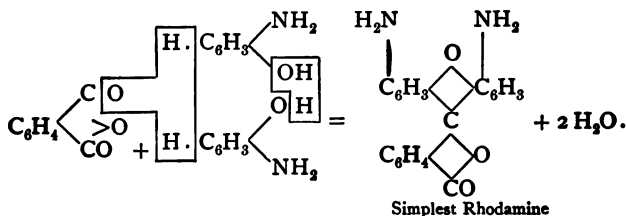
which follows from the fact that eosin in fusion with potassium hydroxide yields di-bromoresorcinol besides phthalic acid. Instead of phthalic anhydride, the di- and tetra-chlor-substitution products are fused with

resorcinol on the large scale; and so there is obtained in the phthalic acid residue, the di- and tetra-chlorfluoresceins from which halogen substitution products, nitro-derivatives, ethers, etc., are prepared on the large scale (Phloxine, Rose Bengal).

Besides fluorescein there is practically only one other phthalein prepared technically, Gallein. This is done by heating phthalic anhydride with the *ν*-trioxybenzene—pyrogallol. In this case the same anhydride formation takes place as in the preparation of fluorescein:



From gallein, a derivative of anthracene, coerulein, a new dye, is obtained by heating with sulphuric acid. Since 1887 the phthaleins have been on the market under the name of rhodamines, which are prepared in a manner similar to that of fluorescein, except that instead of resorcinol, *m*-amidophenol, or amidophenols substituted by alkyls in the amido-group, are used:



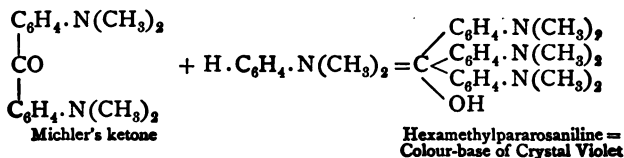
The rhodamine on the market is the tetra-ethyl derivative of this mother-substance.

43. REACTION: CONDENSATION OF MICHLER'S KETONE WITH AN AMINE TO A DYE OF THE FUCHSINE SERIES

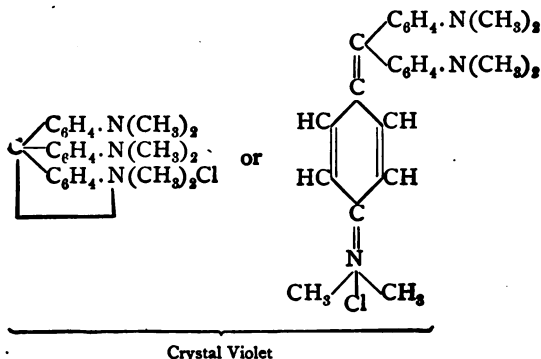
EXAMPLE: Crystal Violet from Michler's Ketone and Dimethyl Aniline

A mixture of 25 grammes of dimethyl aniline, 10 grammes of Michler's ketone (this is on the market), and 10 grammes of phosphorus oxychloride, is heated in an open, dry flask, 5 hours, on an actively boiling water-bath. The blue-coloured mass is then poured into water, made alkaline with a solution of caustic soda, and treated with steam until no drops of the unattacked dimethyl aniline pass over. After cooling, the solidified colour-base remaining in the distillation flask is filtered from the alkaline solution, washed with water, and boiled with a mixture of 1 litre of water and 5 grammes of concentrated hydrochloric acid. The blue solution is filtered while hot from the colour-base, which remains undissolved; the latter is boiled again with a fresh quantity of dilute hydrochloric acid; this operation is repeated until the substance has been almost entirely dissolved. After cooling, the solution of the dye is treated with finely pulverised salt (stirring) until the dye is precipitated. It is then filtered with suction, pressed out on a porous plate, and crystallised from a little water. On cooling, the Crystal Violet separates out in coarse crystals of a greenish colour; these are filtered off and dried in the air on filter-paper.

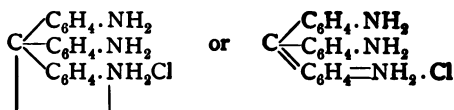
If Michler's ketone is heated with an amine in the presence of a condensation agent (phosphorus oxychloride, POCl_3), addition takes place, in accordance with the following equation:



If this is dissolved in hydrochloric acid, one molecule of this is added, and, as in the formation of Malachite Green, the elimination of a molecule of water immediately takes place and the dye is formed :



It is a derivative of parafuchsine :



indeed, it may be considered as a hexamethyl parafuchsine. It is prepared technically in the same way, and forms the principal constituent of the Methyl Violet obtained by the oxidation of dimethyl aniline.

Dyes can also be prepared in the same way by the combination of other amines with Michler's ketone, of which it is only possible to mention here Victoria Blue and Night Blue.

44. REACTION: CONDENSATION OF PHTHALIC ANHYDRIDE WITH A PHENOL TO AN ANTHRAQUINONE DERIVATIVE

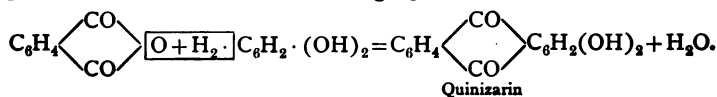
EXAMPLE: Quinizarin from Phthalic Anhydride and Hydroquinone¹

A mixture of 5 grammes of pure hydroquinone and 20 grammes of phthalic anhydride is heated in an open flask with a mixture of 100 grammes of pure concentrated sulphuric acid and 10

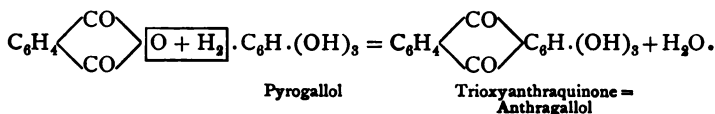
¹ B. 6, 506; 8, 152; A. 212, 10.

grammes of water for 3 hours in an oil-bath to 170–180°, and finally for 1 hour at 190–200°. The directions as to time and temperature must be followed as exactly as possible. The hot solution is poured, with stirring, into about 400 c.c. of water in a porcelain dish, heated to boiling, and filtered hot with the aid of a Büchner funnel. The residue remaining on the filter is again boiled out with water and filtered while hot. In order to separate the quinizarin from carbonaceous decomposition products, the precipitate is boiled with 200 c.c. of glacial acetic acid, filtered hot with suction, the filtrate poured into a beaker, and, while hot, treated with its own volume of hot water. The residue remaining on the filter is again boiled up with 100 c.c. glacial acetic acid, and, after filtering, treated as above. On cooling of the diluted acetic acid solution, the crude quinizarin separating out is filtered off, washed with water several times, dried first on the water-bath, and finally in an air-bath at 120°. Since it is difficult to obtain it pure by crystallisation, after drying it is distilled from a small retort of difficultly fusible glass, and is driven over as rapidly as possible with a large flame. A beaker is used as a receiver. It is more convenient to use a porcelain mortar, and a porcelain dish as a cover. After the distillate in the receiver and that in the neck of the retort (this is broken) has been finely pulverized, it is crystallized from glacial acetic acid, from which, on cooling, the quinizarin separates out in the form of large, orange-yellow leaves; these are filtered off and washed with glacial acetic acid, which is steadily diluted with water, until finally only pure water is used. Better crystals (dark-red compact needles) may be obtained by dissolving the distilled quinizarin in toluene, heated on a water-bath. The filtered crystals are washed first with toluene and then with alcohol.

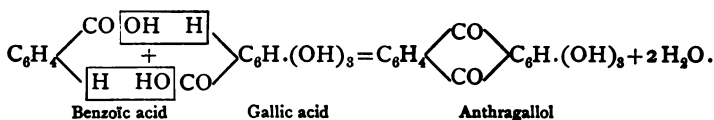
Under the preparation of fluorescein, it has already been mentioned that phthalic anhydride condenses with phenols in certain proportions, to form derivatives of anthraquinone. The reaction just effected takes place in accordance with the following equation :



In an analogous way, mono-acid- as well as poly-acid phenols, condense with phthalic anhydride. It is of theoretical importance that from pyrocatechol (o-dioxybenzene), besides a second isomer, alizarin is obtained, showing that the two hydroxyl groups in alizarin are in the ortho position to each other. Of practical significance is the above reaction for the preparation of anthragallol, which is obtained on the large scale by heating pyrogallol with phthalic anhydride:



It may be mentioned briefly that by the condensation of benzoic acid with oxybenzoic acids, similar compounds are also obtained:



Quinizarin dissolves, like oxyanthraquinones, in alkalis with a violet colouration. (Try it.)

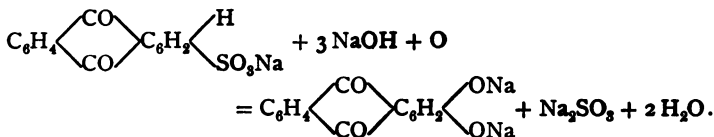
45. REACTION: ALIZARIN FROM SODIUM β -ANTHRAQUINONE-MONOSULPHONATE¹

In an autoclave or an iron pipe with a cap which can be screwed on (see page 69), heat a mixture of 10 parts commercial sodium anthraquinonemonosulphonate, 30 parts of sodium hydroxide, 1.8 parts of finely pulverised potassium chlorate, with 40 parts of water, for 20 hours to 170°. After cooling, the melt is boiled out with water several times, and acidified at the boiling-point of the solution in a large dish with concentrated hydrochloric acid. The alizarin separating out is then filtered off according to the quantity, either with suction or with the aid of a filter-press, washed with water, pressed out on a porous plate, and dried in an air-bath at 120°. In order to obtain it completely pure, it is distilled *rapidly* from a small retort, and is

¹ A. Spl. 7, 300; B. 3, 359; 9, 281.

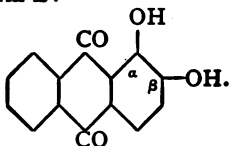
crystallised from glacial acetic acid, or in large quantities from nitrobenzene.

The sodium hydroxide fusion of the sodium anthraquinonemonosulphonate is an abnormal reaction to the extent that besides the replacement of the sulphonic acid group by hydroxyl, a hydrogen atom is also oxidised to a hydroxyl group:



The tendency to the formation of alizarin is so great that even without the addition of an oxidising agent (potassium chlorate or nitrate), it is formed with the evolution of hydrogen. Formerly the oxygen of the air was used as the oxidising agent, the reaction being effected in air.

In order to prepare alizarin on the large scale, anthracene is the starting-point; this is obtained from the highest-boiling fractions of coal tar (anthracene oil). It is oxidised by chromic acid to anthraquinone (see below), and this on heating with sulphuric acid is converted into the monosulphonic acid. The separation of this latter compound is greatly facilitated by the fact that it forms a sodium salt difficultly soluble in water, which, on account of its silvery appearance, is called "Silver salt." If the sulphonation mixture is diluted with water and neutralised with sodium carbonate, the sodium anthraquinonemonosulphonate is precipitated directly, which thus obviates the necessity of removing the excess of sulphuric acid beforehand. On the large scale the alizarin fusion is conducted exactly as on the small scale, except that autoclaves, with stirring attachments, are used. The constitutional formula of alizarin is:



The salts are intensely coloured. The red aluminium salt, the violet ferric salt, and the garnet-brown chromic salt are especially im-

portant in dyeing. With alizarin and all its related compounds the dyeing is effected by mordanting the fibre with a salt of one of the three oxides just mentioned; the thus prepared fibre is heated with a thin dilute water-paste of the free insoluble dye, whereby salts are formed on the fibre (Lakes).

From two disulphonic acids of anthraquinone, two trioxyanthraquinones, flavo- and anthra-purpurin, are prepared in a manner analogous to that by which alizarin is obtained from the monosulphonic acid.

From alizarin there can be prepared, further, by nitration, the α - or β -nitro-alizarin, and from this, by reduction, the corresponding amido-alizarin. From β -nitro- and amido-alizarin, by heating with glycerol and sulphuric acid, the important Alizarin Blue is obtained. Further, by the action of fuming sulphuric acid on alizarin there is obtained a tetraoxyanthraquinone (Bordeaux), etc.

46. REACTION: ZINC DUST DISTILLATION

EXAMPLE: Anthracene from Alizarin or Quinizarin

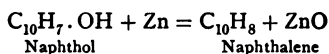
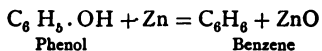
To a paste prepared by rubbing up 100 grammes of zinc dust with 30 c.c. of water, add pieces of porous pumice stone of a size that will conveniently pass into a combustion tube, and stir them around so that they become covered with the zinc dust paste. They are removed from the paste with pincers, heated in a porcelain dish over a free flame (in constant motion) until the water is evaporated. A combustion tube of hard glass 60-70 cm. long is drawn out at one end to a narrow tube, the narrowed end is closed by a loose plug of asbestos, and a layer of zinc dust 5 cm. long is placed next to the plug; then follows a mixture of $\frac{1}{2}$ -1 gramme of alizarin or quinizarin with 10 grammes of zinc dust, and finally, a layer of pumice-zinc dust 30 cm. long. After a canal has been formed over the zinc dust, by placing the tube in a horizontal position and tapping it, the tube is transferred to a combustion furnace inclined at an oblique angle, and dry hydrogen is passed through the tube without heating. In order to test whether the air has been completely expelled from the tube, the

open end is closed by a cork bearing a small glass tube to which is attached a piece of rubber tubing; the gas being evolved is conducted into a soap solution, and the bubbles formed are ignited, during which the greatest care must be taken to keep the flame from coming in contact with the gas issuing from the rubber tubing, otherwise a serious explosion may result. If an explosion accompanied by a report takes place when the bubbles are ignited, the air has not been completely removed, but if they burn quietly, then only pure hydrogen is present.¹ When this is the case, the current of gas is diminished so that only two bubbles per second pass through the wash-bottle; the pumice-zinc dust is then heated with small flames, these are increased in size gradually, and finally, the tiles being placed in position, it is heated as strongly as possible; then the rear layer of 5 cm. of zinc dust is similarly heated, and as soon as this glows, as in the nitrogen determination, the mixture of the substance and zinc dust is gradually heated. The anthracene formed condenses to crystals in the forward cool part of the tube. After the reaction is complete, while the tube is allowed to cool, a moderately rapid current of hydrogen is passed through it; the forward part of the tube containing the anthracene is broken off and the substance removed with a small spatula; it is purified by sublimation in a suitable apparatus (see pages 14 and 15). Melting-point, 213° .

The sublimed anthracene is dissolved by heating in a test-tube with a little glacial acetic acid; it is treated with about double its weight of chromic anhydride, and heated a short time to boiling. The solution is then diluted with several times its volume of water, the anthraquinone separating out is filtered off, washed with some dilute sulphuric acid, then with water, and is finally crystallised in a test-tube from a little glacial acetic acid. Long colourless needles of anthraquinone, which melt at 277° , are thus obtained.

¹ As described under Carbon Monoxide, the test may also be made by filling a test-tube with the gas over water, and applying a match to the mouth of the tube.

Zinc dust is, especially at high temperatures, an excellent reducing agent (Baeyer, A. 140, 205), which can be used for the reduction of almost all aromatic oxygen compounds derived from hydrocarbons, *e.g.*:



Also ketone-oxygen, as the above example shows, can be replaced by hydrogen. The reaction given under Alizarin possesses an historical interest, since, by means of it, Gräbe and Liebermann, in 1868, discovered that alizarin, which had been previously obtained from madder root, was a derivative of anthracene, and could be prepared synthetically from it. (B. 1, 43.)

III. PYRIDINE AND QUINOLINE SERIES

1. REACTION: THE PYRIDINE SYNTHESIS OF HANTZSCH¹

EXAMPLE: Collidine = Trimethylpyridine

Dihydrocollidinedicarboxylic Acid Ester.—A mixture of 25 grammes of acetacetic ester and 8 grammes of aldehyde-ammonia is heated in a small beaker on a wire-gauze, about three minutes, to 100–110°, the mixture being stirred with the thermometer. The warm reaction-mixture is then treated with double its volume of dilute hydrochloric acid, and stirred vigorously without further heating until the liquid mass solidifies. It is then thoroughly triturated in a mortar, filtered, washed with water, and dried, either by pressing out, or by warming on the water-bath. For the further working up of the collidinedicarboxylic acid ester, the crude product can be directly used. In order to obtain the dihydroester in a crystallised condition, 2 grammes of the crude product are dissolved in a small quantity of alcohol in a test-tube, by heat, and allowed

¹ A. 215, 1.

to cool slowly. Colourless tablets with a bluish fluorescence are thus obtained. Melting-point, 131° .

Collidinedicarbonic Acid Ester.—The crude dihydroester is treated in a small flask with an equal weight of alcohol; complete solution does not take place.

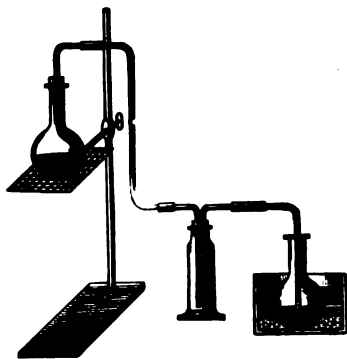


FIG. 81.

Into the mixture cooled by water pass nitrous fumes (Fig. 81), until the dihydroester goes into solution, and a test-portion dissolves to a clear solution in dilute hydrochloric acid. The alcohol is then evaporated by heating on the water-bath, the thick residue is treated with a sodium carbonate solution to alkaline reaction; the oil separating out is taken up with ether.

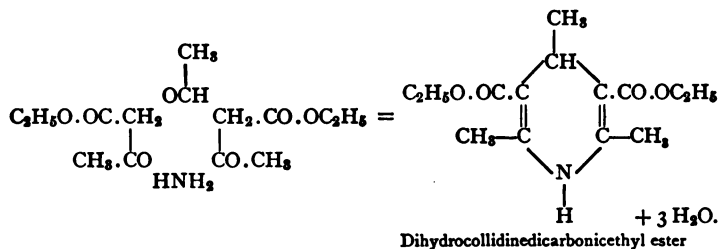
After the ethereal solution has

been dried by a small piece of potassium hydroxide, or potash, the ether is evaporated, and the residue subjected to distillation; on account of the high boiling-point of the ester, a fractionating flask is selected, having the condensation tube as near as possible to the bulb. The fraction passing over between $290-310^{\circ}$ can be used for the following experiment:

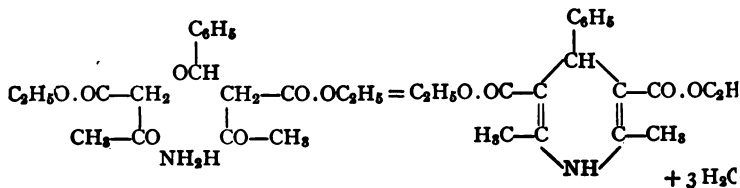
Potassium Collidine Dicarbonate.—The saponification of the ester is effected by boiling with alcoholic potash, prepared in the following manner: Finely pulverised potassium hydroxide (2 parts to 1 part of ester) is moderately heated in a flask on a wire-gauze with 3 times its weight of absolute alcohol, until the greater portion has passed into solution. The alcoholic solution is then poured off from the portion remaining undissolved, treated with the ester to be saponified, and heated 4–5 hours on a rapidly boiling water-bath (with reflux condenser); the potassium salt separates out in crusts. The alcoholic liquid is then poured off from the salt, and the latter washed on the filter with alcohol and finally with ether.

Collidine.—The dried potassium salt is intimately mixed in a mortar with double its weight of slaked lime, and placed in one end of a hard glass tube (about 2 cm. wide and 55 cm. long). In order to prevent the mixture from being carried over into the receiver on heating, a small, loose plug of asbestos is placed in the tube in front of it. After a canal has been made by tapping, the tube is connected with an adapter bent downwards, by means of a cork or asbestos paper; it is then transferred to a combustion furnace, the rear end of which is somewhat elevated and warmed throughout its entire length with small flames, beginning at the closed end. The flames are steadily increased in size until, with the tiles in position, the tube is heated as strongly as possible. The collidine passing over is taken up with ether, dried with potassium hydroxide, and; after the evaporation of the ether, is subjected to distillation. Boiling-point, 172° .

On heating acetacetic ester with aldehyde-ammonia, the following reaction takes place (see A. 215, 8) :



The reaction may be modified by using other aldehydes instead of acet-aldehyde; thus there is obtained from benzaldehyde, acetacetic ester, and ammonia, the dihydrophenyllutidinededicarbonic ester :

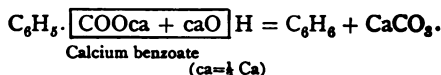


With propionic aldehyde, butyraldehyde, valeraldehyde, cœnanthol, myristic aldehyde, nitrobenzaldehyde, phenylacetaldehyde, furfural, and others, the reaction can be carried out. All the compounds obtained contain the methyl groups of the two acetacetic ester molecules, but the third side-chain is different, depending upon the nature of the aldehyde employed.

By passing nitrous fumes into an alcoholic solution of the dihydro-ester, two hydrogen atoms, and those particular hydrogen atoms in combination with carbon and nitrogen in the methenyl- and imido-groups, respectively, will be oxidised off, and there is formed a derivative of pyridine, containing no ring hydrogen. While the dihydro-esters possess no basic properties, the pyridine derivative dissolves in acid. Therefore, by treating the solution with hydrochloric acid, it can be determined whether any unchanged dihydroester (insoluble in acid) is present.

Concerning the saponification of the ester, refer to what was said under Reaction 36.

The splitting off of carbon dioxide from a carbonic acid, or a salt of a carbonic acid, is generally designated as a "pyro-reaction." For this kind of action a calcium salt is most frequently used; this is mixed with slaked lime and subjected to distillation, *e.g.* :



In poly-basic acids, all the carboxyl groups can be replaced by hydrogen. In this way an acid may be transformed into the hydrocarbon from which it was derived. In the above case, the potassium salt may be used instead of the calcium salt.

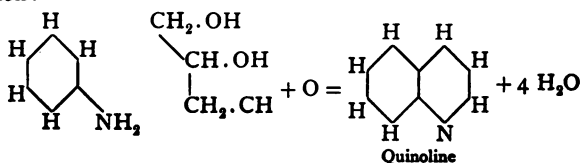
2. REACTION: SKRAUP'S QUINOLINE SYNTHESIS

EXAMPLE : Quinoline

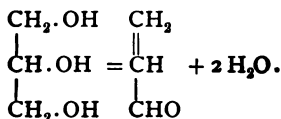
In a flask of about $1\frac{1}{2}$ litres capacity containing a mixture of 24 grammes of nitrobenzene, 38 grammes of aniline, and 120 grammes of glycerol, add, with stirring, 100 grammes of concentrated sulphuric acid. The flask is then connected with a long, wide reflux condenser, and heated on the sand-bath. As soon

as the reaction begins, which is recognised by the sudden evolution of bubbles of vapour ascending through the liquid, the flame is removed, and the energetic reaction is allowed to complete itself without further heating from without. When the reaction-mixture has become quiet, it is again heated for three hours on the sand-bath, diluted with water, and from the acid liquid the unchanged nitrobenzene is removed with steam. As soon as no drops of oil pass over, the distillation with steam is discontinued. The liquid remaining in the distillation flask is allowed to cool somewhat, and then made alkaline with concentrated caustic soda solution, upon which the liberated quinoline, mixed with the unchanged aniline, is distilled over with steam. Since these substances cannot be separated by fractional distillation, their separation must be effected by a chemical method. For this purpose the distillate (oil and water solution) is treated with dilute sulphuric acid until all oil is dissolved and an excess of the acid is present; to the cold solution a solution of sodium nitrite is added until a drop of the liquid will cause a blue spot on potassium iodide-starch paper; if the blue colour does not appear, add more sulphuric acid to the mixture. The aniline (primary amine) is converted into diazobenzenesulphate, while the tertiary quinoline remains unchanged. The mixture is heated for some time on the water-bath, by which, as in Reaction 8, the diazo-sulphate is converted into phenol. The liquid is again made alkaline, upon which the phenol goes into solution, while the quinoline is liberated. The mixture is now distilled with steam, and the quinoline is obtained in a pure condition: it is taken up with ether, the ether evaporated, and the residue distilled. Boiling-point, 237° . Yield, 40-45 grammes. (See Wiener Monatshefte 2, 141.)

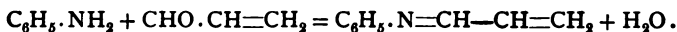
Quinoline is formed in the above reaction according to the following equation:



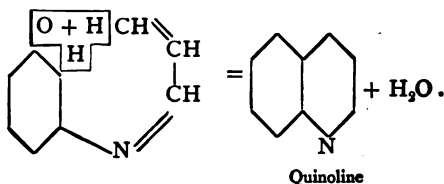
The oxygen necessary for the reaction is taken from the nitrobenzene, which is hereby reduced in a manner that is not wholly clear. It is possible that the reaction may take place in this way: first, acrolein is formed from glycerol, under the influence of sulphuric acid:



Like all aldehydes, this condenses with aniline to form acrolein-aniline.

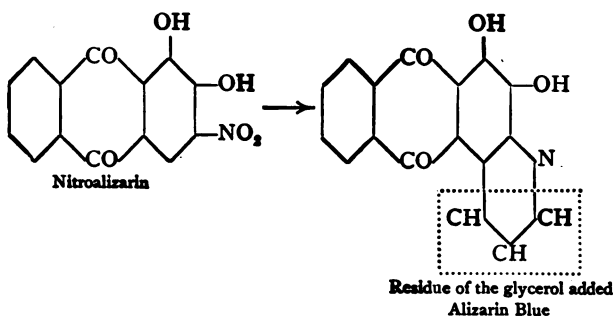


While this, under the influence of the oxidising action of the nitro-compound, loses two atoms of hydrogen, and thus quinoline is formed:



The Skraup reaction is capable of a very many-sided application. If, instead of aniline, its homologues are used, methyl-, dimethyl-aniline, etc., the corresponding quinoline is obtained. Also halogen-, nitro-, etc., substituted amines, yield halogen-, nitro-, etc., substituted quinolines. Amidocarbonic acids, amidosulphonic acids, amidophenols, yield carbonic acid-, sulphonic acid- or oxy-derivatives of quinoline. The reaction is also applicable to the corresponding amido-compounds of the naphthalene series. By starting from the diamines, two new pyridine rings, connected with the benzene ring, are formed; in this way the so-called phenanthrolines, etc., are obtained.

Of technical and historical interest is the discovery which was made by Prudhomme in the year 1877, that β -nitroalizarin, on heating with glycerol and sulphuric acid, yields a blue dye, Alizarin Blue. This gave the impetus to Skraup's synthesis. To Gräbe's investigations we are indebted for the knowledge of the process by which, as above, a quinoline synthesis is effected in the following way:



IV. INORGANIC PART

1. CHLORINE

A flask is one-third filled with manganese dioxide (pyrolusite) in pieces the size of filberts; to this is added a quantity of concentrated hydrochloric acid which is just sufficient to cover it. On heating the mixture on a wire gauze with a free flame, a regular current of chlorine is generated; this is passed through two wash-bottles containing water and concentrated sulphuric acid respectively; the water retains any hydrochloric acid which is carried along with the gas, and the sulphuric acid dries it. (See Figs. 74 and 87.) A piece of thin asbestos-paper is placed on the wire gauze, as is always done on heating large flasks, by which the danger of breaking is essentially diminished. A very regular current of chlorine can also be obtained from finely pulverised potassium dichromate and crude concentrated hydrochloric acid by heating the mixture on the water-bath. To 1 litre of hydrochloric acid, use 180–200 grammes of pulverised potassium dichromate.

Concerning the preparation of chlorine from potassium permanganate and hydrochloric acid see B. 35, 43.

2. HYDROCHLORIC ACID

Gaseous hydrochloric acid, which is frequently needed for the preparation of acid-esters, is generated most conveniently in a

Kipp apparatus charged with fused ammonium chloride in pieces as large as possible, and concentrated sulphuric acid. The operation is conducted in the same way as that for the generation of carbon dioxide or hydrogen from a Kipp apparatus.

If the apparatus is not available, the acid can be generated very conveniently in the following manner :

In concentrated hydrochloric acid contained in a suction flask allow to flow from a separating funnel concentrated sulphuric acid, drop by drop (Fig. 82). The hydrochloric acid evolved is dried by passing it through concentrated sulphuric acid contained in a

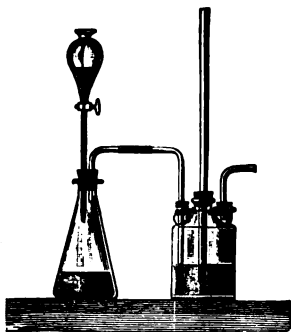


FIG. 82.

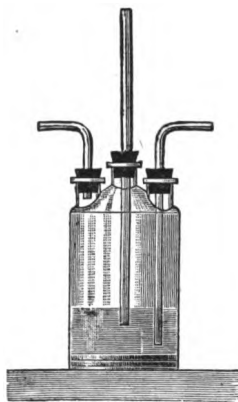


FIG. 83.

safety wash-bottle (Fig. 83) ; this latter is always used, since otherwise, with an irregular gas current, the liquid to be saturated may be easily drawn back into the wash-bottle and then into the generating mixture. In place of a Woulff-flask with three tubulures, a single-neck wash-bottle may be converted into a safety-bottle as follows (see Fig. 84) : Into a two-hole cork place a straight tube as wide as possible ; through this insert a narrow delivery tube, bent at a right angle, which reaches almost to the bottom of the bottle.

The liquid to be saturated cannot flow back into the wash-bottle with this arrangement, since in case there should be a tendency to do so, air would enter the suction-flask through the space between the delivery tube and the wider tube, thus relieving

the pressure. If a wash-bottle having a side-tube is available, it can also be converted into a safety-tube (see Fig. 85).

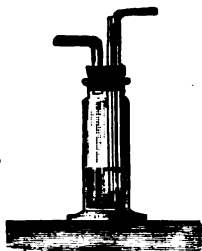


FIG. 84.

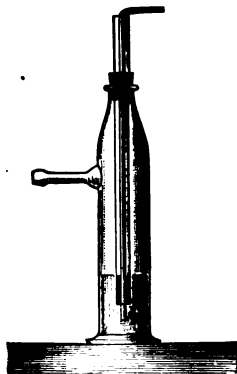


FIG. 85.

Hydrochloric acid gas may also be obtained by warming 10 parts of sodium chloride with a cold mixture of 3 parts of water and 18 parts of concentrated sulphuric acid.

3. HYDROBROMIC ACID (see Brombenzene)

The hydrobromic acid obtained as a by-product in the bromination reactions is purified by distilling it from a fractionating flask. Water first passes over until finally the temperature remains constant at 126° , when a 48 % acid goes over ; this is collected.

In order to prepare potassium bromide for use in the preparation of ethyl bromide, the acid is diluted with some water and then treated with dry potash until there is no further evolution of carbon dioxide and the liquid shows a neutral reaction. To 1 part of hydrobromic acid 0.5 part potassium carbonate is used. The water solution of the potassium bromide is evaporated to dryness on a water-bath. The product thus obtained may be used directly for the preparation of ethyl bromide.

4. HYDRIODIC ACID

To 44 grammes of iodine (not pulverised) contained in a small round flask of about 100 c.c. capacity (Fig. 86), gradually add 4 grammes of yellow phosphorus divided into about 8 pieces under

water ; these are dried just before transferring them to the flask, by pressing between layers of blotting-paper. The first piece of phosphorus added unites with the iodine with an active evolution of heat and light. When the first action is ended, after shaking the contents of the flask, which soon become liquid, the second piece is added. The reaction still proceeds with evident energy, although it is less intense than when the first piece was added. Care is taken to place the phosphorus as nearly as possible in the middle of the flask, and not to allow it to fall on the walls, since otherwise the flask may be easily broken. When all of the phosphorus is added, a fused, dark mass of phosphorus triiodide is obtained which becomes solid on cooling. The hydriodic acid pre-

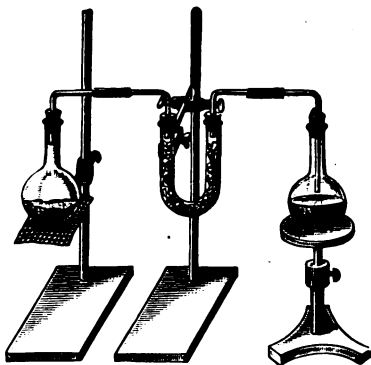


FIG. 86.

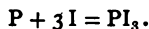
pared from this by warming with water, must be passed over red phosphorus in order to free it from iodine which is carried along with it. Proceed as follows : 5 grammes of red phosphorus are rubbed up to a paste with 2 c.c. of a water solution of hydriodic acid, or in case this is not available, with as little water as possible (1 c.c. at the most). In this is placed glass beads, or bits of broken glass, which on stirring around in the mixture become covered with the paste. They are then transferred to a U-tube. Wide connecting-tubes are used between the generating flask and the U-tube. In order to prepare a water solution of hydriodic acid, the gas issuing from the U-tube is passed into 45 c.c. of water (see Fig. 86). The glass tube is not immersed in the water, but its end must be 1 cm. above the surface ; otherwise, in consequence of the great affinity of water for hydriodic acid, under certain conditions the water may be drawn back.

The hydriodic acid is now obtained by treating the *completely cooled* phosphorus triiodide with 6 grammes of water and warming with a *very small* luminous flame. The contents of the flask steadily become clearer, while in the other flask the heavy layer

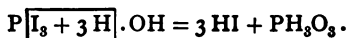
of hydriodic acid sinks to the bottom. The heating is continued until only a clear, colourless liquid remains in the generating flask.

In order to obtain a concentrated solution of hydriodic acid, the liquid in the receiver is distilled. At first a few cubic centimetres of water pass over at 100° , then the temperature rises in a short time to 125° ; the concentrated acid passing over up to 130° is collected separately. This boils for the most part at 127° .

This experiment teaches much concerning the chemistry of phosphorus and iodine. First, it shows that iodine and phosphorus unite directly with a vigorous reaction, to form phosphorus triiodide :



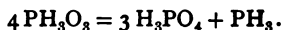
The iodide then decomposes with water, to form hydriodic acid, which is evolved, while the phosphorous acid (H_3PO_3) remains in the flask :



The gaseous hydriodic acid is an intensely fuming substance, which may be easily shown by removing the cork from the receiver containing the aqueous acid, for a moment. Hydriodic acid is absorbed by water with great avidity. The acid, boiling constantly at 127° , contains approximately 50% of anhydrous hydriodic acid.

In this experiment it is observed that the connecting tubes of the apparatus, especially those between the generating flask and the U-tube become coated with crystals of a diamond-like brilliancy. These are crystals of phosphonium iodide, PH_4I , which is formed by the decomposition of phosphorous acid.

It is a common property of all the lower oxidation products of phosphorus, to pass over to the highest oxidation product — phosphoric acid, with the evolution of phosphine on heating. With phosphorous acid, the reaction takes place as follows :



The phosphine thus formed unites, since it possesses weak basic properties, with hydriodic acid, to form phosphonium iodide :



Since this may easily clog the connecting tubes, the tubes selected are as wide as possible. On cleaning the tubes with water, this reacts with the phosphonium iodide with the evolution of phosphine, a gas with a garlic-like odour, and which in this case is not spontaneously inflammable. The phosphonium iodide decomposes with water into its components, in accordance with this equation :



This reaction, as is well known, is employed for preparing pure phosphine which is not spontaneously inflammable.

5. AMMONIA

Gaseous ammonia is prepared most conveniently by heating the *most concentrated* ammonia solution in a flask over a wire gauze with a small flame. In order to dry the gas, it is passed through a drying tower filled with soda-lime. (See Fig. 66.)

6. NITROUS ACID

For the preparation of gaseous nitrous acid, arsenious acid, broken into pieces the size of a pea, is treated with nitric acid, sp. gr. 1.3, and heated gently on a wire gauze with a free flame (under the hood). In order to condense the nitric acid carried along with the gases, an empty wash-bottle, cooled by cold water, is employed. (See Fig. 81.)

7. PHOSPHORUS TRICHLORIDE

Under water, in a porcelain mortar, cut 40 grammes of yellow phosphorus, with a knife or chisel, into pieces which will conveniently pass into the tubulure of a 300 c.c. retort. After the air in the retort has been displaced by dry carbon dioxide (Fig. 87), each single piece of phosphorus is taken from the water by pincers, and dried quickly by pressing it between several layers of filter-paper, and immediately placed in the retort, care being taken to prevent it from becoming ignited by friction in the opening of the tubulure. As soon as all the phosphorus has been transferred

to the retort, the tubulure is connected with a delivery tube which must move easily in the cork, and a moderately rapid current of

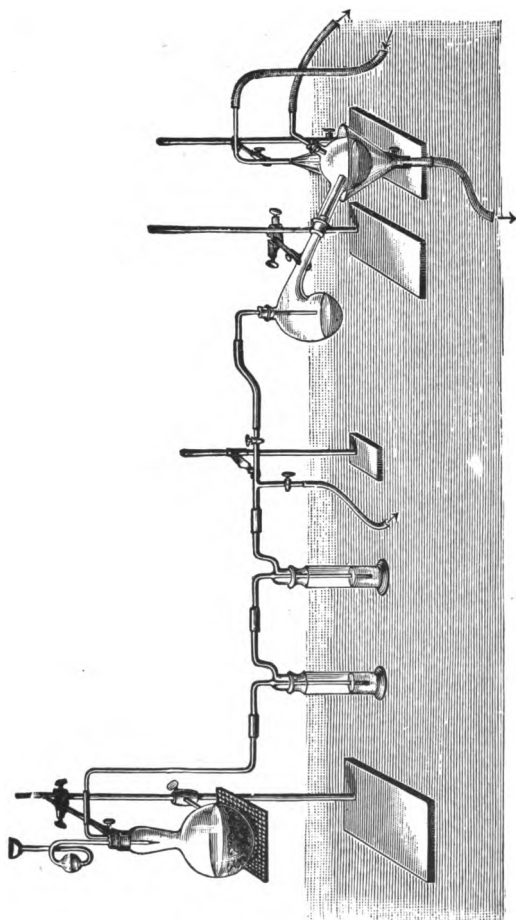


FIG. 87.

dry chlorine passed over the phosphorus ; phosphorus chloride is thus formed with evolution of heat and light. If crystals of phosphorus pentachloride should collect in the neck of the retort, the

delivery tube is pushed somewhat farther into the retort. If, on the other hand, phosphorus distils to the upper part of the retort, the tube is somewhat raised. The phosphorus trichloride condensing in the receiver is distilled from a dry fractionating flask. Boiling-point, 74° . Yield, 125–140 grammes.

8. PHOSPHORUS OXYCHLORIDE¹

To 100 grammes of phosphorus trichloride, contained in a large tubulated retort connected with a condenser, add gradually, in small portions of about 2–3 grammes, 32 grammes of finely pulverised potassium chlorate. After each addition, wait until the liquid bubbles up, before adding a new quantity. If, on the addition of the first portion, no reaction takes place, it is started by a gentle warming. During the addition, no liquid should distil into the receiver, but if this does happen, it is poured back into the retort. After all of the chlorate has been added, the phosphorus oxychloride formed is distilled, by heating the retort in an oil-bath, to 130° , or with a luminous flame. A suction-flask is used as a receiver; this is firmly connected with the end of the condenser, by means of a cork. The distillate is rectified from a fractionating flask provided with a thermometer. Boiling-point, 110° . Yield, 100–110 grammes.

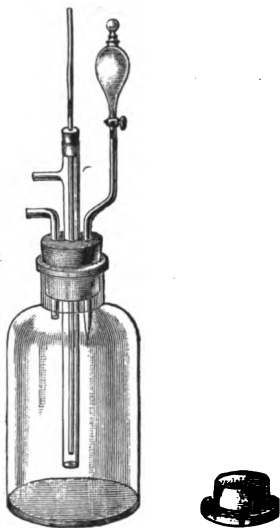


FIG. 88.

9. PHOSPHORUS PENTACHLORIDE

Through the upper delivery tube of an apparatus similar to that represented in Fig. 88, a stream of dry chlorine is admitted, which

¹ J. pr. Ch. 1883, [2] Vol. 23, 382.

passes out of the lower, right-angled tube. From time to time, several cubic centimetres of phosphorus trichloride are allowed to flow into the bottle from a separating funnel, upon which the trichloride unites with the chlorine to form the solid pentachloride. Since this operation can be repeated, as soon as it is evident that the union is completed, any desired quantity of phosphorus pentachloride can be prepared. Should the delivery tube become stopped up, it is cleared by the glass rod with which the apparatus is provided. As the quantity of the pentachloride formed increases, the tube is correspondingly raised. Yield, quantitative.

10. SULPHUROUS ACID

Gaseous sulphurous acid is generated in an apparatus similar to the one represented in Fig. 82, by adding to a concentrated water solution of sodium hydrogen sulphite a cold mixture of equal parts, by volume, of water and concentrated sulphuric acid, drop by drop. The generating flask is shaken frequently, to keep the contents from separating into layers.

11. SODIUM

(a) *To cut Sodium.*—In order to divide sodium into small portions, it can be cut into scales with a knife, or pressed out into a wire with a sodium-press. To cut it into scales, an apparatus similar to that represented in Fig. 89 is convenient. After both sides of the knife and the front part of the table have been coated with a thin layer of vaseline, a long stick of the metal to be cut, the end of which is wrapped in filter-paper, in order that it may be handled, is placed on the table so that it projects somewhat over the front end; it is then cut with a short stroke of the

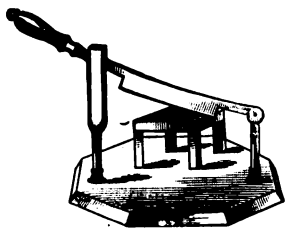


FIG. 89.

knife. On the front part of the lower platform is placed a small dish filled with ether or ligroïn, into which the scales fall. When

using the knife, two points are to be especially observed. The eye is never placed in front of the knife, but always behind it, so that the fingers holding the sodium can always be seen. Only in this way can a wound be prevented. Further, the *cross-section of the piece of sodium must not be too large*, otherwise the metal adheres to the knife. Quadratic scales, the edge of which must not, at most, be more than 5–6 mm. long, are cut. With a little practice, large quantities of the metal can be cut in very thin scales in a short time.

The sodium residues are not thrown into water nor into waste-jars, but are dropped into alcohol contained in a beaker or flask.

(b) *Sodium Amalgam*.—Sodium scales, about the size of a 20-cent piece, are pressed to the bottom of mercury contained in a porcelain mortar, in rather rapid succession, by means of a short, moderately thick glass rod, drawn out to a point and bent at a short right angle. The scales are speared on the glass rod (under the hood; eyes protected by spectacles; hands, with gloves).

The mercury may also be warmed in a porcelain casserole on the water-bath (60–70°), and, without further heating, small pieces of sodium, the size of a half bean, are thrust to the bottom of the vessel with the aid of a glass rod.

12. ALUMINIUM CHLORIDE

A wide tube, diameter $1\frac{1}{2}$ –2 cm., of hard glass drawn out to a narrow tube, is at one end connected by means of a cork with a wide-neck so-called "salt bottle" (Fig. 90). The cork with which this is closed is supplied with a second, smaller hole, bearing a delivery tube of at least 9 mm. diameter, extending to the centre of the receiver. The tube is half filled (half of its cross-section) with aluminium shavings, which have been previously freed from oil by boiling with alcohol and then dried in an air-bath at 120°; an asbestos plug is placed at each end of the layer. A rapid current of hydrochloric acid gas, most conveniently obtained from a Kipp apparatus charged with fused ammonium chloride and concentrated sulphuric acid, is passed

through the apparatus. Care must be taken that the drying flask containing sulphuric acid is not too small, since the acid foams easily. As soon as the air is driven out of the apparatus, — this has been accomplished when the gas evolved is completely absorbed by water (a piece of rubber tubing is attached to the tube, and the gas tested from time to time by immersing the end of the tubing in water in a beaker), — the tube is heated in a combustion furnace throughout its entire length, at first with small flames, which are gradually increased (Fig. 90). When the flames have reached a certain size, white vapours of aluminium chloride, condensing in the receiver, are noticed. The reaction is ended as soon as the aluminium, except for a small, dark-coloured resi-

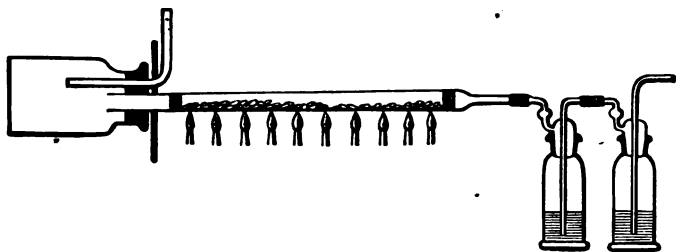


FIG. 90.

due, disappears. For the success of the preparation, the following points are particularly observed : (1) All parts of the apparatus must be perfectly dry. (2) The air must be removed as completely as possible, since, otherwise, an explosion of oxygen and hydrogen may take place. (3) The portion of the tube extending beyond the furnace must be as short as possible, to prevent the aluminium chloride from condensing in it, which results in a stopping up of the apparatus. In order that the cork may not burn, it is protected by an asbestos plate, provided with a circular hole in the centre. (4) The aluminium must not be heated to melting. If this should happen at any particular point, the flames must be immediately lowered. (5) The hydrochloric acid current must be *extremely* rapid. One should not be able to count single bubbles of the gas, but they should follow one another

uninterruptedly. The evolution of a small quantity of a smoky vapour from the outlet-tube will always occur, but the greatest part of the aluminium chloride is condensed even if the hydrochloric acid *rushes through* the wash-bottles. Should the first experiment be unsuccessful, in consequence of a stoppage of the tube, the method for correcting this will readily suggest itself.

Recently it has been shown that it is better to use for the receiver an iron tube 25 cm. long and 4 cm. wide (inner diameter). To one end of this is welded a narrower tube, 2 cm. long; by filing it on the inside the end is given a somewhat conical shape; it is selected of such a diameter that the glass tube can be fastened in it with a few turns of asbestos paper. The end not narrowed is closed by a cork bearing a glass tube as wide as possible leading to the hood. A receiver of this kind is advantageous because the glass tube can be heated strongly to its extreme end, and thus a stopping up of the apparatus may be entirely prevented. If the iron tube should become too hot, a wet towel is placed on it and moistened from time to time.

The aluminium chloride condensing in the receiver is preserved in well-closed bottles, or best, in a desiccator.

13. LEAD PEROXIDE

In a large porcelain dish dissolve, with heat, 50 grammes of lead acetate in 250 c.c. of water, and treat with a solution of bleaching-powder, prepared by shaking 100 grammes of bleaching-powder with $1\frac{1}{2}$ litres of water and filtering, heat not quite to boiling, until the precipitate, bright at first, becomes deep dark brown. A small test-portion is then filtered *hot*, and the filtrate treated with the bleaching-powder solution and heated to boiling; if a dark brown precipitate is formed, more of the bleaching-powder solution is added to the main quantity, and it is heated until a test gives no precipitate with the bleaching-powder solution. The main quantity of the liquid is separated from the heavy precipitate by decantation; the latter is washed several times with water (decantation), and then filtered with suction; the precipi-

tate is washed repeatedly with water. The lead peroxide is not dried, but is preserved in a *closed* vessel in the form of a thick paste.

Value Determination.— In order to determine the value of the paste, a weighed portion is heated with hydrochloric acid, the chlorine evolved is passed into a solution of potassium iodide, and the liberated iodine is titrated with a $\frac{N}{10}$ solution of sodium thiosulphate (refer to a text-book on Volumetric Analysis). The determination, carried out as follows, is sufficiently accurate for preparation work : On an analytical balance weigh off exactly 6.2 grammes of pure, crystallised sodium thiosulphate ; this is dissolved in enough cold water to make the volume of the solution just 250 c.c. In a small flask weigh off 0.5–1 gramme of the peroxide paste ; treat this (with cooling) with a mixture of equal volumes of concentrated hydrochloric acid and water ; the flask is immediately connected with a delivery tube, and this is inserted in an inverted retort, the neck of which has been expanded to a bulb, and which contains a solution of four grammes of potassium iodide in water. When heat is applied to the flask, chlorine is generated, which liberates iodine from the potassium iodide. After the end of the heating, care is taken that the potassium iodide solution is not drawn back into the flask. The contents of the retort are then poured into a beaker and treated with the thiosulphate solution from a burette until the yellow colour of the iodine just disappears. Since a molecule of the peroxide liberates two atoms of iodine, a cubic centimetre of the thiosulphate solution corresponds to $\frac{0.0239}{2} = .012$ gramme pure lead peroxide.

14. CUPROUS CHLORIDE

Heat a solution of 50 grammes of copper sulphate and 24 grammes of salt to 60–70°. Into this conduct a current of sulphur dioxide until the precipitate of cuprous chloride no longer increases. The precipitate is filtered with suction and washed, first with sul-

phurous acid and then with glacial acetic acid until it runs through colourless. The moist preparation is then heated in a shallow porcelain dish or a large watch crystal on the water-bath until the odour of acetic acid cannot be detected. It is preserved in a well-closed flask.

15. DETERMINATION OF THE VALUE OF ZINC DUST

From a weighing-tube pour into a 100 c.c. round flask 0.1 gramme zinc dust (exact weighing) and add a few cubic centimetres of water. The flask is closed by a good three-hole cork. In the middle one is inserted a small dropping funnel; the side holes carry the inlet and outlet tubes (Fig. 91). The stem of the funnel is previously filled with water by opening the cock, immersing the end in water and applying suction. The inlet tube is connected with a Kipp carbon dioxide generator, and the outlet tube with a nitrometer charged with a solution of caustic potash. Carbon dioxide is passed into the apparatus until all the gas escaping from the outlet tube is absorbed by the potash. The current of carbon dioxide is then lessened and from the dropping funnel a mixture of 10 c.c. of concentrated hydrochloric acid and 10 c.c. water containing a



FIG. 91.

few drops of platinic chloride, is allowed to flow in on the zinc dust; the flask is finally heated. From the volume of hydrogen obtained the percentage of zinc in the zinc dust may be calculated. The individual operations of this analysis are conducted as described under "Determination of Nitrogen."

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ABBREVIATIONS

- A.** = Liebig's Annalen der Chemie.
- A. ch.** = Annales de chimie et de physique.
- B.** = Berliner Berichte.
- Bl.** = Bulletin de la société chimique de Paris.
- Ch-Z.** = Chemiker Zeitung.
- J.** = Jahresbericht über die Fortschritte der Chemie.
- J. pr.** = Journal für praktische Chemie.
- P.** = Poggendorff's Annalen.
- R.** = Journal der russischen chemischen Gesellschaft.
- Z.** = Zeitschrift für Chemie.

WEIGHT OF ONE CUBIC CENTIMETRE OF MOIST NITROGEN IN MILLIGRAMMES

t	b	692	694	696	698	700	702	704	706	708	710	712	714	716	718	720 mm
5°	1.108	1.111	1.114	1.117	1.121	1.124	1.127	1.130	1.134	1.137	1.140	1.143	1.146	1.150	1.153	0.6177
6°	0.4439	0.4505	0.4601	0.4817	0.4943	0.5068	0.5192	0.5317	0.5441	0.5564	0.5688	0.5811	0.5933	0.6055	1.145	1.148
7°	1.103	1.106	1.109	1.113	1.116	1.119	1.122	1.125	1.129	1.132	1.135	1.138	1.142	1.145	0.5809	0.5991
8°	0.4251	0.4378	0.4504	0.4630	0.4755	0.4880	0.5005	0.5130	0.5254	0.5378	0.5501	0.5624	0.5747	1.137	1.140	1.143
9°	1.008	1.101	1.105	1.108	1.111	1.114	1.117	1.121	1.124	1.127	1.130	1.133	1.137	1.140	0.5809	0.5991
10°	0.4063	0.4190	0.4316	0.4442	0.4568	0.4693	0.4818	0.4943	0.5067	0.5191	0.5314	0.5437	0.5560	1.132	1.135	1.138
11°	1.093	1.097	1.100	1.103	1.106	1.109	1.112	1.115	1.118	1.121	1.124	1.127	1.130	1.133	0.5497	0.5619
12°	0.3877	0.4003	0.4130	0.4256	0.4382	0.4507	0.4632	0.4757	0.4881	0.5005	0.5128	0.5251	0.5374	1.122	1.125	1.128
13°	1.089	1.092	1.095	1.098	1.101	1.103	1.105	1.108	1.111	1.114	1.117	1.120	1.123	1.126	0.5121	0.5244
14°	0.3691	0.3818	0.3944	0.4070	0.4196	0.4322	0.4447	0.4571	0.4696	0.4820	0.4943	0.5067	0.5190	1.117	1.120	1.123
15°	1.084	1.087	1.090	1.093	1.097	1.100	1.103	1.106	1.109	1.112	1.115	1.118	1.121	1.124	0.4999	0.5121
16°	0.3499	0.3626	0.3752	0.3879	0.4005	0.4130	0.4255	0.4380	0.4505	0.4629	0.4752	0.4876	0.4999	1.114	1.117	1.120
17°	1.079	1.082	1.085	1.088	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	0.4802	0.4925
18°	0.3300	0.3427	0.3554	0.3680	0.3806	0.3932	0.4057	0.4182	0.4307	0.4431	0.4555	0.4679	0.4802	1.109	1.112	1.115
19°	1.074	1.077	1.081	1.084	1.087	1.090	1.093	1.096	1.099	1.103	1.106	1.109	1.112	1.115	0.4612	0.4735
20°	0.3109	0.3236	0.3363	0.3490	0.3616	0.3741	0.3867	0.3992	0.4117	0.4241	0.4365	0.4489	0.4612	1.104	1.107	1.110
21°	1.069	1.073	1.076	1.079	1.082	1.085	1.088	1.091	1.095	1.098	1.101	1.104	1.107	1.110	0.4540	0.4662
22°	0.2912	0.3040	0.3167	0.3293	0.3420	0.3546	0.3671	0.3796	0.3921	0.4046	0.4170	0.4293	0.4417	1.099	1.102	1.105
23°	1.064	1.068	1.071	1.074	1.077	1.080	1.083	1.086	1.089	1.093	1.096	1.099	1.102	1.105	0.4339	0.4461
24°	0.2709	0.2837	0.2964	0.3091	0.3217	0.3343	0.3469	0.3594	0.3719	0.3844	0.3968	0.4092	0.4215	1.094	1.097	1.100
25°	1.059	1.063	1.066	1.069	1.072	1.075	1.078	1.081	1.084	1.088	1.091	1.094	1.097	1.100	0.4138	0.4261
26°	0.2507	0.2635	0.2762	0.2889	0.3016	0.3142	0.3268	0.3393	0.3518	0.3643	0.3767	0.3891	0.4015	1.089	1.092	1.095
27°	1.055	1.058	1.061	1.064	1.067	1.070	1.073	1.076	1.079	1.083	1.086	1.089	1.092	1.095	0.3985	0.4108
28°	0.2305	0.2433	0.2560	0.2687	0.2814	0.2940	0.3066	0.3191	0.3317	0.3442	0.3567	0.3691	0.3815	1.084	1.087	1.090
29°	1.050	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077	1.081	1.084	1.087	1.090	0.3733	0.3856
30°	0.2097	0.2225	0.2353	0.2480	0.2607	0.2734	0.2860	0.2985	0.3111	0.3236	0.3361	0.3485	0.3609	1.077	1.080	1.083

TABLE FOR NITROGEN DETERMINATION

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18°	1.045	1.048	1.051	1.054	1.057	1.060	1.063	1.066	1.069	1.072	1.075	1.078	1.082	1.085	1.088
	01890	02018	02146	02273	02400	02527	02653	02779	02905	03030	03155	03279	03403	03527	03650
19°	1.039	1.042	1.046	1.049	1.052	1.055	1.058	1.061	1.064	1.067	1.070	1.073	1.076	1.079	1.082
	01676	01805	01933	02060	02188	02314	02441	02567	02693	02818	02943	03068	03192	03316	03440
20°	1.034	1.037	1.041	1.043	1.046	1.049	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077
	01457	01585	01713	01841	01969	02096	02222	02349	02475	02600	02725	02850	02975	03099	03223
21°	1.029	1.032	1.035	1.038	1.041	1.044	1.047	1.050	1.053	1.056	1.060	1.063	1.066	1.069	1.072
	01238	01367	01495	01623	01751	01878	02005	02131	02257	02383	02509	02634	02758	02883	03007
22°	1.024	1.027	1.030	1.033	1.036	1.039	1.042	1.045	1.048	1.051	1.054	1.057	1.060	1.063	1.066
	01019	01148	01276	01405	01532	01660	01787	01914	02040	02166	02292	02417	02542	02666	02791
23°	1.018	1.021	1.024	1.027	1.030	1.034	1.037	1.040	1.043	1.046	1.049	1.052	1.055	1.058	1.061
	00788	00917	01046	01174	01302	01430	01557	01684	01811	01937	02063	02189	02314	02439	02563
24°	1.013	1.016	1.019	1.022	1.025	1.028	1.031	1.034	1.037	1.040	1.043	1.046	1.049	1.052	1.055
	00557	00686	00815	00944	01072	01200	01328	01455	01582	01708	01834	01960	02085	02210	02335
25°	1.008	1.011	1.014	1.017	1.020	1.023	1.026	1.029	1.032	1.035	1.038	1.041	1.044	1.047	1.050
	00326	00456	00585	00714	00843	00971	01099	01226	01353	01480	01606	01732	01858	01983	02108
26°	1.002	1.005	1.008	1.011	1.014	1.017	1.020	1.023	1.026	1.029	1.032	1.035	1.038	1.041	1.044
	00083	00213	00342	00472	00600	00729	00857	00985	01112	01239	01366	01492	01618	01743	01868
27°	0.996	0.999	1.002	1.005	1.008	1.011	1.014	1.017	1.020	1.023	1.026	1.029	1.032	1.035	1.038
	99840	99970	00100	00230	00359	00488	00616	00744	00872	00999	01126	01252	01378	01504	01630
28°	0.991	0.994	0.997	1.000	1.003	1.006	1.009	1.012	1.015	1.018	1.020	1.023	1.026	1.029	1.032
	99590	99721	99851	99981	00110	00220	00368	00497	00625	00752	00879	01006	01133	01259	01384
29°	0.985	0.988	0.991	0.994	0.997	1.000	1.003	1.006	1.009	1.012	1.015	1.018	1.021	1.024	1.027
	99341	99472	99603	99733	99863	99994	00121	00250	00378	00506	00634	00761	00887	01014	01140
30°	0.979	0.982	0.985	0.988	0.991	0.994	0.997	1.000	1.003	1.006	1.009	1.012	1.015	1.018	1.021
	99079	99211	99341	99472	99602	99732	99861	99990	00119	00247	00375	00502	00629	00756	00882
t	b 692	694	696	698	700	702	704	706	708	710	712	714	716	718	720 mm

WEIGHT OF ONE CUBIC CENTIMETRE OF MOIST NITROGEN IN MILLIGRAMMES

<i>t</i>	<i>b</i>	732	734	736	738	739	733	734	736	738	740	742	744	746	748	750 mm
5°	1.156	1.159	1.163	1.166	1.169	1.172	1.176	1.179	1.182	1.185	1.188	1.192	1.195	1.198	1.201	07966
6°	06299	06420	06541	06662	06782	06902	07021	07141	07259	07378	07496	07614	07732	07849	07966	07966
7°	1.151	1.154	1.158	1.161	1.164	1.167	1.170	1.174	1.177	1.180	1.183	1.187	1.190	1.193	1.196	07781
8°	06113	06234	06355	06476	06596	06716	06835	06955	07074	07192	07311	07439	07566	07694	07821	07821
9°	1.146	1.149	1.153	1.156	1.159	1.162	1.166	1.169	1.172	1.175	1.178	1.182	1.185	1.188	1.191	07596
10°	05926	06048	06169	06289	06410	06530	06650	06769	06888	07007	07125	07243	07361	07479	07596	07596
11°	1.141	1.145	1.148	1.151	1.154	1.157	1.161	1.164	1.167	1.170	1.173	1.177	1.180	1.183	1.186	07411
12°	05741	05862	05983	06104	06225	06345	06465	06584	06703	06822	06941	07059	07177	07294	07411	07411
13°	1.137	1.140	1.143	1.146	1.149	1.152	1.156	1.159	1.162	1.165	1.168	1.172	1.175	1.178	1.181	07228
14°	05556	05678	05796	05920	06041	06161	06281	06400	06520	06638	06757	06875	06993	07111	07228	07228
15°	1.132	1.135	1.138	1.141	1.144	1.147	1.151	1.154	1.157	1.160	1.163	1.166	1.170	1.173	1.176	07039
16°	05366	05488	05609	05730	05851	05971	06091	06210	06330	06449	06567	06686	06804	06922	07039	07039
17°	1.126	1.130	1.133	1.135	1.139	1.142	1.145	1.149	1.152	1.155	1.158	1.161	1.164	1.168	1.171	06844
18°	05169	05291	05412	05534	05654	05775	05895	06015	06134	06253	06372	06490	06609	06726	06844	06844
19°	1.122	1.125	1.128	1.131	1.134	1.137	1.140	1.144	1.147	1.150	1.153	1.156	1.159	1.163	1.166	06656
20°	04980	05101	05223	05344	05465	05586	05706	05826	05945	06064	06183	06302	06420	06538	06656	06656
21°	1.117	1.120	1.123	1.126	1.129	1.132	1.135	1.139	1.142	1.145	1.148	1.151	1.154	1.157	1.161	06463
22°	04785	04907	05029	05150	05271	05392	05512	05632	05751	05871	05990	06108	06227	06345	06463	06463
23°	1.111	1.114	1.118	1.121	1.124	1.127	1.130	1.133	1.136	1.140	1.143	1.146	1.149	1.152	1.155	06264
24°	04584	04706	04828	04949	05070	05191	05312	05432	05552	05671	05790	05909	06028	06146	06264	06264
25°	1.106	1.109	1.113	1.116	1.119	1.122	1.125	1.128	1.131	1.134	1.137	1.141	1.144	1.147	1.150	06065
26°	04384	04506	04628	04750	04871	04992	05113	05233	05353	05472	05592	05711	05829	05947	06065	06065
27°	1.101	1.104	1.107	1.111	1.114	1.117	1.120	1.123	1.126	1.129	1.132	1.135	1.138	1.142	1.145	05867
28°	04184	04306	04428	04550	04672	04793	04913	05034	05154	05274	05393	05512	05631	05749	05867	05867
29°	1.096	1.099	1.102	1.105	1.108	1.111	1.115	1.118	1.121	1.124	1.127	1.130	1.133	1.136	1.139	05604
30°	03979	04101	04224	04345	04467	04588	04709	04830	04950	05070	05189	05308	05427	05546	05664	05664

TABLE FOR NITROGEN DETERMINATION

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18°	1.091	1.004	1.007	1.100	1.103	1.106	1.109	1.112	1.116	1.119	1.122	1.125	1.128	1.131	1.134
	0.3774	0.3860	0.4019	0.4141	0.4262	0.4384	0.4505	0.4625	0.4746	0.4866	0.4986	0.5105	0.5224	0.5343	0.5461
19°	1.086	1.089	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	1.122	1.126	1.129
	0.3563	0.3686	0.3808	0.3931	0.4053	0.4174	0.4295	0.4416	0.4537	0.4657	0.4777	0.4896	0.5015	0.5134	0.5253
20°	1.080	1.083	1.086	1.089	1.092	1.095	1.099	1.102	1.105	1.108	1.111	1.114	1.117	1.120	1.123
	0.3346	0.3459	0.3592	0.3715	0.3837	0.3958	0.4080	0.4201	0.4321	0.4442	0.4562	0.4682	0.4801	0.4920	0.5039
21°	1.075	1.078	1.081	1.084	1.087	1.090	1.093	1.096	1.099	1.102	1.105	1.108	1.111	1.115	1.118
	0.3130	0.3254	0.3377	0.3499	0.3621	0.3743	0.3865	0.3986	0.4107	0.4228	0.4348	0.4468	0.4587	0.4707	0.4825
22°	1.069	1.073	1.076	1.079	1.082	1.085	1.088	1.091	1.094	1.097	1.100	1.103	1.106	1.109	1.112
	0.2914	0.3038	0.3161	0.3284	0.3406	0.3528	0.3650	0.3772	0.3893	0.4013	0.4134	0.4254	0.4374	0.4493	0.4612
23°	1.064	1.067	1.070	1.073	1.076	1.079	1.082	1.085	1.088	1.091	1.094	1.097	1.100	1.103	1.106
	0.2687	0.2811	0.2934	0.3057	0.3180	0.3302	0.3424	0.3546	0.3667	0.3788	0.3909	0.4029	0.4149	0.4268	0.4388
24°	1.058	1.061	1.064	1.067	1.070	1.073	1.076	1.080	1.083	1.086	1.089	1.092	1.095	1.098	1.101
	0.2459	0.2583	0.2707	0.2830	0.2953	0.3076	0.3198	0.3320	0.3441	0.3562	0.3683	0.3804	0.3924	0.4044	0.4163
25°	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077	1.080	1.083	1.086	1.089	1.092	1.095
	0.2233	0.2357	0.2481	0.2604	0.2727	0.2850	0.2972	0.3094	0.3216	0.3338	0.3459	0.3579	0.3700	0.3820	0.3940
26°	1.047	1.050	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077	1.080	1.083	1.086	1.089
	0.1993	0.2118	0.2242	0.2366	0.2489	0.2612	0.2735	0.2857	0.2979	0.3101	0.3222	0.3343	0.3464	0.3584	0.3704
27°	1.041	1.044	1.047	1.050	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077	1.080	1.083
	0.1755	0.1879	0.2004	0.2128	0.2251	0.2375	0.2498	0.2620	0.2742	0.2864	0.2986	0.3107	0.3228	0.3349	0.3469
28°	1.035	1.038	1.041	1.044	1.047	1.050	1.053	1.056	1.059	1.062	1.065	1.068	1.071	1.074	1.077
	0.1510	0.1635	0.1759	0.1884	0.2008	0.2131	0.2254	0.2377	0.2500	0.2622	0.2744	0.2865	0.2986	0.3107	0.3228
29°	1.030	1.033	1.036	1.039	1.042	1.044	1.047	1.050	1.053	1.056	1.059	1.062	1.065	1.068	1.071
	0.1265	0.1391	0.1516	0.1640	0.1764	0.1888	0.2012	0.2135	0.2258	0.2380	0.2502	0.2624	0.2745	0.2867	0.2987
30°	1.024	1.027	1.029	1.032	1.035	1.038	1.041	1.044	1.047	1.050	1.053	1.056	1.059	1.062	1.065
	0.1008	0.1134	0.1259	0.1384	0.1509	0.1633	0.1757	0.1880	0.2003	0.2126	0.2248	0.2370	0.2492	0.2614	0.2735
ℓ	b 722	724	726	728	730	732	734	736	738	740	742	744	746	748	750 mm

WEIGHT OF ONE CUBIC CENTIMETRE OF MOIST NITROGEN IN MILLIGRAMMES

<i>t</i>	<i>b</i>	752	754	756	758	760	762	764	768	770	772	774	776	778	780 mm
5°	1.205 0.8083	1.208 0.8199	1.211 0.8315	1.214 0.8431	1.218 0.8546	1.221 0.8661	1.224 0.8776	1.227 0.8891	1.230 0.9005	1.234 0.9119	1.237 0.9233	1.240 0.9346	1.243 0.9459	1.247 0.9572	1.250 0.9684
6°	1.199 0.7898	1.203 0.8014	1.206 0.8130	1.209 0.8246	1.212 0.8361	1.216 0.8477	1.219 0.8592	1.222 0.8706	1.225 0.8820	1.228 0.8934	1.232 0.9048	1.235 0.9162	1.238 0.9275	1.241 0.9387	1.245 0.9500
7°	1.194 0.7712	1.198 0.7829	1.201 0.7945	1.204 0.8061	1.207 0.8177	1.210 0.8292	1.214 0.8407	1.217 0.8522	1.220 0.8636	1.223 0.8750	1.226 0.8864	1.230 0.8977	1.233 0.9090	1.236 0.9203	1.239 0.9316
8°	1.180 0.7528	1.183 0.7645	1.186 0.7761	1.189 0.7877	1.192 0.7993	1.195 0.8108	1.198 0.8223	1.201 0.8338	1.204 0.8452	1.207 0.8566	1.210 0.8680	1.213 0.8794	1.216 0.8907	1.219 0.9020	1.222 0.9133
9°	1.174 0.7345	1.177 0.7462	1.180 0.7578	1.183 0.7694	1.186 0.7810	1.189 0.7925	1.192 0.8040	1.195 0.8155	1.198 0.8270	1.201 0.8384	1.204 0.8498	1.207 0.8612	1.210 0.8725	1.213 0.8838	1.216 0.8951
10°	1.169 0.7156	1.172 0.7273	1.175 0.7389	1.178 0.7505	1.181 0.7621	1.184 0.7737	1.187 0.7852	1.190 0.7967	1.193 0.8081	1.196 0.8196	1.199 0.8310	1.202 0.8423	1.205 0.8537	1.208 0.8650	1.211 0.8763
11°	1.164 0.6961	1.167 0.7078	1.170 0.7195	1.173 0.7311	1.176 0.7427	1.179 0.7542	1.182 0.7658	1.185 0.7773	1.188 0.7887	1.191 0.8002	1.194 0.8116	1.197 0.8230	1.200 0.8343	1.203 0.8456	1.206 0.8569
12°	1.158 0.6773	1.161 0.6890	1.164 0.7007	1.167 0.7123	1.170 0.7239	1.173 0.7355	1.176 0.7470	1.179 0.7585	1.182 0.7700	1.185 0.7814	1.188 0.7929	1.191 0.8043	1.194 0.8156	1.197 0.8269	1.200 0.8383
13°	1.153 0.6580	1.156 0.6697	1.159 0.6814	1.162 0.6930	1.165 0.7046	1.168 0.7162	1.171 0.7278	1.174 0.7393	1.177 0.7508	1.180 0.7622	1.183 0.7737	1.186 0.7851	1.189 0.7964	1.192 0.8078	1.195 0.8191
14°	1.148 0.6381	1.151 0.6498	1.154 0.6615	1.157 0.6732	1.160 0.6848	1.163 0.6964	1.166 0.7080	1.169 0.7195	1.172 0.7310	1.175 0.7425	1.178 0.7539	1.181 0.7653	1.184 0.7767	1.187 0.7881	1.190 0.7994
15°	1.143 0.6183	1.146 0.6300	1.149 0.6417	1.152 0.6534	1.155 0.6650	1.158 0.6767	1.161 0.6882	1.164 0.6998	1.167 0.7113	1.170 0.7228	1.173 0.7342	1.176 0.7457	1.179 0.7571	1.182 0.7684	1.185 0.7798
16°	1.138 0.5985	1.141 0.6103	1.144 0.6220	1.147 0.6336	1.150 0.6453	1.153 0.6569	1.156 0.6685	1.159 0.6801	1.162 0.6916	1.165 0.7031	1.168 0.7146	1.171 0.7260	1.174 0.7374	1.177 0.7488	1.180 0.7601
17°	1.133 0.5782	1.136 0.5900	1.139 0.6017	1.142 0.6134	1.145 0.6251	1.148 0.6367	1.151 0.6483	1.154 0.6599	1.157 0.6714	1.160 0.6829	1.163 0.6944	1.166 0.7058	1.169 0.7173	1.172 0.7287	1.175 0.7400

18°	1.137	1.140	1.143	1.146	1.149	1.153	1.156	1.159	1.162	1.165	1.168	1.171	1.174	1.177	1.180
	05579	05697	05814	05931	06048	06165	06281	06397	06512	06627	06742	06857	06971	07085	07199
19°	1.132	1.135	1.138	1.141	1.144	1.147	1.150	1.153	1.156	1.159	1.162	1.166	1.169	1.172	1.175
	05371	05489	05607	05724	05841	05957	06074	06190	06305	06421	06536	06651	06765	06879	06993
20°	1.126	1.129	1.132	1.135	1.138	1.141	1.144	1.148	1.151	1.154	1.157	1.160	1.163	1.166	1.169
	05157	05275	05393	05510	05627	05744	05861	05977	06093	06208	06324	06439	06553	06668	06782
21°	1.121	1.124	1.127	1.130	1.133	1.136	1.139	1.142	1.145	1.148	1.151	1.154	1.157	1.160	1.163
	04944	05062	05180	05298	05415	05532	05649	05765	05881	05997	06112	06227	06342	06457	06571
22°	1.115	1.118	1.121	1.124	1.127	1.130	1.133	1.136	1.139	1.143	1.146	1.149	1.152	1.155	1.158
	04731	04849	04967	05085	05203	05320	05437	05553	05669	05785	05901	06016	06131	06246	06360
23°	1.109	1.112	1.115	1.119	1.122	1.125	1.128	1.131	1.134	1.137	1.140	1.143	1.146	1.149	1.152
	04507	04625	04744	04862	04979	05097	05214	05330	05447	05563	05679	05794	05909	06024	06139
24°	1.104	1.107	1.110	1.113	1.116	1.119	1.122	1.125	1.128	1.131	1.134	1.137	1.140	1.143	1.146
	04282	04401	04520	04638	04756	04873	04991	05108	05224	05341	05457	05572	05688	05803	05917
25°	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	1.122	1.125	1.128	1.131	1.134	1.137	1.140
	04059	04178	04297	04415	04533	04651	04769	04886	05002	05119	05235	05351	05467	05582	05697
26°	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	1.122	1.125	1.128	1.131	1.134
	03823	03943	04062	04180	04299	04417	04534	04652	04769	04886	05002	05118	05234	05349	05465
27°	1.086	1.089	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	1.122	1.125	1.128
	03586	03708	03828	03946	04065	04183	04301	04419	04536	04653	04770	04886	05002	05118	05233
28°	1.080	1.083	1.086	1.089	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116	1.119	1.122
	03348	03468	03587	03706	03825	03944	04062	04180	04297	04415	04531	04648	04764	04880	04996
29°	1.074	1.077	1.080	1.083	1.086	1.089	1.092	1.095	1.098	1.101	1.104	1.107	1.110	1.113	1.116
	03108	03228	03348	03467	03586	03705	03823	03941	04059	04177	04294	04411	04527	04644	04759
30°	1.068	1.071	1.074	1.077	1.080	1.083	1.086	1.089	1.092	1.095	1.098	1.101	1.104	1.107	1.110
	02855	02976	03096	03216	03335	03454	03573	03691	03809	03927	04045	04162	04278	04395	04511
t	b 752	754	756	758	760	762	764	766	768	770	772	774	776	778	780mm

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