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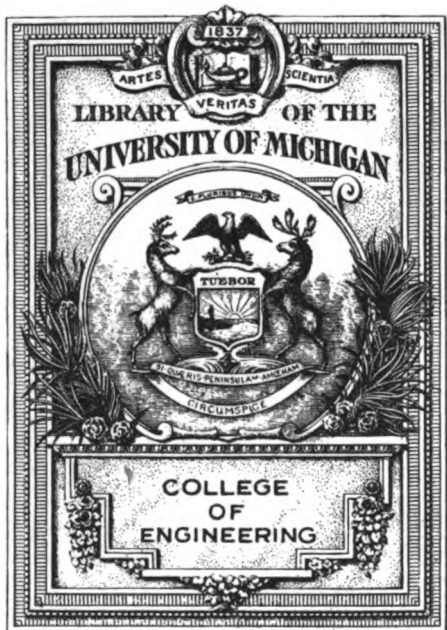
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A method for the identification of pure organic ...

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A METHOD

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VOL. II

CONTAINING CLASSIFIED DESCRIPTIONS
OF ABOUT 4000 OF THE MORE IMPORTANT COMPOUNDS
OF CARBON WITH THE ELEMENTS NITROGEN,
HYDROGEN, AND OXYGEN

BY

SAMUEL PARSONS MULLIKEN, PH. D.

*Associate Professor of Organic Chemical Research at the Massachusetts Institute of Technology,
Boston, Mass.*



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PREFACE.

In sending out this new volume it may not be inappropriate for the author to add a word about certain details of its plan which are believed to represent forward steps in the evolution of the analytical system of which it forms a unit. The improvements to be noted are, however, not of a revolutionary character, time having brought abundant confirmation of the soundness of the fundamental conception which underlies the system of ordinal, generic and specific tests initiated in Volume I. The most important are: a reduction in the number of genera in Order II as compared with Order I; an increase in the proportion of carefully verified specific characterizations; and a more convenient and flexible notation and arrangement for both specific characterizations and numbered tests.

The multiplication of genera in an order, within reasonable limits, is highly desirable, provided the corresponding generic tests can be kept simple and not encumbered with too many alternatives and exceptions. Practically, however, species showing anomalous behavior in any single reaction that may be selected as a generic test occur in most natural genera, and some qualifying provisos are unavoidable. The difficulty thus introduced is not keenly felt when the number of interdependent tests in a consecutive series is short; but it rapidly becomes more discouraging as the number of experiments which the analyst is forced to make in succession increases. In Order II, owing to the influence of the nitrogen atom on the chemical behavior of substituent groups, this difficulty is much less easy to surmount than in Order I. It was therefore found expedient to remain content for the present with a subdivision of the colorless species into three great genera characterized respectively by the predominance of the acidic, basic or neutral character. To provide tests that should be both simple and reliable, even for this degree of subdivision, was no light task, since it could only be accomplished after the titration of a very large number of representative pure compounds of many types. That this was finally brought to a successful conclusion is mainly due to the skillful and indefatigable coöperation of Dr. Heyward Scudder, with whom the author originally hoped to be associated in the joint authorship of the volume, and to whom too much praise cannot be given for the important contribution actually made. Several hundred compounds were titrated, and some years later many additional points of support for testing the accuracy of the classification became accessible through the publication of Dr. Scudder's work on "The Electrical Conductivity and Ionization Constants of Organic Compounds."

Order II is exceptionally rich in compounds whose identification is of occasional urgent practical importance, containing, as it does, all the alkaloids, a majority of the most powerful drugs of other classes, many of the most interesting components of the animal and vegetable organism, proteolytic products, the high explosives, and a considerable fraction of the intermediates of the dyestuff industry. Much effort was accordingly expended in the systematic and critical study of the properties and diagnostic reactions of the more important individual species. Many errors in the description of constants and other properties which

had been long current in the literature were thus discovered and corrected. In selecting tests for use, all procedures that seemed promising were often tried and compared, and only such as were found to be useful or desirable under the circumstances have been recommended. In cases where the records of earlier investigators indicated that descriptions had been prepared with great care and with very pure materials, preference has generally been given to them rather than to original data when based on unanalyzed substances. Whenever the importance of a compound concerning which earlier descriptions conflicted justified the outlay, time was freely spent in critical experimental as well as bibliographical study of the data in the attempt to enable a decision. All parts of specific characterizations which appear in the analytical tables in italics have been experimentally verified in the author's laboratory. Partial experimental verification has also been made of many data that are not italicized.

As in Volume I a preliminary selection of candidates for description was made by a page to page examination of the four volumes of the third edition of Beilstein's Handbuch, and their four supplements. This was followed later by an almost equally thorough search with the same object of the fourth and fifth volumes of Abderhalden's Biochemisches Handlexikon and a few other recent encyclopedic monographs and lists devoted to the description or enumeration of the compounds belonging to important special groups; but no exhaustive canvass of the periodical literature of the last few years was attempted.

Friendly assistance from so many sources has been received in connection with the preparation of this volume that it is scarcely possible to make suitable individual acknowledgments to all who have contributed to the result. Next to Dr. Scudder, the importance of whose aid in establishing the generic classification on a substantial basis has already been emphasized, special thanks are due to Professor Alice F. Blood and Miss Florence Sargent, now of Simmons College, each of whom at different times spent a year as assistants in the solution of many of the most important experimental problems. Among other helpful contributors in the development of the specific characterizations and "numbered tests" during a series of years by theses and minor investigations while students at the Massachusetts Institute of Technology, mention should be made of Messrs. C. L. W. Pettee, Geo. W. Knight, Edw. G. Thatcher, Alden Merrill, J. U. G. Calnan, Walter Burns, Max Cline, Chas. Field, Miss M. J. Ruggles, Messrs. R. E. Drake, Chas. L. Gabriel, W. S. Hughes, T. L. Davis and R. W. Mitchell. Grateful acknowledgment should also be rendered to the American Academy of Arts and Sciences for a grant of \$500 from the C. M. Warren Fund, which was applied toward defraying the expenses of a part of the experimental investigation of the basic species in Genus II of Suborder I.

Experience gained through earlier prefaces has taught the author the uncertainty of predictions concerning future publications. It is, nevertheless, his expectation that this work will be completed in its first edition within a reasonable time by the publication of a fourth volume, which will include procedures for the identification of the more important organic compounds containing combinations of the elements that have not been treated.

MASSACHUSETTS INSTITUTE OF TECHNOLOGY,
April, 1918.

NOTE ON REAGENTS FOR REDUCTION TESTS.

(To face page 17 of "The Identification of Commercial Dyestuffs.")

While the original descriptions of all color discharges obtained in Tests 8 and 11 that are tabulated in Volume III are based on the use of a solution of Rongalite C, practically identical results under the same experimental conditions are secured by the employment of a reducing mixture prepared by the following empirical procedure:

Place in a 300-cc. flask 15 grams of dry sodium bisulphite and 8 grams of zinc dust. Add 75 cc. of distilled water. Shake to dissolve the bisulphite and wet the zinc. Then add from a burette 7.5 cc. of commercial "formalin" (40 per cent formic aldehyde solution). After mixing thoroughly, heat quickly to boiling on a wire gauze over a Bunsen flame, and boil moderately for just 5 minutes. Dilute at once with 45 cc. of cold distilled water, and then cool and filter. Or, double all the quantities mentioned may be taken, and the mixture boiled for the same period of 5 minutes.

The commercial dry sodium bisulphite of the quality placed on the American market by reputable manufacturers as the "U.S.P. viii" grade, if taken from full recently purchased bottles, is generally suitable for the preparation of this reagent, even when not absolutely fresh. The finished reagent should not be preserved for more than a few days, and should not smell of formic aldehyde when warmed. It should be understood that the times required for discharges in reduction tests and the color-returns in oxidation tests as stated in the tables are only approximations, and will be found to vary slightly in independent experiments with the same dyestuff whichever reagent is employed.

In Tests 6, 11, and 21 it is permissible to substitute any good commercial sodium hydrosulphite for Blankit T.

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TABLE OF ABBREVIATIONS.

VERBAL ABBREVIATIONS.

<i>a.</i>	= <i>above</i> . Before a melting-point or boiling-point indicates that the change occurs above the temperature given.	<i>d.</i>	= <i>decomposes</i> . Standing <i>after</i> a number indicates the temperature at which a substance melts or boils with decomposition. Standing <i>before</i> a number indicates the temperature at which decomposition occurs, without necessarily implying that the substance either melts or boils.
<i>A.</i>	= <i>standing after</i> the symbol for a metal represents the acid radical derived from the acid in whose description it occurs by the removal of one hydrogen atom; <i>e.g.</i> , PbA, in a description of benzoic acid would represent lead benzoate.	<i>d. a.</i>	Standing before the name of a compound signifies <i>dextro</i> .
<i>abs.</i>	= <i>absolute</i> .	<i>d. w. m.</i>	= <i>decomposes above</i> . Used in the same way as <i>d.</i>
<i>abt.</i>	= <i>about</i> . Indicates that the value following is only approximate.	<i>deliq.</i>	= <i>decomposes without melting</i> .
<i>ac.</i>	= <i>acid</i> .	<i>dif. fr.</i>	= <i>deliquesces and deliquescent</i> .
<i>Ac.</i>	= <i>acetic acid and acetate</i> .		= <i>differs from</i> . Signifies that one specified compound differs from another in some property or test that has just been mentioned.
<i>alc.</i>	= <i>alcohol</i> (in general 95 per cent ethyl alcohol unless otherwise stated); also, <i>alcoholic</i> .	<i>dil.</i>	= <i>dilute</i> .
<i>ald.</i>	= <i>aldehyde</i> .	<i>dist.</i>	= <i>distils</i> ; also, <i>may be distilled</i> in cases where no definite boiling-point is recorded.
<i>alk.</i>	= <i>alkali and alkaline</i> .	<i>d. s.</i>	= <i>difficultly soluble</i> . Usually indefinite; but, when printed in italics, means soluble in 50 to 150 parts of solvent.
<i>alm.</i>	= <i>almost</i> .	<i>dec.</i>	= <i>decompose</i> ("d" is used instead in certain cases. Cf. above.)
<i>anhyd.</i>	= <i>anhydride and anhydrous</i> .	<i>①</i>	= <i>derivative</i> (used to introduce detailed directions for preparing a derivative in specific characterizations.)
<i>approx.</i>	= <i>approximately</i> .	<i>e. s.</i>	= <i>easily soluble</i> .
<i>aq.</i>	= <i>water and aqueous</i> .	<i>Et.</i>	= <i>ethyl</i> , C ₂ H ₅ .
<i>as.</i>	= <i>asymmetrical</i> .	<i>Eth.</i>	= <i>ether</i> . When referring to a solvent, <i>ethyl oxide</i> .
<i>ammon.</i>	= <i>ammonia, ammonium, or ammoniacal</i> .	<i>exam.</i>	= <i>examine or examination</i> .
<i>Bkn.</i>	= "broken." (Cf. "Color Terminology," Vol. I.)	<i>evap. evapn.</i>	= <i>evaporate, -tion</i> .
<i>b. p.</i>	= <i>boiling-point</i> .	<i>expt.</i>	= <i>experiment</i> .
<i>bril.</i>	= <i>brilliant</i> .	<i>effev.</i>	= <i>effervescence</i> .
<i>bz.</i>	= <i>benzene</i> .	<i>fr.</i>	= <i>from</i> .
<i>B.</i>	= <i>Standing before the symbol of an acid, represents a molecule of the "basic" salt-forming compound in whose specific characterization it occurs. Thus B.HCl in a description of aniline represents aniline hydrochloride, C₆H₅NH₂.HCl.</i>	<i>fil.</i>	= <i>filtrate</i> .
<i>c.</i>	= <i>cold, and (with boiling or melting points) corrected</i> .	<i>floc.</i>	= <i>flocculent</i> .
<i>chl.</i>	= <i>chloroform</i> .	<i>fluor.</i>	= <i>fluoresces, fluorescence or fluorescent</i> .
<i>comp.</i>	= <i>compound or composition</i> .	<i>gran.</i>	= <i>granular</i> .
<i>compl.</i>	= <i>complete</i> .	<i>g. (s)</i>	= <i>gram. (s)</i> .
<i>cryst.</i>	= <i>crystals, crystalline, and crystallizes; crystallization</i> .	<i>h.</i>	= <i>hot</i> .
<i>D. R. P.</i>	= <i>German Patent</i> .	<i>h. c.</i>	= <i>hydrocarbon</i> .
(D)	= <i>dark</i> (following the name of a broken color. Cf. color standard of Vol. III.)	<i>i.</i>	= <i>insoluble; optically inactive</i> (before name of a compound.)
		<i>i. v.</i>	= <i>in vacuo</i> .

TABLE OF ABBREVIATIONS

<i>k.</i>	= ionization constant.	<i>s. cap.</i>	= sealed capillary, fused in. (Cf. foot-note, Vol. I, page 219.)
(<i>L</i>). <i>l-</i> <i>lgr.</i> <i>lig.</i> <i>lft.</i> (<i>M</i>). <i>m.</i> <i>Me.</i> <i>m. p.</i> <i>mic.</i> <i>min.</i> <i>mol. (s).</i> <i>mg.</i> <i>mm.</i> <i>misc.</i> [<i>N</i>]p <i>N. Eq.</i> <i>No.</i> <i>or.</i> <i>o-</i> <i>oxid.</i> ② <i>p-</i> <i>Ph.</i> <i>ppt. (d) (g) (n).</i> <i>pt.</i> <i>pulv.</i> <i>prob.</i> <i>r. h.</i> <i>s.</i> <i>Sap. Eq.</i>	<ul style="list-style-type: none"> = light. (Modifying name of a broken color. Cf. Color standard of Vol. III.) = laevo. = ligroin. = liquid. = leaflet. = medium (modifying name of a broken color. Cf. Color Standard of Vol. III.) = melt. = methyl, CH_3. = melting-point. = microscopic. = minute. = molecule (s). = milligrams. = millimeters. = miscible. = index of refraction, with monochromatic sodium light. = "neutralization equivalent" as defined on page 77, Vol. I. = number. Refers to serial number of a species in this work. = orange. = ortho. = oxide, oxidizing, oxidation. = preliminary test. (Use described on page 5.) = para. = phenyl, C_6H_5. = precipitate, -ed, -ing, -ion. = part and parts. = pulverulent. = probably. = rapidly heated. (Cf. page 220, Vol. I.) = soluble. Usually indefinite; but, when printed in italics, means soluble in 20 to 50 parts of solvent. = symmetrical, when preceding the name of a compound. = saponification equivalent. (Defined on page 113, Vol. I.) 	<ul style="list-style-type: none"> <i>sbl.</i>	= sublimes and sublimate.
<i>sbl. w. m.</i>	= sublimes without melting. (Often followed by specification of the temperature.)		
<i>sec.</i>	= second.		
<i>s. d.</i>	= with slight decomposition. (Signifies that the compound melts or boils with slight decomposition at the temperature given.)		
<i>sf.</i>	= softens. (Often accompanied by temperature of softening.)		
<i>sol.</i>	= solution.		
<i>sl.</i>	= slightly and slight.		
<i>sp. gr.</i>	= specific gravity.		
<i>sat.</i>	= saturated.		
<i>sap. (d) (g) (n)</i>	= saponify, saponifies, -ing, -ion.		
<i>T.</i>	= Test. Followed by a numeral has reference to one of the "numbered tests." (Cf. page 12 of this volume.)		
<i>tbl.</i>	= tables and tabular (crystals).		
(<i>th. i.</i>)	= thermometer immersed in the vapor. Without stem exposure.		
μ (—)	= molar conductivity at a specified concentration.		
<i>u. c.</i>	= uncorrected. May be assumed to mean that the temperature in question has not been corrected for stem exposure, but that any necessary corrections for zero point and calibration of the thermometer have been applied.		
<i>v.</i>	= very, and vicinal or adjacent.		
<i>volat.</i>	= volatile.		
<i>vol. w. st.</i>	= volatile with steam.		
<i>vol. (s)</i>	= volume (s).		
<i>wt. (s)</i>	= weight (s).		
<i>viol.</i>	= violet.		
<i>yel.</i>	= yellow.		
<i>x. s.</i>	= an excess.		
<i>sbl.</i>	= sublimes and sublimate.		
<i>sbl. w. m.</i>	= sublimes without melting. (Often followed by specification of the temperature.)		
<i>sec.</i>	= second.		
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<i>wt. (s)</i>	= weight (s).		
<i>viol.</i>	= violet.		
<i>yel.</i>	= yellow.		
<i>x. s.</i>	= an excess.		

SPECIAL SIGNS.

- † placed before the name of a compound in the tables of this volume indicates that the generic position of the species has been determined by actual titration.
 C (the sign of cyclic bonding) indicates that the terminal atoms in a linear structural formula would be represented in an expanded graphic formula as connected by a valence bond in a closed ring. Thus, the cyclic structure of benzene may be expressed by the formula $\text{C}=\text{CH}:\text{CH}=\text{CH}=\text{CH}:\text{CH}=\text{C}$, which is the linear equivalent of the familiar hexagon formula of Kekulé.

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[The parts of titles here printed in bold type are employed in the text as abbreviations.]

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A METHOD FOR THE IDENTIFICATION OF PURE ORGANIC COMPOUNDS

CHAPTER I.

CLASSIFICATION AND GENERAL ANALYTICAL PROCEDURE IN ORDER II.

The general principles of classification for compounds, and the definitions already adopted in Volume I for the terms *order*, *genus*, *division*, *section*, *species*, and for *ordinal*, *generic*, *specific* and *coördinating tests*, will be retained in this volume.

Order II includes all compounds containing the elements: (a) carbon and nitrogen; (b) carbon, nitrogen and hydrogen; (c) carbon, nitrogen and oxygen; (d) carbon, nitrogen, hydrogen and oxygen. Certain practical considerations, *viz.*, the absence of any direct test for oxygen, the rather inconvenient character of the copper oxide combustion test for hydrogen, and the very small number of the species containing carbon and nitrogen only, make it inadvisable for the present to resolve this large composite order into the four natural and simpler ones corresponding to the letters (a), (b), (c), (d).

In selecting species for description in the tables, the most important intentional omissions among compounds of general interest occur in the case of those whose instability renders their isolation in a condition of assured purity almost impossible, and of those for which neither melting- nor boiling-point has been determined. Ammonium salts, with a few exceptions, are not described as such, but may be identified through their acids, which must first be isolated. Esters are described sparingly, and, aside from the methyl and ethyl esters of the more important acids, are likely to be omitted unless they have been found in natural products, are of commercial interest, or appear serviceable as derivatives for the identification of their acids. Other esters may, of course, be identified through their saponification products.

Order II is divided into two Suborders.

Suborder 1 comprises all species which have colors less saturated than a Tint 3 of the color standard accompanying Volume I. Its species are arranged in three genera, each of which has a "Division A" for its solid, and a "Division B" for its liquid, species. Genus I comprises the compounds of distinctly acidic, Genus II of distinctly basic, and Genus III those of neutral, character. The

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genus in Suborder I is experimentally determined by applying the generic titration tests (Generic Tests I-III) described in Chapter II. The determination of the genus usually requires less time, less material, and fewer successive tests, than in Order I.

Suborder 2 comprises the solid species of the order which are more saturated in color than a Tint 3 of the color standard. It has no Division B for liquid species, the few colored liquids — mostly yellowish — which might have been described in such a division being described with the liquid species of Suborder I.

Species are arranged within their "divisions" in the sequence of the increasing numerical magnitudes of their melting- or boiling-points, and when a number of species in the same division have nearly the same melting- or boiling-point, the relative positions of the individuals are sometimes fixed still more definitely by subordinate coördination tests, such as determination of neutralization equivalent or color. In Suborder II (Colored Compounds), the "coördinating sequence" is the chromatic sequence of the color standard, proceeding from violet-red through yellow, blue, etc., to red-violet. The descriptions of red species of a given melting-point, for example, are made to precede those of yellow, while those of yellow species precede those of blue.

In parts of the tables where the number of species melting or boiling at about the same temperature is small, simple preliminary tests, made conspicuous by the typographical sign \oplus , are often inserted at the beginnings of specific descriptions, and no secondary coördination tests are used. In this volume the most frequently used preliminary tests and general procedures of the type designated as *Semispecific Tests* in Volume I have been massed together in Chapter III as *Numbered Tests* for greater convenience in cross-referencing and indexing.

The final *Specific Tests* relied upon for distinguishing species from others situated near them in the same section of a division are not grouped together after the tables of each genus as was often done in Order I, but are printed in the tables as integral parts of the specific descriptions. To facilitate cross-referencing and indexing, each species described is assigned a *Serial Reference Number* in the tables; and in the Alphabetical Index at the end of the volume the numeral following the name of any compound is this serial number, and not the number of the page on which the description of the compound is to be sought.

DIRECTIONS FOR THE GENERAL PROCEDURE TO BE FOLLOWED IN IDENTIFYING AN UNKNOWN COMPOUND IN ORDER II.

The following numbered paragraphs in bold-faced type provide a complete key to the use of the "Method" in identifying species of Order II. They direct attention in proper succession to each step in the general procedure, and are followed by an illustrative example to make their application clearer.

Before attempting to use this key it is assumed that the reader will have acquainted himself with the principles of classification adopted for the Order to the extent to which they have been outlined in the introductory paragraphs of this chapter.

1. DETERMINATION OF PURITY.

ESTABLISH A PRESUMPTION THAT THE UNKNOWN SUBSTANCE IS REALLY A PURE COMPOUND BEFORE ATTEMPTING TO IDENTIFY IT. IF IT IS NOT HOMOGENEOUS, PURIFY IT. THE CONSTITUENTS OF AN UNKNOWN ORGANIC MIXTURE CAN NOT BE SATISFACTORILY IDENTIFIED PREVIOUS TO THEIR SEPARATION.

The criteria for chemical homogeneity and the problem of dealing with mixtures are discussed under the caption "Purity" on pp. 3-4, Vol. I.

2. DETERMINATION OF PHYSICAL PROPERTIES.

DETERMINE THE COLOR (IN TERMS OF THE COLOR STANDARD* ACCOMPANYING VOL. I), THE ODOR, AND OFTEN TASTE † (WITH EXTREME CAUTION), THE APPROXIMATE SOLUBILITY, IN WATER AT LEAST, AND ANY OTHER SALIENT PHYSICAL CHARACTERISTICS OF THE SUBSTANCE. IF IT IS SOLID, DETERMINE ITS MELTING-POINT; IF A LIQUID, ITS BOILING-POINT AND ITS SPECIFIC GRAVITY. MAKE THESE DETERMINATIONS BY THE METHODS RECOMMENDED UNDER THE CAPTION "PHYSICAL PROPERTIES" ON P. 4 OF VOL. I.

3. DETERMINATION OF ORDER AND SUBORDER.

DETERMINE THE ORDER OF THE COMPOUND BY APPLYING THE ORDINAL TESTS OF VOL. I, CHAP. 2, AFTER NOTING THE DESCRIPTION OF THE IMPROVED ALTERNATIVE TEST FOR NITROGEN GIVEN BELOW. IF IT CONTAINS CARBON AND NITROGEN ALONE, OR THESE ELEMENTS TOGETHER WITH EITHER HYDROGEN OR OXYGEN, OR WITH BOTH THESE LATTER ELEMENTS BUT NO OTHERS, ITS DESCRIPTION SHOULD BE SOUGHT IN THIS VOLUME PROVIDED TEST 2.8 DOES NOT SHOW IT TO BE AN AMMONIUM SALT.

IF THE COMPOUND IS "COLORED" (i.e., HAS A COLOR MORE SATURATED THAN THE THIRD "TINT" OF ANY HUE OF THE COLOR STANDARD), IT SHOULD BE SOUGHT AT ONCE IN THE TABLES OF SUBORDER II, WHICH BEGIN ON P. 237. IF IT IS COLORLESS, PROCEED TO THE DETERMINATION OF ITS GENUS AS DIRECTED IN SECTION 4 OF THIS CHAPTER.

In testing for nitrogen in compounds that are explosive, very volatile, or when only a few milligrams can be spared for the experiment, it will be best to employ the procedure described by Mulliken and Gabriel (Original Communications of Eighth International Congress of Applied Chemistry (1912) 6, 208-211). One centigram or less, according to circumstances, of the substance is mixed with one decigram of pure naphthalene and compressed into pellets weighing about 3 centigrams each. Since the test is often greatly increased in delicacy by using a uniform and intimate mixture, it is best to incorporate the compound with the hydrocarbon by dissolving the former in a couple of drops of some non-nitrogenous solvent like ether or alcohol. Three of these pellets are then decomposed by sodium in the usual manner in the iron ignition tube shown in Fig. 1, Vol. I. Five drops of slightly oxidized and nearly neutral ferrous sulphate solution are added to the concentrated filtered alkaline solution from the fusion, and the

* This Color Standard, consisting of the two cards A and B and a perforated screen, will be mailed to any person owning this work upon receipt of a postal money order for one dollar by the Publishers, John Wiley & Sons, Inc., 432 Fourth Avenue, New York City. It should be replaced when it becomes soiled or faded.

† See Numbered Test 2.29.

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mixture boiled at least two minutes. The precipitate of iron hydroxides is dissolved by a very slight excess of hydrochloric acid, and then allowed to stand for 15 minutes or longer! The solution, which may be clear and without any very noticeable blue or green color, is then filtered through a wet filter not more than 1 inch in diameter, and the filter well washed, first, with hot dilute hydrochloric acid of 1.08 specific gravity, and then with water. After drying, a scanty blue precipitate, or at least a pale bluish stain, will usually be seen on the paper if the substance contained as much as 0.0001 grain of nitrogen. Repeated experiments with naphthalene have shown that atmospheric nitrogen is *not* fixed in sufficient quantity under these conditions — despite the presence of the iron of the tube — to give more than a faint grayish stain to the paper. The presence of nitrogen is thus readily shown in as little as one milligram of morphine, strychnine, sparteine, quinine, ethyl nitrate, nitrated cotton, and nitroglycerine; but the sodium fusion method can not be depended on for the detection of nitrogen in the N₂ group of diazo compounds.

Difficulty occasionally arises in deciding whether a compound shall be considered as colored or colorless when it is accompanied by minute quantities of impurities or decomposition products that defy removal. A somewhat similar difficulty is encountered in Order II in correctly placing species which show the phenomena of phototropy, or are capable of existing in chromoisomeric forms. Thus the white phenylhydrazone of benzaldehyde becomes red on exposure to sunlight, but colorless again after being heated to 100°, or long standing in darkness; various nitrophenols like picric acid crystallize colorless from hot concentrated hydrochloric acid, but yellow from water; while the colors of many nitro compounds as described in chemical literature are unquestionably those of equilibrium mixtures of colored and colorless forms. When there is doubt whether a compound is to be regarded as colored or colorless, the only safe course is to seek for it in both suborders. In cases of known difficulty a mention of such a compound is likely to be found in both places.

4. DETERMINATION OF GENUS.

(FOR SPECIES IN SUBORDER I OF ORDER II.) — FOLLOW THE PROCEDURE GIVEN IN CHAP. II. BEFORE PERFORMING THESE GENERIC TESTS FOR THE FIRST TIME, READ CAREFULLY THE EXPLANATORY REMARKS, AS WELL AS THE DIRECTIONS FOR THE ACTUAL EXPERIMENTS.

5. LOCATION OF A SPECIES AMONG ITS CONGENERS.

TURN TO THE PROPER DIVISION ("A" FOR SOLIDS, AND "B" FOR LIQUIDS), IN THE TABLES OF THE GENUS, OR — IN CASE OF THE COLORED COMPOUNDS WHICH ARE NOT SUBDIVIDED INTO GENERA — TURN SIMPLY TO SUBORDER II. THEN LOCATE THE DESCRIPTION OF THE SPECIES APPROXIMATELY BY APPLYING IN SUCCESSION WHATEVER COÖRDINATING OR SECTIONAL TESTS ARE SHOWN TO BE NECESSARY BY THE TABULAR HEADINGS AND CONTEXT.

Thus in Genus I (Acidic Species) the primary arrangement of species depends on the results of two coördinating tests, the determinations of melting-point or boiling-point and of neutralization equivalent. Compounds of higher melting- or boiling-point follow those of lower, and among those melting or boiling at the same temperature, those having the higher equivalents follow those

having the lower. With acids so weak that their neutralization equivalents cannot be accurately determined by the titration method of Generic Test I, i.e., those having ionization constants so small that the product $k \times 10^6$ is less than unity, the value of the equivalent is not tabulated; but the space which it would fill in the second column of the table is occupied, when possible, by the insertion of the ionization constant, if known, or in some instances by an approximate verbal statement as to the behavior of the substance in the generic titration. With the liquid acidic species in Division B the specific gravity is so frequently recorded that it is sometimes employed as a third coördinating test, although without the formality of reserving a special column for its tabulation.

6. USE OF THE SPECIFIC DESCRIPTIONS OF THE TABLES.

COMPARE THE PROPERTIES AND REACTIONS OF THE COMPOUND WITH THOSE OF ALL SPECIES DESCRIBED IN ITS SECTION OF THE GENUS THAT MELT OR BOIL WITHIN 5° TO 15° OF ITS OBSERVED MELTING- OR BOILING-POINT. IF THERE ARE NUMEROUS SPECIES WHICH CLOSELY RESEMBLE IT IN THE LIST OF MORE SYSTEMATICALLY TABULATED PROPERTIES, TIME WILL BE SAVED BY DIRECTING ATTENTION NEXT TO PRELIMINARY TESTS MARKED \oplus AND TO OTHER CHARACTERISTICS THAT ARE SALIENT OR QUICKLY DETERMINED. THE SIGN \oplus IN THE TABLES PRECEDES DIRECTIONS FOR THE PREPARATION OF PARTICULARLY CHARACTERISTIC DERIVATIVES. IDENTIFICATIONS THAT DEPEND IN PART ON SUCH PREPARATIONS ARE USUALLY THE MOST SATISFACTORY.

ITALICS IN SPECIFIC DESCRIPTIONS ARE USED EXCLUSIVELY TO DENOTE THAT THE PORTIONS ITALICIZED ARE BASED ON EXPERIMENTS MADE OR VERIFIED IN THE LABORATORY OF THE AUTHOR OF THIS WORK. ITALICIZED DATA ARE NOT NECESSARILY MORE RELIABLE THAN OTHERS, BUT THEY HAVE BEEN SECURED BY PROCEDURES SO FULLY EXPLAINED IN THESE PAGES THAT THERE SHOULD RARELY BE ANY DIFFICULTY IN EXACTLY REPRODUCING ALL CONDITIONS ESSENTIAL FOR THEIR SUCCESS. TO INDICATE EMPHASIS THE EXCLAMATION POINT (!) IS EMPLOYED.

All remarks made in section 6 of the "General Directions for the Identification of an Unknown Compound in Order I," on pp. 6-7 of Vol. I, are equally pertinent in this place, and should be read at this point if not already familiar. But, in addition, a word of warning should be spoken on the importance of critically weighing the conclusiveness of the evidence upon which each completed identification rests. When employing the method of exclusion, particularly, one must never inadvertently conclude that an identification is satisfactorily completed merely because a compound is found to differ in some single determined property from all but one of the species that appear in the part of the tables where it would seem that the description of such a substance should be located; for the tables do not describe all the members of any genus; some descriptions are probably inaccurate in some particulars; and others are obviously incomplete in essential details. Until a sufficient number and variety of significant positive coincidences in properties and reactions have been established, therefore, no opinion as to the identity of the compound should be formed, except as a tentative working hypothesis for guidance in the search for other corroborative data.

The following detailed description of the course of an identification in Order II, taken from the data of an actual experiment recorded in a student's notebook, will serve as a concrete example to illustrate the application of the pro-

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cedures of this volume in a rather complicated case where the final steps, although suggested by the text of the tables, are not there described in detail. The unknown substance was a three-gram specimen of diethyl oxamate, $\text{Et}_2\text{N} \cdot \text{CO} \cdot \text{CO}_2\text{Et}$, as received from a German maker. The parenthetical numeral before each paragraph indicates to which of the numbered sections in the "Directions for the General Procedure to be followed in Identifying an Unknown Compound in Order II" (beginning on p. 2) its procedures correspond.

ILLUSTRATIVE EXAMPLE.

(2) The compound, a colorless liquid of faint aromatic odor, boiling at 249.9° to 252.9° C., and insoluble in cold water, was found to have a specific gravity of 1.027 at $20^\circ/20^\circ$ or 1.025 at $20^\circ/4^\circ$, the determination being made by the pipette method of Vol. I, p. 229.

(3) It left no ash on ignition. It contained nitrogen, but no sulphur, halogens or phosphorus. Test 2.8 having shown that it was not an ammonium salt, it was therefore to be sought in Suborder I of Order II.

(4) As 0.10 gram gave no color with 0.5 cc. of decinormal sodium hydroxide, neutralized less than this quantity in the titration of Generic Test I, and neutralized no hydrochloric acid in the titrations of Generic Test II-III, it evidently might be found described in the B Division (Liquid Compounds) of Genus III ("Neutral Species").

(5 and 6) The arrangement of compounds in this part of the tables depending primarily on boiling-point data, it was considered necessary to scrutinize all characterizations of the Division that are described as boiling between 235° and 258° (Serial Numbers 2.2818 to 2.2842), i.e., for a temperature interval extending 5° above the observed corrected boiling-point, and 15° below it. These particular limits were chosen because they were considered sufficiently wide to prevent overlooking any compounds whose boiling-points might have been carelessly published without stem-exposure corrections, or which might be slightly inaccurate for other reasons. Inspection then showed that among the twenty-five compounds thus included, all but seven could be very quickly and simply eliminated from further consideration. No. 2842 was eliminated because it is described as "miscible with water"; Nos. 2819 and 2839, because as nitro compounds they give Test 2.21, which the unknown substance was found not to give; and all remaining species with the exception of Nos. 2820, 2826, 2830, 2834, 2836, 2840, and 2841, because they differed by more than 3 per cent in density from the unknown substance.

Since the remaining seven species are either nitriles, amides, or esters, which could be most conclusively distinguished and identified by an examination of their saponification products (as is indicated by the presence of the sign \oplus \ominus), saponification with aqueous alkali was next applied according to the procedure of T. 2.26-D. The odor after boiling under reflux for an hour (paragraph (a) of T. 2.26-D) was ammoniacal and slightly fishy; but the iodide precipitate obtained in specific test 2.6 for ammonia was too dark and sticky to prove the presence of this compound as a splitting-product. After two hours saponification the odor in the saponification flask was like that in the U-tube, but much more intense. The distillate collected through the inclined condenser in the next step of the procedure (paragraph 6), was accordingly combined with the contents of the U-tube, exactly neutralized with hydrochloric acid, most of the solution distilled off and set aside, and the remaining portion evaporated to dryness. The dry, white residual hydrochloride weighed about a quarter of a gram, and unlike ammonium chloride was found to be completely soluble in 3 cc. of chloroform and with a melting-point of 206.7° u.c. This completed proof of the absence of ammonia eliminated the three nitrile species Nos. 2820, 2840, and 2841. The volatility and solubility of the basic product in water, together with its ammoniacal, fishy odor, indicated that it could not be methylaniline. This eliminated Nos. 2826 and 2834, and suggested that it might be ethylamine, which should be formed from No. 2830, or more probably the less volatile diethylamine that should form from No. 2836. To secure more convincing evidence, the base was therefore liberated from its salt by treating its very concentrated aqueous solution in a narrow tube with an excess of solid potassium hydroxide. The fuming ammoniacal layer, which at once separated, was removed to another narrow tube by a capillary pipette, dried over caustic potash, and the

boiling-point of a few drops determined by Siwoloboff's method (Vol. I, p. 222) to be 55°. This value corresponds to that tabulated for diethylamine (No. 1068-1). (Ethylamine boils at 19°.) A portion of the amine was then tested by Simon's reaction (T. 2.28), and the remainder converted into its picrolonate (T. 2.23). The Simon reaction gave a deep blue color, indicating a secondary amine, while the picrolonate was obtained in long yellow needles, melting about 260° C. when rapidly heated. The amine was therefore diethylamine, and it became quite probable that the unknown substance was ethyl diethyl-oxamate, No. 2836. To prove this assumption beyond reasonable doubt, it was only necessary to establish the formation of ethyl alcohol and oxalic acid as additional saponification products.



This was next successfully accomplished by use of the procedures of paragraphs (c) and (d) of T. 2.26-D applied to the alkaline solution remaining in the saponification flask, after uniting it with the distillate set aside from the distillation of the diethylamine hydrochloride solution mentioned above. The alcohol was recognized by its boiling-point (Siwoloboff's method) after isolation, and by the iodoform reaction; the oxalic acid, by precipitation as calcium oxalate.

Although the number of chemical operations involved in the investigation here described was unusually large, the identification was completed in a much shorter time than would have been required to arrive at an equally certain result by use of the method of the empirical formula.

CHAPTER II.

GENERIC TESTS OF SUBORDER I, ORDER II.

TITRATION TESTS FOR COLORLESS COMPOUNDS CONTAINING THE ELEMENTS CARBON, NITROGEN, (HYDROGEN), (OXYGEN).

To determine whether a colorless species of Order II should be sought among the "Acidic Compounds" (Genus I), the "Basic Compounds" (Genus II), or the "Neutral Compounds" (Genus III), the following titration tests should be applied to the extent, and in the sequence, directed by the text. Test I must always be applied. Test II-III-(A) must be applied to every compound which has been proved by Test I not to be acidic; Test II-III-(B) has to be made only when the substance is not acidic, and has not been proved basic by Test II-III-(A) alone. Colored compounds — *i.e.*, such as after thorough purification show a color more saturated than a Tint 3 of the color standard (Cf. footnote, p. 3) of this work — do not require titration.

Solids must be reduced to impalpable powders before being weighed for titration by either of the procedures. Whenever a substance, owing to difficult solubility, forms a suspension in the solvent, the suspension must be stirred briskly with a short stirring rod to the end of the titration period, or until complete solution ensues.

GENERIC TEST I.

TITRATION TEST FOR ACIDIC SPECIES.

Place 0.10 g. of the compound in a beaker of 25 to 50 cc. capacity with 10 cc. of distilled water and, if it does not readily dissolve, stir for one minute before beginning the titration. Run in, drop by drop, 0.5 cc. of decinormal sodium hydroxide solution, stirring and watching for the development of any color. If no pronounced color develops, add one drop of a 1 : 300 phenolphthalein solution, and continue the drop by drop addition of alkali until a distinct pink end color is obtained which persists after stirring for sixty seconds. Preserve the solution for use in Test (II-III-(A)) in case the compound proves not to be "Acidic."

The compound is to be sought among the species of Genus I (the Acidic Compounds) in either of the following two cases:

- (1) *If a pronounced development of color is observed upon the addition of alkali alone before the addition of phenolphthalein.*

- (2) If more than 1.00 cc. of decinormal alkali is required to produce a permanent pink color when the solution remains colorless in the first part of the experiment and phenolphthalein is afterwards added.

GENERIC TESTS II-III.

TITRATION TESTS FOR BASIC AND NEUTRAL SPECIES.

PROCEDURE A.

(TITRATION IN WATER.)

*If titration has shown the compound not to be acidic, run into the solution preserved from the experiment of Test I a volume of decinormal hydrochloric acid exactly equal to that of the decinormal alkali previously added; and then titrate with decinormal acid added drop by drop, with constant stirring, using red Congo paper * as the indicator. After the addition of each 0.2 cc. of acid (followed by stirring for one minute in case the substance is not completely dissolved at this point), touch the Congo paper with the wet end of the stirring rod so as to produce a small moistened spot. Repeat until a spot is obtained which, instead of being pink after waiting 15 seconds, is a pale but distinct violet or blue, indicating faint acid reaction.*

This titration may lead to one of two results:

- (1) *More than 1.00 cc. of decinormal acid is required to produce the end color. In this case the compound is Basic and to be sought among the species of Genus II.*
- (2) *The end color is obtained before more than 1.00 cc. of acid has been added. In this case it will be necessary to repeat the titration with an alcoholic solution or suspension by the Procedure B before reaching a conclusion.*

PROCEDURE B.

(TITRATION IN ALCOHOL.)

Place 0.10 g. of the compound in a beaker of 25 to 50 cc. capacity with 10 cc. of neutral ethyl alcohol † and, if it does not dissolve readily, stir briskly for one minute before beginning the titration. Titrate with

* Red Congo Paper. — Dissolve one gram of a good commercial Congo red (the sodium salt of the coupling product of diazotized benzidine and naphthionic acid) in one liter of distilled water. Filter, if not clear, into a large shallow porcelain dish. Soak a number of six-inch square pieces of filter paper in this solution for some minutes. Hang the sheets by one edge on a long wire to dry in a room free from ammonia fumes. Cut the paper in strips one quarter-inch in width, and preserve in a wide-mouthed tightly stoppered bottle.

The filter paper, if not of good quality and free from ammonium salts, will yield a product lacking in sensitiveness. Congo paper as it is furnished by dealers is very likely to prove unsatisfactory.

† Neutral Alcohol. — Good commercial 95 per cent alcohol is often suitable for this titration, but it sometimes contains basic impurities which require partial removal. If a blank experiment shows that 10 cc. of the alcohol require only two or three drops of decinormal acid to give the end reaction described, it should be brought to neutrality before beginning the titration by adding this quantity of acid. If the apparent alkalinity is greater, the alcohol must first be purified by slowly distilling after acidification with sulphuric acid, and rejecting the last 10 per cent of distillate. About 3 cc. of concentrated sulphuric acid per liter should be enough.

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decinormal hydrochloric acid and the red Congo paper indicator, exactly following the directions given for Titration A above, until a faint violet or bluish spot is produced on the paper that remains distinctly visible after 30 seconds.

Regardless of whether the substance dissolves or not, this titration leads to one of the following two results:

- (1) *If more than 1.00 cc. of decinormal acid is required to produce a distinctly violet or bluish spot on the Congo paper, persisting for at least 30 seconds, the compound is to be sought among the basic species of Genus II.*
- (2) *If less than 1.00 cc. of acid suffices to produce this end reaction, the substance is to be sought among the Neutral species of Genus III.*

OBSERVATIONS ON GENERIC TEST I.

This test is not identical with Generic Test III for acids in Order I, although the apparatus and reagents used are the same. It differs from the test of Order I: in the quantity of substance taken for titration; in omitting the additional titration in alcohol called for in certain cases; and in the absence of the requirement of "a sharp and normal color transition in the end-reaction." The first two changes are made merely for economy in material and time; the third, because a sharp color transition in the end-reaction is not to be expected from many feebly acidic species of Genus I in which the presence of strongly "positive" nitrogenous radicals is naturally accompanied by very low ionization constants.

If care be taken to powder the substances thoroughly, and to stir constantly and briskly as directed, very few real acids will be separated from their fellows because of their insolubility in water alone, though many may titrate very slowly. The substitution of alcohol for water as the solvent in titrating very feebly acidic amphoteric species would sometimes lead to a largely increased alkali consumption, and thus emphasize their acidic character; but this substitution is expressly *not* legitimized as an alternative.

Whenever an acid has a sufficiently high ionization constant to ensure a sharp end reaction, *i.e.*, when in the course of the titration a single drop of decinormal alkali, added at the moment when the solution is exactly neutral but still colorless, suffices to develop a full strong pink color which is not greatly intensified by an increase in the excess of free alkali, the data secured should be used for the calculation of the "*neutralization equivalent*" of the acid as explained in Test 1.301. When the exact end-point is difficult to determine, the end-color appearing gradually, the attempt to determine this constant can only lead to erroneous conclusions. The neutralization equivalent of acids which have been successfully titrated, and of those whose ionization constants clearly indicate that they may be successfully titrated, are generally given in the tables.

Substances which give colored solutions with alkali alone before phenolphthalein is added are classified with the acids, because there are in the Order many pseudo acids which while colorless in the free state undergo molecular rearrangement in presence of alkali to colored salts corresponding in structure to labile unknown acids, and these colors are liable to obscure the color change of the indicator. Colorless p-nitrophenol, which dissolves to a deep yellow solution, is a familiar case of this kind. Brownish colorations due to oxidation by atmospheric oxygen in alkaline solutions sometimes develop in the course of titrations. When there is reason to suspect that a coloration is due to this cause, the possibility that the compound may be described in Genus II or III should not be overlooked, acidic species not being the only ones that show such a behavior.

OBSERVATIONS ON GENERIC TEST II-III.

Procedure B, the titration in alcohol, is required only when Procedure A, the titration in water, has not proved the substance to be basic. Its use serves to bring into Genus II many additional weak bases and water-insoluble compounds whose chemical affiliations render their tabulation with the "Bases" more desirable than with the "Neutral Species" of Genus III.

Congo red is selected as indicator because it has been found the one on the whole best adapted for titration of weak bases of any of the readily accessible indicators. So far as convenience is concerned the use of a Congo solution would be preferable to that of Congo paper, but experience has shown that the end-reaction given by weak bases in alcoholic solutions is so much more certainly observed on the paper, that its use in all cases had to be prescribed. To arbitrarily substitute any other indicator for Congo, or to use Congo solution instead of the paper in these Generic tests, would lead to results on which no dependence whatever could be placed!

For evident stoichiometric reasons, all acids and bases whose chemical neutralization equivalents are greater than one thousand will be described with the neutral compounds of Genus III, since they neutralize less than 1 cc. of alkali or acid in the generic titrations. When an end-reaction in Test II-III is very "sharp," the chemical neutralization equivalent calculated for a basic substance from the results by a formula like that used for acids (Test 1.301), will give results in good agreement with the theory. But with the weak bases that cause a very gradual development of the violet spot on the Congo paper, it must not be expected that the alkali consumption will stand in any simple relation to the molecular weight of the substance.

Species of Order II whose colorless solutions become distinctly colored during the titration with decinormal acid in Test II-III are likely to be pseudo-bases, *i.e.*, neutral tautomeric substances which form salts colored in presence of hydrochloric acid in consequence of molecular rearrangement involving the development of a chromophoric group. Such species are, however, extremely rare.

The analyst who is making his first acquaintance with these generic titrations is strongly advised to familiarize himself with the color phenomena of the end-reactions on Congo paper by a few practice titrations with representative amines before applying the procedure to the investigation of unknown compounds. Aniline ($k_B \cdot 10^{10} = 3.5$) may be used to represent the moderately weak, but still readily titratable bases, aminoacetic acid * (with the ionization constants, $k_A \cdot 10^6 = 1.8$, and $k_B \cdot 10^3 = 2.8$), to represent the troublesome amphoteric type, and diphenylamine to represent the compounds which titrate like strictly neutral substances in spite of the fact that they are structurally amines.

Although organic salts are rarely described in the tables of this work, it will not be amiss to here note the variable behavior of a few typical ammonium salts when titrated with the Congo paper indicator. Normal ammonium salts of the strong mineral acids, like hydrochloric or sulphuric, titrate as fully neutral substances. Ammonium acetate behaves in Procedure A and B like a weak base, neutralizing much acid before the first signs of a very gradually deepening end-color appear. Aniline acetate titrates as a neutral compound.

Species whose names are preceded in the tables of this volume by the sign † have had their generic positions fixed by actual titrations by the procedures of this chapter. It is these species which constitute the skeleton that is the chief support for the generic classification of the colorless species of Order II. The generic positions of the great majority of the species which have not been titrated, however, may be regarded as almost equally well established, either because we know compounds of the same type near them in the same homologous series which have been titrated, or can compare their ionization constants with those of very similar titrated compounds, or make use of other analogies whose reliability as a basis for prediction have been repeatedly tested. Among the thousands of compounds described it would be strange if some were not misplaced. But the types among which such errors are most liable to occur are now so well known that serious mistakes can usually be guarded against. Such cases, for example, seem liable to occur among the oxypurines. Thus, 1,3-dimethyl-2,6-dioxypurine (Theophyllin, No. 2.747) has been found by titration to belong to Genus I. Its isomer, 3,7-dimethyl-2,6-dioxypurine (Theobromine, No. 2.2651), however, has been proved a species of Genus III.

* Aminoacetic acid in the titration of Generic Test I in aqueous solution with alkali and phenolphthalein shows a perceptible pink color when 0.8 cc. alkali has been added. With 1 cc. of alkali the color is still very pale, but distinctly stronger. In the titration in water of Generic Test II-III (Procedure A), a pale distinct violet spot appears on the Congo paper upon the addition of 0.2 cc. of acid; while in alcoholic solution by "Procedure B" of the same Generic Test, a distinct spot after the addition of about 0.4 cc. of acid. The compound is accordingly described among the "Neutral Species."

CHAPTER III.

NUMBERED TESTS OF ORDER II.

The numbered tests of this chapter — of which an alphabetical list follows — are important general procedures frequently referred to in the specific characterizations of Order II. They correspond in character to the "semi-specific tests" of Order I, but are arranged and numbered according to a slightly different system from that initiated in Vol. I (cf. Vol. I, p. 2). Following the new notation, the volume of this work in which any numbered test is described will be shown by a numeral in bold-faced type preceding a decimal point in the test number. Thus, T. 2.26 is a numbered test of Vol. II, while T. 1.821 will be found in Vol. I. The new notation is thus made to harmonize with that adopted for the "serial reference numbers" of the specific characterizations. (Cf. p. 2.) The general "Alphabetical Index" at the end of the volume will give access to a description of many additional procedures which it was considered unnecessary to place on the numbered list.

ALPHABETICAL LIST OF NUMBERED TESTS.

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Acetylation.....	2.1
Adamkiewicz-Hopkins-Cole Reaction (Tryptophane).....	2.33
Alkaloid Color Reactions.....	2.2-(a and b)
Alkaloid Precipitations.....	2.3-(a to i)
Amines, Distinguishing between Types of.....	2.35
Amino Group, Detection of by use of Nitrous Acid	2.4
α -Amino-acids, Determination of Nitrogen in, by Van Slyke's Method..	2.5
Ammonia, Specific Test for.....	2.6
Aminonia, Formation of, from Compounds readily decomposed by Alkali.	2.7
Ammonium Salts, Specific Test for.....	2.8
Azo Compounds	2.34
Benzene sulphonyl Chloride, Use of in Characterization and Separation of Primary, Secondary, and Tertiary Amines. (Hinsberg's Method.).	2.9
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Hydrazones and Semicarbazones.....	2.43
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	Number
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Saponification Tests for Amides, Nitriles, Anilides, Imino-ethers, Amidoximes, etc.....	2.26 (A)-(D)
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Ureido-acids and Hydantoins, Preparation and Use of.....	2.31
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2.1. Acetylation.

The acetylation of amines, *i.e.*, the replacement of amino hydrogen by acetyl groups, is a prolific source of derivatives in amine characterization. While no universal method for such acetylations can be given, the simple procedure described below has been often used and will prove successful in a large proportion of the cases where the properties of acetyl derivatives are given in the tables without specification of details. If this method should fail from any cause, the original literature of the compound in question, or the compilation of acetylation procedures on pp. 1272-5, Vol. II, of Weyl's work should be consulted for a more suitable procedure.

Mix one or two decigrams of the substances with twice as much acetic anhydride in a small test tube. Heat to boiling over a very small gas flame, and boil for one minute. Cool. Dissolve the product in the smallest possible quantity of a suitable boiling solvent, water, dilute alcohol, or alcohol being most often used. One cubic centimeter of solvent for each drop of an aromatic amine will often give good results. Cool well, with vigorous shaking, to promote rapid crystallization. Rinse the crystals with a little of the solvent. Recrystallize from a slightly smaller volume of the solvent than was first taken until a product of constant melting-point is obtained. Enough material should, of course, be reserved after each crystallization to permit making a melting-point determination. Should any diacetyl derivative be formed, it is quite likely to be destroyed by hydrolysis, or pass into the final mother liquor during the crystallizations.

2.2. Alkaloidal Color Reactions.

In toxicological examinations of alkaloids the quantity of pure material that can be used in individual tests is usually so small that the strictest economy is imperative. It is largely this condition which has led to the multiplication of alkaloidal color reactions in which some color or succession of colors is devel-

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oped by bringing mere traces of the compound into solution in special liquid reagents, or in contact with them. While the analyst should remember that a reaction of this kind is in general a preliminary or semispecific test which must not be regarded as sufficient identification for any compound without the support of other and adequate corroborating evidence, such tests nevertheless constitute a group of great practical importance, and are often also serviceable in the characterization of compounds not belonging to the alkaloid group.

The procedures for a few of these tests which have been applied in a uniform manner to a large number of species in the writer's laboratory will be described in detail below. Numerous special procedures of the same type which are not mentioned in this place will be found scattered through the specific descriptions of the tables.

The presence of small quantities of impurities sometimes seriously interferes with the alkaloidal color reactions. Thus, to cite an extreme case, practically all the striking color reactions attributed to the alkaloid papaverine, which was discovered in opium in 1848 by Merck, were first shown in 1910 by Pictet and Kramers (*Ber.*, **43**, 1329) to be due to the presence in all papaverine prepared and purified by the usual methods of about 4 per cent of a new alkaloid to which the name of kryptopine was given.

GENERAL PROCEDURE.

Unless the italicized directions prescribe otherwise, the numbered tests below should be performed as follows:

Place about one milligram of the substance on the inverted lid of a small porcelain crucible from which the annular handle has been removed. Close beside the substance drop the specified quantity of the reagent. Incline the lid until the reagent flows against the substance, and observe whether any color appears along the line of contact before they are thoroughly mixed. Mix thoroughly with a small dry stirring rod, and watch for any color or succession of colors that may appear at room temperature within fifteen minutes, and compare them with the color standard. When italicized descriptions refer to heating on a water-bath, the experiments were made by placing the crucible on the thin copper cover of a closed water-bath in which water was boiling.

(a) *With Sulphuric Acid.*

Use one or two drops of concentrated sulphuric acid (sp. gr. 1.84), free from nitric and nitrous acids, and from iron salts. An acid of the quality of Baker and Adamson's C. P. guaranteed and analyzed reagent, such as is used in making the Kjeldahl nitrogen determination, is particularly to be recommended.

(b) *With Nitric Acid.*

Use one or two drops of pure nitric acid of specific gravity 1.42.

(c) *With Sodium Hydroxide.*

Use one or two drops of a 10 per cent aqueous solution.

(d) *With Buckingham's Reagent.*

This reagent is to be prepared only as it is required for use. It is then made up by dissolving 0.01 gram of ammonium molybdate in 10 cc. of pure concentrated sulphuric acid (sp. gr. 1.84).

Use four or five drops of the solution.

(e) *With Erdmann's Reagent.*

The reagent is prepared by diluting 4 drops of concentrated nitric acid (sp. gr. 1.42), with 100 cc. of water and dissolving 20 drops of this mixture in 22 cc. of pure concentrated sulphuric acid (sp. gr. 1.84). It was introduced by Erdmann in 1861. (Ann., 120, 188 and 127, 305.)

Use four or five drops of the reagent.

(f) *With Fröhde's Reagent.*

This reagent must be freshly prepared as required by dissolving 0.01 gram of sodium molybdate in 1 cc. of pure concentrated sulphuric acid (sp. gr. 1.84), with gentle heating. It was introduced by Fröhde for use in testing for morphine in 1866. (Arch. Pharm., 176, 54.)

Use four or five drops.

(g) *With Mandelin's Reagent.*

This reagent must be freshly prepared as required by dissolving 0.01 gram of ammonium vanadate in 10 cc. of pure concentrated sulphuric acid. According to the statement of its originator, a slight dilution of the acid often leads to quite different results than are obtained with the strong acid. (Pharm. Z. f. Russland, 1883, 345.)

Use four or five drops.

(h) *With Marquis' Reagent.*

This reagent is prepared by adding two drops of ordinary 40 per cent formalin solution to 3 cc. of pure concentrated sulphuric acid. (Z. anal. Chem., 38, 467 (1899).)

2.3. Alkaloid Precipitations by General Precipitants.

The so-called "general alkaloidal precipitants" include a great number of acids and salts, inorganic and organic, complex and simple, which give precipitates of difficultly soluble salts from alkaloid solutions. Most of them fail to give precipitates with many alkaloids, and some give precipitates with organic compounds which could not be considered as alkaloids even under the broadest definition of the term. The precipitates vary widely in solubility, but are often amorphous and without other properties particularly favorable for use in specific characterization. This class of tests accordingly plays only a subordinate part in the tables of this volume, if we except the use of picric and picrolonic acid, and of auric and platinic chloride, which are so important as to receive separate treatment as Tests 2.23, 2.39, 2.13 and 2.14. It will, however, prove a convenience to enumerate here a few of the other general precipitants which are of greatest interest to toxicologists * with brief directions for their

* When several of these general precipitants selected from different groups fail to give a precipitate from a suitably prepared solution, toxicologists consider it proper to infer that alkaloids are absent. Gadamer (p. 482), in speaking of this matter, says: "If we select for this purpose phosphomolybdic acid, bismuth-potassium iodide, and tannic acid, a mistake is scarcely possible if the tests are properly performed, it being kept in mind that precipitates may be soluble in an excess of precipitant, and that the delicacy of the reaction is often very dependent on the acid originally in combination with the alkaloid. Thus, cocaine hydrochloride gives no precipitate with tannin at a dilution 1 : 10, while the sulphate still gives a distinct turbidity at a dilution 1 : 100."

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preparation. Complete lists of the alkaloids precipitated by all that are mentioned may be found in Witthaus, pp. 165-172.

(a) Phosphomolybdic Acid. (De Vry's or Sonnenschein's Reagent.)

Perhaps the most inclusive of the general precipitants. It gives yellowish precipitates from acidified solutions of ammonia, of simple substituted ammonias, and of complex alkaloids, which are not soluble in dilute mineral acids—except phosphoric—or in cold alcohol or ether, but are soluble in concentrated hydrochloric, hot nitric, acetic, or tartaric acids, and in alkalies.

The reagent is prepared* by completely precipitating with an excess of sodium phosphate solution at 40° the nitric acid solution of ammonium molybdate used in testing phosphoric acid, thoroughly washing the yellow precipitate after standing for 24 hours, dissolving in the smallest possible quantity of warm sodium carbonate solution, evaporating to dryness, igniting *until all ammonia is expelled from the residue*, moistening the latter with nitric acid and again fusing should it appear blue or black, dissolving in hot water, and then adding nitric acid in sufficient excess to dissolve the precipitate that at first appears.

(b) Phosphotungstic Acid. (Scheibler's Reagent.)

A very general precipitant, giving light colored precipitates from acid solutions, but varying in behavior according to its method of preparation. Rosenthaler (p. 952) recommends that it be made by dissolving 1 part of sodium tungstate in 3 parts of water, and then adding one-half part of 25 per cent phosphoric acid.

(c) Mercury-potassium Iodide. (Mayer's Reagent.)

This gives white or yellowish precipitates in solutions of sulphates or chlorides of most alkaloids, which are often crystalline, or tend to become so on standing. It is prepared by dissolving 1.35 grams of mercuric chloride with 4.98 grams of potassium iodide in 100 cc. of water.

(d) Iodine in Potassium Iodide. (Wagner's Reagent.)

This gives brown amorphous precipitates in solutions acidified with sulphuric acid. It is prepared according to Wormley by dissolving 5 parts of potassium iodide and 2 parts of iodine in 100 parts of water.

(e) Bismuth-potassium Iodide. (Dragendorff's Reagent.)

This usually gives orange precipitates when applied to solutions containing four drops of sulphuric acid in 10 cc. It is prepared (Dragendorff, p. 155) by dissolving BiI₃ in a warm concentrated aqueous solution of potassium iodide, or, according to Kraut, by dissolving 80 grams bismuth subnitrate in 200 grams nitric acid (sp. gr. 1.18), pouring into a concentrated solution of 272 grams of potassium iodide in water, pouring after some days from the potassium nitrate crystals that separate, and diluting to one liter. It must be protected from the light, must not be diluted, nor added to solutions containing ether or amyl alcohol.

(f) Cadmium-potassium Iodide. (Marmé's Reagent.)

This usually gives white precipitates from solutions acidified with sulphuric acid, which are soluble in excess of the reagent or alcohol, but insoluble in ether.

* Sonnenschein, Ann. 104 (1857), 45; Autenrieth, p. 214; Dragendorff, p. 154.

The reagent is prepared (*Z. anal. Chem.*, **6** (1867), 123) by saturating a concentrated boiling solution of potassium iodide with cadmium iodide, and then adding an equal volume of cold saturated solution of potassium iodide. It does not keep if diluted.

(g) Mercuric Chloride.

This solution often gives white crystalline precipitates. Many mercuric chloride derivatives of nitrogenous compounds not belonging to the alkaloids in the narrower sense crystallize well, have good melting-points, and have been analyzed.

(h) Tannic Acid.

This usually gives white or yellowish precipitates with alkaloids. These precipitates are often soluble in acids or an excess of the reagent. Tannin is also classed as a general precipitant for glucosides and bitter principles. The reagent solution should always be freshly prepared. Gadamer recommends the employment of a 10 per cent aqueous solution, or of one made by dissolving 1 part of tannin in 8 parts of water and 1 part of alcohol and then shaken out with ether to remove gallic acid, which is likely to exert a disturbing influence.

The use of picric and picrolonic acids, and of auric and platinic chloride as general precipitants may be inferred from the statements made concerning their derivatives under Tests **2.23**, **2.39**, **2.13** and **2.14**, where their general use in specific characterization is discussed.

2.4. The Amino Group, Detection of, by Use of Nitrous Acid.

This important procedure is used to show the presence of the amino group both in amines and amides. It depends on the fact that, with a few exceptions which will later be discussed, all species of Order II containing this radical and soluble in dilute hydrochloric acid evolve their amino nitrogen when heated with nitrous acid at 100° in accordance with a reaction which—disregarding the intermediate steps—may be expressed by the equation, $\text{RNH}_2 + \text{HNO}_2 = \text{ROH} + \text{H}_2\text{O} + \text{N}_2$. Since the reaction is carried out in the presence of an excess of nitrous acid, which spontaneously decomposes in solution to nitric oxide ($3 \text{ HNO}_2 = 2 \text{ NO} + \text{HNO}_3 + \text{H}_2\text{O}$), the procedure has to provide for the separation of this gas from the nitrogen by absorption. Since the object in applying this procedure is a purely qualitative one, many of the precautions that are necessary in the quantitative determination of the amino group—as in the methods of Hans Meyer,* Stanek † and Van Slyke ‡—may be safely omitted.

Dissolve 0.100 gram of the powdered substance in 2.0 cc. of dilute hydrochloric acid (sp. gr. 1.12), in a 5-inch test-tube standing in a small beaker containing ice water. Drop in, while gently shaking, 2.4 cc. of an ice cold solution prepared by dissolving 5.00 grams of sodium nitrite in water and diluting to a volume of 20.0 cc. Insert the rubber stopper (*A*), bearing the glass capillary tube (*B*), firmly. Connect with the Hempel gas absorption pipette (*C*) which

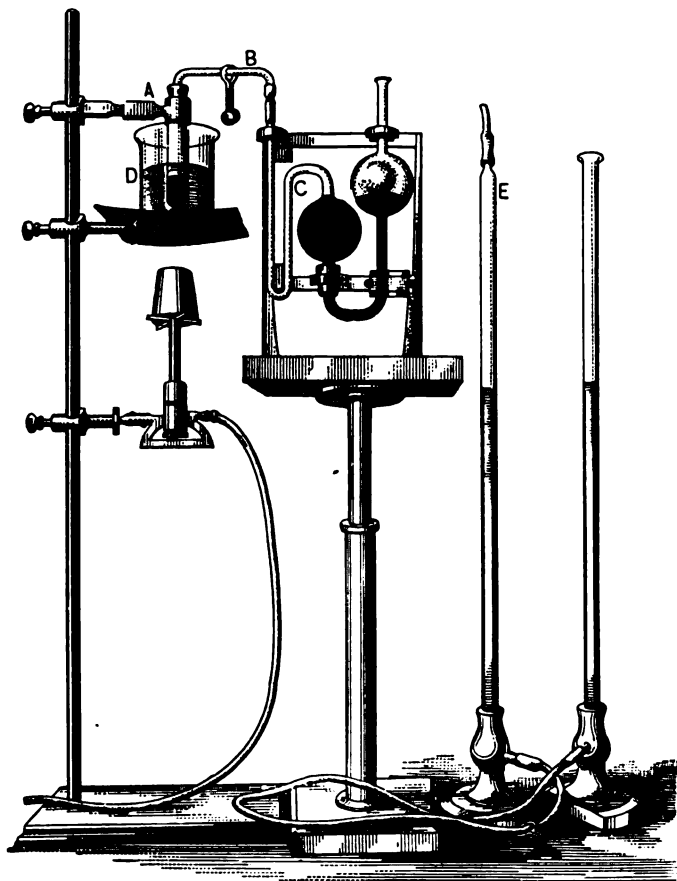
* Anleit. z. quant. Best. d. org. Atomgruppen, 2 Aufl., S. 129. E. Fischer, Ann., **340**, 177 (1905); R. Sachsse & Kohrmann, Z. anal. Chem., **14**, 380 (1875); Euler, Ann., **330**, 287 (1903).

† V. Stanek, Z. physiol. Chem., **46**, 263 (1905).

‡ Cf. Test **2.5**.

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has been filled with a solution * prepared by dissolving 50 grams of potassium permanganate and 25 grams of potassium hydroxide in a liter of water. Immerse the part of the test-tube in which the solution is contained in the small water-bath beaker (*D*). Heat the bath gradually, and finally maintain it at a temperature of 100° until bubbling in the test-tube ceases. Disconnect the absorption pipette and shake until the volume of unabsorbed gas becomes constant. Transfer the gas to the measuring burette (*E*) and determine its volume.



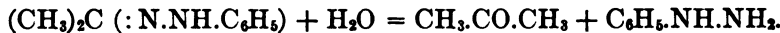
Ascertain by careful blank experiments what volume of air or unabsorbed gases remains in the absorption pipette when the above experiment is repeated in the same test-tube with acid and nitrite in absence of the organic nitrogen compound. Subtract this volume from the volume obtained in the first experiment. If the gas volume thus corrected is not less than 3 cc., the compound probably contains a NH₂ group.

* It has been shown (Van Slyke, J. Biol. Chem., 9, 186 (1911)) that this solution is superior as an absorbent to the saturated solution of ferrous sulphate often used for the same purpose; for besides absorbing nitric oxide much more rapidly than the ferrous solution, it also removes carbon dioxide which is sometimes formed in the test. The ferrous solution is unsuitable for exact quantitative work, because the reaction of nitric oxide upon it is a reversible one. The manganese dioxide which separates from the permanganate solution in absorptions is so finely divided that it does not interfere with the use of the Hempel apparatus. Many absorptions may be made with the same portion of solution, whichever absorbent is used.

The only species of Order II containing an amino group that are somewhat soluble in dilute hydrochloric acid and have molecular weights below 1000 which have been found to fail* to give this reaction, are phenolic amines (o-aminophenol, p-aminophenol, aminothymol, aminosalicylic acid, aminonaphthol (1,5), and aminonaphthol (1,6). m-Aminophenol, unlike its isomers, gives nitrogen. Oxamide being insoluble in dilute hydrochloric acid gives no nitrogen. Dinitroaniline is not soluble enough to give nitrogen unless a little alcohol is added. In other orders hydroxylamine and some sulphonamides have the amino group but yield no nitrogen.

All salts of ammonium give nitrogen freely. This is one reason why it is advantageous to apply the test for ammonium salts (Test 2.8) early in the examination of nitrogenous compounds. Unsymmetrical dimethyl- and diethyl-hydrazines evolve N₂O instead of N₂ (E. Fischer, Ann., 199, 308); but this does not affect the *apparent* result in the test.

Compounds not containing the amino group, but which are readily hydrolyzed by dilute hydrochloric acid to amino compounds, like some of the phenylhydrazone (e.g., acetone-phenylhydrazone), give nitrogen.

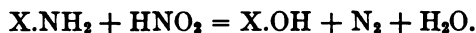


Secondary and tertiary amines, azo and azoxy compounds, and oximes, when they contain no amino group, do not yield nitrogen.

Further remarks on the use of this procedure will be found under Test 2.35.

2.5. α -Amino-acids, Determination of Nitrogen in, by Van Slyke's Method.

This method, now familiar to most physiological chemists, enables the accurate quantitative determination of α -amino nitrogen in all α -amino-acids with an expenditure of only 0.1 gram of substance, in an experiment which may be completed in less than ten minutes. Its introduction has rendered the percentage of α -amino-nitrogen a highly useful numerical constant. The method depends on the fact that these acids when vigorously shaken in glacial acetic acid at room temperature with sodium nitrite solution evolve all of their nitrogen which is present in α -amino groups as nitrogen gas.



Since Van Slyke's apparatus is rather complicated, and adequate working directions for its use do not permit of brief statement, readers wishing to learn the technique of this admirable method should consult Van Slyke's original paper (J. Biol. Chem., 12, 275-284 (1912)).

Van Slyke summarizes the results obtained by the method (J. Biol. Chem., 9, 193 (1911)) in these words: "*Every known amino-acid obtained from proteins by acid hydrolysis reacts quantitatively with one and only one nitrogen atom, excepting lysine ((NH₂)CH₂.CH₂.CH₂.CH(NH₂).CO₂H) which reacts with two.*" All the amino-acids react with all of their nitrogen, except tryptophane (β -indole- α -aminopropionic acid) which reacts with one-half, histidine (β -imidazol- α -amino-propionic acid) with one-third, arginine (δ -guanidine- α -amino-valeric acid) with one-fourth, and proline (α -pyrrolidinecarboxylic acid) and oxyproline with none." The reason for the last results becomes apparent from an inspection

* Amino-groups in the ortho-position to the azo-group are said to be for the most part incapable of reacting with nitrous acid to form diazonium salts. The behavior in this test of compounds having such a structure has not yet been studied.

of the structural formulæ of the substances, tryptophane containing only one-half, histidine one-third, arginine one-fourth, and proline and oxyproline none of its nitrogen in NH₂ groups.

At 20° amino groups in the alpha position to carboxyl react quantitatively in 5 minutes. Ammonium and methylammonium salts require 1.5 to 2 hours. Urea require 8 hours. Amino groups in purines and pyrimidines require 2 to 5 hours. Asparagine reacts only with its α -amino group, the amide nitrogen not being evolved. Peptids, proteins and albumins react with their primary groups only, and give little nitrogen, because most of it is present in peptid linkages. Guanosin anomalously yields one and a quarter instead of one molecule of nitrogen.

Wherever the percentage of amino nitrogen given by any compound in this test is stated in the specific descriptions of these tables, the theoretical value inserted has been verified by the published work of Van Slyke or other investigators.

2.6. Ammonia, Specific Tests for.

In examining the aqueous distillates of ammoniacal odor and alkaline reaction which are so often obtained by the action of heat, alkali, or hydrolyzing agents, on unknown nitrogenous compounds, there is a considerable risk that presence of ammonia may be assumed without sufficient proof; for there are many ammoniacal smelling organic compounds, like the lower alkyl amines, which are formed under these same conditions that give rise to ammonia, and which dangerously simulate its behavior in the ordinary qualitative tests. The present test for ammonia, which was worked out in the author's laboratory some years ago by Mr. E. G. Thatcher, distinguishes it — so far as is now known — from all organic substances with which it is likely to be confused. If other nitrogenous bases are present, they must, however, be first removed, as they may mask the result. A simple and quite general method for effecting this separation, and for distinguishing ammonium chloride from hydrochlorides of all low-boiling amine bases will be found described in Section (a) of Test 2.26-D, and affords by itself a second independent specific test for ammonia that will often be preferred to the Thatcher reaction.

To apply Thatcher's test, add to 3-4 cc. of the aqueous distillate, *which if at all concentrated must first be diluted until the ammoniacal odor becomes only faintly though distinctly perceptible*, 12 drops of 40 per cent formic aldehyde solution. If no precipitate appears at this point,* heat nearly to boiling. Allow to stand for one minute. Then cool to the room temperature, and add 10 drops of a solution prepared by dissolving 12.5 grams of iodine in a solution of 16.5 grams of potassium iodide in 100 cc. of distilled water. Shake well, and allow to stand 4 minutes. Finally, if a precipitate appears, collect it on a very small filter, wash with a little cold water, and note its exact color by comparison, in the moist condition, with the color standard.

If the quantity of ammonia present is large (0.07-0.10 gram), the precipitate will make its appearance at once, and be at first curdy and very voluminous.

* If a precipitate is produced by formic aldehyde at this point, remove it by filtration if possible. If a clear filtrate can not be secured, evaporate the turbid filtrate to dryness on a water-bath in a very small glass evaporator. Add 4 cc. of distilled water, and heat for a few minutes longer on the water-bath. Then add iodine and proceed according to the directions above given.

If the quantity is as small as can be certainly detected (0.0009 g.), the precipitate settles out near the close of the four minute period, showing a powdery microcrystalline appearance. The color of the washed precipitate is a slightly dull orange to yellow-orange (O-YO) to (O-YO) S1. Heavy precipitates settle within about 2 minutes, leaving clear yellowish supernatant solutions. Very dilute solutions appear darker, and become clear only after standing for 10 minutes. *Care must be taken not to regard a deep colored turbid colloidal solution which leaves no precipitate on the paper when filtered as indicating the presence of ammonia.* Many, if not most, amines produce such solutions.

The precipitate formed in this test is hexamethylenetetramine tetraiodide, $C_6H_{12}N_4I_4$. It begins to shrink and soften about 175° , after darkening from 115° , and melts with decomposition at about 202 – 204° to a reddish tarry mass. It is formed in the test by the addition of four atoms of iodine to one molecule of hexamethylenetetramine, the product of the action of ammonia on formic aldehyde. It is unstable in hot solutions. The presence of too large an excess of formic aldehyde tends to prevent complete precipitation. If the test be applied to too concentrated solutions of ammonia, so that there is an excess of free ammonia when the iodine is added, the precipitate will be darkened and its proper color concealed by the formation of black nitrogen iodide.

The following statement concerning the behavior of weak aqueous solutions of a variety of nitrogen compounds when treated with formic aldehyde and iodine according to Thatcher's directions will aid in indicating the limitations of the test. In each experiment 0.1 gram of substance was used if it could be dissolved in 3 cc. of cold water. For less soluble substances 3 cc. of a cold saturated aqueous solution were employed.

Solid precipitates were obtained from: quinoline (after 1 minute, brown to black, curdy, changing to crystalline, and after 24 hours to green needles); ethylenediamine (after 5 minutes, curdy, black); pyridine (2 minutes, nearly colorless crystalline flakes); trimethylamine (2 minutes, curdy OY).

Oily black precipitates were obtained from: nicotine (after 1 minute); coniine (10 minutes); phenylhydrazine (10 minutes); diamylamine, diethylamine, diisobutylamine, ethylamine, ethylaniline, and tripropylamine (all after 5 minutes); methylamine (10 minutes); dimethylamine (15 min.); diethylaniline and morphine (20 min.).

No precipitates were obtained from: acetamide, p-aminophenol, amyłamine, aniline, asparagin, benzamide, benzylaniline, m-bromaline, camphylamine, m-chloraniline, p-chloraniline, cocaine, dipropylamine, diphenylamine, formamide, o-nitraniline, m-bromaniline, p-nitraniline, urea and urethane.

Formic aldehyde alone in the first part of the test gave precipitates with: amyłamine, aniline, m-bromaniline, camphylamine, m-chloraniline, p-chloraniline, dipropylamine, methylaniline, propylaniline, phenylhydrazine.

The special method of fractional condensation and distillation described under Test 2.26-D furnishes another excellent means for partially or completely separating ammonia in aqueous solution from amines boiling below 100° .

For the complete separation of the lower aliphatic amines from ammonia François (Compt. rend., 144 (1907), 857) gives a procedure based on the fact that these amines, unlike ammonia, do not combine with yellow mercuric oxide.

2.7. Ammonia Formation from Compounds easily decomposed by Alkaline Hydroxide Solutions.

The formation of ammonia when a compound is heated with a solution of alkali hydroxide is a much used preliminary test. Tests for ammonia in which

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italicized reference is made to this test number have in general been obtained and tested by one of the following procedures, "(a)" having been employed for alcohol-soluble, and "(b)" for water-soluble species.

(a) Fit a test-tube with a cork stopper carrying a glass tube about 3 inches in length and having an internal diameter of about one-quarter inch. Place in the test-tube 1 gram of powdered potassium hydroxide, 3 cc. of 95 per cent alcohol, and 0.1 gram of the substance to be tested. In the glass tube, place, first, a very thin loose plug of absorbent cotton to stop particles of liquid spray; and, then, above the cotton, a strip of slightly dampened sensitive red litmus paper. (A possibly better arrangement is described by Brach and Lenk [Chem. Ztg., 35, (1911), 1180.]) Boil gently for one minute so that alcohol vapor entirely fills the test-tube without escaping rapidly from the outlet.

(b) The test with water-soluble substances is made as above described under (a) except that a concentrated aqueous solution of potassium hydroxide is substituted for the alcoholic.

In tests whose results have been recorded as positive, the litmus became distinctly blue throughout its length. A faint blueing of the lower end of the paper may be disregarded. A blank experiment should be made with alcohol and potassium hydroxide alone to make sure that the reagents are sufficiently pure. Most substances described as giving this reaction would probably also give it satisfactorily if only one centigram were taken for the experiment. Among the compounds whose behavior in this test has not been so thoroughly investigated as would be desirable are the aromatic nitro and amino compounds. Some aromatic polynitro-compounds and some substituted aromatic amino derivatives are known to yield ammonia.

2.8. Ammonium Salts, Specific Test for.

Ammonium salts are not in general described individually in this work, and can only be identified through its procedure by proving that they contain the ammonium radical, and then identifying their acids. Hence it is necessary to determine at the outset in the examination of every nitrogenous organic compound whether or not it is an ammonium salt. For this purpose the test for ammonium salts usually applied in inorganic qualitative analysis is not entirely reliable; because many organic compounds which are not ammonium salts also evolve ammonia when heated with caustic alkali. The present more discriminating test, which was worked out by Mr. E. G. Thatcher, was suggested by the observation of Remsen and Reid (Am. Chem. J., 1890, 285), that acid amides, unlike ammonium salts, are not rapidly hydrolyzed when boiled with water and magnesia.

Place in a 100-cc. distilling flask provided with a long side tube, 0.1 gram of the substance to be tested, 0.15 gram of finely powdered magnesium oxide, and 15 cc. of distilled water. Support the flask in such a way that its inclined side-tube shall extend to the bottom of a test-tube containing a few drops of distilled water and resting in a beaker of cold water. Distil very slowly until 3 cc. of distillate have been collected. If the distillate is clear, test for ammonia by odor, litmus paper, and by numbered Test 2.6, which distinguishes ammonia from other volatile alkaline amines. If the distillate is turbid through presence of insoluble substances volatile with steam, filter through a wet filter before applying the latter test.

This test does not lay claim to great delicacy. With a salt of the moderate molecular weight of ammonium benzoate, for instance, it is successful with a centigram, but fails with five milligrams. If the attempt were made to increase its delicacy by increasing the concentrations of the reacting substances, the volume of the distillate collected, or by boiling for a considerably longer period under reflux before distilling, the test for ammonia would also be given by urea, and possibly by a few other compounds. But under the prescribed conditions, formamide — which gives off ammonia when boiled with pure water alone — is the only substance which, so far as is now known, would be likely to be mistaken for an ammonium salt.

Among the nitrogen compounds to which the test has been applied with negative results are: acetonitrile, alloxan, aldehyde ammonia, aldehydecyanhydrine, alloxantine, amalic acid, asparagine, benzamide, benzonitrile, biuret, butyronitrile, cyanacetamide, guanidine carbonate, naphthonitrile, oxamide, parabanic acid, phthalimide, propionitrile, propionamide, succinimide, sulphocarbamide, tolunitrile, p-tolnylimidoether, urea, and urethane.

In case it is desired to remove all traces of ammonia in combination as ammonium salts from a mixture before subjecting it to drastic alkaline hydrolysis, as in T. 2.7 and T. 2.26, such ammonia may be most safely expelled by applying the lime and vacuum distillation treatment described by van Slyke (J. Biol. Chem., 10 (1911), 20).

2.9. Use of Benzenesulphonyl Chloride in Identification and Separation of Primary, Secondary and Tertiary Amines.

When a primary amine * is shaken with an excess of potassium hydroxide solution and benzenesulphonyl chloride, the reaction $R.NH_2 + C_6H_5SO_2Cl + 2 KOH = R.NK.SO_2.C_6H_5 + 2 H_2O + KCl$ frequently takes place, and a solution of the water-soluble potassium salt of a benzenesulphonamide is formed. Under the same circumstances secondary amines react generally according to the equation, $R_2NH + C_6H_5SO_2Cl + KOH = R_2N.SO_2.C_6H_5 + H_2O + KCl$, giving a secondary water-insoluble sulphonamide. The free sulphonamide from the primary amine may be precipitated from its alkaline solution by acidification with a mineral acid. The sulphonamides when purified by crystallization are often characteristic derivatives with good melting-points. Tertiary amines do not usually react with benzenesulphonyl chloride. Feebly basic aryl amines, like diphenylamine, and acid amides and nitroamines, also, do not react. The benzenesulphonamides from either primary or secondary amines may be hydrolyzed so as to enable recovery of the original amines by heating in sealed tubes with concentrated hydrochloric acid at 120–160°.

The practical application of these facts in qualitative analysis has been slightly complicated by the discovery † that some primary amines when shaken with benzenesulphonyl chloride and alkali give disulphonyl derivatives, $RN.(SO_2C_6H_5)_2$, which are insoluble in alkali and may be mistaken for the sulphonyl derivatives of secondary amines, and that the sulphonyl derivatives of primary fatty and hydroaromatic amines containing more than six carbon atoms form sodium salts which are insoluble in caustic alkali and easily dissoci-

* Hinsberg, Ber., 23, 2962 (1890); 33, 3526 (1900). Fischer, Ber., 33, 2380.

† Solonino, Compt. rend., 1897 (2), 848; 1899 (2), 867. Marchwald, Ber., 32, 3512 (1899); 33, 765 (1900).

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ated by water to the insoluble free benzenesulphonamides. To overcome these difficulties, as far as possible, the following quite general procedures,* (A) and (B), were developed by Hinsberg and Kessler. They give nearly quantitative yields in most cases, and are valuable methods, though the B procedure is tediously long.

(A)

PROCEDURE FOR PREPARING AND SEPARATING THE BENZENESULPHONAMIDES OF PRIMARY AND SECONDARY AMINES CONTAINING LESS THAN SEVEN CARBON ATOMS.

To about a gram of the amine or amine mixture (1 mol.), add 4 molecules of 12 per cent potassium hydroxide solution; and then, in small portions and with constant shaking, $1\frac{1}{2}$ molecules of benzoyl chloride, warming towards the end to remove the last traces of the acid chloride. If the molecular weight of the amine is not exactly known, estimate the proportions approximately. In case of the lower and more volatile aliphatic amines, cool with ice, and add the acid chloride mixed with the alkali to the amine. Acidify the alkaline liquid with hydrochloric acid. Filter off the precipitate of liberated sulphonamides, or extract it by shaking out with ether. Next, in order to convert any disulphonamide that may have been formed from primary amine into the desired mono derivative, boil the product under reflux for 15 minutes with a solution containing for each gram of the amine 0.8 gram of metallic sodium dissolved in 20 cc. of 96 per cent alcohol. Finally, dilute with water, evaporate off the alcohol to precipitate any alkali-insoluble benzenesulphonamide from the secondary amine, if such is present, and precipitate the benzene sulphonamide of the primary amine by acidification with hydrochloric acid. Any tertiary amine should remain dissolved by the excess of dilute acid.

(B)

PROCEDURE FOR PREPARING AND SEPARATING THE BENZENESULPHONAMIDES OF ALIPHATIC AND HYDROCYCLIC PRIMARY AND SECONDARY AMINES CONTAINING MORE THAN SIX CARBON ATOMS.

Treat the amine or amines with benzenesulphonyl chloride, potassium hydroxide, and sodium alcoholate, exactly as above described under (A). Then dilute the alkaline alcoholic liquid with water, evaporate off the alcohol, acidify with hydrochloric acid, and filter off the sulphonamide or mixture of sulphonamides. Dry, and dissolve in dry ether. Add sodium in small pieces, and warm gently under reflux on a water-bath for 6–8 hours. Allow to cool. Filter. Add ether to the residue in the flask, detach the sodium salt adhering to the pieces of sodium by shaking, and bring the detached salt upon the filter. (Even in the absence of primary amine there may be a slight water-soluble alkaline residue at this point.) Decompose the salt by treatment with hydrochloric acid, thus obtaining the difficultly soluble benzenesulphonamide of the primary amine. The sulphonamide of any secondary amine that may be present may then be isolated by evaporating the ethereal filtrate obtained after shaking out with ether, while any tertiary amine will remain dissolved in the acid aqueous solution.

The benzenesulphonyl derivatives of a very large number of primary and secondary amines have been described by different investigators. Hinsberg † has applied methods (A) and (B) with very satisfactory results to the following compounds among others:

* Ber., 38, 906 (1905).

† Ber., 38, 908 (1905).

methyl, ethyl, propyl, butyl and heptyl amines, pseudocumidine, m-phenylenediamine, butylamine mixed with diethylamine, amylamine with piperidine, pseudocumidine with diethylamine, camphylamine with diethylamine, heptylamine with piperidine, camphylamine with methylaniline, o-xylidine, ethyl-p-xylidine, m-toluidine.

Toluenesulphonylchloride,* 4-nitro-2-sulphonylchloride,† β -naphthalenesulphonylchloride,‡ and β -anthraquinonesulphonylchloride,§ have all often been used for the preparation of characteristic amine derivatives of the benzenesulphonamide type.

2.10. Biuret Reaction.¶

Dissolve 0.01 gram of the substance in a mixture of 2 cc. water and ten drops of 10 per cent sodium hydroxide solution. Then add, drop by drop, the copper sulphate component of Fehling's solution, shaking after each addition until a maximum effect is obtained. Make the test in the cold, and *carefully avoid an excess of copper sulphate*. The result is to be considered positive when a red to violet-blue colored solution is obtained. Thus, with biuret, the first drop of copper sulphate gives a VRT2 color; the second, a VRT1; the third, a VR; the fourth, a RV; and the fifth, a V.

The biuret reaction is a general reaction of the native proteins (blue-violet colorations); of the albumoses and peptones (red to Burgundy red colorations); of some vitellins and histones, and of numerous simpler compounds similar in structure to biuret which form alkali soluble copper compounds. The constitutions of some of these copper compounds have been studied by Schiff.

It is well to convince oneself by a blank experiment with the alkali and copper solution that these reagents alone, in the proportions used, do not, owing to presence of impurities, give a color like that observed in the biuret test. Insoluble substances may be prepared for the test by first boiling with the alkali, and then cooling before adding the copper sulphate. The exact proportion of the substance tested to the reagents is not of very great importance, and in descriptions of the biuret reaction in original publications it is unusual to find any mention made of it.

2.11. Bülow's Reaction.

Dissolve about 0.003 gram of the substance in 5 cc. of pure concentrated sulphuric acid (sp. gr. 1.84). Shake until all is dissolved. Add two drops of a 10 per cent solution of ferric chloride and note the color after 30–40 seconds. Dilution with water will cause the disappearance of the color.

It has been shown by Bülow, Ann., 236, 195 (1886), that phenylhydrazides of the acids ($R.CO.NH.NH.C_6H_5$) give strong red to blue-violet colorations.

2.12. Carbylamine Reaction for Detection of Primary Amines.

Dissolve a crushed fragment of potassium hydroxide of half the size of a pea in 1 cc. of alcohol. Add a centigram or less of the amine and a few drops of

* Hedin, Ber., 23, 3198 (1890). Ssolonina, J. Russ. Phys. Chem. Soc., 29, 405 (1897). Findersen, J. prakt. Chem. (2), 65, 529, 1902.

† Siegfried, H., 43, 69 (1904); Ber., 38, 3054 (1905). E. Fischer, Ber., 39, 540 (1906). Ellinger and Flamand, H., Z. physiol. Chem., 55, 22 (1908).

‡ E. Fischer, Ber., 39, 539 (1906).

§ Hinsberg, Ber., 33, 3527 (1900).

¶ Schiff, Ann., 299, 236 (1898); Ann., 319, 300 (1901). Neumeister, Z. Biol., 26, 324 (1890). Schaar, Z. anal. Chem., 42, 1 (1903).

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chloroform and boil. If a strong carbylamine odor is not given off at once, wait for a minute, and heat again!

Primary amines, unless of very high molecular weight, form isocyanides in this test, according to the equation $XNH_2 + CHCl_3 + 3 KOH = X \cdot N : C + 3 KCl + 3 H_2O$, which are readily recognized by their singularly nauseating penetrating odor. The chief weakness in the test is its excessive delicacy. It is not given by secondary or tertiary amines; but the traces of these isomers which are often present in imperfectly purified preparations sometimes occasion mistakes. The reaction when properly used is very convenient in preliminary testing, and is particularly useful when it is desired to ascertain quickly whether aniline or similar aryl amines have been formed by the breakdown of more complex compounds in saponification experiments. Anilides usually give the carbylamine odor directly, sufficient aniline being liberated by the alkali during the brief period of heating without any special preliminary saponification treatment. The test was first used by A. W. Hofmann (Ber., 3, 767 (1870)). It is superior to the mustard oil reaction which was proposed by this chemist for the same purpose.

The sensation produced by carbylamine vapors when well developed is one which involves the sense of taste as well as that of smell. With most persons the odor produced by the lower arylamines seems more powerful than that by the aliphatic. Ammonia, ammonium salts, and some acid amides also give faint carbylamine-like odors in this test, although not strong enough to interfere with its use, or be easily detected.

2.13. Chloroaurates, Preparation and Use of.

The chloroaurates of basic nitrogen compounds often crystallize well, are not difficult to prepare, and have accordingly been long and widely used in specific characterization. Their melting-points are often of diagnostic interest; but their gold content, which is determined by the simple operation of ignition in a porcelain crucible, has an even greater analytical importance. The following general directions have been followed with success in a number of preparations, but will of course require modification in some cases.

Dissolve 0.10 gram of the substance in 5 cc. of water acidified with five drops of dilute hydrochloric acid (sp. gr. 1.12). Add $1\frac{1}{4}$ times the theoretically calculated quantity of chloroauric acid, $HAuCl_4 \cdot 4 H_2O$, in the form of a 5 to 10 per cent aqueous solution. (In confirmatory testing the formula for the chloroaurate product will be found in the tabulated description.) Wash the precipitate, which is often curdy, by dropping upon it 4 cc. of cold water, and draining by suction. Recrystallize from boiling water, alcohol or dilute alcohol. Wash by suction with 2 cc. of the cold solvent. In drying chloroaurates for melting-points or analysis, remember that they often contain water of crystallization.

2.14. Chloroplatinates, Preparation and Use of.

The chloroplatinates of the nitrogen compounds of basic character are among their most frequently prepared derivatives. When more specific directions for their preparation are not furnished in the tables, the following procedure may be tried.

Dissolve 0.2 gram of the substance, if possible, in 1.0 cc. of water + 1 cc. of dilute hydrochloric acid (sp. gr. 1.12), or in 2 cc. of 95 per cent alcohol + 0.5

cc. of dilute acid. Add a trifle more than the theoretically calculated quantity of a 10 per cent aqueous solution of chloroplatinic acid, H_2PtCl_6 . In confirmatory testing the quantity may be calculated from the formula of the expected product as given in the tables. As some chloroplatinates crystallize out very slowly, allow the mixture to stand for some time, or even to concentrate slightly by evaporation, before filtering by suction on a small hardened filter. Wash with a cubic centimeter of the cold solvent, using suction. Recrystallize slowly from a small volume of the solvent, and wash as before by suction, dropping about 2 cc. of the cold solvent on the crystals. After drying on a bit of porous tile to constant weight, determine the melting-point and the percentage of platinum by ignition in a small porcelain crucible. Many chloroplatinates contain water of crystallization, and the method to be adopted in drying will depend upon the temperature at which this water is lost, and whether it is desired to make a determination of this water while drying.

2.15. Diphenylamine Reaction for Nitrosamines ($XY.N.NO$), Nitrates, Nitrites, and Aliphatic Nitro-compounds.

To 5 cc. of the cold diphenylamine reagent * in a test-tube, add one or two milligrams of the powdered substance to be tested, or, if it is a liquid, as much as will adhere to the tip of a stirring rod. Shake to effect solution. When the result of this test is positive, an intense pure blue color develops within one minute.

The color produced in this reaction is one of great intensity and considerable stability. When diluted with enough concentrated sulphuric acid, to permit comparison with the color standard, it is blue (B). A little of the solution diluted with water may give a violet color. The test *is given* by all compounds, organic or inorganic, which liberate nitric or nitrous acid upon treatment with concentrated sulphuric acid. A good reaction may be expected from any nitrate or nitrite salt or ester, from any nitrosamine having the NO group joined to nitrogen, and from the nitroparaffines. In general, *it is not given* by aromatic nitro or nitroso compounds having the nitro or nitroso group joined directly to a carbon atom, nor by oximes, nor by isonitroso compounds. *Because of the delicacy of the reaction, substances which have been prepared by aid of nitrous or nitric acid need the most careful purification before being tested.*

Since the same blue color is produced by numerous non-nitrogenous oxidizing agents, such as chloranil, thymoquinone, hydrogen peroxide, ferric chloride, potassium manganate, chromate, chlorate and iodate, the reaction must not be accepted as a test for nitrosamines, nitric or nitrous salts or esters, or nitroparaffines, in cases where any of these interfering substances may be present. The introduction of the reaction as a test for nitric and nitrous acids is due to E. Kopp (Ber., 5, 284 (1872)), and the disturbing influence of other oxidizing agents was pointed out by P. Soltien (Chem. Zentr., 1885, 586).

* The Diphenylamine Reagent.—To prepare this reagent mix four volumes of pure sulphuric acid of sp. gr. 1.84 with one volume of distilled water, and dissolve in the mixture, while still warm, 0.02 gram of diphenylamine for each 100 cc. If the sulphuric acid employed is of high purity, nitrogen free, the reagent may be kept in stock for a long period without deterioration. If it contains traces of nitrosoyl sulphate, like most commercial acid prepared by the lead chamber process, it will soon turn blue on standing.

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From experiments made by Mr. Alden Merrill in a preliminary study of the use of this reaction as a coördinating test, the following illustrative cases may be cited.

The blue color was obtained with: the nitrates of amyl, aniline, cellulose and glycerine, and with celluloid; the nitrite of amyl; the nitrosamines of methylaniline, ethylaniline, diphenylamine and dimethylamine; the aliphatic mono-nitro derivatives of methane, ethane, propane and pentane; and with dinitroso-2,5-dimethylpiperazine.

The blue color was not obtained with: the nitro compounds, nitrobenzene, m-dinitrobenzene, 2,4-dinitrotoluene, 1,3,5-trinitrobenzene, 2,4,6-trinitrotoluene, trinitroxylene, o-, m-, or p-nitrophenol, α -nitronaphthalene, 2,4-dinitrophenol, o-, m-, or p-nitrotoluene, o-nitroanisole, picric acid, m- or p-nitrobenzoic acid, dinitrobenzoic acid, dinitronaphthol, o-nitrobenzaldehyde, benzoylnitrotoluide; the nitroso compounds, the nitrosotoluidines, nitrosobenzene, nitrosophenol, α -nitroso- α -naphthol, p-nitrosodiethylaniline, p-nitrosodimethylaniline, nitrosothymol; the isonitroso compounds of acetone and acetoacetic ester; and the oximes of acetone, camphor, methylpropylketone, sanatonin, p-nitrobenzaldehyde, benzophenone, and benzoldioxime.

2.16. Dye Tests in Order II.

Descriptions of commercial dyestuffs for whose identification provision has already been made in Volume III of this work are usually not repeated in this volume. The following tests (a), (b) and (c), based on procedures used and more fully discussed in Volume III, are referred to in the tables in describing some species having dyestuff properties.

(a) Dyeing on Wool. (Cf. Test 3.5.)

Add one square inch of wetted woolen Henrietta cloth which has previously been well boiled out with water to a solution or suspension of 0.01 gram of the powdered substance in 10 cc. of distilled water. Heat the test-tube containing the solution by immersion in a boiling water-bath, stirring the cloth about from time to time. If the solution dyes the cloth, continue the heating until the color of the cloth after rinsing approaches as closely as possible to a "normal tone" of the color standard. Then rinse with cold water and wring out. Bring into a small beaker containing 25 cc. of distilled water actively boiling, and boil for just one minute. Rinse again with cold water. Wring and dry. Compare the color of the wool after this treatment with the color standard.

Cut off a 7-mm. wide strip of the dyed fabric for use in Test (c), saving the remainder for Test (b).

(b) Reduction Test with Rongalite C. (Cf. Test 3.8.)

After wetting with water, boil the piece of colored cloth reserved for the purpose in (a) in a test-tube with 10 cc. of a freshly prepared 5 per cent solution of Rongalite C. until the color is discharged, provided a discharge can be obtained by not more than two minutes of actual boiling. Note the time required to produce a complete discharge, or the color of the cloth at the end of the two minute period when the discharge is incomplete.

If the color is discharged, rinse the cloth with cold water, wring and expose to the air for 10 minutes. Note what color, if any, appears in consequence of oxidation.

(c) Action of Sulphuric Acid and Sodium Hydroxide on Wool Dyeings. (Cf. Tests 3.13 and 3.14.)

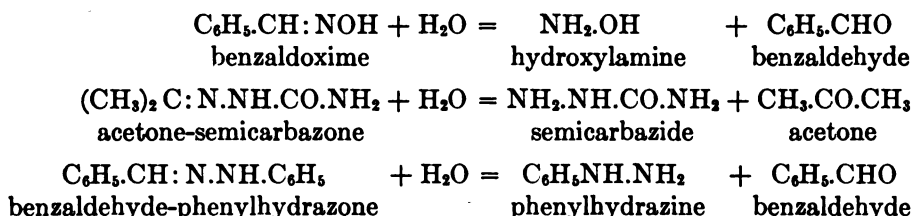
Cut the 7-mm. wide strip of dry dyed fabric reserved from Test (a) into 7-mm. squares. Place one of these squares in the shallow circular depression of one of

the glazed white porcelain tiles used in analytical laboratories for drop testing in volumetric analyses. Cover with 1 cc. of pure concentrated sulphuric acid (sp. gr. 1.84). Noting the time, work the acid into the fiber for *just one minute* by continuous gentle rubbing with the end of a smooth glass rod. Then observe the exact color of the wool.

Repeat the experiment as described, except that a 10 per cent aqueous solution of sodium hydroxide is substituted for the sulphuric acid.

2.17. Hydrolytic Splitting of Oximes, Hydrazones, Semicarbazones, etc.

This test depends on the fact that oximes, hydrazones and semicarbazones may usually be rather easily hydrolyzed when boiled with aqueous hydrochloric acid to the aldehydes or ketones, and the hydroxylamines, hydrazines, or semicarbazide, to which they are genetically related. Thus:



Before applying this test it should be determined whether the original substance will reduce Tollen's reagent in Test 2.30. If it does reduce this reagent directly, neither of the two lettered procedures below — excepting that part of (b) relating to the detection of oximes by the benzhydroxamic acid reaction — can be expected to furnish useful results.

Procedure (a).

Heat for five minutes in a small test-tube immersed in a boiling water-bath 0.05 to 0.10 gram of the substance and 1 cc. of hydrochloric acid (sp. gr. 1.20). Evaporate the mixture to dryness on a watch glass on the water-bath. Treat the residue with 3 cc. of cold water. If a clear solution is not obtained, filter. Pour the clear solution or filtrate into 5 cc. of Tollen's reagent. (Cf. T. 2.30.)

If the substance is a readily hydrolyzed oxime, hydrazone, or semicarbazide, an immediate dark colored precipitate of metallic silver will appear. The characteristic odor of the aldehydes or ketones liberated in such hydrolyses may often be recognized during the experiment. A good silver precipitate may be anticipated whenever the carbon atom of the carbonyl group that would be formed in case of hydrolysis is situated in an open chain; but it may be unsatisfactory in cases when this atom is situated in a cyclic nucleus.

Among the substances which have been found to give good positive reactions in accordance with the above rule may be mentioned: acetaldoxime, methylethylketoxime, methylpropylketoxime, *cenanthaldoxime*, isonitrosobenzaldoxime, benzylideneacetoxime, benzaldehydephenylhydrazone, benzaldehydediphenylhydrazone, salicylaldehydephenylhydrazone (which reduces silver directly in Test 2.30), benzaldehydephenylbenzylhydrazone, phenylgalactosazone, acetonesemicarbazone, benzaldehydesemicarbazone. Camphoroxime (camphor having the carbonyl group in the nucleus) gives only a very slight darkening.

Procedure (b).

(Including a test for hydroxylamine, and applicable to some refractory oximes not split by procedure (a).)

Place 0.2 gram of the substance in a strong 3-inch test-tube with 3 cc. of hydrochloric acid (sp. gr. 1.20). Stopper tightly, securing the cork by strongly wiring with two copper wires crossing at right angles. (See Fig. 3, p. 112, Vol. I.) Suspend the tube in boiling water for 30 minutes. Dilute with two volumes of water. Cool and filter. Evaporate the filtrate on the water-bath to dryness. Dissolve the residue in 12 cc. of cold water. Test 2 cc. of this solution for reducing power with Tollen's solution as in procedure (a).

As a more specific test for hydroxylamine, dissolve in the remaining 10 cc. of solution 0.25 gram of crystallized sodium acetate, add one drop of benzoyl chloride, and shake violently for one minute. Then add four drops of 10 per cent aqueous ferric chloride solution, and, finally, about four drops of concentrated hydrochloric acid, or enough to destroy the turbid yellow appearance due to the action of ferric chloride on the excess of sodium acetate.

The appearance of a red-violet or purple solution at this point is the test for hydroxylamine. The color is due to formation of benzhydroxamic acid according to the equation: $C_6H_5.COCl + NH_3OH = C_6H_5.CO.NHOH + HCl$. This color is much more stable towards a moderate excess of hydrochloric acid than most of the similar colorations given by the phenols. The coloration is quite strong when over one milligram of hydroxylamine is present, but is quite faint in presence of only half a milligram. To ensure delicacy, do not use larger quantities of any of the reagents than is here prescribed. The test is based on a reaction first described by E. Bamberger (*Ber.*, **32**, 1805).

Procedure (b), besides being successful with the oximes mentioned as giving positive results under (a), has been reported as satisfactory with benzophenoneoxime, p-nitrobenzaldoxime, and phenanthrenequinoneoxime; as rather unsatisfactory with camphoroxime and benzildioxime; and as entirely inapplicable in the cases of benzoquinoneoxime (nitrosophenol) and santoninoxime.

Test (b), but not (a), further, gives positive results with primary and secondary nitroparaffines, which are decomposed in the closed tube according to the equation (*Ann.*, **180**, 163), $R.CH_3.NO_2 + H_2O = R.CO_2H + NH_3OH$. This has been verified in the cases of nitromethane, nitroethane, nitropropane, and nitropentane.

2.18. Liebermann's Reaction for Nitrosamines.

As usually performed * this test has apparently been made by dissolving a little (about 0.2 gram) of the nitrosamine together with about an equal quantity of phenol in a cubic centimeter or two of pure concentrated sulphuric acid at the room temperature, pouring the resulting blue solution into several volumes of cold water, and finally supersaturating with sodium hydroxide solution. The dilution with water is followed by the separation of a brownish precipitate, and the addition of alkali by the formation of an intense pure blue solution, which is the most characteristic phenomenon in the test.

So far as known, Liebermann's reaction is not generally given by nitroso compounds in which the nitroso group is joined directly to carbon, by the oximes (with exception of the tautomeric nitrosophenol), or by nitro compounds. Yel-

* V. Meyer and Janny, *Ber.*, **15**, 1529.

lowish and greenish colorations may be produced by colorless compounds belonging to many groups, but should be disregarded. A precise statement of the experimental conditions under which this test has been performed, and of the exact phenomena observed where it has been applied, is usually lacking in the original literature, though the references to it are very numerous. The somewhat similar, less specific, but more thoroughly studied diphenylamine reaction is, hence, at present, a more reliable test for determining whether a compound may be a nitrosamine.

The blue alkali-soluble coloring matter of this test, according to Liebermann,* is produced by the reaction of phenol on nitrosophenol, the latter compound being first formed from a portion of the phenol by the action upon it of nitrous acid generated from the nitrosamine.

2.19. Millon's Reaction.

Heat together in a test-tube about 0.01 gram of the substance to be tested and 2 or 3 cc. of Millon's reagent.† The result of the test is to be considered positive when the substance becomes pink or red, or when a pink or purplish red solution is obtained.

This test has long been used in the examination of proteins and proteolytic products. The colorations are due to the action of the reagent on phenolic components like tyrosin. In testing protein solutions the reagent is added in excess, and the mixture heated. In this case a colorless precipitate will at first appear, which will later become colored and dissolve as described above. For a fuller discussion of the test, see papers by Vaubel (*Z. angew. Chem.*, 1900, 1125), and by Nasse (*Pfluger's Arch.*, 83, 361 (1901)).

2.20. Murexide Reaction.

Mix in a very small porcelain evaporating dish, or on the lid of a porcelain crucible, a couple of milligrams of the powdered substance, a minute crystal of potassium chlorate and two drops of dilute hydrochloric acid (sp. gr. 1.12). Evaporate to dryness on a boiling water-bath, and then continue the heating for several minutes longer. If the residue at this point should remain entirely colorless, support the porcelain on wire gauze, and cautiously heat to a somewhat higher temperature with a small flame until the residue colors slightly pinkish or brownish. Then cool and treat with two drops of dilute ammonia. The result of the test is positive when a strong purple-red colored solution is obtained. The color given by uric acid, caffeine, theobromine, and xanthine is intense VR.

The murexide test is known to be given by many purine derivatives, and its literature is quite extensive. The purple product formed when it is applied to uric acid is murexide, the ammonium salt of purpuric acid, ($\text{NH}_4\text{C}_8\text{H}_4\text{O}_6\text{N}_5$), while the corresponding product from caffeine, murexoin, is tetramethylmurexide. Instead of potassium chlorate and hydrochloric acid, other oxidizing agents, like nitric acid, bromine water, and chlorine water, have been

* Liebermann, *Ber.*, 7, 248, 1098.

† **Millon's Reagent.** — This reagent is prepared by treating mercury with twice its weight of nitric acid of specific gravity 1.42, first, in the cold, and then gently warming until solution is complete; then, adding two volumes of water, and after standing for some hours, decanting off as reagent the clear solution from the crystalline precipitate which separates.

often recommended. The present procedure with chlorate, which is practically identical with what is known as the "Weidel test" for xanthin, is, however, according to our experience, much the most reliable. The addition of a little strong potassium hydroxide to the colored solution has been observed in some cases to cause the color to become bluer (uric acid and theobromine give an approximately RV color), but in other cases (caffeine and theophylline) to entirely discharge it. It is sometimes recommended to expose the evaporated residue to the fumes of strong ammonia rather than to treat it with ammonia solution.

2.21. Nitro Group, Tests for.

Dissolve about 0.2 gram of the substance in about 3 cc. of hot 50 per cent alcohol. Add five or six drops of 10 per cent calcium chloride solution and a pinch of zinc dust, and heat until violent boiling begins. Chemical action accompanied by frothing will then often continue for some time without further heating. Allow to stand from two to five minutes. Filter into a 10 per cent solution of silver nitrate in concentrated ammonia. Nitro compounds thus treated give an immediate black or gray precipitate of metallic silver, or a silver mirror. This test must be accompanied by a blank experiment with an alcoholic solution of the substance and ammoniacal silver nitrate; for it is, of course, useless to apply it to compounds that reduce silver before treatment with zinc dust. A white or light yellow precipitate in the test is without significance.

The usual reduction products of nitro compounds by the procedure of the test are hydroxylamines. Thus, nitrobenzene gives phenylhydroxylamine, $C_6H_5NO_2 + 2 H_2 = C_6H_5.NH(OH) + H_2O$. The only important classes of nitrogenous carbon compounds not containing the nitro radical, which, while unable to reduce ammoniacal silver nitrate in the cold, are known to gain the power to do so after the treatment with zinc dust, are nitroso, azo and azoxy substances. Their reduction gives hydroxylamines, hydrazines, or hydrazo compounds. Nitraniline gives aminophenol, which reduces silver solution. The addition of calcium chloride is made to accelerate the reaction.

In the aliphatic series this test has been successfully used, for example, with the mono-nitro-derivatives of methane, ethane, pentane and with chlorpicrin; in the aromatic series with mononitroderivatives of benzene, toluene, naphthalene, chlorbenzene, brombenzene, anisole, cinnamic acid dibromide, benzoic acid, benzenesulphonic acid and with p-nitrophenylpropionic acid; with dinitro compounds of benzene, aniline and phenol; and with trinitro derivatives of benzene, toluene, phenol and triphenylmethane.

For a fuller account of the test see Mulliken and Barker (Am. Chem. J., 1899, 271).

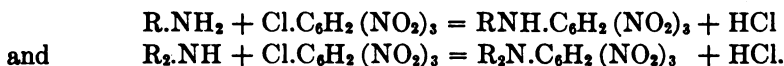
2.22. Picramides from Primary and Secondary Amines.

Dissolve 0.1 gram of the amine in 1 to 2 cc. of alcohol. Add 10 cc. of a 2 per cent solution of picryl chloride in strong alcohol. If a precipitate appears, filter after shaking and allowing to stand for a few minutes. If no precipitate appears at once, heat to boiling. Then cool well with running water, scratching the wall of the test-tube with a glass rod, and again shake after adding enough water — when it is necessary — to produce a slight permanent turbidity. After standing, wash the precipitate with a few cubic centimeters of cold alcohol. Dis-

solve the precipitate in 10 cc., or, if possible, in less hot alcohol or hot glacial acetic acid, the more suitable solvent being specified in the text of the tables, and allow it to crystallize out slowly on cooling. Note the color of the original precipitate as well as that of the recrystallized product. Dry the crystals at 110°, unless they melt at a lower temperature, and determine their melting-point.

Liquid amines should be dropped into the picryl chloride solution until present in slight excess. This point is reached when the solution shows a deep red color that does not disappear on shaking.

Picramides are formed from primary and secondary amines by the reactions:



As derivatives they have the advantage over many of the corresponding picrates of being more stable and less soluble, and in melting more sharply and with less decomposition. They also equal or excel the picrates in their tendency to crystallize well and quickly, and with brilliant colors. The differences between the colors of the picramides of isomeric amines are frequently striking. (See characterizations of the isomeric anisidines, naphthylamines, and toluidines.)

Picryl chloride may be obtained from the larger general dealers in organic chemicals, or may be very easily prepared from picric acid and phosphorus pentachloride as described by Jackson and Gazzolo (Am. Chem. J., 23, 384). It is soluble in 45 parts of cold, or in 4 parts of boiling alcohol. The alcoholic picryl chloride solution should not be prepared in large quantities, but may be kept for over a month, at least, without spoiling.

The directions for the test as here given are based on experiments made on over thirty amines by Mr. Max Cline. In special cases slight modifications in the procedure would probably give better yields. The preparations may often be made successfully on a smaller scale than is recommended.

2.23. Picrates and Picrolonates, Preparation of, as Derivatives in Order II.

The picrates of the species of Order II are of extraordinary importance as derivatives for confirmatory testing. Their importance is due to the singular ease with which they may be prepared, purified, and crystallized; to the wide range in their solubilities, melting-points, colors, and crystalline forms; and to the surprisingly large number of cases in which they have been carefully described in the chemical literature. Appreciation of these circumstances has made it desirable to mention in the tables the picrates of practically all included species, so far as access to published data regarding melting-point and color rendered it possible. A clue to the identity of species whose physical properties make them difficult to isolate in a condition of purity is thus often provided in the preparation and examination of their picrates.

Detailed directions for the preparation on the small scale of a large number of picrates for specific test purposes are given in the tables. (For examples see Tests 2.797, 2.1060, 2.1546.) In addition to these special directions, the following general information concerning picrate preparations will be found useful.

Various solvents are employed. When water is applicable, the precipitation is effected by about one and a quarter times the theoretical quantity of a saturated aqueous solution of picric acid. (100 parts of cold water dissolve a

little more than one part of the acid.) When the substance is not soluble in water, the solution may often be prepared, in the case of basic compounds, by adding 1 to 3 drops of dilute hydrochloric acid (sp. gr. 1.12) to each 2 or 3 cc. of water. If a heavy precipitate does not form at once, the solution should be allowed to stand for some time, and well shaken before being filtered. The precipitates should be collected and washed on very small filters, recrystallized, washed and dried with observance of the technique described on pp. 234-235 in Vol. I. Organic solvents have to be substituted for water in certain cases.

Working on the small scale recommended, 2 cc. of solvent will sufficiently remove the excess of picric acid, provided the precipitate is first brought into the point of a very small filter. When the directions are not explicit, boiling water, boiling alcohol of 95, 66 or 50 per cent, boiling 10 per cent glacial acetic acid, and boiling benzene, are the solvents that are to be tried in the order here given. The crystallization may be promoted, when necessary, by cooling and vigorous shaking. In working with glacial acetic acid, excellent results are sometimes obtained when the picrate is very soluble, by adding water until the solution begins to cloud, warming to remove the cloudiness, and then cooling. The crystals are usually washed with 1.0 cc. of cold solvent. Most picrates melt above 100°, and may therefore be dried completely on a bit of porous tile in a 100° drying-oven in 15 minutes.

The *picrolonates*, or colored salts of 1-p-nitrophenyl-3-methyl-4-nitropyrazolone(5) with the basic nitrogen compounds, are often *less soluble* than the corresponding picrates, have higher melting-points, crystallize equally well, and are sometimes more characteristic derivatives, although the melting-points of adjacent picrolonates of compounds in the same homologous series are usually less widely separated. They are prepared in the same general manner as picrates by precipitating solutions of the amines or their salts with an aqueous or alcoholic solution of the acid. Some of the more soluble and strongly basic of the amines have been precipitated from aqueous solutions after saturation with carbon dioxide (Cf. No. 2.1059), others from solutions slightly acidified by hydrochloric acid, while some of the alkaloidal picrolonates may be quantitatively precipitated from ether-chloroform solutions.

Picrolonic acid is described as No. 2.3184, and the identification of picrolonates under Test 2.39. A partial bibliography of picrolonic acid and the picrolonates will be found in the Archiv der Pharmacie, 245 (1907), 112.

2.24. Pine Splinter Reaction for Pyrrole Derivatives.

(a) *Test with Volatile Substances.*

Suspend a freshly cut splinter of soft pine wood which has been soaked for a few minutes in concentrated hydrochloric acid in the vapor rising from a few drops of the substance boiling by itself, or mixed with a little water, in a test-tube.

With pyrrole, the splinter is quickly colored a pale red, soon changing to deep carmine. The result of the reaction may be considered as positive whenever a color between red and violet is obtained.

(b) *Test with Non-volatile Substances.*

Boil a freshly cut pine-splinter for a minute or two in an alcoholic solution containing about 0.001 gram of the substance to each cubic centimeter of 95 per cent alcohol. Then soak the splinter in cold hydrochloric acid (sp. gr. 1.20).

Substances like carbazole which contain the pyrrole ring, but are too difficultly volatile and water-insoluble to be tested by procedure (a), may give very distinct red stains in procedure (b), but the colorations are seldom, if ever, so intense as those produced by pyrrole vapor.

The colored compounds formed in both procedures are said * to be condensation products of pyrrole or its derivatives with an aldehyde, "hadromal," which is present in varying quantities in woods. The field of usefulness for these tests has been considerably widened by their indirect application in the characterization of substances from which pyrrole derivatives may be prepared by simple methods. Neuberg's publication, cited above, includes a discussion of these additional applications, arranging the compounds which may be thus tested in the following three groups:

(1) *Nitrogenous substances which evolve pyrrolic vapors, giving Test (a) on being decomposed by simple ignition in a glass tube.* — Among these are many ammonium salts and amino acids, including glutaminic acid, serin, glucosaminic acid, glycocoll, alanine, leucine, tyrosine, phenylalanine, asparagine, pyrrolidinecarboxylic acid, arginine, lysine, histidine, diaminopropionic acid, cystin, diaminosuccinic acid, and taurin.

(2) *Nitrogenous substances which give pyrrolic vapors when ignited with zinc dust.* — Among these are ammonium oxalate and malonate, which do not thus react alone, and a group, including glycocoll, alanine, leucine, tyrosine, phenylalanine, asparagine, which give the reaction rather feebly without the addition. The result is considerably influenced by the manner of ignition; and, if the conditions are such that sublimation is prevented, the carbon liberated will often take the place of the zinc dust as a reducing agent. The reaction may be obtained in one way or another from the ammonium salts of all the carbonic acids of the carbohydrate group, as well as from other nitrogenous derivatives of the sugars, like glucosoxime. (In illustration see No. 2.2037.)

(3) Non-nitrogenous compounds like the γ -diketones which yield pyrrole derivatives by the procedure of L. Knorr (Ber., 19, 46 (1886)), which has already been described in T. 1.703.

2.25. Rimini's Test † for Primary Aliphatic Amines.

To a solution or suspension of one drop of the compound (or an equivalent quantity of a solution) in 5 cc. of water, add 1 cc. of pure acetone, and one drop of a 1 per cent aqueous solution of sodium nitroprusside.

If the compound is an aliphatic primary amine and somewhat soluble in water, a violet-red (VR-VRT2) color will develop within one minute. Secondary and tertiary amines and primary aromatic amines should not show this behavior.

These conclusions have been verified in experiments made with primary methyl, ethyl, isopropyl, butyl, isobutyl, isoamyl, hexyl, heptyl and benzyl amines, and pentamethylenediamine, among which ethylenediamine alone gave a color less intense than VRT2; and with dimethyl, diethyl, dipropyl, diisopropyl, diisoamyl, triethyl, and tripropyl amines, aniline, and ammonia, all of which gave no color.

* C. Neuberg. *Festschrift für E. Salkowski, Beitr. wiss. Med. Chem.*, 1904, 20.

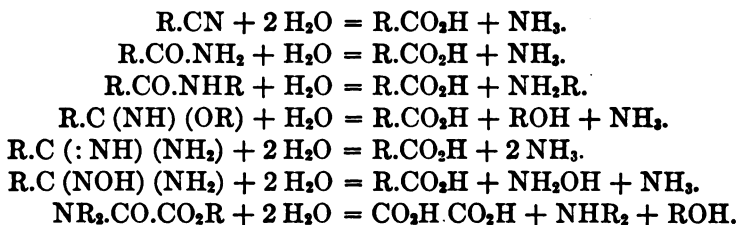
† E. Rimini, *Chem. Zentr.*, 1898 (2), 132.

36 METHOD FOR THE IDENTIFICATION OF PURE ORGANIC COMPOUNDS.

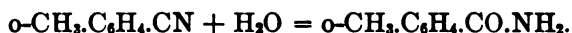
The rapid development of a strong color is greatly favored by the large excess of acetone prescribed. Reduction of the acetone to one drop increases the time required for the development of the color in the case of higher homologues to five or ten minutes. Owing to the delicacy of the test, and the consequent danger that traces of primary amines accidentally present with a secondary or tertiary compound may cause slight colorations, the appearance of no color less saturated than VRT1 should be accepted as evidence indicating the presence of a primary amine. Crude acetone contains acetic aldehyde; and this, in presence of a secondary amine, may develop a blue color and thus interfere with the reaction. The use of this test with mixtures of primary and secondary amines will be more fully discussed under T. 2.28 (Simon's test for secondary amines).

2.26. Saponification of Nitriles, Amides, Anilides, Iminoethers, Amidines, Amidoximes, etc.

The procedure more frequently applied than any other in the analytical examination of species of the natural genera enumerated in the caption is saponification or hydrolysis. An acid, or its salts, and ammonia, or a substituted ammonia, are the normal products when the compound saponified is of simple type; but additional products (*e.g.*, alcohol from a cyano ester) may be formed and require identification when the substance is of mixed type and susceptible to hydrolytic attack at more than one point. Thus:



In using saponification procedures it should be remembered that the velocity of saponification varies widely among species of the same type; that aromatic nitriles having two radicals substituting in the ortho positions about the cyanogen group are probably unaffected by either of the treatments to be described; and that some nitriles simply add water to form amides without evolving ammonia, as in the case of ortho-tolunitrile,



The three sub-procedures of this test, designated by the letters A, B, and C, provide for the identification of the acid products only, and can not be used interchangeably for all saponifiable compounds. Saponification by one of them may be rapid and complete, while by another it may fail entirely. Which method should be used in a particular case will sometimes be found stated in the tables, but more frequently the selection has to be left to the analyst. Procedure D makes the most systematic provision for the conduct of saponifications yielding several products, and will generally be preferred to either of the other procedures for compounds that are readily attacked by normal aqueous alkali. Any difficulty that may have to be overcome in first mastering the technique

of this method will be well repaid by the definite conclusiveness of the solutions that often result from its application to the fundamental problem of determining the intimate molecular structure of saponifiable species.

A.

(SAPONIFICATIONS WITH HYDROCHLORIC ACID.)

This procedure should be usually tried before **B** or **C** when the substance under examination is soluble in hot hydrochloric acid and there is no recommendation in the text or other special reason for giving preference to another. It is less likely to give dark colored decomposition products than the procedure with sulphuric acid, and is much more rapid in its hydrolysis of anilides than the procedure with alkali.

Boil 0.5–1.0 gram of the substance in a small round-bottomed flask fitted with reflux condenser with 10–15 cc. of hydrochloric acid of specific gravity 1.12 for one-half hour or longer.

If one saponification product is an insoluble solid (acid or amide), collect on a small filter and wash by suction with about 5 cc. of water. Recrystallize, dry, and identify the acid or amide. In the case of the simpler aromatic compounds the crystallization may often be best made from 10 to 15 cc. of boiling water.

If the saponification product is an amine hydrochloride (e.g., aniline hydrochloride), or ammonium chloride, evaporate the acid solution to dryness in a small glass dish on the water-bath, transfer the residue to a test-tube, and treat with 5 cc. of 10 per cent sodium hydroxide. Separate the liberated amine by pipetting, filtration, or distillation, according to its physical properties, and identify it by suitable tests. The formation of ammonia may, of course, have been already shown by preliminary Test 2.7.

B.

(SAPONIFICATION WITH SULPHURIC ACID.)

Boil 0.5 gram of the substance for half an hour under reflux as in **A** with 10 cc. of a mixture of equal volumes of concentrated sulphuric acid and water. Dilute with 5 cc. of water, and, if a precipitate forms, cool and filter. The precipitate may often be purified for identification by washing with a little water and recrystallizing from 10 cc. of boiling water.

This procedure is sometimes successful with compounds which do not dissolve in hydrochloric acid of sp. gr. 1.12. It has been found, for example, to give good results with benzanilide, benzoylxylylide and the ortho and para benzotoluides, which are not satisfactorily decomposed by procedure **C**.

C.

(SAPONIFICATION WITH ALCOHOLIC POTASH.)

Boil under reflux on the water-bath for one-half hour or longer 0.3 gram of substance with 10 cc. of 20 per cent alcoholic potash. Evaporate the alcohol. Acidify with 10 cc. dilute hydrochloric acid of 1.12 sp. gr. Add 3 cc. of water. Cool. Shake. Filter. Recrystallize any separated solid product from about 20 to 25 cc. of boiling water, boiling a couple of minutes with 0.1 gram bone-black if the solution should be colored. Filter. Wash. Dry and identify the crystals of acid or amide.

These directions apply more particularly to the examination of difficultly soluble aromatic compounds. The acids from fatty acid nitriles or amides would, of course, often remain dissolved after acidification, and the procedure for their identification would have to be modified accordingly. The fatty acid nitriles and amides are generally easily decomposed by this treatment. It is a very satisfactory method for many aromatic compounds, such as benzonitrile, *p*-tolunitrile, benzamide and β -naphthonitrile. With *o*-tolunitrile and α -naphthonitrile it yields the corresponding amides instead of the acids. As has been indicated, it can not be depended upon to give satisfactory saponifications with anilides and toluides.

D.

ALKALINE SAPONIFICATIONS YIELDING MORE THAN TWO ORGANIC COMPOUNDS WITH PROCEDURES FOR SEPARATION OF THE LATTER.

Many species of Order II that yield ammonia and an acid on saponification give at the same time other products, and it is sometimes desirable to identify

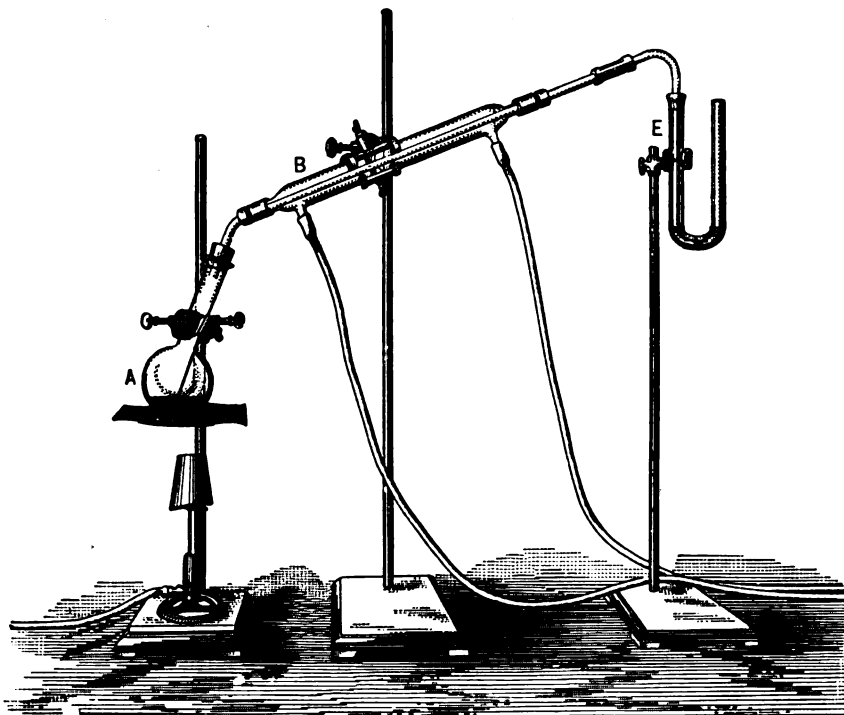


Fig. a.

all of them. It is impracticable to anticipate all cases that may occur, but the following suggestions will point the way to a successful solution for many such problems.

Fit a strong 250-cc. round-bottomed flask, "A," with a clean sound stopper perforated to receive the lower end of the reflux condenser, "B," mounted as shown in Fig. a. The bent inner tube of the condenser should have a total length of 80 cm. and a diameter of 1 cm. Connect it at the upper end with a U-tube containing just enough distilled water (about 10 cc.) to barely seal the bend. Introduce into the flask about 2 grams of the substance to be saponified, and 50 cc. of normal aqueous sodium hydroxide solution. Then drop in a long

ebullator tube (Vol. I, p. 223), to prevent bumping, and boil gently for about two hours while maintaining a circulation of cold water through the condenser jacket.

(a) **Aminonia.** — After having continued the boiling for nearly an hour, disconnect the U-tube, and note whether there is any odor at the end "E." If the odor is strongly ammoniacal, and a strip of red litmus when introduced is promptly blued, close the ends of the tube, and shake. Then withdraw 4 cc. of the solution and apply Test 2.6 for ammonia.

If the odor of the vapors and the color of the precipitate in Test 2.6 do not exactly correspond with those which pure ammonia should give, replace the water removed from the U-tube, and reconnect the latter with the condenser. After the conclusion of the saponification: transfer the contents of the U-tube to a small dish; add a drop of aqueous Congo red solution (1 : 500); titrate with decinormal hydrochloric acid; evaporate to dryness on the water-bath; and dry the saline residue, which may be a mixture of ammonia and amine hydrochlorides, at 100°.

To isolate and identify ammonium chloride in such a mixture, triturate it thoroughly with 3 cc. of chloroform. Decant the clear solution. Repeat the trituration with a fresh 3-cc. portion of chloroform, and again decant. Substituting ether for chloroform, extract twice with this solvent, and then twice again with 3-cc. portions of a cold saturated solution of ammonium chloride in absolute alcohol. Finally, rinse the residue once with 3 cc. of cold absolute alcohol, and determine whether it is really ammonium chloride by heating the dried salt in a glass tube over a small gas flame. *Ammonium chloride sublimes without previously melting or darkening, and is the only hydrochloride of an amine of low boiling-point that shows this behavior;* or tests 2.8, 2.6, determination of the chlorine content, or determination of the platinum and chlorine content of the corresponding chloroplatinate, may be resorted to. These latter tests, however, add very little to the certainty of a conclusion based on the simpler sublimation test.

When 1 decigram of NH₃ is liberated in one of these saponifications, although a little of the gas always remains in the condenser, more than three-quarters will pass on to be absorbed in the U-tube. A characteristic ammonia reaction in T. 2.6 will still be obtained when the saponified mixture contains only 15 milligrams of ammonia, and has been boiled for only half an hour. The fractional distillation and condensation involved in this procedure is in itself enough to free ammonia almost completely from contamination with aromatic and other higher boiling amines; but gives, at the best, only very imperfect separations from amines boiling below 100°. Successful separation from amines of the latter group is, however, generally effected by the successive washings with chloroform, ether and alcoholic ammonium chloride solution. Ammonium chloride is almost entirely insoluble in either of these solvents, while the amine hydrochlorides here concerned, so far as is known, are all soluble in one or more of them. Alcoholic ammonium chloride is used rather than alcohol, because the solubility of ammonium chloride is by no means negligible, 10 grams of absolute alcohol dissolving 0.062 gram at 19°. (Rec. trav. chim., 1892, 156.)

(b) **Volatile Amines.** — Having removed most of the ammonia, and possibly some of the lower aliphatic amines, by the distillation procedure of (a), drop a fresh ebullator tube into the distilling flask, rearrange the apparatus as shown in Fig. b, and distil off 30 cc. of liquid. Note whether the odor of the distillate is ammoniacal, fishy or like aniline, pyridine, or piperidine. If solid particles separate, filter them off for examination. If an oily layer forms, sep-

erate by aid of a thin-stemmed pipette; dry in a very narrow tube over a bit of solid potassium hydroxide; determine the boiling-point, if desired, by Siwoloboff's method (as in (i), p. 115, Vol. I); or use the material for other special identification tests.

If no solid or oily matter separates from the distillate, and the aqueous liquid has a strong basic odor, neutralize it exactly with normal hydrochloric acid, evaporate on the water-bath to obtain the dry hydrochloride; purify this if possible by crystallization from organic solvents as suggested under (a);

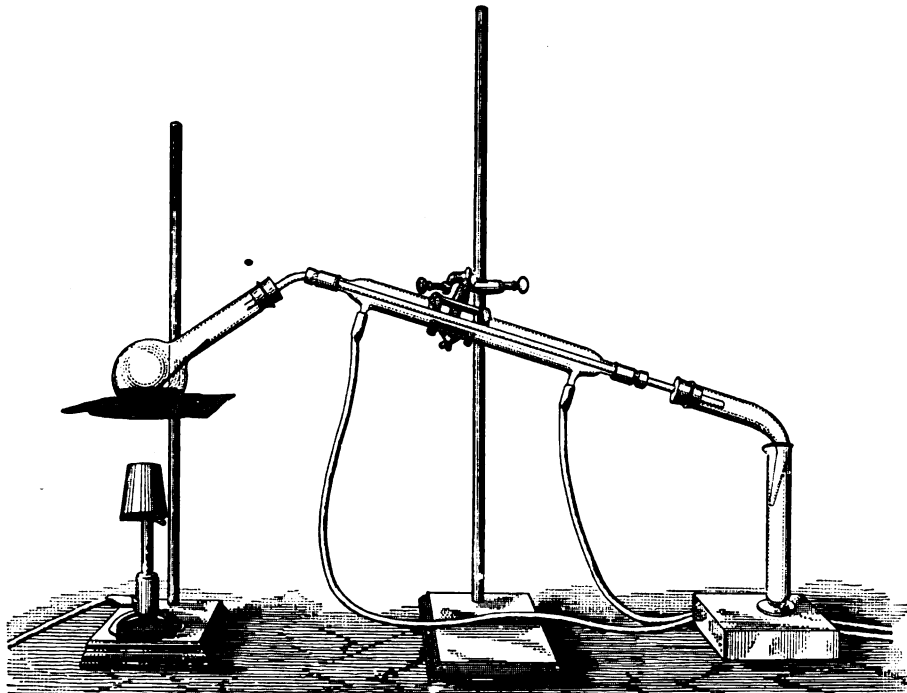


Fig. b.

and then continue to an identification, by conversion to picrate, picrolonate, chloroplatinate, chloroaurate, or by use of special tests suggested by the context of the tables and other circumstances.

(c) **Alcohols (from Esters).** — If there is reason to suspect that the compound saponified may be an ester, the aqueous solutions containing ammonia or amine hydrochlorides obtained in (a) and (b) when the basic distillates are neutralized with hydrochloric acid are to be placed in a 50-cc. distilling flask, and about 35 cc. of liquid distilled off. The distillate will contain any volatile alcohol that may have been formed in the saponification, and should be examined by the procedure described on p. 113 of Vol. I, under the caption "**B (Examination of the Neutral Saponification Products).**"

(d) **Acid Saponification Products and Non-volatile Bases.** — Extract the strongly alkaline solution remaining in the saponification flask after the conclusion of the distillation in (a) or (b) with ether or other appropriate organic solvent, to separate any basic saponification products which are not volatile with steam, taking pains not to lose any of the alkaline solution during the extraction. To isolate and identify the acids in this solution, first determine the exact quantity of normal alkali consumed in saponification by titrating the

entire quantity with normal hydrochloric acid and phenolphthalein. (This will be 50 cc. of normal sodium hydroxide solution less the number of cubic centimeters of normal acid used in the titration.) Then proceed as directed on p. 116 of Vol. I, under the caption "C (Examination of the Acid Saponification Products)."

2.27. The Schotten-Baumann Type of Procedure * for Benzoylating or Acylating Alcohols, Phenols, and Amines.

The principle underlying this important group of procedures for replacing hydrogen in the hydroxyl of alcohols or phenols, or in the amino group of amines by the radicals of organic acids by the reactions $\text{ROH} + \text{R.COCl} = \text{R.CO}_2\text{R} + \text{HCl}$ and $\text{R.NH}_2 + \text{R.COCl} = \text{R.NH.CO.R} + \text{HCl}$ is, that such reactions take place more readily, and may be most conveniently applied to making pure derivatives on the small scale, when carried out in presence of an excess of free caustic alkali or some weakly alkaline substance. The four special procedures (a), (b), (c), and (d), below, provide for most of the more important cases where benzoylation or acylation is suggested in the tables. The original literature of a proposed derivative should be generally consulted, however, since these procedures do not make the best possible provision for all cases.

A.

BENZOYLATION OF ALCOHOLS OR PHENOLS WHICH ARE UNAFFECTED BY CAUSTIC ALKALI.

Place half a gram of the substance with more than the chemically equivalent quantity of 10 to 20 per cent aqueous sodium hydroxide solution and of benzoyl chloride, in a small glass-stoppered bottle, and shake vigorously until the odor of the chloride entirely disappears. This may take as long as fifteen or twenty minutes. There must be alkali enough to leave a strong alkaline reaction at the end of the operation, or somewhat more than enough to decompose the whole of the benzoyl chloride. Unless the benzoyl derivative formed is of acidic character, it may then be directly filtered off and recrystallized or otherwise purified for examination.

Skraup (Monatsh., 10, 390 (1891)) found that very satisfactory results are obtained in preparing benzoates from phenols when 5 molecules of benzoyl chloride and 7 molecules of 10 per cent sodium hydroxide are used for each phenolic hydroxyl group. But so large an excess of the reagent is not always necessary or advantageous.

The mixture heats up considerably during the shaking. This heating hastens the reaction and is often desirable, though cooling is recommended in some cases. When the benzoylated product is acidic and not soluble in cold ether or petroleum ether, it may often be separated from benzoic acid (which will precipitate with it upon acidification with mineral acid) by shaking out with one of these solvents, in either of which benzoic acid dissolves easily.

* For a more complete list and bibliography of such procedures, introduced by Schotten in 1884 (Ber., 17, 2545), see Th. Weil's "Die Methoden d. organ. Chemie," pp. 572, 580, 748, 846, and 1278.

B.**BENZOYLATION OR ACYLATION IN PRESENCE OF PYRIDINE.**

When an alcohol or phenol is sensitive to alkali, or when acylating with an aliphatic acid chloride yields a product that would be rapidly saponified by caustic alkali, Einhorn's procedure (Ann., 301, 95 (1898)) may be substituted for the foregoing with much advantage.

Following this procedure, dissolve the substance in 5 to 10 parts of pyridine, and drop in gradually, cooling, the chemically equivalent quantity of the acid chloride. After 6 to 8 hours, pour into a slight excess of well cooled dilute sulphuric acid, which will cause the separation of the derivative. Ethyl acetate, for example, may thus be prepared even in an acid solution containing acetic acid as well as pyridine.

C.**BENZOYLATION OF AMINES.**

In benzoylating amines two equivalents of the amine and one of benzoyl chloride may be dissolved together in a neutral solvent like benzene and warmed. The excess of amine takes the place of the caustic alkali employed in procedure A for neutralizing the liberated hydrochloric acid. Or, the amine and benzoyl chloride may be shaken with one equivalent of alkali hydroxide in dilute aqueous solution. Or, if the benzoylated product is sensitive to caustic alkali, alkali carbonate, bicarbonate, or acetate may be substituted for the alkali hydroxide. (Cf. Lossen, Ann., 265, 148 (1891); E. Fischer, Ber., 32, 2454 (1899); Claisen, Ber., 27, 3182; Ann., 291, 58 (1896)).

The two latter procedures enable a more complete utilization of the organic material.

D.**BENZOYLATION OF AMINO-ACIDS.**

It has been shown by E. Fischer that while the Schotten-Baumann reaction in its more usual form gives very poor results in benzoylating the amino-acids formed in proteolyses, the pure benzoyl derivatives of these acids may be obtained in excellent yields by conducting the reaction with certain precautions in the presence of large excesses of sodium bicarbonate and benzoyl chloride. (Ber., 32, 2453 (1899); 33, 2370-2373 (1900)). The method is one of much importance to the physiological chemist. It is well illustrated in its application by F. Ehrlich to d-isoleucine (Ber., 37, 1827 (1904)).

"3 grams (1 mol.) of pure d-isoleucine were dissolved in 23 cc. of normal sodium hydroxide (1 mol.) and 60 cc. of water. 11.5 grams of sodium bicarbonate were next added, and then, with constant shaking, in small portions, in the course of 4½ hours, 9.6 grams (6 mol.) of benzoyl chloride. After brief treatment with boneblack the solution was filtered, acidified with sulphuric acid and set aside for a considerable time in the cold. The crystalline mass that soon formed from the emulsion which at first precipitated was drained by suction, washed with cold water and air-dried over night between filter papers. These crystals, consisting of a mixture of the benzoyl compound and benzoic acid, were well shaken with cold benzene, when the benzoic acid passed into solution with the greatest ease. After washing several times with cold benzene, the residue was dissolved in much hot water and the solution boiled up with boneblack and filtered. On cooling the solution yielded an emulsion, from which, after scratching the walls of the container, a mass of long, lustrous colorless needles separated. The yield of pure benzoyl-d-isoleucine obtained for analysis was 3 grams."

2.28. Simon's Test * for Secondary Amines.

To a solution or suspension of one drop of the compound (or an equivalent quantity of a solution) in 3 cc. of water, add 1 cc. of acetic aldehyde solution † and one drop of a 1 per cent aqueous solution of sodium nitroprusside.

Secondary amines that are somewhat soluble in water cause the appearance in this test, in between 30 seconds and 5 minutes, of a blue (B-BT1) coloration, which then slowly changes through greenish blue to a pale yellow.

This test has been verified with very satisfactory positive results in the cases of secondary dimethyl, diethyl, dipropyl, diisobutyl, and benzylmethyl amines; and with equally satisfactory negative results with a series of aliphatic primary and tertiary amines, and aniline, pyridine, and quinoline. Coniine, a secondary heterocyclic amine with its nitrogen atom in the nucleus, gives an orange-red instead of a blue color.

In general, it appears that mixtures of primary and secondary amines which contain only a small quantity of the primary may be expected to give both the Rimini and Simon reactions; but that when the quantity of primary amine is greater, the Simon reaction is not given. Thus we have found that in a mixture of methyl and dimethyl amines containing over 5 per cent of the primary compound, the blue color of the Simon reaction does not make its appearance, while as little as 2 per cent of methylamine causes a considerable decrease in the intensity of the color. On the other hand, the application of Rimini's test for primary amines to the 5 per cent mixture gives a good VRT1 color. Again, working with a mixture of isobutylamine and dipropylamine, it was found that 10 per cent of the primary amine did not suppress the blue color of the Simon reaction, though it considerably weakened it. A mixture of equal parts of these two amines, however, gave a strong Rimini reaction, but no Simon reaction.

2.29. Tastes.

It has been found possible to give a roughly quantitative expression to the terms *bitter*, *sweet*, and *pungent* when used in the descriptions of the tastes of substances, by establishing arbitrary scales of bitterness, sweetness, and pungency analogous to the familiar scale of hardness used in descriptive mineralogy. The method has not been applied to a large number of species, but will be occasionally referred to in the tables. It is based on the use of the following series of standard aqueous solutions, of which only the most concentrated for each scale should be kept on hand.

BITTERNESS SCALE.

No.

4. A solution of one part of quinine sulphate in 1000 parts of water.
3. Dilute one volume of No. 4 to 5 volumes.
2. Dilute one volume of No. 4 to 25 volumes.
1. Dilute one volume of No. 4 to 125 volumes.

* Simon Compt. rend., 125, 536 (1897).

† Acetic Aldehyde Solution.—This special aldehyde solution is readily prepared as required in the following manner:

Wind a piece of rather light copper wire around a lead pencil so that the closely coiled spiral shall form a cylinder 2 cm. in length, while 20 cm. of the wire is left unwound to serve as a handle. Oxidize the spiral superficially by holding it in the upper part of a Bunsen flame; and then, *while still at a red heat*, plunge it into 5 cc. of 50 per cent ethyl alcohol in a six-inch test-tube. Withdraw the spiral *at once*. Cool the test-tube with running water. Repeat the oxidation with the heated wire once or twice, and use the cooled solution at once.

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SWEETNESS SCALE.

No.

5. A solution of 85 grams of cane sugar in water diluted to 100 cc. (This is the "Syrupus" of the United States Pharmacopeia.)
4. Dilute one volume of No. 5 to 3 volumes.
3. Dilute one volume of No. 5 to 9 volumes.
2. Dilute one volume of No. 5 to 27 volumes.
1. Dilute one volume of No. 5 to 81 volumes.

PUNGENCY SCALE.

No.

4. A half normal solution of ammonium hydroxide.
3. Dilute one volume of No. 4 to 2 volumes.
2. Dilute one volume of No. 4 to 4 volumes.
1. Dilute one volume of No. 4 to 8 volumes.

In using these standards in comparative experiments, rinse out the mouth with water, drop 1 cc. of the standard solution on the tongue, bring the tongue in contact with the roof of the mouth, and make careful mental note of the sensation at the end of half a minute. Then, after rinsing out the mouth again with water repeat the experiment after a very short interval with a solution of the substance made up to have the concentration prescribed for that particular compound in the tables. It is impossible to make very satisfactory comparisons when the scale number is as high as No. 4. Comparisons are preferably made at dilutions giving tastes weaker than correspond to No. 3.

Remember in making taste tests that Order II contains many highly poisonous species! Always eject the solution remaining on the tongue at the conclusion of a test, and rinse out the mouth thoroughly with water! Whenever the compound tasted *may* be poisonous, and no specific directions for the tasting test are given, proceed with due caution, and do not bring more than one milligram of substance into the mouth!

An elaborate discussion of the subject of the tastes of organic compounds, and of the known relations between taste and chemical structure, may be found in "Die organischen Gesmacksstoffe," by George Cohn, Berlin, 1914.

2.30. Tollen's Reagent, The Reduction Test with.

In Order II many compounds of the most diverse type reduce Tollen's reagent. The procedure is identical with that given for Test 1.101.

Place 1 to 2 drops, or about 0.05 gram of finely powdered substance, in a clean test-tube with 2 to 3 cc. of the reagent (whose preparation is described with Test 1.101). Shake, and then allow to stand for five minutes. *Never warm!*

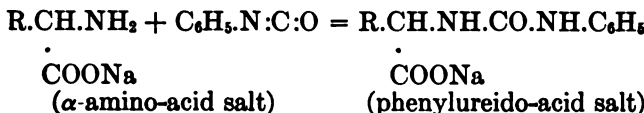
The result of the test is to be considered positive when a black, grayish, or brownish-black precipitate, or an adherent silver mirror, forms.

2.31. Ureido-acids and Hydantoins of the α -Amino-acids.

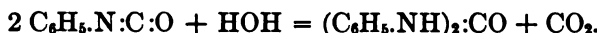
The phenyl and naphthyl ureido acids, and their anhydrides, the hydantoins, have been shown * to be among the more characteristic derivatives of the α -amino-acids.

* Paal, Ber., 27, 530. Mouneyrat, Ber., 33, 2393 (1900). E. Fischer, Ber., 39, 530; Z. physiol. Chem., 83, 151.

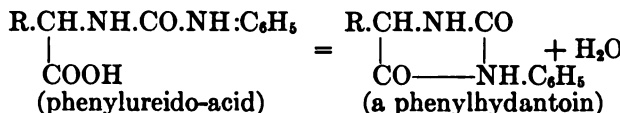
They are not difficult to prepare when the amino-acids used are pure. They crystallize well, and the melting-points of the hydantoins are fairly sharp. The phenylureido acids are addition products of phenyl isocyanate, being formed according to the reaction:



Any excess of isocyanate will react on the water in which the amino salt is dissolved during the reaction to give insoluble carbanilide, which may be removed by filtration.



The free phenylureido-acids yield their hydantoins upon being boiled for a short time with hydrochloric acid of sp. gr. 1.12, and then evaporating to dryness.



Before preparing these derivatives of any particular α -amino-acid, it is best to consult the original papers in which they were first described. The following procedure for preparing the phenylureido acids and hydantoins of amino-acetic acid and its homologues will, however, according to Miss M. J. Ruggles, require only slight modification in special cases.

Dissolve not less than half a gram of the α -amino-acid in slightly more than the chemically equivalent quantity of carefully measured aqueous normal sodium hydroxide solution. Add to the cold solution in a small glass stoppered bottle a quantity of phenyl isocyanate just equivalent to the sodium hydroxide. Shake vigorously with good cooling until the odor of the isocyanate has disappeared. Filter off any carbanilide that may separate. Precipitate the phenylureido acid in the filtrate by the addition of a measured quantity of hydrochloric acid chemically just equivalent to the sodium hydroxide used in dissolving the amino-acid. Filter by suction, and wash with a very little cold water. Recrystallize from the smallest possible quantity of boiling water until the dried product shows a constant melting-point.

To obtain the hydantoin, boil the ureido-acid for fifteen or twenty minutes with eighty parts of hydrochloric acid of sp. gr. 1.12, replacing the evaporated acid if necessary. Then evaporate to complete dryness on a steam-bath. Dissolve the residue in the smallest possible quantity of boiling water. Allow the hydantoin to crystallize out slowly from the cooling solution. Recrystallize until the melting-point is constant.

The hydantoins usually melt more sharply than the ureido acids, and both should be prepared, since some of the compounds in each class are not far separated in their melting-points. In chemical literature the phenylureido-acids and hydantoins are frequently described in the same place with the corresponding amino-acids under the designation of their "phenylisocyanates" or "phenyl-isocyanate compounds," and the hydantoins as the "anhydrides" of the latter.

2.32. Xanthoproteic Reaction.

On a small porcelain crucible lid resting on the hot copper cover of a boiling water-bath, mix a couple of milligrams of the substance to be tested and one drop of nitric acid (sp. gr. 1.42). After one minute add enough sodium hydroxide solution (1 : 10) to produce a strong alkaline reaction (usually two or three drops).

The result of this test is to be considered positive when the nitric acid develops an approximately OY, and the caustic alkali an approximately O, color.

This test, like Millon's reaction, is often applied to protein material and proteolytic products; and, as in the Millon reaction, the colorations are attributable to presence of aromatic complexes which in the xanthoproteic reaction undergo nitration. Substances belonging to other classes may give similar colorations.

2.33. Adamkiewicz-Hopkins-Cole Reaction.

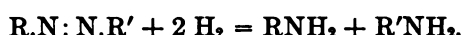
Boil a few milligrams of the substance to be tested with a mixture of two volumes of the Hopkins and Cole glyoxylic acid solution * and one volume of concentrated sulphuric acid.

The result of the test is to be considered as positive when a fine violet-colored solution showing feeble fluorescence is obtained. Suitably diluted, the solution shows an absorption band in its spectrum between the Fraunhofer lines *b* and *F*. In applying the test to albumin solutions, the solution may be mixed with an equal volume of the glyoxylic acid reagent and then treated with concentrated sulphuric acid and warmed.

In the form described, the test is due to Hopkins and Cole (Proc. Roy. Soc. London, **68**, 21 (1901)). It is a reaction for tryptophane, a common protein component.

2.34. Azo and Azoxy Compounds.

Azo compounds contain the grouping N:N; are all colored; reduce ammoniacal silver solution like nitro compounds after the reduction prescribed in Test 2.21; and split upon stronger reduction at the double bond between each pair of azo-nitrogen atoms, giving derivatives containing an amino group in place of each of these atoms.



Test 2.21 should, therefore, always be applied as a preliminary test for azo compounds; and a complete reduction (Test 2.40) with tin and hydrochloric acid, or with sodium hydrosulphite, followed by identification of one or more of the reduction products, is usually the best procedure for concluding the examination.

No cases are recalled in which Test 2.21 gives a negative result when applied to an azo compound of Order II. The complete reduction of compounds having nitro groups ortho to azo groups may, however, lead to the formation of additional anomalous products. Thus, o-nitro-azobenzene gives besides the normal

* Glyoxylic Acid Reagent. — Allow 15 grams of sodium amalgam to act on 250 cc. of a saturated aqueous solution of oxalic acid in a tall glass cylinder as long as hydrogen continues to be evolved. Then filter and dilute with two or three volumes of water.

products, aniline and o-phenylenediamine, the heterocyclic azimidobenzene, $^1\text{N-C}_6\text{H}_4.\text{NPh,}$ ² and its oxide. (J. pr. Chem. (2), **76**, 134.)

All that is said above of the azo compounds is probably equally true of the azoxy compounds, except that they are not necessarily distinctly colored.

2.35. Amines Distinguishing between Primary, Secondary, and Tertiary.

Amines of these three types are derived from ammonia by the substitution of one, two, or three of its hydrogen atoms by hydrocarbon radicals. So many ways have been proposed for determining to which type an amine belongs, that some suggestions to aid in the selection of suitable methods for use in particular cases are desirable.

(a) *Primary Amines.* — The most reliable of all tests for primary amines is the test with nitrous acid (for the NH_2 group), which has been fully described as Test 2.4.

If the amine is probably aliphatic, and somewhat soluble in water, Rimini's test (Test 2.25), which is shorter, may be tried first, or may be sufficient alone.

Too much weight should not be given to the results of the carbylamine test (Test 2.12), or to the mustard oil reaction; since they are so delicate with many amines of low molecular weight that the odors produced may be due to traces of impurities; and since they are liable to fail with primary amines of very high molecular weight. They have a certain value, however, as simple confirmatory tests.

(b) *Secondary Amines.* — It will usually be most satisfactory to test amines which have been found by the above methods *not* to be primary by Hinsberg's method with benzenesulphonyl chloride (Test 2.9), although this procedure is often comparatively long and complicated. Amines which are not primary, but which are somewhat soluble in water, may be tested by Simon's reaction (Test 2.28). Test 2.36 with nitrous acid is most valuable when it happens to give a nitroso derivative whose properties are already known and are described in the tables.

(c) *Tertiary Amines.* — An amine is to be regarded as tertiary if it does not give the reactions of a primary or secondary amine. Its tertiary nature may be most directly determined by the outcome of the Hinsberg reaction (Test 2.9). The iodomethylates are often prepared (Test 2.37) as tertiary amine derivatives.

2.36. Nitrosamines from Secondary Amines.

Nitrosamines, when solid and easily crystallized, are important derivatives in the characterization of the secondary amines. They are easily obtained by the reaction: $\text{R}_2\text{NH} + \text{HNO}_2 = \text{R}_2\text{N}(\text{NO}) + \text{H}_2\text{O}$. The following procedure for their preparation will usually prove successful where more specific directions are lacking.

Prepare a concentrated solution of a few decigrams of the amine in water containing one or two equivalents of hydrochloric or sulphuric acid. Drop in slowly, cooling and shaking, one equivalent of a nearly saturated aqueous solution of sodium nitrite, or enough to produce a distinct blue spot when a drop of the mixture, after standing for two minutes, is brought upon iodo-starch paper. The nitrosamine will usually soon precipitate, or may be extracted by ether, and should be recrystallized.

A single equivalent of hydrochloric or sulphuric acid has been recommended for some of these preparations, but a considerable excess has been employed in other cases without apparently injurious effects. Any considerable excess of nitrous acid is to be avoided, since it is known to cause gradual nitration of some aromatic nitrosamines. Very feebly basic amines which will not give concentrated solutions in dilute aqueous acid (like diphenylamine, No. 2.1568) may sometimes be most successfully treated in alcoholic solution.

2.37. Iodomethylates.

Tertiary amines usually form addition products, iodomethylates, with methyl iodide, according to the equation: $R_3N + MeI = R_3NI$. The reaction sometimes takes place at once with evolution of heat on mixing the compounds, as described for Nos. 2.1356 and 2.1365, but prolonged heating in concentrated methyl alcohol solution, or with an excess of methyl iodide under reflux, is often necessary to ensure combination.

Unless the percentage composition of the amine and its derivative are also determined, the mere fact that an amine reacts with methyl iodide gives no indication as to whether it is a primary, secondary, or tertiary compound; for all three types may react. The preparation of previously described iodomethylates is, however, a method of considerable value in completing the identification of many tertiary amines.

2.38. Methoxyl Group, Determination of.

The methoxyl group in most organic compounds may be quantitatively determined by heating the compound with hydriodic acid, and precipitating the methyl iodide formed by silver nitrate, as silver iodide. The method has been extensively used in the study of the constitution of the alkaloids. For a detailed description of the method see T. Weyl's "Die Methoden d. organischen Chemie," Vol. II, p. 596.

2.39. Picrates, Picrolonates, etc., Use of.

The picrates, and to a lesser extent the picrolonates and addition products of other nitro compounds with amines, alkaloids, hydrocarbons, or phenols, are numerously represented among the colored species of Order II. Since they are most frequently prepared as an aid to the resolution of mixtures, or the purification or identification of substances, the fact that they are picrates or picrolonates will usually be apparent to the chemist who has to deal with them without special testing. To completely identify them, they should be resolved into their proximate components, and the latter separately examined by the usual procedures.

Picrates of amines or alkaloids are to be decomposed by adding a moderate excess of dilute sulphuric acid to their concentrated boiling aqueous solutions or suspensions. If the sulphate of the base formed is readily soluble, the picric acid will separate after cooling in small crystals, or it may be shaken out from the cold solution with benzene or ether, and afterwards purified by recrystallization from hot water containing hydrochloric or sulphuric acid, while the basic component may be liberated from its combination with the sulphuric acid by treatment with alkali. Picrates of the hydrocarbons may in general be decomposed by simply boiling with water in a distilling flask. If the hydrocarbon is

volatile with steam, it may be condensed in the aqueous distillate, the non-volatile picric acid remaining in the flask. Otherwise, the suspension after short boiling should be filtered through a wet filter, and the hydrocarbon freed from all traces of picric acid by washing with hot water. Picric acid is described as No. 2.3168, and the use of picrates as derivatives under Test 2.23.

Picrolonates of nitrogenous bases are decomposed and identified by the same kind of treatment as the picrates. Ether is recommended as the solvent for extracting the acid. Picrolonic acid is described as No. 2.3184, and the use of picrolonates as derivatives under Test No. 2.23.

2.40. Reduction Leading to Amino Compounds.

Reductions of this class are frequently necessary, especially in the identification of nitro and azo compounds, and can be profitably made in many instances where no definite suggestion or directions for such procedure are given in the tables. The purpose of this section is to indicate by special examples how such reductions may be made on a very small scale by a few of the most serviceable methods.

(a) *Reductions with Tin and Hydrochloric Acid.* — The familiar method of warming with an excess of tin and hydrochloric acid is a very general one, reducing all nitro groups to amino groups, and splitting each azo group so as to give two amino groups, as shown by the equation under Test 2.34. Its use and advantages in examining azo dyestuffs are discussed by Witt (Ber., 21 (1888), 3471). In the cases described in the following examples, the quantity of organic compound taken for an experiment varies from one drop to half a gram. Nos. 2.1562 and 2.1651 illustrate its use when the amine formed is separated from tin salts by alkali and ether; Nos. 2.164 and 2.139, when the tin is removed by hydrogen sulphide.

(b) *Reductions with Sodium Hydrosulphite ($\text{Na}_2\text{S}_2\text{O}_4$).* — This method is usually preferable to the foregoing when azo compounds (cf. Test 2.34) are being examined, being very rapid, and furnishing very pure reduction products when one of them is insoluble in water or volatile with steam. It is carried out by slowly adding dry sodium hydrosulphite in small portions to a boiling concentrated solution or suspension of the azo compound in water, dilute alkali, or dilute alcohol, until all color is discharged. One or more of the reduction products may then separate out, or, if volatile, may be distilled off. Its general use is illustrated in Nos. 2.3187 and 2.3349, and is thoroughly discussed by E. Grandmougin (J. prak. Chem., 76 (1907), 124), and its use in dyestuff identification under Test 3.21. It is probably applicable to many nitro and nitroso compounds, though comparatively few cases belonging to this class have been recorded.

(c) *Reductions with Hydrogen Sulphide and Ammonia.* — This mild reduction method is exemplified under No. 2.3016 (dinitrobenzene), and is occasionally advantageous when it is desired to reduce only a single nitro group in a polynitro compound.

2.41. Konowalow's Reaction.

According to Konowalow (Ber., 28 (1895), 1850), colorations are given by primary and secondary, but not by tertiary nitro compounds when treated as follows: Shake a little of the compound several minutes with a slight excess of

potassium hydroxide solution, or for a moment with sodium alcoholate. Extract with a little water. Add ether to the solution, and then ferric chloride until the ether becomes colored. Nitroethane gives a red color. Nitropropane gives first a red, and with more ferric chloride, a green and blue.

2.42. Pettenkofer's Test for Gall Acids.*

Add concentrated sulphuric acid gradually to a little of the gall acid, or its aqueous solution or suspension in a test-tube, warming or cooling so as to keep the temperature between 60° and 70° until any precipitate formed dissolves in the excess of acid. Stir into the solution a 10 per cent cane-sugar solution, added a drop at a time. A fine cherry, changing later to a persistent permanaganate red color showing a characteristic spectrum (cf. No. 2.119), appears. The test being a furfural reaction,† may also be made by adding 1 drop of 0.1 per cent furfural solution to 1 cc. of an alcoholic solution of the gall acid, and then mixing, with cooling, with 1 cc. of concentrated sulphuric acid.

Presence of albuminous matter or oxidizing agents disturbs the reaction. Albumin and some organic compounds may also give somewhat similar purple colorations with sulphuric acid, it is said.

2.43. Hydrazones and Semicarbazones.

As a preliminary reaction for hydrazones and semicarbazones, apply Test 2.17. In cases in which the carbonyl derivatives formed by this method of hydrolysis are stable in the presence of hot acid, it will often be desirable to repeat the hydrolysis on a somewhat larger scale than is recommended in this test, in order that an attempt to isolate and identify them may be made.

* Pettenkofer, Ann., 52, 1844, 90.

† Mylius, Z. physiol. Chem., 2 (1887), 492.

SUBORDER I OF ORDER II.
COLORLESS COMPOUNDS CONTAINING C, N, H, AND O.

GENUS I, ACIDIC COMPOUNDS.

DIVISION A, SOLID SPECIES.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	SPECIFIC NAMES. — Tests and Miscellaneous Properties.
1	-21	"Reacts acid"	Propylnitramine, $\text{Pr.NH}(\text{NO}_2)_2$. — \textcircled{P} Gives color react. described under No. 2.6 — D.s. aq.; v.s. alc., eth. Sp. gr. 1.103 (15 $^{\circ}$). B.p. 128 $^{\circ}$ (40 mm.).
2	0	"Reacts acid"	Butylnitramine, $\text{Bu.NH}(\text{NO}_2)_2$. — \textcircled{P} Gives color react. described under No. 2.6 — Sp. gr. 1.066 (15 $^{\circ}$). D.s. aq.; e.s. alc. — BaA_2 , ppt.; lust. lft. fr. h. aq.
3	6	"Strongly acid"	Ethylnitramine, $\text{Et.NH}(\text{NO}_2)_2$. — \textcircled{P} Gives color react. described under No. 2.6. — Sp. gr. 1.167 (15 $^{\circ}$). — BaA_2 melts at 228 $^{\circ}$ and like other salts explodes at higher temperatures.
4	5.5-6.5		Hexylnitramine, $\text{C}_6\text{H}_{13}\text{NH}(\text{NO}_2)_2$. — V.d.s. aq.; misc. alc., eth.
5	15	$\mu(32) = 327$	Trinitromethane, Nitroform, $\text{CH}(\text{NO}_2)_3$. — Colorless cryst. e.s. aq. w. intense yel. color! Color discharged by large x.s. conc. HCl. (Cryst. yel. unless v. dry.) The acid and many of its salts explode on rapid heating. Salts unstable, even when solid, changing to nitrates.
6	38	$k \cdot 10^7 = 7$	Methylnitramine, $\text{Me.NH}(\text{NO}_2)_2$. — \textcircled{P} Dissolve 0.05 g. substance + 0.05 g. α -naphthylamine in 5 cc. gl. ac. ac. Add abt. 1 g. zinc. An intense V-R-R color quickly develops. — Flat ndl. fr. eth. V.s. aq., alc., bz., chlf. — Odorless. Taste v. sour. Gives nearly theoretical value for N. Eq. on titration. — Dissolve 0.1 g. substance + 0.1 g. NaNO_2 , each in 1 cc. aq. Mix the solutions. Brisk evolution of nitrogen occurs. Warm. Peculiar nauseous odor sl. suggestive of carbylaniline will be noticed.
—	38		Diacetanilide. — Cf. No. 2.1500.
8	abt. 45	113	γ -Cyanobutyric Ac., $\text{CN.CH}_2(\text{CH}_2)_2\text{COOH}$. — Deliq. lft. v.s. aq., alc., eth. — Saponify to glutaric ac. by boiling w. dil. aq. NaOH! (T. 2.26 & 1.316.)
9	45		Ethylparabanic Ac., $\text{C}_6\text{H}_4\text{O}_2\text{N}_2$. — Ndl. e.s. aq.; s. alc., eth. — Saponification yields NH_3 (T. 2.6); EtNH_2 (No. 2.1062); CO_2 and oxalic ac.
10	46-6.5	$k \cdot 10^8 = 2.3$	Phenylnitramine, $\text{Ph.NH}(\text{NO}_2)_2$. — \textcircled{P} Rapid heating explodes. — Pearly lft. fr. lgr. mod. s. c. aq.; v.s. alc.; d.s. lgr. — At 97-8 $^{\circ}$ dec. to o- & p-nitraniline, o- & p-nitrophenol, nitrobenzene, HNO_2 , CO_2 & N. — Stable towards 10% boiling NaOH. — $\text{BaA}_2 \cdot 2\text{Aq.}$, lust. tbl. — AgA , ppt., ndl. fr. h. aq.
II	47	abt. $\frac{3}{4}$ theoret. N. Eq.	Phenylcyanamide, Cyananilide, $\text{Ph.NH}(\text{CN}) + \frac{1}{2}\text{H}_2\text{O}$. — D.s. aq. Evaporation of sol. leaves syrup which becomes cryst. after long contact w. alc. Ndl. fr. eth. — Aq. added to alc. sol. is followed by separation of phenylurea. — Polymerizes in time, quickly on water-bath, to triphenylisomelamin, ndl. fr. alc., m.p. 185 $^{\circ}$, v.d.s. h. aq.

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
I2	49-50		Diethylparabanic Ac., $C_7H_{10}O_2N_2$. — Ndl. e.s. acetone. — E. saponified to $EtNH_2$ (No. 2.1062), oxalic ac., etc.
I3	49.5		Isonitrosomethylbutylketone, $Me.CO.C(NOH).CH_2.CH_3.Me$. — Lust. lft. sol. (probably w. yel. color) in dil. NaOH.
I4	50		Methylethylioxazolone, $C_6H_8O_2N$. — Ndl. s. h. aq. w. strong acid react.; e.s. Na_2CO_3 sol. Alc. sol. red-brown w. $FeCl_3$. — $Ba(C_4H_6O_2N)_2 \cdot 5\frac{1}{2} H_2O$. — Aq., ndl. s. aq. w. "neutral reaction."
I5	50		Cyanoacetylacetone, $CH(CN).(CO.Me)_2$. — Lft. fr. dil. alc.; i. aq.
	51		Nitrosoacetanilide. — Cf. No. 2.1541.
I7	52-3		Malonyldiethylurea, Diethylbarbituric Ac., $C_8H_{12}O_2N_2$. — Rhomb. cryst. fr. h. bz. + pet-eth., s. h. aq. — Saponification gives $EtNH_2$ (No. 2.1062), malonic ac. (Vol. II), etc.
I8	53-5	$k.10^{10} = 3.1$	Isonitrosomethylpropylketone, $Me.CO.C(NOH).Et$. — \textcircled{P} E.s. alk. w. yel. color. — D.s. aq.; e.s. alc., eth.
I9	56	$k.10^6 = 8.6$	Ethyl Isonitrosoacetacetate, $Me.CO.C(NOH).CO.Et$. — \textcircled{P} Soluble in alk. w. yel. color. — Pr. f. chlf. v.s. alc.; eth.
—			Trinitroethane, $Me.C(NO_2)_2$. — Cf. No. 2.1573-1.
I9-I	56-8		Isonitrosomethylnonylketone, $Me.C(NOH).C_9H_{17}$. — Lust. cryst. fr. alc. S. alk. w. yel. color.
22	57	141	Diethylcyanoacetic Ac., $Et_2C(CN).CO_2H$. — Boils undec. — At 160° w. conc. HCl gives diethylacetic ac.
23	59-60	151 $k.10^6 = 4.2$	β -Anilinopropionic Ac., Phenyl- β -alanine, $PhNH.(CH_2)_2.CO_2H$. — Lust. lft. fr. chlf. + lgr.
23-I	abt. 63d.		1'-Nitromesitylene (iso form), $Me_2C_6H_4.CH:NO.OH$. — S. in Na_2CO_3 sol. & repprd. by dil. H_2SO_4 . Silky ndl. fr. bz. E.s. alc., eth., bz.; v.d.s. lgr. Stable in cold only, isomerizing easily to true nitro isomer (No. 2.1527).
24	63		Phenylbenzoylglycine, $PhN(Ph.CO).CH_2.CO_2H$. — Amorph. V.s. alc., eth., chlf., bz. Saponify, and identify the benzoic ac. (T. 1.312).
25	64	Alk. sol. colored	Methylnitrolic Ac., $HC(NOH)(NO_2)$. — \textcircled{P} Caustic alk. dissolves to deep orange sol. — Ndl. e.s. aq., eth. — V. unstable. Boiled w. dil. H_2SO_4 gives N_2O and formic ac.
26	64	154	Nitrourethane, $NO_2.NH.CO.Et$. — Lft. fr. lgr. e.s. aq. Warming w. aniline gives phenylurethane (No. 2.1544). ($k.10^6 = 4.8$)
27	65	$k.10^6 = 3.8$ †	Isonitrosoacetone, $Me.CO.CH(NOH)$. — \textcircled{P} Sol. in 0.1 N $NaOH$, pale yel-brown gradually deepening. — Silvery lft. or pr. s. aq., eth. E. vol. w. st. — Ammon. $AgNO_3$ gives gold yel. ppt., $AgC_6H_4NO_2$.
28	66	Alk. sol. colored	Propynitrolic Ac., $EtC(NOH)(NO_2)$. — Alkalies give red sol. — Ndl. fr. pet-eth. V.s. aq.; sol. tastes sweet and pungent.
29	66-7	119 $k.10^6 = 1.6$	β -Nitropropionic Ac., $(NO_2)CH_2.CH_2.CO_2H$. — Scales fr. chlf., v.s. aq., alc., eth.
30	68	Alk. sol. colored	Allylnitrolic Ac., $CH_2:CH.C(NOH)(NO_2)$. — \textcircled{P} Dil. $NaOH$ gives red sol. — Ndl. fr. eth. — Explodes at 95°.
31	68-9	Alk. sol. blue †	α -Nitrophenyllacticacidketone, $NO_2.C_6H_4.CH(OH).CH_2.CO_2H$. — \textcircled{P} Sol. becomes indigo-blue in Gen. T. 2.II! — Pr. mod. s. h. aq., e.s. alc., eth., chlf.; i. lgr.
33	69-70	85 $k.10^6 = 3.7$	Cyanoacetic Ac., $CN.CH_2.CO_2H$. — E.s. aq. — Action of heat gives methyl cyanide (No. 2.2676) and CO_2 . — $Ag\ddot{A}$, yel. unstable ppt.
34	72	155.5 †	Citric-acid-p-phenetidide, $EtO.C_6H_4.NH.CO.C_6H_3O(CO_2H)_2$. — Cryst. fr. aq.; mod. s. h. aq. (D.R.P. 87,428.)
35	74	$k.10^{10} = 1.3$	Isonitrosomethylethylketone, $Me.CO.CMe(NOH)$. — \textcircled{P} Sol. in alk. yel.

(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
36	74 u.c.	Alk. sol. yellow	8-(o), or Bz-1-Hydroxyquinoline, Quinophenol, $\text{HO.C}_6\text{H}_4.\text{C}_6\text{H}_3\text{N}$. — \textcircled{P} Add 1 drop 10% FeCl_3 sol. to 10 cc. c. sat. aq. sol. Intense green color (approx. GS1-BGS1) develops at once. — Long lust. ndl. fr. h. alc. sol. on cooling. V.d.s. in c. aq.; sol. yellow w. aromatic-phenolic taste and odor. E.s. alc.; d.s. eth.; e.s. bz. and dil. NaOH. Solution in alkali or acids, yellow. — B.p. 267° c. — Vol. w. st. — Dissolve 0.05 g. hydroxyquinoline in 5 cc. aq. to which 3 drops HCl (sp. gr. 1.12) have been added, warming if necessary. Treat the c. sol. w. a slight x.s. Br-Aq. (8 cc.—10 cc.). The ppt. at first pale yel. (YT2), becomes (Y) as more Br. is added. Add aqueous sulphurous acid sol. until the sol. is colorless and the ppt. YT2-YT3. Allow to settle. Decant most of the aq. Filter. Wash w. 5 cc. aq. Dry on porous tile. Cryst. fr. 3 cc. h. benzene, cooling well to separate crystals. Dry 15 min. at 100°. Dibromoxyquinoline, the product in this test, forms nearly colorless ndl. which melt without decomposition at 195.1–6.1°(u.c.), 199.4–202.4°(c), (above 175° the bath should be rapidly heated).
37	74	"Decomposes carbonates"	α -Cyanobenzoylacetone, $\text{Me.CO.CH(CN).CO.C}_6\text{H}_4$. — Long ndl. fr. alc. E.s. alc., eth., bz., alk. — $\text{Ag}\ddot{\text{A}}$, cryst. ppt.
38	77–8	Alk. sol. yellow	Isonitrosoacetylacetone, $\text{Me.CO.C(NOH).CO.Me}$. — Ndl. or pearly scales fr. acetic eth.; i. lgr.
39	77–8	152 $k.10^4 = 6.5$	β -Nitrosobenzylhydroxylamine, $\text{Ph.CH}_2.\text{N}(\text{NO})(\text{OH})$. — Pr. fr. eth. + lgr. V.d.s. aq., e.s. alc., eth., alk. — $\text{Ag}\ddot{\text{A}}$, ppt., m.p. 108–9° d.
—	78		Diacetamide. — Cf. No. 2.1710.
—	78–9		Triacetamide. — Cf. No. 2.1711.
—	82 u.c.		2,4,6-Trinitrotoluene. — Cf. No. 2.1733 (0.1-N NaOH in Gen. T. 2. I. gives faint reddish coloration, appearing slowly).
41	abt. 82–3	225	Opionic-acid-oxime, $(\text{MeO})_2\text{C}_6\text{H}_3(\text{CH}:\text{NOH}).\text{CO}_2\text{H}$. — Ndl. e.s. alc., warm aq., eth., bz. — Prolonged heating at 80° gives hemipinimide.
42	82.5–84	Alk. sol. blue-violet	α -Nitrobenzyl Cyanide, $\text{NO}_2\text{C}_6\text{H}_4.\text{CH}_2\text{CN}$. — \textcircled{P} Sol. becomes blue-violet on addition of little NaOH! Flat lust. ndl. fr. aq. Mod. s. h. aq. — Sap. w. HCl (Cf. T. 2.26, A) gives o-nitrophenylacetic ac.
43	84	193 $k.10^4 = 5.3$	α -Toluidinobutyric Ac., $\text{C}_7\text{H}_5\text{NH.CHEt.CO}_2\text{H}$. — Pr. fr. lgr. d.s. c. chlf., bz., or lgr.
43-I	85	$k.10^4 = 3$	Propionhydroxamic Ac., $\text{Et.C}(:\text{NOH}).\text{OH}$. — \textcircled{P} Sol. gives red color w. FeCl_3 . — Pr. fr. alc. V.s. aq., alc.; i. eth.
44	84.5		m-Aminohydrocinnamic Ac., $\text{NH}_2\text{C}_6\text{H}_4.\text{C}_6\text{H}_3\text{CO}_2\text{H}$. — Cryst. fr. aq. E.s. aq., alc., eth. — B.HCl , scales e.s. aq.
—	85		Acetacetanilide. — Cf. No. 2.1751. \textcircled{P} Aq. sol. dark violet w. FeCl_3 .
—	86		Dinitromesitylene. — Cf. No. 2.1761.
47	86	179 $k.10^4 = 2.2$	β -(p)-Toluidinopropionic Ac., $\text{C}_7\text{H}_5\text{NH}(\text{CH}_2)_2.\text{CO}_2\text{H}$. — Pearly scales fr. bz. D.s. bz., lgr.; s. in 33 pts. c. aq.
48	87		Isonitrosocyanobutyric Ac., $(\text{HON})(\text{CN})\text{C}(\text{CH}_2)_2.\text{CO}_2\text{H}$. — \textcircled{P} Warmed 1–2 min. w. little conc. H_2SO_4 at 60° & then cautiously treated w. aq. gives blue color. — Pr. fr. eth. + bz. V.s. aq.
48-I	87–8	$k.10^4 = 2.8$	Acethydroxamic Ac., $\text{Me.C}(:\text{NOH}).\text{OH}$. — \textcircled{P} Reduces ammon. AgNO_3 sol. FeCl_3 colors sol. dark cherry-red! — V.s. aq., alc.; i. eth. ("Reacts neutral").
49	86–8d.	Alk. sol. colored $k.10^4 = 6$	Ethylnitrolic Ac., $\text{MeC}(\text{NOH})(\text{NO}_2)$. — \textcircled{P} Dissolves w. O-YO color in aq. NaOH. — Lust. pale yel. orthorhomb. cryst. fr. aq. or alc. Taste v. sweet. Unstable, decg. to acetic ac. & N oxides on keeping. The red potassium salt is easily exploded.

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
49-I	89		Diacethydroxamic Ac., $\text{Me.C(OH)(NO.CO.Me)}$. — Ndl. V.s. aq. Dec. easily to ac. ac. & acethydroxamic ac. — Dil. aq. sol. on standing w. $\frac{1}{2}$ mol. Na_2CO_3 gives <i>s</i> -dimethylurea (No. 2.1869).
50	abt. 90d.	"V. weak acid"	(Iso-) Nitrodiphenylmethane, $\text{PhC:NO}_2\text{H}$. — \oplus Gives Konowalow's react. (T. 2.41). — Pr. fr. eth. V. unstable, changing even at room temp. to its isomer, benzophenone & benzoephoneoxime.
51	90	†	Acetobenzyl Cyanide, Me.CO.CHPh(CN) . — \oplus Gives pale moss-green coloration w. FeCl_3 . — Cryst. fr. dil. alc., e.s. alc., eth. Vol. w. st.
52	92	111	α -Cyanocrotonic Ac., $\text{MeCH:C(CN).CO}_2\text{H}$. — Deliq. ndl. e.s. aq. Heating w. aq. KOH gives malonic ac. (Vol. I).
53	92	†	Antipyrine Salicylate, Salipyrine, $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O} + \text{C}_7\text{H}_4\text{O}_3$. — \oplus Gives dark fiery-red fluid, changing to violet & blue, when warmed w. HNO_3 (sp. gr. 1.185), while a black greasy substance appears in the sol.! Cryst. powd. s. in 200 pts. aq. at 15°, or in 25 pts. at 100°. V.s. chlf. less s. eth. — Cf. Flückiger for numerous reactions.
54	94		δ -Benzoylaminovaleric Ac., $\text{Ph.CO.NH(CH}_2)_4\text{CO}_2\text{H}$. — \oplus Remelts after fusion and solidification at 104–5°. E.s. dil. Na_2CO_3 sol.; alc.; d.s. eth. — Boiled w. acetic anhydride gives an anhydride, pearly lsf. fr. alc., m.p. 112°.
55	95–6	131 $k.10^4 = 2.7$	γ -Oximinovalerianic Ac. (Oxime of Levulinic Ac.), $\text{Me.C(NOH)-(CH}_2)_2\text{CO}_2\text{H}$. — Long thick pr. v.s. aq., less s. alc., eth. — Boiling w. dil. HNO_3 readily oxidizes to succinic (Cf. Vol. I) & acetic acids.
56	96	Alk. sol. colored	3,4,6-Trinitrophenol, $(\text{NO}_2)_3\text{C}_6\text{H}(\text{OH})$. — \oplus Taste v. bitter! "Colorless" (?) ndl. or scales fr. alc., s.c. aq., mod. s. h., v.s. alc., eth., bz. — Warmed w. fuming HNO_3 gives styphnic ac. — $\text{K}\ddot{\text{A}}$, bright red.
57	99–100		Hexahydro-p-dimethylaminobenzoic Ac., $\text{Me}_2\text{N.C}_6\text{H}_{10}\text{CO}_2\text{H}-2\frac{1}{2}\text{H}_2\text{O}$. — Tbl. fr. alc. + chlf. Solidifies after melting at 130° & remelts at 218–20°. E.s. aq., alc.; i. chlf., lgr. — $\text{B}_2\text{H}_2\text{PtCl}_6$, yel-red tbl., m.p. 232°.
58	99–101	145	Diethyloxamic Ac., $\text{Et}_2\text{N.CO.CO(OH)}$. — Pr. e.s. aq., alc. Boiled w. NaOH sol. gives Et_2NH (No. 2.1058–1).
59	102		ab-Phenylmethylureidoacetic Ac., $\text{PhNH.CO.NMe.CH}_2\text{CO}_2\text{H}$. — Ndl. fr. aq., e.s. h. aq., alc.; alm. i. eth., chlf. (From sarcosine + phenyl isocyanate.)
60	103	201	2,6-Dimethyl-3-oximinoctanoic(8) Ac., Menthoximic Ac., $\text{Me}_2\text{CH.C(NOH).(CH}_2)_2\text{CHMe.CH}_2\text{CO}_2\text{H}$. — Cryst. s. in 24 pt. c. aq.; e.s. alc., eth. — $\text{Ag}\ddot{\text{A}}$, floe ppt. — Probably gives T. 2.17.
61	104–6u.c.		Benzoylleucine, $\text{Me}_2\text{CH.CH}_2\text{CH}(\text{C}_6\text{H}_5\text{CO.NH}).\text{CO}_2\text{H}$. — $[\alpha]_D^{20} = +6.59^\circ$ (1.0463 g. in 6 cc. normal NaOH sol.) for deriv. fr. l-leucin; and -6.44° ($p = 9.57$) for the optical antipode.
62	108	206	Levulinic-acid-phenylhydrazone, $\text{PhNH.N:CM}_2\text{(CH}_2)_2\text{CO}_2\text{H}$. — \oplus Gives Ag. ppt. in T. 2.17-a. — Pr. fr. bz. E.s. h. aq., h. bz., eth.
63	109 w. efferv.	$k.10^4 = 7.7$	Aminomalonic Ac., $\text{NH}_2\text{CH}(\text{CO}_2\text{H})_2$. — \oplus In melting evolves CO_2 (cf. T. 1.303), leaving residue of glycine (No. 2.2568)! Same result when aq. sol. is boiled. — Crysts. w. 1 H_2O (lost in desiccator). S.c. aq.; d.s. alc.
64	109–10	131	Propyloxamic Ac., $\text{PrNH.CO.CO}_2\text{H}$. — Ndl. e.s. aq., alc., eth. — Sap. gives propylamine (No. 2.1067) & oxalic ac. (Vol. I).
65	110		α -p-Toluidinoisovaleric Ac., $\text{C}_6\text{H}_5\text{NH.CH}(\text{CO}_2\text{H})\text{CHMe}_2$. — Cryst. fr. 25% alc.; e.s. alc., eth., chlf., lgr., h. aq.
66	111		3-Methylpyridinecarbonic(2) Ac., $\text{Me.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Pr. fr. h. alc. V.s. aq.

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
67	111.5	161	Nitrocaproic Ac., $C_6H_{10}(NO_2)CO_2H$. — \textcircled{P} Add KNO_2 to sol. of $K\bar{A}$ & acidify w. H_2SO_4 . Sol. becomes blue, color passing into ether on shaking w. latter! ("Delicate"). — Should give T. 2.21. — 4 sided pr. e.s. aq., alc. Volat. below 100°.
68	112	151 $k \cdot 10^4 = 8$	m-Ethylaminobenzoic Ac., $EtNH \cdot C_6H_4 \cdot CO_2H$. — Pr. d.s. h. aq.; misc. alc., eth. Alm. tasteless. — $Ba\bar{A}_2 \cdot 2H_2O$, v.s. c. aq.
69	113		p-Hydroxybenzonitrile, $HO \cdot C_6H_4 \cdot CN$. — Rhomb. tbl. D.s. c. aq.; e.s. alc., eth., chlf. — Sap. (T. 2.26) to obtain p-hydroxybenzoic ac.
70	113-4	$k \cdot 10^4 = 3.6$	Biliverdic Ac. (Imide of tribasic Hematinic Ac.), $CO_2H \cdot C_6H_4 \cdot (CO)_2 \cdot NH$. — Ndl. fr. aq.; pr. fr. alc. 100 pt. c. aq. dis. 4 pts.; 100 pts. eth., 6 pts. — Reduces c. alk. $KMnO_4$ immediately.
71	113	"Alk. sol. intensely yel."	Isonitrosomethylphenylketone, $Me \cdot C(NOH) \cdot CO \cdot Ph$. — Ndl.
72	114 u.c.	$Alk. sol. yel.$ $k \cdot 10^4 = 6.4$	p-Nitrophenol, $NO_2 \cdot C_6H_4 \cdot OH$. — \textcircled{P} 5 drops 3% $FeCl_3$ sol. added to 5 cc. said. aq. sol. of compound gives VRS_2-RS_2 color. 0.03 g. comp. dissolved in 10 cc. 10% aq. $NaOH$ gives $YT1$ sol. — Colorless pr. fr. h. aq. or h. conc. HCl . D.s. c. aq., e.s. h. aq.; v.s. eth. B.p. 275° c. $\pm 5^\circ$ d. Not vol. w. st. (dif. fr. ortho). Odor, faint aromatic. Taste of said. aq. sol. (T. 2.29) at first sweet (No. 2 of scale), then pungent (No. 2). — \textcircled{P} Place 0.2 g. of the substance w. 2 cc. 10% $CaCl_2$ sol. and 0.5 g. zinc dust in a 25 cc. flask. Connect w. a return-flow condenser and boil (abt. 5 min.) until colorless. Filter hot. Cool filtrate by shaking under running water. Filter. Wash ppt. w. 2 cc. cold aq. Remove to a porous tile. When nearly dry dissolve in 15 cc. boiling benzene. Filter. Cool under running water. Collect the ppt. on a very small filter. Dry on tile at 75° for 15 min. The product, p-amidophenol, cryst. from the benzene in imperfect prs. of a "broken" YO color, and melts w. decn. at 184° (u.c.) after becoming nearly black at abt. 175° . It crysts. in lfl. fr. aq., but the crysts. are liable to be v. dark colored through oxidation.
73	114	96	4-Nitrophthalic Anhydride, $NO_2 \cdot C_6H_4 \cdot (CO)_2 \cdot O$. — Subl. D.s. c. aq.; e.s. eth. Gives corresponding acid (m.p. 161°) on evapn. w. aq.
74	115	235	d, l-Benzoylisoleucine, $C_6H_7 \cdot CH(NH \cdot CO \cdot Ph) \cdot CO_2H$.
75	114; 116		p-Nitrobenzyl Cyanide, $NO_2 \cdot C_6H_4 \cdot CH_2CN$. — \textcircled{P} Alc. KOH gives intense carmine red sol.! Consumes alm. no alk. in Gen. T. 2.I, but suspended powder turns yel., or., and brown.
76	116-7	235	Benzoyl-d-isoleucine, $C_6H_7 \cdot CH(NH \cdot CO \cdot Ph) \cdot CO_2H$. — Lust. ndl. v.d.s. c. aq.; more s. h.; v.s. alc., eth.; e.s. h. bz.; d.s. lgr. — Sinters at 114° . — $[\alpha]D^{20} = +26.36^\circ$ (1.4612 g. in 9 cc. N. NaOH dild. to 19.654 g.).
77	117	Alk. sol. colored	2,3,6-Trinitrophenol, $(NO_2)_3C_6H_2(OH)$. — "Colorless" ndl. d.s. c. aq.; mod. s. h. aq. — Boiled w. fuming HNO_3 gives styphnic ac. — $K\bar{A}$, bright red ndl., v.s. aq.; alm. i. alc.
—	118		Trinitromesitylene. — Cf. No. 2.2560.
79	118	221 $k \cdot 10^4 = 1.1$	α -Acetylanilinobutyric Ac., $(C_6H_5O)NPh \cdot CHEt \cdot CO_2H$. — Pr. fr. h. bz.; d.s. lgr.
80	118	179 $k \cdot 10^4 = 3.9$	α -(o)-Toluidinopropionic Ac., $(C_6H_5NH)CHMe \cdot CO_2H$. — Ndl. e.s. aq., alc., eth.
80-I	118-9		1-Phenyl-5-pyrazolone, $[NPh \cdot N:CH \cdot CH_2 \cdot CO]$. — \textcircled{P} $FeCl_3$ gives blue ppt. w. h. aq. sol. — Cryst. e.s. acids or alkalies. — $B \cdot HCl$. ndl. v.s. aq.; m.p. 165°.
81	119	Alk. sol. yellow	β -Isonitrosolevulinic Ac., $MeCO \cdot C(NOH) \cdot CH_2 \cdot CO_2H$. — E.s. aq., alc., eth. — Boiled w. dil. H_2SO_4 gives diacetyl (Cf. Vol. I).

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
82	119	269	Diphenylsuccinamic Ac., $\text{Ph}_2\text{N.CO.(CH}_2)_2\text{CO}_2\text{H}$. — Lust.tbl. fr. 1 pt. alc. + 3 pt. aq. V.s. alc., bz.; less s. eth. — Ag \bar{A} , floc. ppt., soon becoming cryst.
83	119-20d.	250	d- α -Phenylureido- β -methyl- β -ethylpropionic Ac., $\text{MeEtCH-CH(NH.CO.NHPH).CO}_2\text{H}$. — (Deriv. of d-isoleucine.) Lft. fr. dil. alc. i. c. aq.; e.s. h. aq.; v.s. alc., eth. $[\alpha]_D^{20} = +14.92^{\circ}$ (in alk. sol.). — Corresponding hydantoin (Cf. T. 2.31), silky ndl. fr. lgr. d.s. aq.; m.p. 78-9° (Ber. 37, 1829).
84	120	145	Ethyloxamic Ac., $\text{EtNH.CO.CO}_2\text{Et}$. — 6-Sided tbl. Sublimes. E.s. aq., alc., eth. — Distil w. NaOH sol. Distillate will contain ethylamine (No. 2.1062); residue, sodium oxalate! This reaction begins in the cold.
85	120	165 $k.10^4 = 1.7$	p-Toluidinoacetic Ac., p-Tolylglycine, $\text{C}_7\text{H}_5\text{NH.CH}_2\text{CO}_2\text{H}$. — Cryst. fr. h. aq. or alc.; d.s. c. eth., chlf., bz. Unstable in air. — Gives Ag mirror w. ammon. AgNO_3 sol.
86	120-1c.		d-Benzoyl- α -aminobutyric Ac., $\text{C}_6\text{H}_5\text{CONH.CHEt.CO}_2\text{H}$. — Cryst. s. in 93 pt. aq. at 20°. — $[\alpha]_D^{20} = +30.7^{\circ}$ (1.1 g. disd. in 1 mol. aq. NaOH, total wt. 14.306 g.).
87	120-1	Alk. sol. yellow	Ethyl Benzoylisotrosoacetate, $\text{C}_6\text{H}_5\text{CO.C(NOHO).CO}_2\text{Et}$. — Ndl. fr. dil. alc., e.s. alc., eth., chlf., bz.; e.s. alk.
88	121 (frothing)	89.5 $k.10^4 = 1.2$	Anilinomalononic Ac., $\text{PhNH.CH(CO}_2\text{H})_2$. — Ndl. fr. alc. + lgr.; e.s. h. aq., alc.; less sol. eth. — FeCl_3 gives deep red-brown color. Boiled in aq. sol. decomposes to anilinoacetic ac. & CO_2 .
	121-1.5u.c.	Alk. sol. yellow	Picric Ac. — Cf. No. 2.3168.
90	abt. 121	Alk. sol. colored	$\text{o-Nitrophenylpyruvic Ac.}, \text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{CO.CO}_2\text{H}$. — \oplus Aq. sol. + FeCl_3 becomes emerald green! — Sl. yellowish ndl. fr. bz., sintering at 115°. S. h. aq.; v.s. alc., eth.; d.s. bz.; v.d.s. lgr. — Salts v. unstable; those of alkalies, deep brown. — (D.R.P. 92,794.)
91	Sbl. 122-3d.		Indoxylic Ac., $[\text{NH.C}_6\text{H}_4\text{C(OH) : C.CO}_2\text{H}]$. — \oplus When dissolved in dil. NaOH and air is bubbled through sol., ppt. of indigo blue forms.
92	122	Alk. sol. prob. yel. $k.10^4 = 2.1$	3,5-Dinitrophenol, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{OH}$. — Silky lft. fr. h. dil. HCl. — Me. \bar{A} , long ndl., m.p. 105°.
93	123-4		Dimethylisoxalone, $\text{C}_6\text{H}_5\text{O}_2\text{N}$. — White ndl. e.s. alk. carbonates; pptd. by acids. S. h. aq.; e.s. alc., eth. — Alc. sol. + FeCl_3 , red-brown.
94	123.5-4	Alk. sol. yellow	Isonitrosobenzoylacetone, $\text{Ph.CO.C(NOHO).CO.Me}$. — Ndl. i. c. aq., lgr.; e.s. eth., chlf., bz.
95	124-5	$k.10^4 = 7.5$	Benzhydroxamic Ac., PhC(NOHO).OH . — \oplus Sol. + FeCl_3 gives dark red ppt. s. in x.s. FeCl_3 w. intense dark cherry red color. Rhomb. tbl. s. in 44 pt. aq. at 6°. V.s. alc.; d.s. eth.; i. bz. — Hydrolyze w. h. HCl & test for hydroxylamine (T. 2.17-b).
96	125d.		Anthracene Nitrate, $\text{C}_{14}\text{H}_{10}\text{NO}_3$. — \oplus Should give blue color in T. 2.15! Colorless ndl. e.s. h. bz. Unstable in presence of moisture. "Imparts yellow color to NaOH sol." Evolves nitrous fumes in melting.
97	124.5-5	209	$\text{o-Nitrophenylglycidic Ac.}, \text{NO}_2\text{C}_6\text{H}_4(\text{O.CH.CH}^2)\text{CO}_2\text{H}$. — \oplus Decomposes to CO_2 & indigo on heating! — Cryst. fr. h. aq. in ndl. (w. cryst. aq.), m.p. 94°. — E.s. alc., h. aq.; i. lgr. Colored red by conc. H_2SO_4 . — Ag \bar{A} , stable, cryst., ppt.
98	126		$\text{o-Nitrophenyllactic Ac.}, \text{NO}_2\text{C}_6\text{H}_4\text{CH(OH).CH}_2\text{CO}_2\text{H}$. — \oplus Warmed w. conc. H_2SO_4 gives blue sol.! Monclin. pr. fr. aq. E.s. aq., alc., eth.

(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
99	126-7; 125	151 $k \cdot 10^4 = 3.8$	Anilinoacetic Ac., Phenylglycine, $\text{PhNH} \cdot \text{CH}_2 \cdot \text{CO}_2 \text{H}$. — \textcircled{P} Prepare CaA_2 , & ignite w. calcium formate. Gives fecal odor of indole! — Mix w. powdered moist KOH & fuse cautiously at low temp. Dissolve mass in aq. & expose to air. A coloration or ppt. due to indigo will appear. — Cryst. mod. s. aq.; d.s. eth.
100	126d.	146 $k \cdot 10^4 = 2$	Benzoylcyanamide, $\text{PhCO} \cdot \text{NH} \cdot \text{CN}$. — \textcircled{P} Should give NH_2 easily in T. 2.7. — Rhomb. pr. fr. acetone. E.s. alc.; eth.; d.s. bz.; i. lgr. — Distn. should give benzonitrile (No. 2.2781). AgA_2 , floc. ppt.
101	125-7		Suberic Ac., $\text{NH}_2 \cdot \text{CO} \cdot (\text{CH}_2)_4 \cdot \text{CO}_2 \text{H}$. — \textcircled{P} Should give NH_2 in T. 2.7. — Saponify to suberic ac. (T. 2.26).
102	126.5	240	2,4-Dinitrohydrocinnamic Ac., $(\text{NO}_2)_2 \cdot \text{C}_6\text{H}_3 \cdot (\text{CH}_2)_2 \cdot \text{CO}_2 \text{H}$. — \textcircled{P} Should give T. 2.21. — Nearly colorless ndl. fr. h. aq. S. h. aq., alc.; i. eth.
103	125-30		Adipamic Ac., $\text{NH}_2 \cdot \text{CO} \cdot (\text{CH}_2)_6 \cdot \text{CO}_2 \text{H}$. — \textcircled{P} Should give NH_2 in T. 2.7. — Ndl. fr. aq. — Saponify to adipic ac. (Vol. I) by T. 2.26.
104	abt. 127	163	Anilpyruvic Ac., $\text{PhN} \cdot \text{CMe} \cdot \text{CO}_2 \text{H}$. — \textcircled{P} Dis. w. violet-red color in conc. H_2SO_4 ! — E.s. aq.; d.s. bz., chlf. — BaA_2 , v.s. aq. Aq. sol. boiled gives off aniline.
105	127 compl. d.	$k \cdot 10^2 = 1.5$	anti-Phenylglyoxylic-acid-oxime, $\text{PhC}(\text{CO}_2 \text{H}) : \text{NOH}$. — Pr. fr. eth. V.s. aq., eth., alc. V. unstable, changing on contact w. aq. after 12 hr. to syn. compound.
106	127	†	3-Methyl-l-phenylpyrazolone(5), $[\text{NPh.N} : \text{CMe.CH}_2 \cdot \text{CO}]$. — \textcircled{P} Alc. sol. gives intense wine red color w. FeCl_3 . — Thick pr. fr. aq. Alm. i.c. aq., eth., lgr.; e.s. h. aq., dil. HCl, alk. or Na_2CO_3 sol. Boiled w. aq. + little FeCl_3 gives dull violet-blue ppt. (Color observed on filter after washing.) — B.p. 287° (th. i. 205 m.m.).
107	127-8	179	β -Anilinobutyric Ac., $(\text{PhNH}) \cdot \text{CHMe} \cdot \text{CH}_2 \cdot \text{CO}_2 \text{H}$. — Ndl. d.s. c. aq.; v.s. alc., eth. — BaA_2 (at 100°), scales.
108	129	Alk. sol. yellow	Isonitrosobenzyl Cyanide, $\text{PhC}(\text{NOH}) \cdot \text{CN}$. — Lft. e.s. h. aq.; e.s. alc., eth. — AgA_2 , floc. ppt.
109	129u.c.		Benzoyl-d-glutaminic Ac., $\text{CO}_2 \text{H} \cdot \text{CH}(\text{NH.CO.Ph}) \cdot (\text{CH}_2)_4 \cdot \text{CO}_2 \text{H}$. — Lft. s. in 21 pt. aq. at 20°, or in less than 2 pt. h. aq. — $[\alpha]_D^{20} \text{ in aq. sol. } (D = 1.0114) = +13.34^\circ$. — Sep. (T. 2.26) to benzoic & glutaminic ac., No. 2.357.
110	129	179	(stable)-3-Aminocuminic Ac., $(\text{NH}_2)(\text{Me}_2 \cdot \text{CH}) \cdot \text{C}_6\text{H}_3 \cdot \text{CO}_2 \text{H}$, $[(\text{Me}_2 \cdot \text{CH}) : \text{CO}_2 \text{H} = 1.4]$. — \textcircled{P} Mixed w. baryta and distilled gives cumidine (No. 2.1334). — Cryst. d.s. c. aq.; s. h. aq.; e.s. alc., eth. — Long heating w. 20 pt. aq. at 100° gives labile form, m.p. 104°. — AgA_2 , floc. ppt.
111	127-32		3,4-Dimethyl-l-phenylpyrazolone(5), $[\text{CHMe} \cdot \text{MeC} : \text{N.NPh-CO}]$. — Cryst. powd. V.d.s. aq., eth.; e.s. alc. — (Fr. phenylhydrazine + ethyl methylacetacetate.)
112	130d.	117	Dimethyloxamic Ac., $\text{Me}_2\text{N.CO.CO}_2 \text{H}$. — \textcircled{P} Should evolve dimethylamine (No. 2.1061) in T. 2.7-a. — Flat ndl. fr. bz.; v.s. aq., alc.; d.s. eth., bz.
113	130	179	o-Tolyloxamic Ac., $\text{NH}(\text{C}_6\text{H}_4)_2 \cdot \text{CO.CO}_2 \text{H}$. — \textcircled{P} Ignition w. Zn dust gives indole (fecal odor). — Ndl. fr. aq. w. 1 mol. aq. of crystn.; m.p. 83-4°. — D.s. c. aq.; s. alc.; less s. chlf., eth.; i. lgr. — Sapon. should give o-toluidine (No. 2.1262) & oxalic ac.
114	131		p-Aminohydrocinnamic Ac., $\text{NH}_2 \cdot \text{C}_6\text{H}_4 \cdot (\text{CH}_2)_2 \cdot \text{CO}_2 \text{H}$. — Cryst. fr. aq. — Prepare acetyl deriv. by boiling 1 pt. w. 5 pt. acetic anhydride for 5 min. Cryst. fr. aq. M.p. of product 143°. — “Metallic salts v. unstable; salts w. acids cryst. well.”

(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
I15	130-2	211	p-Nitrophenyllactic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$. — ⊖ Should give T. 2.21. — Cryst. r.d.s. c. aq.; e.s. h. aq., h. alc. or h. eth. — Protracted boiling w. dil. H_2SO_4 gives p-nitrocinnamic ac., No. 2.488.
I16	130.8 u.c.	221	d, l- α -Benzoylaminovalericianic Ac., $\text{PrCH}(\text{PhCO.NH})\text{CO}_2\text{H}$. — Lft. fr. eth. + lgr. V.d.s. h. aq.; mod. s. alc., eth.; alm. i. lgr.
I17	132d.	179 $k.10^4 = 2$	Malonanilic Ac., $\text{PhNH.CO.CH}_2\text{CO}_2\text{H}$. — ⊖ Should give carbylamine odor in T. 2.12. — Cryst. s. aq. — Dec. quantitatively on fusion to CO_2 & acetanilide (No. 2.1975).
I18	132	151	2-Amino-m-toluic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — Pr. s. h. aq.
I19	132-4 (138-40)	465 $k.10^4 = 1.3$	Glycocholic Ac., $\text{C}_2\text{H}_5\text{O}_4\text{NH.CH}_2\text{CO}_2\text{H}$. — ⊖ Pettenkofer's gall-acids react. (T. 2.42) gives coloration, at first deep red, but gradually changing towards blue-violet and showing an absorption band between D & E near F, and a second near F on diluting w. alc. — Fine ndl. S. in 300 pt. c. aq.; more s. h. aq.; e.s. alc.; alm. i. eth., bz. Taste bitter-sweet. Dextrorotatory. — Aq. sol. of alk. salts froths like soap sol. when shaken.
I20	132-4		Xylylglycine, $\text{Me}_2\text{C}_6\text{H}_3\text{NH.CH}_2\text{CO}_2\text{H}$, [Me : Me : NH = 1:3:4]. — Pr. fr. dil. alc.; i. aq.; e.s. alc.
I21	132-5	†	Acetylphthalimide, $\text{C}_8\text{H}_4\text{(CO)}_2\text{N}(\text{C}_6\text{H}_5\text{O})$. — Octahedra fr. bz. or c. alc. I. c. aq.; alm. i. c. alc. — Saponified even by c. aq. NaOH sol. to phthalimide (No. 2.2555) & ac. ac. — (Gen. T. 2.I positive only by use of fine powder & vigorous stirring.)
I22	134	Alk. sol. yellow $k.10^4 = 4.3$	3,4-Dinitrophenol, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{OH}$. — "Colorless" silky ndl. gradually crumbling to sandy powd. Under aq. melts at 50-60°. — Alk. salts colored. — MeA , yel. ndl., m.p. 70°.
I22-I	134.5		Quinolinic (Pyridine-2,3-dicarbonic) Ac. Anhydride, $[\text{CO.C}_6\text{H}_4\text{N.CO}_2]$. — CCl_4 ppts. cryst. fr. sol. in gl. ac. ac. I. aq. — ⊖ Fusion w. phenol & v. little conc. H_2SO_4 at 120°, or w. resorcinol alone at 200°, gives products s. dil. NaOH w. intense colors similar to those of alk. phenolphthaleine or fluoresceine sol.
I23	134		Indole-Pr-3-propionic Ac., Skatoleacetic Ac., $\text{C}_8\text{H}_7\text{N}(\text{CH}_3)_2\text{CO}_2\text{H}$. — Lust. pr. fr. h. aq. V.s. alc., eth. — Sol. + KNO_2 + ac. ac. gives cryst. ppt. of nitroso deriv.; i. aq.; m.p. 135°. — (Fr. putrefaction of albumin.)
I24	134-5	201 $k.10^4 = 5.9$	β -Naphthylaminoacetic Ac., β -Naphthylglycine, $\text{C}_{10}\text{H}_7\text{NH.CH}_2\text{CO}_2\text{H}$. — Mic. cryst. fr. aq.
I25	134-5		Methylaceturic Ac., $\text{MeN}(\text{CH}_2\text{CO})\text{CH}_2\text{CO}_2\text{H}$. — Cryst. e.s. aq. — AgA , ndl.
—	134-5d.		5-Nitroso-o-cresol. — Cf. No. 2.3243.
I27	135	193	α -Anilinoisovalericianic Ac., $\text{Me}_2\text{CH.CH}(\text{NHPh})\text{CO}_2\text{H}$. — Lft. fr. h. aq. (1000 pt. aq. at 100° dis. 6-7 pt.). V.e.s. alc., eth. — Reduces ammon. AgNO_3 in the cold. B.HCl , ndl. e.s. alc., aq.
I28	135d.		p-Nitrobenzoylacetic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CO.CH}_2\text{CO}_2\text{H}$. — Mic. ndl. fr. bz. E.s. alc., eth., chlf. — FeCl_3 colors alc. sol. deep red-brown. — Boiled w. dil. H_2SO_4 or alkalies dec. to p-nitroacetophenone & CO_2 . — Salts v. unstable.
I29	135	195	2-Nitro-1,3-dimethylbenzoic(4) Ac., $\text{NO}_2\text{C}_6\text{H}_3\text{Me}_2\text{CO}_2\text{H}$. — Cryst. e.s. alc.
I30	136	$k.10^4 = 5 \dagger$	2-Pyridinecarbonic Ac., Picolinic Ac., $\text{C}_5\text{H}_4\text{N.CO}_2\text{H}$. — Ndl. fr. aq. or alc. E.s. aq.; alm. i. eth., bz., chlf. — Sbl. in ndl. — Ignition w. CaO gives pyridine & dipyridine. — CuA (dried at 100°), small blue-violet ndl. w. metallic luster. ("Characteristic Salt.") Determine the Cu.)
I31	135-9 u.c.	235	d,l-Benzoylleucine, $\text{Me}_2\text{CH.CH}_2\text{CH}(\text{Ph.CO.NH})\text{CO}_2\text{H}$. — ⊖ Cryst. s. in 200 pt. h. aq., or 690 pt. at 19°. Separates at first in oily drops fr. h. sol.! — E.s. alc., eth., chlf., Na_2CO_3 sol.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
I32	137-8d.	k.10 ⁴ = 1	Nitrosoacetic Ac., HON:CH.CO ₂ H. — Ndl. fr. alc. V.s. aq., alc.; less s. eth. — Colored red by FeCl ₃ . — Dec. to HCN, H ₂ O & CO ₂ in melting.
I33	137d.	139 k.10 ⁴ = 2.1	2,4-Dimethylpyrrole-5-carbonic Ac., Me ₂ C ₄ HNH.CO ₂ H. — \textcircled{P} Strongly heated in test-tube w. few drops aq. should give pyrrole react., T. 2.24. — V.d.s. c. aq. — Partly dec. on boiling w. aq. to CO ₂ & dimethylpyrrole (No. 2.2759).
I34	137-8d.		Aminosuccinuric Ac., C ₄ H ₆ O ₂ N ₂ CO ₂ H. — \textcircled{P} T. 2.7 should give NH ₃ . — S.c. aq.; alm. i. alc., eth.
I35	138; 139.9c.	†	Salicylamide, o-HO.C ₆ H ₄ CONH ₂ . — \textcircled{P} Should give NH ₃ in T. 2.7. — Sbl. in lft. Boils at 270° w. decn. to phenol, CO ₂ , aq. & the nitrile. S. in Na ₂ CO ₃ sol.; crystg. out unchanged on evapn. — Saponify to salicylic ac. (Vol. I) by T. 2.26-C.
I36	137; 140	Alk. sol. prob. yel.	6-Nitrothymol, NO ₂ .C ₆ H ₅ Me(OH), [NO ₂ :Me:OH = 6:1:3]. — Nearly colorless or faintly yellowish ndl. w. bluish fluorescence fr. lgr.
I37	139-40	179 k.10 ⁴ = 3.2	α -Anilinobutyric Ac., MeCH ₂ .CH(NH.Ph).CO ₂ H. — Lust. granules d.s. c. aq.; v.s. alc., eth. — Reduces Ag. salts.
I38	139	193	m-Toluric Ac., Me.C ₆ H ₄ .CO.NH.CH ₃ .CO ₂ H. — Thin lft. fr. aq.
I39	140u.c.	167 †	m-Nitrobenzoic Ac., No.C ₆ H ₄ .CO ₂ H. — \textcircled{P} Gives brown ppt. in T. 2.21. — Lft. fr. h. aq. Odorless. Nearly tasteless. — 10 cc. of each of following solvents dissolves: aq. at 20°, 0.031 g.; eth. at 11°, 2.51 g. V.d.s. chlf. lgr., bz. — Convert into m-aminobenzoic ac. by procedure given for reduction of o-nitrobenzoic ac. (No. 2.164), except that a little more aq. may be required in recrystn. of product. The m-amino acid is obtained as faintly yellowish cryst. powd., m.p. 170-1° u.c.
I40	141; 137-8	181	o-Nitro- α -tolylic Ac., NO ₂ .C ₆ H ₄ .CH ₃ .CO ₂ H. — Ndl. fr. h. aq. — Amide melts at 109-10°.
I41	141	Alk. sol. prob. col. k.10 ⁴ = 1.6	Dimethylvioluric Ac., C ₆ H ₅ O ₂ N ₂ . — \textcircled{P} Colored intense dark blue by FeSO ₄ ! — Cryst. w. 1H ₂ O in pearly ndl., melting at 124°. D.s. c. aq.; e.s. h. aq. — Boiled w. Ba(OH) ₂ gives methylamine. — AgA blue-violet ppt.
I42	141-2d.	Alk. sol. reddish	Oximinomethyl-syn-oxazolone, C ₆ H ₅ O ₂ N ₂ . — \textcircled{P} Sol. in alk. w. intense red color. — Colored deep red by FeCl ₃ sol. — Strongly heated, explodes. Cryst. w. 1H ₂ O. V.s. aq., alc. eth.
—	141-2	(?)	α -Anilinopalmitic Ac. — Cf. No. 2.2147-1.
I44	142d.	179 k.10 ⁴ = 3.9	α -Anilinoisobutyric Ac., MeC(NHPh).CO ₂ H. — Pr. fr. chlf. — Dec. on distn. to isopropylaniline (No. 2.1302) & CO ₂ .
I45	143	193 k.10 ⁴ = 2	Phenaceturic Ac., Ph.CH ₂ .CO.NH.CH ₃ .CO ₂ H. — Cryst. fr. alc. S. in 136.2 pt. aq. at 11.2°; e.s. alc.; v.d.s. abs. eth.; d.s. h. bz. Boiling w. conc. HCl hydrolyzes to phenylacetic ac. (Vol. I) & glycine No. 2.2568. — PbA ₂ .2H ₂ O, pr. d.s. h. aq. [In urine of herbivorous animals.]
I46	143	207 k.10 ⁴ = 1.2	α -Acetylanilinopropionic Ac., Me.CH[PhN(Me.CO)].CO ₂ H. — Pr. fr. h. aq. D.s. c. aq. or bz.
I47	143-4u.c.	207	Benzoyl-d,l- α -aminobutyric Ac., (Ph.CO)NH.CHEt.CO ₂ H. — Cryst. fr. 25 pt. h. aq. Sinters abt. 140°. S. in 225 pt. aq. at 20°. V.s. alc., chlf.; d.s. eth.; alm. i. lgr.
I48	144-5u.c.	137 k.10 ⁴ = 1.0	α -Aminobenzoic Ac., Anthranilic Ac., NH ₂ .C ₆ H ₄ .CO ₂ H. — \textcircled{P} Aq. sol. shows pale bluish fluorescence and tastes sweet. — Place abt. 0.01 g. in a dry t.t. & heat sharply w. small flame to decompose to CO ₂ & aniline. Apply T. 2.12 to the oily drop of aniline that condenses on walls of t.t. Strong carbylamine odor will be observed. — Long flat ndl. fr. h. aq. — Odorless. — 10 cc. of each of following solvents dissolve: aq., 0.035 g. at 13.8°; 90% alc., 1.07 g. at 9.6°; eth., 1.605 g. at 6.8°; bz., 0.018 g. at 10°. — Add 1 drop sat. aq. sol. bleaching powder to a cold (1:1000) aq. sol. of compound. A VR color appears,

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
			<i>changing to ROSI (by direct transmitted light fr. sky). — ① Add 5 cc. of alc. picryl chloride sol. (Cf. T. 2.22) to sol. of 0.05 g. compound in 1 cc. h. alc. Boil gently for 1 min. Add 4 cc. c. aq. Dissolve ppt. in boiling bz. & allow to cool slowly. Wash cryst. Ppt. w. 5 cc. c. alc. Recryst. fr. 3 cc. boiling gl. ac. ac., allowing to stand 30–40 min. after cooling and shaking. Wash w. 1 cc. c. gl. ac. ac. Dry 15 min. at 100°. The product, o-picrylaminobenzoic ac., $\text{O}-\text{CO}_2\text{H.C}_6\text{H}_4\text{NH.C}_6\text{H}_4(\text{NO}_2)_2$, is obtained in fine Y-YO ndl., m.p. 270–1° u.c.</i>
149	143–5		Glyoxylic-acid-phenylhydrazone, $\text{Ph.NH.N:CH.CO}_2\text{H}$. — Yellowish ndl. fr. aq.; v.d.s. c. aq.; e.s. alc. — PbA_2 , yellow ppt.
150	144	177	Methylenephthalamic Ac., $\text{CH}_2:\text{N.CO.C}_6\text{H}_4\text{CO}_2\text{H}$. — Pr. fr. bz. I. aq.; e.s. alc.; mod. s. eth.; v.d.s. c. bz. — Heating w. aq. dec. to formic aldehyde & phthalimide.
151	144	183 $k.10^3 = 1.6$	3-Nitrosalicylic Ac.(1), $\text{NO}_2\text{C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})$. — ② Gives blood red color w. FeCl_3 . — Ndl. w. H_2O of crystn., m.p. 125°. — S. in 770 pt. c. aq.; e.s. alc., eth., chlf., bz. — Ignition w. CaO gives o-nitrophenol. — BaA_2 , gold-yel. lft., d.s. h. aq.
152	144		3-Methyquinolinecarbonic(2) Ac. — Pr. fr. eth. + alc. Warts fr. aq. — Dec. at 160° to CO_2 & 3-methylquinoline (No. 2.1388).
153	144–5		α,β -Dibenzoylcaproic Ac., Dibenzoyl-d-lysine, Lysuric Ac., ($\text{Ph.CO.NH(CH}_2)_4\text{CH(NH.CO.Ph).CO}_2\text{H}$). — Lust. lft. v.d.s. c. aq. or eth.; e.s. alc. (M.p. of d, l-derivative said to be 145–6°.)
154	Deflagrates at 145	"Dec. carbonates"	Fulminuric Ac., $\text{C}_2\text{H}_3\text{O}_3\text{N}_2$. — Pr. fr. alc. Dec. by boiling w. dil. HCl (less readily w. alk.) giving oxalic ac., NH_3 & CO_2 .
155	145		Cholestrophan, Dimethylparabanic Ac., $[\text{CO.NMe.CO.NMe-CO}]$. — ② Readily sapd. by boiling w. NaOH sol. (T. 2.26-D) to methylamine (No. 2.1059), oxalic & carbonic acids! — Pearly lft. s. at 20° in 53.4 pt. aq.; d.s. c. alc.
156	145		α -Triazobenzoic Ac., $[\text{N:N.C}_6\text{H}_4(\text{CO}_2\text{H})\text{N}]$. — Ndl. mod. s. h. aq.; e.s. alc., h. bz. — Heated w. NaOH sol. gives o-azoxybenzoic ac. & anthranilic ac. (No. 2.148).
157	145d.	$k.10^3 = 1.8$	syn-Phenylglyoxylic-acid-oxime, $\text{Ph.C}(\text{CO}_2\text{H}): \text{NOH}$. — Ndl. fr. eth. Mod. s. aq.; alc., eth. — Boiling w. dil. HCl slowly changes to benzonitrile (No. 2.2781).
158	145	181	6-Nitrotoluic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$, [$\text{Me:CO}_2\text{H} = 1:2$]. — ② Should give T. 2.21. — Ndl. fr. dil. alc. D.s. h. aq.; v.s. alc.
159	145–6		Methyloxamic Ac., $[\text{MeNH.CO.CO}_2\text{H}]$ — Vol. w. st. — Boiling w. NaOH sol. gives methylamine (No. 2.1059) & oxalic ac.
160	145	"Sol. yel."	Pyruvic-acid-diphenylhydrazone, $\text{Ph.N:NCMe.CO}_2\text{H}$. — Ndl. fr. alc. E.s. bz., chlf. — Convert into l-phenylindolecarboxylic ac. (Ber. 17, 567.)
161	145–6		Dibenzoyl-d,l-lysine, $\text{Ph.CO.NH(CH}_2)_4\text{CH(Pr.CO.NH).CO}_2\text{H}$. — Plates d.s. aq., eth., bz., chlf.; e.s. alc.
162	146	193 $k.10^3 = 2.1$	Succinanilic Ac., $\text{PhNH.CO(CH}_2)_2\text{CO}_2\text{H}$. — ② Gives carbylamine odor in T. 2.12. — Ndl. fr. h. aq. V.d.s. c. aq.; e.s. alc., eth., h. aq. — Sapn. gives aniline & succinic ac.
163	146	Alk. sol. yellow	Isonitrosodibenzoylmethane, $(\text{Ph.CO})_2\text{C:NOH}$. — Powd. e.s. alk.
164	146–7u.c.	167 † $k.10^3 = 6.5$	α -Nitrobenzoic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$. — ② Gives dark brown ppt. in T. 2.21. Aq. sol. tastes intensely sweet. — 10 pt. of each of following solvents dissolve: aq. at 20°, 0.068 g.; 90% alc. at 10°, 2.8 g.; eth. at 11°, 2.16 g.; chlf. at 15°, 1.06 g. Alm. i. lgr., CS_2 . — Odorless. — ① Add 5 cc. HCl (sp. gr. 1.20) to 0.5 g. of substance + 1 g. gran. tin. Start

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
			<i>action by warming & allow to continue until tin dissolves. Evaporate sol. to abt. 1 cc. on water-bath in small glass dish. Dissolve residue in 25 cc. aq. Saturate the boiling hot sol. w. H₂S. Filter off pptd. tin sulphide. Evap. filtrate to dryness on water-bath. Dissolve residue in 5 cc. aq. + 1.5 cc. conc. ammonia. Acidify by adding several drops gl. ac. ac. Wash the crystals that separate on cooling (!) w. 2 cc. aq. Recryst. fr. 3 cc. boiling aq. after boiling the sol. w. 0.1 g. boneblack, & filtering hot. Wash cryst. which separate on cooling w. 1 cc. aq. Dry 10 min. at 100$^{\circ}$. The product, o-aminobenzoic ac., is obtained in long ndl., m.p. 144-4, 5$^{\circ}$ u.c.</i>
165	148u.c.	193 k.10 ⁴ = 2	Benzoyl-d-alanine, Me.CH(Ph.CO.NH).CO ₂ H. — Lust. plates fr. aq. S. in 85 pt. aq. at 20 $^{\circ}$. — [α] _D ²⁰ = +3.3 $^{\circ}$ in 0.99% aq. sol. (Optical antipode gives same m.p.)
166	147-8		Diphenylphthalamic Ac., Ph ₂ N.CO.C ₆ H ₄ .CO ₂ H. — Lust. cryst. fr. alc. I. aq.; v.s. alc.; d.s. eth. — Ag \ddot{A} , cryst. pulv. ppt. fr. NH ₄ \ddot{A} sol. + AgNO ₃ .
167	148-9	151	3-Amino- α -toluyllic Ac., NH ₂ .C ₆ H ₄ .CH ₃ .CO ₂ H. — Ndl. fr. h. bz.
168	148-9	165	Phthalamic Ac., o-NH ₂ .CO.C ₆ H ₄ .CO ₂ H. — (P) Gives NH ₃ in T. 2.7-2. — Mod. s.c. aq., alc.; d.s. eth., bz.; i. lgr. — Heating at 155 $^{\circ}$ dec. completely to aq. & phthalimide (No. 2.2555). V. easily saponified to phthalic ac. (No. 1.318).
169	148d.	189	Pyrotartaranic Ac., Me.C ₆ H ₅ (CO.NHPh).CO ₂ H. — (P) T. 2.12 probably gives carbylamine odor. — Ndl. d.s. aq.; e.s. alc.; d.s. chlf. — Ag \ddot{A} , pulv. ppt.
170	148d.		3-Ethylquinolinecarbolic(2) Ac., Et.C ₆ H ₅ .CO ₂ H. — Lust. ndl. fr. aq. D.s. eth.; more s. aq. — Dec. to CO ₂ & 3-ethylquinoline in melting. — Cu \ddot{A} , blue-green ppt. of mic. ndl.
171	149	128	Methylparabanic Ac., $^{[}\text{CO.NMe.CO.NMe.CO}^{]}\text{.}$ — (P) Evolves methylamine in T. 2.7 for NH ₃ , oxalic acid also being formed. — Lust. ndl. d.s. c. aq.; e.s. h. aq.; s. alc. eth.
172	149	165 k.10 ⁴ = 1.2	Oxanilic Ac., Ph.NH.CO.CO ₂ H. — (P) T. 2.12 gives strong carbylamine odor! Addition of K ₂ Cr ₂ O ₇ to sol. in conc. H ₂ SO ₄ on crucible cover gives transient VR color. — Cryst. fr. aq. w. 1H ₂ O. S. h. aq.; less s. h. bz.; e.s. alc., chlf.; d.s. lgr. — Sapn. gives aniline (No. 2.1235) & oxalic ac.
173	149-50	193 k.10 ⁴ = 6.8	α -p-Toluidinoisobutyric Ac., Me.C ₆ H ₄ .NH.CMe ₂ .CO ₂ H. — Tbl. fr. acetone d.s. lgr.
174	149u.c.	209	β -Benzoylamino- α -hydroxypropionic Ac., C ₆ H ₅ .CO.NH.CH ₂ .CH(OH).CO ₂ H. — Pr. fr. h. aq.; e.s. h. aq., alc.; d.s. eth., chlf., bz.
175	150-1		3,5-Dimethylpyridinecarbolic(2) Ac., Me ₂ .C ₆ H ₃ N.CO ₂ H. — Cryst. mass. E.s. aq., alc.
176	150-1d.	k.10 ⁴ = 2.7	Phenyliminodiacetic Ac., PhN.(CH ₂ .CO ₂ H) ₂ . — Silky ndl. e.s. h. aq., alc.; less s. eth.
177	151d.	k.10 ⁴ = 8	α -Isonitrosobutyric Ac., Et.C(NO ₂).CO ₂ H. — Silky ndl.; e.s. alc.; less s. aq., eth.
178	151.5	181	p-Nitro- α -toluyllic Ac., NO ₂ .C ₆ H ₄ .CH ₃ .CO ₂ H. — (P) Should give T. 2.21. — Silky ndl. d.s. aq.; e.s. alc. — Me \ddot{A} , ndl. fr. lgr., m.p. 54 $^{\circ}$.
179	151		2-Methylquinolinecarbolic(8), Ac. Me.C ₆ H ₄ N.CO ₂ H. — (P) Distillation gives quinaldine (No. 2.1376) & CO ₂ . — Ndl. d.s. c. aq.; e.s. h. aq., alc. — B ₂ H ₅ PtCl ₆ .2Aq., red pr. d.s. c. aq.; e.s. h. aq. (T. 2.14). — Cu \ddot{A} .1½Aq., dark green ndl.; losing 1H ₂ O at 100 $^{\circ}$.
180	152	149 k.10 ⁴ = 5.5	3-Phenylisoxazolone(5), $^{[}\text{CPh.NH.O.CO.CH}^{]}\text{.}$ — (P) Reduces ammon. AgNO ₃ . Ndl. v.d.s. aq., alc., eth. Heated w. conc. H ₂ SO ₄ gives sulphanilic ac. ac. ac., CO ₂ .

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
181	152	181	5-Nitro-o-toluidic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — (P) Should give T. 2.21. — Lust. ndl. fr. aq. V.d.s. c. aq.; v.s. h. aq., c. alc. — $\text{BaA}_2\cdot 5\text{H}_2\text{O}$, flat pr. e.s. aq.
182	153; (165d.)	151	5-Amino-o-toluidic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — Ndl. fr. alc. — Sbl.
183	153		2,4-Dimethylpyridinecarboxylic (2) Ac., $\text{Me}_2\text{C}_6\text{H}_3\text{N.CO}_2\text{H}\cdot \frac{1}{2}\text{H}_2\text{O}$. — Cryst. v.s. aq., alc.; less s. eth. — Ignition w. CaO gives dimethylpyridine.
184	153-4u.c.		Benzoyl-d,L-glutaminic Ac., $\text{CO}_2\text{H}(\text{NH.CO.Ph})\cdot (\text{CH}_2)_2\text{CO}_2\text{H}$. — Cryst. w. $1\text{H}_2\text{O}$; then s. in 124 pt. c. aq. E.s. alc. — Ag. salts d.s. aq.
185	153		Methylglyoxime, $\text{MeC}(\text{NOH}).\text{CH}(\text{NOH})$. — Pr. fr. alc. Sbl. in ndl. D.s. c. aq.; e.s. h. aq.; e.s. NaOH (sol. colorless). — Ammon. AgNO_3 gives ppt. $\text{AgC}_6\text{H}_5\text{O}_2\text{N}_3$, somewhat sol. aq.
186	154		Succinamic Ac., $\text{NH}_2\text{CO.CH}_2\text{CH}_2\text{CO}_2\text{H}$. — (P) T. 2.7-a gives NH_3 . — D.s. c. aq. — Salts gradually decd. by boiling aq. — Sap. to succinic ac. (Vol. I). — AgA_2 , cryst. ppt.
187	153-6d.		Hydantoic Ac., $\text{NH}_2\text{CO.NH.CH}_2\text{CO}_2\text{H}$. — Monoclin. pr., s. c. aq. alc.; v.d.s. eth. — Colored red by FeCl_3 sol.
188	153-6	190 $k.10^4 = 1$	α -p-Toluidinobutyric Ac., $\text{MeC}_6\text{H}_5\text{NH.CH(CH}_2\text{Me).CO}_2\text{H}$. — Lft. fr. eth. E.s. alc.; d.s. chlf., bz.
189	154-5d.		Malonhydroxamic Ac., $\text{CH}_2[\text{C}(\text{NOH})(\text{OH})]_2$. — Pr. fr. aq. — Melts w. brisk efferv. — I. eth.
190	154d.		Malonphenylhydrazidic Ac., $\text{PhNH.NH.CO.CH}_2\text{CO}_2\text{H}$. — Ndl. s. aq.
191	155		2,4,6-Trimethylpyridinecarboxylic (2) Ac., $\text{Me}_2\text{C}_6\text{H}_3\text{N.CO}_2\text{H}$. — Cryst. w. $2\text{H}_2\text{O}$ in ndl. or cube, m.p. 110°. V.s. aq. — KA_2 , ndl. fr. alc. — B.HCl , ndl. e.s. aq.
192	d. suddenly at 155-6	191 $k.10^2 = 1.1$	σ -Nitrophenylpropionic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{C}_2\text{H}_4\text{CO}_2\text{H}$. — Ndl. fr. h. aq. E.s. h. aq.; v.d.s. chlf.; alm. i. CS_2 , lgr. — (P) (D) Add a few cgs. finely powdered ac. & 2 drops 10% NaOH sol. to 10 cc. 10% aq. sol. of grape sugar. Sol. becomes green, soon darkening to blue with ptn. of indigo when boiled, or will show this change after addition (while hot) of 1 to 3 more drops of alkali. Much x's of alkali must be avoided.
193	155	$\mu(32) = 3.9$	Tetrazole, Penta-1,2,3,4-tetrazadiene (2, 4), $[\text{N}=\text{N}:\text{N}=\text{N}:\text{-CH}]$. — Lft. fr. alc. — Sbl. — E.s. aq., alc.; d.s. eth., bz. — Heating w. conc. HCl gives CO_2 , NH_3 & N_2 . Salts are explosive.
194	155; d. 155-8	Alk. sol. colored	6-Nitrosocresol (3), Toluquinone-o-oxime, $\text{Me.C}_6\text{H}_4\text{(O)(NOH)}$, [$\text{Me} = 1$]. — White or pale greenish-yel. ndl. fr. eth. or dil. alc. Aq. sol. greenish yel. Conc. sol. of alk. salts red; but dil. sols. greenish-yel.
195	abt. 156d.	$\mu(32) = 9.3$	Methylalloxan, $[\text{CO.NMe}(\text{CO})_2\text{NH}]$. — Cryst. fr. aq.
196	156	173 $k.10^4 = 1.3$	α -Quinolinecarboxylic Ac., Quinoline-2-carboxylic Ac., Quinaldinic Ac., $\text{C}_8\text{H}_5\text{N.CO}_2\text{H}$. — (P) Heated above m.p. dec. to quinoline (No. 2.1356), (unpleasant, characteristic odor) & CO_2 . — Asbestos-like ndl. fr. aq. w. $2\text{H}_2\text{O}$ of crystn. lost at 100°. D.s. c. aq.; s. h. bz. — $\text{B}_2\text{H}_6\text{PtCl}_6\cdot 2\text{H}_2\text{O}$, red triclin. tbl., d.s. c. aq.; e.s. h. aq.
—	157-8	286	2,6-Dinitro-p-toluidic Ac. — Cf. No. 2.3389-1.
196	158	179 $k.10^4 = 6.9$	α -(p)-Toluidinopropionic Ac., $\text{Me.CH}(\text{NH.C}_6\text{H}_5)\text{CO}_2\text{H}$. — Silvery lft. fr. dil. alc. E.s. alc., eth., chlf., alk., mineral acids; d.s. lgr., bz.
199	158	190	Citrodiamic Ac., $\text{HO.C}_6\text{H}_4(\text{CO}_2\text{H})(\text{CONH}_2)_2$. — Lft. e.s. aq.; alm. i. alc., eth.
200	156-7; 159	209 $k.10^4 = 2.1$	3-Nitrocuminic Ac., $(\text{NO}_2)^2(\text{Me}_2\text{CH})^1\text{C}_6\text{H}_3\text{(CO}_2\text{H})^4$. — (P) Should give T. 2.21. — Cryst. (sl. yellowish) fr. alc. I. aq.; e.s. alc., eth. — Exposed to sunlight in bz. sol. yields red substance s. w. intense violet-red color in ammonia. — MeA , m.p. 64°.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
201	159-9d.		1-Hydroxyindolecarboxylic(2) Ac. , $\text{C}_8\text{H}_7\text{CH}:\text{C}(\text{CO}_2\text{H})\text{N}(\text{OH})$. — \textcircled{P} Warmed w. conc. H_2SO_4 gives deep blue sol. which remains clear on dilution w. aq.! — Pr. fr. dil. acetone. Mod. s. c. sq., e.s. alc., eth.; d.s. bz., chlf. — Unstable. Alc. sol. on long exposure to air becomes emerald-green & finally indigo-blue. Conc. HNO_3 gives deep cherry-red sol., becoming yellow on standing.
202	158-62d.	223 $k \cdot 10^4 = 2.1$	o-Tolyliminodiacetic Ac. , $\text{MeC}_6\text{H}_4\text{N}(\text{CH}_2\text{CO}_2\text{H})_2$. — Cryst. d.s. eth., c. bz.; s. alc.
203	160d.	119	Tartronamic Ac. , $(\text{HO})\text{CH}(\text{CONH}_2)(\text{CO}_2\text{H})$. — \textcircled{P} Should give NH_3 in T. 2.7. — Ndl. d.s. c. sq.; s. alc.; v.d.s. eth. — Sapn. gives CO_2 & glycollic ac. — $\text{Pb}\bar{\text{A}}_2\cdot 1\frac{1}{2}\text{H}_2\text{O}$ (charac.).
204	160		2,6-Dimethylpyridinecarboxylic(3) Ac. , $\text{Me}_2\text{C}_6\text{H}_3\text{N.CO}_2\text{H}\cdot \frac{1}{2}\text{H}_2\text{O}$. — Ndl. s. aq.
205	160		o-Tolylaminoacetic Ac. , $\text{C}_7\text{H}_7\text{NH.CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Reduces ammon. AgNO_2 on warming. — Cryst. i. c. aq.; v.s. alc., eth. — Distn. of $\text{Ca}\bar{\text{A}}$ gives pure o-toluidine (No. 2.1262)!
206	160	197	Salicyluric Ac. , $\text{o-HO.C}_6\text{H}_4\text{CO.NH.CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Colored violet by FeCl_3 . — Thin ndl. D.s. c. aq.; e.s. alc.; mod. s. eth. — Sapn. (Cf. T. 2.26-a) gives salicylic ac. (Vol. I) & glycine.
207	160	226	2,4-Dinitro-α-toluyllic Ac. , $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq. — Na amalgam gives black acids & NH_3 at once. Alk. salts on boiling w. aq. dec. quickly to carbonates & 2, 4-dinitrotoluene!
208	160-1	237	p-Ethoxyphenylsuccinamide Ac. , $\text{EtO.C}_6\text{H}_4\text{NH.CO.C}_6\text{H}_4\text{CO}_2\text{H}$. — \textcircled{P} Is colored violet by Cl-aq. — Pearly lft. fr. h. aq. S. alc. — $\text{Na}\bar{\text{A}}$ is the antipyretic, "pyrantin."
209	abt. 160d. (r.h.)		Benzoylformic-acid-phenylhydrazone , $\text{Ph.NH.N:CPh}(\text{CO}_2\text{H})$. — \textcircled{P} Boiled w. alc. gives odor of benzaldehyde. — Ndl. fr. gl. ac. ac. Alm. i. c. aq.; v.s. alc., eth.
210	160u.c. d.	250	α-Phenylureidoisobutylic Ac. , $\text{Me}_2\text{CH.CH}_2\text{CH}(\text{NH.CO-NHPh}).\text{CO}_2\text{H}$. — Ndl. fr. warm dil. alc. s. in 300 pt. boiling aq. or 2 pt. boiling alc.; v.s. eth., chlf., bz. — Corresponding hydantoin (Z. physiol. Chem., 33, 177-92), m.p. 125°.
211	161-2d.		Pyrrolecarboxylic(3) Ac. , $\text{C}_4\text{H}_5\text{N.CO}_2\text{H}$. — \textcircled{P} T. 2.24-a gives pyrrole reaction. — Ndl. Dec. on keeping or boiling w. aq. to CO_2 & pyrrole.
212	161	118.5	o-Nitrobenzalmalonic Ac. , $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{C}(\text{CO}_2\text{H})_2$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq.; e.s. aq.; v.s. eth.; i. bz. — Continued boiling w. aq. dec. to o-nitrobenzaldehyde, malonic ac., & a little o-nitrocinnamic ac. — $\text{Ba}\bar{\text{A}}\cdot 2\frac{1}{2}\text{H}_2\text{O}$, cryst. pptd. fr. $(\text{NH}_4)_2\bar{\text{A}}$ sol. + BaCl_2 .
213	161; 164-5c.	181	3-Nitro-p-toluic Ac. , $\text{NO}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq. — Sbl. D.s. h. aq.; v.s. alc., eth. — $\text{Ag}\bar{\text{A}}$, ndl. i. c. aq. — $\text{Ba}\bar{\text{A}}\cdot \text{XH}_2\text{O}$, lft. v.s. aq.
214	161-1-5	193 $k \cdot 10^4 = 2$	p-Toluric Ac. , $\text{Me.C}_6\text{H}_4\text{CONH.CH}_2\text{CO}_2\text{H}$. — Lft. e.s. h. aq.; e.s. alc.; d.s. eth. — $\text{Ba}\bar{\text{A}}\cdot 5\text{H}_2\text{O}$, ndl. e.s. h. aq.
215	161	211	4-Nitrophthalic Ac. , $\text{NO}_2\text{C}_6\text{H}_4(\text{CO}_2\text{H})_2$. — \textcircled{P} Should give T. 2.21. — Cryst. in ndl. w. $1\text{H}_2\text{O}$ (lost at 100°). S. aq., alc., eth.; i. chlf., bz. Reduction w. $\text{Sn} + \text{h. HCl}$ gives CO_2 & m-aminobenzoic ac. (No. 2.259). — $\text{Me}_2\bar{\text{A}}$, m.p. 65-6°.
216	161	215	α-Naphthylaminopropionic Ac. , $\text{C}_{10}\text{H}_7\text{NH.CHMe.CO}_2\text{H}$. — Scales fr. bz. E.s. alc., eth., chlf.
217	160.7u.c.	236	d,L-Phenylureidovalerenic Ac. , $\text{Pr.CH}(\text{NH.CO.NHPh})\text{-CO}_2\text{H}$. — Lft. fr. 130 pt. h. aq. Mod. s. h. alc.; d.s. eth.; e.s. alk. carbonates.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
218	161	"Reacts ac."	Benzhydroxamic-acid-benzoate, Dibenzhydroxamic Ac., Ph.C-(NO ₂ C ₆ H ₄ O)(OH). — \textcircled{P} FeCl ₃ gives red-yel. ppt. w. aq. sol. of K salt. — Ndl. alm. i. aq.; r.d.s. alc.; d.s. eth.; i. bz. — Sapd. by alk. (T. 2.26) to benzoic & benzhydroxamic acids.
219	162		d- and L-Pyroglutaminic Acids, Pyrrolidinecarboxylic Ac., $^{\text{L}}\text{CH}(\text{CO}_2\text{H})\text{CH}_2\text{CH}_2\text{CO.NH}^{\text{L}}$. — \textcircled{P} Vapors evolved on ignition in glass tube give pyrrole-red splinter reaction (T. 2.24). Tbl. fr. aq. s. in 2.1 pt. aq. at 13°. — $[\alpha]_D = \pm 7^{\circ}$ for sol. of 6.33 g. in 50 cc. aq.
220	162	165 $k.10^4 = 4.2$	α -Anilinopropionic Ac., Me.CH(Ph.NH).CO ₂ H. — \textcircled{P} Probably gives carbalamine odor in T. 2.12. — Lft. fr. h. aq.; d.s. c. aq.; e.s. alc.; d.s. eth., chlf., bz. — Sbl. undec.
221	162.5	193 $k.10^4 = 1.9$	α -Toluric Ac., Me.C ₆ H ₄ .CO.NH.CH ₃ .CO ₂ H. — Cryst.
222	162	224	m-Nitrohippuric Ac., (NO ₂).C ₆ H ₄ .CO.NH.CH ₂ .CO ₂ H. — \textcircled{P} Should give T. 2.21. — Ndl. s. in 271 pt. c. aq.; e.s. h. aq., alc., eth. — PbA ₂ .5H ₂ O, cryst. ppt.
223	abt. 162 u.c. d.	250	d,l-Phenylureidoisobutylic Ac., Me ₂ CH.CH ₂ .CH(NH.CO-NHPh).CO ₂ H. — Lft. fr. alc. S. in 300 pt. boiling aq., or 2 pt. boiling alc.; progressively less sol. in eth., chlf., bz., lgr. — Corresponding hydantoin (Cf. T. 2.31) melts at 125°.
225	163 u.c.	225 † $k.10^4 = 2$	d,l-Benzoylalanine, Me.CH(Ph.CO.NH).CO ₂ H. — \textcircled{P} Taste sweet. — Lft. fr. eth. S. in 250 pt. c. aq.; e.s. alc.; d.s. eth. — AgA, lft., v.d.s. alc.
226	163-4	195	p-Nitrohydrocinnamic Ac., NO ₂ .C ₆ H ₄ .CH:CH.CO ₂ H. — \textcircled{P} Should give T. 2.21. — Flat ndl. fr. h. aq. Alm. i. c. aq.; v.d.s. h. aq.; e.s. h. alc. — Oxidation by CrO ₃ mixture gives o-nitrobenzoic ac. (No. 2.164). — BaA ₂ .2H ₂ O, ndl. r.d.s. c. aq.
226-1	163-4	212	3,4-Dinitrobenzoic Ac., (NO ₂) ₂ C ₆ H ₃ .CO ₂ H. — Taste intensely bitter! Cryst. aggregates. Sbl. 100 pt. aq. at 25° dis. 0.673 pt.; e.s. alc., eth., h. aq.
226-2	163-4		3-Nitrophthalic Anhydride, $^{\text{L}}\text{C}_6\text{H}_4(\text{NO}_2)\text{CO.O.CO}^{\text{L}}$. — Warm w. alk. sol. & convert to corresponding ac. (No. 2.389).
227	164-5	151	2-Amino-p-tolanic Ac., NH ₂ .C ₆ H ₄ .Me(CO ₂ H). — \textcircled{P} Should give N in T. 2.4. — Ndl. mod. s. aq. — BaA ₂ .1½H ₂ O, v.s. aq.
226	164	195	4-Nitro-L-ethylbenzoic Ac., NO ₂ .C ₆ H ₄ .Et.CO ₂ H. — Cryst. e.s. alc., eth.; i. lgr.
229	164		Geronic-acid-semicarbazone, Me.C(:N.NH.CO.NH).CH ₂ .CMe ₂ (CH ₃) ₂ .CO ₂ H. — \textcircled{P} Should give Ag. ppt. after hydrolysis in T. 2.17. — Cryst. fr. acetic-eth.
230	165	†	Pyrazolone, $^{\text{L}}\text{NH.N:CH.CH}_2\text{CO}^{\text{L}}$. — \textcircled{P} Aq. sol. is colored OR-RO by FeCl ₃ sol. — Ndl. fr. toluene. E.s. aq.; v.s. alc.; v.d.s. eth. — Tasteless & odorless. — Convert to 4-isonitroso-pyrazolone (Cf. Ber., 1896, 256).
231	165	175(?)	Indole-Pr-3-acetic Ac., "Skatolecarboxylic" Ac., C ₉ H ₈ N.CH ₂ .CO ₂ H. — \textcircled{P} Heated above m.p. yields fecal odor of skatole. — Lft. fr. aq. or bz. — V.d.s. c. aq.; e.s. alc., eth. — Upon adding a few drops of HNO ₃ (sp. gr. 1.12) to a 1% sol., and then a few drops of 2% KNO ₃ , the sol. quickly becomes cherry red & then turbid, depositing a red coloring matter s. in ac. eth. Color in ac. eth. is changed to yel. by NaOH & restored by acid. — [A product of protein putrefaction.]
232	165		Pyromykuric Ac., C ₉ H ₈ O.CO.NH.CH ₂ .CO ₂ H. — Pr. fr. aq. Boiled w. baryta sol. gives glycine & pyromucic ac. [Fr. urine of dogs fed w. furfural.]
233	165-6 u.c.	224	α -Phenylureidohydroxymethylacetacetic Ac., CH ₂ (OH).CH-(CO ₂ H).NH.CO.NHPh. — Fine ndl. fr. aq. V.s. h. aq.; mod. s. c. aq.; e.s. alc. Fr. action of d,L-serin on phenylisocyanate.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS. — Colorless and Solid.
234	165	228	Hydrazobenzene- α -carbonic Ac., Ph.NH.NH.C ₆ H ₄ .CO ₂ H. — Pr. fr. dil. alc. E.s. alc., eth.; d.s. lgr., bz.
235	166d.	187	N-Phenylpyrrole- α -carbonic Ac., Ph.NC ₆ H ₅ .CO ₂ H. — \textcircled{P} T. 2.24-b gives violet coloration on splint. — Odorless, tasteless ndl. fr. bz. or alc. I. c. aq.; v.s. alc., bz.
236	165-7d.	195 k.10 ⁴ = 4.7	Pr-3-Methylindole-Pr-2-carbonic Ac., Me.C ₆ H ₅ N.CO ₂ H. — \textcircled{P} Heated above m.p. gives fecal odor of skatole. — Warmed w. conc. H ₂ SO ₄ gives purple-red color. — Ndl. or lft. fr. h. aq. D.s. aq.; e.s. alc., eth.; less s. bz.; i. lgr. — FeCl ₃ gives deep red color w. alc. sol. — Ag \ddot{A} , pulv. i. ppt.
237	164-8d.	227	Phenylanilinoacetic Ac., Ph.CH(NHPh).CO ₂ H. — Lft. Sbl. at 173-5° in ndl. — I. aq.; e.s. alc.; d.s. eth., bz. — Rapidly heated gives aniline & benzylamine.
238	166	k.10 ⁴ = 2.9	Nitroopianic Ac., (MeO) ₂ ^{2,3} (NO) ₂ ⁴ (CHO) ⁶ .C ₆ H ₄ (CO ₂ H) ¹ . — Pr. fr. h. aq. D.s. c. aq. Salts e.s. aq.
239	166		β -Indole- α -phenylureidopropionic Ac., C ₁₂ H ₁₇ O ₂ N ₃ . — Ndl. d.s. c. aq.; e.s. alc., eth. — Reddens in the light. [Deriv. fr. l-tryptophane.]
240	167	151	6-Amino-m-toluic Ac., NH ₂ C ₆ H ₄ Me.CO ₂ H. — \textcircled{P} Should give N in T. 2.4. — Ndl. fr. aq. D.s. aq.; e.s. h. aq. — Ba \ddot{A} , 10H ₂ O, tbl. v.s. aq.
241	167	181	5-Nitro-m-toluic Ac., NO ₂ C ₆ H ₄ Me.CO ₂ H. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq. Mod. s. h. aq.; v.s. alc., eth. Vol. w. st. — Ba \ddot{A} , 4H ₂ O, warts s. in 325 pt. c. aq.
242	167	192	5-Nitro-3-hydroxybenzoic Ac., NO ₂ C ₆ H ₄ (OH)(CO ₂ H).H ₂ O. — \textcircled{P} Should give T. 2.21.
243	167 u.c. d.	222	d,l- α -Phenylureidobutyric Ac., Et.CH(NH.CO.NHPh). — \textcircled{P} Should give carbylamine odor in T. 2.12. — Ndl. fr. 50 pt. h. aq. E.s. alc.; d.s. eth. — M.p. of corresponding hyantoin (Cf. T. 2.31) 126-7° c. d.
244	168 u.c.	208	α -Phenylureidopropionic Ac., Ph.NH.CO.NH.CHMe.CO ₂ H. — \textcircled{P} Should give carbylamine odor in T. 2.12. — Lust. cryst. fr. h. aq. [A deriv. of α -alanine.]
245	167-70	179	p-Tolioxamic Ac., p-Toluidinoxalic Ac., Me.C ₆ H ₄ .NH.CO ₂ H. — \textcircled{P} Probably gives carbylamine odor in T. 2.12. — Ndl. fr. h. bz. D.s. c. aq.; e.s. h. aq., alc. — Sapn. gives p-toluidine (No. 2.566) & oxalic ac. — Ba \ddot{A} , d.s. scales.
246	169-70d.	k.10 ⁴ = 5	Methyl-syn-oxazolone, β -Oximinobutyric Anhydride, $[\text{ON}:\text{CMe.CH}_2\text{CO}]$. — \textcircled{P} FeCl ₃ sol. gives inky coloration. — Silky ndl. e.s. h. aq., c. alc.
247	169-9.5; 158	241	Phenylphthalamic Ac., Ph.NH.CO.C ₆ H ₄ .CO ₂ H. — \textcircled{P} Probably gives carbylamine odor in T. 2.12. — Lft. alm. i. c. aq.; e.s. alc.; i. eth., chlf.
248	169; 168-9	ROTI sol. in Gen. T. 2.I w. 0.2 cc. alk.	Diphenylcarbazid, CO(NH.NHPh) ₂ . — \textcircled{P} Ppts. Ag slowly in T. 2.30. — Cryst. d.s. h. aq.; e.s. h. alc.; i. eth. — Dil. sol. gives violet-red color w. v. dil. CuSO ₄ sol. [Used as indicator in determination of Cu. — Sample fr. Kahlbaum of Berlin.]
249	170		Sebamic Ac., NH ₂ .CO.C ₆ H ₄ .CO ₂ H. — \textcircled{P} Probably gives NH ₂ odor in T. 2.7. — S. h. aq.; e.s. alc. — Sap. to anilin & sebacic ac. (Vol. I).
250	170c.	222	α -Phenylureidobutyric Ac., Ph.NH.CO.NH.CHEt.CO ₂ H. — \textcircled{P} Should give carbylamine odor in T. 2.12. — Ndl. fr. 50 pt. h. aq.
251	abt. 170-2d.	175	Pr-2-Methylindole-Pr-3-carbonic Ac., Methylketolecarboxylic Ac., Me.C ₆ H ₅ N.CO ₂ H. — \textcircled{P} Vapors evolved on heating in tube above m.p. should give splinter reaction. T. 2.24. — Cryst. v.d.s. aq.; d.s. bz.; e.s. alc., eth.; alm. i. lgr. — Dec. on melting, or even in boiling aq., to 2-methylindole (No. 2.1601).

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
252	171-2d.	208	β -Phenylureidopropionic Ac., Ph.NH.CO.NH.(CH ₂) ₂ .CO ₂ H. — (P) Should give carbalamine odor in T. 2.12. — Cryst. fr. aq. D.s. c. aq. or eth.; v.d.s. chlf., bz.; e.s. alc. — Heating w. aq. in sealed tube at 140° gives CO ₂ & carbanilide (No. 2.2580).
253	172	151	4-Amino-m-toluic Ac., NH ₂ .C ₆ H ₄ .CO ₂ H. — (P) Should give N in T. 2.4. — Thin lft. V.d.s. c. aq.; mod. s. h. aq.; v.s. alc., eth.
254	172-4 w. efferv.		α -Phenylhydrazinepropionic Ac., Ph.NH.NH.CHMe.CO ₂ H. — Ndl. fr. dil. alc. V.d.s. aq., c. alc., eth.; e.s. conc. HCl. — Easily transformed by ammon. Cu sol. to pyruvic-acid-phenylhydrazone.
255	173	201	2-Ethylquinolinecarbonic(4) Ac., Et.C ₆ H ₄ N.CO ₂ H. — Cryst. d.s. c. aq.; e.s. alc., eth. — Ignition w. soda-lime gives 2-ethyl-quinoline (No. 2.1398). — Ag \ddot{A} , d.s. ppt. — BaCl ₂ , cryst. e.s. aq., alc.
256	173-4d.		Dioxytartaric-acid-phenylhydrazone, Phenylizidindioxytartaric Ac., Ph.N ₂ H:C(CO ₂ H).CO.CO ₂ H. — (P) Aq. sol. is colored red by FeCl ₃ . — Ndl. alm. i. c. aq.; v.d.s. eth.; d.s. bz.; e.s. h. alc. — Ag \ddot{A} , or-yel. ppt.
256-I	173	237	p-Nitrophenaceturic Ac., NO ₂ .C ₆ H ₄ .CH ₂ .CO.NH.CH ₂ .CO ₂ H. — (P) Prob. gives T. 2.21. — Long hair-like ndl. fr. h. aq. V.d.s. c. aq., alc., chlf.; e.s. h. aq. or alc.; i. eth., bz. — (P) Sapn. by boiling conc. HCl (T. 2.26) gives p-nitrophenylacetic ac. (No. 2.178), etc. — Ag \ddot{A} , ndl. fr. h. aq., v.d.s. c. aq.
257	173	288	3,5-Dinitrosalicylic Ac., (NO ₂) ₂ .C ₆ H ₃ (OH) ² (CO ₂ H) ¹ . — (P) FeCl ₃ colors sol. blood-red. — Lust. cryst., crystg. w. 1H ₂ O which become yellowish w. time & lose aq. quickly at 110°. M.p. of hydrated & anhydrous acid identical; but changes to 157-8° when remelted. — E.s. c. aq.; v.s. alc., eth. — K \ddot{A} , yellowish cryst. d.s. h. aq.
258	174r.h. d.		Mesoxalic-acid-phenylhydrazone, Ph.NH.N:C(CO ₂ H) ₂ . — (P) Probably gives hydrazone reaction in T. 2.17. — Cryst. v.s. eth.; d.s. chlf.; i. lgr. — Ag \ddot{A} , light yel. ppt.
259	174u.c.	137 k.10 ⁶ = 1.6	m-Aminobenzoic Ac., NH ₂ .C ₆ H ₄ .CO ₂ H. — (P) Gives intense carbalamine odor in T. 2.12! — Cryst., warty aggregates fr. h. aq., coloring sl. w. time. 10 cc. of each of following solvents dis. at 10°; aq., 0.056 g.; 90% alc., 0.220 g.; eth., 0.170 g.; bz. 0.000 g.; chlf., 0.007 g. Odorless. Taste of sat. aq., sol. sweet! — Add 1 drop c. sat. bleaching powd. sol. to 10 cc. of a c. (1:1000) aq. sol. An OYS ² color develops within 30 sec., becoming OS ² (sky light) after 5 min! — (P) Prepare m-picrylaminobenzoic acid, by the procedure given for the corresponding comp. of anthranilic acid (No. 2.148); except that 3 cc. aq. instead of 4 cc. are used in diluting, and that 1 cc. of h. gl. ac. ac. is employed for solvent, and 0.5 cc. for washing. The picryl deriv. is obtained as a Y powd., m.p. 233.0-4.0° u.c.
260	175	k.10 ⁶ = 2.3	N-Dimethylanthranilic Ac., o-NMe ₂ .C ₆ H ₄ .CO ₂ H. — (P) Solution fluoresces blue. — Ndl. fr. bz. s. in 500 pt. c. or 250 pt. h. aq.; e.s. alc., eth.
261	177d.		3-Amino-p-toluic Ac., Homoanthranilic Ac., NH ₂ .C ₆ H ₃ Me.CO ₂ H. — (P) Gives CO ₂ in T. 1.303, & probably N in T. 2.4. — Ndl. fr. alc. D.s. c. aq.; v.s. h. aq., eth., bz. — Boiling w. acetic anhydride gives acetyl deriv., ndl. fr. dil. alc., m.p. 183°.
262	177		4-Aminonaphthoic(1)Ac., NH ₂ .C ₁₀ H ₇ .CO ₂ H. — (P) T. 2.17 probably gives N. — Ndl. fr. h. aq.; d.s. c. aq., bz., lgr. — Dec. v. easily on heating w. HCl to CO ₂ & α -naphthylamine (No. 2.589)!
263	177	212 k.10 ⁶ = 2.6	2,5-Dinitrobenzoic Ac., (NO ₂) ₂ .C ₆ H ₃ .CO ₂ H. — (P) Probably gives T. 2.21. — Separates fr. sat. h. aq. sol. as oil, gradually solidifying to ndl. Mod. s. h. aq. Reduction w. Sn & HCl gives 2,5-diaminobenzoic ac. — Ba \ddot{A} , 4H ₂ O, long 6-sided lft., d.s. h. aq.

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
264	177 (181c.)	241	o-Benzoylamino-benzoic Ac., $(\text{Ph.CO.NH})\text{C}_6\text{H}_4\text{CO}_2\text{H}$. — Ndl. fr. alc. I. aq.; e.s. alc., eth. — Heated w. 25% HCl at 160° gives aniline, benzoic ac. & CO ₂ .
265	177-8	"Behaves as acid"	ab-Dibenzoylphenylhydrazine, $(\text{Ph.CO})\text{NH.N(Ph.CO)(Ph)}$. — (P) Reduces Ag sol. in T. 2.30. — Pr. v.d.s. aq.; s. h. alc. — Readily hydrolyzed by fuming HCl at 100° to benzoic ac. & phenylhydrazine. — $\text{NaC}_6\text{H}_4\text{N}_2\text{O}_2$ (fr. Na + alc. sol. of compound), lust. lft. e.s. aq.
266	178		Methylaspartic Ac., $\text{CO}_2\text{H.CH}(\text{NHMe})\text{CH}_2\text{CO}_2\text{H}$. — Cryst. w. $1\text{H}_2\text{O}$ in monoclinc. pyramids (m.p. 122-3°). 100 pt. c. aq. dis. 2.6 pt. anhydrous acid. — Optically i.
267	178	†	Glyoxime, HO.N:CH.CH:NOH . — Rhomb. tbl. fr. aq. E.s. h. aq., alc., eth. — Sbl. — Ammon. AgNO ₃ gives ppt. of Ag salt, $\text{AgC}_6\text{H}_4\text{N}_2\text{O}_2$. — Try T. 2.17.
268	d.w.m. abt. 178	$k.10^4 = 5$	α -Nitrosopropionic Ac., $\text{Me.C:(NOH).CO}_2\text{H}$. — (P) Dec. by heat w. vigorous evolution of gas without melting. — Reduction by Sn & HCl gives alanine.
269	178	183	2-Nitro-3-hydroxybenzoic Ac., $\text{NO}_2\text{C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})$. — (P) Should give T. 2.21. Taste intensely sweet! — "Yellowish-white" lft. fr. h. aq. — $\text{Ba}\bar{\text{A}}.\frac{1}{2}\text{H}_2\text{O}$, v.s. aq. — Et $\bar{\text{A}}$, pr. v.s. alc., m.p. 124°.
270	179	151 $k.10^4 = 4.6$	N-Methylanthranilic Ac., o-NHMe.C ₆ H ₄ CO ₂ H. — Lft. fr. lgr.
271	179	181	4-Nitro-o-toluic Ac., $\text{Me.C}_6\text{H}_4(\text{NO}_2)(\text{CO}_2\text{H})$. — (P) Should give T. 2.21. — Ndl. fr. dil. alc. V.d.s. h. aq.; v.s. alc. — Ca $\bar{\text{A}}.2\text{H}_2\text{O}$, ndl. v.s. h. aq.
272	179	195	2-Nitro-1, 3-dimethylbenzoic(5) Ac., $(\text{NO}_2)\text{Me}_2\text{C}_6\text{H}_3\text{CO}_2\text{H}$. — (P) Should give T. 2.21. — Cryst. fr. h. aq. V.d.s. h. aq.; e.s. h. alc. — After recrystn. fr. abs. alc. melts at 223° (214-20°)! — Et $\bar{\text{A}}$, e.s. alc., m.p. 72°.
273	179-80u.c. d.	210	β -Phenylureido- α -hydroxypropionic Ac., $\text{Ph.NH.CO.NH.CH}_2\text{CH}(\text{OH}).\text{CO}_2\text{H}$. — Rosettes fr. aq. S. in abt. 200 pt. aq. at 20°, or 16 pt. at 100°; e.s. alc.; alm. i. eth. (Gives no hydantoin on evapn. w. HCl.)
274	179u.c.	212 † $k.10^4 = 3.8$	2,4-Dinitrobenzoic Ac., $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{CO}_2\text{H}$. — (P) Gives heavy dark brown ppt. in T. 2.21. — Taste, v. bitter! — Nearly colorless cryst. fr. aq. — 100 pt. aq. at 25° dis. 1.849 pt. S. in 140 pt. bz. at 30°. — Melts under boiling aq. — Sbl. — Reduction w. Sn & HCl gives p-phenylenediamine (No. 2.877) & CO ₂ .
275	179		Citric-acid-diphenetidid, $(\text{EtO.C}_6\text{H}_4\text{NH.CO})_2\text{C}_6\text{H}_4\text{O.CO}_2\text{H}$. — Powd. d.s. aq. [D.R.P. 87,428.]
276	180-1d.		4-Isonitrosopyrazolone, $[\text{NH.N:CH.C(NOH).CO}] + \frac{1}{2}\text{H}_2\text{O}$. — Cryst. fr. aq. — Ag $\bar{\text{A}}$, deep red ndl. deflagrating at 241°.
277	180-1u.c.	118.5 $k.10^4 = 1.9$	Benzoyl-l-aspartic Ac., $\text{CO}_2\text{H.CH}_2\text{CH}(\text{NH.CO.Ph})(\text{CO}_2\text{H})$. — Cryst. fr. aq. S. in 3-4 pt. h. or 261 pt. aq. at 20°; e.s. alc.; i. eth. — $[\alpha]_D^{20}$, (dis. in 2 mol. KOH sol., D: 1.0592) = +37.4°. Dioxindole. — Cf. No. 2.2366.
—	180 to viol. liq.		1 ^o - α -Cyanocinnamic Ac., $\text{Ph.CH:C(CN)(CO}_2\text{H)}$.
278	180	173	Pyroglutaminic Ac., $\text{CO}_2\text{H}^+ \text{HC.NH.CO.C}_6\text{H}_4^-$. — (P) Strongly heated in tube above m.p. give pyrrole splinter reaction (T. 2.24). — Pr. s. in 19 pt. aq. at 13.5°. — Ag $\bar{\text{A}}$, cryst.; alm. i. aq., m.p. 176-80°.
279	182d.	129	2-Nitro-m-toluic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — (P) Should give T. 2.21. Cryst. — Ba salt alm. in c. aq.; d.s. h. aq.
280	182	181	α -Phenylureido-d, 1- β -phenylpropionic Ac., $\text{Ph.CH}_2\text{CH}(\text{Ph.NH.CO.NH})\text{CO}_2\text{H}$. — Boiled w. 400 pt. dil. HCl give hydantoin of m.p. 173-4° c. (Cf. T. 2.31.)
281	182d.	284	

GENUS I, DIV. A.
(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
282	183	"Alk. sol. deep yel."	5-Nitrocoumarin, $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}:\text{CH.CO}_2$. — Ndl. d.s. h. aq., alc., eth.
283	183-4 u.c.	224	Benzoyl-d, l-phenylalanine, $\text{Ph.CH}_2\text{CH}(\text{NH.CO.Ph}).\text{CO}_2\text{H}$. — Lust. lft.
284	183-4 u.c.		d,l-Ornithuric Ac., α,δ -Dibenzoyl-d,l-ornithin, $\text{Ph.CO.NH.}(\text{CH}_2)_2\text{CH}(\text{CO}_2\text{H})\text{NH.CO.Ph}$. — Mic. ndl. fr. alc. E.s. warm cons. HCl.
285	183-4		Paraglycocholic Ac., $\text{C}_2\text{H}_4\text{NO}_6$. — Pearly lft. alm. i. h. aq. Taste v. bitter! Sol. in alk. or alc. gives glycocholic ac.
—	184 u.c. d.		p-Aminophenol. — Cf. No. 2.963. (Brownish coloration in Gen. T. 2.I.)
287	184		Phenylglycine-o-carbonic-acid-exo-nitrile, $\text{CO}_2\text{H.C}_6\text{H}_4\text{NH.CH}_2\text{-CN}$. — Cryst. fr. dil. alc. Sol. in alk. is pptd. by acids.
288	184-5	177	β -Anilinoisobutyric Ac., $\text{Ph.NH.CH}_2\text{CHMe.CO}_2\text{H}$. — Ndl. s. aq.; e.s. alc., eth.; i. lgr. — Sbl. — Aniline is formed on destructive distn.
289	184-5	"Acid to litmus"	d-Ornithuric Ac., α,δ -Dibenzoyl-d-aminovaleric Ac., $\text{Ph.CO.NH.}(\text{CH}_2)_2\text{CH}(\text{CO}_2\text{H})\text{NH.CO.Ph}$. — (P) Strongly heated emits odor of benzaldehyde & gives woolly sbl. — Ndl. v.d.s. h. aq.; i. eth.; s. acetic-eth. — $[\alpha]_D = +9.2^\circ$ to 9.3° in 10% sol. of Na salt.
290	184.5-5.5 u.c.	137 † $k.10^4 = 1.1$	p-Aminobenzoic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$. — (P) Gives strong carbamyline odor in T. 2.12. — One drop sat. aq. bleaching powd. sol. added to (1:1000) c. aq. sol. of acid gives VR-R color within 30 sec., changing to alm. opaque OS ² (by transmitted light fr. sky) after 5 min. — Odorless ndl. fr. h. aq. 10 cc. of each of following solvents at 10° dis.: aq., 0.033 g.; 90% alc., 1.13 g.; eth., 0.611 g.; chlf., 1.13 g.; bz., 0.006 g. — (P) Prepare the picryl derivative as directed for anthranilic ac. (No. 2.148), except that no water should be added after boiling w. the picryl chloride, & that a second recrystn. fr. 2 cc. boiling gl. ac. ac. should follow. The o-picrylaminobenzoic ac. is obtained in v. lust. Y scales which melt to a red lig. at 287-8° u.c. after darkening & beginning to soften at 283-4°.
291	180-90d.	146	Methyloxaluric Ac., $\text{NH}_2\text{CO.NMe.CO.CO}_2\text{H}$. — (P) Should give NH_2 , MeNH_2 & oxalic ac. on sapon. (T. 2.26).
292	180-90	147	o-Cyanobenzoic Ac., $\text{CN.C}_6\text{H}_4\text{CO}_2\text{H}$. — Ndl. fr. alc.; alm. i. aq., bz.; e.s. h. alc. — Heating above m.p. converts to phthalimide (No. 2.2555)! — Sol. in conc. H_2SO_4 gives phthalamic ac.
293	185	179 $k.10^4 = 2.4$	o-Acetylamoно-benzoic Ac., $\text{Me.CO.NH.C}_6\text{H}_4\text{CO}_2\text{H}$. — Flat ndl. fr. gl. ac. ac. D.s. c. aq.; e.s. eth., bz., h. aq., h. alc. — Easily sapon. by h. HCl (Cf. T. 2.26).
294	185	183	3-Nitro-p-hydroxybenzoic Ac., $\text{NO}_2\text{C}_6\text{H}_4(\text{OH}).\text{CO}_2\text{H}$. — (P) Should give T. 2.21. — Ndl. d.s. h. aq. (Nearly colorless when bone-blackened.) Cryst. w. aq. of crystn. — $\text{BaA}_2\text{H}_2\text{O}$, deep red warts, v.d.s. h. aq.; and also $\text{BaA}_2\text{.4H}_2\text{O}$ as yel. lft. — EtA , light red pr. fr. alc., m.p. 69°.
295	185	199 $k.10^4 = 5$	3-(β)-Phenylpyridine-Bz-2-Carbonic Ac., $\text{C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Ndl. fr. aq. Dist. undecd. D.s. c. aq.; more s. alc. — Ignition w. soda-lime gives 3-phenylpyridine. Boiled w. CrO_3 & H_2SO_4 gives nicotinic ac. (No. 2.410).
296	abt. 186d.		4-Methylpyridine-2,3-dicarbonic Ac., $\text{Me.C}_6\text{H}_4\text{N}(\text{CO}_2\text{H})_2$. — (P) Aq. sol. gives yel. color w. FeSO_4 . — Cryst. s. in 118.6 pt. aq. at 10°; d.s. alc., eth., bz. — $\text{Ba(C}_2\text{H}_5\text{O}_2)_2$ gives cryst. ppt.
297	186d.	$k.10^4 = 7.5$	2,4-Dimethylpyrrole-3-carbonic Ac., $\text{Me}_2\text{C}_4\text{H}_3\text{N.CO}_2\text{H}$. — (P) Vapors evolved on ignition in tube probably give pyrrole splinter reaction, T. 2.24. — Cryst. flocks. — Gives CO_2 & dimethylpyrrole (No. 2.2759) in melting.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
298	186d.		m-Hydrazinobenzoic Ac., $\text{NH}_2\text{NH.C}_6\text{H}_4\text{CO}_2\text{H}$. — \textcircled{P} Reduces Fehling's sol. — Faintly yellowish lft. D.s. h. aq., alc. (Reacts strongly acid.)
299	186-7 u.c.	165	4-Aminomesitylenic Ac., $(\text{NH}_3)^4(\text{Me}_2)^{1,2}\text{C}_6\text{H}_3\text{CO}_2\text{H}$. — Ndl. fr. alc.
300	186w. frothing		Pyruvic-acid-phenylhydrazone, $\text{Ph.NH.N:CM}_e\text{CO}_2\text{H}$. — \textcircled{P} Sol. in conc. H_2SO_4 at first yel., later deep red. — Hydrolyzed by boiling HCl to phenylhydrazine (Cf. T. 2.17). [M.P. given obtained by starting with bath at 160°. Variable w. rate of heating.]
—	187		Tetramethylsuccinimide. — Cf. No. 2.2405.
302	186-7.5, 183	173 $k.10^4 = 1.3$	Quinoline-8-carbonic Ac., Cinchoninic Ac., $\text{C}_8\text{H}_5\text{N.CO}_2\text{H}$. — \textcircled{P} FeSO_4 added to sol. of $\text{NH}_4\bar{A}$ gives dark purple-red color which soon disappears, a purplish red or brown cryst. powd. pptg. — Ndl. D.s. c. aq., alc.; e.s. ac., alk. — $\text{Ag}\bar{A}$, cryst. ppt. fr. $\text{NH}_4\bar{A}$ by AgNO_3 .
302-1	187		β -Naphthoquinolinecarboxylic Ac., $\text{C}_{12}\text{H}_8\text{N}(\text{CO}_2\text{H})$. — Mic. ndl. fr. gl. ac. — \textcircled{P} Heating at 200° dec. to CO_2 & β -naphthoquinoline (No. 2.716). — $\text{Ba}\bar{A}.4\text{H}_2\text{O}$, floc. ppt. i. aq., becoming cryst. after boiling w. aq.
303	187d.	173	Levulinic-acid-semicarbazone, $\text{NH}_2\text{CO.NH.N:CM}_e(\text{CH}_2)_2\text{CO}_2\text{H}$. — \textcircled{P} Probably gives T. 2.17. — Ndl. fr. much alc.
304	187	179 † $k.10^4 = 2.2$	Hippuric Ac., Benzoylglycine, $\text{Ph.CO.NH.CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Heated rapidly above m.p. in glass tube reddens, gives subl. (benzoic ac.), & emits odor of bitter almonds. — 4-sided pr. w. 2 or 4 terminal pyramid planes (rhombic system) fr. h. sol. E.s. h. aq.; s. in 600 pt. c. aq. at 0°; e.s. alc., acetic eth. (used for extraction fr. urine); i. bz., CS_2 or pet-eth. (Last named solvent used in separation fr. benzoic ac.) Odorless. Not vol. w. st. — \textcircled{P} Heat together on the water-bath for 30 min., 1 mol. of substance, 1 mol. benzaldehyde, 1 mol. cryst. sodium acetate, & 3 mol. anhydrous sodium acetate. After cooling, wash the cryst. paste w. alc. and recryst. fr. alc. The product, the lactimide of benzoylaminocinnamic ac., $\text{Ph.CO}^{\text{C}}\text{N.C:CHPh.CO}^{\text{C}}$, melts at 165-6°. (Spiro, Z. physiol. Chem., 28, 177 (1899).) [Abundant in urine of herbivorae.]
305	187	267	Opianic-acid-semicarbazone, $(\text{MeO})_2\text{C}_6\text{H}_3(\text{CH:N.NH.CO-NH}_2)\text{CO}_2\text{H}$. — \textcircled{P} Should give T. 2.17. — Cryst. fr. gl. ac. ac. E.s. alc.; v.d.s. bz., eth.
306	186-7; 188-9	197	3-Nitro-4-methoxybenzoic Ac., $\text{NO}_2\text{C}_6\text{H}_3(\text{OMe}).\text{CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Cryst. v.d.s. h. aq.; e.s. h. alc.; e.s. eth. — $\text{Pb}\bar{A}_2$, ndl., exploding violently on ignition. — $\text{Me}\bar{A}$, m.p. 108° or 109-10°.
307	188r.h., complete d.	209	Kyanuric Ac., Oxalyianthranilic Ac., o-CO ₂ H ₂ C ₆ H ₄ NH.CO.CO ₂ H. — \textcircled{P} FeCl_3 colors dil. aq. sol. carmine red. — Ndl. fr. eth. (Loses 1 mol. cryst. aq. at 100°) S. in 890 pt. aq. at 10°; v.s. h. alc., eth. — Ignition w. CaO gives aniline. Boiling w. dil. ac. or alk. gives anthranilic (No. 2.148) & oxalic acids.
308	188	224	o-Nitrohippuric Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CO.NH.CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Lft. e.s. alc., h. aq.; d.s. eth.
309	189-90	181	2-Nitro-p-toluic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Monoclin. pr. fr. alc. D.s. c. aq.; e.s. alc., h. aq. — $\text{Ba}\bar{A}.4\text{H}_2\text{O}$, ndl. v.d.s. c. aq., e.s. h. aq.
310	191	151	6-Amino-o-toluic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — \textcircled{P} Should give N in T. 2.4. — Glassy ndl. fr. aq. Mod. s. c. aq.
311	190-1; 194-5	193 $k.10^4 = 2.6$	Acetylphenylglycine, $\text{CH}_3\text{CO.NH.CH}_2\text{CO}_2\text{H}$. — Pearly lft. fr. aq. D.s. c. aq., eth., chlf., lgr., bz.; e.s. alc. — $\text{Cu}\bar{A}_2$, green cryst. powd. i. aq.

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
312	188; 194	226	2,5-Dimtro-p-toluic Ac., $(NO_2)_2C_6H_3Me.CO_2H$. — \textcircled{P} Probably gives T. 2.21. — Pr. V.d.s. c. aq.; v.s. alc., eth.; alm. i. chlf., bz., lgr.
313	192-3	228	Hydrazobenzene-p-carbonic Ac., $Ph.NH.NH.C_6H_4.CO_2H$. — Ndl. fr. dil. alc. — Treated in alc. sol. in the cold w. $SnCl_4$ & HCl gives benzidine (No. 2.840) & CO_2 .
315	193-5d.		α -Naphthylglyoxylic-acid-oxime, $C_{10}H_7.C(NOH).CO_2H$. — On distn. in <i>vacuo</i> dec. to aq., CO_2 , & α -naphthyl cyanide.
316	195d. abt. 210r.h.	160 $k.10^6 = 3.1$	Succinic Ac., $NH_2.CO.NH.CO.(CH_2)_2.CO_2H$. — \textcircled{P} Should give NH_2 in T. 2.7-a. — Scales fr. aq. Alm. i. c. aq., alc., eth.; s. without decn. in conc. H_2SO_4 . — Sap. (Cf. T. 2.26) to succinic ac. (Vol. I).
317	194-6	193 $k.10^7 = 4.1$	β -p-Toluidinoisobutyric Ac., $Me.C_6H_4.NH.CH_2.CHMe.CO_2H$. — Pr. fr. alc.; d.s. lgr., eth., bz.
318	195u.c. d.	194	Phenylureidoacetic Ac., $Ph.NH.CO.NH.CH_2.CO_2H$. — Cryst. fr. aq.; S. in 70 pt. boiling aq.; mod. s. h. alc.; alm. i. eth., chlf., bz. [Deriv. of glycine in T. 2.31.]
319	195	195	6-Nitro-1,3-dimethylbenzoic(4) Ac., $(NO_2)Me_2C_6H_3.CO_2H$. — \textcircled{P} Should give T. 2.21. — Ndl. d.s. c. aq.; e.s. alc., eth., chlf. — $BaA_2.9H_2O$, ndl. e.s. aq.
320	192; 198-9	201 $k.10^6 = 3.3$	α -Naphthylglycine, $C_8H_7.NH.CH_2.CO_2H$. — Silky ndl. fr. dil. alc. V.d.s. aq., eth., lgr. — Heating at 130° gives anhydride, scales fr. alc., m.p. 274-5°.
321	196	151	4-Amino-o-toluic Ac., $NH_2.C_6H_4.Me.CO_2H$. — Pr. fr. aq. D.s. c. aq.; e.s. h. aq.; v.s. h. alc.
322	196-7	193	m-Nitrocinnamic Ac., $NO_2.C_6H_4.CH:CH.CO_2H$. — \textcircled{P} Should give T. 2.21. — [Nearly colorless when freshly boneblacked & recrystd. Specimen fr. Kahlbaum, YT3.]
323	197-8		Benzylaminoacetic Ac., $Ph.CH_2.NH.CH_2.CO_2H$. — Ndl. fr. aq. — $Cu\bar{A}$, (dried over H_2SO_4), ppt.; small dark blue pr. fr. h. aq.
324	197-8d.	$k.10^2 = 1.7$	Isonitrosodiketohydrindene, $C_6H_4:(CO)_2:C:NOH$. — Lft. fr. gl. ac. ac. S. alk.
325	195-200d.	189 $k.10^6 = 2.2$	2-Methylindole-3-acetic Ac., $[C_6H_5.NH.CMe:C(CH_2.CO_2H)]$. — \textcircled{P} Vapors evolved on ignition in tube (2,3-dimethylindole) should give splinter reaction, T. 2.24. — Cryst. fr. acetone. D.s. h. aq., chlf.; s. h. alc.
326	198	$k.10^6 = 4$	i-Glutaminic Ac., $CO_2H.CH_2.CH_2.CH(NH_2).CO_2H$. — \textcircled{P} Gives the pyrrole splinter test as described for the d-acid (No. 2.357). — Tetrahedral cryst. or spheroids fr. aq. S. in 66.7 pt. aq. at 20°; d.s. alc., eth., lgr. — Is resolved into the l- & d- acids by repeated recryst. T. 2.5 gives 9.52% amino N.
—	198-9	187	5-Aminonaphthoic(1) Ac. — Cf. No. 2.367.
328	198; 181d.	191	p-Nitrophenylpropionic Ac., $NO_2.C_6H_4.C:CO_2H$. — \textcircled{P} Probably gives T. 2.21. — Ndl. fr. alc. D.s. aq., bz., chlf.; e.s. h. alc. — Adds 2 atoms of Br.
325	197-9u.c.	227	2-Nitroveratric Ac., $(NO_2)^2C_6H_3(MeO)^{3,4}(CO_2H)^1$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq. — D.s. c. aq., bz., pet-eth.; otherwise e.s.
330	198		Isogeronicsemicarbazone, $Me:N.NH.CO.NH_2.(CH_2)_2.CMe_2-CO_2H$. — \textcircled{P} Probably gives T. 2.17. ("Slowly split by alc. H_2SO_4 ") — Lft. fr. much alc.
331	199-200d.		4-Amino- α -toluyllic Ac., $NH_2.C_6H_4.CH_2.CO_2H$. — \textcircled{P} Should give N in T. 2.4. — Pearly lft. I. c., mod. s. h. aq. — Warmed w. 2.5 pt. acetic anhydride at 100° gives acetyl deriv. of m.p. 168-70°.
332	200; 203-4d.		6-(p)-Hydroxyquinolinecarboxylic Ac., $HO.C_6H_4.C_6H_3.N.CO_2H$. — \textcircled{P} Colored blood red by $FeCl_3$. — Yellowish white mic. cryst. flocks fr. much h. aq. Dec. to 6-hydroxyquinoline & CO_2 in melting. — $Ag\bar{A}$, ppt. — $B_2H_2PtCl_6.2H_2O$, dark yell. cryst. powd., d.s. h. aq.

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
333	200d.	223	Anhydro-o-aminohemipinic Ac., $[C_6H(MeO)_2(CO_2H).CO.NH]^2$. — Ndl. fr. h. aq. S. in 80% alc.; s. in conc. H_2SO_4 & reproto. by aq. — $Ba\bar{A}_2 \cdot 6H_2O$, fine ndl. — $Me\bar{A}$ (fr. MeOH & HCl), m.p. 127°.
334	200	256	3,6-Dinitrophthalic Ac., $(NO_2)_2.C_6H_3.(CO_2H)_2$. — \oplus Should give T. 2.21. — Fine ndl. fr. eth. + lgr. E.s. aq., alc., eth. — $Ba\bar{A}$, cryst. ppt. fr. h. aq. sol. + $Ba(CH_3CO_2)_2$.
335	197-9sl.d.; 200-3	"Alk. sol. bright yel."	Disalicylamide, $(HO.C_6H_4.CO)_2.NH$. — \oplus Should give NH ₂ in T. 2.7. — $FeCl_3$ colors alc. sol. red! — Yellowish white asbestos-like ndl. I. aq.; s. h. alc.; d.s. eth. — Sap. to salicylic ac. (Vol. I).
336	abt. 200-2 u.c.	298	d,l- α,β -Dibenzoyldiaminopropionic Ac., $Ph.CO.NH.CH_2.CH-(NH.CO.Ph).CO_2H$. — Ndl. fr. alc. — V.d.s. aq.; e.s. alc.; alm. i. eth.; i. pet-eth.
337	201	153	3-Amino-4-hydroxybenzoic Ac., $NH_2.C_6H_3(OH).CO_2H$. — \oplus Should give N in T. 2.4. — Pr. w. 1H ₂ O fr. aq. S. h. aq.; d.s. h. alc.; alm. i. eth., chlf., bz. — [An azo-color component.]
338	201.5 (206.5c.)		Hippurylglycine, $Ph.CO.NH.CH_2.CO.NH.CH_2.CO_2H$. — Cryst. fr. aq. D.s. c. alc.; more s. dil. alc. than in aq.; i. c. eth., chlf., bz. — Sapd. to benzoic ac. (Vol. I) & glycine (No. 2.2568) by boiling dil. acids or conc. KOH (T. 2.28).
339	202		α -Naphthindolecarboxylic Ac., $[NH.C_6H_4.CH:C](CO_2H)$. — Silvery lft. fr. aq. V.d.s. h. aq.; e.s. alc., eth.; d.s. bz.; lgr. (Does not give pine splinter pyrrole react.)
340	202	212 $k.10^4 = 8.5$	2,6-Dinitrobenzoic Ac., $(NO_2)_2.C_6H_3.CO_2H$. — \oplus Should give T. 2.21. — Ndl. e.s. h. aq. — \oplus Distr. gives m-dinitrobenzene (No. 2.3016) & CO ₂ . — $Ba\bar{A}_2 \cdot 2H_2O$, ndl. v.s. c. aq.
—	202-3		\leftarrow -Aminocaproic Ac. — Cf. No. 2.2464.
341	203-5; d.w. efferv. abt. 205.r.h.	"Weak acid react. on litmus"	d,l-Pyrrolidine- α -carbonic Ac., $C_6H_5N.CO_2H$. — \oplus When strongly heated in tube, vapors give red coloration in pyrrole react (T. 2.24). — Ndl. fr. alc. + eth. (Must be well dried to remove cryst. aq. before m.p. determination.) — V.s. c. aq.; e.s. c. alc.; d.s. chlf., bz.; i. eth. Taste, sweet. — $Cu\bar{A}_2 \cdot 2H_2O$, blue lft. becoming violet on drying; e.s. h.; d.s. c. aq.
342	abt. 203d.	161 $k.10^4 = 1.8$	Pr-2-(α)-Indolecarboxylic Ac., $C_6H_5N.CO_2H$. — \oplus Vapors evolved on ignition in t.t. (above 230°) give pyrrole splinter react. (T. 2.24). — Ndl. fr. h. aq.; silky lft. fr. bz.; r.d.s. h. aq., bz.; e.s. alc., eth. — $Ba\bar{A}_2$, r.d.s. h. aq., pptg. in lust. lft. on cooling.
343	203-4u.c.	212 † $k.10^4 = 1.6$	3,5-Dinitrobenzoic Ac., $(NO_2)_2.C_6H_3.CO_2H$. — \oplus Gives heavy brownish ppt. in T. 2.21. — Thin faintly yellowish tbl. fr. h. aq. S. in 53 pt. boiling aq.; v.s. alc., gl. ac. ac.; d.s. eth. — Odorless. Taste slightly bitter. — Evaporation of 2 drops of h. sat. aq. sol. on microscopic slide gives transparent trapezia (depicted in Rec. trav. chim. 15, 278-80). — \oplus Prepare the methyl ester, m.p. 107.5°, following directions of T. 1.819-c!
344	203-4		Camphoranilic Ac., $CO_2H.C_6H_4.CO.NH.Ph$. — \oplus T. 2.12 probably gives carbylamine odor. — Ndl. fr. h. chlf., dil. alc.; d.s. h. aq.; e.s. alc., eth.
345	203d.	Alk. sol. prob. colored	2,4,6-Hexanitrocarbanilide, $CO_2[NH.C_6H_4(NO_2)_2]_2$. — \oplus E.s. in ammonia w. red color. — Colorless mic. cryst. e.s. h. nitrobenzene; d.s. h. gl. ac. ac. Warming w. ammonia quickly yields cryst. of trinitroaniline, m.p. 186°. — K salt, scarlet pr. v.s. aq.; explosive; gives trinitroaniline (No. 2.3574) on boiling w. aq.
347	205-6	$k.10^4 = 1$	o-Phthalylamino-acetic Ac., $C_6H_4(CO_2)_2:N.CH_2.CO_2H$. — Lust. lft. fr. aq. V.d.s. c. aq.; s. h. aq.; i. c. alc., eth., chlf., lgr. — $Ag\bar{A}$, ppt., cryst. fr. h. aq. — V.e. sapd. by 2 mol. NaOH sol. to glycinephthaloylic ac., $CO_2H.C_6H_4.CO.NH.CH_2.CO_2H.H_2O$, lft., m.p. 105-6°.

GENUS I, DIV. A.
(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
348	205.5 u.c.	240	2,4-Dinitromesitylenic Ac., $(NO_2)_2^{\text{Me}}(\text{Me}_2)_1^{\text{C}_6\text{H}_4}(\text{CO}_2\text{H})^4$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq. E.s. alc.; d.s. aq., lgr. — Reduced by Sn & HCl to 2,4-diamino-m-xylene (No. 2.643).
349	205	272	o-Hydrazobenzoic Ac., $\text{CO}_2\text{H.C}_6\text{H}_4.\text{NH.NH.C}_6\text{H}_4.\text{CO}_2\text{H}$. — Lft. fr. alc. — Oxidize to yellow o-azobenzoic ac. by adding NaNO_2 in x.s. to warm sol. in gl. ac. ac. Product pptd. by aq. melts at 267–8°.
350	205	319	Nitropodocarpic Ac., $\text{C}_{17}\text{H}_{21}\text{O}_5.\text{NO}_2$. — \textcircled{P} Should give T. 2.21. — Lust. cryst. fr. alc. I. aq.; d.s. c. alc.; v.d.s. chlf., bz. — $\text{K}_2\text{A}.5\frac{1}{2}\text{H}_2\text{O}$, red-yel. ndl. w. green metallic reflections, e.s. aq., alc.
350-I	abt. 205		Dianilinosuccinic Ac., $\text{CO}_2\text{H.CH(NH.Ph).CH(NH.Ph).CO}_2\text{H}$. — \textcircled{P} Mixed w. ZnCl_2 , moistened w. HCl, & fused, gives fuchsine-red mass. — Sinters fr. abt. 190°. Alm. i. aq., bz.; more s. alc., warm eth., chlf. Lft. fr. h. gl. ac. ac. Unstable.
351	206	117 $k.10^4 = 2.3$	Acetylglycine, Aceturic Ac., $\text{Me.CO.NH.CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Colored red by FeCl_3 . — Long pointed cryst. fr. aq. 100 pt. aq. at 15° dis. 2.7 pt.; s. alc.; i. eth. — Conc. HNO_3 evolves N_2O in cold. — $\text{CuA}.4\frac{1}{2}\text{H}_2\text{O}$, sky-blue ndl. e.s. aq., alc.; loses $3\frac{1}{2}\text{H}_2\text{O}$ at 105°.
352	206	226	4,6-Dinitro-o-toluic Ac., $(NO_2)_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$. — \textcircled{P} Should give T. 2.21. — Ndl. fr. aq.; mod. s. h. aq. — $\text{BaA}.2\text{H}_2\text{O}$, v.s. aq. — MeA , ndl. fr. alc., m.p. 73–4°.
353	207	$k.10^4 = 1.5$	3-Phenylpyridine-Bz-2, Py-2-dicarboxylic Ac., $\text{CO}_2\text{H.C}_6\text{H}_4\text{C}_6\text{H}_3\text{N}(\text{CO}_2\text{H})_2$. — \textcircled{P} Aq. sol. is colored or.-red by FeSO_4 ; also gives light yel. cryst. w. Br-Aq. — Cryst. fr. aq. in clear short pr. w. $1\text{H}_2\text{O}$ (lost at 100°). D.s. c. aq., alc. v.d.s. eth., bz. — Distn. gives CO_2 & phenylpyridinecarboxylic ac. (No. 2.295).
354	207		2-Methylpyridine-5-carboxylic Ac., $\text{Me.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Pr. v.s. aq., alc. — Ignition w. Ca(OH)_2 gives methylpyridine (No. 2.1153). — Oxidation w. KMnO_4 gives cinchomeronic ac. (No. 2.470).
355	207	†	Gallaniide, $\text{CH}_2(\text{OH})_2\text{CO.Ph}$. — Lust. lft. w. $2\text{H}_2\text{O}$ fr. aq. containing SO_2 . S. in 1500 pt. c. aq. (Sol. deep yellow in Gen. T. 2.I.)
356	d.208.5; 191.5d.	131 $k.10^4 = 4$	Pyrrole-2-carboxylic Ac., $\text{C}_4\text{H}_5\text{N.CO}_2\text{H}$. — \textcircled{P} Gives splinter pyrrole react. (T. 2.24) when ignited in tube. — Lft. fr. aq. showing metallic green color after drying in air. S. aq., alc.; mod. s. eth.
357	208 u.c. r.h. d.w. efferv.	$k.10^4 = 4.1$ †	d-Glutaminic Ac., $\text{CO}_2\text{H}(\text{CH}_2)_2\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$. — \textcircled{P} Ignite 0.05 g. sharply in small dry l.t. & insert into vapors a splinter of soft pine that has been soaked for 30 sec. in conc. HCl. Splinter becomes red (T. 2.24) fr. action of pyrrole vapors. — Cryst. s. in 100 pt. c. aq.; alm. i. alc. — Odorless. Taste, sour. — $[\alpha]_D^{25} = +10.2^\circ$ for aq. sol. when $p = 2$. — Gives 9.52% amino nitrogen in T. 2.5. Behaves as if monobasic in titration of Gen. T. 2.I. [Important proteolytic product.]
358	208		Phenylparabanic Ac., $^{\text{OCO.NPh.CO.CO.NH}^2}$. — Silky ndl. e.s. alc., eth., h. aq.
359	208.4 u.c. w. efferv.	297	d,l- α,β -Diphenylureidopropionic Ac., $\text{Ph.NH.CO.NH.CH}_2\text{-CH}(\text{Ph.NH.CO.NH}).\text{CO}_2\text{H}$. — I. c. aq.; s. h. alc.
360	210 u.c. d. w. efferv.	89 † $k.10^4 = 8.5$	Oxamic Ac., $\text{NH}_2\text{CO.CO}_2\text{H}$. — \textcircled{P} Aq. sol. gives no ppt. w. CaCl_2 ; but if first made strongly alk. w. NaOH & boiled some seconds evolves NH_3 (T. 2.7-a), & after acidification w. ac. ac. & addition of CaCl_2 gives abundant ppt. of calcium oxalate. — Cryst. powd., on cooling aq. sol. saturated at 70–80°. S. in 71 pt. aq. at 14°; alm. i. abs. alc. — Odorless. Intensely sour.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
361	210d.		o-Nitrosobenzoic Ac., $\text{NO.C}_6\text{H}_4.\text{CO}_2\text{H}$. — \oplus Solutions in h. alc., ac. ac., or ammonia are green! — Cryst. fr. abs. alc.; v.d.s. eth., bz.
362	210-1d.		3,4-Diaminobenzoic Ac., $(\text{NH}_2)_2\text{C}_6\text{H}_3.\text{CO}_2\text{H}$. — Lft. d.s. c. aq.; s. h. aq. Dec. on distn. to CO_2 & o-phenylenediamine (No. 2.751). — $\text{B}_2\text{H}_2\text{SO}_4$, lft. v.d.s. h. aq.
363	210d.	Alk. sol. yellow	2-Isonitrosohydindone, $[\text{CO.C}_6\text{H}_4.\text{CH}_2.(\text{NOH})\text{C}]$. — Ndl. fr. h. alc. — NaA , light canary yel. pr. fr. c. alc., changed by heat to labile scarlet modification.
364	abt. 210d.	231 $k.10^4 = 1$	Acetylphenylglycine-o-carbonic Ac., $(\text{Me.CO})\text{N}[\text{C}_6\text{H}_4(\text{CO}_2\text{H})].-\text{CH}_2.\text{CO}_2\text{H}$. — \oplus Heated at 100 $^{\circ}$ w. fuming H_2SO_4 gives blue sol. of indigosulphonic ac. — Cryst. fr. aq. or MeOH.
365	210u.c.	150	4,6-Dinitromesitylenic Ac., $(\text{Me}_2)^{1,2}(\text{NO}_2)^{4,6}.\text{C}_6\text{H}_2(\text{CO}_2\text{H})$. — \oplus Should give T. 2.21. — Colorless pr. e.s. alc., eth.; mod. s. aq., chlf. — Reduced by Sn & HCl to 4,6-diamino-m-xylene (No. 2.762).
366	211	181	6-Nitro-m-toluic Ac., $\text{NO}_2.\text{C}_6\text{H}_4.\text{Me.CO}_2\text{H}$. — \oplus Should give T. 2.21. — Cryst. alm. i. c. aq.; v.d.s. h. aq. — $\text{BaA}_2.4\text{H}_2\text{O}$, ndl. e.s. aq. — MeA , ndl. fr. alc., m.p. 72 $^{\circ}$.
367	210; 211-2		5-Aminonaphthoic(1) Ac., $\text{NH}_2.\text{C}_{10}\text{H}_7.\text{CO}_2\text{H}$. — \oplus Prob. gives N in T. 2.4. — Ndl. fr. alc. D.s. h. aq.; v.d.s. eth.; more s. alc. — Sbl. in ndl., m.p. 196 $^{\circ}$. — Becomes violet in melting.
368	210-2	195	4-Nitromesitylenic Ac., $(\text{NO}_2)^4(\text{Me}_2)^{1,2}.\text{C}_6\text{H}_2(\text{CO}_2\text{H})^2$. — \oplus Should give T. 2.21. — Ndl. fr. h. aq. D.s. c. aq.; s. alc., eth. — $\text{BaA}_2.4\text{H}_2\text{O}$, ndl. v.s. c. aq. — EtA , tbl. fr. alc., m.p. 64-5 $^{\circ}$.
369	210-2	207 $k.10^4 = 2.2$	Acetyl-o-tolylglycine, $\text{MeC}_6\text{H}_4.\text{N}(\text{Me.CO}).\text{CH}_2.\text{CO}_2\text{H}$. — Tbl. fr. dil. alc. D.s. c. eth., lgr., bz.
370	211	389	Dibenzoyl-l-tyrosine, $\text{Ph.CO.O.C}_6\text{H}_4.\text{CH}_2.\text{CH}(\text{Ph.CO.NH}).-\text{CO}_2\text{H}$. — Ndl. fr. gl. ac. ac. I. c. aq.; v.d.s. h.aq.; e.s. alc.; d.s. l.z., eth. Dextrorotatory. — $\text{K}\bar{A}$, ndl. e.s. aq.
371	abt. 212d.	$k.10^4 = 3$	Anildiacetic-o-carbonic Ac., $\text{CO}_2\text{H.C}_6\text{H}_4.\text{N}(\text{CH}_2.\text{CO}_2\text{H})_2$. — Lft. fr. aq.
372	d.210-3	$k.10^4 = 1.1$	2,5-Dimethylpyrrole-3-carbonic Ac., $\text{Me}_2\text{C}_5\text{H}_3.\text{CO}_2\text{H}$. — \oplus Ignition in t.t. yields vapors of dimethylpyrrole which should give T. 2.24. — $\text{NH}_2\bar{A}$ is colored red by FeCl_3 .
373	212-3d.		Phenyltetrazole, Benzenyltetrazotic Ac., $[\text{PhC:N.N:N.NH}]^+$. — Long ndl. fr. aq. Alm. i. c. aq.; s. alc.; d.s. eth.; v.d.s. bz., lgr. — Heated w. conc. HCl at 220 $^{\circ}$ gives aniline, NH_2 , N, & CO_2 .
374	211-3d.		Phenyliminodiacetmonoanilide, $\text{CO}_2\text{H.CH}_2.\text{NPh.CH}_2.\text{CO.NH}-\text{Ph}$. — Silky ndl. fr. dil. alc. Alm. i. aq.; e.s. alc. — Acetic anhydride in x.s. gives diphenyldiacipiperazine (No. 2.2612), m.p. 158 $^{\circ}$.
375	d.213-5w.m.		i-Asparagine, $\text{NH}_2.\text{CO.CH}(\text{NH}_2).\text{CH}_2.\text{CO}_2\text{H.H}_2\text{O}$. — \oplus Should give NH_2 in T. 2.7. — Tasteless triclin. tbl. S. c. aq.; i. alc. or eth.
376	214-5d.		3,2'-Bipyridyl-2,3'-dicarbonic Ac., $[-\text{C}_6\text{H}_4(\text{CO}_2\text{H})]_2$. — \oplus FeSO_4 colors aq. sol. red. — Tbl. w. $2\text{H}_2\text{O}$ (lost at 110 $^{\circ}$) fr. aq. — D.s. c. aq.; e.s. c. alc.; v.d.s. eth., bz. — Ignition w. CaO gives 3,2'-bipyridyl (No. 2.1438-1).
377	213d.		3,3'-Bipyridyl-2,2'-dicarbonic Ac., $[-\text{C}_6\text{H}_4(\text{CO}_2\text{H})]_2$. — \oplus Aq-sol. colored dark or yellow by FeSO_4 . — Granules fr. aq. Anhydrous at 105 $^{\circ}$. — D.s. c. aq., alc.; e.s. h. aq.; alm. i. eth. — Gives CO_2 & 3,3'-bipyridyl (No. 2.1444) in melting.
378	215		3-Methylpyrazolone, $[\text{NH.N:CM}_2\text{CH}_2\text{CO}]^+$. — Glassy pr. fr. aq. Mod. s. c. aq.; d.s. h. alc., dil. acids & Na_2CO_3 , sol. — Yellow isonitroso deriv., m.p. 194 $^{\circ}$, may be prepared by passing nitrous fumes into aq. sol. (J. prakt. Chem. [2] 50, 512.)

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
379	215	128	Dinitroisophthalic Ac., $(NO_2)_2C_6H_3(CO_2H)^{1,2}$. — \textcircled{P} Should give T. 2.21. — Cryst. fr. aq. w. 5H ₂ O. D.s. c. aq.; more s. alc. — Ba \bar{A} .7H ₂ O, lft., r.d.s. c. aq. — Ag \bar{A} , voluminous ppt.
380	abt. 215d.	$k \cdot 10^4 = 2.3$	Phenylglycine- α -carbonic Ac., $CO_2H \cdot C_6H_4 \cdot NH \cdot CH_2 \cdot CO_2H$. — Ndl. fr. MeOH. S. h. aq.; i. alc., eth., gl. ac. ac.; alm. i. chlf., bz. — Rub together in mortar 0.1 g. substance, 0.3 g. KOH, & 6 drops aq. Fuse cautiously! over v. small flame in t.t. until color of fusion has passed through lemon yellow to bright orange. Diss. in 15 cc. c. aq. & pass current of air through sol. A ppt. of indigo blue will appear.
381	215	217	8-Nitronaphthoic(1) Ac., $C_{10}H_8(NO_2) \cdot CO_2H$. — \textcircled{P} Prob. gives T. 2.21. — Pr. fr. alc. S. in 2590 pt. c. aq. or 21.5 pt. alc.; e.s. h. gl. ac. ac.; d.s. eth., bz. — Ba \bar{A} .6H ₂ O, fine yel. ndl., v.s. aq. — Pb \bar{A} .H ₂ O, yel. pr. s. in 248 pt. c. aq. — Et \bar{A} , yel. octahedra fr. alc., m.p. 68—9°.
382	215	262	β -Dinitro- α -naphthoic Ac., $(NO_2)_2 \cdot C_{10}H_8 \cdot CO_2H$. — \textcircled{P} Prob. gives T. 2.21. — Silky cryst. fr. alc. D.s. h. aq.; v.d.s. bz., lgr.; e.s. alc., h. eth. — Et \bar{A} , ndl. fr. alc., m.p. 137°.
383	216	"Reacts ac."	Nitroacetonitrile, $(NO_2)CH_2CN$. — \textcircled{P} Prob. gives T. 2.21. — Cryst. fr. h. aq.; i. c. aq., alc., eth. — Boiled w. Ba(OH) ₂ gives NH ₃ . — Hg(N ₂ O ₂) ₂ ppts. Hg(C ₂ HN ₂ O ₂) ₂ fr. aq. sol.
384	216	153	4-Amino-3-hydroxybenzoic Ac., $NH_2 \cdot C_6H_4(OH) \cdot CO_2H$. — \textcircled{P} FeCl ₃ gives dark blue color or brown ppt. in sol. — Lft. fr. dil. alc. E.s. alc. Darkens before melting.
385	217	$k \cdot 10^4 = 2$ 147	m-Cyanobenzoic Ac., $CN \cdot C_6H_4 \cdot CO_2H$. — \textcircled{P} Prob. gives NH ₃ in T. 2.7. — Mic. ndl. fr. aq. E.s. alc., eth., h. aq. — Sbl. undec. — Ignition of Ca \bar{A} w. CaO gives benzonitrile (No. 2.2781). — \textcircled{P} Sapon. by boiling NaOH to isophthalic Ac. (T. 2.26 & 1.318).
386	abt. 218d. in sealed tube		Pr-3(β)-Indolecarboxylic Ac., $C_8H_7N \cdot CO_2H$. — \textcircled{P} Fecal odor of indole is noticed on boiling w. aq. Gives splinter pyrrole react. slowly in T. 2.24-a. — Lft. fr. h. aq. D.s. h. aq., bz.; more s. alc., eth.; alm. i. lgr. (M.p. varies w. mode of heating). [Among indole derivatives in urine of herbivore.]
387	218	204	Acetylamino-salicylic Ac., $Me \cdot CO \cdot NH \cdot C_6H_4(OH) \cdot (CO_2H) \cdot H_2O$. — Thick cryst. S. aq., alc., ac. ac.
388	218d. in sealed tube		1-Tyrosinehydantoic Ac., $HO \cdot C_6H_4 \cdot CH_2 \cdot CH(NH \cdot CO \cdot NH_2) \cdot CO_2H$. — \textcircled{P} Sol. boiled w. Millon's reagent gives intense red color & then a dark red ppt. (Cf. T. 2.19). — Glassy pr. fr. aq. Ndl. s. in 36 pt. alc.; s. alc.; i. eth. (Begins to soften at 154—5° in open tube, but is not melted at 170°.)
389	219—20; 218 in closed tube	$k \cdot 10^4 = 1.3$ 105.5	3-Nitrophthalic Ac., $NO_2 \cdot C_6H_4 \cdot (CO_2H)_2$. — \textcircled{P} Should give T. 2.21. — Yellowish monoclin. pr. fr. eth. Ndl. s. h. aq.; v.s. alc.; e.s. eth.; alm. i. chlf. Prolonged heating gives anhydride, m.p. 163—4°. — Me \bar{A} , m.p. 67—8°. — Ba \bar{A} .H ₂ O, mic. lft. alm. i.c. aq., s. h. aq.
390	219	181	4-Nitro-m-toluic Ac., $NO_2 \cdot C_6H_4 \cdot Me \cdot CO_2H$. — \textcircled{P} Should give T. 2.21. — Pr. fr. alc. E.s. h. alc.
391	219—20	218	Bz-Nitroquinolinecarbonic Ac., $NO_2 \cdot C_6H_4 \cdot C_6H_5 \cdot N \cdot CO_2H$. — \textcircled{P} Should give T. 2.21. — Cryst. fr. aq. D.s. c. aq.; e.s. h. aq. — Ag \bar{A} , cryst. powd. d.s. aq.
392	212—3d.; 229d.(r.h.)		Urocanic Ac., $C_6H_5 \cdot O_2H_4$. — Ndl. w. 4H ₂ O aq. of crystn. (lost at 105°). E.s. h. aq.; d.s. c. aq.; i. alc., eth. Reacts acid & dis. BaCO ₃ giving Ba \bar{A} .8H ₂ O, cryst. fr. aq. on adding alc.; but also gives B.2HCl, ndl. d.s. HCl, v.s. aq. [Found in dogs' urine.]
393	d. abt. 220	$\mu(170.5) =$ 367	4-Hydroxypyridine-2,6-dicarbonic Ac., Ammonchelidonic Ac., $HO \cdot C_6H_4 \cdot N(CO_2H)_2 \cdot H_2O$. — Rhomb. pr. s. in 637 pt. aq.; v.d.s. alc.; alm. i. eth. — Ignition w. Zn. dust gives pyridine (No. 2.1125).
394	220	198	5-Nitro-3-aminosalicylic Ac., $(NO_2)_2(NH_2) \cdot C_6H_3(OH)^2(CO_2H)$. — \textcircled{P} Should give T. 2.21. — Cryst.

(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
395	220-4 ± 2d. u.c. w. efferv.	k.10 ^s = 2.7 Alk. sol. viol. red †	Violuric Ac., $[NH.CO.C(NOH).CO.NH.CO]$. — ⊖ Sol. in dil. NaOH is violet-red, VR! — Cryst. in symmetrical trimetric octahedra. E.s. warm aq.; s. alc. Odorless. Baryta sol. in x.s. gives fr. aq. sol. ppt. of VRT ² color (after collection on filter). Dif. fr. alloxantine & amalic ac. — Melts completely w. brisk efferv. to or.-red liq. after softening fr. abt. 210°.
396	abt. 221 ± 5 u.c. d.	†	Amalic Ac., Tetramethylalloxantine, $[CO.NMe.CO.NMe.CO-C(OH)^{-}]$. — ⊖ Place abt. 0.005 g. on porcelain crucible cover. Mix w. 1 drop conc. ammonia. Heat over small flame until most of ammonia is removed. Residual sol. shows intense crimson coloration (Murexine). — Cryst. fr. h. aq. similar to those of alloxantine, which amalic ac. resembles in many properties & reactions. Alm. i. c. aq., alc.; d.s. h. aq. Melts to or.-red liq. after softening at lower temperature. Odorless. — Dissolve abt. 0.005-0.01 g. in 1 cc. conc. ammonia & add 1 drop $FeSO_4$ sol. An intense, but not very permanent dark blue coloration appears at once! (Alloxantine does not give this react.) — Amalic ac., although changed to deep violet color when moistened w. baryta sol., does not give the heavy violet ppt. described in test for alloxantine (No. 2.398).
397	220	217	4-Nitronaphthoic Ac., $NO_2.C_{10}H_8.CO_2H$. — ⊖ Should give T. 2.21. — Ndl. fr. alc. E.s. alc., gl. ac. ac., chlf.; d.s. bz. — Et \bar{A} , m.p. 54°.
399	abt. 222 ± 5 u.c. d. w. efferv.	$\mu(256) = 46$ †	Alloxantine, $C_6H_4N_2O_5 \cdot 3H_2O$. — ⊖ Place abt. 0.005-0.01 g. on porcelain crucible cover. Mix w. 1 drop conc. ammonia. Heat over small flame until most of ammonia is removed. Residual sol. shows intense VR color of much greater purity than in color standard! (Murexide.) — Cryst. fr. h. aq. sol. (slowly cooled) in minute transparent pr. somewhat resembling those of allantoin (No. 2.413), becoming pink on continued exposure to air & light. E.s. h. aq.; v.d.s. h. alc. Odorless. — Dissolve 0.005 g. in 10 cc. h. aq. Cool. Add baryta water in x.s. A voluminous ppt. of VR color (best observed on filter) is formed at once! — Cold sat. aq. sol. gives immediate ppt. of Ag fr. $AgNO_3$ sol. This react. & the above color react. are not given by allantoin (No. 2.413). — Changes to or.-red 10-20° below m.p. Loses 11.18% aq. when dried at 110°.
—	223	195	2-Nitro-1,3-dimethylbenzoic(5) Ac. — Cf. No. 2.272.
400	220-5d. w. efferv.		p-Hydrazinobenzoic Ac., $NH_2.NH.C_6H_4.CO_2H$. — Cryst. fr. aq. V.d.s. c. aq.; more s. h. aq. — In melting dec. to CO_2 & phenylhydrazine (Cf. No. 2.1369).
401	abt. 225	"Strongly acid"	Iminoacetic Ac., $NH.(CH_2CO_2H)_2$. — Rhomb. pr. — 100 pt. aq. at 5° dis. 2.43 pt.; i. alc., eth. — Cu $\bar{A}.2H_2O$, deep blue pr., d.s. h. aq.
402	255d.		m-Aminophenoxyamidic Ac., $NH_2.C_6H_4.NH.CO.CO_2H$. — Mic. ndl. d.s. h. aq.; e.s. alk. — Ag \bar{A} , ndl. fr. h. aq. giving CO & CO_2 at 170°.
403	225	"Dec. carbonates"	5-Nitrosalicylamide, $HO.C_6H_4(NO_2).CO.NH_2$. — ⊖ Aq. sol. colored blood red by $FeCl_3$. — Ndl. e.s. alc., h. aq. — Ba \bar{A} , $4H_2O$, yel. ndl., mod. s. c. aq. — Sapp. should give corresponding acid of m.p. 228°.
404	226d. w. efferv.		Pyridine-2,6-dicarboxylic Ac., Dipicolinic Ac., $C_6H_4N(CO_2H)_2$. — ⊖ Aq. sol. gives yel.-red color w. $FeCl_3$. — Cryst. fr. c. aq. w. $1\frac{1}{2}H_2O$ in hair-like nndl. D.s. c. aq.; v.d.s. alc. — Decn. by heat gives CO_2 & pyridine (No. 2.1125). — PCl_5 gives acid chloride of m.p. 61°.
405	226	105	3,5-Dinitrophthalic Ac., $(NO_2)_2.C_6H_3.(CO_2H)_2$. — ⊖ Should give T. 2.21. — Pr. fr. aq. V.s. aq., alc., eth.; i. bz., lgr. — Ba \bar{A} , ppt. i. aq. or dil. ac. ac.
406	226d. w. efferv.		β -Naphthindolecarboxylic Ac., $[NH.C_{10}H_8.CH:C(CO_2H)]$. — Alm. i. aq.; d.s. eth.; more s. alc.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
407	226	226	3,5-Dinitro-p-toluic Ac., $(NO_2)_2.C_6H_3Me.CO_2H$. — (P) Should give T. 2.21. — Cryst. fr. aq. V.d.s. c. aq.; v.s. alc., eth. Sbl. in ndl. Taste intensely bitter! — $Ba\bar{A}_2.H_2O$, intensely yel. pr., mod. s. c. aq.
408	227d.		Pyridine-2,4,6-tricarbonic Ac., $C_5H_4N(CO_2H)_3$. — (P) Gives violet color w. $FeSO_4$. — Cryst. w. $2H_2O$ fr. dil. H_2SO_4 , losing aq. at 110°. D.s. c. aq.; eth. — $Ba\bar{A}_2.6H_2O$, floc. ppt. — $Cu\bar{A}_2.12H_2O$, sky-blue cryst., d.s. aq.
409	227d.		cis-Piperidine-2,3-dicarbonic Ac., $C_5H_5N(CO_2H)_2$. — Glassy mass. V.s. aq.; i. alc. — B_2HCl , cryst. ppt. fr. HCl , m.p. 239° d. — W. $NaNO_2$ & HCl gives nitroso deriv. (T. 2.36), m.p. 138—9° d.
—	228.5 u.c.		Phthalimide. — Cf. No. 2.2555.
410	228-9	123	Nicotinic Ac., Pyridine-3-carbonic Ac., $C_5H_4N.CO_2H$. — Fine ndl. d.s. c. aq., e.s. h. aq., h. alc.; alm. i. eth. — Aq. sol. not pptd. by $Pb(CH_3COO)_2$. — $Ag\bar{A}$, long ndl. fr. h. aq. — B_2HAuCl_4 , lft., m.p. 207°.
411	228; 236 (r.h.)	$k.10^{\circ} = 5$	3,5-Diaminobenzoic Ac., $(NH_2)_2.C_6H_3.CO_2H$. — (P) V. dil. aq. sol. is colored yel. by trace HNO_2 . — Cryst. w. $1H_2O$ in long ndl. (aq. lost at 110°). 1000 pt. aq. at 8° dis. 11 pt. S. alc., eth. — Ignited w. BaO gives m-phenylenediamine, No. 2.634. — $Ag\bar{A}.2H_2O$, ppt. of mic. ndl. — B_2HCl , ndl., e.s. aq., alc.
—	228-30	Alk. sol. yel.	Hemipinimide. — Cf. No. 2.2556.
413	$229 \pm 2d.$ w. efferv.	$k.10^{\circ} = 1.2$	Allantoin, $[NH.CO.NH.CO.CH^2].NH.CO.NH_2$. — (P) A few drops of h. 10% aq. sol. allowed to cool on glass slide deposit v. perfect mic. pr., shown in Table 2 of Neubauer & Vogel's Harnanalyse. — S. in abt. 160 pt. c. aq.; e.s. h. aq.; alm. i. c. alc., eth. Odorless & nearly tasteless. Faintly acid to litmus. Consumes very little more than 1 cc. 0.1 N $NaOH$ in Gen. T. 2.I, w. v. indefinite end reaction. Turns red-brown about 220°. — Boil about 0.01 g. w. abt. 5 cc. $NaOH$ sol. (1 : 10) for several min. NH_3 is slowly evolved. Dilute sol. w. 10 cc. aq. Acidify strongly w. ac. ac. Add $CaCl_2$ sol. & heat to boiling. A considerable powdery ppt. of CaC_2O_4 is formed.
414	$230 \pm 5d.$ in 1 mm. sealed tube 5 mm. long	$k.10^{\circ} = 1.3 \dagger$	1-(ordinary)Asparagine, $CO_2H.C_2H_5(NH_2).CO.NH_3$. (Dried at 110°). — [In open capillary d.w.m. between 220 & 300°]. — (P) Cryst. on mic. slide v. readily fr. 15% sol. in short clear rhombic pr. of characteristic appearance, showing l-hemihedral planes, & containing $1H_2O$ of crystn. — S. in 55.8 pt. aq. at 10.5°, or in 1.89 pt. at 100°; i. c. abs. alc., eth. Taste of sat. aq. sol. astringent & then faintly sour. Sapn. gives NH_3 & aspartic ac. — Loses 12.0% aq. when cryst. are heated 1 hour at 120°! — Gives 9.34% α -amino nitrogen in T. 2.5. — $[\alpha]_D = -6.14^{\circ}$ in aq. sol., but +37.27° in HCl sol. [d-Asparagine differs fr. its optical antipode in having a sweet taste. For isolation & determination of asparagine in plants, Cf. Abderhalden's Handb. d. Biochem. Arbeitsmethoden, 2. 511.]
415	230d.	"Sol. in alk. red." "An acid"	Phthalylhydroxylamine, $C_6H_4:(CO)_2.NOH$. — Cryst. somewhat s. aq.; e.s. h. alc.; i. eth. — $Ag\bar{A}$, dark red ppt. — Heating gives phthalic anhydride.
416	230; 228	183	5-Nitrosalicylic Ac., $(NO_2)^2.C_6H_3(OH)^2(CO_2H)^2$. — (P) $FeCl_3$ gives blood red color w. aq. sol. — Thick cryst. fr. gl. ac. ac. S. in 1475 pt. aq. at 15.5°, much more s. h. aq.; e.s. alc. — Boiling w. HNO_3 gives picric ac., No. 2.3168. Ignition w. CaO gives p-nitrophenol.
418	230d.		Diphenylasparagine, $CO_2H.C_6H_4(NH_2).CO.NPh_3$. — Warts d.s. alc., i. eth.
419	231d.	$k.10^{\circ} = 3$	Quinolinic Ac., Pyridine-2,3-dicarbonic Ac., $C_5H_4N.(CO_2H)_2$. — Sinters and burns at 190—5°. Lust. monoclin. pr. S. in 183 pt. aq. at 6.5°; d.s. alc.; v.d.s. eth., i. bz. — $AgNO_3$ + HNO_3 added to h. aq. sol. gives ppt. of $Ag\bar{A}$, lust. ndl. which recryst. fr. aq. w. $1H_2O$.

(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
420	227-35d. w. efferv.	k.10 ⁷ = 7.5 †	Parabanic Ac., Oxalylurea, $[CO.NH.CO.CO.NH]^2$. — (P) Sapp. by aq. NaOH sol. gives NH ₃ & oxalic ac.! — Thin, clear 4-sided plates fr. aq. or alc. Odorless. S. in 21.2 pt. aq. at 8°; e.s. c. alc., d.s. eth. Becomes or-yel. in melting. — (D) Dis. a few mg. of acid in 5 cc. of CaCl ₂ sol. Add 10 drops conc. ammonia. No ppt. occurs in the cold. Boil. A copious ppt. of CaC ₂ O ₄ appears just before boiling begins.
421	232		1-Pinonicsemicarbazone, CO ₂ H.C ₆ H ₅ :N.NH.CO.NH ₃ . — Cryst. fr. alc. — Prob. gives T. 2.17.
422	abt. 230-5		2-Phenylpyridine-Bz-2,Py-3-dicarbonic Ac., CO ₂ H.C ₆ H ₄ .C ₆ NH-CO ₂ H. — (P) Heated to 240-1° loses CO ₂ , giving dark blue comp., e.s. chlf. — Cryst. Froths strongly at 236°. — Ignition w. CaO gives 2-phenylpyridine (No. 2.1422).
—	d.233 ± 2		Glycine. — Cf. No. 2.2568.
424	233d.	k.10 ² = 1	Papaveric Ac., (MeO) ₂ .C ₆ H ₅ .CO.C ₆ H ₅ N(CO ₂ H) ₂ .H ₂ O. — Cryst. w. 1H ₂ O. HCl ppts. oily & then resinous fr. conc. sol. of salts. V.d.s. c. aq., alc., eth., chlf., bz. Dis. unchanged in conc. H ₂ SO ₄ . — CaA.1½H ₂ O, cryst. powd., by pptn. of ac. by CaCl ₂ + NH ₄ OH; once pptd., v.d.s. aq.
425	234-5u.c.	167 † k.10 ⁴ = 4	p-Nitrobenzoic Ac., NO ₂ .C ₆ H ₄ .CO ₂ H. — (P) Gives dark brown ppt. in T. 2.21. — GYT3 lft. fr. h. aq. Odorless. Taste bitter. — 10 cc. of each of following solvents dis.: — aq. at 20°, 0.004 g.; alc. at 10°; 0.009 g.; eth. at 11°, 0.22 g. V.d.s. chlf., bz.; i. lgr. — (D) Reduce 0.6 g. to p-aminobenzoic ac. by procedure given under No. 2.164, except that 10 cc. of boiling water is used in the recrystn. The product is obtained in slender ndl. of m.p. 185-6° u.c.
426	234d.		2-Methylquinoline-3-carbonic Ac., Me.C ₆ H ₅ N.CO ₂ H. — Ndl. fr. alc. Dec. in melting to CO ₂ & 2-methylquinoline, No. 2.1376! — AgA, cryst. ppt.
427	d. abt. 235		Pyridine-2,4,5-tricarbonic Ac., Berberonic Ac., C ₆ H ₅ N.(CO ₂ H) ₃ . — (P) FeSO ₄ gives blood red coloration, & Pb(CH ₃ .CO) ₂ , i. ppt. — Trichlin. pr. Cryst. w. 2H ₂ O, losing 1H ₂ O in air. D.s. c. aq.; e.s. h.; v.d.s. h. alc.; i. eth., bz. — Becomes red at 215°. Loses CO ₂ above m.p.
428	235d.	153	3-Aminosalicylic Ac., NH ₂ .C ₆ H ₄ (OH)(CO ₂ H). — Alm. i. alc. Aq. sol. reduces h. Fehling's sol. — B.HCl, m.p. 150°. — W. benzoyl chloride at 190° gives benzoyl deriv., m.p. 189°.
429	235	165 k.10 ⁶ = 9.2	p-Dimethylaminobenzoic Ac., Me ₂ N.C ₆ H ₄ .CO ₂ H. — Ndl. fr. alc.; i. ac. ac. — MeA, silvery lft. fr. alc., m.p. 102°.
430	235	165	2-Aminomesitylenic Ac., (NH ₂)(Me) _{1,2} .C ₆ H ₃ (CO ₂ H) ² . — Ndl. fr. alc. D.s. aq.; e.s. h. alc.
431	236	k.10 ³ = abt. 4.3	Pyridine-2,5-dicarbonic Ac., Isocinchomeric Ac., C ₆ H ₅ N-(CO ₂ H) ₂ . — (P) Aq. sol. gives yel.-red color w. FeSO ₄ (dif. fr. 2,4-isomer, No. 2.433) & ppt. w. Pb(Me.CO) ₂ . — Mic. lft. fr. h. aq. + little HCl. Cryst. w. 1H ₂ O fr. h., or w. 2H ₂ O fr. c. sol. Alm. i. c. aq., alc., eth., bz.; e.s. upon addition of few drops HCl. — Dec. at 245° to CO ₂ & nicotinic ac. (No. 2.410). — CuA.H ₂ O, light blue cryst. ppt. fr. h. sol.; i. aq. after pptn.
432	236d.	†	3-Phenylpyrazalone(5), $[CH_3.CO.NH.N:CPh]^2$. — On account of insolubility neutralization of alk. in Gen. T. 2.I is v. slow. Lft. fr. alc. Alm. i. c. alc., eth., bz. — CaA, cryst. ppt. — B.HCl, silky ndl., m.p. 196°. — 4Br added to gl. ac. ac. sol. after neutralization w. NaOH gives dibromo deriv., yel. lft., m.p. 196°.
433	239-40	k.10 ³ = 6	Pyridine-2,4-dicarbonic Ac., C ₆ H ₅ N.(CO ₂ H) ₂ . — (P) Gives intense blood red color w. Fe ₂ SO ₄ . — Lft. fr. aq. Mod. s. c. aq.; v.s. h. aq.; e.s. alc.; i. eth., bz. — Decn. by heat gives CO ₂ & pyridine-4-carbonic ac. — PCl ₅ gives acid chloride, m.p. 203°. — CuA, blue-green cryst. ppt., i. aq., crystg. w. 3H ₂ O unless pptd. boiling hot. — PbA, ppt. i. aq.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
434	239-40	270	Di-o-toluidinoacetic Ac., $(\text{Me.C}_6\text{H}_4.\text{NH})_2.\text{CH.CO}_2\text{H}$. — Yellowish ndl. fr. alc. I. aq.; v.d.s. alc.; e.s. acids or alk.
—	abt. 240d.		Isatoic Anhydride. — Cf. No. 2.2588. (Sol. in 1% alk. shows transient bluish fluorescence.)
436	240u.c.		v-(1,2,3)-Hemimellitimide, $\text{NH}:(\text{CO})_2.\text{CO}_2\text{H}$. — Ndl. v.d.s. aq.; d.s. alc., gl. ac. ac. — $\text{Ag}\bar{A}$, ppt.
437	240u.c.	193 †	o-Nitrocinnamic Ac., $\text{NO}_2.\text{C}_6\text{H}_4.\text{CH}:\text{CH.CO}_2\text{H}$. — \oplus A few mg. heated for some sec. over small flame w. conc. H_2SO_4 gives deep purple color (indigo)! — Pearly scales or flat ndl. fr. h. alc. I. c. aq.; d.s. c. alc. Tasteless. Odor faintly aromatic. — \oplus Heat 0.5 g. in t.t. in boiling water-bath w. 30 cc. of the KMnO_4 sol. of T. 1.905(1). Add 10 cc. boiling aq. Filter hot. Wash w. 5 cc. h. aq. Evaporate filtrate to 5-6 cc. Add 2 cc. HCl (sp. gr. 1.12). Cool. Filter. Recryst. fr. 5 cc. boiling aq. Add 0.1 g. boneblack. Boil 1-2 min. Filter hot. Dry the lust. white ndl. of o-nitrobenzoic ac., which separate on cooling, for 15 min. at 100° . The product melts at 146° u.c. after softening at 144.5° . (Yield excellent.)
438	241-2d.		Apophylenic Ac., Cinchomeronicmethylbetaine, $\text{CO}_2\text{H}.\text{C}_6\text{H}_4.\text{Me.CO.O}^+$. — Octahedra + H_2O (aq. lost at 120°) fr. c. aq. sol. D.s. c. aq.; i. alc., eth.
439	241-2	217	(β)-5-Nitronaphthoic(1) Ac., $\text{NO}_2.\text{C}_{10}\text{H}_8.\text{CO}_2\text{H}$. — \oplus Prob. gives T. 2.21. — Fine ndl. Sbl. in lust. spangles. S. in 4820 pt. c. aq., or in 187 pt. alc.; e.s. h. alc., eth., gl. ac. ac., bz. — Oxidation by KMnO_4 gives 3-nitrophthalic ac. (No. 2.389). — $\text{Ba}\bar{A}.3\frac{1}{2}\text{H}_2\text{O}$, yel. ndl. mod. s. c. aq. — $\text{Pb}\bar{A}.5\frac{1}{2}\text{H}_2\text{O}$, ppt. of mic. ndl. — $\text{Me}\bar{A}$, m.p. 110° , yel. ndl., e.s. alc.
440	242-3u.c. (r.h.)	†	"Lycetol," 2,5-(α)-Dimethylpiperazine Tartrate, $[\text{NH.CHMe.-CH}_2.\text{NH.CHMe.CH}_2]_2.\text{C}_6\text{H}_4.\text{O}_2$. — \oplus 5-6 drops sat. aq. sol. of picric ac. added to 0.05 g. lycetol disd. in 2 cc. c. aq. gives YTI (dry) ppt. of picrate, i. aq., alc., bz., gl. ac. ac., which begins to darken, without melting, at 240° u.c. — Mic. cryst. fr. h. 50% alc. Cryst. fr. h. aq. sol. w. $3\text{H}_2\text{O}$ in t.b.l. efflorescing over H_2SO_4 or on water-bath. — T. 1.302 gives Y-YTI color. — \oplus Dis. 0.1 g. in 2 cc. c. aq. + 1 cc. HCl (sp. gr. 1.12). Add slowly 1 cc. sat. aq. sol. of NaNO_2 . Recryst. scaly ppt. fr. 2 cc. boiling aq., cooling & shaking. The dinitroso deriv. formed, $\text{C}_6\text{H}_{12}\text{N}_2(\text{NO})_2$, is obtained in faintly yellowish lust. ndl., m.p. 172° u.c.
441	245-7d.	167	p-Hydroxyphenylglycine, $\text{HO.C}_6\text{H}_4.\text{NH.CH}_2.\text{CO}_2\text{H}$. — \oplus Colored blood red by FeCl_3 . — Lft. d.s. aq., alc.; i. eth. Browns at 200° . Softens fr. 220° . Dec. to CO_2 & methylaminophenol in melting.
442	245-6	228	3,5-Dinitro-p-hydroxybenzoic Ac., $(\text{NO}_2)_2.\text{C}_6\text{H}_4(\text{OH}).\text{CO}_2\text{H}$. — \oplus Prob. gives T. 2.21. — Tbl. d.s. c. aq.; mod. s. h. aq.; e.s. alc. eth. — $\text{Ba}\bar{A}.5\text{H}_2\text{O}$, yel. ndl.
443	246; 258-9	105.5	4-Nitroisophthalic Ac., $\text{NO}_2.\text{C}_6\text{H}_4(\text{CO}_2\text{H})_2$. — \oplus Prob. gives T. 2.21. — Cryst. fr. aq. w. 1 or $3\text{H}_2\text{O}$. Mod. s. c. aq.; v.s. alc. eth. — $\text{Ba}\bar{A}.4\text{H}_2\text{O}$, lust. ndl.
444	246; 240-1		2-Methylquinoline-4-carbonic Ac., Aniluvitonic Ac., $\text{Me.C}_6\text{H}_4\text{N}-(\text{CO}_2\text{H})$. — \oplus Dec. to CO_2 & quinaldine (No. 2.1376) in melting, or, better, on distn. w. CaO . — Cryst. D.s. c. aq.; alm. i. h. chlf.; v.s. dil. mineral ac. — $\text{Ba}\bar{A}$, ndl. d.s. c. aq. — $\text{Ag}\bar{A}$, lft. — BPk, green-yel. ndl. fr. alc., m.p. $190-1^{\circ}$.
445	248-9 sl.d.	105.5	5-Nitroisophthalic Ac., $\text{NO}_2.\text{C}_6\text{H}_4(\text{CO}_2\text{H})_2$. — \oplus Prob. gives T. 2.21. — Lft. s. at 15° in 685 pt. aq., or in 1.23 pt. at 99° ; v.s. alc. — $\text{Me}\bar{A}$, lust. felted ndl. w. anise odor when warmed, m.p. 121.5° . — $\text{Ba}\bar{A}.2\frac{1}{2}\text{H}_2\text{O}$, ndl. s. in 117 pt. c. aq. Turns rose red in sunlight.
446	248-9; 247	173	Quinoline-7-carbonic Ac., $\text{C}_6\text{H}_4\text{N.CO}_2\text{H}$. — \oplus Above m.p. dec. to quinoline (No. 2.1356) & CO_2 . — Ndl. fr. aq. Sbl. in woolly flocks. D.s. c. aq., i. eth.; v.d.s. bz.; e.s. alc. — $\text{Ag}\bar{A}$, floc. ppt.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. ^o).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
447	248d. w. efferv.	179 $k \cdot 10^6 = 8.5$	m-Acetylamino-benzoic Ac., $\text{Me.CO.NH.C}_6\text{H}_4\text{CO}_2\text{H}$.—D.s. h. aq., h. alc., eth. — Ag \bar{A} , ndl. fr. h. aq. — Ba \bar{A} , 3H ₂ O, fine ndl. v.s. aq.
448	249-50d. (r.h.)	$\mu(256) = 337$	Pyridine-2,3,4-tricarbonic Ac., Carbocinchomeric Ac., $\text{C}_5\text{H}_5\text{N}(\text{CO}_2\text{H})_3$.— \oplus Gives pale red color w. FeSO ₄ . Clear rhomb. tbl. w. 1 $\frac{1}{2}$ H ₂ O (lost at 115-20°). Cryst. s. in 83.9 pt. aq. at 15°. E.s. h. aq.; r.d.s. alc., alm. i. eth., bz. — Ba \bar{A} , 16H ₂ O (ppt. fr. Ammon. salt & Ba(MeCO ₂) ₂ sol.). — Ba \bar{A} , 12H ₂ O (pptn. of free acid by Ba(MeCO ₂) ₂ alm. i. ppt.)
449	249	226	2,3-Dinitro-p-toluic Ac., $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me.CO}_2\text{H}$.— \oplus Prob. gives T. 2.21. — Lust. rhomb. pr. fr. alc. — Taste intensely bitter! — Ba \bar{A} , w. 3 or 4H ₂ O of crystn., ndl. d.s. aq.
450	250d.	139	β -Hydroxypicolinic Ac., $\text{HO.C}_6\text{H}_4\text{N.CO}_2\text{H}$.—Lft. e.s. h. aq., alc.; i. eth. — Strongly heated dec. to CO ₂ & hydroxypiperidine of m.p. 148°. — Ba \bar{A} , 2H ₂ O, tbl. r.d.s. aq.
451	250 ± 5	173 † $k \cdot 10^6 = 1.3$	Quinoline-4-carbonic Ac., 4-Cinchoninic Ac., $\text{C}_9\text{H}_8\text{C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Cryst. w. 1 or 2H ₂ O in tbl. or pr. on slow evapn. of aq. sol. (Aq. lost at 100°). Softens fr. 135-6°. V.d.s. aq., alc.; i. eth. — Ignition w. CaO gives quinoline (No. 2.3156) & some β -biquinolyl. — Ag \bar{A} , cryst. ppt.
452	250d.	179 † $k \cdot 10^6 = 5.2$	p-Acetylamino-benzoic Ac., $\text{Me.CO.NH.C}_6\text{H}_4\text{CO}_2\text{H}$. — Ndl. d.s. aq., eth.; more s. alc. — Sapn. (Cf. T. 2.26) by boiling HCl gives p-aminobenzoic ac., No. 2.290. — Ag \bar{A} , ndl., mod. s. aq.
453	250-1d.	$\mu(1316) = 324$	2,5-Dimethylpyrrole-3,4-dicarbonic Ac., $\text{Me}_2\text{C}_4\text{NH}(\text{CO}_2\text{H})_2$. — \oplus Vapors (dimethylpyrrole) evolved on ignition in t.t. should give splinter react., T. 2.24. — Flat ndl. fr. alc.
455	250d.	261	3-Oxyindole-5-carbonic Ac., $\text{C}_9\text{H}_7\text{O}_2\text{N}$. — \oplus Warmed w. Na ₂ CO ₃ sol. gives intense blue sol. of indigocarbonic ac. — Sl. yellowish mass. V.d.s. aq., alc.; more s. gl. ac. ac.
456	252	257	5-Benzoylamino-salicylic Ac., $\text{Ph.CO.NH.C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})$. — Alm. amorph. powd. Alm. i. h. aq.; v.s. ac. ac.; i. chlf., bz., lgr.
457	253d		cis-trans-Piperidine-2,3-dicarbonic Ac., $\text{C}_9\text{H}_8\text{N}(\text{CO}_2\text{H})_2$. — Cryst. s. aq.; v.d.s. alc. — B.HCl, m.p. 221°, s. aq. — B.HAuCl ₄ .H ₂ O. Cryst. m.p. 184° (when anhydrous). — Gives cryst. nitroso deriv., s. aq., m.p. 154° d.
458	254		3-Methylquinoline-4-carbonic Ac., $\text{Me.C}_9\text{H}_7\text{N.CO}_2\text{H}$. — Glassy mic. lft. fr. aq. D.s. acetone; i. eth., lgr., bz. — Distn. w. soda-lime gives 3-methylquinoline, No. 2.1388.
459	255d.	128	3,5-Dinitrotetraphthalic Ac., $(\text{NO}_2)_2\text{C}_6\text{H}_2(\text{CO}_2\text{H})_2$. — \oplus Prob. gives T. 2.21. — Glassy cryst. fr. aq. D.s. c. aq. — Ba \bar{A} (at 100°), ppt.
460	254-6		Methylasparagine, $\text{NH}_2\text{CO.CH}_2\text{CMe}(\text{NH}_2)\text{CO}_2\text{H}$ (?). — Cryst. w. 1H ₂ O in plates which effloresce; also anhydrous. Aq. sol. tastes sweetish.
461	254-6d.		d,L-Glutamine, $\text{CO}_2\text{H.CH}(\text{NH}_2)_2\text{CH}_2\text{CONH}_2\text{H}_2\text{O}$. — \oplus Should give N in T. 2.4. Effloresces easily. Becomes yel. at 240°. Reacts faintly ac. & tastes sweetish.
462	255	$k \cdot 10^7 = 4$	Methylnitouracil, $[\text{CO.C}(\text{NO}_2)_2\text{CH.NH.CO.NMe}_2]$. — \oplus Evolves methylamine (No. 2.1059) on sapn. (Cf. T. 2.26). — Cryst. fr. alc. s. c. aq. v.d.s. alc. — Ag \bar{A} , ppt., long ndl. fr. aq.
463	255	173	o-Cyanocinnamic Ac., $\text{C}_6\text{H}_5\text{CN.CH}: \text{CH.CO}_2\text{H}$. — Fine ndl. fr. alc.
464	256		2-Hydroxypyridine-3-carbonic Ac., $\text{HO.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — \oplus Dec. on distn. to CO ₂ & 2-oxypyridine. — Lust. ndl. fr. aq. V.d.s. c. aq. — Ag \bar{A} , silky ndl.
465	256; 265sbl. w.m.	151	Phenylaminoacetic Ac., $\text{Ph.CH}(\text{NH}_2)\text{CO}_2\text{H}$. — \oplus Distn. gives CO ₂ & benzylamine (No. 2.1236). — Pearly lft. V.d.s. solvents. — Ag \bar{A} , cryst. ppt. alm. i. aq.

GENUS I, DIV. A.
(ORDER II, SUBORDER I.)

No.	Melting-point (C. ^o).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
466	256d.		Piperidine-3,4-dicarbonic Ac., C ₆ H ₅ N.(CO ₂ H) ₂ . — E.s. aq. "Monobasic ac." — Ca \bar{A} . ₂ 5H ₂ O. — B.HCl, m.p. 237° d., v.s. aq.
467	256d.	207	γ -Hydroxyquinolinecarbonic Ac., C ₉ H ₇ ON.CO ₂ H ₂ O. — ⊖ Aq. sol. colored red-brown by FeCl ₃ . Silky ndl. S. aq., alc.; alm. i. eth., chlf., bz. — Dec. to CO ₂ & 8-oxyquinaline in melting.
468	257d.		2,3,4-Trimethylpyridine-5-carbonic Ac., Me ₃ C ₆ H ₃ N.CO ₂ H. — Pr. fr. dil. alc. E.s. aq. — Ag \bar{A} , white ppt.
469	258sl.d.	139	γ -Hydroxypicolinic Ac., HO.C ₆ H ₃ N.CO ₂ H. — ⊖ Aq. sol. colored yel.-red by FeCl ₃ . — Tbl. fr. aq. Cryst. w. 1H ₂ O (lost at 110°). D.s. aq.; more s. alc.; i. eth. — Ba \bar{A} , mic. pr., r.d.s. aq.
470	abt. 258d. w. efferv.	k.10 ⁸ = 2.1 †	Cinchomeronic Ac., Pyridine-3,4-dicarbonic Ac., C ₆ H ₅ N(CO ₂ H) ₂ . — ⊖ Ignition w. CaO gives pyridine (No. 2.1125). — Pr. fr. aq. + little HCl. V.d.s. h. aq.; d.s. alc.; alm. i. eth.; i. chlf. — Distn. gives mixture of nicotinic & isonicotinic acids. — Not colored by FeSO ₄ . — Hot. sol. gives floc. ppt. w. Cu(Me.CO ₂) ₂ sol. which redissolves on cooling, but on long boiling gives i. characteristic, cryst. blue salt. — Ag \bar{A} , ppt. — Ag \bar{A} , cryst. ppt. w. AgNO ₃ , fr. sol. containing HNO ₃ .
471	259		2-Methylquinoline-6-carbonic Ac., Me ₂ C ₉ H ₇ N.CO ₂ H. — Browns at 240°. Ndl. fr. alc. V.d.s. h. aq.; e.s. h. alc. — Ag \bar{A} , gelatinous ppt., cryst. after boiling. — B.H ₂ PtCl ₆ . ₄ H ₂ O, monoclin. tbl. d.s. c. v. dilute HCl.
472	d.w.m. 260	k.10 ⁸ = 2.1	2,4-Dimethylpyrrole-3,5-dicarbonic Ac., Me ₂ C ₄ NH ₂ (CO ₂ H) ₂ . — ⊖ Gives splinter pyrrole react. (T. 2.24), decomposing to CO ₂ & dimethylpyrrole (No. 2.2759) on ignition, or boiling w. HCl. — Cryst. flocks. — Reduces ammon. AgNO ₃ sol. on boiling.
473	261 Carbonizes	μ (256) = 278	Pyridine-3,4,5-tricarbonic Ac., C ₆ H ₅ N.(CO ₂ H) ₃ . — Lft. crystg. w. 3H ₂ O (lost at 115°). D.s. c. aq.; e.s. h. aq. — Ag \bar{A} . ₂ H ₂ O, cryst. ppt.
473-1	266-7; 257-8 (w. frothing)		4-Hydroxyquinolinecarbonic(3) Ac., Kynurenic Ac., ⁷ C ₉ H ₇ N ₂ CH ₂ CO ₂ H :C(OH) ² . — [In dog urine.] — Ndl. w. H ₂ O (anhydrous at 150°). — ⊖ Warmed w. HCl & KClO ₃ & evapd. to dryness gives emerald-green color w. ammon. — Slowly fused gives CO ₂ & kyanurin (No. 2.2456). — Ignition w. Zn dust gives quinoline (No. 2.1356), & CO ₂ . — Ag \bar{A} . ₂ H ₂ O, white ppt.
474	264u.c.	†	Theophylline, 1,3-Dimethyl-2,6-dioxypurine, "Theocin," C ₈ H ₁₀ O ₄ N ₄ H ₂ O. — ⊖ Gives the murexide reaction, T. 2.20! Ndl. fr. h. aq. S. in 226 pt. aq. at 15°, or in 75 pt. at 37°. d.s. c. alc., eth., CCl ₄ ; more s. h. alc. Chlf. extracts fr. acid, but not fr. alk. sol. (Gadamer, p. 462). — Tannic ac. gives ppt. in aq. sol. s. in x.s. of reagent. — AgNO ₃ added to 1% ammon. sol. gives ppt. in cold. — Gives François' reaction. — 0.1 g. neutralizes 2.6 ± 0.2 cc. 0.1 N NaOH in Gen. T. 2.1. [Occurs in tea.]
475	262-3; 264d.	177 k.10 ⁸ = 1.1	1-Phenylurazole, C ₉ H ₇ O ₂ N ₂ . — ⊖ FeCl ₃ gives violet coloration. — Lust. lft. fr. aq. Alm. i. c. aq.; e.s. h. aq.; d.s. c. alc., eth.; e.s. alk. — Ag \bar{A} , m.p. 252° d. — Boiling w. acetic anhydride & crystn. fr. aq. gives acetyl deriv., m.p. 170°.
476	265	262	4,5-Dinitro- α -naphthoic Ac., (NO ₂) ₂ C ₁₀ H ₆ CO ₂ H. — ⊖ Prob. gives T. 2.21. — Ndl. fr. alc. D.s. eth., bz., h. aq.; e.s. h. alc. — Sbl. — Ba \bar{A} . ₂ 4H ₂ O, yel. pr., e.s. h. aq. — Et \bar{A} (fr. Ag \bar{A} + ETI), fine ndl. fr. alc., m.p. 143°.
477	267	k.10 ⁸ = 5.7	α -Hydroxypicolinic Ac., HO.C ₆ H ₃ N.CO ₂ H. — Ndl. w. 1 or 2H ₂ O (aq. lost at 130°). E.s. h. aq., alc.; i. eth. — Ba \bar{A} . ₂ H ₂ O, ndl. fr. h. aq.; d.s. c. aq.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
478	270; 262-3	211	Nitrotetraphthalic Ac., $\text{NO}_2\text{C}_6\text{H}_4(\text{CO}_2\text{H})_2$. — (2) Prob. gives T. 2.21. — Sometimes cryst. w. $1\text{H}_2\text{O}$. E.s. h. aq., h. alc. — $\text{Me}_2\bar{A}$, pr. fr. eth., m.p. 70° or $74-5^{\circ}$.
478-I	263-5; 271-3; (d.w. efferv.)	$k_A \times 10^6 =$ 2.5 $k_B \times 10^{12} =$ 1.3	d,l-Phenylalanine, Phenyl- α -aminopropionic Ac., $\text{Ph.CH}_2\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$. — (2) Stellate groups of short lust. pr. fr. aq. D.s. c. aq. or h. alc.; i. eth. Sbl. w. sl. decon. Taste sweet. Opt. i. Gives same results as No. 2.481-1 in prelim. T. (1) & (2), & in α -amino N test. — B_2Pk (fr. aq. sol.), sulphur-yel. ndl. (100 pt. c. aq. dis. 2.55 pt.; 100 pt. alc., 1.3 pt.), m.p. 173° (darkening at 170°). — $\text{B.C}_{10}\text{H}_8\text{O}_4\text{N}_4$ (picrolonate, yel. cryst. fr. aq., m.p. 238° , browning fr. 220° (100 pt. c. aq. dis. 0.19 pt.). — Benzoyl deriv., m.p. $187-8^{\circ}$ c. — Phenylisocyanate deriv., m.p. abt. 182° d.; corresponding hydantoin, ndl., v.d.s. aq., m.p. $173-4^{\circ}$ c. [d,l-Et \bar{A} , thick oil of faint odor, d.s. aq., b.p. 143° (10 mm.).]
479	269-70; 265		m-Ureidobenzoic Ac., $\text{NH.CO.NH.C}_6\text{H}_4\text{CO}_2\text{H}$. — (2) Should give NH_3 in T. 2.7. — Cryst. w. $1\text{H}_2\text{O}$ (lost at 100°). Cryst. s. in 98.5 pt. aq. at 100° ; in 139 pt. alc. or 786 pt. eth. at room temperature. — Sapn. (T. 2.26) w. boiling KOH gives m-aminobenzoic ac. (No. 2.259) & NH_4 .
480	abt. 270	"Monobasic acid."	p-Urazin, Diurea, $\text{CO}(\text{NH.NH})_2\text{CO}$. — (2) Reduces ammon. AgNO_3 sol., but not Fehling's sol. Gives red color w. FeCl_3 . — Monoclin. pr. fr. aq. D.s. c. aq., h. alc. — $\text{Ag}\bar{A}$ (dried at 100°), ppt. — W. conc. HCl at 150° gives CO_2 & hydrazine.
481	272d.	173	Isoquinoline-5 or 8-carbonic Ac., $\text{C}_8\text{H}_7\text{N.CO}_2\text{H}$. — Ndl. fr. alc. V.d.s. h. aq. — B.Pk , yel. fr. alc., m.p. $212-3^{\circ}$.
481-I	abt. 273u.c. d.(283c.)		† l-Phenylalanine, β -Phenyl- α -aminopropionic Ac., $\text{Ph.CH}_2\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$. — [Proteolytic product & in germinating plants.] Lust. lft. fr. h. conc. aq. sol., or ndl. w. aq. fr. aq. S. in 32.4 pt. aq. at 25° ; more s. h. aq. alm. i. most neutral solvents. Taste, sl. bitter. $[\alpha]_D^{20} = -35.1^{\circ}$ (in aq. sol.). Partly sbl. on heating & partly changes to lactimide. — (2) (1) Vapors on ignition in tube give pyrrole splinter react., T. 2.24. — (2) Give charac. odor of phenylacetic aldehyde on warming w. H_2SO_4 & $\text{K}_2\text{Cr}_2\text{O}_7$. — Sol. is pptd. by HgN_3O_6 . — $\text{Cu}\bar{A}$, pale blue scales, alm. i. aq. (fr. h. aq. sol. w. $\text{Cu}(\text{OH})_2$). — $\text{B.C}_8\text{H}_7\text{O}_4\text{N}_4$ (picrolonate), cryst. s. in 294 pt. aq. at 20° . — Phenylisocyanate deriv. (T. 2.31), ndl., d.s. aq., m.p. 182° ; corresponding hydantoin, ndl. v.d.s. aq., m.p. $173-4^{\circ}$ c. — (2) Gives 8.49% α -amino N by T. 2.5.
482	274d.		o-Aminohexahydrobenzoic Ac., $\text{NH.C}_6\text{H}_{10}\text{CO}_2\text{H}$. — (2) Should give N in T. 2.4. — Lust. ndl. fr. dil. alc. Alm. i. abs. alc., eth. Taste bitter. — $\text{Cu}\bar{A}_2\text{H}_2\text{O}$, blue ppt.
483	274d. w. efferv.		2-Methylpyridine-4,6-dicarbonic Ac., Uvitonic Ac., $\text{Me.C}_6\text{H}_4\text{N}-(\text{CO}_2\text{H})_2$. — Cryst. powd. alm. i. c. aq.; e.s. h. mineral ac. — Ignition w. CaO gives 2-picoline (No. 2.1153). — $\text{Cu}\bar{A}_3\frac{1}{2}-4\text{H}_2\text{O}$, blue green cryst. ppt.
484	275el.d.		3-(β)-Quinolinecarbonic Ac., $\text{C}_8\text{H}_7\text{N.CO}_2\text{H}$. — Cryst. fr. dil. alc. D.s. c. aq.; s. h. aq.; e.s. alc. — Ignition w. CaO gives quinoline (No. 2.1356). — Picrate, long fine ndl. d.s. c. alc., m.p. 216° .
485	278	241	Benzoylamino-benzoic Ac., $\text{Ph.CO.NH.C}_6\text{H}_4\text{CO}_2\text{H}$. — Ndl. fr. alc. D.s. aq.; s. alc., eth., gl. ac. ac. — Ag salt, silvery scales fr. h. aq.
486	280d.	192	Quinic Ac., $\text{MeO.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — (2) Conc. alc. sol. shows a blue, & dil. sol. a violet fluor. by H_2SO_4 . — Thin yellowish white pr. fr. dil. HCl. D.s. c. or h. aq.; eth., bz.; e.s. mineral ac. w. yel. color. — $\text{Ag}\bar{A}$, powdery ppt.
487	285d.		2-Methylquinoline-7-carbonic Ac., $\text{Me.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Browns at 270° . — Silky ndl. fr. alc. Sbl. w. partial decon. Alm. i. aq.; e.s. alc. — $\text{Ag}\bar{A}$, voluminous ppt. — $\text{B}_2\text{H}_2\text{PtCl}_6$, monoclin. pr.

No.	Melting-point (C. $^{\circ}$).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
488	285 \pm 1	193 †	p-Nitrocinnamic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CH}(\text{CO}_2\text{H})\text{CH}_2\text{CO}_2\text{H}$. — (D) T. 2.21 gives Dark Bkn. Y ppt. — GYT3-T4 mic. pr. fr. h. alc. (after boneblacking). D.s. h. alc. & still less s. in h. aq. or eth.; i. lgr., CS ₂ . — Hot conc. H_2SO_4 colors orange. — (D) Oxidize 0.5 g. w. KMnO_4 as directed in case of No. 2.437. Evaporate clear filtrate to 15–20 cc. Add 2 cc. HCl (sp. gr. 1.12). Cool. Filter. Recryst. fr. 40 cc. h. aq. containing 0.1 g. boneblack, boiling 2 min. p-Nitrobenzoic ac. cryst. out on cooling in alm. white (GYT3) lft., m.p. 234–4.5° u.c., after softening at 232°.
499	286		8-Methylquinoline-5-carbonic Ac., $\text{Me.C}_6\text{H}_4(\text{CO}_2\text{H})\text{C}_6\text{H}_5\text{N}$. — Powd. Ignition w. soda-lime gives 8-methylquinoline (No. 2.1382). — CaA, cryst. ppt. — $\text{B}_2\text{H}_2\text{PtCl}_6\text{H}_2\text{O}$, long yel. ndl.
490	285–9		Phenylisocyanuric Ac., $\text{C}_6\text{H}_5\text{O}_2\text{N}_3$. — Flat ndl. fr. aq.; e.s. ammonia or h. aq. — AgA, amorph. ppt., gradually changing to lust. quadratic lft.
491	287–9d.		6-Hydroxypyridine-2,5-dicarbonic Ac., $\text{HO.C}_6\text{H}_4\text{N}(\text{CO}_2\text{H})_2$. — (D) FeSO_4 gives intense yel. color. — Glassy grains alm. i. c. aq., alc., eth. — BaA, silky ndl. fr. BaCl_2 & sol. of free acid, alm. i. aq.
492	290	"Reddins litmus"	2,3-Diphenyl-5-hydroxytriazole, $[\text{NPh.N:C(OH).N:CPh}]$. — Ndl. fr. alc. D.s. alc., eth., e.s. dil. alk. or warm dil. Na_2CO_3 . — AgA. H_2O , amorph. ppt. — Benzoyl deriv., flat ndl., m.p. 134°.
493	291d.		Dimethylasparagine, $\text{Me.NH.CO.CHMe.CH}_2\text{CO}_2\text{H}$. — Lust. tbl. fr. dil. alc. — Opt. i.
494	291–2		Quinoline-6-carbonic Ac., $\text{C}_6\text{H}_5\text{N.CO}_2\text{H}$. — Cryst. powd. Softens abt. 280°. V.d.s. h. aq.; more s. h. alc.; e.s. dil. ac. or alk.
495	292 \pm 2u.c. d.; (s. cap.)	61.5 †	l-Aspartic Ac., $\text{CO}_2\text{H.CH}(\text{NH}_2)\text{CH}_2\text{CO}_2\text{H}$. — (D) Gives 10.54% amino N by Van Slyke's method (T. 2.5)! — Cryst. fr. h. aq. in thin lust. lft. Odorless. Taste sour. S. in 256 pt. aq. at 10° or in 18.6 pt. at 100°; i. abs. alc. — $[\alpha]_D = +4.36^\circ$ fr. c. aq. sol., decreasing w. rise in temperature & becoming -1.86° at 90° C. — No definite m.p. or temperature of decomposition can be observed in open tubes. The m.p. here given was obtained using a 5-mm. layer of substance in s. cap. 1 mm. in diameter & 5 cm. in length. (Cf. Ber. 28, 1632, & foot-note, Vol. I, 219, for additional information on this point.)
497		"Dec. car- bonates"	Glutazine, $\text{NH:C:(CH}_2\text{CO}_2)_2:\text{NH}$. — (D) Aq. sol. colored red by FeCl_3 , changing to green on warming. — Tbl. fr. aq. D.s. c. aq.; i. alc.; s. alk., dil. mineral acids.
498	301–2d.	139	6-Hydroxypyridine-3-carbonic Ac., $\text{HO.C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Ndl. fr. aq. D.s. h. aq.; alm. i. alc., eth., chlf., bz. — Ignition w. Zn dust gives pyridine (No. 2.1125).
499	303–4		p-Aminohexahydrobenzoic Ac., $\text{NH}_2\text{C}_6\text{H}_{10}\text{CO}_2\text{H}$. — Lft. fr. alc. Alm. i. alc., eth., lgr.
500	315, 298–9, (s. cap.); 309.5; 317	123 $k.10^4 = 1.1$	Isonicotinic Ac., 4-(γ)-Pyridinecarbonic Ac., $\text{C}_6\text{H}_4\text{N.CO}_2\text{H}$. — Sbl. in tbl. before melting. Ndl. fr. aq. D.s. c. aq.; e.s. h. aq.; alm. i. h. alc.; d.s. eth., bz. — Ignition w. CaO gives pyridine. — CaA. $4\text{H}_2\text{O}$, silky ndl., mod. s. aq. — CuA. $4\text{H}_2\text{O}$, blue green lft.
502	abt. 315–20d.		1,3,9-Trimethyluric Ac., 1,3,9-Trimethyl-2,6,8-trioxypurine, $\text{C}_8\text{H}_{10}\text{O}_3\text{N}_4$. — (D) Gives murexide react. (T. 2.20). — Ndl. fr. h. aq. V.d.s. c. aq.; s. in abt. 30 pt. h. aq.; d.s. h. alc., chlf.; i. eth. Reduces ammon. AgNO_3 sol. on boiling.
503	316	$k.10^4 = 3.7$	2,6-Dimethylpyridine-3,5-dicarbonic Ac., Lutidinedicarbonic Ac., $\text{Me}_2\text{C}_6\text{H}_3(\text{CO}_2\text{H})_2\text{H}_2\text{O}$. — Ndl. v.d.s. c. aq., alc., eth. — Ignition w. CaO gives 2,6-dimethylpyridine (No. 2.1169).

(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
504	320 browning		Isocarbostyryl-3-carbonic Ac., $C_6H_4\cdot CH\cdot C(O_2H)\cdot NH\cdot CO^+$. — Silky ndl. fr. acetone. Sbl. Dec. below m.p. to CO_2 & isocarbostyryl (No. 2.2486-1). D.s. alc., eth., chlf., bz. — Dist. w. Zn dust gives isoquinoline (No. 2.1365). — $Ag\bar{A}$, gelat. ppt.
505	323	$\mu(256) = 327$	Pyridine-2,3,5-tricarbonic Ac., $C_6H_5N(CO_2H)_3$. — \oplus Colored red by $FeSO_4$. — Spherical aggregates d.s. c. aq.; e.s. alc. — Ignition w. CaO gives pyridine (No. 2.1125).
506	323 w. efferv.	$k\cdot 10^3 = 1.5$	Pyridine-3,5-dicarbonic Ac., $C_6H_5N(CO_2H)_2$. — Alm. i. aq. Salts generally i. — Heat gives CO_2 & nicotinic ac.
507	"Browns at 340 & dec. in melting"	Reacts weakly acid	3,9-Dimethyluric Ac., $C_8H_{10}O_4N_4$. — \oplus Reduces ammon. $AgNO_3$ on boiling. — NH_4 salt separates free ac. on boiling aq. sol. — Small obliquely truncated pr. fr. h. aq. S. in 195 pt. boiling, or 1885 pt. c. aq.; alm. i. alc.; i. eth.; e.s. conc. H_2SO_4 . — Heated w. alk. gives NH_3 , NH_2Me , & CO_2 .
508	abt. 340c. w. efferv.		1,7,9-Trimethyluric Ac., $C_8H_{10}O_4N_4$. — \oplus Does not reduce ammon. $AgNO_3$ on boiling. — Lust. ndl. fr. aq. or alc.; s. in abt. 19 pt. h. aq.; s. h. alc.; e.s. conc. HCl or dil. alk. — Ag salt, felted ndl., e.s. ammon.
—	d. abt. 300		Citrazinic Ac. — Cf. No. 2.3932.
509	a. 360d.		3-Methyluric Ac., $C_6H_7O_3N_4 \cdot H_2O$ (at 100°). — \oplus Gives murexide react. (T. 2.20). — Pr. fr. h. aq.; s. in 262 pt. boiling aq.; alm. i. c. aq., alc.; e.s. $NaOH$, conc. H_2SO_4 . — Conc. HCl at 170° gives CO_2 , NH_3 , NH_2Me & glycine.
510	370-80w. efferv. (r.h.)		3,7,9-Trimethyluric Ac., $C_8H_{10}O_4N_4$. — \oplus Gives murexide react. (T. 2.20). — Ndl. softening fr. 350°. S. in 130 pt. boiling aq.; d.s. alc., chlf.; e.s. conc. HCl , dil. ammon. or alk. — Gives ppt. of $Ag\bar{A}\cdot 2H_2O$ w. ammon. $AgNO_3$; ndl. unstable to light when moist.
511	d.w.m. 370- 80(r.h.)		7-Methyluric Ac., $C_6H_7O_3N_4$. — \oplus Gives murexide react. (T. 2.20). — Lft. fr. h. aq. w. $1H_2O$ (air-dry). — S. in abt. 80 pt. boiling aq.; s. alk. or ammon. — Ammon. sol. w. little $AgNO_3$ gives gelat. ppt. of Ag salt, stable on boiling.
512	abt. 390d.		1,7-Dimethyluric Ac., $C_7H_9O_3N_4$. — Cryst. fr. h. aq. S. in 105-14 pt. boiling aq. — Ammon. sol. w. little $AgNO_3$ gives gelat. ppt. strongly colored by boiling.
513	abt. 400 w. efferv.		1,9-Dimethyluric Ac., $C_7H_9O_3N_4$. — \oplus Gives murexide react. (T. 2.20) & reduced ammon. $AgNO_3$ on warming. — Tbl. fr. aq. S. in 360 pt. boiling aq.; e.s. dil. alk. or ammon.
514	Browns at 400; carbon- izes higher		1-Methyluric Ac., $C_6H_7O_3N_4$. — \oplus Gives murexide react. (T. 2.20). — Mic. ndl. s. i. 2050 pt. boiling aq.
515	abt. 410d. (r.h.)		1,3-Dimethyluric Ac., $C_7H_9O_3N_4$. — Ndl. w. $1H_2O$ fr. h. aq. S. alc., h. aq.; alm. i. chlf.; i. eth.; e.s. ammon. — Ammon. sol. w. $AgNO_3$ gives gelat. ppt. which dec. w. separation of Ag on boiling.
516	alm. un- changed at 360(v.s.l. browned)	$k\cdot 10^3 = 1.5$ †	Uric Ac., 2,6,8-Trioxypurine, $C_6H_4O_3N_4$. — \oplus Gives the murexide react. (T. 2.20)! — Prepare a sat. sol. in boiling aq. Filter hot. Cool & shake. The scanty ppt. viewed under microscope will consist exclusively of minute, transparent, rectangular plates. (The "whetstone-shaped" cryst. & rosetted aggregates figured in works on urine analysis are characteristic of the slightly impure acid only.) — Odorless, cryst. powd. S. in 10,000-40,000 pt. c. aq., or 1600 pt. h. aq.; i. alc., eth.; s. alk. or alk carbonates. Taste of sat. aq. sol. v. faintly sour. — Dis. a few mg. in 1 cc. Na_2CO_3 sol. Bring a drop of sol. on filter paper that has been moistened w. $AgNO_3$ sol. An immediate black or dark gray stain of reduced Ag will appear. — [Fr. urine.]

No.	Melting-point (C.).	Neut. Equiv., etc.	ACIDIC COMPOUNDS.—Colorless and Solid.
517	alm. un-changed at 360 in open or closed capillary	$k \cdot 10^w = 1.2 \dagger$	Xanthine, 2,6-Dioxypurine, $C_6H_4O_2N_4$. — (1) Gives VR color in murexide react. (T. 2.20)! — Fine, odorless, tasteless powd. Appears as lust. mic. plates when warm v. dil. alk. sol. is acidified w. ac. ac. & allowed to cool slowly. S. in 1336–1498 pt. aq. at 100°, or in 14,151–14,583 pt. at 16°; 100 pt. alc. dis. 0.033 pt. at 17°; v.s. alk., but pptd. by CO_2 ; s. conc. H_2SO_4 & not pptd. by aq. — (1) ("Xanthine Test.") Mix 0.001 g. w. 2 drops HNO_3 (sp. gr. 1.42). Evaporate on water-bath to dryness. Add 1 drop 10% $NaOH$ to yel. residue after cooling. A YO-O color appears, changing to VRT1 after heating 1 or 2 min. on steam-bath, and finally fading. — (2) A white ppt. appears when 5–6 drops of 5% aq. $HgCl_2$ sol. is added to sol. of 0.001 g. in h. aq. — (3) $AgNO_3$ sol. added to sol. of 0.001 g. in 1 cc. ammon. (sp. gr. 0.90) + 9 cc. aq. gives gelat. white ppt. — (4) Scattered on crucible cover w. a little bleaching powd. & NaOH sol., dark green ring forms about particles, soon changing to brown & then disappearing. [Hoppe-Seyler Handb., p. 114.]
518	Unchanged at 360	$k \cdot 10^w = 1.8 \dagger$	Cyanuric Ac., $C_3H_3O_3N_3$. — (1) Ignite abt. 0.005 g. in narrow glass ignition tube closed at lower end, and cautiously (!) observe the intensely sharp odor (like formic or gl. ac. ac.) of the cyanic ac. fumes evolved. A white sublimate will also be formed. — Cryst. fr. aq. w. $2H_2O$ in monoclin. pr. efflorescing in air. S. in abt. 400 pt. c. aq. or 300 pt. c. alc.; alm. i. eth.; s. (undec.) conc. H_2SO_4 . — Odorless. Taste of sat. aq. sol. faintly sour. — (1) Dis. abt. 0.1 g. in 5 cc. h. aq. + 6 drops conc. ammonia. Add to h. sol. $CuSO_4$ sol., drop by drop, boiling a few sec. after each addition until a distinct blue color (which, until there is x.s. of reagent, will disappear) makes its permanent appearance. Allow to cool. A compact glistening ppt. of peculiar amethystine color (RV-VR), which is best observed after collecting on small filter & washing w. a little aq., is obtained. The product $[C_3O_3N_3]_2Cu \cdot 2NH_3$ will be found under the microscope to consist of minute but well-developed cryst. of charac. appearance!

SUBORDER I OF ORDER II.
COLORLESS COMPOUNDS CONTAINING C, N, H, AND (USUALLY) O.

GENUS I, ACIDIC COMPOUNDS.

DIVISION B, LIQUID SPECIES.

No.	Boiling-point (C°).	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
519	25.2	Hydrocyanic Ac., Prussic Ac., Formonitrile, HCN.—Sp. gr., 0.6969 (18°). Solidifies at -15°; m.p., -10° to 12°.— $k.10^{\circ} = 7.2$ at 25°.—Odor characteristic, like bitter almonds. (Vapors highly dangerous! Smell the dil. sol. only!) — Misc. w. aq., alc.; e.s. eth., acetic eth. — (1) [Prussian Blue Test.] Make 2 cc. of a dil. aq. sol. of the acid strongly alk. w. NaOH, and then proceed as directed in the Ordinal Test for nitrogen. (Cf. Vol. I., p. 12.) — (2) [Sulphocyanate Test.] Add to a few cc. of a dil. aq. sol. 3 drops of KOH sol. & 10 drops yellow ammon. sulphide sol. Evap. to dryness on water-bath. Dissolve residue in 5 cc. aq. & filter to remove S. Add 1 drop FeCl_3 sol. A blood red color will appear. — [Consult Rosenthaler, Authenrieth, Gadamer, etc., for toxicology, bibliography & additional tests.]
—	101.5	Nitromethane.—Cf. No. 2.2688. (Odor ethereal.)
—	114.5	Nitroethane.—Cf. No. 2.2698. (Odor ethereal.)
—	166	Phenyl Isocyanate.—Cf. No. 2.2750. (Odor powerfully irritating!)
521	185-6c.	Dinitroethane, $\text{Me.CH(NO}_2)_2$. — [$k.10^{\circ} = 5.8$]. Sp. gr. 1.350 (23.5°). "Somewhat s. aq." Strongly acid.—KA, yel. monoclin. cryst. s. in 14.51 pt. aq. at 20°; explodes w. violence on percussion.
—	187	p-Tolyl Isocyanate.—Cf. No. 2.2777. (Odor powerfully irritating!)
523	189c.	Dinitropropane, $\text{Et.CH(NO}_2)_2$. — ["Strong monobasic ac."] Sp. gr. 1.258 (22.5°). Heated w. dil. H_2SO_4 gives NO & propionic ac.
524	197	1,1-Dinitrobutane.—Sp. gr. 1.205 (15°/4°). — KA, yel. lft. — AgA, deep yel. lft. fr. h. aq. showing blue-violet dichroism. — Salts not explosive.

SUBORDER I OF ORDER II.
COLORLESS COMPOUNDS CONTAINING C, N, H, AND (OFTEN) O.

GENUS II, BASIC COMPOUNDS.

DIVISION A, SOLID SPECIES.

No.	Melting-point (C.).	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
—	12	Triethylenetetramine. — Cf. No. 2.1412. B.p. 266–7°.
—	15	2,5-Dimethylpyrazine. — Cf. No. 2.1196–1. B.p. 155°.
527	abt. 16	Benzalhydrazine, Ph.CH:N.NH ₂ . — (P) Reduces ammon. AgNO ₃ sol. — Leafy cryst. mass. B.p. 140° (14 mm.). Dec. quickly in moist air.
528	16.5	1-Aminoundecane, C ₁₁ H ₂₃ .NH ₂ . — B.p. 233–4°.
529	17	1-Aminodecane, C ₁₀ H ₂₁ .NH ₂ . — B.p. 216–8°.
—	17.5	Phenylhydrazine. — Cf. No. 2.1369. B.p. 243.5°.
—	20	2,5,6,8-Tetramethylquinoline. — Cf. No. 2.1446. B.p. 297–300°.
—	20–2	Pyrimidine. — Cf. No. 2.1138. B.p. 123.5–4° c.
531	23–4	4-Amino-1,2,3,5-tetramethylbenzene, Isouridine, NH ₂ .C ₈ H ₁₄ .Me ₄ . — B.p. 255°. — Prim. amine.* — Acetyl deriv. (T. 2.1), ndl. e.s. alc., m.p. 210–1°, 215°.
532	26	† aa-Phenylbenzylhydrazine, Ph.N(C ₆ H ₅).N.NH ₂ . — (P) Reduces Tollen's reagent (T. 2.30) easily. — Yellowish oil solidifying w. some difficulty. Dist. in vacuo only. V.d.s. aq.; e.s. most organic solvents. — (D) Add 1 drop benzaldehyde to sol. of 0.05 g. substance in 7 cc. alc. + 3 cc. aq. Boil 30 sec. Cool & shake. Filter. Wash cryst. w. 5 cc. dil. alc. Recryst. fr. 20 cc. alc. + 10 cc. aq. Wash w. 3 cc. dil. alc. Dry 30 min. at 100°. The product, benzaldehydephenylbenzylhydrazone, forms YT'4 ndl. of m.p. 110° u.c.
533	27–8	1-Aminododecane, C ₁₂ H ₂₅ .NH ₂ . — B.p. 134–5° (15 mm.). — Prim. amine.* — B ₂ .H ₂ PtCl ₆ , § yel. lft. fr. alc., d. 215° (T. 2.14).
534	27	1-Aminotridecane, C ₁₃ H ₂₇ .NH ₂ . — B.p. 265°. — Prim. amine.* — Lust. fatty mass w. basic odor. E.s. alc., eth. — B ₂ .H ₂ PtCl ₆ , § lust. yel. lft. fr. alc., d. 233°. (T. 2.14).
535	27	Damascenine, C ₁₀ H ₁₁ O ₂ N. — (P) Solutions all show strong blue fluorescence! — Yellowish, sl. fluorescent cryst. of narcotic odor. Alm. i. c. aq.; e.s. alc., eth., chlf. — B.Pk,† ndl., r.s. h. aq.; m.p. 180–90°, becoming blue. — [Fr. seeds of Nigella damascena.]
536	27	Quinoxoline, "C ₈ H ₆ .N:CH.CH:N". — B.p. 225–6°. — Cryst. V.s. c. aq., alc., eth., bz. Separates fr. aq. sol. on warming. — B ₂ .H ₂ PtCl ₆ (100°), § ppts. in fine yel. ndl.
—	abt. 27	Putrescene. — Odor basic. S. aq. — Cf. No. 2.1201.
538	28	Iminoethyl Alcohol, NH(CH ₂ .CH ₂ .OH). — B.p. 270° c. (748 mm.). — Fumes in air; v. deliq.; caustic. Misc. w. aq., alc., eth. Not volat. w. st. — B.HAuCl ₄ , ndl., m.p. abt. 122° (T. 2.13).

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. ‡ = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
539	28-9	1,7-Diaminoheptane, $C_7H_{14} \cdot (NH_2)_2$. — B.p. 223-5°. — Prim. amine.* — M.p. picrate, † 175° d.
540	31	Oxyheptaisobutylideneamine, $C_{23}H_{24}ON_6$. — V.d.s. c. aq.; e.s. alc., eth. — Split at once to isobutyric aldehyde & NH_2 by dil. H_2SO_4 .
541	32	† Phenylbenzylamine, $Ph.NH_2C_6H_5$. — B.p. 298-300°. — (P) Sec. amine.* — 4-Sided pr. fr. alc. — $B.HCl$, lft., m.p. 197°. — $B_2H_2PtCl_6$, § yel.-red lft., mod. s. aq., m.p. 155°.
542	32	2-Amino-1,3-dimethyl-5-tert-butylbenzene, $(NH_2)(Me)_2(C_6H_4)C_9H_{18}$. — B.p. 256°. — (P) Prim. amine.* — Pr. fr. lgr. V.s. alc., eth. — Acetyl deriv. (T. 2.1), m.p. 81°. — Thiourea deriv., m.p. 234°.
543	33-0.5	Methyl-o-anisidine, $MeO.C_6H_4.NHMe$. — B.p. 218-20°. — (P) Sec. amine.* — Reduces ammon. $AgNO_3$. Aq. sol. colored brown-red by oxidizing agents.
544	34	† Pilocarpine, $^2CH_2CO.O.CH_2CH_2$ — CH_2 — $^2C:CH.N:CH.NMe$. — [Narcotic alkaloid fr. leaves of Pilocarpus Jaborandi, P. microphyllus & P. pennatifolius. Usually obtained as deliq. syrup, difficult to cryst.] — (P) Add to sol. of 0.001 g. $B.HCl$ in 5 cc. aq., 2 cc. of 3% H_2O_2 & 1 drop HCl (sp. gr. 1.12). Then add 1 cc. bz. and, without shaking!, 4 drops of $K_2Cr_2O_7$ sol. (1 : 300). A VB ring forms at junction of layers. It soon fades on standing, or immediately on shaking. The color develops even at a dilution of 1 : 50,000. — E.s. aq., alc., chlf.; d.s. eth.; i. pet. eth. May be shaken out from weak ammon., but not from NaOH sol., w. chlf. — $[\alpha]D = +101.6^\circ$ (in 7.24% sol.). N. Eq., 208 (Congo indicator). — Odorless. Taste, bitter. — (P) Dis. 3 small drops B in 3 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 4 cc. sat. aq. sol. of picric ac. Heat to boiling to dis. ppt. & allow to cryst. by slow cooling. Recryst. fr. 5 cc. boiling 50% alc. Filter. Wash w. 1 cc. 50% alc. Dry the broad, lust. Y ndl. 15 min. at 100°. The picrate obtained softens at 156° & melts at 158-60° u.c. to a clear yel. liq. — $B.HAuCl_4 \cdot H_2O$, lemon-yel. ndl., m.p. 100° (T. 2.13). — 1 or 2 drops of 1% sol. on the eye contracts the pupil. Resembles nicotine in toxic action.
545	34	† Ethyl β-Aminocrotonate, $MeC(NH_2):CH.CO_2Et$. — B.p. 210-15° d. — (P) Prim. amine.* — Thick lust. pr. i. aq.; e.s. alc., eth. — When fused yields isomer, m.p. 23-4°, which reverts to original compound on contact w. cryst. of latter.
546	35	1',4'-Diaminoxylene, $(NH_2.CH_2)_2.C_6H_4$. — (P) Prim. amine.* — Cryst. — B_2PK , † or. ndl.
547	36	6-Amino-1,2,4-trimethylbenzene, $NH_2.C_6H_3(Me)_3$. — (P) Prim. amine.* — Cryst.
548	36.5	n-Dioctylamine, $(C_8H_{17})_2.NH$. — B.p. 297-8°. — (P) Sec. amine.* — Odor like tallow. — I. aq.; e.s. alc., eth. — $B_2H_2PtCl_6$, § scales fr. alc.
549	36.5	1-Aminopentadecane, $C_{15}H_{31}NH_2$. — B.p. 298-301°. — (P) Prim. amine.* — Waxy. Odor, peculiar. — $B.HCl$, fatty feel, e.s. alc., d.s. aq.
550	37	1-Aminotetradecane, $C_{13}H_{29}NH_2$. — B.p. 162° (15 mm.). — (P) Prim. amine.*
551	37-7.5	1,9-Diaminononane, $NH_2.(CH_2)_9.NH_2$. — B.p. 258-9°. — (P) Prim. amine.* — Absorbs CO_2 & aq. — $B_2H_2PtCl_6$, §
552	38	6-Methyl-1,2,3,4-tetrahydroquinoline, $C_{10}H_{12}N$. — B.p. 262° (712 mm.). — (P) Sec. amine.* — Nitroso deriv., tbl. fr. eth., m.p. 65°. — $B.HCl$, silky ndl., v.s. aq., alc.; i. eth.; m.p. 189°.
553	39	Tricytylamine, $(C_{15}H_{31})_3N$. — (P) Tert. amine.* — Ndl. — $B_2H_2PtCl_6$, § yel. ppt.
554	39.6; 35	Triacetonamine, $^2NH.CMe_2.CH_2.CO.CH_2.CMe_2$. — Ndl. w. ammon. camphorous odor fr. dry eth. E.s. aq., eth. Cryst. fr. moist eth. w. $1H_2O$ & m.p. 59°. "Reacts feebly alk." — $B_2H_2PtCl_6 \cdot 3H_2O$, § cryst. fr. h. aq. in golden ndl., of which 9.57 g. dis. in 100 pt. aq. at 14°. — Gives nitroso deriv., ** ndl. fr. dil. alc., m.p. 72-3°.
555	38-40	Ephedrine, $C_{10}H_{15}ON$. — [Mydriatic alkaloid fr. Ephedra vulgaris L.] — B.p. 255°. $[\alpha]D^{20} = -6.3^\circ$ (in abt. 3.5% alc. sol.). Cryst. mass. S. aq., alc., chlf., eth. — Secondary base. — $B.HCl$, ndl., s. aq., m.p. 210°. (Trade name "Mydrin.") — Alkaloidal color reactions not characteristic. — $B_2H_2PtCl_6$, § long felted e.s. ndl. fr. conc. aq. sol., m.p. 183-4°.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
556	39-40	1,6-Diaminohexane, $\text{NH}_2(\text{CH}_2)_5\text{NH}_2$. — B.p. abt. 200°. — \textcircled{P} Prim. amine.* — Odor, basic like piperidine. Sbl. in ndl. — $\text{B}_2\text{HCl} \cdot 4\text{HgCl}_2$, d.s. lft. (charac.). — Picrate † melts abt. 220° d.
557	40-1	p-Methylbenzylhydrazine, $\text{Me.C}_6\text{H}_4\text{CH}_2\text{NH.NH}_2$. — B.p. 135° (18 mm.). — \textcircled{P} Prim. amine.* — Cryst. mass. — $\text{B.Pk.} \ddagger$ ndl., m.p. 144°.
558	40-1	3,7-Dimethyl-2-ethylquinoline, $\text{Me.C}_6\text{H}_4\text{C}_2\text{HN(Me, Et)}$. — B.p. 288-92°. — \textcircled{P} Tert. amine.* — Trimet. lft. fr. lgr. — $\text{B.Pk.} \ddagger$ yel. ndl. fr. alc., alm. i. aq.; e.s. h. alc., m.p. 219-20°. — $\text{B}_2\text{H}_4\text{PtCl}_4 \cdot 2\text{H}_2\text{O}$, § yel.-red cryst. d.s.c. aq.
559	40-1	β -Coniceine, 1- α -Allylpiperidine, $\text{C}_6\text{H}_5\text{NH.C}_6\text{H}_4$. — B.p. 168-9°. — \textcircled{P} "Strong" sec. base.* — Ndl. d.s. aq.; e.s. alc., eth. — $[\alpha]_D^{25} = -50.64^\circ$. — $\text{B.HAuCl}_4 \parallel$ ppt. oily & solidifies to tbl., m.p. 123°.
560	41	† Dimethyl-p-phenylenediamine, $\text{Me}_2\text{N.C}_6\text{H}_4\text{NH}_2$. — B.p. 262° c. — Rather unstable & when sl. impure liable to separate fr. sol. of its salts upon addition of alk. as a dark oil, v.d. to solidify. — Ndl. fr. bz. + lgr. E.s. aq.; v.s. alc., eth. — \textcircled{P} Dis. abt. 0.001 g. in 10 cc. aq. + 1 cc. HCl (sp. gr. 1.2). Add 3 or 4 drops sat. H_2S sol., & then 1 drop 10% FeCl_3 sol. A color, at first GT1, changing rapidly to BG, and within 4-5 min. to intense pure blue (B), appears, and on longer standing becomes too deep for comparison. — \textcircled{D} Dis. abt. 0.06 g. in 3 cc. alc. Add 5 drops benzaldehyde & 2-3 drops 10% NaOH sol. A heavy ppt. of golden scales separates. Filter. Wash w. 1 cc. alc. Cryst. fr. h. 50% alc. Dry on porous tile at room temp. for 30 min. The product, benzylidene-dimethyl-p-phenylenediamine, $\text{Me}_2\text{N.C}_6\text{H}_4\text{N:CHPh}$, melts at 97.8-98° u.c.
561	41-2	Tropinone (Tropinol), $\text{C}_6\text{H}_5\text{ON}$. — B.p. 224-5°. — \textcircled{P} Strong tert. base,* fuming w. HCl . — Vapors evolved on strongly heating in tube give splinter react. T. 2.24. — V.s. aq., alc.; less s. lgr. — Reduces ammon. AgNO_3 sol. — B.HCl , pr. fr. alc., m.p. 188-9° d. — $\text{B.Pk.} \ddagger$ lust. yel. ndl., m.p. 220° d.; v.d.s. c. aq.; alm. i. c. alc., eth. — $\text{B.Mel.} \ddagger \ddagger$ m.p. 263-5° d. Is prepared fr. components in dil. alc. sol. (Ber. 29, 400, 947), & warmed w. aq. KOH sol. evolves dimethylamine in violent react.
562	41-2	Benzoyltropeine, $\text{C}_{10}\text{H}_{11}\text{O}_2\text{N}$. — \textcircled{P} Tert. base.* — Cryst. mass of narcotic odor. — $\text{B}_2\text{H}_2\text{O}$, silky lft., m.p. 58°. — $\text{B}_2\text{H}_4\text{PtCl}_4 \cdot 2\text{H}_2\text{O}$, §, yel. rhomb. ft., d.s. c. aq.
563	42	Cetylaniline, $(\text{C}_{10}\text{H}_{11})_2\text{NHPH}$. — Scales fr. alc.; e.s. alc., eth.; i. aq.
564	42 (r.h.)	β -Methylhydroxylamine, Me.NH(OH) . — \textcircled{P} Boiling w. Fehling's sol. gives NH_3 & Me.NH_2 . — Pr. v.s. aq., alc.; d.s. eth., lgr. — Dec. on keeping. — B.HCl , m.p. 88-90°. — $\text{B.Pk.} \ddagger$ yel. tbl., v.s. aq., m.p. 128-30°.
565	42	Trimethyleneethylenediamine, $\text{NH.CH}_2\text{CH}_2\text{CH}_2\text{NH.CH}_2\text{CH}_2\text{NH}_2$. — B.p. 168°. — \textcircled{P} Sec. base.* — Deliq. Fumes in moist air. V.s. aq.; s. alc.; less s. eth. — $\text{B}_2\text{HCl} \cdot 4\text{HgCl}_2$, s. h. aq., m.p. 243°. — Gives nitroso deriv., ** m.p. 92°.
566	42.9	† p-Toluidine, $\text{Me.C}_6\text{H}_4\text{NH}_2$. — B.p. 200.5° c. — Prim. amine giving N in T. 2.4. — $k_{10} = 2.2$. — Odor, peculiar, like aniline. S. in 135 pt. aq. at 20.8°. Taste of c. sat. aq. s., burning (Pungency No. 2-3, T. 2.29). — \textcircled{P} (1) Dis. fragment of size of grain of wheat in 10 cc. c. aq. + 1 drop HCl (sp. gr. 1.2). Add 1 drop 10% FeCl_3 sol. & heat gently, so that boiling shall begin in 45 sec. Keep just below boiling temperature for 1 min. more. Cool without delay. The color will now be red-orange (RO). After standing 1 hour it changes to RT1 and later approaches VRT1. — (2) Drop small crushed fragment into 10 cc. of mixture of 2 vol. conc. H_2SO_4 + 1 vol. aq. Add 2 drops HNO_3 (sp. gr. 1.42) without shaking. A distinct blue ring appears at once. Shake and view sol. by direct transmitted light from sky. Color changes within 60 to 90 sec. fr. BV through V, RV & VR to R. — (1) To 0.05 g. in small t.t., add 12 drops acetic anhydride & heat to boiling. Dis. product by boiling in 6 cc. aq. Unless sol. is clear, filter rapidly while hot through small filter. Cool. Shake vigorously. Filter. Wash w. 2 cc. aq. Recryst. fr. 2 cc. h. aq. Dry 15 min. on porous tile at 100°. The product, p-acet toluidine, $\text{Me.C}_6\text{H}_4\text{NH.CO.Me}$, is obtained in lust. white ndl., m.p. 145.5-6° u.c. (147.8-8.3° c.).
567	42.9	† Thalline, p-Methoxytetrahydroquinoline, $\text{MeO.C}_6\text{H}_4\text{N}$. — Sec. base* of cumarin-like odor. Cryst. v.d.s.-i. aq.; e.s. alc., eth., bz. Taste of c. sat. aq.

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(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
		<i>sol. burning (Pungency No. 3, T. 2.29) w. persistent unpleasant after-taste.</i> — (P) Dis. abt. 0.002 g. + 2 drops HCl in 10 cc. aq. Add 1 drop 10% FeCl ₃ sol. After 1 min. the color becomes GS2 by "reflected light" (p. 232, Vol. I), but soon becomes deep red by direct transmitted "sky light," though still green in <i>v.</i> thin layers. Heat destroys this final color! — (P) Add 3 cc. c. sat. aq. picric ac. sol. to sol. of 0.1 g. thalline in 3 cc. alc. Shake vigorously & <i>v.</i> persistently! (There is a tendency for the sol. to remain supersaturated, but this once overcome, the ppt. formed is heavy.) Filter. Wash <i>w.</i> 2 cc. alc. Recryst. fr. 2 cc. h. alc., chilling & shaking persistently. Dry 15 min. at 100°. The product, thalline picrate, is obtained in minute yel. (OYT ₁ -YT ₁) ndl., m.p. 157-60° d. u.c. (159.5-62.5° c.)
568	43	Acetylacetonamine, Me.CO.CH:C(NH ₂)(Me). — B.p. 209°. — (P) Prim. amine.* — Deliq. cryst. e.s. c. aq. — Warming <i>w.</i> dil. acids or aq. dec. to NH ₂ & acetylacetone (Vol. I).
569	42-4	1-Dimethyl-1,2,4-triaminobenzene, Me ₂ N.C ₆ H ₃ (NH ₂) ₂ . — (P) Soon turns dark blue in the air. — Asbestos-like ndl. fr. lgr. V.s. aq.
570	44	5-Methylamino-1,2,4-trimethylbenzene, Methylpseudocumidine, NHMe.C ₆ H ₃ Me. — B.p. 237°. — (P) Sec. amine.* — B ₂ H ₅ PtCl ₆ .§
571	44	d-Lupanine, C ₁₅ H ₂₂ ON ₂ . — [Fr. seeds of blue lupine, Lupinus angustifolius.] — (P) Tert. base.* — Ndl. or syrup. S. aq., alc., eth., chlf.; less s. lgr. Odor, faint, coniine-like. Taste bitter. Fumes <i>w.</i> HCl. — B.HCl.2H ₂ O, rhomb. cryst. fr. aq.; anhydrous (dried 12 h. over H ₂ SO ₄), m.p. 127°. — B.MeI,†† cryst., m.p. 239-41°.
572	44	3,8-Dimethyl-2-ethylquinoline, Me ₂ C ₆ H ₄ C ₆ H ₃ MeEt. — B.p. 279-80° (717 mm.). — Monoclin. cryst. fr. lgr. — (P) Tert. base.* — BPk,† thick yel. ndl. fr. alc., alm. i. aq., m.p. 187°.
573	45	2,4'-Diaminobiphenyl, Diphenyline, β-Benzidine, NH ₂ .C ₆ H ₄ .C ₆ H ₄ .NH ₂ . — B.p. 363°. — (P) Prim. amine.* — Long ndl. Alm. i. aq.; e.s. alc., eth. — B.H ₂ SO ₄ , unlike benzidine sulphate, v.s. aq. — Diacetyl deriv.¶ (fr. acetic anhydride), cryst., m.p. 202°.
574	45.5	o-Aminobiphenyl, Ph.C ₆ H ₄ .NH ₂ . — B.p. 299°. — (P) Prim. amine.* — E. vol. w. st. — B ₂ H ₅ PtCl ₆ .4H ₂ O,§ lust. or. lft. d.s. aq. — Acetyl deriv.¶ (fr. acetyl chloride) pr. fr. alc., m.p. 117.5° c.
575	45.6	Cetylamine, C ₁₈ H ₃₈ .NH ₂ . — B.p. 330°. — (P) Prim. amine.* — Silvery lft.
576	45	o-Dimethylaminophenol, NMe ₂ .C ₆ H ₄ .OH. — B.p. 199-200°. — (P) Gives red-violet color <i>w.</i> FeCl ₃ sol. — Odor penetrating, tar-like. E. vol. w. st. Rhomb. pr. v.d.s. h. aq.; e.s. alc., eth., NaOH sol.
577	45-6	Methylenedibenzylamine, (Ph.CH ₂ .NH).CH ₂ . — B.p. 225-30° d. — (P) Sec. amine.* — Cryst. i. aq.; s. alc., eth., bz. — B ₂ HCl, lft., m.p. 240-2°.
578	abt. 46	2,6-Dimethylphenylhydrazine, Me ₂ C ₆ H ₃ .NH.NH ₂ . — Ndl. fr. lgr. Unstable. Prob. reduces Fehling's sol.
579	abt. 46	β-Propylhydroxylamine, Pr.NH(OH). — (P) Reduces Fehling's sol. easily. — "Reacts basic." Ndl. fr. eth. E.s. except in lgr.
580	46.5	Tetrahydro-α-naphthoquinoline, C ₁₅ H ₁₂ N. — (P) Sec. amine.* — Alc. sol. fluor. intense blue! — Lust. lft. fr. eth. D.s. c. aq.; e.s. alc. — B.HCl, glassy pr., m.p. 260-1°. — Nitroso deriv.** C ₁₅ H ₁₂ N.NO, glassy yel. pr. fr. lgr.; v.s. aq., m.p. 59.5°.
581	46	2,6,8-Trimethylquinoline, Me ₂ .C ₆ H ₃ C ₆ H ₃ N.Me. — B.p. 260° (719 mm.). — Tert. amine.* — Monoclin. pr. fr. lgr. I. aq.; v.s. alc., eth., lgr. V. vol. w. st. — B.Pk,† long yel. ndl. fr. dil. alc., m.p. 185°.
582	46	Dimethyl-β-naphthylamine, C ₁₀ H ₇ .NMe ₂ . — B.p. 305°. — (P) Tert. amine.* — B ₂ H ₅ PtCl ₆ ,§ d.s. h. alc.
583	47	Pyrazine, Hexadiazia(1,4)-triene(1,3,5), ^c CH:CH.N:CH.CH:N ^c . — (P) B.p. 118° c. — Pr. fr. aq.; tbl. fr. eth. Misc. aq.! e.s. alc., eth., etc. "Weak base." Fumes <i>w.</i> HCl. Na + alc. gives piperazine (No. 2.765). — B.HCl, lust. deliq. pr. — B.HgCl ₂ , lust. monoclin. pr. fr. aq., or cubes fr. HCl, m.p. 273° <i>w.</i> efferv. — B.Pk,† yel. pr. v.d.s. c. aq., m.p. 156°.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
584	47-8	2,6-Dimethylpyrazine, $\text{^TCH:CM}\text{e.N:CM}\text{e.CH:N^T}$. — Monoclin. pr. V.s. aq. alc., eth. Volatile at ordinary temperature. — B.Pk,‡ cryst. fr. aq., m.p. 175-6°. — [Produced by action of ammonia on grape sugar.]
585	48.2-0.5	Decahydroquinoline, Hexanohexazane(2,3), $\text{C}_8\text{H}_{17}\text{N}$. — B.p. 204° (th.i., 714 mm.). — ⊖ Sec. amine.* — Pr. fr. lgr. E.s. h. aq.; v.s. alc. Vol. w. st. Odor like coniine. "Strongly alkaline." Reduces ammon. AgNO_3 , sol. — B.Pk,‡ lust. ndl. fr. chlf. + lgr., m.p. 151-2°.
586	48	Pseudopelletierine, N-Methylgranatone, $\text{C}_8\text{H}_{15}\text{ON}$. — B.p. 246°. — ⊖ Tert. amine.* — Tbl. fr. aq. or pet.-eth. V.s. aq., alc., eth., chlf.; d.s. pet.-eth. "V. strong base." — $\text{K}_2\text{Cr}_2\text{O}_7 + \text{H}_2\text{SO}_4$ gives green coloration. — KHg iodide gives light yel., & tannin, muddy white, ppt. — B.MeI,‡‡ colorless cubes fr. dil. alc., not melted at 280°.
587	49	† 1-Aminoheptadecane, $\text{C}_{17}\text{H}_{34}\text{NH}_2$. — B.p. 335-40°. — ⊖ Prim. amine.* — Odorless cryst. i. aq. — B.HCl, e.s. alc., but i. aq.; d.w.m. — B. H_2PtCl_6 , § ppt. of yel. cryst. spangles.
588	49	† 4-Amino-1,2-xylene, $\text{Me}_2\text{C}_6\text{H}_3\text{NH}_2$. — B.p. 225° c. — ⊖ T. 2.2-b w. HNO_3 gives v. fugitive, but rather strong green color, changing almost immediately to light greenish brown. — Prim. amine.* — Cryst. D.s. c. aq.; e.s. lgr. — (1) Heat to boiling mixture of 0.1 g. amine + 1 cc. acetic anhydride. Dis. product in 9 cc. boiling aq. Cool & shake vigorously. An oil separates & soon solidifies. Filter. Wash w. 1 cc. c. aq. Dis. in smallest possible quantity c. alc. & ppt. w. 3 cc. c. aq. Dry 30 min. at 60°. o-Acetylalide, the product, is obtained in colorless cryst., m.p. 95.3-6° u.c. (96.1-96.8° c.). — (2) T. 2.22 gives the corresponding picramide, w. good yield, as O colored ndl., fr. gl. ac. ac., m.p. 178-9°. — B.HCl, m.p. 256°.
589	49-9.5	† α-Naphthylamine, $\text{C}_{10}\text{H}_7\text{NH}_2$. — B.p. 301° c. — ⊖ (1) Highly disagreeable, persistent, fecal indole odor! — (2) Dissolve exactly 0.001 g. in 15 cc. alc. Add 1 small drop KNO_3 , sol. & then 1 small drop HCl (sp. gr. 1.12). A VRT2-VRT1 color develops after a few seconds, becoming redder & more intense later, & finally fading rapidly. W. an x.s. of amine the color is deep red at once, dilution giving a brownish color. — (3) Heat sol. of 0.001 g. amine in 10 cc. aq. + 1 drop HCl (sp. gr. 1.12) w. 2 drops FeCl_3 , sol. Bring to boiling in abt. 1½ min. & keep near b.p. 45 sec. Collect floc. ppt. on small filter & wash w. 3 cc. aq. Ppt. is RVS1 while moist, but may be bluer (VBS1) if heating is shorter. (Ann. 183, 265.) — Fine ndl. fr. alc., usually pinkish on keeping. 100 cc. c. aq. dis. 0.167 g.; v.s. alc., eth. — (4) Convert into the picrylamide by T. 2.22. This deriv. is obtained fr. gl. ac. ac. as heavy cryst. ppt. of OR-R colored pr., m.p. 198° u.c.! — The picrate,‡ darkens fr. abt. 150°, m.p. 166° u.c. d. — The acetyl deriv.¶ cryst. fr. h. aq., m.p. 159°.
590	49-50	p-Aminobenzylaniline, $\text{NH}_2\text{C}_6\text{H}_4\text{CH}_2\text{NHPh}$. — ⊖ Prim. amine.* — Difficult to cryst. Alm. i. aq.; dil. acids; e.s. alc., eth., bz. — Dist. in vacuo splits alm. quantitatively to aniline (No. 2.1235).
591	49	† Tropacocaine, Benzoylelpseudotropine, $\text{C}_8\text{H}_{14}\text{ON.CO.Ph}$. — [Local anaesthetic fr. Javan coca leaves.] — ⊖ 3% aq. sol. brought on tongue on filter paper in manner directed in test for cocaine (No. 2.741) produces alm. immediate sensation of numbness which persists several min., but is less striking than in cocaine test. — Odorless cryst. of greasy luster fr. eth. D.s. aq.; v.s. alc., eth., chlf., lgr., bz.; s. in ammon., unlike other coca alkaloids. — Taste bitter. Opt. i. — Color react. (a), (e), (f) in T. 2.2 give colorless sol. Gives the same results as l-cocaine (No. 2.741) in the Giesel & Greither's reactions, & the tests w. chromic ac., I in KI sol., & the ethyl benzoate test. — B.HCl, m.p. 276-7° d. — (1) Dis. 0.05 g. B.HCl in 2 cc. aq. Add 3 cc. sat. aq. picric ac. sol. Wash curdy ppt. w. 2 cc. aq. Recryst. fr. 4 cc. boiling alc. Filter. Wash lust. Y cryst. w. 1 cc. alc. Dry 15 min. at 100°. The picrate, B. $\text{C}_6\text{H}_3(\text{NO}_2)_2\text{O}$ melts to dark brown liquid at 234-9° u.c. — B.HAuCl ₄ ,¶ yel. ndl., m.p. 203° d.
592	abt. 50	Pulegoneamine, $\text{^TCH}_2\text{CH}_2\text{CM}\text{e}_2\text{CH}(\text{NH}_2)\text{CH}_2\text{CHM}\text{e}^2$. — B.p. 205-10°. — ⊖ Prim. amine.*
593	50	Atroscine-Hesse, $\text{C}_{17}\text{H}_{21}\text{O}_4\text{N}$. — [Mydriatic alkaloid fr. root of Scopolia atropoides.] Cryst. w. $2\text{H}_2\text{O}$ (lost over H_2SO_4), m.p. 36-7°. E.s. alc., eth., chlf., bz., h. aq. Opt. i. — B.HAuCl ₄ ,¶ yel. lft., m.p. 201-2°.

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. ‡ = T. 2.23. ¶ = T. 2.14. || = T. 2.13. § = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§§ = T. 2.38.

No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
594	50-1	Triacetonehydroxylamine, $[\text{CH}_3\text{CO.CH}_2\text{CMe}_2\text{N}(\text{OH}).\text{CMe}_2]$. — Tbl. fr. lgr. D.s. c. aq.; e.s. conc. acids. — Reduces Fehling's sol. only on heating. — Oxalate, pr. e.s. aq., alc.; i. eth.; m.p. 85°. — Benzoyl deriv. (by boiling w. benzoyl chloride), ndl. fr. pet.-eth., m.p. 117°.
595	51	Tetramethyl-p-phenylenediamine, $\text{NMe}_2\text{C}_6\text{H}_4\text{NMe}_2$. — B.p. 280° (th.i.). — \textcircled{P} Aq. sol. deep violet-blue after short exposure to air! K_4FeC_2 gives indigo-like ppt. fr. aq. sol. of sulphate. — Lft. fr. dil. alc.; d.s. c. aq.; e.s. lgr., alc., eth.
596	51	1-Aminonaphthol(2)-ethylether, $\text{NH}_2\text{C}_1\text{H}_4\text{OEt}$. — B.p. 300-2°. — Prim. amine.* — \textcircled{P} FeCl_3 gives intense blue color w. aq. sol.! — Cryst. fr. lgr. E.s. alc. — Gives acetyl deriv., m.p. 145°. [An azo-component in dye manufacture.]
597	51.5-2	Tetrahydro- β -naphthoquinaldine, $\text{C}_1\text{H}_4\text{N}$. — \textcircled{P} Sec. amine.* Alc. sol. fluor. blue! — Glassy pr. fr. lgr. E.s. alc. — B.HCl , m.p. 239-40°; e.s. aq. — Nitroso deriv.,** straw yel. lft. fr. alc., m.p. 69-9.5°.
598	51; 52	α -Naphthoquinoline, $\text{C}_1\text{H}_4\text{N}$. — B.p. 351° c. — \textcircled{P} Tert. amine.* — Monoclin. cryst. fr. eth. Alm. i. aq.; e.s. alc., eth., bz. Faint charac. odor. Remains long liquid after fusion. — $\text{B}_2\text{H}_6\text{PtCl}_6\cdot 2\text{H}_2\text{O}$, § v.d.s. aq.; m.p. (water-free) 224°.
599	51.2	1-Phenylpyrazoline, $[\text{NPh.N:CH.CH}_2\text{CH}_2]$. — B.p. 273-4° (th.i.). — \textcircled{P} $\text{K}_2\text{Cr}_2\text{O}_7$ sol. added to v. dil. H_2SO_4 or HCl sol. soon gives a red-violet to pure blue coloration that shows presence of 1 pt. in 25,000. (Ann. 239, 196). — Oblique trimet. tbl. fr. lgr. Somewhat s. h. aq.; e.s. alc., eth., bz. E.s. conc. HCl ; ptdt. by diln. w. much aq. ("V. weak base.") — [Fr. action of phenylhydrazine on acrolein.]
600	52	1,8-Diaminooctane, $\text{NH}_2\text{(CH}_2)_6\text{NH}_2$. — B.p. 240°. — \textcircled{P} Prim. amine.* — Absorbs CO_2 fr. air. — Picrate,† m.p. 180° d.
601	53	p-Aminobiphenyl, Xenyamine, $\text{Ph.C}_6\text{H}_4\text{NH}_2$. — B.p. 302° c. — \textcircled{P} Prim. amine.* — D.s. h. aq.; e.s. alc., eth., chlf. Vol. w. st. — B.HCl , lft. d.s. c. aq.; e.s. h. aq.
602	53-4	m-Aminobenzonitrile, $\text{NH}_2\text{C}_6\text{H}_4\text{CN}$. — B.p. 288-90°. — \textcircled{P} Prim. amine.* — D.s. aq.; v.s. eth., alc., chlf. Ndl. fr. dil. alc. — Sapn. (T. 2.26) prob. gives m-aminobenzoic ac. (No. 2.259). — B.HCl , v.s. pr.
603	53	N-Phenylmorpholine, $[\text{CH.NPh.CH}_2\text{CH}_2\text{O.CH}_3]$. — B.p. 270°. — \textcircled{P} Tert. amine.* — E.s. alc., eth. E. vol. w. st.
604	54	2-Amino-4-dimethylaminotoluene, $\text{Me.C}_6\text{H}_4\text{(NH)}_2\text{(NMe}_2)$. — \textcircled{P} Prim. amine.* — Pr. fr. lgr. D.s. aq.; e.s. alc., eth., bz. [An intermediate for azine colors.]
605	54	3,6-Dimethyl-2-ethylquinoline, $\text{Me.C}_6\text{H}_4\text{C}_2\text{HN(Me.Et)}$. — B.p. 287-8° (720 mm.). — \textcircled{P} Tert. amine.* — Cryst. fr. lgr. Difficultly vol. w. st. — B.Pk ,‡ yel. ndl., m.p. 177°.
606	54.5-6.5	3,5-Diamino-4-dimethylaminotoluene, $(\text{NH}_2)_2\text{(Me).C}_6\text{H}_4\text{NMe}_2$. — \textcircled{P} Prim. amine.* — Pr. fr. lgr. — B_2HCl , pr. fr. HCl , m.p. 221-5°. — B.Pk ,‡ S-yel. ndl. fr. MeOH , d.s. c. aq., m.p. 142-3°.
607	50-60	Granatinine, Hexanohexazane(2,4), $\text{C}_6\text{H}_{11}\text{N}$. — \textcircled{P} Sec. amine.* — Ndl. of unpleasant penetrating odor. Absorbs CO_2 fr. air. — B.HAuCl_4 , yel. lft. fr. aq., m.p. 225°. — Nitroso deriv.,** scales fr. lgr., m.p. 225°.
608	55	2,6-Dimethylquinoline, $\text{Me}_2\text{C}_6\text{H}_4\text{N}$. — B.p. 259-61°. — \textcircled{P} Tert. amine.* — D.s. h. aq. — Odor like anise. — $\text{B}_2\text{H}_6\text{PtCl}_6$, § ndl. i. c. aq.; s. h. aq.
609	55	Hydrocotarnine, Methoxymethylenedioxy-N-methyltetrahydroisoquinoline, $\text{C}_{15}\text{H}_{18}\text{O}_2\text{N}_2\text{H}_2\text{O}$. — \textcircled{P} S. in conc. H_2SO_4 w. yel. color, changing to carmine red, blue-violet, & violet on warming! — Monoclin. pr. fr. eth. Opt. i. V.s. alc., eth., bz., chlf. — $\text{B}_2\text{H}_6\text{PtCl}_6$, § amorph. ppt. soon changing to or.-red pr. — [In opium.]
610	56-7	† Hyoscine, 1-Scopolamine, Sikermanine, "Hyoscine-Hesse," Ester of 1-Tropic Ac. & i-Scopoline, $\text{C}_{17}\text{H}_{21}\text{O}_2\text{N}$. — [An alkaloid of more powerful but less lasting mydriatic action than atropine or hyoscyamine. Usually obtained as amorph. varnish-like residue on evapg. sol. obtained by shaking out w. chlf. an aq. sol. of its salts to which NaHCO_3 has been added.] — Mod. s. aq.; e.s. alc., eth., chlf.; less s. pet.-eth., bz. Levo-rotatory. — \textcircled{P} Vittali's

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		react. applied as directed for Atropine (No. 2.797) gives same colors: a VR, changing alm. immediately to R, which disappears within 45 min.—Gives physiological test (1) & odor test (3) (Gulielmo's T.) as described for atropine, but does not give atropine test (4).—① (a) Prepare picrate by adding 4 cc. sat. aq. sol. of picric ac. to 0.05 g. B.HCl disd. in 2 cc. aq. Allow curdy ppt. to settle for 30 min. Filter. Wash w. 2 cc. c. aq. Recryst. by slow cooling fr. 5 cc. boiling aq. Filter. Wash the beautiful lust. Y cryst. w. 1 cc. aq. Dry 15 min. at 100°. The product, $\text{BC}_6\text{H}_4(\text{NO}_2)_3\text{O}$, melts to clear yel. liq. at 187.5° u.c.—(b) Prepare chloroaurate by adding 3 cc. gold chloride T.S. to sol. of 0.05 g. B.HCl in 2 cc. aq. Filter abundant yel. ppt. Wash w. 2 cc. c. aq. Dis. in 12 cc. boiling aq. Large lust. indented \leftarrow OY lft. separate on slow cooling. Filter. Wash w. 1 cc. aq. Dry 15 min. at 100°. The product BHAuCl_4 , melts w. efferv. to red-brown liquid at 198–9° u.c.
611	56	α -Tolylhydrazine, $\text{Me.C}_6\text{H}_4.\text{NH.NH}_2$.—② Prob. reduces Ag. in T. 2.30 instantly.—Oblique tbl. fr. lgr. E.s. alc., eth., chlf.; d.s. c. lgr.—Acetyl deriv. lft. fr. aq., m.p. 104°.—Benzoyl deriv., §§ ndl., m.p. 180°.
612	56	β -Aminopyridine, $\text{NH.C}_6\text{H}_4\text{N}$.—B.p. 204°.—Lft. fr. lgr. D.s. lgr.; v.s. alc.—B.Pk, ‡ yel. ndl., v.d.s. m.p. 216–7°.— $\text{B}_2\text{H}_2\text{PtCl}_4\text{H}_2\text{O}$, § (fr. conc. HCl), yel. triclin. pr., m.p. 231°; 227–8° d.
613	56–7	3-Methyl-2-ethylquinoline, $(\text{Me.Et})\text{C}_6\text{H}_4\text{N}$.—B.p. 268–9° (711 mm.)—③ Tert. amine.*—Monoclin. pr. fr. eth. R.d.s. aq.; e.s. alc., eth., bz.— $\text{B}_2\text{H}_2\text{PtCl}_4$, §, or w. $2\text{H}_2\text{O}$, cryst., m.p. 238° d.
614	57.7 u.c.	† p-Anisidine, $\text{MeO.C}_6\text{H}_4\text{NH}_2$.—B.p. 245° c.; 239.7° c.— $k.10^9 = 1.5$.—④ (1) Warm sol. of abt. 0.002 g. in 10 cc. aq. + 1 drop HCl w. 1 drop 10% FeCl_3 sol. for 15 sec. Cool. An intense VR–RV color appears. (Avoid too long heating.)—(2) T. 2.2–b w. HNO_3 gives streaks of ORT 2 & RVT 2 , becoming RVT 2 on stirring, & after standing some time, B.—Thick sheaves of white ndl. fr. h. aq. on cooling. Odor aniline-like when strongly heated.—⑤ Mix 0.1 g. w. 6 drops acetic anhydride. Heat to boiling. Dis. product in 3 cc. boiling aq. Cool. Shake vigorously. Filter. Recryst. fr. 2 cc. h. aq. Dry 15 min. at 100°. The product, p-acetanisidine, is obtained in white scales, m.p. 125–6° u.c. (126.6–7.6° c.).
615	57–8	β -Naphthylpiperidine, $\text{C}_{10}\text{H}_7\text{C}_6\text{H}_5\text{N}$.—Pr. fr. lgr. E.s. alc., eth., bz., lgr., chlf.—B.Pk, ‡ yel. cryst. ppt. m.p. 188°, d.s. h. alc.
616	57	β -Benzylhydroxylamine, $\text{Ph.CH}_2\text{NH(OH)}$.—⑥ Reduces Fehling's sol. in the cold.—Ndl. fr. lgr. S. aq.—⑦ Convert into nitroso deriv., ** m.p. 77–8°, by adding to sol. of 1 g. B.HCl in 25 cc. aq. at 0°, 0.45 g. 90% NaNO_2 in c. aq. sol. Dis. ppt. in eth. & ppt. by addition of lgr. (Ann. 263, 217).
617	57	Isovalerianic-aldehyde-ammonia, $\text{C}_6\text{H}_5\text{CH(OH)(NH}_2\text{)}\cdot 7\text{H}_2\text{O}$.—⑧ Dil. H_2SO_4 liberates isovalerianic aldehyde (fruity odor) in the cold. Separates in 2 layers, anhydrous compound & aq., in melting.
618	58–9 (Sinters at 50)	Benzhydrylhydrazine, $\text{Ph}_2\text{CH.NH.NH}_2$.—⑨ Reduces c. ammon. AgNO_3 or Fehling's sol. slowly.—Cryst. mass. Odor, basic. Vol. w. st.—B.Pk, ‡ green-yel. ndl. fr. alc., m.p. 160° d.
619	58–9	Cinchoninic, Cinchotoxine (?), $\text{C}_{15}\text{H}_{24}\text{ON}_2$.—Pr. s. in 564 pt. aq. at 18°. E.s. alc., eth., chlf., bz.— $[\alpha]_D$ in 1% alc. sol. = +57.60°.
620	58	Meconidine, $\text{C}_{12}\text{H}_{20}\text{O}_2\text{N}$.—[Small quantity in opium.] “Brownish yel.” amorph. mass fr. eth. V.s. alc., eth., chlf., bz.; d.s. ammon.; e.s. NaOH. Eth. extracts fr. ammon., but not fr. NaOH sol. Reacts strongly alk. Acids dec. w. red color.
—	59	Triacetonamine (hydrated).—Cf. No. 2.554. (Odor ammon. & camphorous.)
621	59–60d.	β -Ethylhydroxylamine, Et.NH(OH) .—⑩ Reduces Fehling's sol.—V.s. aq., alc.; d.s. eth., lgr., bz. V. vol.—Oxalate, $\text{B}_2\text{C}_2\text{H}_4\text{O}_4$, lft., m.p. 95°.—⑪ Convert into phenylisocyanate addition product of m.p. 98°. (Cf. Ber. 26, 2381.)
622	58–60	Salicyltropeine, $\text{C}_{15}\text{H}_{14}\text{O}_2\text{N}$.—Lust. silky ndl. D.s. c. aq.; v.s. alk. (“Strong base.” Not a mydiatic.)

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(ORDER II, SUBORDER I.)

No.	Melting-point (C. ^o).	BASIC COMPOUNDS.—Colorless and Solid.
623	60	m-Aminobenzylamine, NH ₂ .C ₆ H ₄ .CH ₂ .NH ₂ . — ⊖ Prim. amine.* — Ndl. — B ₂ H ₂ PtCl ₆ . §
624	60	Quinicine, Quinotoxine, C ₂₀ H ₂₄ O ₂ N ₂ . — [Fr. Cinchona bark.] — Yellowish oil solidifying in desicator. D.s. aq.; e.s. alc., eth., chlf. [α] _D in chlf. ($p = 2$) = +44.1°. "Reacts alk." Taste, bitter. Alc. sol. w. Cl-aq. + amonon. gives greenish coloration.
625	60-2	Apostropine, Tropine Atropate, C ₁₇ H ₂₁ O ₄ N. — [Dehydration product of atropine. Not a mydriatic.] — Pr. fr. eth.; little s. aq.; more s. lgr.; e.s. alc., eth., chlf., bz. — B ₂ HAuCl ₄ , ndl., m.p. 110-2° fr. h. aq.
626	61.5	1,10-Diaminodecane, C ₁₀ H ₂₀ .(NH ₂) ₂ . — B.p. 140° (12 mm.). — ⊖ Prim. amine.*
627	61.2	2,3-Toluylenediamine, Me.C ₆ H ₄ (NH) ₂ . — B.p. 255°. — ⊖ Prim. amine.* — Cryst.
628	61	2,7-Dimethylquinoline, Me ₂ C ₆ H ₃ N. — B.p. 264-5°. — ⊖ Tert. amine.* — B ₂ H ₂ PtCl ₆ , § yel. ndl. fr. h. aq.
629	61-2	Py-4-Phenylquinoline, Ph.C ₆ H ₄ N. — ⊖ Tert. amine§. — Ndl. fr. eth. E.s. neut. solvents except aq. — The v. dil. sol. of B.HCl fluor. blue-violet! — Picrate, † m.p. 224°.
630	62	Glycocolanilide, NH ₂ .CH ₂ .CO.NH.Ph.1½H ₂ O. — ⊖ Prim. amine.* T. 2.12 prob. gives carblyamine odor. — Ndl. fr. aq. E.s. aq., alc.; d.s. eth., bz., lgr. — "Strong base." M.p. on remelting, 55°. — [D.R.P. 59,874.]
630-1	62	C-Methylpiperazine, [NH.CHMe.CH ₂ .NH.CH ₂ .CH ₃]. — B.p. 151°; 155-0.5° (th.i.). Thin deliq. lft. E.s. aq., alc., chlf., bz. — ⊖ Sec. amine.‡ — B.Pk, † lust. yel. lft., v.d.s. h. aq., or alc., d. 276-8°. — B ₂ H ₂ PtCl ₆ , § lust. red pr., d.w.m. — Dinitrosamine, ** lft. fr. aq., m.p. 71°.
631	62	m-Ethylaminophenol, EtNH.C ₆ H ₄ (OH). — Feathery cryst. fr. bz. + lgr.
632	62-3	Naphthalanmorpholine, C ₁₂ H ₁₅ ON. — B.p. 312° (754 mm.). — ⊖ Sec. amine.* — Pr. d.s. aq.; e.s. organic solvents. — Aq. sol. strongly alk. & gives ppt. w. most alkaloid reagents (T. 2.3). — B.HCl, m.p. 275°, e.s. aq. — B ₂ H ₂ PtCl ₆ , § pr. dec. abt. 225°. — Picrate, † d. abt. 250°. — Nitrosamine, ** ndl. fr. alc., m.p. 161°! — [Hypnotic, — D.R.P. 105,498.]
633	62	Tetraethyldiaminotriphenylmethane, Ph.CH(C ₆ H ₄ .NEt ₂) ₂ . — ⊖ Tert. amine.* — Lust. ndl. fr. alc. V.d.s. h. aq.; e.s. alc., eth., bz.
634	63 u.c.	m-Phenylenediamine, C ₆ H ₄ (NH) ₂ . — B.p. 282-4° (th.i.). — Prim. amine.* — [V. unstable on exposure to air & light, becoming dark colored. When separating in oily state fr. sol., v. difficult to solidify.] — ⊖ (1) Sol. in dil. H ₂ SO ₄ colors yel. on adding NaNO ₂ sol., & then gives ppt. of Bismarck brown on saturating w. salt. — (2) Br-ag. in dil. sol. gives violet ppt. which soon becomes cryst. (Behrens). — Clear plates on slow evapn. of bz. sol. E.s. aq., alc., eth. — ⊖ To 0.1 g. add 5 drops acetic anhydride & heat to boiling. Dis. product in 4 cc. h. aq. Cool & add Br-ag., a few drops at a time until sol. remains yel. Filter. Wash the heavy white ppt. w. 3 cc. aq. Recryst., once fr. 3 cc. & once fr. 2 cc. h. alc. Dry on porous tile 30 min. at room temp. Take m.p. in bath w. temp. rising v. rapidly (40° per min. up 240°). The product C ₆ H ₅ Br.(NH.CO.Me) ₂ is white & granular, m.p. 256-7° d. u.c. (264-5° c.).
635	63	† Tropine, Tropanol, C ₉ H ₁₁ ON. — B.p. 229°. — ⊖ N. Eq., 141. (k.10 ⁴ = 2.7). Tert. amine.* — Hygroscopic ndl. V.s. aq., alc., separating out oily on evapn. Opt. i. — Ignition w. soda-lime gives methylamine (No. 2,1059), etc. — B.Pk, † yel. ppt., tbl. fr. h. aq., d.w.m. abt. 275°. — B ₂ H ₂ PtCl ₆ , § or.-yel. monoclin. cryst., m.p. 198-200°. — B.HAuCl ₄ , yel. tbl. (on evapn.), m.p. 210-2° d.
636	64.5	1'-Amino-1,2,4,5-tetramethylbenzene, NH ₂ .CH ₂ .C ₆ H ₃ .Me ₂ . — ⊖ Prim. amine.* — Ndl. fr. dil. alc.; i.c. aq. — B.HCl, cryst., m.p. 240-2° d. — B ₂ H ₂ PtCl ₆ , red-yel. ppt., d. 208-9°.
637	64	3-Aminopyridine, NH ₂ .C ₆ H ₄ N. — B.p. 250-2°. — ⊖ Prim. amine.* — Lft. fr. bz. + lgr. V.s. aq., alc., eth.; i. lgr. — B ₂ HCl, glassy tbl., m.p. 175° d. — B ₂ H ₂ PtCl ₆ , § yel.-red cryst., d. 225°. — Acetyl deriv., ¶ m.p. 131°.
638	64	2,5-Toluylenediamine, (NH) ₂ . ^{2,5} C ₆ H ₃ (Me) ¹ . — B.p. 273-4°. — ⊖ Sol. + little o-toluidine gives intense green color w. FeCl ₃ (Ber. 12, 2237). Prim. amine.* — Lit. fr. bz. E.s. aq., alc., eth., h. bz. — Heated w. dil. H ₂ SO ₄ & MnO ₂ gives pungent odor of toluquinone.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
639	65-6; 61	p-Tolylhydrazine, $\text{Me.C}_6\text{H}_4.\text{NH.NH}_2$. — B.p. 240-4° sl. d. — \textcircled{P} Should reduce Fehling's sol. w. evolution of N. — Lft. E.s. alc., eth., bz. — Acetyl deriv., lft., m.p. 121°.
640	64	8-Quinolinehydrazine, $\text{NH}_2.\text{NH.C}_6\text{H}_4\text{N}$. — \textcircled{P} Prob. reduces Fehling's sol. w. evolution of N. — Fine "sl. yellowish" ndl. — B_2HCl , thick lust. yel. pr. Sol. of $\text{B}_2\text{HCl} + \text{KCNO}$ sol. gives ppt. of the semicarbazide, $\text{C}_6\text{H}_4\text{N.NH.CO.NH}_2$, lft. fr. alc., m.p. 235° d.
641	64-5	4-Methyl-2-phenylquinoline, Flavoline, $\text{Me.Ph.C}_6\text{H}_4\text{N}$. — B.p. 373-5°. — \textcircled{P} Tert. amine* of quinoline-like odor. — Thick 4-cornered tbl. fr. lgr. — $\text{B}_2\text{HCl.2H}_2\text{O}$, colorless pr. e.s. aq.
642	65	p-Aminobenzyl Alc., $\text{NH}_2.\text{C}_6\text{H}_4.\text{CH}_2\text{OH}$. — \textcircled{P} Prim. amine.* Lft. fr. bz.
643	65-6	2,4-Diamino-m-xylene, $(\text{NH}_2)_2.\text{C}_6\text{H}_4(\text{Me})_2$. — \textcircled{P} Prim. amine.* — Ndl. fr. lgr. E.s. aq., alc. bz. — Dibenzoyl deriv., m.p. 232°, 227° c., ndl. fr. alc.
644	63; 65	Ethylenediphenyldiamine, $\text{Ph.NH}(\text{CH}_2)_2.\text{NH.Ph}$. — \textcircled{P} Sec. amine.* — Alc. sol. w. alc. sol. HgCl_2 gives ppt. of B_2HgCl_4 , rhomb. plates, m.p. 129° c. — Cryst. E.s. alc., eth. — \textcircled{D} Dis. 1 g. in 5 g. HCl + 30 cc. aq. Add slowly sol. of 2 mol. NaNO_2 . Cryst. pptd. dinitroso deriv. fr. gl. ac. ac. Lft., m.p. 157°, i. c. aq.
645	65	3,4-Dimethylquinoline, $\text{Me}_2.\text{C}_6\text{H}_4$. — B.p. 291° (th.i.). — Cryst. mass. — $\text{B}_2\text{Pk.}$ † yel. ndl. fr. abs. alc., m.p. 205°. — $\text{B}_2\text{H}_4\text{PtCl}_6.2\text{H}_2\text{O}$, § cryst. fr. dil. HCl, d. 234-40°. — B_2HAuCl_4 , cryst. fr. dil. HCl, m.p. 177°.
646	66 u.c.	† s-Pseudocumidine, $\text{Me}_2.\text{C}_6\text{H}_4.\text{NH}_2$. — B.p. 234-5°. — \textcircled{P} Prim. amine.* — Ndl. fr. dil. alc. Odor mild-aromatic, somewhat like aniline. Taste of c. sat. aq. sol. burning (Pungency, No. 3, T. 2.29). — \textcircled{D} Heat together to boiling 0.04 g. amine & 6 drops acetic anhydride. Dis. product by boiling w. 10 cc. aq. Cool, & shake vigorously. Filter. Wash w. 2 cc. c. aq. Recryst. fr. 4-5 cc. h. aq. Dry 15 min. on porous tile at 100°. The product, acet pseudocumidine, $\text{Me}_2.\text{C}_6\text{H}_4.\text{NH.CO.CH}_3$, is obtained in colorless ndl., m.p. 162.2-3.2° u.c. (165.0-166.0° c.).
647	66-7	p-Aminodiphenylamine, $\text{NH}_2.\text{C}_6\text{H}_4.\text{NH.Ph}$. — \textcircled{P} FeCl ₃ added to sol. gives red color, soon changing to green, or in more conc. sol., green ppt. s. in conc. H_2SO_4 w. carmine red color. Prim. amine.* — Lft. which melt at 75° after melting & resolidifying, or after cryst. fr. lgr. — D.s. aq.; e.s. eth., abs. alc.
648	66-7	3,5-Diaminoxylylene(1,2), $(\text{NH}_2)_2.\text{C}_6\text{H}_4.\text{Me}$. — \textcircled{P} Prim. amine.* — NaNO_2 gives dark brown ppt. in ac. ac. sol. Sol. in warm dil. H_2SO_4 becomes cherry red. — Ndl. fr. alc. V.s. aq., alc., eth., bz. Diacetyl deriv., ndl. d.s. aq., m.p. 240-1°.
649	66.5	1,8-Diaminonaphthalene, $(\text{NH}_2)_2.\text{C}_{10}\text{H}_8$. — \textcircled{P} Prim. amine.* — Sol. gives chestnut brown ppt. w. FeCl_3 . — Cryst. fr. dil. alc. Sbl. D.s. aq.; v.s. alc., eth. — B_2HCl , lft., m.p. abt. 280°.
650	66	Hydrohydrastinine, 2-Methyl-6,7-methylenedioxytetrahydroisoquinoline, $\text{Me.-C}_6\text{H}_3:\text{O}_2:\text{CH}_2$. — \textcircled{P} Tert. amine.* — Cryst. v.s. alc., eth., bz. — B_2HCl , cryst. powd., m.p. 273-4°. — $\text{B}_2\text{H}_4\text{PtCl}_6$, § yel. lft., m.p. 216°; 207-8°. — B_2MeI , †† ndl. fr. aq., m.p. 227-8°.
651	67.5-8 u.c.	† Lupinine, $\text{C}_{10}\text{H}_{15}\text{ON}$. — B.p. 255-7°. — Rhomb. pr. fr. pet.-eth. More s. c. than h. aq.; e.s. alc., eth., chlf., bz. Levorotatory. Taste, intensely bitter & lasting. T. 2.2-(a), (b), (f), (g) give no colorations. — B_2HAuCl_4 , ndl. d.s. aq.; v.s. alc. — [Fr. seeds of <i>Lupinus Luteus</i> .]
652	68	Emetine, $\text{C}_{20}\text{H}_{24}\text{O}_2\text{N}_2$. — [Powerful emetic alkaloid fr. root of <i>Cephaelis Ipecacuanha</i> .] — \textcircled{P} Froehde's reagent, T. 2.2-(f), gives dirty green color, changing to light grass green w. a drop conc. HCl or granule of NaCl! — Amorph. crusts fr. lgr.; lft. fr. alc., eth. Usually amorph. S. in 1000 pt. c. aq.; e.s. alc., eth., bz., chlf., pet.-eth. Can be shaken out by eth. fr. alk., but not acid, solutions. Unlike cephaline, not s. x.s. NaOH sol. — Taste, harsh & bitter (poisonous).
653	68	Benzyl-β-naphthylamine, $\text{Ph.CH}_2.\text{NH.C}_10\text{H}_7$. — \textcircled{P} Sec. amine.* — Pr. fr. eth. S. bz. lgr. — Dil. H_2SO_4 & NaNO_2 gives nitroso deriv., ** ndl. fr. alc., m.p. 111-2°.

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. † = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
654	68	3-Methylisoquinoline , $\text{Me.C}_6\text{H}_4\text{N}$. — B.p. 240°. — \textcircled{P} Tert. amine.* — Picrate, † m.p. 197–8°. — $\text{B}_2\text{H}_2\text{PtCl}_6\text{H}_2\text{O}$, § or.-yel. ndl. fr. aq., m.p. abt. 195° w. frothing.
655	68	2,3-Dimethylquinoline , $\text{Me}_2\text{C}_6\text{H}_4\text{N}$. — B.p. 262°. — \textcircled{P} Tert. amine.* — Tbl. D.s. aq.; v.s. alc.; e.s. eth. — Picrate, † m.p. 225°. — $\text{B}_2\text{H}_2\text{PtCl}_6\text{H}_2\text{O}$, § or.-yel. ndl., blackens at 230°.
—	68	3,3'-Bipyridyl . — Cf. No. 2.1444. (Usually an oil.)
657	69–70	Nortropinone , $\text{C}_7\text{H}_11\text{ON}$. — \textcircled{P} Sec. amine.* — "Strong base." Thin deliq. cryst. fr. bz. + lgr. V.s. aq., alc., bz.; v.d.s. lgr.; e.s. eth. — B.Pk , † yel. pr., v.d.s. c. alc., m.p. 159–60°.
658	69–70	Dimethylamino-benzhydrol , $\text{NMe}_2\text{C}_6\text{H}_4\text{CH}(\text{OH})\text{Ph}$. — \textcircled{P} Tert. amine.* — Ndl. fr. dil. alc. I. aq.; e.s. alc. — Heated w. ZnCl_2 + dimethylaniline gives leucomalachite green (No. 2.727)!
—	69.5–70	Pyrazole . — Cf. No. 2.1650–1.
659	70	5-Amino-1,2,3,4-tetramethylbenzene , Prehnidine , $\text{NH}_2\text{C}_6\text{H}_4\text{Me}_4$. — B.p. 259–60°. — \textcircled{P} Prim. amine.* — S. h. aq.; e.s. alc., eth., lgr. — Acetyl deriv., ¶ ndl. e.s. alc., m.p. 169.5°; 172°.
660	70	aa-Benzoylphenylhydrazine , Ph.CO.NPh.NH_2 . — \textcircled{P} Reduces Fehling's sol. on boiling. — Ndl. fr. aq. D.s. c. aq.; v.s. alc., eth., chlf. — B.HCl , r.d.s. aq., m.p. 202°. — B.Pk , † m.p. 122°.
661	70	Dibenzylaniline , $(\text{Ph.CH}_2)_2\text{N.Ph}$. — \textcircled{P} Sec. amine.* — Ndl. d.s. c. alc.; e.s. eth., bz. — B.Pk , † yel. ndl., m.p. 131–2° d. — \textcircled{P} Prepare 2,4-dinitrophenyldibenzylamine by treating sol. of 1 g. amine in 10 cc. gl. ac. ac. at 30° w. 0.8 cc. HNO_3 (sp. gr. 1.52). Product, crystd. fr. gl. ac. ac., melts at 106° (Ber. 32, 913).
662	70	Ethylenediyldiphenylamine , $(\text{PhEtN})\text{CH}_2\text{CH}_2\text{(NEtPh)}$. — \textcircled{P} Tert. amine.* — Cryst. — $\text{B.H}_2\text{PtCl}_6$, § ndl.
663	70	Cinnamyltropeine , $\text{C}_{11}\text{H}_{19}\text{O}_2\text{N}$. — \textcircled{P} Tert. amine.* — Lft. V.d.s. aq.; e.s. alc., chlf. — V. toxic, but not mydriatic.
—	74 u.c.	8-(o)-Hydroxyquinoline . — Cf. No. 2.36. (Sol. in alk. yellow.)
665	74 u.c.	† 8-Hydroxyquinaldine , $\text{HO.C}_6\text{H}_4\text{C}_6\text{H}_3\text{N}(\text{Me})$. — B.p. 266–7°. — Tert. amine* — \textcircled{P} Unlike 8-hydroxyquinoline (above), sol. in dil. NaOH is colorless instead of yellow, and FeCl_3 test described under No. 2.36 gives faint blue-green color destroyed by heat. Well-developed orthorhombic pr. fr. dil. alc. Odor, & taste of sat. aq. sol., aromatic-phenolic, like guaiacol but less intense. V.d.s. c. aq.; e.s. eth., bz., h. alc. Vol. w.st. Begins to subl. abt. 100°. — \textcircled{D} To 0.05 g. dissd. in 3 cc. alc., add 3 cc. c. sat. alc. sol. of picric ac. Warm. Shake vigorously & allow to stand for some time. Filter off abundant ppt. of yel. ndl. Wash w. 3 cc. c. alc. Cryst. fr. 3–4 cc. 50% alc. Dry 15 min. at 100°. The resulting picrate, B.Pk (analyzed), is obtained in Y ndl. softening abt. 215° & melting w. decn. at 219–20° u.c. (224.8–5.8° c.).
667	74	4-Phenyl-1,2,3,4-tetrahydroquinoline , $\text{Ph.C}_6\text{H}_4\text{N}$. — \textcircled{P} Sec. amine.* — Lust. lft. fr. dil. alc. — Picrate, † m.p. abt. 183° — Nitroso deriv., ** yel. ndl. fr. alc., m.p. 72°.
668	74–5	Lactyltropeine , $\text{C}_{11}\text{H}_{19}\text{O}_2\text{N}$. — \textcircled{P} Tert. amine.* — E.s. cryst.
669	75	5-Amino-1,2,3-trimethylbenzene , $\text{NH}_2\text{C}_6\text{H}_3\text{Me}_2$. — B.p. 240°. — \textcircled{P} Prim. amine.* — Ndl. i. c. aq. — Acetyl deriv., ¶ m.p. 163–4°.
670	75	2,3-Diaminoxylene(1,4) , $(\text{NH}_2)_2\text{C}_6\text{H}_2\text{Me}_2$. — \textcircled{P} Prim. amine.* Gives cherry red color w. FeCl_3 . — Sbl. in ndl. E.s. aq., alc., bz. — NaNO_2 added to ac. ac. sol. gives yel. color followed by white ppt.
671	75	Benzoinhydrazine , $\text{Ph.CH(OH).CPh:N.NH}_2$. — \textcircled{P} Prob. reduces alc. AgNO_3 sol. — Pr. fr. alc. I. aq., alk.; v.s. h. alc. — Decd. at once by acids to benzoin (Vol. I) & hydrazine.
672	75–6	Ethylenedi-o-tolyldiamine , $\text{C}_6\text{H}_5\text{NH.CH}_2\text{CH}_2\text{NH.C}_6\text{H}_5$. — \textcircled{P} Sec. amine.* Ignition w. Zn dust gives indole (No. 2.1546)! — Tbl. fr. lgr. Alm. i. aq.; e.s. gl. ac. ac.; s. in 10 pt. eth. or c. alc.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
673	75	Ditamine, $C_{16}H_{19}O_2N$. — [Fr. Apocine Echites scholaris L., a Philippine febrifuge.] — Amorph. powd. E.s. alc., eth., chlf., bz.; v.s. dil. ac., reptd. by NH_3 in flocks.
674	abt. 76	8-Hydroxy-1-ethyltetrahydroquinoline, $HO.C_8H_7NEt$. — \textcircled{P} Is colored violet & then dark muddy brown by little $FeCl_3$, and then on addition of little conc. H_2SO_4 , purple red. Colored blood-red by fuming HNO_2 . Aq. sol. w. $FeSO_4$ gives ppt. of black-green flocks. — Lft. d.s. h. aq.; e.s. alc., eth., bz.; r.d.s. lgr. — Boiled for 6 hr. w. acetic anhydride + anhydrous Na acetate gives acetyl deriv., pr fr. eth., i. aq., m.p. $63\text{--}4^{\circ}$. — $\bar{B}.HCl$ (the antipyretic Kairin A), trimet. pr., e.s. aq. — [Cf. also Dragendorff, p. 313.]
675	77	2-Amino-4'-methyl diphenylamine, $NH_2.C_6H_4.NH.C_6H_4.Me$. — \textcircled{P} Colored purple red by $FeCl_3$. — Prim. amine.* — Lft. fr. aq., or lgr. E.s. alc., eth., chlf., bz.
676	77-8	4,5-Diaminoxylenne(1,3), $(NH_2)_2.C_6H_4.Me_2$. — \textcircled{P} Colored red by $FeCl_3$. — $K_2Cr_2O_7$ gives brown ppt. in cold, & red sol. w. pungent quinone odor on heating. — Cryst. s. h. aq.; e.s. alc., eth.; d.s. lgr., bz.
677	77	1,2,3,4-Tetrahydronaphthalenediamine(1,5), $C_{10}H_{10}(NH_2)_2$. — B.p. 264° (60 mm.). — \textcircled{P} $FeCl_3$ colors <i>only hot</i> sol. red-brown. Odor basic piperidine-like. — Glassy pr. fr. eth. V.s. alc., eth.; d.s. lgr. — $\bar{B}_2H_2PtCl_6\ddagger$ (dried over H_2SO_4), ocher-yel. cryst. ppt., alm. i. aq.; e.s. mineral a.c.
678	77.5	1,6-Diaminonaphthalene, $(NH_2)_2.C_10H_8$. — \textcircled{P} Aq. sol. fluor. blue! Prim. amine.* — Ndl. fr. aq. V.d.s. c. aq., eth.; e.s. alc., bz., h. aq. — Diacetyl deriv., ¶ m.p. abt. 257° .
679	77-80	Benzylamidine, $Ph.C(:NH).NH_2$. — \textcircled{P} T. 2.7 gives NH_3 ! — Deliq. alk. cryst. Mod. s. aq.; v.s. alc.; d.s. eth. — Heated above m.p. dec. to NH_3 , benzonitrile (No. 2.2781) & kyaphenine. — $\bar{B}.Pk,\ddagger$ yel. ndl., m.p. 228° . — $\bar{B}_2H_2PtCl_6.2H_2O,\ddagger$ ndl., d.s. aq., m.p. $209\text{--}10^{\circ}$ d.
680	77	4-(γ)-Phenylpyridine, $Ph.C_6H_4N$. — B.p. $274\text{--}5^{\circ}$. — \textcircled{P} Tert. amine.* — Lust. lft. fr. h. aq. D.s. h. aq. Boiled w. $KMnO_4$ sol. gives pyridine-4-carbonic ac. — Picrate, \ddagger fine yel. ndl., m.p. $195\text{--}6^{\circ}$, v.d.s. h. aq., alc. — $\bar{B}_2H_2PtCl_6,\ddagger$ light yel. mic. ndl., alm. i. aq.
681	78	Dimyricylamine, $(C_{10}H_{11})_2NH$. — \textcircled{P} Sec. amine.* — Cryst.
682	78-9	o-Aminodiphenylamine, $NH_2.C_6H_4.NHPh$. — \textcircled{P} Prim. amine.* — Ndl. fr. aq. E.s. chlf., bz.; less s. lgr. [Used in manufact. of rosinduline colors.]
683	78	m-Diethylamino-phenol, $Et_2N.C_6H_4.OH$. — B.p. $276\text{--}80^{\circ}$. — [An intermediate in dyestuff industry.]
684	78-8.5	Phenanthroline, $C_{10}H_8N_2$. — B.p. much above 360° . — \textcircled{P} Bitert. amine.* — Cryst. w. $2H_2O$ (m.p. 65.5°) in soft ndl. Alm. i. c. aq.; s. h. aq.; alm. i. eth. bz., lgr.; e.s. dil. acids; v.s. alc. — $\bar{B}.Pk,\ddagger$ mic. yel. pr., v.d.s. alc., m.p. $238\text{--}40^{\circ}$. — \bar{B}_2Br_2 , mic. cryst. ppt., m.p. 149° , forms when Br_2 (as Br -aq) is added to aq. sol. of $\bar{B}.2HCl$.
685	78	Isonicotine, $C_{10}H_{14}N_2$. — B.p. much above 260° d. — \textcircled{P} Tert. amine.* Hydroscopic ndl. Acts caustic like KOH on skin. — V.s. aq.; e.s. eth., lgr., bz. — $\bar{B}_2H_2PtCl_6.H_2O,\ddagger$ lust. or.-yel. lft.
686	78-80d.	2,4-Diaminophenol, $(NH_2)_2.C_6H_4(OH)$. — \textcircled{P} Sol. in ammon. becomes deep blue on exposure to air! — Ppt. in lft. fr. aq. sol. of HCl by Na_2SO_4 . Becomes quickly brown-black in air. — $\bar{B}.Pk,\ddagger$ lemon-yel. ndl. s. in 33 pt. c. aq.; m.p. abt. 120° . — [Dyestuff for feathers & an intermediate in manufact. of sulphur colors.]
687	79	Cusparidine, $C_{17}H_{21}O_4N$. — [Tert. amine* fr. Angustura bark.] — \textcircled{P} Gives red sol. in conc. H_2SO_4 . — Mic. ndl. fr. pet.-eth. E.s. alc., eth., chlf. — $\bar{B}_2H_2PtCl_6,\ddagger$ m.p. 182° . — $\bar{B}.HAuCl_4,\parallel$ m.p. 167° . — $\bar{B}.MeI,\ddagger\ddagger$ light yel. powd., m.p. 149° .
688	80	d,l-Cocaine, $C_{17}H_{21}O_4N$. — \textcircled{P} Action on tongue like l-cocaine as described under (1), No. 2.741. — 6-Sided lft. fr. lgr. I. aq.; v.s. alc., eth. — $\bar{B}.HCl$, 6-sided tbl. fr. alc.

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. \ddagger = T. 2.23. $\ddot{\ddagger}$ = T. 2.14. \parallel = T. 2.13. $\ddagger\ddagger$ = T. 2.1. $\ddot{\ddot{\ddagger}}$ = T. 2.36. $\ddot{\ddot{\ddagger\ddagger}}$ = T. 2.37. $\ddot{\ddot{\ddot{\ddagger}}}$ = T. 2.38.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
690	82	o-Aminobenzyl Alc., $\text{NH}_2\text{C}_6\text{H}_4\text{CH}_2(\text{OH})$. — B.p. 270–80° sl. d. — \textcircled{P} Prim. amine.* — Ndl. fr. bz. Mod. s. aq., eth.; e.s. alc., bz., chlf. — B.HCl , tbl. fr. eth.-alc., m.p. 108°. — Picrate, \ddagger fine yel. ndl., m.p. 110°.
691	82	β -Naphthoquinaldine, $\text{C}_{11}\text{H}_{11}\text{N}$. — B.p. above 300°. — Tert. amine.* — Ndl. fr. dil. alc. D.s. aq.; e.s. alc., eth. — B.Pk , \ddagger mic. ndl. i.c. aq.; e.s. gl. ac. ac.; m.p. 220–1° d. — $\text{B}_2\text{H}_2\text{PtCl}_4\cdot 2\text{H}_2\text{O}$, \ddagger yel. ndl. d.s. aq.
692	82(u.c.) sl.d.	\dagger Taxine, $\text{C}_{17}\text{H}_{24}\text{O}_4\text{N}(\text{N})?$. — [Specimen fr. Th. Schuchardt. Alkaloid fr. yew tree, <i>Taccus baccata</i> L.] — \textcircled{P} Evap. eth. sol. on crucible cover. Place cover on crucible containing conc. HNO_3 . Residue becomes green-blue after few min. Then place cover on crucible containing HCl. The rose color which appears is destroyed by NH_3 . (Vreven, Ann. de Pharm., Louvain, 1896.) — Tert. amine.* — Amorph. I. aq., pet.-eth.; s. alc., eth., bz., chlf. Odorless. Taste, v. bitter (poisonous). Alk. ppt. fr. sol. in dil. acids as voluminous white mass. — Gives muddy reddish violet coloration w. H_2SO_4 , <i>Buckingham's reag.</i> , & <i>Froehde's reag.</i> in <i>T. 2.2-(a), (d), (f)</i> . — B.Mel , \ddagger (by mixing bz. sol. w. MeI), white amorph. powd., m.p. abt. 121°.
693	82–3	i-Scopolamine, (d,l)-Tropate of i-Scopoline, $\text{C}_{17}\text{H}_{21}\text{O}_4\text{N}$. — Crystg. w. $1\text{H}_2\text{O}$, m.p. 56–7°. S. in 38 pt. aq. at 18°. — Picrate, \ddagger m.p. 192–4°.
694	83	6-Methylisoquinoline, $\text{Me.C}_6\text{H}_4\text{N}$. — B.p. 263–4°. Tert. amine.* — Picrate, \ddagger m.p. 212°.
695	83–4	\dagger Oxyparteine, $\text{C}_{17}\text{H}_{24}\text{ON}_2$. — Tert. amine.* Ndl. e.s. aq., alc., eth., chlf. Unchanged after heating w. conc. H_2SO_4 to 250° on warming w. fuming HNO_3 . — Picrate, \ddagger m.p. 176–8°. — $\text{B}_2\text{H}_2\text{PtCl}_4\cdot 4\text{H}_2\text{O}$, \ddagger lust. ruby-red cryst., m.p. 209° w. efferv., & $\text{B.H}_2\text{PtCl}_4\cdot 2\text{H}_2\text{O}$, lust. ndl., m.p. 221–3° d.
696	83	Dimethyl-tetramethyldiamino-diphenylmethane, $\text{Me}_2\text{C}(\text{C}_6\text{H}_4\text{NMe}_2)_2$. — \textcircled{P} Iodine gives emerald green color. — Tert. amine.* — Silky ndl. fr. dil. alc. I. boiling aq.; d.s. c. alc., e.s. h.; e.s. eth.
697	84	5,6,7,8-Tetrahydro-1,4-diaminonaphthalene, $\text{C}_{10}\text{H}_{16}(\text{NH}_2)_2$. — \textcircled{P} Reduces ammon. AgNO_3 sol. Prim. amine.* — B.p. 220° (81 mm.). — E.s. alc., eth., h. aq. — Diacetyl deriv., \ddagger silky ndl.; d.s. eth., lgr.; v.s. alc.; m.p. 245°.
698	84; 83; 86	2-(α)-Phenylquinoline, $\text{Ph.C}_6\text{H}_4\text{N}$. — Dist. undec. a. 300°. — \textcircled{P} Tert. amine.* — Silky ndl. fr. dil. alc. D.s. aq.; e.s. eth., h. alc. — B.Pk , \ddagger yel. lt. fr. alc. m.p. 187–8°.
699	85	1,3,4-Xylylhydrazine, $\text{Me}_2\text{C}_6\text{H}_3\text{NH.NH}_2$. — \textcircled{P} Prob. reduces AgNO_3 sol. — Ndl. fr. eth. V.s. eth. — Sol. in bz. becomes deep green on long standing.
700	85.5	β -Bisethylideneaniline, $\text{Ph.N:CH.CH}_2\text{CHMe.NHPh}$. — \textcircled{P} Sec. amine.* Silky ndl. fr. alc. S. c. alc.; e.s. eth., chlf., bz. — Dinitroso deriv. ** (Ber. 29, 2977) lust. cryst. powd. fr. eth. + bz., m.p. 102°.
701	85	Ethyldieneimine, $[\text{NH.CHMe.NH.CHMe.NH.CHMe}]$. — \textcircled{P} B.p. 123–4°! Odor acetamide-like. — Lust. cryst. s. aq., alc., bz.
702	85 u.c.	\dagger m-Dimethylamino-phenol, $\text{Me}_2\text{N.C}_6\text{H}_4\text{OH}$. — B.p. 265–8°. — Ndl. fr. lgr. E.s. alc., eth., bz., 10% aq. NaOH sol.
703	85.5 (77.5)	Anhalonine, $\text{C}_{12}\text{H}_{14}\text{O}_2\text{N}$. — [Tetanic poison fr. Mexican "mescal buttons," <i>Anhalonium Lewinii</i> .] — \textcircled{P} Sec. amine* containing one methoxyl group. — Ndl. fr. pet.-eth. E.s. alc., eth., chlf., lgr. — Levorotatory. — $\text{B}_2\text{H}_2\text{PtCl}_4$, \ddagger yel. mic. pr. d.s. aq. — Nitroso deriv., ** $\text{C}_{12}\text{H}_{14}\text{O}_2\text{N}(\text{NO})$, cryst., sinter & melt at 59°.
704	86	p-Aminobenzonitrile, $\text{NH}_2\text{C}_6\text{H}_4\text{CN}$. — \textcircled{P} Prim. amine.* — Dist. w. sl. dec. — E.s. alc., eth., h. aq. — $\text{B}_2\text{H}_2\text{PtCl}_4$, \ddagger d.s. ndl.
705	86–7	o-Aminobenzylaniline, $\text{NH}_2\text{C}_6\text{H}_4\text{CH}_2\text{Ph}$. — \textcircled{P} Prim. amine.* — Glassy ndl. fr. alc. D.s. lgr.; e.s. alc.
706	86	Tetramethylpyrazine, Tetramethyl-1,4-diazatriene, Dimethylketine, $[\text{N:CMc-CMe:N.CMe:CMc}]$. — B.p. 189°. — Tert. amine.* — Ndl. fr. aq. w. $3\text{H}_2\text{O}$, m.p. 74–7°, changing to anhydrous octahedra in desiccator. Vol. w. st. E.s. alc., eth. — ["Reacts neut." — $\text{B.HCl.2H}_2\text{O}$, e.s. aq., alc. — NaOH sol. ppts. a hydrate fr. sol. of salts.] — B.2Pk , \ddagger ndl. d.s. c. aq., alc., m.p. 191–2°. — $\text{B.Mel.2H}_2\text{O}$, \ddagger light yel. ndl. fr. aq., losing aq. over H_2SO_4 & then melting at 216° w. dec. to components.

No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
707	88.5	3,4-Toluylenediamine, $\text{Me.C}_6\text{H}_4(\text{NH}_2)_2$. — B.p. 265°. — Prim. amine.* — \textcircled{P} Aq. sol. soon blackens in air. — Scales s.c. aq.
708	88	4,4'-Diaminodiphenylmethane, $\text{CH}_3(\text{C}_6\text{H}_4\text{NH}_2)_2$. — \textcircled{P} Prim. amine.* — Pearly lft. E.s. alc., bz. — Sulphate, lft. v.d.s. alc. — Diacetyl deriv., \ddagger lust. tbl. fr. alc., m.p. 228°.
709	88-90d.	o-Methylamino-phenol, $\text{Me.NH.C}_6\text{H}_4(\text{OH})$. — \textcircled{P} Sec. amine.* — Lft. Sol. in dil. HCl gives deep red-brown color w. FeCl_3 . — Nitrosamine, ** $\text{Me}(\text{NO})\text{N-C}_6\text{H}_4\text{OH}$, ndl. fr. alc., e.s. h. aq., d. 121°.
710	88-9	Imidazole, Glyoxaline, $[\text{NH.CH: N.CH: CH}]_2$. — B.p. 255°. — $k \cdot 10^7 = 1.2$. — \textcircled{P} Not acted on by chromic ac., acetyl chloride or acetic anhydride. — Thick pr. "Reacts strongly alk." E.s. aq., alc., eth. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or-red pr., e.s. h. aq.; contains cryst. aq. lost at 100°.
711	87-8	α -Cocaine, $\text{NMe.C}_7\text{H}_{10}(\text{Ph.CO}_2)(\text{CO}_2\text{Me})$. — Tert. amine.* Glassy pr. fr. lgr. Alm. i. c. aq.; v.s. alc., eth., chlf., bz. — B.Pk , \ddagger yel. pr. fr. wood alc. (v.d.s.), m.p. 195°.
712	89	3,4-Diaminoxylene(1,2), $(\text{NH}_2)_2\text{C}_6\text{H}_2\text{Me}_2$. — Prim. amine.* — \textcircled{P} Sol. gives red color w. FeCl_3 . — Tbl. e.s. alc., eth., bz.; d.s. lgr.
713	89	4,6'-Diamino-3,3'-dimethyldiphenylmethane, $[\text{Me.C}_6\text{H}_4(\text{NH}_2)]_2\text{CH}_2$. — \textcircled{P} Prim. amine.*
714	89u.c.	† d-Laudanosine, N-Methyltetrahydropapaverine, $\text{Me.C}_1\text{H}_{12}(\text{OMe})_4$. — [Fr. opium in small quantity.] — Tert. amine.* — \textcircled{P} Mandellin's reag. in T. 2.2-(g) gives RS1-VRS1 sol., changing in $\frac{1}{2}$ hr. to OS2. Froehde's reag. in T. 2.2-(f) gives VRT2 color, changing after more than 1 hr. to RT1. H_2SO_4 in T. 2.2-(a) gives v. faint reddish hue which on gently heating w. full flame changes to RV Bkn. (L). — Ndl. fr. bz. I. aq., alk.; e.s. h. bz., lgr.; v.s. alc., chlf.; s. in 19.3 pt. eth. at 16°. — $[\alpha]_D^{25} = +103.23^\circ$ in alc. sol. ($p = 2$). — Odorless. Taste, bitter; salts v. bitter. — $\text{B}_2\text{H}_2\text{PtCl}_6 \cdot 3\text{H}_2\text{O}$, § yel. amorph. ppt., i. c. aq.
715	90-1	Phthalazine, Phenohexa-2,3-diazadiene, $\text{C}_8\text{H}_4\text{N}_2$. — B.p. 315-7° w. partial d. — Flat ndl. fr. eth. V.s. aq.; e.s. alc., bz.; less s. eth.; i. lgr. — B.HCl , ndl. fr. abs. alc., m.p. 231° d. — B.Pk , \ddagger ppt., m.p. 208-10°. — B.Mel , $\ddagger\ddagger$ yel. ndl., m.p. 235-40°. (By allowing sol. in 4 pt. wood alc. + x.s. Mel to stand for 12 hr.)
716	90u.c.	† β -Naphthoquinoline, $\text{C}_{11}\text{H}_8\text{N}$. — B.p. 349.5-50° (th.i., 721 mm.). — \textcircled{P} Tert. amine* of basic odor. Alc. sol. colored brown by FeCl_3 . — Lust. scales fr. h. aq. V.d.s. aq.; v.e.s. alc., eth., bz. — B.Pk , \ddagger light yel. cryst. ppt., m.p. 251-2° (darkening). — $\text{B}_2\text{H}_2\text{PtCl}_6 \cdot 2\text{H}_2\text{O}$, § red-yel. cryst. ppt., i. aq.; v.d.s. HCl.
717	90	Cusparine, $\text{C}_{20}\text{H}_{28}\text{O}_2\text{N}$. — Tert. amine.* [Fr. Angustura bark.] — \textcircled{P} Conc. H_2SO_4 dis. w. dull red, changing to cherry-red color. Conc. HNO_3 dis. yellow. Froehde's reag. (Cf. T. 2.2) gives deep blue color. — Ndl. fr. lgr. E.s. alc., eth., chlf., bz.; less s. lgr. — $\text{B}_2\text{H}_2\text{PtCl}_6 \cdot 6\text{H}_2\text{O}$, § m.p. 179°.
718	90-1	† p-Tetramethylaminodiphenylmethane, $(\text{C}_6\text{H}_4\text{NMe}_2)_2\text{CH}_2$. — Tert. amine.* Colored emerald-green by trace of I. — Dist. unded. Lust. lft. fr. alc. D.s. c. alc.; e.s. eth.; bz. — B.2Pk , \ddagger cryst., m.p. 178°.
719	90.5-1	o-Stilbazole, $[\text{CH: N.C(CH:CHPh):CH.CH:CH}]_2$. — B.p. 324-5° (th.i.). — Tert. amine.* — Cryst. fr. alc. Vol. w. st. "Alc. sol. reacts neut. to litmus." S. alc., lgr.; e.s. eth. — Br, dropped into CS_2 sol. gives dibromide, ndl. fr. alc., m.p. 166-7°. — $\text{B.HCl} \cdot 4\text{H}_2\text{O}$, ndl., m.p. 177°.
720	91	β -Euclidean, Benzoylvinylacetonealkamine, $[\text{NH.CHMe.CH}_2\text{CH}(\text{Ph.CO}_2)\text{C-H}_2\text{CMe}_2]$. — [A synthetic local anesthetic used like cocaine as hydrochloride. Is extracted fr. sol. made slightly alk. w. ammon. by eth. or pet.-eth.] — \textcircled{P} Gives Giesel's test described under No. 2.741, for cocaine, but the color of sol. & ppt. soon begins to turn brown, while in case of pure cocaine the original purple color persists for fully half an hour. — Cryst. e.s. eth., pet.-eth., bz., chlf., amyl alc. — Ammon. in x.s. readily dis. ppt. obtained fr. sol. of salts, as in case of l-cocaine, but unlike α -euclidean. — B.HCl , fern-like or tabular cryst. fr. conc. sol. S. in abt. 33 pt. c., or in less than 16 pt. h. aq.; s. in 9 pt.

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(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
		alc., m.p. 268° d. The hydrochloride is little changed when moistened w. equal vol. aq.; while under same circumstances cocaine hydrochloride dissolves at once! — KI sol. (1:10) gives no ppt. w. mod. dil. B.HCl, or cocaine hydrochloride. α-Euaine under these circumstances gives white, silky, glistening ppt. — [Cf. Parsons, J. Am. Chem. Soc., 23, 885 (1901), for fuller discussion.]
721	92	5-(or 7-)Nitroquinidine, $\text{NO}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_3\text{NMe}$. — Tert. amine.* — Ndl. fr. dil. alc. D.s. aq.; e.s. eth. Vol. w. st. — B.HCl, pr. fr. dil. HCl, e.s. aq. without decn.
722	91.5-2	† Tribenzylamine, $(\text{Ph.C}_6\text{H}_5)_3\text{N}$. — Dec. on distn. — ② Tert. amine.* — Lust. white scales or ndl. fr. alc. I. aq.; d.s. c., e.s. h. alc.; e.s. eth. — Heated above m.p. has mild aromatic odor. Tasteless. Gives no colorations in Tests 2.2-(a), (b), (d), (e). — ③ Dis. 0.10 g. in 5 cc. alc. Add 0.5 cc. HCl (sp. gr. 1.12) & evap. on water-bath in small round bottomed glass dish until cryst. begin to appear at edges. Cool w. persistent stirring. Filter. Wash w. 1 cc. alc. Dry 15 min. on porous tile at 100°. The product, B.HCl, is obtained in pr., m.p. 231.8° u.c. (238.3° c.).
723	92	Dibenzylpiperazine, $[\text{N}(\text{Ph.C}_6\text{H}_5)\text{CH}_2\text{CH}_2\text{N}(\text{Ph.C}_6\text{H}_5)\text{CH}_2\text{CH}_2]$. — ② Tert. amine.* — Ndl. fr. alc.; i. aq.; s. alc., eth., chlf., bz., lgr. — B.MeI,‡‡ (fr. components at 100°), pr. fr. alc., m.p. 217°.
724	93	p-Tolylhydrylamine, $(\text{Me.C}_6\text{H}_4)_2\text{CH}(\text{NH}_2)$. — B.p. 317-8°. — ② Prim. amine.* — Cryst. fr. lgr. E.s. alc., eth., h. lgr. — Acetyl deriv.,¶ ndl. fr. alc., m.p. 159°.
725	93.5	Tetrahydro-β-naphthoquinoline, $\text{C}_{10}\text{H}_{12}\text{N}$. — ② Sec. amine.* Sol. in mineral ac. fluor. blue. — Cryst. fr. lgr. — B.HCl, lust. tbl., m.p. 230.5-1°. — Nitroso deriv.,** $\text{C}_{10}\text{H}_{12}\text{N.NO}$, lust. tbl. fr. alc., m.p. 105.5°.
726	93	Codethyline, Morphineethyllether, $\text{C}_{17}\text{H}_{21}\text{O}_2\text{N(OEt).H}_2\text{O}$. — ② Marquis's reag. [Cf. T. 2.2-(h)] gives: first, a distinct green; then blue; and finally, a fine & rather permanent blue-violet! Otherwise its alkaloid color reactions are similar to those of codeine (No. 2.903). — Lust. pr. s. in 290 pt. aq.; e.s. alc., eth., chlf.; less s. bz.; s. 35-40 pt. boiling aq. — B.HCl.2H ₂ O ("Dionin"), microcryst. powd.; s. in abt. 7 pt. c. aq., or 1½ pt. alc.; i. eth., m.p. 123-5° d.
727	93-4	† Leucomalachite-green, p,p-Tetramethyldiamino-triphenylmethane, $\text{Ph.C}_6\text{H}_4[\text{C}_6\text{H}_4\text{NMe}]_3$. — ② Dis. abt. 0.001 g. in several cc. water acidified by small fraction of a drop HCl taken from glass rod. Add abt. 0.001 g. PbO, and stir. An intense BGS1 to GS1 color develops at once. X.s. of HCl changes to brownish yel., the green being restored by dilution. — White plates fr. h. alc., the sol. coloring dark green by oxidation. Is said to cryst. fr. bz. in ndl. of m.p. 102°. — I. aq.; e.s. alc., eth., bz.
728	94	Caffeidine, $\text{C}_7\text{H}_{12}\text{ON}_2$. — Sec. amine. — Oily, solidifying to alk. cryst. mass. V.s. aq., alc., chlf.; d.s. eth. — Prob. reduces Tollen's reag. (T. 2.30). — Unstable in sol., decomp. to NH ₃ , MeNH ₂ , & cholestophane.
729	95	Trimethyl-2,4,2',4'-tetraamino-diphenylmethane, $\text{Me}_3\text{N.C}_6\text{H}_4(\text{NH}_2)_2\text{CH}_2\text{C}_6\text{H}_3(\text{NHMe})(\text{NH}_2)$. — Aq. sol. boiled gives yel. acridine dyestuff.
730	95-6	1,2-Diaminonaphthalene, $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$. — ② Aq. sol. of B.HCl colored green by FeCl ₃ . — Silvery rhomb. lft. fr. h. aq. E.s. alc., eth., chlf.; less s. h. aq. E. oxidized. — Diacetyl deriv.¶ (fr. ac. anhydride + NaC ₂ H ₅ O ₂), ndl. fr. alc., m.p. 234°.
731	96	1,3-Diaminonaphthalene, $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$. — Lt. — B.2HCl, e.s. aq.; less s. alc. — Diacetyl deriv.,¶ ndl. fr. gl. ac. ac., d.s. alc., m.p. 263°.
732	95-7d.	8-Aminonaphthol(1), $\text{NH}_2\text{C}_{10}\text{H}_4(\text{OH})$. — ② Greenish flocks separate fr. ammon. sol. on shaking w. air. — Ndl. fr. bz + lgr. [A "first component" for azo colors.]
733	96-6.5u.c.	† Aldehydeammonia, $\text{Me.CH(OH)(NH}_2)$. — Dist. w. efferv. & partial decn. to ammonia & acetaldehyde at 90-110°, the distillate solidifying in condenser to clear colorless cryst. of the pure compound. Colorless pr., decomposing & soon becoming yel. on keeping if not perfectly pure originally. Odor, peculiar, faintly pungent. Taste of 1% aq. sol. aldehyde-like, burning, <1 on pungency scale (T. 2.29). V.s. aq.; d.s. eth. Sol. strongly alk. to litmus. — ② Dis. 0.05 g.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
		in 5 drops dil. H_2SO_4 . Add 5 cc. $NaOH$ sol. (1:10). Boil. Within 30–40 sec. the sol. becomes turbid & yel., & the characteristic odor of aldehyde resin appears. (T. 1.111). — \oplus Dis. 0.1 g. in 3 cc. alc. Add 5 drops CS_2 . Allow to stand some time w. occasional shaking. After 10–20 min. a ppt. of colorless rhombic plates begins to separate. After 30 min., filter. Wash w. 2 cc. alc. Dry on porous tile 15 min. at room temperature. The product, carbothialdine, is obtained in rhombic plates, m.p. $129^\circ \pm 1^\circ$ u.c.
734	96	† Semicarbazide, $NH_2CO.NH.NH_2$. — \oplus Gives Ag ppt. immediately in T. 2.30! Reduces Fehling's sol. in the cold. — Pr. fr. abs. alc. E.s. aq., bz., chlf. — Dec. on keeping. — $B.HCl$ (dried at 100°), m.p. 175° w. efferv. — \oplus Add 3 drops benzaldehyde to 0.05 g. $B.HCl$ in 5 cc. h. aq. Filter off ppt., after cooling & vigorous shaking. Recryst. fr. 5 cc. h. alc. After long standing ndl. of the semicarbazone separate. Filter. Wash w. 1 cc. alc. Dry 15 min. at 100° on tile. The product, benzalsemicarbazone, $NH_2CO.NH.N:CH.Ph$, melts w. efferv. to clear liquid at $213-4^\circ$ u.c.
735	95; 97.5	Ethylenedi-p-tolyl diamine, $(Ph.CH_2)CH_2CH_2(CH_2Ph)_2$. — Sec. amine.* — V.s. alc. — $B.HgCl_2$, m.p. 133° c. (J. Chem. Soc., 77, 1022). — Dinitrosamine deriv.,** yel. cryst., i. alc., m.p. 183° .
736	96	p,p'-Tetramethyldiamino-benzhydrol, $(MeN.C_6H_4)_2CH(OH)$. — Tert. amine.* — \oplus S. in ac. w. blue color! — Triclin. cryst. fr. eth. or bz. E.s. alc., eth. — $B.Pk$; dark-green cryst. granules, i. eth.; d.s. bz. — [An intermediate in dyeestuff manufacture.]
737	97	m-Aminobenzyl Alc., $NH_2.C_6H_4.CH_2.OH$. — \oplus Prim. amine.* — Browns slowly in air. Tbl. fr. bz. V.s. alc., chlf., mineral acids, h. aq.; s. eth.; d.s. bz. — Diacetyl deriv.¶ (by heating w. ac. anhydride), ndl. fr. lgr. + bz., m.p. 67° .
738	97	Flavainiline, 4-Methyl-2-p-aminophenylquinoline, $C_{12}H_{14}N_2$. — \oplus Prim. amine* giving N in T. 2.4. — Long colorless cryst. fr. bz., becoming yel. on keeping. Vol. without decon. V.d.s. aq.; e.s. alc. — $B.2HCl$, colorless ndl. d.s. conc. HCl , e.s. aq. Addition of $NaC_2H_3O_2$ & $NaCl$ to aq. sol. ppts. $B.HCl$. $1\frac{1}{2}H_2O$ in yel.-red pr. e.s. aq. w. yel. color & moss-green fluor. — Acetyl deriv.,¶ pr. fr. dil. alc., m.p. $162-3^\circ$.
739	96.5-7	Tetrahydroquinoxaline, $C_8H_8NH.CH_2CH_2.NH^2$. — \oplus Conc. aq. sol. colored violet by $FeCl_3$. — Lft. fr. eth. D.s. aq.; e.s. chlf., bz. — Dinitroso deriv.,** (fr. sol. B in HCl at 0° + x.s. $NaNO_2$), light-yel. mic. ndl., m.p. 168° d.
740	96.5-7.5 u.c.	† Homatropine, Phenylglycolyltropeine, $C_8H_{11}ON.CO.CH(OH).Ph$. — [Synthetic mydriatic alkaloid used chiefly as hydrobromide. A drop of 1% sol. on the eye causes dilation within abt. 8 min., reaching a maximum after 1 hr., the effect disappearing after 20 hr. Dilation by atropine, even w. weaker solutions, is much more persistent. (Ann. 217, 86.)] — Glassy cryst. fr. abs. eth. or CS_2 . S. alc., eth., chlf.; less s. aq. Odorless. Taste, faintly bitter (poisonous), without producing numbness like cocaine. \oplus (1) Apply Guelmo's react. as directed for atropine (No. 2.797); but drop in a small cryst. of $K_2Cr_2O_7$ before decomposing w. aq. The odor of bitter almonds will be observed. — (2) Vitali's react. as described under atropine, gives only a YT1 instead of V color. (This coloration is said to be sometimes orange or yellow-brown, & it is claimed that a very feeble violet coloration is sometimes noticed on the margin as the alc. evaporates.) — (3) Gerard's react. gives same results as with atropine. — (4) Test (5) w. Br. + HBr described under atropine gives broad flat mic. ndl. similar to those fr. atropine. — \oplus Add 4 cc. sat. aq. sol. of picric ac. to sol. of 0.05 g. B in 5 cc. aq. + 2 drops HCl (sp. gr. 1.12). Filter off ppt. Wash w. 2 cc. aq. Recryst. fr. 10 cc. h. aq., allowing to cool slowly. Filter. Wash w. 1 cc. aq. Dry 15 min. at 100° . The picrate thus obtained forms fine ← YT1 lft., melting to clear liquid at $180.5-1.5^\circ$ u.c.
741	98u.c.	† 1-Cocaine, Methyl Ester of Benzoylecgonine, $C_{17}H_{21}O_3N$. — [Local anesthetic & mydriatic alkaloid fr. Erythroxylon coca.] — Large, clear monoclin. pr. fr. c. alc. S. at 25° in: — 600 pt. aq.; 5 pt. alc.; 38 pt. eth.; v.s. chlf., h. alc.; s. bz., pet.-eth., CS_2 . Odorless. $[\alpha]D^{20} = -16.4^\circ$ (in 10–20% chlf. sol.). \oplus [A sol. of $B.HCl$ should be prepared for the following tests by carefully neutralizing a little of the base w. dil. HCl and evapn. to dryness on water-

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No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
		bath. (1) Physiological T. — Place a drop of 3% B.HCl sol. on 1 sq. cm. filter paper. Lay this on upper surface of the tongue near the tip. After 10 or 15 sec. add a second drop of solution to the paper, and after 30 sec., a third. The paper may be removed and mouth rinsed after abt. a minute. The taste produced is slightly saline & bitter, followed by a peculiar sensation of tingling and numbness of the part of the tongue in contact with the sol. which reaches a maximum after 2 or 3 min. and nearly disappears within 10 min.! (Sensitiveness of individuals in this test is variable.) — (2) Giesel's React. (Z. anal. chem., 47, 447). — Add 1% KMnO_4 sol. drop by drop to sol. of abt. 5 mg. B.HCl in 1 cc. aq. A VR cryst. ppt. of cocaine permanganate separates. Examine some of this ppt. on a slide under the microscope. It will be found to consist of well-developed rhombic plates of characteristic appearance! (It is said that the plates are sometimes arranged in rosettes, and that presence of impurities is liable to interfere w. react., although the test is usually delicate.) — (3) Greitherr's T. (Pharm. Ztg., 34 (1899), 617). — Add to 3 drops 5% aq. sol. of B.HCl 3 cc. chlorine aq. & 3 drops palladious chloride sol. An OSI-O ppt. appears gradually decd. by aq., i. alc. & eth., s. in sod. thiosulphate. (This react. while not v. delicate, is said not to be given by other alkaloids.) — (4) Ethyl Benzoate React. — Dis. 0.01 g. B.HCl in 1 cc. alc., & in this sol. dis. by stirring 0.25 g. KOH. Dil. in a small open glass dish w. 3 vol. h. aq. The sweet aromatic odor of ethyl benzoate will be produced. — (5) Chromic Acid T. — Add 5 drops of 5% aq. Cr_2O_7 sol. to 1 cc. 2% aq. sol. B.HCl . Each drop causes a ppt. which disappears on shaking. Add 1 cc. HCl. A cryst. YO_2 ppt. of cocaine chromate appears. — (6) I in KI. — Add 2 drops of the reagt. to sol. of 1 mg. B.HCl in 10 cc. aq. A ppt. of OSI-YOSI color (on filter) appears. (Delicacy according to Dragendorff, 1:100,000.) ① Heat 0.1 g. B 3 min. on boiling water-bath w. 2 cc. conc. H_2SO_4 . Cool. Dilute w. 5 cc. aq., cooling. Filter off ppt. of benzoic ac. Wash w. few drops aq. Recryst. fr. 1 cc. boiling aq. Filter. Wash w. 0.5 cc. aq. Dry 15 min. at 100°. The benzoic ac. melts at 120° (u.c.). [For discussion of differences between cocaine & the eucaines, cf. Parsons, J. Am. Chem. Soc., 1901, 885.]
742	98	Benzophenonehydrazone , $\text{Ph}_2\text{C:NH}_2$. — B.p. 225–30° (55 mm.). — ② T. 2.17 splits to hydrazine & benzophenone. — Pr. fr. alc. D.s. alc.; e.s. eth., bz.; e.s. dil. H_2SO_4 & ptd. fr. sol. by alk. — Benzoyl deriv., ¶¶ m.p. 116.5°; e.s. alc., eth.
743	99–100	Amylenenitrolamine , $\text{Me.C(NOH).CMe}_2\text{NH}_2$. — ② Gives dark red-violet color to conc. aq. CuSO_4 sol. — Sbl. in long silky ndl. V.s. aq., alc.; less s. eth.; i. lgr. "Strongly alk." — B.HCl , cryst. powd.; v.s. aq.; m.p. 186° d.
744	99 u.c.	† 2,4-Toluylenediamine , $(\text{NH}_2)_2\text{C}_6\text{H}_3\text{Me}$. — B.p. 280°. — E. crystd. in colorless, odorless ndl. fr. lgr. S. c. aq.; e.s. alc., eth. Taste of c. sat. aq. sol. sl. burning & bitter. — ② Treat 0.05 g. w. 5 drops acetic anhydride. Heat mass gently over small flame. Boil w. 5 cc. aq. Cool well & shake persistently. A heavy ppt. of colorless ndl. finally separates. Filter. Wash w. 3 cc. aq. Recryst. fr. 3 cc. h. aq., cooling & shaking persistently. Dry filtered cryst. 15 min. at 100°. The product, diacetyltoluylenediamine, is obtained in white ndl., m.p. 220–0.4° u.c. (226.8–6.2° c.).
745	99–100	(m)- 3,3'-Hydrazodimethylaniline , $\text{Me}_2\text{N.C}_6\text{H}_4\text{NH.NH.C}_6\text{H}_4\text{NMe}_2$. — Ndl. fr. eth., v.s. bz.
746	99	d,l-Lupanine , $\text{C}_{16}\text{H}_{22}\text{ON}_2$. — [Fr. seeds of white & blue lupine.] — Monoclin. pr. fr. pet.-eth. E.s. aq., alc., eth., chlf.; alm. i. lgr. Aq. sol. becomes cloudy on boiling. Taste v. bitter. — B.HAuCl_4 , lust. yel. cryst. fr. dil. alc., m.p. 182–3° d. — B.Mel , ¶¶ cryst. e.s. aq., m.p. 239–40°.
746-I	99	Carbazolin , $\text{C}_{12}\text{H}_{18}\text{N}$. — Silky ndl. Alm. i. aq.; v.s. alc., chlf., eth. E. vol. w. st. ("Weak base" giving e.s. salts w. min. acids.)
747	100	Piperazylidihydrazine , $[\text{N}(\text{NH}_2)\text{CH}_2\text{CH}_2(\text{NH}_2)\text{N.CH}_2\text{CH}_2]^+$. — B.p. 228°. — ② Reduces Fehling's sol. on boiling. — Ndl. fr. eth.-alc. V.s. aq., alc.; d.s. eth.
748	abt. 100	Leucoaniline , Triaminodiphenyltolylimethane , $(\text{NH}_2\text{C}_6\text{H}_4)_3\text{CH.C}_6\text{H}_3\text{Me}(\text{NH}_2)_3$. — Cryst. fr. h. aq. D.s. h. aq., eth.; v.s. alc. — ② Warmed w. dil. ac. ac. + trace PbO_2 gives red sol. — Boiled w. acetic anhydride gives triacetyl deriv., ¶ ndl. fr. g. ac. ac., m.p. 168°.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
748-I	100	Triphenylrosaniline, $(\text{Ph.NH.C}_6\text{H}_4)_2\text{C(OH).C}_6\text{H}_4\text{Me.NHPh}$. — \textcircled{P} Becomes bluish during washing & drying. — Indistinctly cryst. mass fr. pptn. of salt (aniline blue) by ammon. E.s. alc., eth. — Dry dist. gives diphenylamine, (No. 2.1568)! — $\text{C}_{18}\text{H}_{14}\text{N.Cl}$, bluish-brown cryst. grains, d.s. alc. w. blue color.
749	100	Aspidosamine, $\text{C}_{22}\text{H}_{28}\text{O}_2\text{N}_2$. — [Alkaloid fr. bark of Aspidosperma Quebracho.] — \textcircled{P} Sol. in conc. H_2SO_4 bluish, becoming dark blue w. a little $\text{K}_2\text{Cr}_2\text{O}_7$. — Alm. i. aq., alk.; v.s. alc., eth., chlf., bz.; v.d.s. lgr. “Alc. sol. reacts strongly basic.” Taste bitter. — $\text{B}_2\text{H}_2\text{PtCl}_6\cdot 3\text{H}_2\text{O}$, § pale yel. floc. ppt.
750	100	Benzylidenequinaldine(2), $[\text{C}_6\text{H}_4\text{N}:\text{C(CH:CHPh).CH:CH}]^2$. — Cryst. Sbl. I. aq.; e.s. alc., chlf. — Adds Br_2 giving iridescent lft. fr. alc., m.p. 173-4 $^{\circ}$.
751	100.5 u.c. (101.2 c.)	† o-Phenylenediamine, $\text{C}_6\text{H}_4(\text{NH}_2)_2$. — B.p. 256-8 $^{\circ}$ (th.i.). — Cryst. fr. chlf. (best solvent for purification) in plates; d.s. c. aq.; e.s. h. aq., alc., eth., chlf. — Odorless. Alm. tasteless. \textcircled{P} Dis. 0.005 g. in 3 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 2 drops 10% FeCl_3 sol. Shake & allow to stand. After 2 or 3 min. shimmering red ($R-O\text{R}'$)ndl. separate! — \textcircled{D} Treat 0.1 g. B w. 6 drops acetic anhydride & heat to boiling. Boil reaction product w. 2½ cc. aq. Cool & shake persistently. Filter. Wash w. 1 cc. aq. Cryst. fr. 1½ cc. h. aq., cooling & shaking (v. persistently!). Dry 15 min. at 100 $^{\circ}$. The resulting diacetyl deriv., whose quantity is small, melts at 182-3 $^{\circ}$ u.c. (185.5-6.5 $^{\circ}$ c.).
752	100.5	Aminoacetyl-p-phenetidine, $\text{EtO.C}_6\text{H}_4\text{NH.CO.CH}_2(\text{NH}_2)$. — [The antipyretic “Phenocoll” is B.HCl.I .] — Cryst. w. 1 mol. H_2O , m.p. abt. 95 $^{\circ}$.
753	101-2	p-Toluenamidine, $\text{Me.C}_6\text{H}_4\text{C}(\text{NH})(\text{NH}_2)$. — \textcircled{P} Should give NH_3 in T. 2.6; & p-toluic ac. on sapon. — Pearly lft. fr. bz. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yel. ndl., d.s. aq., m.p. 225 $^{\circ}$.
754	101 u.c.	† Hydrobenzamide, $(\text{Ph.CH})_2\text{N}_2$. — \textcircled{P} Warmed w. dil. HCl gives odor of benzaldehyde. — Pr. fr. abs. alc. Tasteless & odorless when fresh & pure, but soon developing odor of benzaldehyde on exposure to moist air. I. aq.; e.s. alc., eth. — \textcircled{D} Heat just to boiling sol. of 0.05 g. B in 10 cc. 50% alc. + 4 drops HCl (sp. gr. 1.2). Add 1 drop pure phenylhydrazine & boil 30 sec. Cool & shake. Filter. Wash ppt. w. 5 cc. 50% alc. Dis. in 5 cc. 50% alc. On cooling a bulky ppt. of fine white ndl. separates. Dry 15 min. at 100 $^{\circ}$ & determine m.p. without delay! The product, benzylidenephenylhydrazone, melts at 155.3 $^{\circ}$ u.c. (157.9 $^{\circ}$ c.).
—	101	Amarine. (Air dried). — Cf. No. 2.853.
—	102	Leucomalachite-green. (Crystd. fr. bz.) — Cf. No. 2.727.
756	102	2,6-Diaminoxyleno, $(\text{NH}_2)_2\text{C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Warmed w. dil. H_2SO_4 + little FeCl_3 or $\text{K}_2\text{Cr}_2\text{O}_7$ gives cherry-red color & pungent quinone odor. — Sbl. in ndl. Pr. fr. bz. + lgr. E.s. alc., h. aq.
757	103	1,2,3-(v)-Triaminobenzene, $\text{C}_6\text{H}_3(\text{NH}_2)_3$. — B.p. 336 $^{\circ}$ c. — \textcircled{P} Gives dark blue color when disd. in conc. H_2SO_4 + trace HNO_3 ! — Aq. sol. reduces AgNO_3 sol. in cold, & produces violet, followed by brown ppt. w. FeCl_3 sol. — Cryst. (usually colored fr. impurities) soften before melting. E.s. aq., alc., eth. “Reacts strongly alk.” — B.2HCl , ndl. v.s. aq.
758	103-4	2,5-Diaminoxyleno(1,3), $(\text{NH}_2)_2\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} FeCl_3 gives green color in neut. or ac. ac. sol. — Lft. fr. bz. + lgr. E.s. except in lgr.
759	102; 103; 105	Triphenylmethylamine, $\text{Ph}_3\text{C.NH}_2$. — Ndl. fr. alc. I. aq.; e.s. alc., eth., bz. — $\text{B}_2\text{H}_2\text{PtCl}_6\cdot 7\frac{1}{2}\text{H}_2\text{O}$, § yel. ndl. fr. aq.; loses all aq. at 120 $^{\circ}$; d.s. aq.; e.s. alc.
760	103; 104.5	α -Euaine, Methyl 1,2,2,6,6-pentamethyl-4-benzyloxy(piperidine)carbonate, $\text{C}_{18}\text{H}_{28}\text{O}_2\text{N}$. — [The hydrochloride is a local anesthetic resembling β -euaine (No. 2.720) & cocaine (No. 2.741). Cf. J. Am. Chem. Soc., 1901, 885. Extracted by eth. or pet.-eth. fr. alk. sol.] — Cryst. E.s. eth., pet.-eth., chlf., bz., amyl alc. \textcircled{P} (1) Phenomena in Giesel's react. as described for β -euaine No. 2.720. — (2) Conc. ammon. added to sol. of salts does not readily re-dissolve ppt. formed as in case of cocaine & β -euaine. — (3) One drop 1% sol. w.

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No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
		10% KI sol., unlike β -eucaine or cocaine, gives silky cryst. ppt. — (4) One drop 1% sol. w. 5% HgCl ₂ sol., like cocaine & unlike β -eucaine, gives ppt. — (5) Conc. K ₂ Cr ₂ O ₇ sol. added to 0.5% sol., unlike cocaine or β -eucaine, gives lemon-yel. cryst. ppt. — B.HCl, unlike cocaine salt, is s. at once in an equal vol. aq. or alc.
761	103.5-5	2,6-Diaminotoluene, Me.C ₆ H ₄ (NH ₂) ₂ . — (P) Sol. colored deep blue by FeCl ₃ ! — Cryst. V.s. aq. 100 pt. c. aq. sol. contain 18 pt. — Diacetyl deriv. [T. 2.1], m.p. 202-3°. — [An intermediate in dyestuff manufacture.]
762	104	4,6-Diaminoxyleno(1,3), Me ₂ C ₆ H ₄ (NH ₂) ₂ . — (P) Neut. sol. gives or.-yel. color w. FeCl ₃ on warming. — Ndl. fr. lgr. E.s. aq., alc., eth., h. bz.; s. h. lgr.
763	104	Phenanthridine, ^c C ₆ H ₄ .C ₆ H ₅ .CH:N ² . — B.p. a. 360°. — (P) Aq. sol. of base & dil. aq. sol. of salts fluor. blue. Alm. odorless, but warm vapors provoke sneezing. Taste pungent. — Long hair-like ndl. fr. dil. alc. S. in 3500 pt. c. aq. E.s. alc., eth., chlf., bz. Salts yellow. — B.Pk, long ndl. not melting at 220°. — B.HCl.HgCl ₂ , pptd. by HgCl ₂ , fr. sol. in dil. HCl, m.p. 197°.
764	105	Lysidine, Methylglyoxalidine, ^c CH ₂ .CH ₂ .N:CMe.NH ⁺ . — [Solvent for uric ac. like piperazine.] B.p. 195-8°. — Deliq. cryst. v.s. aq., alc.; i. eth. — "Strong base." KI + I gives red-brown ppt. — (P) Shake dil. aq. sol. persistently w. x.s. benzoyl chloride & large x.s. NaOH sol. Wash w. aq. & recryst. fr. h. alc. The dibenzoyl deriv. formed (Ber. 28, 3068) melts at 244° & is obtained even fr. a 0.1% of B.
765	105.6 u.c.	† Piperazine, ^c NH.CH ₂ .CH ₂ .NH.CH ₂ .CH ₂ . — B.p. 145-6°. — [Being v. deliq., commercial product must be treated for some time in fused state w. solid KOH, or will melt many degrees too low.] Softens 2° or 3° below m.p. Hygroscopic cryst. E.s. aq., alc. Odorless. Taste, bitter. — (P) A cold glass rod brought into vapors of boiling compound occasions charac. ppt. of shimmering iridescent scales! Gives Simon's react. (T. 2.28) feebly. No colorations in T. 2.2-(a), (b),(d),(e). — (P) Add 2 cc. c. sat. aq. sol. NaNO ₂ to sol. of 0.2 g. B in 3 cc. aq. + 1 cc. HCl (sp. gr. 1.12). Warm gently. Shake. A gradual pptn. of yellowish scales appears. When complete, cool, filter, & wash w. 1 cc. c. aq. Dry 15 min. at 100°. The product, dinitrosopiperazine, is obtained in pearly white scales, m.p. 157° u.c.
766	105-6	Pseudoconhydrine, C ₈ H ₁₁ ON. — B.p. 236-6.5° [α] = +10.98°. — KI + I sol. (T. 2.3) gives amorph. ppt. at conc. 1:1000. Picric ac. gives no ppt. at conc. 1:100. — B.HCl, not deliq., m.p. 212-3°. — B.(HAuCl ₄) ₂ cryst. (separating oily), m.p. 133-4°. — B ₂ H ₂ PtCl ₆ , § v.s. aq., m.p. 185-6°.
—	105-6	Ethylsemicarbazide, EtHN.NH.CO.NH ₂ . — Cf. No. 2.1899.
767	105-6	† Physostigmine, Eserine, C ₁₈ H ₂₁ O ₂ N ₃ . — [Toxic alkaloid fr. seeds of Physostigma venenosum (Calabar bean). Causes pronounced contraction of pupil, persisting 10-24 hr. Even 0.1 mg. brought into conjunctival sac suffices to produce marked contraction in physiological test!] — Flat trimet. pr. fr. c. bz. Odorless. Tasteless. Levorotatory. D.s. c. aq.; e.s. alc., eth., bz., chlf. Sol. unstable, turning red on exposure to air & light. Alk., ammon., & alk. carbonates ppt. fr. sol. of salts, sol. becoming red. — (P) (1) Evap. to dryness on water-bath sol. of 0.001 g. in 1 drop HCl (sp. gr. 1.12). Add 0.5 cc. h. conc. ammon. An OT1 color appears. Evap. to dryness on water-bath. A BT1 residue is obtained. Add 0.3 cc. alc. A BVS1 sol. results. Add 2 cc. 10% ac. ac. A fine fluor. sol., RT1 by reflected light against dark background, & VRT1 by transmitted light w. sky background, will be observed! W. 0.05 mg. B the test is still applicable, though color tones are much paler. — (2) Add 1 drop gold chloride reagl. to sol. of 0.001 g. B in 5 cc. aq. + 1 drop dil. HCl & warm. An orange-red color ("rubreserin") appears, but there is no evident ppn. or reduction at first. Gradual reduction, more or less rapid according to conditions, occurs on heating. — B ₂ H ₂ PtCl ₆ , § (fr. sol. of B in conc. HCl by addition of 10% platinic chloride sol.), or.-yel. ndl. in stellate groups, d. fr. 180°. — B.(HAuCl ₄) ₂ (fr. sol. of B in conc. HCl by adding 5% gold chloride sol. drop by drop w. shaking), at first oily, then fine yel. lft., m.p. 163-5° d. — [Cf. Rosenthaler, p. 754 for additional react.]
768	105-6	Benzoylcinchonine, C ₂₁ H ₂₁ ON ₂ (Ph.CO). — Amorph. I. aq. Misc. alc., eth. E. sapd. by alc. KOH. [α] _D ²⁴ in abs. alc. (p = 1) = -22.26°.

No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
769	105	Guanylurea, Dicyandiamidine, HN:C(NH₂).NH.CO(NH₂)₂. — Strongly basic hygroscopic cryst. — \textcircled{P} . T. 2.7 gives NH ₃ freely. — E.s. aq.; d.s. alc.; i. t.z., chlf., eth. — B.Pk, \ddagger yel. lft. fr. aq., m.p. 265°, d. abt. 285°. — B ₂ H ₄ SO ₄ .2H ₂ O sol. added in x.s. to ammon. sol. of Ni salts, followed by addition of NaOH sol. in x.s., gives yel. cryst. ppt. of Ni(C ₂ H ₄ N ₂ O) ₂ .2H ₂ O. [Cf. Die Chemische Analyse, XVI, 37, for bibliography of use of compound in determination of Ni.]
770	107	p-Tolyltolylenediamine, 4-Amino-3-p-toluidinotoluene, Me.C₆H₄(NH₂).NH-C₆H₄CH₃. — \textcircled{P} Solid NaNO ₂ added to sol. in conc. H ₂ SO ₄ gives deep blue color. — Prim. amine.* — Lft. of fatty luster fr. dil. alc. V.s. alc., eth., bz.
771	108-9	Ethyl Ester of L-Tyrosine, p-HO.C₆H₄.CH₂(NH₂).CO₂Et. — \textcircled{P} Should give Millon's react. (T. 2.19)! — Prim. amine.* — Flat ndl. fr. acetic eth. V.d.s. c. aq.; e.s. alc.; v.d.s. eth. — $[\alpha]_D^{20} = +20.4^\circ$ for 5% alc. sol.
772	108	Aminoacenaphthene, NH₂.C₁₂H₈. — Prim. amine.* — Ndl. — Acetyl deriv. \ddagger (fr. acetyl chloride), dark yel. lft. fr. alc., m.p. 176°.
773	108	o-Aminoacetophenonephenylhydrazone, NH₂.C₆H₄CMe:N.NH.Ph. — \textcircled{P} Prim. amine.* — Fine ndl. fr. alc.; r.s. h. aq.; e.s. h. alc., eth., chlf., bz.; d.s. lgr. \dagger Acridine. — Cf. No. 2.3102. (Color, YT3.)
—	108 u.c.	
775	108	Acetylquinine, C₂₀H₂₁O₂N.(Me.CO). — \textcircled{P} Sol. in dil. H ₂ SO ₄ fluor. blue! Gives color react. w. Cl-aq. & ammon. like quinine (No. 2.947), — Pr. fr. eth. E.s. alc., chlf., dil. acids; less s. eth. Taste v. bitter. — E. sapd. to quinine by alc. KOH.
776	107-8 u.c.	\dagger Pyramidone, 4-Dimethylamino-antipyrine, "NPh.NMe.CMe:CNMe₂.CO". — [Synthetic antipyretic. Is extracted fr. alk. sol. on shaking out w. eth. or chlf.] — Colorless cryst. fr. ethyl acetate + lgr. S. in abt. 30 pt. c. aq., more s. h.; v.s. alc., chlf.; e.s. eth. <i>Odorless. Taste, faintly bitter.</i> — \textcircled{P} (1) 4 drops 10% FeCl ₃ added to sol. of a few mgr. substance in 10 cc. aq. gives following colors: VBS2 (30 sec.); RV (2 min.); RVT2 (15 min.); colorless ($\frac{1}{2}$ hr.). — (2) 5 drops HNO ₃ added to 5 cc. conc. aq. sol. gives VBS2 color. — (3) Gives blue color w. AgNO ₃ in aq. sol. After some time sol. blackens w. separation of Ag. (Gadamer.)
777	107-8 u.c.	\dagger 1-Hyoscyamine, Ester of L-Tropine and L-Tropic Ac., C₁₇H₂₁O₂N. — [Mydriatic alkaloid fr. Hyoscyamus niger L.]. — Plates fr. chlf. D.s. c. aq.; more s. h. aq.; e.s. alc., amyl alc., chlf., bz.; i. pet.-eth. Tends to change to its stereoisomer atropine & to hydrolyze when kept long time in sol. $[\alpha]_D^{25} = -20.3^\circ$ (for 3% sol. in abs. alc.). \textcircled{P} Gives reactions (1), (2), (3), (4), described for Atropine (No. 2.797). Gerrard's react. (Cf. (5) under Atropine) gives no ppt., while the ppt. w. atropine is heavy. The physiological test, (1), under atropine, is more delicate w. hyoscyamine (according to Blyth, p. 385) than w. atropine. — \textcircled{P} Add 5 cc. gold chloride sol. to 0.05 g. B. disd. in 5 cc. aq. + 2 drops HCl (sp. gr. 1.12). Wash yel. powdery ppt. on filter w. 2 cc. aq. Recryst. fr. 10 cc. boiling aq. Fine, large, lust. Y lft. separate. Wash w. 2 cc. c. aq. Recryst. fr. 8 cc. boiling aq. Wash w. 1 cc. aq. on filter. Dry 15 min. at 100°. The product, B.HAuCl ₄ , after softening at 159-60°, melts at 161.5° u.c.! — B.Pk, \ddagger m.p. 161-3°. — B ₂ H ₄ PtCl ₆ , § orange pr., m.p. 206°.
778	108	Pseudotropine, ψ-Tropanol, C₈H₁₁ON. — B.p. 240-1°. Ndl. fr. lgr. V.s. aq., alc. “Sol. strongly alk.” Not vol. w. st. Opt. i. — B.Pk, \ddagger m.p. 257-8° d. (“Charac. deriv.” Ber. 33, 1172.)
779	109-10	5-Aminoquinoline, NH₂.C₉H₇N. — Prim. amine.* — Silky ndl. fr. alc. Sbl. D.s. c. aq.; e.s. alc., eth.; alm. i. lgr. — Picrate, \ddagger long red ndl. fr. alc., alm. i. eth. — Acetyl deriv., \ddagger m.p. 178°.
780	abt. 109	Oscine, Scopoline, Oxytropine, C₉H₁₁O₂N. — [Alkaloid accompanying atropine.] — B.p. 241-3°. Tert. amine.* Clear hygroscopic cryst., soon becoming brown & dull. — Benzoyl deriv., ndl. fr. chlf., m.p. 59° (Ann. 271, 119).
781	110 u.c.	\dagger β-Naphthylamine, C₁₀H₇.NH₂. — B.p. 304.5° c. <i>Odorless (unlike α-isomer).</i> Prim. amine.* — Pearly scales fr. h. aq. or lgr.; alm. i. c. aq. — $[\alpha]_D^{20} = 2.$ — \textcircled{P} Convert into the corresponding picramide by T. 2.22. The product, C ₁₀ H ₇ .NH.C ₆ H ₂ (NO ₂) ₂ , is obtained as a heavy ppt. which cryst. fr. ac. ac. in YO-O pr., m.p. 232° u.c.!

Explanation of typographical signs in this Division: * = T. 2.35. \dagger = Generic position established by actual titrations. \ddagger = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
782	110	Pellotine, $C_{12}H_{16}O_2N$. — [Alkaloid fr. Mexican "mescal buttons," Anhalonium Williamsi.] — Cleartbl. fr. alc. Alm. i. aq.; e.s. aq.-alk.; sol. alc., eth., chlf.; less s. pet.-eth. Taste very bitter & persistent. — \textcircled{P} Addition of little HNO_3 to colorless sol. in conc. H_2SO_4 gives intense permanganate-like color! Aq. sol. of salts give blue color w. $FeCl_3$, changing to green & disappearing on warming. — $B.Mel$, \ddagger easily formed fr. components in conc. $MeOH$ sol., pr., crystg. w. $1H_2O$, m.p. 198° .
783	110	Cusconine, $C_{20}H_{32}O_2N_2$. — [In Cusco barks.] — Cryst. fr. eth. in dull lft. w. $2H_2O$ (lost at 100°). Alm. i. aq.; s. in 35 pt. eth.; more s. alc.; v.s. chlf.; v.d.s. bz., lgr. "Alc. sol. feebly alk." — \textcircled{P} Colored dark green by conc. HNO_3 , dissolving w. greenish yel. color. — $B_2H_2SO_4$ (at 100°) gelat., yel., horny after drying.
784	110-1	Methyl 3-Amino-4-hydroxybenzoate, $HO.C_6H_4(NH_2).CO_2Me$. — Dimorphic. Cryst. fr. chlf. white. Orange ndl., m.p. 142° , fr. chlf., bz. E.s. h. aq., alc., eth. $FeCl_3$ gives muddy green ppt. or color. — $[B.HCl]$, m.p. 225° , synthetic anesthetic, "Orthoform-neu." D.R.P. 97,333, 97,334.]
785	108-13	Euphthalmine, $^{\textcircled{P}}NMe.CHMe.CH_2CH[Ph.CH(OH).CO_2]CH_2.CMe_2$. — $B.HCl$ is the synthetic mydriatic known by this name. — 6-sided pr. fr. h. pet.-eth. Tasteless; odorless; not a local anesthetic; only slightly toxic. Alm. i. c. aq.; mod. s. h. E.s. org. solvents, except pet.-eth. — Gives ppts. (T. 2.3) w. alkaloid reagt.; w. bismuth-potassium-iodide sol. at dilution 1:20,000, a distinct red ppt. — $B.HCl$, spheroidal aggregates fr. abs. alc. + little eth., v.s. aq., m.p. $183-4^{\circ}$.
786	110	2,2'-Dihydrazinobiphenyl, $NH_2.NH.C_6H_4.C_6H_4.NH.NH_2$. — Lft. fr. bz. V.s. chlf., alc., h. bz.; s. eth., h. aq.; i. lgr. — Diacetyl deriv. (by boiling w. gl. ac. ac.), cryst. fr. ac. ac., m.p. $250-60^{\circ}$ d.
787	111-2	Ethylphenylsemicarbazide, $NHET.NH.CO.NHPh$: — \textcircled{P} Easily hydrolyzed in T. 2.17. — Lft. r.d. s. h. aq.; e.s. alc.; e.s. dil. ac. & repptd. by alk.
788	111-2	4,4'-Bipyridyl, $NC_6H_4.C_6H_4N$. — B.p. 305° c. — \textcircled{P} Few drops K_4FeCy_4 sol. added to not too dil. sol. $B.2HCl$ gives light-colored ppt. soon changing to dull indigo blue, & disg. in h. aq. w. purple red color. — Tbl. w. fatty luster. Alm. i. c. aq.; mod. s. h. Cryst. fr. aq. w. $2H_2O$ (lost on distn.), m.p. 73° . V. stable. Tastes bitter. Oxidation w. $KMnO_4$ gives pyridine-4-carbonic ac. (No. 2.500). — $B.2HCl$, cryst. v.s. aq.
789	112.5-13; 108	o-Hydroxybenzylaniline, $HO.C_6H_4.CH_2.NH.Ph$. — \textcircled{P} Sec. amine.* Ndl. fr. alc. D.s. aq., lgr.; e.s. alc., eth.; s. acids or alkalis. — Nitrosamine,** v. pale yel. pr. fr. dil. $MeOH$, m.p. 131.5° .
—	112.5	Benzoylhydrazine. — Cf. No. 2.1953.
790	113	Gallipidine, $C_{19}H_{21}O_2N$. — [Angustura bark.] — Tert. amine.* Lft. e.s. alc., eth. chlf., bz. — $B.Mel$, \ddagger cryst. powd., m.p. 142° .
791	114	6-Aminoquinoline, $NH_2.C_6H_4N$. — \textcircled{P} Aq. sol. strongly yel.! — Prim. amine.* Cryst. w. $2H_2O$ (lost over H_2SO_4). Turns brown on exposure. Sbl. E.s. alc., eth., ammon.; s. aq., lgr. — $B.2HCl$, clear, glassy, cryst., e.s. aq. w. intense yel. color. — $B.2Pk$, \ddagger woolly ndl. — $B_2H_2PtCl_6.2H_2O$, \ddagger yel. cryst. ppt.
792	114-5	2,5(β)-Dimethylpiperazine, $^{\textcircled{P}}NH.CHMe.CH_2.NH.CHMe.CH_2.NH^{\textcircled{P}}$. — B.p. 162° (th.i.). Sec. amine.* Tbl. fr. chlf. V.s. aq., alc., chlf.; v.d.s. eth. — Picrate \ddagger alm. i. aq. — Dinitroso deriv.,** m.p. $95-6^{\circ}$ (J. prakt. Chem., 1893, 513).
793	114	1-Methyl-8-hydroxytetrahydroquinoline, $MeN.C_6H_4.C_6H_4(OH)$. — \textcircled{P} A drop $FeCl_3$ colors alc. sol. deep brown. $NaNO_2$ added to dil. H_2SO_4 sol. gives intense red-yel. dyestuff. — Trimet. tbl. fr. eth. D.s. aq.; e.s. KOH sol., bz., warm alc. Strong tertiary base. — $B.HCl.H_2O$, lust. monoclin. cryst., the obsolete antipyretic "Kairin M."
794	114-5	Lycopodine, $C_{12}H_{16}O_2N_2$. — [Fr. Lycopodium complanatum.] Resinous ppt. fr. sol. by alk. slowly changing to monoclin. pr. Taste, pure bitter. E.s. aq., eth.; v.s. alc., chlf., bz. — $B.2HCl.H_2O$, monoclin. cryst. fr. aq., losing aq. at 100° , not melted at 200° . — $B.2HAuCl_4.H_2O$, \parallel lust. yel. ndl.
796	114.5-5	1',1'-Diaminotoluene, $(NH_2)_2CH.Ph$. — Dist. unded. Cryst. fr. dil. alc. I. aq.; v.s. alc. — $B.HCl$, pr. fr. aq., m.p. $223.5-4.5^{\circ}$.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
797	115-0.5 u.c.	<p>† Atropine, Daturine, i-Tropine r-Tropate, $C_{17}H_{21}O_3N$. — [Mydriatic alkaloid fr. Atropa belladonna, Datura stramonium, etc. Extracted by eth. or chlf. ndl. on spontaneous evapn. c. sat. sol. in 66% alc. — S. at 25° in: 450 pt. aq.; 1.46 pt. alc.; 16.6 pt. eth.; 1.56 pt. chlf. S. at 80° in 86.7 pt. aq.; in 0.9 pt. alc. at 60°. V.s. amyl alc.; v.d.s. pet.-eth., CS. — Odorless. Taste (v. toxic!!), disagreeable, bitter, somewhat metallic, persistent.—Opt. i. when pure. — $k \cdot 10^7 > 1$.</p> <p>② (1) Physiological T. — Prepare a sol. of $B_2H_4SO_4$ by disg. abt. 0.003 g. B in abt. 3 cc. 0.01 normal H_2SO_4, taking care to have clear sol. exactly neutral to litmus! One drop of this sol. brought into conjunctival sac by medicine dropper gives marked dilation of pupil that persists several hours. (According to Fedderson, Diss., Berlin, 1888, p. 88, the minimum quantity required to produce dilation in healthy pupil of adult is 0.0002 mg.; but half as much suffices w. 42% of subjects.) — (2) Vitali's T. — [Z. anal. chem., 20 (1887), 563.] — Dis. abt. 0.5 mg. in 3 drops conc. or fuming HNO_3. Evap. to dryness on water-bath. Moisten c. residue w. 3 drops sol. of 1 g. KOH in 5 cc. 95% alc. A VR color appears, & almost immediately, or within 10 min. (according to concentration & other conditions), changes to R, disappearing after abt. 45 min. to 1 hr! Even 0.001 mg. may give colorations in this test although the hue as above specified will not then be clearly distinguishable. — (3) Gulielmo's T. — Heat 0.01 g. in small dry t.t. until white vapors appear. An aromatic odor is produced. Add 1 cc. conc. H_2SO_4 & heat again until mixt. begins to turn brown. Then add at once, cautiously, without cooling, 2 cc. aq. An intense sweetish odor recalling honey & orange blossoms will be noticed. This odor is most pronounced if observed after the mixture has been allowed to stand until entirely cold. It changes to odor of bitter almonds when a v. small cryst. of $KMnO_4$ is added. — (4) Br + HBr T. — Add ½ cc. sat. sol. Br in aq. HBr to 0.001 g. B in 10 cc. aq. A YO amorph. ppt., i. ac. ac., d.s. large x.s. mineral ac., forms. In time ppl. becomes cryst. w. charac. appearance under microscope (magnification 75 diameters. Cf. cut in Wormley.) Ppt. is still distinct w. as little as 0.001 g. in 20 cc. — (5) Gerrard's T. — Add 2 cc. 1% $HgCl_2$ sol. in 50% alc. to 0.005 g. B in t.t. & heat gently. A Y ppt., mercuric oxide, appears at once, changing w. time to OS1-YOS1.</p> <p>③ (a) Picrate. — Add 3 cc. sat. aq. picric ac. sol. to sol. 0.05 g. B in aq. + 3 drops HCl (sp. gr. 1.12). Shake. Filter curdy ppt. Wash w. 1 cc. aq. Cryst. fr. 7 cc. dil. acetone (1:5) by disg. h. & allowing to cool slowly. Filter. Wash w. 2 cc. solvent. Dry 15 min. at 100°. Picrate is obtained in lust. Y-GY scales, m.p. 174.5-5° u.c. w. good yield. — (b) $B_2H_4PtCl_6$. — Ppt. 0.2 g. B disd. in 1 cc. HCl (sp. gr. 1.12) + 1 cc. aq. by adding calculated quantity conc. chloroplatinic ac. sol., & allow to stand over night. Recryst. the prismatic O cryst. fr. 4 cc. h. aq. Cool. Shake. Filter. Wash w. 2 cc. c. aq. Dry at 100°. Product melts at 206-8° u.c. & gives calculated quantity Pt on ignition. — (c) B_2HAuCl_6. — Proceed as for chloroplatinate, substituting 3 cc. $HAuCl_6$ sol., containing 0.1359 g. Au for chloroplatinic ac. Recryst. ppt. fr. 15 cc. boiling aq. Cryst. (Cf. plate XIII, Fig. 2, Wormley); dried at 80° melts at 195-6° u.c. & gives calculated % Au on ignition.</p>
798	115	Anhaline, $C_{10}H_{17}ON$. — [Fr. Mexican "mescal buttons," Anhalonium fissuratum, which produces an intoxication accompanied by dilation of pupil.] — Pr. d.s. c. aq., more s. h.; v.s. alc., eth., chlf., pet.-eth. Becomes brownish in crystg. — ② One drop HNO_3 added to colorless sol. B in conc. H_2SO_4 gives green color. — Most alkaloid reagt. give amorph. ppt. s. in mineral acids. — $B_2H_4SO_4 \cdot 2H_2O$, lust. tbl. v.s. aq.; d.s. c. alc., m.p. 197°.
799	115	d,l-Laudanosine, N-Methyltetrahydropapaverine, $C_{17}H_{21}O_3N$. — Lust. ndl. fr. alc. or pet.-eth. D.s. h. aq.; e.s. h. alc., bz., acetic eth.; v.s. chlf.; d.s. pet.-eth.; i. alk. Alc. sol. strongly alk. w. bitter taste. — ② Gives color reactions w. alkaloid reagt. as described for No. 2.714. — Picrate, † yel. tbl. fr. alc., m.p. 175°. — Chloroaurate, fine red-yel. tbl. fr. dil. alc., m.p. 163°.
800	115.5	Galipine, $C_{20}H_{21}O_3N$. — [Fr. Angustura bark.] — Silky ndl. fr. pet.-eth. Tert. base.* — B_2HAuCl_6 , m.p. 174-5°.
801	115-6	Kyanpropine, $C_{12}H_{21}N_3$. — [Fr. action of Na on propyl cyanide.] — Pr. s. in 1572 pt. aq. at 23°. — Aq. sol. weakly alk.

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No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
302	115-6	Pentamethylleucoaniline, $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{CH.C}_6\text{H}_4\text{NHMe}$. — Cryst. fr. dil. alc. — W. MeI gives B.3MeI, \ddagger m.p. 185°.
—	116	ms-Ethylacridine. — Cf. No. 2.3148. (Probably yellowish.)
304	116u.c.	α -Naphthylhydrazine, $\text{C}_{10}\text{H}_8\text{NH.NH}_2$. — Leafy cryst. fr. h. aq. V.d.s. c. aq.; v.s. h. alc., chlf., bz. — B.p. 203° (20 mm.). — \textcircled{D} Prepare corresponding hydrazone of benzaldehyde, m.p. 144-5°. (Cf. Ber. 33, 751.)
304-1	116(r.h.)	Mesitylhydroxylamine, $\text{Me}_3\text{C}_6\text{H}_3\text{NH(OH)}$. — \textcircled{D} Reduces Tollen's reag. (T. 2.30)! Dec. in air. — Ndl. S. h. aq., lgr. — \textcircled{D} To sol. of 1.5 g. in 50 cc. bz., add 1.2 g. phenyl isocyanate. A ppt. of interlacing ndl. of phenyl-mesityloxyurea, m.p. 116°, appears within a few min.
305	116	3,6-Diaminoxylene(1,2), $(\text{NH}_2)_2\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{D} Gives green color w. FeCl_3 in neut. sol.; or dark green w. NaNO_2 , in ac. ac. sol., becoming yel. w. pungent quinone odor on warming. — Faintly yellowish ndl. fr. bz. V.s. aq., alc.; d.s. eth.
306	116	† Furfurine, $\text{C}_{11}\text{H}_{12}\text{O}_2\text{N}_2$. — [Product of alk. on furfuramide.] — Ndl. fr. aq. S. in 4800 pt. aq. at 8°, or 135 pt. at 100°; v.s. alc., eth. — B.HNO_3 , pr. fr. alc., m.p. 154°.
307	116-7	† Hydrastinine, $\text{C}_{11}\text{H}_{12}\text{O}_2\text{N}$. — [Oxidation product fr. alkaloid hydrastine.] — Broad colorless ndl. fr. lgr. becoming yel. on keeping. Commercial preparations if yellow may be decolorized by repeated boneblacking & recrystn. fr. bz. D.s. aq.; v.s. alc., eth., chlf. Aq. sol. tastes bitter. Odorless. — \textcircled{D} Very dil. B.HCl sol. shows blue fluor. (seen during titration)! Sol. in abs. eth. colorless; but yel. & fluor. in aq. or alc. — Gives Ag ppt. in T. 2.30 after 1 min. shaking. — Gives the alkaloid color reactions designated (1)-(7) inclusive in description of hydrastine, No. 2.858. — B.HCl , m.p. 212°. — $\text{C}_{11}\text{H}_{12}\text{O}_2\text{N}(\text{CO.Ph})$, (prepared by Schotten-Baumann react., T. 2.27), cryst. fr. dil. alc., m.p. 98-9°.
308	117	Diphenetidine, o-Diaminodiphenetole, $\text{NH}_2\text{C}_6\text{H}_4(\text{OEt})\text{C}_6\text{H}_4(\text{OEt})(\text{NH}_2)$. — \textcircled{D} Should give N in T. 2.4. — Aq. sol. of salts colored red by FeCl_3 . — Reduces Ag & Au salts. — Pptd. by ammon. fr. sol. of salts in ndl. Dist. undec. I. c. aq.; d.s. h. aq.; e.s. alc., eth., chlf. — B.2HCl , cryst. mod. s. aq.; d.s. alc.
309	117.5	1,7-Diaminonaphthalene, $(\text{NH}_2)_2\text{C}_10\text{H}_6$. — Prim. amine.* Ndl. fr. aq.; lft. fr. bz. E.s. alc., bz.; less s. aq.; v.d.s. eth., lgr. — Diacetyl deriv., \ddagger rhombohedra fr. alc., m.p. 213°.
310	117-8	2,4,6-Triaminomesitylene, $(\text{NH}_2)_3\text{C}_6\text{Me}_3$. — Prim. amine.* Mic. ndl. fr. xylene. D.s. c. aq., eth.; alm. i. bz., lgr.; e.s. h. xylene.
311	117u.c.	† Holocaine (base), Ethenyl-p,p'-diethoxy-ab-diphenylamidine, $\text{EtO.C}_6\text{H}_4\text{NH-CMe:N.C}_6\text{H}_4\text{OEt}$. — [Holocaine is B.HCl ; is used as local anesthetic on eye but not as mydriatic.] — Long silky ndl. on cooling h. sol. in 50% alc. 100 pt. aq. at 15° dis. 2.218 pt. E.s. alc., eth., bz. Odorless. Taste, faintly bitter.
		\textcircled{D} (1) Froehde's Reagt. [T. 2.2-(f)] gives GBT1 color changing in 1 to 3 min. to BS1. — (2) Bleaching Powder T. — Add 3 drops sat. aq. sol. bleaching powd. to sol. of 0.001 g. B in 1 cc. 50% alc. OT2 color develops. — (3) Iodic Ac. T. — Dis. 0.001 g. B in 3 drops conc. H_2SO_4 & add small cryst. iodic anhydride or KIO_3 . An OS1-OYS1 color, not changed on warming, develops. — (4) KMnO_4 , T. — 5 or 6 drops of holocaine sol. produced in generic titration discharge color of 2-3 drops 1% KMnO_4 sol. — \textcircled{D} Evaporate 5 cc. of aq. sol. fr. generic titration test to vol. of 0.5 cc. B.HCl separates in pr. or plates of m.p. 190-2° u.c. after drying 15 min. at 100°.
312	117.5	Pseudoephedrine, $\text{C}_{10}\text{H}_{12}\text{ON}$. — [Fr. Ephedra vulgaris L.] — Faint, agreeable odor. Cryst. d.s. c., more s. h. aq.; s. alc., eth. — $[\alpha]_D^{20} = +51.24^\circ$ (in abt. 0.6% alc. sol.). — B.HCl , ndl. fr. eth.-alc., v.s. aq., alc.; m.p. 176°. — Picrate, f. oily.
313	117.8	Hordenine, p-Hydroxyphenyldimethylmethylethylamine, $\text{HO.C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{NMe}_2$. — [From malt.] — \textcircled{D} Prim. amine.* Neutralized sol. colored violet by FeCl_3 . — Colorless, nearly tasteless pr. D.s. aq.; less s. lgr.; e.s. alc., chlf., eth.; less s. bz. — B.p. 173-4° (11 mm.). — For derivatives w. nitro comp. & microchemistry, Cf. Z. anal. Chem., 49, 340. — B.MeI^{**} (fr. $\text{B} + \text{MeI}$ in cold), d.s. c. aq., m.p. 230-1°.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
814	118-9	2,5-(α)-Dimethylpiperazine, $[NH.CHMe.CH_2.NH.CHMe.CH_2.NH]^2$. — B.p. 162° (th.i.). — \textcircled{P} Sec. amine.* — Monoclin. pr. fr. chlf. Sbl. at room temp. V.s. aq.; alc.; d.s. eth. Reacts strongly alk. — \textcircled{D} Convert into dinitroso deriv.,** as described under Lycetol, No. 2.440.
815	118	4-Benzylisoquinoline, $Ph.CH_2.C_6H_3N$. — \textcircled{D} Tert. amine.* — Cryst. fr. lgr. or aq. D. vol. w. st. — $\textcircled{B}.\text{Pk}, \textcircled{t}$ ndl. fr. h. alk.; r.d.s. alc.; m.p. 190-1°.
816	119	Leucodiphenylene-green, Tetramethyldiamino-diphenylamine, $NH.(C_6H_4-NMe_2)_2$. — Large quadratic tbl. fr. CS. Treatment of dil. ac. ac. sol. w. trace PbO, prob. gives green color.
817	119-20	Pseudoatropine, Atrolaktyltropeine, $C_{17}H_{24}O_3N$. — Lust. ndl. fr. aq. V.d.s. c. aq.; v.e.s. alc. — Dilates the pupil like atropine (No. 2.797). — $\textcircled{B}.\text{HAuCl}_4, \textcircled{ }$ yel. ndl. fr. aq., m.p. 112-4°. (Cf. No. 2.797-(c).)
818	119-20	Glaucine, $C_{11}H_{22}O_4N$. — [Fr. Glaucium luteum.] — Slightly yellowish pr. or tbl. Tasteless; salts sl. bitter. D.s. c., more s. h. aq.; v.s. alc., chlf. $[\alpha]_D = +113.3^\circ$ (in alc., c = 5.0449). — \textcircled{D} Conc. H_2SO_4 gives colorless sol. (after first showing transient yel. coloration); but after some hours sol. becomes sky-blue, or becomes rapidly dark blue or violet on heating at 100°. — $\textcircled{B}.\text{MeI}, \textcircled{t}$ (formed fr. x.s. MeI + 1 g. B in 4 cc. MeOH at ordinary temperature), cryst. mod. s. h. aq., alc., m.p. 216° d.
819	119.6 u.c.	Carpine, $C_{11}H_{22}O_4N$. — [Fr. leaves of Carica Papaya L. (papaw).] — Sec. base.* Alc. sol. alk. to litmus. — Silky ndl. fr. alc. I. aq.; s. alc.; v.s. eth., chlf., bz. — \textcircled{P} Taste v. bitter & distinct at dilution 1:100,000! — $[\alpha]_D^{20} = +21.55^\circ$ (in abs. alc., p = 9.236). — $\textcircled{B}.\text{H}_2\text{PtCl}_6, \textcircled{g}$ ocher-yel. floc. ppt.; i. aq., alc.
820	120	1,4-Diaminonaphthalene, $C_{10}H_8(NH_2)_2$. — Unstable, especially if moist. — Diacetyl deriv., $\textcircled{ }$ cryst. fr. gl. ac. ac., m.p. 303-4°.
821	120	1,2,4-Trimethylphenylhydrazine(5), $Me_3C_6H_3.NH.NH_2$. — \textcircled{D} Prob. reduces Tollen's reagt. & Fehling's sol. easily. — Ndl. fr. eth. I. aq., e.s. alc., eth., chlf.; s. lgr., bz.
822	d.w.m. 120	† Delphinine, $C_{10}H_{16}O_3N$ or $C_{10}H_{18}O_3N$ (?). — [Asphyxiating poison fr. seeds of Delphinium Staphisagria L.] — \textcircled{D} Rubbed together w. 1-2 vol. malic ac. & treated w. conc. H_2SO_4 becomes orange, after some hours dark rose red, & finally dingy cobalt-blue. — Rhomb. cryst. S. in 50,000 pt. c. aq.; 20.8 pt. 98% alc.; 11.1 pt. eth.; 15.8 pt. chlf. Opt. i.
823	120.6	† Conhydrine, d- α -Ethylpiperidylalkin, $C_8H_9NH.CH(OH).CH_2.CH_3$. — [Toxic alkaloid accompanying coniine in seeds of Conium maculatum.] — B.p. 226°. — \textcircled{D} Odor like mouse urine & coniine. — Mod. s. aq.; e.s. alc., eth. — $\textcircled{B}.\text{AuCl}_4$, rhomb. pr., m.p. 133-4°. Its sol. gives ppt. with (T. 2.3): Mayer's reagt., amorph. (1:100); phosphotungstic ac., cryst. (1:1000); I in KI, amorph. (1:1000); picric ac., $H_2\text{PtCl}_6$, or HAuCl_4 , no ppt. (1:100). — Benzoyl deriv., cryst., m.p. 132° (Ber. 15, 2315).
824	121 u.c.	† Furfamide, $(C_5H_8O)_2N_2$. — Flat, colorless ndl., notched at ends, or rosettes fr. h. dil. alc. Rather unstable, usually being yellowish & having slight furfural odor fr. decomposition. I. c. aq.; e.s. alc., eth. When brought on tongue nearly tasteless at first. — \textcircled{D} T. 2.2-(a), w. conc. H_2SO_4 , gives VR coloration, becoming brown on stirring. — Add 6 drops dil. H_2SO_4 (sp. gr. 1.4) to suspension of 0.1 g. B in 4 cc. aq. & heat just to boiling. Apply test 1.115-(2) for furfural by holding strip of aniline acetate in vapors. The paper will become intensely red.
825	121	† Codamine, $C_{10}H_{22}O_4N$. — [An opium alkaloid.] — 6-sided pr. fr. eth. S. h. aq.; e.s. eth., bz., chlf.; v.s. alc.; s. in KOH when freshly pptd. — \textcircled{D} S. conc. HNO_3 w. dark green color. Colored dark green by $FeCl_3$. — $\textcircled{B}.\text{H}_2\text{PtCl}_6, 2H_2O, \textcircled{g}$ yel. amorph. ppt., v.d.s. aq.
826	121	† 1-Cinnamylcocaine, Cinnamylecgoninemethyleneester, $C_{19}H_{22}O_4N$. — [Alkaloid fr. coca leaves.] — Ndl. fr. h. bz. + lgr. $[\alpha]_D^{16} = -4.7^\circ$ (in chlf. sol., p = 10). — $\textcircled{B}.\text{HCl}.2H_2O$, m.p. 176° (after dehydration). — $\textcircled{B}.\text{H}_2\text{PtCl}_6, \textcircled{g}$ cryst. ppt., m.p. 217°.

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. \ddagger = T. 2.14. $\ddot{\delta}$ = T. 2.13. $\textcircled{||}$ = T. 2.1. \textcircled{q} = T. 2.36. ** = T. 2.37. $\textcircled{\ddagger}$ = T. 2.38.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
827	122	Phenylcarbamic-acid-hydrazide, Ph.NH.CO.NH.NH_2 . — \textcircled{P} Reduces Tollen's reagt. (T. 2.30) or Fehling's sol. on warming. — Ndl. fr. h. bz.; d.s. h. aq.; e.s. alc., chlf., h. bz., dil. ac. or alk.; i. eth. — \textcircled{D} Convert into corresponding acetonecarbazone by dissolving in aq. acidulated w. HCl, adding sodium acetate & acetone, & shaking vigorously. Product, ndl., d.s. aq., m.p. 155-6 $^{\circ}$.
828	122	† Conessine, Wrightine, $\text{C}_{24}\text{H}_{40}\text{N}_2$. — [Toxic alkaloid fr. Wrightia antidysenterica & Holarrhena africana.] — Hair-like ndl. by spontan. evapn. of sol. in 66% alc. Sbl. w. part. decn. Taste "sharp & harsh." — \textcircled{P} Recryst. preparation fr. Schuchardt (m.p. 121-2 $^{\circ}$ u.c.) gave following colors in T. 2.2: (a) w. conc. H_2SO_4 , YT1, changing in 15-20 min. to RT3 Bkn.; (b) w. conc. HNO_3 , colorless; (f) w. Froehde's reagt., GY gradually darkening to permanent GYS2; (g) w. Mandellin's reagt., GYS1 changing to YGS1 in 15 min. & BG-BGS1 in 30 min. — $\text{B.H}_2\text{PtCl}_6 \cdot \frac{1}{2}\text{H}_2\text{O}$, $\frac{1}{2}$ ndl. fr. alc. + HCl. — B.2MeI, \ddagger tbd. fr. aq.
—	122-3	2,9-Dimethylacridine. — Cf. No. 2.3176. (Yellowish.)
830	123	m-Aminophenol, $\text{NH}_2\text{C}_6\text{H}_4\text{OH}$. — [An intermediate in dyestuff manufacture.] Pr. fr. toluene. S. h. aq.; e.s. eth., alc., amyl alc.; d.s. bz.; v.d.s. lgr. — B.HCl , pr., m.p. 229 $^{\circ}$. — W. alc. sol. picryl chloride (Ber. 33, 433) gives picramide (Cf. T. 2.22), deep red ndl., m.p. 203-4 $^{\circ}$.
831	123	1'-Amino-1,3,4,5-Tetramethylbenzene, $\text{NH}_2\text{CH}_3\text{C}_6\text{H}_3\text{Me}_3$. — \textcircled{P} Should give N in T. 2.4. — B.HCl , m.p. 270 $^{\circ}$, d.s. c. aq., alc. — B.HAuCl_4 , \parallel garnet red pr., m.p. 162-5 $^{\circ}$ d. — B.Pk , \ddagger ndl. fr. dil. alc., m.p. 239.5 $^{\circ}$ d.
832	123; 121	Conquinamine, $\text{C}_{10}\text{H}_{14}\text{O}_2\text{N}_2$. — [In cinchona barks.] — Triclinic cryst. fr. 80% alc. V.d.s. aq.; e.s. alc., chlf. 100 pt. bz. at 18 $^{\circ}$ dis. 24.4 pt.; 100 pt. eth. at 15 $^{\circ}$ dis. 13.5 pt. $[\alpha]_D^{15} = +204.6^{\circ}$ (in 97% alc., p = 2). Alc. sol. reacts strongly alk.'
833	124-5	β -Naphthylhydrazine, $\text{C}_{10}\text{H}_7\text{NH.NH}_2$. — \textcircled{P} Prob. reduces Tollen's reagt. (T. 2.30). — Lft. E.s. h. alc., chlf., bz.; r.d.s. eth. — \textcircled{D} Prepare the picramide (Cf. T. 2.22) fr. h. alc. picryl chloride sol. (J. prakt. Chem. [2], 43, 179); red pr., d. 175 $^{\circ}$.
834	123-5	Cinchene, $\text{C}_{10}\text{H}_{20}\text{N}_2$. — Lft. fr. lgr. Dextrorotatory. Adds Br_2 .
835	125-6	4,5-Diaminoxylene(1,2), $(\text{NH}_2)_2\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} Gives blue-green color w. FeCl_3 . NaNO_2 added to ac. ac. sol. gives yel. sol. & then white ppt. — Lft. mod. s. aq.; v.s. alc., lgr.
836	125	Isobenzidine, $\text{C}_{12}\text{H}_{14}\text{N}_2$. — Iridescent lft. fr. aq. — \textcircled{P} Cl-aq. produces green color followed by brown ppt in aq. sol. — B.2HCl , lft., r.d.s. — Sulphate, d.s.
837	125	Benzoyl-m-phenylenediamine, $\text{NH}_2\text{C}_6\text{H}_3\text{NH.CO.Ph}$. — Cryst.
838	124-8	6-Aminocresol(2), $\text{NH}_2\text{C}_6\text{H}_5\text{Me(OH)}$. — Ndl. d.s. c. aq., eth.
839	126.5	Phenylglycinehydrazide, $\text{Ph.NH.CH}_2\text{CO.NH.NH}_2$. — Lft. fr. alc.; e.s. h. aq. or alc.; alm. i. eth.
840	126.5-7 u.c. 128.2-0.7 c.	† Benzidine, 4,4'-Diaminodiphenyl, $\text{NH}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NH}_2$. — [Important intermediate in dye mfg.] — Lust. pearly scales fr. h. aq. Ndl. fr. bz. Becomes colored in air. S. in 2447 pt. aq. at 12 $^{\circ}$, or in 106.5 pt. at 100 $^{\circ}$. — B.p. 400-1 $^{\circ}$ (740 mm.). \textcircled{P} (1) To 5 cc. c. sat. aq. sol., add 2 drops conc. H_2SO_4 . Within a few sec. sol. becomes turbid fr. pptn. of the v. i. $\text{B.H}_2\text{SO}_4$. — (2) To 3 cc. c. sat. aq. sol., add 5 cc. v. dil. Br sol. made by mixing 1 drop sat. Br-aq. w. 20 cc. aq. An intense green-blue (GB) color develops at once. Further addition of 10-20 cc. Br sol. changes color to green (G) and then to yel.-orange. [Tests (1) & (2) are also given by No. 2.849.] \textcircled{D} Add 3 drops benzaldehyde to sol. 0.05 g. \bar{B} in 4 cc. alc. Filter heavy ppt. & wash w. 2 cc. alc. Recryst. 3 times fr. h. bz., using successively 4, 3, & 2 cc. of solvent. Dry 15 min. at 100 $^{\circ}$. The product, dibenzylidenebenzidine, is obtained in lust. YT2 scales, m.p. 232.5-0.8 $^{\circ}$ u.c. (238.9-4 $^{\circ}$ c.).
841	126	(α)-Bisethyldeneaniline, $\text{Ph:NCH}_2\text{CH}_2\text{CHMe.NHPh}$. — B.p. 300 $^{\circ}$. — Rhomb. cryst. fr. eth. I. aq., lgr.; e.s. eth., chlf., bz. — Dinitroso deriv., \ddagger m.p. 161 $^{\circ}$ (Ber. 27, 2977). — Benzoyl deriv., silky lft. (by shaking w. benzoyl chloride & bz.), alm. i. h. alc., bz., m.p. 218 $^{\circ}$.
842	126-7	Py-2,4-Dimethyl- β -naphthoquinoline, $\text{Me}_2\text{C}_1\text{H}_7\text{N}$. — B.p. a. 300 $^{\circ}$ sl. dec. Ndl. fr. eth. V.d.s. h. aq.; e.s. chlf. — B.Pk , \ddagger yel. ppt., ndl. fr. alc., m.p. 215 $^{\circ}$.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
843	126-7; 125	β -Isocinchonine, $C_{18}H_{20}ON_2$. — Pseudomorphic pr. I. aq., alk.; e.s. alc., eth. $[\alpha]D^{16} = -53.7^{\circ}$ (in abs. alc., $p = 1$).
844	128	β -Naphthyl-m-phenylenediamine, $NH_2.C_8H_4.NH.C_6H_4.H_2$. — Asbestos-like ndl. fr. alc. B.p. abt. 320° (40 mm.). V.s. eth., chlf. — B_2HCl (110°), ndl., m.p. 210° , d.s. h. aq. — B_2Pk , \ddagger ppt., golden lft., m.p. 180° d.
845	128	\dagger 5-(or 8-)Aminoisoquinoline, $C_9H_8N.NH_2$. — Cryst. Sbl. — $B_2H_2PtCl_6$, \ddagger or. ppt.
846	125d.; 132-3	Cotanine, $C_{12}H_{14}O_2N$. — [Oxidation product of narcotine.] — Ndl. D.s. c. aq.; e.s. alc., eth. "Reacts alk." Sbl. Taste, v. bitter. — \oplus Scattered upon Froehde's reag. gives brown-red color changing to green, violet streaks appearing later. — $B_2HCl.H_2O$, a styptic known as "stypticin," yellowish cryst., s. aq., alc. NaOH, but not NH_4OH , ppt. B fr. aq. sol. — B_2Pk , \ddagger yel. ndl. (by ptn. fr. aq. sol. & recrystn. fr. h. aq.), softens at 133° , m.p. 143° .
847	129	1'-Aminoresol(2), Salicylamine, $HO.C_6H_4.CH_2.NH_2$. — Ndl. fr. eth. Sbl. V.s. aq. — $B_2H_2PtCl_6$, \ddagger golden ndl., m.p. 197° d.
848	129	4,4'-Diamino-3-methyldiphenylmethane, $NH_2.C_6H_4.CH_2.C_6H_3Me(NH_2)$. — Lust. lft. fr. alc. E.s. alc., eth., chlf. — B_2HCl , aq. sol. becomes strawberry-red w. $FeCl_3$.
849	129-30u.c.	\dagger o-Tolidine, 4,4'-Diamino-3,3'-dimethylbiphenyl, $(NH_2)Me.C_6H_3.C_6H_3Me(NH_2)$. — [Important intermediate in dye manufacture.] — Pearly scales fr. boiling aq. Becomes colored in air. D.s. aq.; e.s. alc., eth. — \oplus Gives the two preliminary tests described for benzidine, No. 2.840. — \ominus Dis. 0.10 g. B in 6 cc. alc. Add 10 drops benzaldehyde. Ptn. of fine light-yel. ndl. begins within abt. 10 min. Let stand 10 min. longer w. occasional shaking. Recryst. the heavy ppt. fr. 6 cc. alc. + 1 cc. bz. Recryst. again fr. 3 cc. of same solvent. The product, dibenzylideneditolidine, is obtained in YT \ddagger ndl., m.p. $150-0.5^{\circ}$ u.c. ($152.6-3.1^{\circ}$ c.).
850	129c.	2-Aminoquinoline, $C_9H_8N.NH_2$. — Lft. fr. aq. Alm. i. c. aq.; e.s. alc., eth., chlf., h. aq.; d.s. bz., lgr. — $B_2H_2PtCl_6.2H_2O$, \ddagger ppt., lust. or.-yel. ndl. fr. conc. HCl. — B_2MeI , \ddagger (fr. components at 100°), pr. e.s. h. aq., alc., m.p. 247° .
851	130	Dimethylamino-p-phenylenediamine, $Me_2N.C_6H_4.NH_2Ph$. — \oplus Salts colored blue by $FeCl_3$. — Ndl. fr. lgr. E.s. dil. HCl. — Nitroso deriv., \ddagger yel. ndl. fr. alc., m.p. 116° d.
852	130	2,4-Diaminodiphenylamine, $Ph.NH.C_6H_4(NH_2)_2$. — Ndl.
853	131u.c.	\dagger Amarine, $C_{11}H_{12}N_2$. — B_2H_2O , colorless cryst. fr. alc., m.p. 101° (air dried); m.p. 131° u.c. when anhydrous (dried at 115°). Powder, at first tasteless (poisonous) & then faintly bitter (No. 1 in scale, T. 2.29). E.s. alc., eth.; alm. i. c. aq. Gives no colorations in T. 2.2-(a),(b),(d),(e). — \ominus Dis. 0.10 g. B in 3 cc. alc. + 4 drops HCl (sp. gr. 1.2). Add 1 cc. c. sat. aq. $NaNO_2$, sol. Heat to boiling. Cool & shake very persistently! Filter the YT \ddagger ppt. of lust. scales & wash w. 1 cc. dil. alc. (3 vol. alc. : 1 vol. aq.). Recryst. fr. 5 cc. of same dil. alc. Filter, & wash w. 1 cc. dil. alc. Dry at 100° for 15 min. The product, nitrosoamarine, forms nearly colorless scales, decg. quite sharply at 152.9° u.c. (154.7° c.) without entirely melting. Recrystd. by spontaneous evapn. of conc. alc. sol. gives beautiful diamond shaped plates.
854	130-1	Tetraphenylguanidine, $HN:C(NPh)_4$. — Rhomb. pyramids fr. lgr. I. aq.; e.s. alc., eth. Salts d.s. aq. — $B_2H_2PtCl_6$, pale-yel. ppt.
855	131	uns-Triphenylguanidine, $HN:C(Ph.NH)(Ph.NH)_2$. — Tbl. e.s. alc., eth. Heated w. conc. H_2SO_4 (prob. containing trace nitrous ac.) gives violet sol.
856	131.5	o-Dianisidine, 4,4'-Diamino-3,3'-dimethoxybiphenyl, $(NH_2)(MeO).C_6H_3.C_6H_3(MeO)(NH_2)$. — [Important intermediate in dye manufacture.] — Lft. Becomes violet in air. E.s. alc., eth., bz.; s. h. aq. — Diacetyl deriv., \ddagger (by boiling w. acetic anhydride), i. aq., eth., alc., bz.; m.p. 231° . — $B_2H_2SO_4$, ndl. fr. aq. Aq. at 100° dis. 4.7%. — $B_2H_2CrO_4$, brown cryst. ppt. i. aq.
857	131-2	Morphinebenzylether, $C_{17}H_{18}O_2N(OC_6H_5)_2$. — [Dangerous cardiac poison.] — Pr. or tbl. v.d.s. aq.; e.s. alc., eth., bz., chlf.; i. KOH; v.d.s. NH_4OH . Taste of salts, bitter & burning. — \oplus (1) Conc. H_2SO_4 , dis. w. pale red-yel. color,

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(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		changing to dark red on warming. — (2) Marquis' reagt. gives fine carmine-red, gradually becoming slightly violet. — (3) Froehde's reagt. gives red violet sol. at once, soon changing to brown-green. — $\bar{B} \cdot HCl$ ("Peronin"), colorless ndl. s. aq.
858	132 u.c.	<p>† Hydrastine, $C_{11}H_{21}O_6N$. — [Alkaloid fr. root of <i>Hydrastis canadensis</i>.] — Large, clear, lust., rhomb. pr. on spontan. evapn. of alc. sol. Alm. i. c. aq.; s. i. 4000 pt. aq. at 80°; s. at 25° in 135 pt. alc., 124 pt. eth., or 2 pt. chlf.; s. at 60° in 17 pt. alc.; e.s. bz. — $[\alpha]D^{25} = -67.8^{\circ}$ (for 1.2759 g. in 50 cc. chlf. — k.10⁷ = abt. 1. — Odorless. Taste slightly bitter.</p> <p>① (1) T. 2.2-(b) w. HNO_3 gives $\leftarrow OY$ sol. Diln. w. 30 cc. aq. then gives sol. w. strong blue fluor.! — (2) Dis. 0.001 g. \bar{B} in 1 cc. dil. H_2SO_4 (sp. gr. 1.175) & add 1 drop 1% $KMnO_4$ sol. Color is discharged. Diln. to 25 cc. now gives sol. w. marked blue fluor.! — (3) T. 2.2-(a) w. conc. H_2SO_4 gives $YT1$ sol., changing to RV Bkn. [D] after heating for some sec. on water-bath. — (4) T. 2.2-(f) w. Froehde's reagt. gives $YS2$, changing almost immediately on stirring to BG Bkn. [D]. — (5) T. 2.2-(g) w. Mandellin's reagt. gives $OR-RO$, gradually fading. — (6) $[H_2SO_4 + MnO_2 \cdot T]$ — Scatter a few particles powd. MnO_2 on sol. 0.001 g. \bar{B} in 2 drops conc. H_2SO_4. An $OS1$ color appears, changing to $ORS1$ within 10 min., & disappearing within 4–5 hr. — (7) Dis. 0.001 g. \bar{B} in 5 drops conc. HNO_3 & heat to boiling. Evap. to dryness on water-bath. Add 2 drops 25% KOH sol. in alc. Again evap. to dryness. The $OYS1$ residue becomes $VRS1$ on adding 2 drops conc. H_2SO_4.</p> <p>② Dis. \bar{B} 0.05 g. in 3 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 4 cc. sat. aq. sol. picric ac. Filter off the floc. ppt. Wash w. 2 cc. aq. Recrys. fr. 1/4 cc. boiling alc., allowing to stand for some hr. Filter. Wash w. 1 cc. alc. Dry at 100° on porous tile for 15 min. The picrate formed, $\bar{B} \cdot Pk \cdot H_2O$, is obtained in fine lust. Y → ndl., m.p. 165–70°, decomposing to dark brown liquid.</p>
858-I	132.5	1- Canadiane , Tetrahydroberberine, $C_{19}H_{21}O_6N$. — [Fr. root of <i>Hydrastis canadensis</i> .] — Silky ndl. I. aq.; mod. s. alc.; v.s. eth. ("Reacts neutral.") $[\alpha]D^{20} = -298.05^{\circ}$ (0.2517 g. in 24.9 cc. chlf.). — Contains 2 methoxyl groups. Oxidn. w. alc. I. sol. gives berberine. Colors vanadium sulphuric ac. reagt. olive-green & then black-brown. — $\bar{B} \cdot H_2SO_4$, monoclin. tbl., mod. s. c. aq.
859	132	4,4'- Diaminodiphenylethane , $NH_2 \cdot C_6H_4 \cdot CH_2 \cdot CH_2 \cdot C_6H_4 \cdot NH_2$. — Lust. scales. Sbl. Alm. i. c. aq.; more s. h. aq.; v.s. alc. — $\bar{B} \cdot 2HCl$, cryst. e.s. aq., alc. — $\bar{B} \cdot H_2SO_4$, cryst. powd. d.s. aq.
860	132	Tetramethyl-4,4'- diaminotriphenylcarbinol , Malachite-green-leucohydrate, $[NMe_2 \cdot C_6H_4]_2 \cdot CPh(OH)$. — Colorless cube-like cryst. fr. bz. Alm. i. aq.; e.s. alc., giving green sol.; e.s. h. bz.; d.s. eth. when in cryst. — ② Colorless sol. in dil. acids becomes blue-green on standing or warming.
861	133–4	5-Amino-1,3- xylenol(4) , $NH_2 \cdot C_6HMe_2(OH)$. — Lft. fr. alc. Alm. i. aq., bz.; e.s. alc., eth.
862	133	2,4,4'- Triaminodiphenylmethane , $(NH_2)_2 \cdot C_6H_3 \cdot CH_2 \cdot C_6H_4 \cdot NH_2$. — [Pat.]
863	133.5	Acetylcodeine, $MeO \cdot (C_{17}H_{17}NO) \cdot O \cdot C_6H_5O$. — Pr. fr. eth. D.s. h. aq.; s. alc., eth., chlf., bz. — $\bar{B} \cdot H_2PtCl_6$, § amorph. yel. ppt.
864	134	2-Aminobenzidine, $C_6H_5(NH_2)_2 \cdot C_6H_4(NH_2)_2$. — Ndl. — $\bar{B} \cdot 3HCl$, cryst.
865	134	2,5-Diamino-4-anilinotoluene, $Me \cdot C_6H_4(NH_2)_2 \cdot (Ph \cdot NH)$. — Lft.
866	134–5	2-Hydrazinoquinoline, α -Quinolylhydrazine, $NC_6H_4 \cdot NH \cdot NH_2$. — Cryst. fr. aq., bz. E.s. alc.; d.s. eth., lgr. — $\bar{B} \cdot Pk$, † v.d.s., m.p. 187° d. — $\bar{B} \cdot H_2PtCl_6$, § cryst. ppt., m.p. 170° d.
867	134.5 u.c.	† Corydaline , $C_{11}H_{21}O_6N$. — [Alkaloid fr. root of <i>Corydalis cava</i> .] — Colorless pr. fr. alc. tending to become yellowish during cryst. I. aq.; s. alc.; e.s. eth., chlf. Powder tasteless (poisonous). — ② T. 2.2-(b) w. HNO_3 gives permanent $OY-OYS1$ color after thorough mixing. — ③ Rub well together in v. small glass capsule 0.05 g. finely powd. \bar{B} & 4 cc. dil. HNO_3 (1 vol. acid, sp. gr. 1.42 + 20 vol. aq.). Decant after 10 min. Drain residue on porous tile. Dissolve in 3 cc. h. aq. & filter hot. Allow nitrate to cryst. by spontan. evapn. Wash cryst. on tile w. alc. Dry. The product, $\bar{B} \cdot HNO_3$, melts w. vigorous frothing at $198 \pm 5^{\circ}$ u.c.
868	134	Laurotetanine , $C_{10}H_{21}O_6N$. — [Tetanic poison fr. <i>Tetanthera citrata</i> & other E. Indian plants.] — Ndl. fr. eth. V.d.s. aq.; e.s. alc., chlf., acetic eth.;

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		d.s. eth., pet.-eth., bz. — \textcircled{P} S. in conc. H_2SO_4 w. blue color, changing to violet on warming. — Reduces AgNO_3 & Fehling's sol. — Froehde's reagt. gives indigo-blue sol. changing to yel. on adding a drop aq. — Erdmann's reagt. gives a blue & then brown color. — A sec. monacid base.
869	135	3-Aminocresol(4), $\text{NH}_2\text{C}_6\text{H}_3\text{Me}(\text{OH})$. — Scales fr. eth. Sbl. in lft. Alm. i. c. aq.; e.s. alc., eth., chlf.; less s. bz. — \textcircled{P} Aq. sol. of BHCl (but not of free B) colored red by FeCl_3 . [An intermediate in color mfg.]
870	135	Leucoauramine, $[\text{NMe}_2\text{C}_6\text{H}_4]_2\text{CH.NH}_2$. — Cryst. fr. alc. Covered w. HCl becomes greenish & dis. colorless. Alm. i. aq.; r.d.s. alc. — \textcircled{P} Gl. ac. ac. dis. w. intense blue color!
871	135-6d. u.c.	† Chelidone, $\text{C}_{10}\text{H}_{11}\text{O}_4\text{N.H}_2\text{O}$. — [Alkaloid fr. Chelidonium majus L., celandine poppy.] Loses aq. above 100°. Softens fr. 132°. Fuses to pale yel. liquid. Glassy monoclin. tbl. or ndl. I. aq.; s. alc., eth. $[\alpha]_D^{20} = +115.24^{\circ}$ (in 96% alc., $p = 2$). Taste (very slowly developed), bitter & persistent. Odorless. \textcircled{P} (1) 0.001 g. B + 0.5 cc. conc. H_2SO_4 + 1 drop quiacol gives brilliant VRT1-R color! — (2) T. 2.2-(a) w. conc. H_2SO_4 gives Y Bkn [L]. — YOS1, changing to VS1 within 5 min. — (3) T. 2.2-(e) w. Erdmann's reagt. gives GYT1, quickly changing to RV Bkn[M]. — (4) T. 2.2-(f) w. Froehde's reagt. gives BGS1, darkening to BGS2. — (5) T. 2.2-(g) w. Mandelin's reagt. gives GS1, changing within 3 min. to BGS2. — (6) Furfurol react. (Cf. Z. anal. Chem. 24, 165) gives VRT1 within 1 min., then fading to pale pink. — BHCl , cryst. s. in 325 pt. aq. at 18°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yellowish ppt., m.p. 155°. — Benzoyl deriv., $\text{C}_{20}\text{H}_{15}\text{O}_4\text{N}(\text{Ph.CO})$, colorless cryst., m.p. 217°.
872	136	2-Methylglyoxaline, $[\text{CH}:\text{CH.NH.CMe:N}]$. — B.p. 267°. — Ndl. fr. bz. V.s. aq., alc.; less s. c. bz. Pptd. by tannin, picric ac., & salts of heavy metals. — 6 Br added to aq. sol. gives $\text{C}_4\text{H}_4\text{N.Br}_3$, cryst. fr. alc., m.p. 258° d.
873	136.5; 138	2-p-Aminophenylquinoline, $\text{C}_{15}\text{H}_{12}\text{N}_2$. — Felted hair-like ndl. fr. aq. Dist. undec. Alm. i. c. aq.; mod. s. h. aq.; e.s. alc., eth., bz. — BHCl , fine yel. ndl. — Acetyl deriv. (T. 2.1) (by boiling w. ac. anhydride), lft. fr. alc., m.p. 189°.
874	137	Apoconchinine, $\text{C}_{18}\text{H}_{22}\text{O}_2\text{N}_2$. — Amorph. Cryst. w. $2\text{H}_2\text{O}$, lost at 120°. E.s. alc., eth. "Reacts alk." Sol. in conc. H_2SO_4 not fluor. — \textcircled{P} Gives green color in quinine T. w. Cl-aq. & ammon. (No. 2.947). — $[\alpha]_D = +155.3^{\circ}$ (in 97% alc., $p = 2$).
875	139	Benzoylquinine, $\text{C}_{20}\text{H}_{22}\text{O}_2\text{N}_2(\text{Ph.CO})$. — Pr. fr. moist eth. I. aq.; v.s. eth., alc., chlf., bz., lgr.; s. acids, & repptd. by alk.
876	139	4,4'-Diaminotriphenylmethane, $(\text{NH}_2\text{C}_6\text{H}_4)_2\text{CH.Ph}$. — Spheroidal aggregates fr. dry eth. Alm. i. aq.; e.s. alc., eth., chlf., lgr. — $\text{B}\text{C}_6\text{H}_5$, cryst., m.p. 106° w. sl. deecn.
877	140 u.c.	† p-Phenylenediamine, $\text{C}_6\text{H}_4(\text{NH}_2)_2$. — B.p. 267°. — Unstable, soon becoming dark violet or black by oxidation on exposure, especially in aq. sol. Rather difficult to purify. Best crystd. fr. bz., separating fr. sol. in colorless scales on rapid cooling. E.s. aq., alc., eth. \textcircled{P} Dis. exactly 0.0025 g. B in 10 cc. aq. Add 1 drop 10% FeCl_3 sol. Sol. becomes deep green (GS2 by direct transmitted light fr. sky). Add 1 more drop FeCl_3 . The green changes rapidly to brown, & after abt. 1 min. appears RV by same illumination, & much darker after 5 min. Diluted w. equal vol. aq., color, w. white background, becomes V or VB. \textcircled{P} Mix 0.10 g. B w. 8 drops acetic anhydride. Heat to boiling. Boil vigorously w. 3 cc. aq., breaking up lumps w. stirring rod. Cool. Filter. Wash w. 3 cc. aq. Dry on porous tile. Dis. in 2 cc. h. gl. ac. ac. Ppt. w. 2 cc. alc. Filter. Wash w. alc. Dry 15 min. at 100° on tile. The product, diacetyl-p-phenylenediamine, melts at 298.7-9.7° u.c. (309.5-10.5° c.).
878	abt. 140	Aconine, $\text{C}_2\text{H}_4\text{O}_2\text{N}$. — [Fr. decn. ofaconitine & fr. Aconitum Napellus.] Amorph. V.s. aq., alc.; i. abs. eth., lgr. Reduces ammon. AgNO_3 or Fehling's sol. on heating. Taste, v. bitter (poisonous). Dextrorotatory; in acid sol. levorotatory. — Dibenzoyl deriv., ndl. fr. eth., m.p. 265°. (Fr. 1 mol. B + 1 mol. benzoic anhydride in chlf. sol.)

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. ‡ = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
879	141-2	Meteloidine , $C_{12}H_{21}O_4N$. — [Alkaloid fr. <i>Datura Meteloides</i> .] — Ndl. fr. bz. D.s. aq., eth., bz.; e.s. alc., chlf. — B.Pk., \ddagger yel. hexag. tbi. fr. alc., m.p. 177-80 $^{\circ}$, d.s. aq., alc.
880	144.5	2-Aminocresol(4) , $NH_2C_6H_4Me(OH)$. — [An intermediate in color mfg.] Cryst. — B.HCl, e.s. aq., the sol. giving blue floc. ppt. when treated w. ammon. & exposed to air.
881	144	Concusconine , $C_{22}H_{34}O_4N_2H_2O$. — [In bark of <i>Remijia purdieana</i> .] — Monoclin. cryst. After melting & solidifying, remelts at 206-8 $^{\circ}$. — \textcircled{P} Sol. in conc. H_2SO_4 , blue-green; olive-green after warming. Sol. in ac. ac. or HCl green upon addition of conc. HNO_3 . — I. aq.; v.d.s. alc.; e.s. bz., v.e.s. eth., chlf. $[\alpha]_D^{25} = +40.8^{\circ}$ (in 97% alc., $p = 2$). — $B_2H_2SO_4$, pr. alm. i. c. aq., alc.
882	145c.	† α-Triphenylguanidine , $PhN:C(NH.Ph)_2$. — <i>Cryst. easily in ndl. & rosettes fr. h. alc.</i> Alm. i. h. aq.; s. in 22 pt. alc. at 0 $^{\circ}$. — \textcircled{P} To sol. 0.05 g. B in 1 cc. warm alc., add 3 cc. c. sat. aq. sol. picric ac. After standing some min. the fine ppt. becomes granular &, under microscope, is seen to consist of single & rosetted ndl. Filter. Wash w. 2 cc. dil. alc. (1:1). Dry at 100 $^{\circ}$ for 15 min. • The picrate obtained is yellow, <i>YT1</i> , m.p. 180-1 $^{\circ}$ u.c. w. sl. dec. (183.4-4.4 $^{\circ}$ c.).
—	145	4,4'-Diaminohydrazobenzene . — Cf. No. 2.3317. (Yellow.)
884	d.140-50	2,4,6-Triamino-1,3-Xylene , $Me_2C_6H(NH_2)_2$. — Browns in air. Sbl. in ndl.
885	146-7u.c.	† Papaverine , Tetramethoxybenzylisoquinoline , $C_{20}H_{21}O_4N$. — [Narcotic alkaloid fr. opium.] — Odorless, tasteless ndl. fr. alc. Alm. i. h. aq.; e.s. chlf., alc.; mod. s. h. bz.; s. in 258 pt. eth. at 10 $^{\circ}$. — Opt. i. — [Commercial B usually contains kryptopine & the color reactions given in its literature are often due to this impurity. The reactions below were obtained fr. a preparation thoroughly purified by repeated recrystn. of the oxalate.] • \textcircled{P} No colorations in T. 2.2 w. conc. H_2SO_4 , or w. Erdmann's, Froehde's or Mandelin's reagts. — Marquis' reagt. [T. 2.2-(h)] gives slowly a very faint pink, changing to brown. • Dis. 0.05 g. B in 3 cc. h. alc. Add 3 cc. c. sat. alc. sol. picric ac. Cool. Filter. Dis. ppt. in 7 cc. boiling alc. The picrate obtained separates fr. the cooled sol. in small lust. scales which, after drying at 100 $^{\circ}$, melt at 179-81 $^{\circ}$ u.c. d. — $B_2C_10H_8O_6N_4\ddagger$ (picrolonate, fr. components in dil. alc. sol.), fine ndl., v.d.s. alc., m.p. 220 $^{\circ}$.
886	147	Diphenylguanidine , $HN:C(NH.Ph)_2$. — Monoclin. ndl. fr. alc. D.s. c. aq.; s. in 11 pt. c. 90% alc. Conc. HCl at 250 $^{\circ}$ splits to aniline, CO_2 & NH_3 . — $B_2HAuCl_4\parallel$ lft.
887	148 (to reddish liq.)	Tris-(p)-4',4"-Aminophenylmethane , p-Leucoaniline, $CH(C_6H_4NH_2)_3$. — Lft. — \textcircled{P} Ac. ac. sol. + little PbO_2 gives violet coloration. Heated w. little chloranil in alc. sol. gives bright red color of parafuchsine (No. 3.140).
888	148	p-Dimethylaminobenzaldehydophenylhydrazone , $NMe_2C_6H_4CH:N.NH.Ph$. — Ndl. fr. alc.
889	148	6-Hydroxyquinolinetetrahydride , $HO.C_9H_8N$. — [D.R.P. 42,871.] — S. acids or alk. — Acetyl deriv., \parallel white ndl., m.p. 82 $^{\circ}$.
890	149	4,4'-Diamino-3,3'-dimethylidiphenylmethane , $CH_2[C_6H_4Me(NH_2)]_2$. — Lft. fr. aq., alc. — Acetyl deriv., \parallel m.p. 198 $^{\circ}$.
890-I	149	a-Diethylsemicarbazide , $Et_2N.NH.CO.NH_2$. — Reduces boiling Fehling's sol. only v. slowly. — Pr. fr. alc. V.s. alc., h. aq.; alm. i. eth.
891	146-52	5-Aminoresorcinol, Phloramine , $NH_2C_6H_4(OH)_2$. — Silky ndl. D.s. c. aq.; e.s. alc.; alm. i. eth. Oxidizes quickly in air! Long boiling w. aq. dec. to phloroglucine & NH_3 . — Triacetyl deriv., cryst. powd., alm. i. lgr., m.p. 119-21 $^{\circ}$.
892	150	2,5-Diamino-p-xylene , $Me_2C_6H_4(NH_2)_2$. — Ndl. D.s. c. aq.; e.s. alc., h. aq. — $B_2H_2SO_4$, cryst. meal, alm. i. h. aq.
893	150	3,4',4"-Triaminotriphenylmethane , Pseudoleucoaniline , $(NH_2C_6H_4)_3CH$. — Rosettes w. adamantine lust. fr. eth. + lgr. Cryst. fr. bz. w. $1C_6H_4$, m.p. 145 $^{\circ}$. — Heated at 150 $^{\circ}$ w. HCl gives violet dyestuff (dif. fr. p-leucoaniline).
894	151-2	Aminopentamethylbenzene , $NH_2C_5Me_4$. — B.p. 277-8 $^{\circ}$. — Ndl. fr. dil. alc. I. h. aq.; e.s. alc., eth. — Acetyl deriv., \parallel ndl. fr. alc., m.p. 213 $^{\circ}$.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
895	151-2	Tetramethylleucoaniline, $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{CH.C}_6\text{H}_4\text{NH}_2$. — Cryst. w. adamantine lust. fr. alc. R.d.s. alc. — Little PbO_2 w. dil. sol. in ac. ac. gives violet-red color.
896	150-2	4,4'-Diaminobiphenyleneoxide, $[\text{O.C}_6\text{H}_4(\text{NH}_2)\text{C}_6\text{H}_4(\text{NH}_2)]^2$. — Ndl. fr. h. aq.
897	152	Carbohydrazide, $\text{CO}(\text{NH.NH}_2)_2$. — \oplus Reduced Fehling's sol. in cold. — Lust. ndl. fr. dil. alc.; i. eth., bz. — Acid or alk. hydrolysis (T. 2.17) gives hydrazine & CO_2 . — $\text{B.H}_2\text{SO}_4$ lust. pr. e.s. h. aq., m.p. 218° w. efferv.
898	152	Ibogaine, $\text{C}_{21}\text{H}_{29}\text{O}_2\text{N}_2$. — [Alkaloid of cocaine-like action fr. Tabernoe montana, Congo region.] — Cryst. alm. i. aq.; e.s. alc., eth., chlf., bz. Taste similar to cocaine (No. 2.741). $[\alpha]_D^{16} = -12.88^{\circ}$ (0.4851 g. in 25 cc. bz.)
899	152-3	Dimethylanilineoxide, $\text{Ph.NMe}_2\text{O}$. — Glassy pr., deliq. in moist air. V.s. aq., alc., chlf.; alm. i. eth., pet.-eth. — \oplus Warmed on water-bath w. benzaldehyde gives malachite green (No. 3.177). — Heated above m.p. gives dimethylaniline (No. 2.1247) & gas. — B.HCl , ndl. fr. acetone + bz., m.p. $124-5^{\circ}$. — $\text{B.Pk}, \ddagger$ (by pptn. fr. aq. sol.) yel. ndl. w. violet reflections. M.p. $137-8^{\circ}$ d.
900	152-3	Cytisine, Ulexine, $\text{C}_{11}\text{H}_{14}\text{ON}_2$. — [Toxic alkaloid fr. seeds of Cytisus Laburnum L. Causes vomiting.] — Clear cryst. fr. lgr. V.s. aq., alc., bz., chlf.; mod. s. eth., amyl alc.; v.d.s. lgr. Sbl. in ndl. — $[\alpha]_D^{17} = -119.57^{\circ}$ (in 1.99% sol.). Reacts strongly alk. May be extracted fr. acid sol. by chlf. — \oplus FeCl_3 gives blood-red color which disappears on dilution w. aq. or acidification. Addition of a few drops H_2O_2 sol. to red sol. discharges color; but sol. then becomes blue if warmed on water-bath. Results best w. proportions, 1 mol. B , 1 atom Fe, & 2 atoms O. (Cf. Arch. Pharm. 229, 57 & 233, 525; Z. anal. Chem., 34, 648. — K-Bi-iodide sol. (T. 2.3) gives brown-red ppt. in dil. sol. — Ignition w. soda-lime gives pyrrole (Cf. T. 2.24), etc. — \oplus Warm on water-bath w. 2 pt. conc. HNO_3 , & ppt. nitronitroso deriv., $\text{C}_{11}\text{H}_{14}\text{ON}_2$ (NO_2) (NO), by dilution of sol. w. aq. Recryst. fr. 50% alc. Product cryst. in yellowish scales, m.p. $242-4^{\circ}$ (Rosenthaler, p. 750). — $\text{B.HAuCl}_4, \parallel$ brown-red ndl., m.p. $212-3^{\circ}$ w. frothing.
901	154	4-Aminoquinoline, $\text{C}_9\text{H}_7\text{N.NH}_2$. — Ndl. w. $1\text{H}_2\text{O}$, m.p. $69-70^{\circ}$, fr. aq. (losing aq. at 100°). V.d.s. lgr.; e.s. chlf. — $\text{B}_2\text{H}_6\text{PtCl}_6\text{.2H}_2\text{O}$, § yel. ppt.; or-yel. ndl. fr. conc. HCl , m.p. $268-70^{\circ}$ d. — $\text{B.MeI}, \ddagger$ ndl. fr. abs. alc., m.p. 224° .
902	154	Phenylleucoauramine, $\text{Ph.NH.CH(C}_6\text{H}_4\text{.NMe}_2)_2$. — Pr. V.s. bz., chlf.
903	155 u.c.	\ddagger Codeine, Morphinemethylether, $\text{C}_{18}\text{H}_{21}\text{O}_2\text{N}$. — [Fr. opium.] — Cryst. w. $1\text{H}_2\text{O}$ in clear rhombic octahedra by spontan. evapn. of sat. sol. in moist eth. Odorless. Taste, slightly bitter. S. at 25° in 120 pt. aq., 1.6 pt. alc., 12.5 pt. eth., or 0.66 pt. chlf. S. at 80° in 59 pt. aq.; at 60° , in 0.92 pt. alc. 100 pt. fusel-oil dis. 15.68 pt.; 100 pt. bz. dis. 9.6 pt.; s. anisole; i. lgr. Sol. in aq. ammon. abt. as in aq.; e.s. dil. acids & alm. quantitatively repptd. by caustic alk. $[\alpha]_D = -137.75^{\circ}$ (in 80% alc.). \oplus (1) Oxidation T. — Dis. 0.001 g. B in 6 drops conc. H_2SO_4 on crucible cover. Stir into sol. abt. 0.01 g. powd. cryst. sodium arsenite (an x.s. of this oxidant does not interfere), placing on steam-bath. A RVT3 color appears, changing in a few sec. to BV, & gradually darkening to BVS2 which persists for long time. — (2) T. 2.2-(f) w. Froehde's reagl. gives at first a G Y, changing in 1 min. to YGS1-2, in 10 min. to GB-GBT1, & in 40 min. to YGS1 which persists for an hr. or more. — (3) Pellagrini's & Husemann's reactions give results described under morphine (No. 2.1024). — (4) T. 2.2-(h) w. Marquis' reagl. gives a VS1, changing after 10-15 min. to VR. — (5) FeCl_3 T. described under morphine gives YOS1, unchanged after 15 min. — (6) T. 2.2-(b) w. conc. HNO_3 gives Y sol. — (7) Treat 0.001 g. B w. 5 cc. conc. H_2SO_4 + 1 drop 10% aq. sugar sol. on crucible cover, stirring. A brilliant VRT1 color appears, changing after 10 sec. (not longer) heating on water-bath to deep R. — (8) Does not give the iodic acid or Prussian blue tests described under morphine. \oplus Add 4 cc. c. sat. aq. sol. picric ac. to sol. of 0.1 g. B in 3 cc. boiling aq. + $\frac{1}{2}$ cc. HCl (sp. gr. 1.12). Cool & shake. Filter off ppt. Wash w. 2 cc. aq. + 2 drops HCl (sp. gr. 1.12). Recryst. fr. 6 cc. boiling 10% ac. ac. by slow cooling, allowing to stand an hr. or two. Filter. Wash w. 2 cc. solvent. Dry 15 min. at 100° on tile. The resulting picrate is obtained in short Y lusterless ndl., m.p. $195.5-6^{\circ}$ u.c. d.

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. \ddagger = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
904	156	Paytine, $C_{11}H_{24}ON_3$. — [Fr. white Payta bark.] — \textcircled{P} Bleaching powd. cautiously added to acid sol. produces dark red to blue color that quickly disappears. — Cryst. w. $1H_2O$ (lost at 130°) fr. alc. V.s. alc., eth., bz., lgr., chlf. — Levorotatory. — $B.HCl$ pr. fr. aq.
905	156	5,8-Diaminoquinoline, $C_9H_4N(NH_2)_2$. — Yellowish ndl. Sbl. w. decn. Not vol. w. st. V.s. alc.; s. eth. — $B_2H_2PtCl_6$, § dark red ndl.
906	154; 158	4-Aminopyridine, $NH_2.C_6H_4N$. — Ndl. fr. bz. E.s. aq., alc.; s. bz., eth.; d.s. lgr. — $B_2H_2PtCl_6$, § m.p. 190–200° d. — $B.HAuCl_4$, m.p. 195–200°. — Acetyl deriv., cryst. w. aq., lost at 110°, m.p. then 150°.
907	158	4,4'-Diaminodiphenylamine, $NH(C_6H_4.NH_2)_2$. — [Intermediate in dye mfg. & oxidation black for coloring hair.] — \textcircled{P} Aq. sol. colored intense dark green by $FeCl_3$! — Lft. fr. aq. Oxidizing agents give benzoquinone (Vol. I). — $B_2H_2SO_4$ (120°), thin ndl. v.d.s. aq. — Diacetyl deriv. (fr. acetic anhydride), ndl. e.s. alc., m.p. 239°.
908	159.5	† p-Aminoacetanilide, Acetyl-p-phenylenediamine, $NH_2.C_6H_4.NH.CO.Me$. — Ndl. fr. aq. D.s. c. aq.; mod. s. h. aq.; v.s. alc., eth. — E. sapd. to p-phenylenediamine (No. 2.877) by boiling conc. HCl (Cf. T. 2.26).
909	159–60	β -Homochelidonine, $C_2H_2O_2N$. — [Fr. Chelidonium majus, etc.] — Pr. fr. acetic eth. E.s. chlf.; less s. alc., CS_2 , bz. — \textcircled{P} Conc. H_2SO_4 gives yellowish & then rose or red-violet color. — Froehde's reagt. gives violet, then blue, & finally moss-green colors. Erdmann's reagt. gives yellow, violet, & dirty violet colors. — $B.HAuCl_4$, blood red warts fr. alc., m.p. 187°.
910	159–61	4-Aminocresol(2), $NH_2.C_6H_4Me(OH)$. — [Pat.] Colorless lft. or ndl. Sbl. Mod. s. c., e.s. h. aq.; v.s. alc., eth. — Acetyl deriv., pr. fr. dil. alc., m.p. 224–5°; & diacetyl deriv. (using acetic anhydride in x.s.), m.p. 132.5°.
911	159	2,7-Diaminonaphthalene, $(NH_2)_2C_{10}H_6$. — Lft. fr. aq.
912	160	p-Hydroxyphenylethylamine, $HO.C_6H_4.CH_2.CH_2.NH_2$. — [Fr. ergot.] B.p. 179–81° (8 mm.). S. in 10 pt. boiling alc., less s. aq.; d.s. h. xylene. — \textcircled{P} Gives T. 2.19 w. Millon's reagt. — Picrate, † fr. aq., m.p. 200°.
913	160; 159 softening at 155	Anhalonidine, $C_9H_{11}O_2N$. — [Fr. Anhalonium Lewinii, Mexican mescal buttons.] — \textcircled{P} FeCl ₃ gives blue color w. aq. sol. of salts, changing to green & disappearing. — Ndl. e.s. aq.; sol. strongly alk. Opt. i. — $B_2H_2PtCl_6$, § d.s. aq. — $B.HAuCl_4$, unstable, d.s. aq., m.p. 152°. — Benzoylation (w. benzoyl chloride in alk. sol., Cf. T. 2.27) gives dibenzoyl deriv., pr. fr. alc., m.p. 125–6°.
914	161	6-Amino-1,3-xylenol(4), $NH_2.C_6H_4Me_2(OH)$. — Cryst. E.s. alc., eth.
915	161 u.c.	2,7-Diaminofluorene, $(NH_2)_2C_{12}H_8$. — Ndl. fr. alc. V.d.s. aq.; e.s. alc. — Sulphate, lft., d.s. aq.
916	161	4-Dimethylamino-4'-hydroxyphenylamine, $Me_2N.C_6H_4.NH.C_6H_4.OH$. — Pr. fr. bz. D.s. c. aq., alc., eth., bz., chlf. — Heated w. Na_2S + S gives pure blue sulphur dyestuff, Immedial Pure Blue.
917	161	Nortropine, Tropigenine, $C_7H_{11}ON$. — Ndl. by sbln. E.s. aq., alc.; less s. eth. Absorbs CO, fr. air. Alc. sol. boiled w. MeI gives tropine. — $B.HAuCl_4$, gold-yel. lft., m.p. 215–6° d. — Benzoyl deriv. (T. 2.27), pr., m.p. 125°.
918	161	2,4,2',4'-Tetraaminodiphenylmethane, $[(NH_2)_2.C_6H_4]_2CH_2$. — Ndl. fr. h. bz. S. aq.; v.s. alc.; d.s. bz.
919	162	m-Aminobenzaldehyde-phenylhydrazone, $NH_2.C_6H_4.CH:N.NH.Ph$. — Ndl. fr. alc., e.s. alc., eth.; alm. i. lgr.; i. aq.
920	162–3	6,8-Diaminoquinoline, $(NH_2)_2C_9H_4N$. — Cryst. Not vol. w. st. E.s. aq., alc.; less s. eth., bz.; v.d.s. chlf., lgr. — $B_2H_2PtCl_6$, § light yel. cryst. powd.
921	162	Retamine, $C_{15}H_{22}ON$. — [Fr. plants of Retama sphaerocarpa family.] — Ndl. fr. pet.-eth.; lft. fr. alc. 100 g. sat. alc. sol. at 17° contains 2.462 g. E.s. aq., eth. Taste bitter. Not toxic. Titrates as diacid base w. phenolphthalein. $[\alpha]_D^{25} = +43.11$ to 43.15° (in alc., sp. gr. 0.799).
922	162	Aspidospermatine, $C_{21}H_{22}O_2N_2$. — [Fr. bark of Aspidosperma Quebracho.] Ndl. fr. lgr. Freshly pptd., mod. s. aq. V.s. alc., eth., chlf. Alc. sol. reacts alk. — $[\alpha]_D^{25} = -73.3^\circ$ (in 97% alc., $p=2$). — $B_2H_2PtCl_6.4H_2O$, § pale yel. ppt.

No.	Melting-point (C. ^b).	BASIC COMPOUNDS.—Colorless and Solid.
923	163	r-Bornylamine, 1-Aminocamphane, $\text{NH}_2\text{C}_{10}\text{H}_{17}$. — Camphor-like mass. Odor piperidine-like! I. aq. — $[\alpha]_D = +23.2^\circ$ (2% sol. in alc.). — B.Pk, \ddagger S.yel. ndl., m.p. 257° d.
923-1	163-4 (bath fr. 155)	† Cupferron, Ammonium Salt of Nitrosophenylhydroxylamine, Ph.N(NO)(ONH ₄). — Silvery lft. fr. alc. E.s. c. aq.; v.d.s. eth., bz. In light becomes yellowish & acquires pungent nitrosobenzene odor. A preparation fr. Kahlbaum, which became black without melting at 131–9° u.c., neutralized little alkali in Gen. T. 2.I, but much more in the titration in alc. sol. of Gen. T. 2.II–III, gave following reactions: NH ₃ evolution in T. 2.7 w. NaOH sol.; Ag ppt. in T. 2.21, but not w. Tollen's reagt. alone; a heavy floc. YOS1 ppt. on adding dil. aq. sol. to dil. FeCl ₃ sol., or heavy GB Bkn [L] ppt. w. CuSO ₄ sol. — [Used as reagent in Fe & Cu analyses. Cf. Chem. Ztg., 1909, 1298, & J. Ind. Eng. Chem. 1911, 629, for fuller description.]
924	165	Dimethylethylenediphenyldiamine, Ph.NMe.CH ₂ CH.NMe.Ph. — Lft.
925	165	2,4',4"-Triaminotriphenylmethane, $\text{NH}_2\text{C}_6\text{H}_4\text{CH}(\text{C}_6\text{H}_4\text{NH}_2)_2$. — Sl. brownish cryst. fr. alc.
926	165; 166	2,2',4,4'-Tetraaminobiphenyl, $(\text{NH}_2)_2\text{C}_6\text{H}_4\text{C}_6\text{H}_4(\text{NH}_2)_2$. — Lft. fr. ammon. — Tetraacetyl deriv. (by boiling w. gl. ac. ac. & ac. anhydride), ndl., d.s. aq., m.p. 284°.
927	165-7d.	4,4'-Dihydrazinobiphenyl, $\text{NH}_2\text{NH.C}_6\text{H}_4\text{C}_6\text{H}_4\text{NH.NH}_2$. — Lust. lft. D.s. h. aq.; v.d.s. alc., eth., chlf. — Dinitroso deriv.** (fr. B.2HCl + 2NaNO ₂), d.s. bz., m.p. 112–3° d.
928	160-70d.	Apomorphine, C ₁₇ H ₁₇ O ₂ N. — [B.HCl is powerful emetic & expectorant. B is pptd. as powd. by adding NaHCO ₃ to sol. of B.HCl, but is v. difficult to isolate pure, becoming colored to a broken green, so that m.p. is a rather unreliable constant.] Odorless, alm. tasteless powd. quickly becoming greenish. E.s. alc., chlf.; d.s. aq.; s. in x.s. alk. [Being a dehydration product of morphine many of its color reactions are related more or less closely to those of the latter alkaloid (No. 2.1024). The corresponding morphine tests are however not usually identical, it being often necessary to prescribe conditions that will bring about a preliminary dehydration to apomorphine.] ② (1) FeCl ₃ T., as described for morphine, gives RV Bkn. [D] color, changing in 5 min. to BV Bkn. [D]. — (2) T. 2.2-(f) w. Froehde's reagt. gives YGS2. — (3) Pellagri's React.: — Dis. 0.001 g. B in 1 cc. dil. HCl. Neutralize w. NaHCO ₃ . Add 5 cc. aq. + 3 drops alc. T.S. of I. Shake several min. Then shake w. 2 cc. eth. The aq. sol. becomes BG-BGS2 & the eth. sol. R. — (4) Husemann's React.: — Dis. 0.001 g. B in 1 drop conc. H ₂ SO ₄ . Add fraction of drop conc. HNO ₃ , fr. stirring rod. A VR color, changing within 1 or 2 sec. to OR, & after 15 min. to O, appears. — (5) T. 2.2-(b) w. conc. HNO ₃ , gives VR color, changing within a min. to RO. — (6) Iodic Ac. T.: — Test & results are as described for morphine. — (7) Marquis's react. applied as directed under morphine gives fleeting V Bkn. [M] color, changing almost immediately to permanent green-black. — (8) Prussian Blue T.: — Test & results are as described for morphine. — (9) T. 2.2-(a) w. conc. H ₂ SO ₄ gives colorless sol. as w. morphine. — (10) Wangerin's React. (Pharm. Zt. 48 (1903), 688): — Dis. 0.01 g. B.HCl in 1 cc. c. aq. Add 4 drops 0.3% K ₄ Cr ₇ O ₇ sol. Shake 1 min. Sol. becomes BGS2. Shake w. 10 cc. ethyl acetate. A VS2 sol. is obtained. Add 3 drops SnCl ₄ sol. [prepared by dissolving 1 g. SnCl ₄ ·2H ₂ O in 50 cc. HCl (sp. gr. 1.12) + 50 cc. aq.]. Upon shaking, the color of ac.-eth. layer changes to G. If 10 cc. chlf. be substituted for ac.-eth. in test, & 1 drop SnCl ₄ sol. be added, chlf. layer becomes deep blue, BS1).
929	165.5-6	Tetramethyldiaminobenzidine, NMe ₂ C ₆ H ₄ (NH ₂) ₂ C ₆ H ₄ (NH ₂) ₂ NMe ₂ . — ② Gives violet color w. FeCl ₃ or drop of NaNO ₂ sol. — Ndl. fr. bz. D.s. c. alc., lgr.; v.s. bz. — B.H ₂ PtCl ₆ , § light yel. powd.
930	166	Laudanine, C ₂₀ H ₂₂ O ₂ N. — [Fr. opium.] — Pr. fr. dil. alc. D.s. c. alc.; e.s. chlf., bz.; s. in 647 pt. eth. at 18°. Opt. i. Reacts alk. Tasteless. Salts bitter. Pptd. by KOH fr. sol. of salts, redissolves in x.s. alk. Chlf. extracts fr. ammon., but not fr. KOH sol. — ② Dis. w. pale rose-red color in conc. H ₂ SO ₄ , changing to dark violet on warming. — B.H ₂ PtCl ₆ ·2H ₂ O, § yel. cryst. ppt.

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No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
931	166-7d.	Hydroquinidine, $C_{20}H_{28}O_2N_2$. — [Fr. Cinchona bark.] — Cryst. w. $2\frac{1}{2}H_2O$ in ndl. fr. dil. alc. Dextrorotatory. E.s. alc., chlf.; r.d.s. eth. Alc. sol. reacts alk. — \textcircled{P} Sol. in x.s. dil. H_2SO_4 , fluor. blue. — $B_2H_4PtCl_6 \cdot 2H_2O$, $\frac{1}{2}$ short or. ndl.
932	167-8d.	3,4-Diaminophenol, $(NH_2)_2C_6H_4OH$. — V. unstable. — B_2HCl , lft. e.s. aq.; sol. gives blood-red color w. bleaching powd. sol.
933	168	2,2',5,5'-Tetraaminobiphenyl, $(NH_2)_2C_6H_4C_6H_4(NH_2)_2$. — Ndl. fr. toluene. E.s. aq., alc., chlf.; d.s. eth., h. bz.; i. lgr.
934	168	Di-p-tolylguanidine, $NH_2C(NH_2C_6H_4)_2$. — Ndl.
935	168-8.5 u.c.	† Quinidine, Conchinine, $C_{20}H_{28}O_2N_2$. — [In Cinchona barks.] — Cryst. fr. aq., alc., eth., bz., etc., w. varying quantities of solvent (lost in drying). Clear cryst. fr. c. alc. S. at 15° in 2000 pt. aq.; at 10° in 35 pt. eth.; at 20° , in 26 pt. 80% alc.; d.s. chlf., lgr., bz. $[\alpha]D^{25} = +289.57-3.90^{\circ}$ c. (in 97% alc., c = 1-3). Odorless. Taste, bitter. \textcircled{P} Gives the blue fluorescence described under "(1)" in characterization of quinine (Cf. No. 2.947). — Is a stereoisomer of quinine and gives the same results as quinine in all the numbered sub-tests (1) to (4) under No. 2.947, if possible slight differences in the appearance of the ppt. in thalleioquin & herapathite tests be excepted. It is most readily distinguished fr. quinine by the strong dextrorotation of its alc. sol.
936	169	γ -Homochelidonine, $C_{21}H_{28}O_2N$. — [Fr. root of Sanguinaria canadensis.] — Tbl. fr. ac.-eth., requiring drying at 100° . — B_2HAuCl_4 , blood-red warts fr. alc., m.p. 187° .
937	168-70	3,5-Diaminophenol, $(NH_2)_2C_6H_4OH$. — Pr. E.s. aq.; d.s. eth.
938	170	o-Aminophenol, $NH_2C_6H_4OH$. — Scales, soon turning brown in air. Sbl. S. at 0° in 59 pt. aq. or 23 pt. alc.; much more s. eth. — B_2HCl , ndl. s. in 1.25 aq., or 2.36 pt. alc. at 0° . — Me-ether (o-anisidine, No. 2.1332), b.p. 225° (th.i.).
939	170d.	4,6-Diaminocresol(3), $(NH_2)_2C_6H_4Me(OH)$. — Ndl. V. unstable.
940	170	Anthraquinoline, $C_{17}H_{11}N$. — B.p. 446° . — Lft. I. aq.; e.s. alc., eth., bz. — \textcircled{P} Sol. show intense blue fluor. — Salts yel., their dil. alc. sol. fluorescing intense green! — Oxid. w. CrO_3 in gl. ac. ac. sol. gives anthraquinone (Vol. I).
941	170u.c.	† Heroine, Diacetylmorphine, $C_{18}H_{21}O_3N$. — [Synthetic narcotic.] — Small colorless pr. fr. eth. Said to also cryst. w. $\frac{1}{2}$ mol. or more of aq. of cryst. Alm. i. c. aq.; s. alc.; e.s. chlf., bz. Pptd. by NH_4OH or alk. fr. aq. sol. of salts & rediss. by x.s. reagt. May be extracted fr. alc. sol. by eth. or chlf. E. sapd. to morphine, even by long boiling w. aq. Odorless. Taste, faintly bitter. \textcircled{P} Color reactions described (Cf. No. 2.1024) for morphine give following results: (1) w. $FeCl_3$, & (2) w. iodic acid, negative. (8) Prussian blue T., is also negative, though a heavy blue ppt. appears after an hr. (3) Pelagri's, (4) Husemann's, (9) H_2SO_4 , & (10) $AgNO_3$ tests, give same results as morphine. (7) Marquis' T., is as w. morphine, but somewhat slower. — (2) Froehde's reagt., gives VRS1, changing to BGS1 within 10 min., & to GY after 40 min. — much as w. morphine. (Delicacy 0.000005 g. according to Dragendorff.) (5) HNO_3 , T., gives first GYT1, changing on steam-bath after 1-3 min. to G, & then gradually fading to Y. — 0.005 g. B warmed w. 2 cc. H_2SO_4 (sp. gr. 1.175) & then w. 0.5 cc. alc. gives odor of ethyl acetate. — Picrate, † (fr. hydrochloride + aq. pic. ac. sol. & recrystd. fr. 50% alc.), hexag. plates, m.p. $200-5^{\circ}$ u.c. d.
942	172	Quinamine, $C_{18}H_{21}O_2N_2$. — [Fr. Cinchona barks.] — Pr. fr. h. dil. alc. S. at 16° in 1516 pt. aq.; at 20° , in 105 pt. 80% alc.; at 20° , in 32° pt. eth.; e.s. h. bz., lgr. $[\alpha]D^{20} = +104.5^{\circ}$ (in 97% alc., p = 2). "Reacts alk." — \textcircled{P} Writing made w. quill pen & sol. of a salt of B strongly acidified w. H_2SO_4 on paper becomes olive-green when placed over watch glass containing conc. H_2SO_4 and few cryst. $KClO_3$; and, after exposure to air, azure blue, or if sol. is conc., dark black-blue; & after moistening w. aq., rose-red. (Ann. 197, 56). — B_2HCl reduces gold chloride sol.
943	172.3c.	Hydroquinine, $C_{20}H_{28}O_2N_2$. — [Fr. Cinchona barks.] — Cryst. w. $2H_2O$. Ndl. fr. chlf., eth. D.s. aq.; e.s. org. solvents, ammon.; i. $NaOH$. Reacts basic. Taste, bitter. $[\alpha]D^{20} = -142.2^{\circ}$ (in 95% alc., p = 2.4). — \textcircled{P} S. in dil. H_2SO_4 fluor. blue & gives same color reacts. as quinine w. Cl-aq. & ammon. (Cf. No. 2.947). — $B_2H_4PtCl_6 \cdot 3H_2O$, yel. ppt., d.s. aq., alc. — Cf. also Chem. Zentr. 1897, II, 915.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
944	172; 160	† Gelsemine ("Gelseminine"), $C_{22}H_{28}O_2N_2$. — [Alkaloid of strychnine-like action fr. Gelsemium sempervirens or yellow jasmine. The name "gelseminine" should perhaps be reserved for another jasmine alkaloid, but there is some confusion on this point. Descriptions, also, are not particularly satisfactory.] — Silky ndl. fr. bz., softening abt. 100° & melting clear at 160° when dried in desiccator, but at 172° when dried at temperature of boiling toluene & xylene. D.s. aq.; e.s. alc., eth., chlf.; s.v. alkali. Taste, bitter. (Caution!) — ② Sol. in conc. H_2SO_4 , colorless, becoming light red, then brown-red, & finally intense green, w. crystal of $K_2Cr_2O_7$. — W. little MnO_2 powder, H_2SO_4 sol. becomes fine wine-red, increasing for a time in intensity, but color later disappearing except in scattered points. — Sol. in conc. HNO_3 reddens on warming, but finally becomes dark green. — Is rptd. by general alkaloidal precipitants (T. 2.3). — $B. MeI \cdot 2H_2O$; (by warming alc. sol. w. MeI on water-bath, lust. tbl. fr. aq., m.p. 286°).
945	173	Hexamethyleucoaniline, $[NM_eC_6H_4]_3CH$. — ② Melts to blue liquid! — Silvery lft. I.c. aq.; d.s. c. alc.; e.s. eth., chlf., bz.
946	173	Pseudophenanthroline, $C_{12}H_8N_2$. — Cryst. w. $4H_2O$ (lost at 100°) in thick ndl. fr. aq. Dist. undec. S. h. aq.; e.s. alc.; d.s. eth., bz.; v.s. chlf. Alc. sol. "feeble alk." Alc. sol. colored red-yel. by $FeCl_3$. — BI, red-brown ppt. giving sol. in warm ammon. which shows characteristic color changes on heating & cooling. Dark blue-gray ndl. fr. warm alc.
947	abt. 173u.c.	† Quinine, $C_{20}H_{24}O_2N_2$. — [Fr. Cinchona barks.] — [Is usually more or less hydrated. $B.3H_2O$, m.p. 57°, losing $2H_2O$ at 100°, & $3H_2O$ at 125° (U.S.P.). Commercial preparations fr. three reputable makers melted at 168–70° u. c., which is the m.p. of anhydrous quinine given by Flückiger! Rosenthaler accepts m.p. 174.6°.] Silky ndl. fr. bz. — S. at 25° dis. in: 1750 pt. aq.; 0.6 pt. alc.; 4.5 pt. eth.; 1.9 pt. chlf.; 120 pt. bz.; 3450 pt. KOH (1:20); 1810 pt. ammon. aq. S. at 80° in 810 pt. aq. (U.S.P.). — $B.3H_2O$ dis. at 25° in: 1550 pt. aq.; 0.6 pt. alc.; 1.3 pt. eth.; 1.6 pt. chlf.; 166 pt. bz.; 3450 pt. KOH (1:20); 1810 pt. ammon.-aq. S. in 775 pt. aq. at 80°. (U.S.P.) — Odorless. Taste, v. bitter. — $[\alpha]D^{20} = -158.2^\circ$ (in abs. alc., $c = 2.136$). ② (Among following tests (1), (2), (3), are of first importance.) (1) Fluorescence T. — Dis. 0.001 g. B in 10 cc. dil. H_2SO_4 . A pronounced blue fluor. (best observed in strong light w. black background) appears, & is still distinguishable on further dilution to 100 cc. HCl sol. does not fluor. — (2) Thallequin T. — Dis. 0.001 g. B in 6 drops 10% ac. ac. Add 10 drops sat. Cl -aq., & then 2 cc. ammon. (sp. gr. 0.90). The clear sol. becomes BG-G. Or, rub together 0.001 g. B , 0.005 g. bleaching powd., 2 cc. aq., & 2 drops conc. HCl . Add 2 cc. ammon. (sp. gr. 0.90). A BG-G sol. & BGS_1 floc. ppt. appears. Urea interferes w. the test, & antipyrine & caffeine do under certain conditions (Cf. Authenreith). — (3) Herapathite T. — Add 20 drops of mixl. of 30 drops ac. ac. + 20 drops alc. + 1 drop H_2SO_4 (sp. gr. 1.176) to 0.01 g. B & heat to boiling. Add 1 drop of 1% alc. I. sol. Allow to stand for some time. Examine the shimmering green lft. of "herapathite" ($4B \cdot 3H_2SO_4 \cdot 2HI \cdot I_2 \cdot 6H_2O$) which precipitate. These cryst. are small transparent rhombic bl. which when viewed separately under microscope appear nearly colorless; but, since they polarize light strongly like tourmaline, many of them which overlap look perfectly black on the slide! Gadamer recommends to always begin w. this test in forensic cases, since B may be recovered for other tests by addition of sulphurous ac. followed by treatment w. alk. & extraction w. eth. or chlf. — (4) Dis. 0.1 g. B in 0.5 cc. dil. H_2SO_4 (sp. gr. 1.176) + 10 cc. aq. Neutralize exactly w. ammon. Add 1 drop 3% H_2O_2 + 1 drop $CuSO_4$ T.S. Boil. Sol. becomes red, slowly changing to blue, & finally to yellowish green. According to U.S.P. quinine & quinidine are only alkaloids responding to this test. ③ Ppt. sol. of 0.05 g. B in 3 cc. aq. + 3 drops HCl (sp. gr. 1.12) w. 4 cc. sat. aq. picric ac. sol. Wash curdy Y ppt. w. 2 cc. c. aq. Dis. in 1 cc. h. gl. ac. ac. Add 6 cc. aq. Heat to boiling & allow to cool slowly. After standing over night, wash ppt. of mic. YT1 ndl. on filter w. 1 cc. 10% ac. ac. sol. The product, $B.Pk$, dried at 100°, melts at 125–6° u.c. (Yield small.)
948	abt. 174	5-Aminocresol(2), $NH_2C_6H_4Me(OH)$. — Lft. fr. bz. Sbl. E.s. alc., eth., less s. aq., bz. — $B.HCl$ sol. not colored by $FeCl_3$. — Oxidizing agents give toluquinone easily.

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No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
949	174d.	6-Aminocresol(3), $\text{NH}_2\text{C}_6\text{H}_4\text{Me}(\text{OH})$. — Warts fr. bz. — Oxidation w. CrO_3 gives toluquinone, m.p. 67–8°. — Acetyl deriv., lft. fr. aq., m.p. 80°, or 125° when anhydrous.
—	171c.; 176	2,7-Dimethylacridine. — Cf. No. 2.3491. (Colored compounds.)
951	175	Phenylguanazole, $\text{PhN.NH.C(:NH).NH.C(:NH)}^2$. — Cryst. v.s. aq. (pptd. by NaOH); e.s. alc.; d.s. eth., bz. — B_2H_2 , ndl. e.s. aq., alc., m.p. 240°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or.-yel. ndl. — B_2AgNO_3 , floc. ppt.
952	176; (174u.c.)	† Narcotine, Opianine, $\text{C}_{20}\text{H}_{22}\text{O}_2\text{N}$. — [Weak narcotic alkaloid fr. opium.] — Lust. ndl. fr. alc. 1. aq.; 100 pt. 85% alc. dis. 1 pt. cold, or 5 pt. boiling; 100 pt. eth. dis. 0.77 pt. c., or 2.1 pt. boiling; 100 pt. c. bz. dis. 4.61 pt. (separation fr. morphine, which is i.); i.c. NaOH ; v.d.s. NH_4OH ; e.s. chlf. $[\alpha]_D = -207.35^\circ$ (in chlf. sol.). Odorless. Tasteless.
		① (1) Husemann's T. — Dis. 0.001 g. in 5 drops aq. + 1 drop conc. H_2SO_4 on crucible cover. Evap. to dryness on steam-bath. Residue, at first ORT_1 darkens to ORS_1 . Heat cautiously over small flame until white fumes appear. Residue becomes $VR-RV$ Bkn. [D]. (Recommended by Dragendorff as most charac. test, w. limit of 0.1 mg.) — (2) T. 2.2-(f) w. Froehde's reag. gives YGS_2 color; but if applied w. reagent containing 0.05 g. sodium molybdate in 1 cc. conc. H_2SO_4 this color gradually changes to GBS_1 , & then on heating on steam-bath, if removed after 1 min., gives deep R w. VB border, gradually changing to VB which persist more than 2 hr. — (3) Bloxam's T. — Dis. 0.01 g. in 2 cc. aq. + 1 drop conc. HCl . Add 1 drop Br-aq. A YT_2 ppt. appears. Heat to boiling & add 6 drops Br-aq., boiling for a moment after each drop. An OT_1 sol. is obtained. Add 8 cc. aq. + 1 drop Br-aq. & boil again. Color of sol., now ROT_2 , becomes ORT_2 on being again boiled after addition of 20 cc. aq. — (4) T. 2.2-(a) w. conc. H_2SO_4 gives YT_1 color, changing on water-bath & stirring to OS_1 , RT_1 & RV Bkn. [M].
		② Add 3 cc. sat. aq. picric ac. sol. to sol. 0.05 g. B in 3 cc. aq. + 1 drop conc. HCl . Wash curdy yel. ppt. on filter w. 2 cc. aq. Dis. in 2 cc. c. gl. ac. ac. Add 6 cc. aq. & boil. Allow to cool slowly. Filter, & wash the Y-poud. w. 2 cc. dil. ac. ac. Dry 15 min. at 100°. B . Pk melts to clear yel. liq. at 141° u.c.
953	176	p-Aminophenoylethyleneether, $(\text{NH}_2\text{C}_6\text{H}_4\text{O})_2\text{C}_2\text{H}_4$. — [Intermediate of dye mfg.] — ① Sol. in conc. H_2SO_4 w. blue color. — Ndl. fr. alc. D.s. h. aq.; s. h. alc.; e.s. h. bz.; d.s. eth. — FeCl_3 gives cherry-red color.
954	176–7	† 6-Aminothymol, $\text{NH}_2\text{C}_6\text{H}_4\text{Me}^{(1)}(\text{C}_6\text{H}_5)^{(6)}(\text{OH})^{(3)}$. — Lft. fr. fusel-oil. Boiled w. FeCl_3 sol. gives thymoquinone (Vol. I).
955	177	Laudanidine, $\text{C}_{20}\text{H}_{22}\text{O}_4\text{N}$. — [Fr. opium.] — Cryst. fr. alc. Resembles laudanine in solubilities & reactions. — $[\alpha]_D^{20} = -87.8^\circ$ (in chlf., $p = 5$). — $\text{B}_2\text{H}_2\text{PtCl}_6$. — $4\text{H}_2\text{O}$, § brownish yel. amorph. ppt.
956	178u.c.	† Brucine, $\text{C}_{20}\text{H}_{22}\text{O}_4\text{N}_2$. — [Tetanic poison accompanying strychnine in Strychnos nux vomica.] — Cryst. w. $4\text{H}_2\text{O}$ fr. dil. alc. in clear lust.tbl., m.p. 105° (r.h.). May also cryst. w. $2\text{H}_2\text{O}$ fr. stronger alc. Effloresces at 100°. Sbl. Odorless. Taste intensely & persistently bitter. Nagelvoort (Cf. Fückiger, p. 24) claims that 5 cc. of aq. scl. 1:500,000 was found sufficient to produce slight sensation of bitterness after swallowing. Hydrated cryst. dis. in 320 pt. c., or 150 pt. boiling aq. Anhydrous B dis. in 850 pt. c., or 500 pt. boiling aq.; 1.5 pt. alc.; 2 pt. chlf.; 64 pt. bz.; e.s. amyl alc., pet.-eth.; i. eth. $[\alpha]_D^{20} = -80.1^\circ$ (in abt. 2% abs. alc. sol.).
		① (1) HNO_3 , T. — Dis. 0.001 g. B in 3 drops conc. HNO_3 . Sol. is O. Add 7 drops c. sat. aq. sol. sodium thiosulphate & keep sol. at 40–50°. Within 2 or 3 min. a VR color, accompanied by yel. ppt., develops. — (2) HNO_3 + SnCl_4 , T. — Dis. 0.001 g. in small drop conc. HNO_3 . Heat on steam-bath until the color, at first OR, changes to OY. Remove fr. bath & add 5 drops freshly prepared SnCl_4 , T.S. Color changes to VR. — (3) T. 2.2-(e) w. Erdmann's reag. gives RT_2 , changing quickly through ORT_2 & OT_2 within 5 min. to YT_2 . — (4) T. 2.2-(f) w. Froehde's reag. gives RT_1 , changing quickly to OS_2 & slowly, within an hr., to YG . — (5) Pure brucine does not give the color phenomena described in test (1) w. H_2SO_4 + MnO_2 under description of strychnine (No. 2.1047).
		② Add 4 cc. picrolonic ac. T.S. to sol. of 0.05 g. B in 10 cc. h. aq. Allow to stand several hr. Filter off the fine Y ppt. Wash w. 2 cc. alc.-eth. (1:3). Boil ppt. w. 3 cc. alc. Filter. Dry 15 min. at 100°. The resulting picrolonate, $\text{B}_2\text{C}_{10}\text{H}_{14}\text{O}_4\text{N}_2$, consists of small micro cryst. of charac. appearance, which begin to darken abt. 214° & melts to dark brown mass at 256–7° u.c.

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
957	178-80	Pseudocodeine, $\text{MeO.C}_1\text{H}_{18}\text{O}_2\text{N}$. — [Dehydration product fr. codeine.] Cryst. w. $1\text{H}_2\text{O}$ fr. dil. alc., or anhydrous fr. lgr. D.s. eth. $[\alpha]_D = -91.4^\circ$. — $\text{B.Pk}, \ddagger \text{d. } 209-10^\circ$.
958	180	4,4'-Diamino-o-hydrazotoluene, $\text{NH}_2\text{C}_6\text{H}_4(\text{Me})\text{NH.NH}(\text{Me})\text{C}_6\text{H}_4\text{NH}_2$. — Rhomb. tbl. I. aq., c. alc.; d.s. h. alc. — B E. oxidized in sol. to orange colored diaminoazotoluene.
959	180-1; 183	6-Amino-2,4-dimethylpyrimidine, Kyanmethine, $[\text{N:C}(\text{NH}_2)\text{CH:CM}\text{e}]$. — Ndl. fr. alc. — B S. in 0.64 pt. aq. at 18° . — S. in 5.25 pt. alc. — B.HCl , ndl. s. aq.; sbl. w. m. at $200-250^\circ$. — $\text{B}_2\text{H}_2\text{PtCl}_6$, \ddagger yel. ndl. fr. h. aq. — $\text{B.Pk}, \ddagger$ ndl., m.p. 214° .
960	182	α -Homochelidonine, $\text{C}_{11}\text{H}_{21}\text{O}_4\text{N}$. — [Fr. root of Chelidonium majus.] — Trimet. pr. fr. ac.-eth. E.s. chlf.; less s. alc., ac.-eth.; v.d.s. eth. — B.HAuCl_4 , yel.-red ndl. fr. alc.
961	182-3	Leucoanisidine, Triaminodihydroxytriphenylmethane-dimethyleneether, $[\text{NH}_2\text{C}_6\text{H}_5(\text{OMe})_3\text{CH.C}_6\text{H}_4(\text{NH}_2)_2]$. — Plates fr. abs. alc. Alm. i. aq.; v.s. alc.
962	182	Tritopine, $\text{C}_{14}\text{H}_{14}\text{O}_7\text{N}_2$. — [Fr. opium.] — Pr. fr. alc. E.s. chlf.; d.s. eth. Pptd. fr. sol. of salts by ammon.; ppt. s. in NaOH . A diacidic base.
—	183-4	γ -Aminobutyric Ac. — Cf. No. 2.2382.
963	184 <u>c. d.</u>	† p-Aminophenol, $\text{NH}_2\text{C}_6\text{H}_4\text{OH}$. — White scales by pptn. of B salt sol. by Na_2SO_4 sol. Oxidizes so quickly that ppt. formed by other methods quickly becomes brown or black. — S. in 90 pt. aq. at 0° , or in 22 pt. abs. alc. — B Heat 0.05 g. dry B.HCl in t.t. until it melts giving white sublimate & odor of phenol. Treat cold dark-colored residue w. 5 cc. aq. + 3 cc. conc. ammon. An intensely blue (B) sol. is obtained. The color is changed to red by acidification, & restored to blue by alk. — 0.002 g. B or its hydrochloride dissolved in 5 cc. aq. + 5 drops conc. ammon. gives intense RV color after short exposure to air. — B Dis. 0.1 g. B in 3 cc. dil. ac. ac. (3 vol. acid : 7 vol. aq., the proportions are important), without warming. Add 5 drops benzaldehyde to sol. & shake until sol. is complete. Continue the shaking longer and a heavy ppt. of pale yel. cryst. will appear. Allow to stand 5 min. Filter. Wash w. 3 cc. dil. ac. ac. Cryst. fr. 3 cc. h. dil. alc. (3 vol. alc. : 7 vol. aq.). The product, benzylidene-p-amino-phenol, $\text{HO.C}_6\text{H}_4\text{N:CHPh}$, is obtained in faintly yellowish rhombic plates, m.p. $180.5-1.5^\circ$ u.c. (183.9-4.9° c.).
964	184	1(or Neo-)Bornylamine, $\text{C}_{10}\text{H}_{17}\text{NH}_2$. — Powd. I. aq. $[\alpha]_D = -43.7^\circ$ (in 4% alc. sol.). — Acetyl deriv., \ddagger ndl. fr. pet.-eth., m.p. 144° . — B picrate, \ddagger ndl., m.p. 248° d.
965	184-5	† Cinchonamine, $\text{C}_{12}\text{H}_{24}\text{ON}_2$. — [Toxic alkaloid fr. Cuprea bark, Remijia purdieana Wedd.] — Lust. orthorhombic ndl. fr. alc. I. c. aq. S. at 17° in 31.6 pt. 90% alc., or in 100 pt. eth. E.s. h. chlf., bz.; v.d.s. lgr. Alc. sol. v. bitter. $[\alpha]_D^{18} = +121.1^\circ$ (in 97% alc., p = 2). Salts do not show fluor. B (1) T. 2.2-(g) w. Mandelin's reagt. gives BV color, changing to RV on stirring, & after 5-10 min. to BGS1 or BGS2. — (2) T. 2.2-(b) w. HNO_3 gives Y sol. — (3) T. 2.2-(f) w. Froehde's reagt. gives G or GS1, changing after some min. to YGS2. — (4) T. 2.2-(e) w. Erdmann's reagt. gives GYT1, gradually fading. — (5) T. 2.2-(a) w. conc. H_2SO_4 , colorless. B HNO_3 is obtained as charac. ppt. of micro cryst. on adding 1 drop HNO_3 to 0.01 g. B disd. in 1 cc. aq. + 1 drop dil. HCl . Short pr., m.p. 195° . 100 pt. aq. sol. contain 0.21 pt. at 11.5° . (Its formation has been proposed as mic. test for nitric ac. & nitrates.)
966	185d.	4-Aminonaphthol (2), $\text{NH}_2\text{C}_1\text{H}_8\text{OH}$. — Ndl. fr. alc. Alm. i. bz., chlf.
967	185	m-Aminobenzoylamino-2,4-diaminobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{CO.NH.C}_6\text{H}_4(\text{NH}_2)_2$. — [Pat.] — Ndl. fr. aq. E.s. alc.
968	185-6	Phenyl-bis-p-aminotolylmethane, $\text{Ph.CH[C}_6\text{H}_4(\text{NH}_2)\text{Me}]_2$. — B.p. $427-33^\circ$ sl. d. — Ndl. fr. alc. Pr. w. $1\text{C}_6\text{H}_4$ (lost at 120°) fr. bz. — Mod. s. eth., h. alc., h. bz.; e.s. chlf. — Diacetyl deriv., \ddagger lft. fr. alc., m.p. $217-8^\circ$.
969	184-6.5 <u>c.</u>	Dibenzoylmorphine, $\text{C}_{17}\text{H}_{21}\text{O}_3\text{N}(\text{Ph.CO})_2$. — Cryst. I. aq.; d.s. c. alc. — Protracted boiling w. aq. saps. — B.HCl , amorph.; v.d.s. alc.

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No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
970	186	4-Amino-2,6-dimethylpyridine , $\text{NH}_2\text{C}_6\text{H}_3\text{NMe}_2$. — B.p. 246°. — Ndl. fr. sq. Mod. s. c. aq.; e.s. alc.; d.s. eth., c. bz. — B_2HCl , ndl. e.s. aq. — B_2P_k , \ddagger ndl. d.s. c. aq., alc., m.p. 194–5°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § ppt. of red-yel. pr., d. 250° w. m.
971	186	Anhalamine , $(\text{MeO})_2(\text{HO})\text{C}_6\text{H}_4\text{N}$. — [Fr. Mexican meecal buttons, Anhalonium Lewinii.] Spherical aggregates of mic. ndl. fr. alc. — D_2 Aq. sol. gives blue color w. FeCl_3 , becoming green & then disappearing on warming. — S. aq.; e.s. h. alc., alk.; d.s. chlf., bz.; d.s. pet.-eth., eth. Opt. i. “Strong base.” — $\text{B}_2\text{H}_2\text{PtCl}_6$, § ndl. e.s. h. aq.
972	188–9 u.c. sl. d.; 192–3 u.c. (r.h.)	† Aconitine, Acetyl-benzoyl-aconine , $\text{C}_{14}\text{H}_{17}\text{O}_3\text{N}$. — [Highly toxic alkaloid fr. Aconitum Napellus L.] — Authorities differ widely in regard to m.p. The m.p. is much influenced by rate of heating & by traces of impurities liable to be present. Values for slow heating as low as 182°, and for rapid heating as high as 197–8°, will be found. <i>The value here given was obtained fr. a commercial product brought to constant m.p. by two recrystn. fr. alc.</i> Clear mic. pr. fr. c. alc. S. at 25° in: 3200 pt. aq.; 22 pt. alc.; 44 pt. eth.; 5.8 pt. bz.; 3580 pt. pet.-benzene; v.s. chlf. Odorless. $[\alpha]_D^{25} = +11^{\circ}$ (in 3% alc. sol.). — Unstable, being spdt. even by long heating w. aq. to ac. ac., benzoylaconine, benzoic ac., & aconine. B is extracted fr. sol. of salts by adding NaHCO_3 & shaking out w. eth. — Color reactions of decisive diagnostic value are unknown. <i>Unlike pseudoaconitine (No. 2.991) gives negative results in Vitali's react. (Cf. 2.797(2))!</i> D_2 (1) Physiological T. — Dis. 0.005 g. (5 mg.) B in 12 cc. aq. + 1 drop dil. HCl (sp. gr. 1.12) + 3 drops alc. Bring 1 (one) cc., reserving remainder, into anterior part of mouth & retain there for 1 min. Do not swallow!! (1 mg. has proved fatal!). Rinse out the mouth. After 5 to 15 min. an unpleasant tingling burning sensation will be felt on the tongue, lips & throat! If more than a trace of sol. reaches the posterior portion of mouth, there will also be more or less feeling of strangulation & difficulty in swallowing. (Cf. Flückiger.) — (2) Add 2 drops 1 T.S. in KI to 1 cc. of reserved sol. prepared in (1) above. A floc. yel.-brown ppt. forms. — (3) Add 5 drops KMnO_4 T.S. to sol. of 0.001 g. B in 1 cc. aq. + 1 drop HCl (sp. gr. 1.12). A ppt. of B permanganate is produced which, moist on filter, has OrS_1-RO_1 color. — (4) Dis. 5 mg. of B or of a salt in 0.5 cc. HNO_3 (sp. gr. 1.20). Evap. to dryness. Add a few drops alc. KOH sol. (1 : 5) to c. residue. The sweetish aromatic odor of ethyl benzoate will be observed. Dis. 0.05 g. B in 3 cc. aq. + 2 drops HCl (sp. gr. 1.12). Add 0.5 cc. gold chloride T.S. Filter off curdy yel. ppt. Wash w. 2 cc. aq. Recryel. fr. 5 cc. 50% alc., shaking well & cooling. Filter. Wash w. 1 cc. 95% alc. Dry 15 min. on tile at 90°. The product $\text{B}_2\text{HAuCl}_4\cdot 3\text{H}_2\text{O}$ consists of mic. pr., m.p. 155–6° u.c. (The anhydrous salt is said to melt at 145°.)
973	188d.	Aricine , $\text{C}_{12}\text{H}_{14}\text{O}_2\text{N}_2$. — [Fr. Cusco barks.] — D_2 Colored dark green by conc. HNO_3 . — Pr. fr. dil. alc. I. aq.; v.s. chlf.; s. in 20 pt. eth., or 235 pt. 80% alc. Alc. sol. “scarcely alk.” Taste, not bitter. $[\alpha]_D = -58.18^{\circ}$ (for alc. sol.). — $\text{B}_2\text{H}_2\text{C}_6\text{O}_4\cdot 2\text{H}_2\text{O}$, cryst. ppt. changing to rhombohedra; s. in 2025 pt. aq. at 18°.
974	188–90	7-Aminoquinoline , $\text{NH}_2\text{C}_6\text{H}_4\text{N}$. — Ndl. fr. alc. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yel. cryst. ppt., m.p. abt. 228° d.
975	189.5	1,5-Diaminonaphthalene , $(\text{NH}_2)_2\text{C}_{10}\text{H}_6$. — D_2 FeCl_3 colors aq. suspension blue-violet. — Pr. fr. eth. Sbl. alm. undecd. Alm. i. c. aq.; mod. s. h. aq.; e.s. eth., chlf. — $\text{B}_2\text{H}_2\text{SO}_4$, ndl. alm. i. dil. H_2SO_4 .
976	189	6-Amino-5-methyl-2,4-diethylpyrimidine , Kyanethine , $\text{N}:\text{C}(\text{NH}_2)\text{CMe}:\text{CEt}-\text{N}:\text{CEt}^2$. — Cryst. fr. aq. or alc. S. at 17° in 1365–1380 pt. aq., or 17.6 pt. 90% alc. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § ruby-red octahedra.
977	188–90	4-Dimethylamino-2,2',4'-triaminodiphenylmethane , $\text{Me}_2\text{N.C}_6\text{H}_3(\text{NH}_2)_2\text{CH}_2\text{C}_6\text{H}_3(\text{NH}_2)_2$. — [Pat.]
978	189d.	Vellosine , $\text{C}_{12}\text{H}_{14}\text{O}_2\text{N}_2$. — [Toxic alkaloid fr. bark of Geissospermum Vellosii.] — Tbl. fr. alc. Alm. i. aq.; s. c. chlf., eth.; s. h. alc., bz., lgr. $[\alpha]_D^{25} = +22.8^{\circ}$ (in chlf., conc. 2.703 : 25). A diacidic base w. 2(MeO) groups. — B_2MeI , \ddagger cryst. fr. h. aq., m.p. 264°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yel.-brown cryst., m.p. abt. 80°.
979	191	2,3-Diaminonaphthalene , $(\text{NH}_2)_2\text{C}_{10}\text{H}_6$. — I.f.t. fr. eth. E.s. alc.; mod. s. eth. — Diacetyl deriv., \ddagger m.p. 247°.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
980	190-1	2-Amino-4,4',4''-hexamethyltriaminotriphenylmethane, $(\text{NH}_2)(\text{NMe}_2)_2\text{C}_6\text{H}_4\text{C}_6\text{H}_2(\text{NMe}_2)_2$. — [Pat.] Ndl. fr. toluene. Oxidation of acetyl deriv. by PbO_2 gives blue dyestuff.
981	190-1	Dimethylanilinephthaleine, $[\text{C}(\text{NMe}_2)\text{C}_6\text{H}_4]_2\text{C}_6\text{H}_4\text{CO.O}^{\ominus}$. — Pr. fr. bz. or alc. I. aq.; e.s. bz.; s. eth.; v.d.s. lgr. Evapn. w. conc. HNO_3 gives phthalic ac. (Cf. T. 1.318). — B.2HCl , cryst., losing 1HCl at 100 $^{\circ}$.
982	193 u.c.	† Thebaine, $(\text{MeO})_2\text{C}_6\text{H}_3\text{N}$. — [Tetanic poison fr. opium having narcotic action in small doses.] — <i>Lust. pr. fr. slowly cooling alc. sol.</i> Alm. i. aq.; pet.-eth.; s. in 140 pt. eth. at 10 $^{\circ}$; i. caustic alk.; somewhat s. ammon.; s. in 10 pt. c. alc., in 60 pt. amyl alc., in 18 pt. chlf., in 20 pt. bz. $[\alpha]_D^{25} = -218.64^{\circ}$ (in 2% alc. sol.). Extracted by chlf. or eth. fr. sol. made alk. by NaOH, but not fr. acid sol. Sol. in dil. ac. unstable. — (1) $T. 2.2-(a)$ w. conc. H_2SO_4 gives <i>ROS1-RO</i> color, changing to <i>YO</i> after 1 hr. Erdmann's, Froehde's & Mandelin's reagts. ($T. 2.2-(e,f,g)$ gives same initial color as H_2SO_4). — (2) $T. 2.2-(b)$ w. conc. HNO_3 gives <i>YT1</i> color at once. — (3) $\text{Dis. } 0.05 \text{ g. in } 2 \text{ cc. boiling aq. + 4 drops HCl (sp. gr. 1.12)}$. Add 3 cc. c. sat. aq. picric ac. sol. Wash yel. ppt. on filter w. 1 cc. aq. Recryst. fr. abt. 4 cc. h. 10% ac. ac. Wash w. 1 cc. c. dil. ac. ac. Dry 15 min. at 100 $^{\circ}$ on tile. The resulting picrate is a yel. powd., m.p. 189-91° u.c. d.
983	192-3	Morphothebaine, $\text{C}_{12}\text{H}_{19}\text{O}_3\text{N}$. — [Product of decn. of thebaine by h. conc. HCl.] — May be obtained colorless by crystn. fr. nitrobenzene, but is usually somewhat bluish. Rhomb. cryst. fr. bz. D.s. h. aq.; e.s. alc., eth.; s. alk. — (2) Distinguished fr. thebaine by giving colorless sol. in conc. H_2SO_4 .
984	193.5; 195	Tetramethylbenzidine, $\text{NMe}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_2\text{NMe}_2$. — B.p. a. 360 $^{\circ}$. D.s. c. alc., eth., e.s. h. bz. — NaNO_2 colors HCl sol. yel.-red., & NaOH then gives ppt. of green flocks soon becoming brown-yel.
985	195	N-Methyldiphenyleneimidazole, Epiosin, $\text{C}_{10}\text{H}_{12}\text{N}_2$. — [Lessens sensibility to pain like morphine.] — Pr. e.s. alc. — B.HCl , pr. e.s. aq.
986	198	Carbotriphenylamine, Diphenylamino-p-aminobenzylamidine, $\text{NH}_2\text{C}_6\text{H}_3\text{C}(\text{:N.Ph})(\text{NH.Ph})$. — 4-sided tbl. I. aq.; d.s. eth. Dec. on distn. to NH_2 , HCN , Ph.CN , & Ph_2NH . — (2) Test distillate for diphenylamine (No. 2.1568). — B.HCl , cryst. s. h. aq.
987	198 (dried at 125)	Cupreine, $\text{C}_{18}\text{H}_{22}\text{O}_2\text{N}_2$. — [Fr. bark of Cinchona cuprea or Remijia pedunculata.] Cryst. w. $2\text{H}_2\text{O}$ in pr. fr. eth. I. aq.; d.s. eth., chlf.; s. alc. Pptd. by alk. fr. salts, but s. x.s. KOH & not extracted fr. the sol. by eth. $[\alpha]_D^{25} = -175.5^{\circ}$ (0.2354 g. in 19 cc. alc.). — (2) Gives "Thalleioquin react." like quinine (No. 2.947), but unlike quinine does not give blue fluor. in dil. H_2SO_4 sol. — For color react. w. H_2O_2 & ammon., cf. Denigès, Compt. rend., 151, 1354. — Neut. reacting salts, B.HX , give yel. sol. in h. aq.; salts, B.2HX , are colorless.
988	199	Acetylbenzidine, $\text{NH}(\text{CO.Me})\text{C}_6\text{H}_4\text{C}_6\text{H}_2\text{NH}_2$. — Ndl. fr. dil. alc. E.s. alc.; i. eth. "Combines w. acids."
989	199	† Bulbocapnine, $(\text{MeO})(\text{HO})_2\text{C}_6\text{H}_3\text{N}$. — Cardiac poison fr. root of <i>Bulbocapnus cavus</i> . — Rhomb. cryst. fr. alc. S. in usual solvents except aq. S. alk. (in NaOH w. greenish color) & repptd. by CO_2 . May be extracted fr. alk. sol. by eth. $[\alpha]_D = +237.1^{\circ}$ (for 0.6722 g. in 15 cc. chlf.). — A commercial preparation fr. Schuchardt (m.p. 195° u.c.) in $T. 2.2-(a)$ gave a colorless sol., quickly changing to <i>OY</i> bordered by <i>BVT1</i> , changing in 5-10 min. to <i>V1</i> , & in 30 min. <i>RV1</i> — B.Mel , lust. ndl. fr. h. aq., m.p. 257°.
990	abt. 200	Lanthopine, $\text{C}_{18}\text{H}_{22}\text{O}_2\text{N}$. — [Fr. opium.] — Mic. pr. D.s. alc., eth., bz.; mod. s. chlf.; s. NaOH, but not in ammon. — Sol. in H_2SO_4 colorless. No coloration w. FeCl_3 . — $\text{B.2H}_2\text{PtCl}_6\cdot 2\text{H}_2\text{O}$, lemon-yel., i., cryst. powd.
—	201	4-Hydroxyquinoline. — Cf. No. 2.2456.
991	201-2 sl. d.; (210-2)	Pseudoaconitine, Nepaline, Acetylveratroylpseudoaconine, $\text{C}_{20}\text{H}_{26}\text{O}_2\text{N}$. — [Fr. root of Aconitum ferox, Indian aconite. Perhaps the most poisonous of alkaloids.] — Ndl. fr. eth. Cryst. w. aq. lost at 80 $^{\circ}$. Alm. i. aq.; s. alc., eth.; little s. KOH. Extracted by eth. fr. ammon. mixt. — B , dextro-, & salts levo-rotatory. — (2) (1) Like aconitine causes intense tingling sensation when v. dil. sol. is brought in contact w. tongue & lips in "Physiological T.

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(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		(1) " described under No. 2.972! (Test only w. greatest caution, the dose being only 0.4 that of aconitine!) — (2) Unlike aconitine gives purple-red color in Vitali's react.! (Cf. No. 2.797). — (3) Cautiously warmed w. conc. H_2SO_4 gives violet-red color (Z. anal. Chem., 23, 238). — $B.HNO_3$, $3H_2O$, cryst., d.s. aq., m.p. 185–6°. — $B.HAuCl_4$, yel. ndl., m.p. 236–8°.
992	202	Stylopine, $C_{16}H_{21}O_4N$. — [Fr. Stylophorum diphyllo.] — Ndl. fr. eth. V.s. gl. ac. ac. $[\alpha]_D = -315^{\circ}$, 12' (in abs. alc.). — $B_2H_4PtCl_6$, §
993	202.4; (201 u.c.)	† Cinchonidine, $C_{18}H_{21}ON_3$. — [Fr. Cinchona bark.] Clear lust. trimetric pr. fr. h. alc. S. in 5263 pt. aq. at 11.5°; in 1782 pt. boiling aq.; in 21.1 pt. 98% alc. at 11.5°; in 188 pt. eth. (sp. gr. 0.72). Odorless. Taste, bitter. $[\alpha]_D^{25} = -107.5^{\circ} + 0.297 c$. (1 < c < 5), in 97% alc. — (2) Apply "Herapathite T." described for quinine (No. 2.947). The YGT% ppt. of mic. ndl. is without metallic luster. Unlike quinine gives no fluorescence in "Fluorescence T." & no color in "Thalleioquin T." — T. 2.2-(a,e,f), w. H_2SO_4 , w. Erdmann's or w. Froehde's reagts. give no colors. — (3) Prepare the picrate (not analyzed) as directed for quinine (No. 2.947). Recryst. by slow cooling fr. sol. in 6 cc. boiling 10% ac. ac., & wash w. 1 cc. solvent. The product is obtained as Y-ndl., m.p. 208–9° u.c. d., darkening fr. abt. 200°.
994	202–3	Indaconitine, $C_{21}H_{21}O_{10}N$. — [Highly toxic alkaloid of aconitine-like action fr. Aconitum Chasmantum.] — Ndl. I. aq., pet.-eth.; s. alc., eth., chlf. $[\alpha]_D^{25} = +18^{\circ}$, 17' (in alc., c = 2.1–2.3). Contains 4MeO groups. — $B.HNO_3$, pr. fr. alc. + eth., m.p. 202–3°.
995	203–4 (browning)	3,3'-Dimethyl-4,6,4',6'-tetraaminodiphenylmethane, [Me.C ₆ H ₄ (NH ₂) ₂ .CH ₃ . — Lst. fr. aq. D.s. alc., toluene, h. aq.
996	d. 204 after darkening fr. 200	2-Amino-1,6-dihydropurine, Desoxyguanine, $C_9H_{11}N_5$. — Ndl. fr. aq. — (2) V.s. aq. — D.s. alc. Reacts alk. Absorbs CO_2 fr. air. $FeCl_3$ colors sol. brown-red. — $B.2Pk$, † or-yel. rhombohedra s. in 200–250 pt. h. aq.
997	202–4; 208–14; 204–5 u.c.	† Oxyacanthine, $C_{18}H_{21}O_4N$. — [Fr. Berberis vulgaris.] — Ndl. fr. alc. or eth. E.s. chlf., bz.; v.d.s. lgr. $[\alpha]_D^{25} = +131.6^{\circ}$ (in chlf., p = 4). — (2) (Based on a commercial preparation.) (1) T. 2.2-(a) w. H_2SO_4 gives colorless sol. — (2) T. 2.2-(b) w. HNO_3 gives YOSI color. — (3) T. 2.2-(f) w. Froehde's reag. gives RV at once, changing to B in 15 min. — (4) T. 2.2-(g) w. Mandelin's reag. gives v. dark brown at once. — (5) Prussian blue T. as described for morphine (No. 2.1024) gives heavy blue ppt.
998	203.5–4.5	Japaconitine, $C_{21}H_{21}O_{11}N$. — [Highly toxic alkaloid of aconite-like action fr. Aconitum japonicum.] Ndl. fr. alc., eth., chlf. V.s. alc., chlf.; alm. i. pet.-eth. $[\alpha]_D^{25.5} = +23.6^{\circ}$ (in alc., p = 0.605). — (2) Physiological test (great caution!!) gives results described under aconitine (No. 2.972) & pseudoaconitine (No. 2.991). — $B.HAuCl_4$, amorph. ppt. fr. $B.HCl$, when disd. in alc. & allowed to stand for short time, m.p. 231°; or in β -form, m.p. 154–60°, when chlf. sol. is left to spontaneous evapn.
999	205	Lappaconitine, $C_{21}H_{21}O_8N_2$. — [Toxic alkaloid fr. Aconitum septentrionale.] — Hexag. cryst. D.s. eth., giving dextrorotatory sol. w. strong red-violet fluor.! Taste, bitter, but not sharp. Colors vanadium-sulphuric ac. yel.-red & then green.
1000	205; (203–4 u.c.)	† Cevadine, "Veratrinum purissimum crystallisatum," $C_{21}H_{21}O_8N$. — [Highly toxic alkaloid fr. sabadilla seeds, Schoenocaulon officinale, Asa Gray. Commercial "Veratrine" is a mixture containing this alkaloid.] — Fine ndl. fr. 66% alc. on spontan. evapn.; sometimes w. 2 mol. alc. of cryst. (then efflorescing in air). S. in 1000 pt. boiling aq.; i. c. aq.; s. in 10–12 pt. c. alc.; s. eth., chlf., amyl alc., bz.; d.s. pet.-eth. Taste, (greatest caution, v. poisonous!) acid, burning, leaving tingling sensation. Odorless, but traces of the dust provoke sneezing. Opt. i. — (2) (1) T. 2.2-(a) w. conc. H_2SO_4 gives YO sol., changing in 5 min. to OR, & in 30 min. to brilliant VRT1–RT1. — (2) Weppen's T. — Rub well together in small mortar 0.002 g. B + 0.010 g. conc. sugar sol. Place on crucible cover & moisten w. 2 drops conc. H_2SO_4 . A Y color appears, soon changing to bright G & BG on stirring. Within 5 min. mixt. appears BS ² , or bright B, according to thickness of layer. — (3) HCl T. — Heat 0.001 g. B w. 2 drops HCl (sp. gr. 1.20) on crucible cover on steam-bath for 1 min. A stable ORT1 color develops. (Gadamer states it persists for a week.) 0.001 g. B heated w. 1 cc. conc. HCl for 15 min. on boiling water-bath gives OR-R sol. — (4)

No.	Melting-point (C°).	BASIC COMPOUNDS.—Colorless and Solid.
		Fluorescence T. — Dis. 0.001 g. in 1 cc. conc. H_2SO_4 . An olive-green fluor. will be observed, as well as colors mentioned under (1) above. — (5) Vitali's React., applied as directed for atropine (No. 2.797), except that 0.002 g. \tilde{B} is used, gives VR color, changing almost immediately to R. Evapn. of the alc. on water-bath gives coniine-like odor.
		① (a) $B.HCl.HgCl_2$: Dis. 0.05 g. \tilde{B} in 2 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 5 cc. sat. aq. $HgCl_2$ sol. Wash heavy ppt. on filter w. 2 cc. aq. Recryst. fr. 4 cc. boiling 50% alc. White silvery lf. separate slowly. Filter. Wash w. 1 cc. dil. alc. Dry on tile 15 min. at 100°. Product melts at 171.5–2.5° u.c. to light yel. pasty mass. — (b) $\tilde{B}.HAuCl_4 $ (dried at 100°). Cryst. w. $2H_2O$. Dis. 0.05 g. \tilde{B} in 2 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 4 cc. $HAuCl_4$ T.S. Filter. Wash ppt. w. 1 cc. aq. Recryst. fr. 5 cc. boiling 50% alc. Wash the short slender ndl. on filter w. 1 cc. 50% alc. Dry on tile 15 min. at 100°. Product melts to dark brown liq. at 182° u.c. after darkening fr. 178–9°.
1001	205–6	† Aspidospermine , $C_{22}H_{30}O_4N_2$. — [Fr. Aspidosperma Quebracho.] — Ndl. fr. alc. S. in 6000 pt. aq. at 14° (taste of sol. bitter); s. in 48 pt. abs. alc. at 14°; less s. eth., lgr., pet.-eth.; e.s. bz., chlf. "V. weak base." May be extracted fr. sol. of salts by eth. or chlf. $[\alpha]D^{15} = -100.2^\circ$ (in 97% alc. p = 2). — ② Gives colorless sol. in conc. H_2SO_4 which on addition of drop $K_2Cr_2O_7$ sol. shows brown zone, slowly changing to olive-green.
1002	206	Guanazole , $^2NH.C(:NH).NH.NH.C(:NH)^2$. — Monoclin. pr. fr. aq. — ② V.s. aq. — $FeCl_3$ gives intense red coloration. — S. alc.; i. eth., chlf., bz. "Reacts feebly alk." — $B.Pk$, † lust. yel. tbl., d.s. aq., alc., m.p. 245°.
1003	206u.c. (r.h.)	Echitamine , $C_{22}H_{30}O_4N_2H_2O$. [Toxic alkaloid fr. Echites scholaris L., Philippine dita bark.] — Glassy pr. w. $4H_2O$ fr. alc. or acetone + aq. $B.H_2O$, for which m.p. is given, was obtained by drying at 80° (Ann. 203, 147). — S. aq.; more s. alc.; v.d.s. bz.; i. lgr.; mod. s. chlf., eth., when freshly pptd. "Reacts alk." Taste v. bitter (Caution!). $[\alpha]D^{15} = -28.8^\circ$ (in 97% alc. sol., p = 2). — ② Sol. in conc. H_2SO_4 , intense purple-red! Sol. in conc. HNO_3 , red, soon fading & changing to intense green. — $B_2H_2PtCl_6.3H_2O$, § yel. floc. ppt., v.d.s. c. aq.
1004	206	Xanthaline , $C_{27}H_{38}O_4N_2$. — [Fr. opium.] — Cryst. powd. I. aq., alk.; d.s. h. alc. — ② S. in conc. H_2SO_4 w. deep red color! Sol. in dil. acids give salts of yel. color. — $B.2HCl.4H_2O$, yel. ndl.
1005	207 204 or 208c.	Protopine , Macleyine , $C_{20}H_{31}O_4N$. — [Fr. opium, Macleya cordata, etc.] — Ndl. fr. eth. I. aq.; d.s. alc., ac.-eth. Cryst. s. in 1000 pt. eth. Opt. i. — ② H_2SO_4 dis. w. fine blue-violet color, changing to muddy violet, green on margin. — Erdmann's reag. gives yel., blue-violet, blue, green, & yel. colors. — $B.HAuCl_4 $ red brown amorph. powd., m.p. 198°.
1006	207.6	Homocinchonidine , $C_{16}H_{21}ON_2$. — [Fr. Cinchona bark.] — Cryst. fr. alc. S. in 20.5 pt. 97% alc. at 13°; in 216 pt. eth. (sp. gr. 0.72) at 15°; alm. i. aq.; v.s. chlf. "Alc. sol. strongly alk." $[\alpha]D = -107.3^\circ$ (in 97% alc. p = 2). — Acid sol. not fluor. & does not give green color in Cl-aq. & ammon. test for quinine (No. 2.947). For micro-chem. tests cf. Z. anal. Chem., 35, 134.
1007	d.207–7.5	d-Arginine , δ-Guanidine-α-aminovaleric Ac. , $HN : C(NH_2).NH.(CH_2).CH-(NH_2).CO_2H$. — [Proteolytic product & in germinating plants.] — Rosettes, tbl. & thin pr. E.s. aq.; alm. i. alc. Reacts alk. ($k.10^7$ less than 1.0). Odorless. Taste faintly bitter. Dextrorotatory. — ① (a) Picrolonate, $B_2C_{10}H_{14}O_4N_2H_2O$, sulphur-yel. ndl. fr. h. aq., s. in 1124 pt. aq., or 2885 pt. 96% alc., m.p. 225°. (Z. physiol. Chem., 44, 157, 49, 215). — (b) Basic copper nitrate compound, $B_2Cu(NO_3)_2.3$ or $3\frac{1}{2}H_2O$, is prepared by boiling sol. of nitrate w. $CuCO_3$. Spheroidal aggregates of dark blue monoclinic ndl. or thin pr. s. in 95.2 pt. c. aq. M.p. (hydrated) 112–14°; (anhydrous) 232–4° d. — $B.Pk.2H_2O$, † silky yel. ndl., m.p. 205–6°. 100 pt. aq. at 16° dis. 0.5 pt.
1008	209	α-Aminoacridine , $NH_2C_12H_8N$. — Ndl. fr. aq. (prob. yellowish). E.s. alc., eth. w. green fluor.
1009	210d.	Apoquinine , $C_{19}H_{22}O_2N_2$. — [Fr. action of h. aqueous HCl on quinine.] — Cryst. w. $2H_2O$. Ndl. fr. eth. D.s. c. aq., e.s. h.; s.c. alc.; e.s. eth., chlf., bz.; v.s. KOH. — Unlike quinine (No. 2.947) does not give fluor. sol. in x. s. dil. H_2SO_4 , or green color in Thalleioquin reaction. — $[\alpha]D = -178.1^\circ$ (in 97% alc. p = 2).

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No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
1010	211	2-o-Aminophenylbenzimidazole, $[C(C_6H_4NH_2) : N.C_6H_4.NH]$. — Pr. fr. alc. I. c. aq.; e.s. h. alc., chlf.; less s. bz., eth. — $\bar{B}.2HCl$, d.s. h. aq., alc., m.p. 275° d.
1011	211; 211-3	† Sanguinarine, $C_{20}H_{15}O_4N$. — [Fr. Sanguinaria canadensis & Chelidonium majus.] — Cryst. fr. eth. in colorless plates which may contain aq. or alc. Reddens w. decn. on exposure to air. S. alc., eth., chlf. — \oplus Salts are blood-red! — Solutions, particularly of sl. impure \bar{B} , show blue-violet fluor. — Color reactions of T. 2.2 are: w. conc. H_2SO_4 , dark red-yel. color; w. conc. HNO_3 , brown-yel.; w. Erdmann's reagt., fine or.-red, slowly becoming turbid; w. Froehde's reagt., dark brown-yel., red-yel., & finally muddy brown. — $\bar{B}.HCl.5H_2O$, red ndl. — $\bar{B}.HAuCl_4$, brown-red, heavy floc. ppt.
1012	d.212u.c	Adrenaline, Suprarenine, 3,4-Dihydroxyphenyl-1-methylaminoethanol(2), $(HO)_2C_6H_3.CH(OH).CH_2.NHMe$. — [Fr. the adrenal capsule.] — Colorless ndl. Aq. at 20° dis. 0.0268%, somewhat more s. h.; less s. alc.; i. chlf., pet.-eth.; e.s. ac., alk. $[\alpha]D^{19.8} = -51.40^{\circ}$ (in aq. + HCl). — \oplus $FeCl_3$ in acid sol. gives green, or in alk. sol., carmine-red color. — In presence of sulfanilic ac. $FeCl_3$ gives intense red-brown color at high dilution. (Biochem. Z., 20, 178 (1909)). — Reduces ammon. $AgNO_3$ in the cold. — Warmed w. chloral & KOH sol. gives nauseous carbylamine odor. — Added to dil. aq. sol. of iodic ac. w. chlf., the chlf. becomes rose colored on shaking. (Biochem. Z., 12, 131 (1909)).
1013	214	Bebeanine (Beberine), Pelosine, $C_{18}H_{21}O_4N_2$. — [Toxic alkaloid of curari-like action fr. bark of Nectandra Rodiei.] — Glassy cryst. fr. MeOH. Separates amorph. fr. chlf. or acetone w. m.p. 180°.
1014	214-6d.	Quebrachine, $C_{21}H_{28}O_4N_2$. — [Fr. bark of Aspidosperma Quebracho.] — Colorless ndl. fr. alc., becoming yel. on keeping. Alm. i. aq., NaOH, ammon.; e.s. h. alc., chlf.; d.s. c. alc., eth., lgr. Alc. sol. "reacts alk." & tastes v. bitter. Dextrorotatory. — \oplus Sol. in conc. H_2SO_4 at first alm. colorless, later bluish, & on adding little PbO_2 becomes deep blue! — $\bar{B}_2.H_2PtCl_6.5H_2O$, § yel. amorph. ppt.
1015	d.w.m.212-8	8-Aminonaphthol(2), $NH_2.C_10H_8(OH)$. — Ndl. fr. eth. E.s. aq., alc., eth. — \oplus Colored green-blue by little $FeCl_3$. — Salts fluor. violet-blue.
1016	210-20	† Cyanoaniline, $C_6H_5N_2$. — Lft. i. aq.; d.s. alc., eth. Decd. by boiling acids to aniline, NH_3 , oxanilide, etc.; but not attacked by aq. alk.
1017	216; 216-8	2,6-Diaminonaphthalene, $(NH_2)_2C_10H_6$. — Lft. fr. aq. V.d.s. h. aq.; d.s. alc., eth. — \oplus Aq. sol. colored green in the cold, or blue when hot, by $FeCl_3$.
1018	217u.c.	Cryptopine, $C_{21}H_{22}O_4N$. — [Fr. opium.] — Pr. or tbl. fr. alc. I. aq., alk.; alm. i. eth., bz., lgr.; e.s. chlf.; s. in 80 pt. 95% boiling alc., or in 455 pt. at 15°. Opt. i. — \oplus In color reactions of T. 2.2: H_2SO_4 gives dark blue-violet, soon changing to green & later to yel.; Erdmann's reagt., violet-pink, changing to gray & yel.; Froehde's reagt., intense violet, changing to blue-green, green, & after 2 hr., yel.; Marquis' reagt., violet, changing to brown. — $\bar{B}.Pk.H_2O$, § hair-like ndl., v.d.s. h. aq., m.p. 215°. — $\bar{B}_2.H_2PtCl_6$, § v. pale yel. ndl. w. 1 or $6H_2O$, m.p. 204°.
1019	219d. (browning & sintering fr. 210)	Ergotinine, $C_{20}H_{20}O_4N_4$. — [Fr. ergot.] — Colorless ndl., sensitive to light, soon turning yel.-brown. I. aq.; s. in 200 pt. 95% alc. at 20°, or 50-60 pt. at boiling temp.; s. in 150 pt. boiling bz. "Weak base." $[\alpha]D = +334-6^{\circ}$ (in alc. sol.). — \oplus Acid solutions fluor. violet & soon becomes red. — When treated w. ethyl acetate & conc. H_2SO_4 gives or.-red color which changes through violet to blue!
—	abt. 221d.	o-Aminobenzaldehydephenylhydrazone. — Cf. No. 2.3751.
—	225-6	Cyanphenylhydrazine. — Cf. No. 2.2549 (S. in conc. H_2SO_4 w. indigo-blue color).
1022	229	Aribine, $C_{22}H_{20}N_4$. — [Fr. bark of Brazilian Arariba rubra.] — Rhomb. pyramids fr. dry eth.; or in 4-sided pr. w. $8H_2O$ fr. wet eth. S. in 7762 pt. aq. at 23° more s. h.; v.s. alc.; less s. eth. Opt. i. — $\bar{B}.2HCl$, pr. e.s. aq. — $\bar{B}_2.H_2PtCl_6$, § ppt. of pale yel. ndl.
1023	229-30	Cinchamidine, Hydrocinchonidine, $C_{19}H_{24}ON_2$. — 6-sided lft. fr. dil. alc. Alm. i. aq.; d.s. eth.; v.d.s. h. chlf.; i. NaOH. Alc. sol. reacts alk. $[\alpha]D^{16} = -98.4^{\circ}$ (in 97% alc., p = 2). Salts taste v. bitter. Sol. in dil. H_2SO_4 not fluor.

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
1024	230 ± 2 u.c., d.	<p>† Morphine, $C_{17}H_{21}O_3N.H_2O$. — [Fr. opium.] Pr. or ndl. fr. h. alc., permanent in air (losing cryst. aq. abt. 100°). The rhombic pr. obtained fr. amyl alc. are figured in Witthaus, p. 944. The m.p. or decn. temp. here given was determined by the usual method w. temperature rising 2° per min. from some degrees below fusion point. The constant is much influenced by the rate of heating, so that, starting with a hot bath, and heating towards the last at a 5° per min. rate, the m.p. lies at $252^{\circ} \pm 2^{\circ}$ u.c. S. in 3330 pt. aq., 168 pt. alc., 100 pt. lime-aq., 4464 pt. eth., 1800 pt. chlf., 113.5 pt. amyl alc., or 525 pt. ac.-eth. at 25°. S. in 1040 pt. aq. at 80°, in 76 pt. alc. at 60°; i. bz. (U.S.P.). S. in 117 pt. ammon. (sp. gr. 0.97). E.s. caustic alk. sol. Aq. sol. alk. to red litmus. Odorless. Taste, v. bitter. $[\alpha]D^{25} = -130.9^{\circ}$ (in MeOH, $c = 2.292$).</p> <p>⊕ Witthaus (p. 984) states the opinion, that the first seven of the following color reactions "Are the most prominent, and are sufficient, we believe, when yielding the reactions distinctly & unmistakably, to identify morphine."</p> <p>(1) Ferric Chloride or Robiquet T. — Add to 1 cc. 1% sol. of B. HCl in a t.t., or to a small fragment of B on white porcelain, 1 drop 1% aq. sol. of sublimed $FeCl_3$ (i.e., a chloride free fr. x.s. HCl). A GBS1-B Bkn. color appears, which is destroyed by heat, alc., acids, or caustic alk. (Wormley found the $FeCl_3$ react. distinct w. 0.0065 mg. solid B, but unsatisfactory w. sol. containing 0.13 mg.) — (2) Froehde's Reagt. T. [T. 2.2-(f)]. — Performed on porcelain w. 0.2–0.3 mg. B gives first a VRS1, changing after 10 min. to broken G w. YG margin, & after 40 min. to G Y. [Delicacy 0.005 mg. (Dragendorff), or 0.0065 mg. (Wormley).] — (3) Pellarri's T. — Dis. 0.002 g. in 6 drops conc. HCl (sp. gr. 1.20). Add 3 or 4 drops conc. H_2SO_4. Evap. on water-bath, & continue heating 15 min. when dry. Dis. the sometimes purplish residue in 2–3 cc. aq. Add 3 drops dil. HCl (sp. gr. 1.12). Neutralize w. abt. 3 cc. sat. aq. $NaHCO_3$ sol. Add 6 drops of I sol. (25 gr. I in 500 cc. alc.). Shake several min. A GS2-BGS2 sol. is obtained. Extract w. eth. The eth. becomes VRT3, while aq. sol. remains green. — (4) Husemann's T. — Treat 0.002 g. B on porcelain crucible cover w. 4 or 5 drops conc. H_2SO_4. Heat 10 min. in 100° oven. Cool. Touch center of liquid w. stirring rod moistened w. dil. HNO_3 (sp. gr. 1.20). An intense R-VR, persisting for some min. & then changing to OR fr. center outward, and after 15 min. to O, appears. [Delicacy 0.01 mg. (Ann. 128, 305).] — (5) Nitric Ac. T. [T. 2.2-(b)]. — Gives at first O, and after a min. or two, YO. — (6) Iodic Ac. or LeFort's T. — Dis. 0.002 g. in 2 cc. dil. H_2SO_4 (sp. gr. 1.175) & shake w. 2 cc. chlf. & a few drops c. sat. aq. iodic ac. sol. I will be liberated, coloring the chlf. VRT1. — (7) Marquis' or Formaldehyde-Sulphuric Ac. T. — Mix 2 or 3 drops 40% formaldehyde sol. w. 3 cc. pure conc. H_2SO_4. Add 2 or 3 drops of this reag., freshly prepared, to 0.2 to 0.3 mg. B on porcelain & stir. The color is at first VR, but becomes bluer, and after 15 min. is RV-RVS1. — (8) Prussian Blue T. — Add 1 drop 1% $K_3Fe(C_6H_5)_2$ sol. & 1 drop 10% $FeCl_3$ sol. to a few mg. B suspended in 1 cc. aq. An immediate pptn. of Prussian blue is obtained. — (9) T. 2.2-(a) w. conc. H_2SO_4 gives colorless sol. — (10) Furfurol T. — This test [the test w. H_2SO_4 & sugar, described as (6) under No. 2.903] gives same colors as codeine.</p>
1025	231	4-Hydroxyquinidine, $HO.C_{10}H_8N$. — Lust. pr. fr. aq. Cryst. w. $2H_2O$, lost at 110° . S. in 100 pt. c., or 10 pt. boiling aq.; e.s. alc.; v.d.s. eth., bz., lgr. — ⊕ Aq. sol. colored intensely red by $FeCl_3$; — Picrate, † light yell. ndl. fr. aq., m.p. 200° . — $B_2H_6PtCl_6$, § yell. ndl. d.s. c. aq., m.p. 215° d.
1026	232d.	Rhoadine, $C_{11}H_{12}O_2N$. — [Fr. Papaver Rhocas & opium.] — Pr. alm. i. aq., chlf., bz., ammon.; s. in 1100 pt. c. 80% alc., or 1280 pt. eth. at 18° . Sol. of B or salts do not taste bitter. — ⊕ S. in "moderately conc." HCl or H_2SO_4 w. purple-red color. (Delicate!) Conc. H_2SO_4 gives olive-green sol.
1027	232–3	Gnoskopine, $C_{20}H_{21}O_3N$. — [Fr. opium.] — Cryst. by adding several volumes alc. to sat. chlf. sol. S. in 1500 pt. c. alc., e.s. chlf., bz.; i. alk., fusel-oil. Blues red litmus in alc. suspension. — ⊕ S. in conc. H_2SO_4 w. yell. color, changed to carmine-red by trace HNO_3 . — $B_2H_6Cl_3H_2O$, flat pr., m.p. 238° d. — Picrate, † yell., m.p. 185° . — Picrolonate, yell.-brown, m.p. 232° d.
—	233 ± 2 u.c.	Glycocol. — Cf. No. 2.2568. (V.s. aq. Taste, sweet.)
1028	234	3-Aminonaphthol(2), $NH_2.C_{10}H_8(OH)$. — Ndl. fr. aq.; e.s. alc.; d.s. eth., bz.
1029	234–4.5	Yohimbine (Anhydroyohimbine), $C_{19}H_{21}O_3N_2$. — [Aphrodisiac alkaloid fr. bark of Corynanthe Yohimbe.] — Sometimes cryst. w. $1H_2O$. Ndl. v.d.s. aq.;

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No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		v.s. alc., eth., chlf.; s. bz. Causes brief sensation of numbness on tongue. $[\alpha]_D = +1.5^{\circ}$ (in 1% aq. sol.). — (1) Dis. in conc. in H_2SO_4 , on porcelain, & draw rod moistened w. dil. $K_2Cr_2O_7$ sol. quickly through sol. A blue-violet color, changing to green appears. — (2) Freshly prepared Froehde's reagt. [T. 2.2-(f)] gives at once a gray-blue, changing to dark blue! — (3) Millon's reagt. gives deep brown-red. — $B.HNO_3$, cryst. easily, s. to abt. 0.9% in c., or 5% in boiling aq.; m.p. 276°. — [For detection in tablets, cf. Ber. d. deutsch. pharm. Gesell., 22, 380 (1912)].
1030	235; 236	p,p'-Diaminotolane, $NH_2.C_6H_4.C : C.C_6H_4.NH_2$. — Pale yellowish ndl. fr. h. abs. alc. — Boiled w. acetic anhydride gives diacetyl deriv., m.p. 270°, which gradually turns blue in light.
1031	237-8	Corybulbine, $(MeO)_2C_6H_4ON$. — [Fr. root of Corydalis cava.] — Cryst. powd. fr. alc. Turns yel. in light. I. aq., eth.; d.s. alc.; e.s. chlf., h. bz.; s. NaOH. $[\alpha]_D^{20} = +303.3^{\circ}$ (0.35-6 g. in 25 cc. chlf.). — $B.HCl$, colorless pr., m.p. 245-50° d.
1032	238d.	† Harmaline, $C_{12}H_{14}ON_2$. — [Fr. Russian Peganum harmala.] — Thick colorless cryst. fr. alc.-bz. Yellow to honey-yellow in thick layers. Taste, bitter. — (1) Sol. in conc. H_2SO_4 intense yel. without fluor. Alc. sol. of salts fluor. green. — $B.HCl.2H_2O$, yel. ndl. — Acetyl deriv. (fr. B disd. in 3-4 pt. ice-cold pyridine + acetyl chloride), $C_{12}H_{14}ON_2(Me.CO)$, ndl. fr. alc., m.p. 204-5°.
1033	238-40 u.c. d.	Sabadine, $C_{20}H_{24}O_4N$. — [Fr. sabadilla seeds.] — Ndl. fr. eth. S. in eth. when freshly pptd.; cryst. d.s. eth. — (1) 0.001 g. boiled w. 5 cc. conc. HCl gives VT1 sol. w. greenish fluor.! — 0.0005 g. gives Y sol. in 1 cc. conc. H_2SO_4 w. greenish fluor., becoming red after some hr. (Test made w. commercial preparation.) — $B.HNO_3$, ndl. fr. h. aq., s. in 131 pt. aq. at 13°, m.p. 308°.
1034	241	Jervine, $C_{20}H_{27}O_3.2H_2O$. — [Toxic alkaloid fr. Veratrum album.] — Ndl. fr. alc. S. alc., amyl alc., chlf.; v.d.s. eth.; i. bz., pet.-eth. Ammon. ppts. in delicate ndl. fr. sol. of salts. — (1) Sol. in conc. H_2SO_4 yellowish, then green & impure green. — $B.HAuCl_4$, yel. ppt.
1035	240.6; 236c.	Rubijervine, $C_{20}H_{24}O_4N.H_2O$. — [Fr. Veratrum album.] — Cryst. fr. h. alc. — (1) Sol. in conc. H_2SO_4 w. yel. color, changing to orange-red & dark red. — $B.HAuCl_4$, yel. ppt.
1036	243-4	Calycanthine, $C_{11}H_{14}N_2$. — (Dried at 120°.) — [Fr. seeds of Calycanthus glaucus.] — Brilliant orthorhombic bipyramids fr. aq.-acetone w. $\frac{1}{2}H_2O$ (m.p. 216-8°). Alm. i. aq.; s. eth., chlf.; d.s. bz. — (1) HNO_3 , dis. w. fine green color. — (2) Froehde's reagt. gives first yel., & after 1 hr. a nearly red color. — (3) 1 or 2 drops of 5% gold chloride sol. added to trace of B disd. in v. dil. HCl , & sol. made alk. w. Na_2CO_3 gives fine purple color at once. (Delicacy 1:1,000,000). — (4) Mandelin's reagt. gives fine blood-red color, turning green on margin after a few min. — $B.Pk\frac{1}{2}H_2O$ is prepared by adding x.s. sat. aq. picric ac. sol. to 1% sol. of B in 1% H_2SO_4 & enough aq. to give clear sol. on heating, setting aside for 48 hr., washing, & drying at 25°. The picrate forms long, silky yel. ndl. which, dried in vacuo over H_2SO_4 , melt at 186-7°.
1037	247d.	3-Aminocarbazole, $NH_2.C_9H_9N$. — Ndl. fr. aniline. Blackens abt. 240°. — Prim. amine.* — $B.HCl$, s. aq.; i. conc. HCl , colors lignin red.
1038	245-50d.	Protoveratrine, $C_{20}H_{24}O_4N$. — [Fr. Veratrum album.] — Thin 4-sided plates fr. abs. alc. S. chlf., h. abs. alc.; v.d.s. c. eth.; i. bz., aq., pet.-eth. Tasteless (v. poisonous!), numbing the tongue. Dust provokes sneezing! — (1) Conc. H_2SO_4 gives successively yel., green, greenish blue, blue, & blue-violet colorations. — Conc. HCl on warming gives rose-red & red color. — $B.HAuCl_4$, unstable gold-yel. ppt.
1039	d. abt. 250°	Lycorine, $C_{16}H_{22}O_3N_2$. — [Toxic alkaloid fr. Japanese Lycoris radiata Herb.] — Cryst. fr. alc. Becomes yel. at 235°. V.d.s. aq., alc., eth., chlf.; e.s. acids. — (1) Sol. in conc. H_2SO_4 colorless, and then ochre-yel. — Conc. HNO_3 gives brown-yel. sol. — Conc. H_2SO_4 + sodium molybdate gives green, & then blue color. — $B.2HCl.2H_2O$, ndl. s. aq., m.p. 208°.
1039-I	250 sl. d.; 255.4	Cinchonine, $C_{16}H_{22}ON_2$. — [Fr. Cinchona barks.] — Clear pr. or ndl. Sbl. fr. below 200°! — S. at 20° in: 3670 pt. aq. (2500 pt. boiling); 100 pt. 90% alc.; 371 pt. eth.; 280 pt. chlf. I. pet.-eth., caustic alk. or ammon. $[\alpha]_D = +223.3^{\circ}$ (0.1-0.15 g. in 20 cc. abs. alc.) — Diacid base. $[k_B \times 10^3 = 1$;

No.	Melting-point (C.).	BASIC COMPOUNDS.—Colorless and Solid.
		second $k_B \times 10^{10} = 3.3$. — The general alkaloid reagents (T. 2.3) ppt. fr. v. dil. sol. Thus, in sulphate sol. ppt. still forms at following dilutions: 1:400,000, w. reagent (a); 1:250,000, w. (c); 1:200,000, w. (d) or (e); 1:160,000, w. gold chloride. Simple salts are often more s. in aq. than those of quinine. Thus: $B_2H_2SO_4 \cdot 2H_2O$, pr., s. in 65 pt. aq. at 13° ; $B_2H_2SO_4 \cdot 4H_2O$, e.s. aq.
		Unlike quinine (No. 2.947), contains no methoxyl group, and does not give the thalleioquin reaction, nor a fluor. sol. w. dil. H_2SO_4 .
	252 ± 2 u.c. d. (r.h.)	Morphine. — Cf. No. 2.1024.
1040	254d.	Imperialine , $C_{13}H_{20}O_4N$ (?). — [Fr. tubers of <i>Coronaria imperialis</i> .] — Ndl. D.s. aq., eth., bz., pet.-eth., amyl alc.; more s. alc. $[\alpha]_D = -35.4^\circ$ (in 5% chlf. sol.). — \oplus Rubbed w. sugar & moistened w. conc. H_2SO_4 gives successively, yel.-green, pale brown, flesh, cherry-red, & after long standing, dark violet, colorations. — B_2HCl , opalescent cryst., e.s. aq., alc.
1041	d. 254 ± 10 u.c.; 250-60	† Solanine , $C_{11}H_{18}O_1N$ (?), $C_{12}H_{20}O_1N$ (?). — [Alkaloid fr. <i>Solanum nigrum</i> , potatoes, etc.] — (Commercial preparation began to yellow at 225° , was much contracted & softened at 254° , but showed no definite m.p.) Fine silky ndl. alm. i. aq.; d.s. c. alc.; e.s. h.; i. bz., lgr., chlf., eth. Reduces Ag sol. Taste (poisonous!), burning & bitter. May be extracted by amyl alc. fr. either ac. or alk. sol. — \oplus (1) Add 2 drops mixt. of 6 cc. conc. H_2SO_4 + 9 cc. alc. to 0.001 g. B on porcelain. ORT1 color appears after heating few sec. on water-bath. — (2) Conc. H_2SO_4 [T. 2.2-(a)] gives O; changing to OS3 after heating 10 sec. on steam-bath. — (3) Sol. in selenium-sulphuric ac. (6 cc. conc. H_2SO_4 + 8 cc. aq. + 0.3 g. Na_2SeO_4) warmed until appearance of pale red, gives fine red on cooling. (Gadamer.) — (4) Evapn. w. a drop or two platinic chloride sol. gives purple to violet color, disappearing on cooling, & reappearing on again warming. (Missaghi, through Gadamer.)
1042	257-9d.	Harmine , $C_{12}H_7ON_2$. — [Fr. seeds of Russian Peganum Harmala, L.] — Rhomb. pr. fr. MeOH. Alm. i. aq.; d.s. alc., eth. Alc. sol. faintly bitter. Salts colorless, but fluor. indigo-blue in dil. sol.! — \oplus S. in conc. H_2SO_4 w. green fluor. — $B_2H_2PtCl_6$, § floe. ppt. becoming cryst. on heating under sol.
—	abt. 260d. (slow heating)	Strychnine. — Cf. No. 2.1047.
1043	263-4	Chelerythrine , $C_{20}H_{17}O_4N$. — [Narcotic alkaloid fr. <i>Sanguinaria canadensis</i> , etc.] — Colorless cryst. fr. ac.-eth. (containing, as usually prepared, 1 mol. EtOH; m.p. 203°). E.s. chlf.; d.s. alc., eth., ac.-eth. Powder provokes sneezing. Salts are egg-yel. Ammon. ppts. B fr. salt sol. colorless; but it turns yel. in air. — \oplus Conc. H_2SO_4 [T. 2.2-(a)] gives greenish yel. sol., changing to dirty yel. — Erdmann's reagt. [T. 2.2-(e)] gives yellow, dark olive-green, chlorophyll-green, & muddy yellow colors. — $B_2H_2SO_4 \cdot 2H_2O$, gold-yel. ndl., d.s. c. aq. — B_2HAuCl_4 , long lust. brown ndl. fr. alc., m.p. 233° d.
1044	265	Protoveratridine , $C_{19}H_{14}O_3N$. — [Fr. sabadilla seeds.] — 4-sided plates. V.d. s. most solvents. Best crystd. fr. chlf. Sol. of salts v. bitter. Not toxic. — \oplus Contact w. conc. H_2SO_4 gives violet & then cherry-red color. Sol. in conc. H_2SO_4 is blood-red, becoming carmine-red on warming. — $B_2H_2PtCl_6 \cdot 6H_2O$, § is only ptd. upon adding alc. to sol., 6-sided plates dehydrated at 100° .
1045	265	Acetoguanamine , $[N : C(NH_2).N : C(NH_2).N : CMe^2]$. — Rhomb. lft. D.s. c., e.s. h. aq.; s. alc. "Reacts weakly alk." — \oplus Boiling w. aq. KOH sol. gives NH ₃ [Cf. T. 2.7-(a)]. — $B_2H_2PtCl_6$, § yel. cryst. ppt., e.s. aq. — B_2AgNO_3 , crystg. in rhomb. tbl. fr. h. aq.
1046	268u.c.	Cinchotine , Hydrocinchonine , $C_{19}H_{22}ON_2$. — [Fr. Cinchona barks.] — Ndl. fr. alc. S. at 16° in 1360 pt. aq.; at 20° in 534 pt. eth. 1 liter 90% alc. at 15° dis. 7.25 g. $[\alpha]_D^{25} = +204.5^\circ$ (in abs. alc., $p = 0.6$). — B_2MeI , pale yel. cryst. fr. MeOH, m.p. $234-5^\circ$.
1047	abt. 268d. (r.h.)	† Strychnine , $C_{21}H_{22}O_2N_2$. — [Tetanic poison fr. <i>Nux vomica</i> .] — Clear 4-sided orthorhombic pr. w. 4-sided pyramidal points fr. alc. Octahedra fr. bz. (Cf. cut, Witthaus, p. 1012.) <i>Odorless</i> . <i>Taste of dil. sol. intensely & persist-</i>

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(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	BASIC COMPOUNDS.—Colorless and Solid.
		ently bitter! A few drops of 1:100,000 aq. sol. will be found faintly bitter to most persons after being held on the tongue for some seconds. S. at 25° in 6400 pt. aq., 110 pt. alc., 5500 pt. eth., 6 pt. chlf., 150 pt. bz., 180 pt. amyl alc. S. in 3000 pt. aq. at 80°, & in 28 pt. alc. at 60° (U.S.P.). $[\alpha]_D^{20} = -114.7$ (in 0.25% alc. sol.). B is extracted fr. alk., but not fr. ac. sol. by shaking w. chlf., or chlf. + eth.
		(P) (1) $H_2SO_4 + KMnO_4$ T. — Dis. 0.5 mg. B completely in 2 drops conc. H_2SO_4 on porcelain. Scatter on the colorless sol. a trace of finely powdered MnO_2 . The colors produced are: at first, V; after 45–50 sec., VR; after 4 min., R; after 7 min., OR; after 12 min., O, persisting for 2 hr. or more. Other oxidants, especially $K_2Cr_2O_7$, & CeO_2 , are often substituted for MnO_2 . Wittaus recommends that the test should be repeated w. each of the three oxidants here mentioned if the quantity of material permits. The most rapid succession of color changes results fr. employment of $K_2Cr_2O_7$. — (2) Euchlorine T. [Bloxam, Chem. News, 155 (1887)]. — Dis. 0.5 mg. B in 1 drop HNO_3 (sp. gr. 1.20) on crucible cover. Warm gently on steam-bath & add abt. 1 mg. solid $KClO_3$. A RO-O color appears immediately, changing to YOS2 upon addition of 2 drops ammon. (sp. gr. 0.90). Evap. to dryness. Dis. the dull greenish residue in 3 drops aq. The GS2 sol. resulting is changed to OYS1 by 3 drops 10% $NaOH$ sol., dil. HNO_3 restoring the GS2 color. — (3) Physiological T. — In medico-legal cases tetanic spasms may be produced in frogs by hypodermic injections. Cf. Wittaus, p. 1063, for details. — (4) T. 2.2 w. conc. H_2SO_4 , Buckingham's & Froehde's reagents. give colorless sol.; & conc. HNO_3 , a YT1, darkening to YS1.
		(D) Dis. 0.05 g. B in 10 cc. h. aq. + 5 drops H_2SO_4 . Add 3 cc. sat. aq. picric ac. sol. Filter. Boil ppt. (which does not dis.) w. 5 cc. alc. Filter. Boil residue again w. 5 cc. chlf. Filter. Dry 15 min. at 100° on tile. The picrate, B.Pk, \ddagger is obtained as a Y microcryst. powd., of unusual insolubility in most organic solvents, M.p. 275–86°. — B.Picrolonate, \ddagger d. 286°.
1048	271d.	2,7-Diamino-3,6-dimethylcarbazole, $(NH_2)_2C_6H_4N$. — Ndl. fr. alc. D.s. c. alc. — Diacetyl deriv., \ddagger m.p. a. 300°; ndl. fr. h. gl. ac. ac.
1049	abt. 270 u.c. d., darkening & softening fr. 260; d.w.m. 235	† Creatinine, $[C(: NH)NMe.CH.CO.NH]^+$. — [In urine & meat extract.] — Lust. pr. fr. aq. S. in 11.5 pt. aq. at 16°; in 625 pt. abs. alc., more s. h. k. 10 ¹¹ = 3.7 at 40°. — (P) Jaffé's React. — Dis. 0.001 g. B in small t.t. in 1 cc. aq. Add 1 drop $NaOH$ sol. (1:10) & 10 drops c. sat. aq. picric ac. sol. The color soon becomes O, changing to RO within abt. one-half min. After longer standing it becomes a permanent OR. [Z. physiol. Chem., 399 (1886)]. — (D) Dis. 0.03 g. B in 1 cc. aq. Add 2 cc. c. sat. aq. sol. picric ac. A ppt. of Y-YT1 ndl. appears at once on shaking. Filter. Wash w. 2 cc. c. aq. Dry 15 min. on porous tile at 100°. The product, B.Pk, \ddagger melts at 212–3° u.c.
1050	275	† Lophine, Triphenylimidazole, $[NH.CPh:N.CPh:CPh]^+$. — Ndl. fr. alc. I. c. aq. 100 pt. abs. alc. dis. 0.88 pt. at 21°, or 2.72 pt. at 78°; alm. i. c. eth. Odorless. Tasteless. Dist. undecd. — (P) Suspend 0.01 g. B in porod. in 5 cc. conc. alc. KOH sol. Heat to 65° & observe appearance in a perfectly dark room. A brilliant phosphorescence which does not entirely disappear even on cooling will be visible. The experiment may be many times repeated w. same material. — B.HCl. \ddagger H ₂ O, m.p. 155°.
1051	280.4 u.c. (in s. cap., 1×30 mm., heating 3° per min. fr. 250°)	† Hexamethylenetetramine, Urotropine, $C_6H_{12}N_4$. — Small charac. hexag. cryst. fr. cooling h. sat. alc. sol. S. in 1.2 pt. aq. at 12°, or in 31 pt. abs. alc.; alm. i. eth. Odorless, when pure. Taste of powd. at first slightly burning, then sweet (No. 1–2 of scale, T. 2.29). — (P) Mix 1 cc. of 1% aq. sol. of B w. 5 cc. dil. H_2SO_4 & boil gently 1 min. Cool & add 2 drops 0.5% aq. resorcinol sol. Pour sol. so as to form layer over conc. H_2SO_4 in t.t. The results described in T. 1.114–(1) for formic aldehyde will be obtained, w. copious separation of pink flocks.
		(D) (a) Dis. 0.1 g. B in 5 cc. aq. Add 5 drops HCl of sp. gr. 1.12 (the proportions are important); then 1 cc. of sol. of 2 g. $NaNO_2$ in 5 cc. aq. Allow to stand some minutes w. occasional shaking. Filter off the ppt. of faintly yellowish ndl. which begins to appear within 5 min. Wash w. 3 cc. c. aq. Dry 30 min. at room temp. on porous tile. The product, dinitrosopentamethylene-tetramine, is obtained in nearly white ndl. which dec. rather violently at 209.5° u.c. — (b) Mix 1 cc. of 1% aq. sol. of B w. 5 cc. sat. aq. $HgCl_2$ sol. Add 10 cc. aq. & boil until heavy ppt. redissolves. Filter, & allow filtrate to cool slowly. Wash the ppt. of long ndl. w. aq., & dry 15 min. on tile at 100°. The product darkens & softens abt. 215°, & melts abt. 233° u.c. (The ppt. forms at high dilutions. Cf. Z. anal. Chem., 36, 44.)

Explanation of typographical signs in this Division: * = T. 2.35. † = Generic position established by actual titrations. ; = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

SUBORDER I OF ORDER II.
COLORLESS COMPOUNDS CONTAINING C, N, H, AND OFTEN O.

GENUS II, BASIC COMPOUNDS.

DIVISION B, LIQUID SPECIES.

No.	Boiling-point (C°).	Neut. Equiv., etc.	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
1059	-6.7°	31 $k \cdot 10^4 = 5$	† Methylamine, MeNH_2. — Odor intensely ammoniacal! 1 vol. aq. at 12° dis. 1150 vol. gaseous \bar{B} . Sp. gr. 0.699 (-10.8°). — ⊖ Sol. gives Rimini's react. (T. 2.25)! — Unlike NH_3 , gives oily black ppt. in T. 2.6., & in x.s. does not dis. freshly ptdt. $\text{Ni}(\text{OH})_2$. ⊖ (a) Mix in a t.t. 8 drops of a 33% aq. \bar{B} sol. (or equivalent quantity not weaker than 5%) & 5 drops ethyl oxalate. In case of strong sol., a stiff mass of white ndl. forms at once. Dil. to 2 cc. & dis. by boiling. Cool. Filter. Dry 15 min. at 100° on porous tile. The product, dimethyloxamide, melts at 211.5-2° u.c. (216.9-7.4° c.) — (b) Pass CO_2 into aq. sol. until only slightly alk. Add x.s. sat. aq. sol. of picrolonic ac. Allow to stand 24 hr. Filter. Wash w. little alc. Recryst. the fine light yel. ndl. fr. little aq. The picrolonate, $\bar{B} \cdot \text{C}_10\text{H}_8\text{O}_3\text{N}_4$, is s. in 1073 pt. c., or 369 pt. h. aq., d. 244°. (Z. physiol. Chem., 43, 308). — $\bar{B} \cdot \text{HCl}$, deliq. lft. i. chlf., m.p. 226°. — $\bar{B} \cdot \text{Pk}$, † yel. lft., s. in 75 pt. aq., m.p. 215°; 207°. — $\bar{B} \cdot \text{H}_2\text{PtCl}_6$, § hexag. yel. tbl., s. in 50 pt. aq., i. abs. alc., m.p. 224°.
1060	+3.5°	59 $k \cdot 10^4 = 5.9$	† Trimethylamine, Me_3N. — Odor intensely ammon. & fishy! V.s. aq. Sp. gr. 0.662 (-5.2°). — ⊖ Sol. does not give Rimini's react. (T. 2.25). — Gives curdy OY ppt. after 2 min. in T. 2.6. ⊖ Mix 5 drops aq. \bar{B} sol. (not weaker than 15%) w. 3 cc. c. sat. aq. picric ac. sol. Filter off the heavy ppt. of yel. ndl. after short interval. Wash w. 1 cc. c. alc. Dry 15 min. on porous tile at 100°. The product, $\bar{B} \cdot \text{Pk}$, † dis. in 77 pt. aq., m.p. 215-6° u.c. In starting w. a weaker sol., use a larger quantity & concentrate by evapn. after adding the picric ac. — $\bar{B} \cdot \text{HCl}$, m.p. 271-5° d. — \bar{B} picrolonate, † $\text{B} \cdot \text{C}_10\text{H}_8\text{O}_3\text{N}_4$, yel. tbl. s. in 1121 pt. c., or 166 pt. boiling aq., d. 250-2°. — $\bar{B} \cdot \text{H}_2\text{PtCl}_6$, § or. red cryst. s. in 3400 pt. boiling abs. alc., d. 242-3°.
1061	7.2°	45 $k \cdot 10^4 = 5.3$	† Dimethylamine, Me_2NH. — Odor intensely ammon.! V.s. aq. Sp. gr. 0.6865 (-5.8°). — ⊖ Sol. gives Simon's react. (T. 2.28), but not Rimini's! — ⊖ Prepare the picrate, $\bar{B} \cdot \text{Pk}$, † by directions given under No. 2.1060. The product, $\bar{B} \cdot \text{Pk}$, cryst. in lust. yel. scales or ndl., m.p. 155.5°-6.5° u.c., s. in 155 pt. aq. — $\bar{B} \cdot \text{HCl}$, e.s. chlf. (unlike NH_4Cl or MeNH_2Cl). — \bar{B} Picrolonate, $\text{B} \cdot \text{C}_10\text{H}_8\text{O}_3\text{N}_4$, fine yel. ndl. s. in 764 pt. c., or 33 pt. boiling aq., d. 222°. — $\bar{B} \cdot \text{H}_2\text{PtCl}_6$, § 6-sided plates, m.p. 206°.

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¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C°).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
1062	abt. 19°	45 $k.10^4 = 5.6$	† Ethylamine, EtNH_2 . — Odor intensely ammon.! Misc. w. aq. Sp. gr. 0.689 (15°). — (P) Gives Rimini's react. (T. 2.25)! — (D) Mix 8 drops 33% aq. B sol. w. 5 drops ethyl oxalate. Mixt. solidifies w. heat evolution. Add 2 cc. c. aq. & crush lumps. Filter. Wash by suction w. 0.5 cc. aq. Dry 15 min. at 100° on porous tile. Diethyloxamide, the product, is obtained in minute white ndl., m.p. 178–9° u.c. (Procedure has been used w. sol. containing as little as 7%, but pptn. is not then immediate, & dilution w. aq. is to be omitted.) — B.HCl, deliq. lft. s. in 0.42 pt. aq. at 17°; s. alc.; m.p. 76–80°. — B.Pk,† yel. cryst. fr. alc. S. in 66.7 pt. aq., or 30.7 pt. alc., at 16°; m.p. 165°. — B. Picrolonate, $\text{B.C}_1\text{H}_8\text{O}_4\text{N}_4$, pale-yel. tbl., v. slowly separating, s. in 93 pt. boiling, or 3846 pt. c. aq., m.p. 244°, browning abt. 220°.
1063	32.2°	59 $k.10^4 = 5.3$	† Isopropylamine, $\text{Me}.\text{CH}(\text{NH}_2).\text{Me}$. — Odor strongly ammon.! Misc. w. aq. Sp. gr. 0.690 (18°). — (P) Gives Rimini react. (T. 2.25)! — Prim. amine.* — (D) Mix 5 drops B w. 5 drops ethyl oxalate. Shake, & allow to stand. Dis. ppt. in 1 cc. boiling 50% alc. Cool. Filter. Wash w. few drops dil. alc. Dry 15 min. on porous tile at 100°. The product, diisopropyl-oxamide, forms white ndl., m.p. 213–4° u.c. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § d.s. yel. cryst., m.p. 227–8°.
1064	34–5°	59	Methylethylamine, $\text{Me}.\text{NHET}$. — Odor ammon.! E.s. aq. — (P) Sec. amine.* Should give Simon's react. (T. 2.28)! — $\text{B}_2\text{H}_2\text{PtCl}_6$, § pr., m.p. 207–8°. — $\text{B.C}_2\text{H}_2\text{O}_4$, ndl., d.s. abs. alc., m.p. 154–5°.
1065	42.4°	61	Dimethylhydroxylamine, MeNH.O.Me . — (P) Odor sweetish, not ammon.! Does not reduce AgNO_3 , or Fehling's sol. — B.HCl, non-hydroscopic tbl., m.p. 115–6°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § red pr., m.p. 180° d.
1066	45	73 $k.10^4 = 3.4$	tert.-Butyl-amine, Me_3NH_2 . — S. aq. Sp. gr. 0.7054 (8°). — (P) Tert. amine.* — B.HCl, tbl. fr. abs. alc., m.p. 270–80°.
1067	49	59 $k.10^4 = 4.7$	n-Propylamine, Pr.NH_2 . — Odor ammon. S. aq. Sp. gr. 0.7186 (20°). — (P) Should give Rimini's react.! Prim. amine.* — B.HCl, m.p. 157–8°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § d.s. c. aq., m.p. 214°. — B.Pk,† m.p. 135°.
1068	53.3; 56–6.5	57 $k.10^4 = 4.6$	† Allylamine, $\text{C}_2\text{H}_5\text{NH}_2$. — Odor irritating & ammon.! V.s. aq. Sp. gr. 0.769 (15°). — (P) Gives Rimini's react. (T. 2.25)! — Prim. amine.* — (D) Mix 2 drops B w. 2 drops ethyl oxalate (avoid x.s.). Dis. in 3 cc. h. 33% alc. Cool. Filter. Dry at 100°. The product, diallyloxamide, is obtained in white scales or ndl., m.p. 154–4.5° u.c. (156.5–7° c.). — B.Pk,† lemon-yel. ndl. fr. aq., m.p. 140–1° (after sintering).
1068-I	55.5	73 $k.10^4 = 1.3$	Diethylamine, Et_2NH . — Odor ammon.! V.s. aq. Sp. gr. 0.7116 (15°). M.p. abt. –50°. — (P) Sec. amine* giving Simon's react. (T. 2.28)! — (D) Mix 3 drops w. 3 cc. c. sat. alc. sol. oxalic ac. & add 5 cc. eth. Filter. Wash ppt. w. 2 cc. eth.-alc. (1:1). Dry 15 min. at 110° on tile. The product $\text{Et}_2\text{NH.C}_2\text{O}_4\text{H}_2$, is obtained in minute colorless cryst., m.p. 206.5–7.5° u.c. — B.HCl, non-deliq. lft. fr. eth.-alc., v.s. aq.; mod. s. abs. alc.; e.s. chlf., m.p. 215–7°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, orange-yel. monoclin. cryst. — B. Picrolonate, $\text{B.C}_1\text{H}_8\text{O}_4\text{N}_4$, pale yel. ppt., s. in 402 pt. boiling or 3788 pt. c. aq., d. 260°.
1069	50–60		s-Dimethylhydrazine, MeNH.NHMe . — $\text{B.C}_2\text{H}_2\text{O}_4$, cryst. s. in 60 pt. abs. alc., m.p. 132°.
1070	55–6	43	Vinylamine, $\text{CH}_2:\text{CH.NH}_2$. — Odor ammon.! Fumes in air. Misc. w. aq. Prim. amine.* Stable to c. KMnO_4 sol. — (P) Sp. gr. 0.8321 (24°). — B.Pk,† monoclin. tbl., m.p. 142°, c.s. warm aq. — $\text{B.C}_2\text{H}_2\text{O}_4$, ndl. e.s. aq., d.s. abs. alc., m.p. 115° d.

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 ¶ = 2.1. ↔ = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C.).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I071	63	57	Trimethyleneimine, $\text{[CH}_2\text{CH}_2\text{CH}_2\text{NH}]$. — Odor strongly ammon. Fumes in air. Misc. w. aq. Sec. amine.* Sp. gr. 0.843 (20°). — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or.-yel. ndl., m.p. abt. 200°.
I072	63		Dimethylhydrazine, $\text{Me}_2\text{N.NH}_2$. — Odor strongly ammon. E.s. aq. Sp. gr. 0.7914 (22.3°). HNO_3 gives N_2O . — $\text{B}_2\text{C}_2\text{H}_4\text{O}_2$, plates fr. alc., m.p. 142–3°.
I073	63	73	sec.-Butyl-amine, $\text{EtCH}(\text{NH}_2)\text{Me}$. — Sp. gr. 0.718 (20°). — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.*
I074	62–4	73	Methylpropylamine, MeNH.Pr . — Odor fishy-ammon. S. aq. Sp. gr. 0.720 (17°). — \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § e.s. aq., d.s. h. alc., m.p. 200° d.
I075	63–5	87 $k.10^4 = 2.7$	Methyldiethylamine, MeNEt_2 . — E.s. aq. — \textcircled{P} Tert. amine. §
I076	65	71	Methylallylamine, $\text{Me.NH.C}_2\text{H}_4$. — Odor ammon. Misc. w. aq. — \textcircled{P} Sec. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yel. cryst. d.s. c. aq., i. alc., m.p. 164°.
I077	68	61	α -Ethylhydroxylamine, EtO.NH_2 . — Strong, but not ammon. odor. Misc. aq., alc., eth. Sp. gr. 0.8827 (7.5°)! Gives white ppt. w. AgNO_3 , w. evolution of gas & metallic Ag on boiling.
I078	68c.	73 $k.10^4 = 3.1$	† Isobutyramine, $\text{Me}_2\text{CH.CH}_2\text{NH}_2$. — Odor ammon. Misc. w. aq. Sp. gr. 0.7345 (15°). — \textcircled{P} Gives Rimini's react. (T. 2.25). Prim. amine.* — \textcircled{P} Mix 2 drops B w. 3 drops ethyl oxalate. Recryst. ppt. fr. 4 cc. h. 50% alc. Wash on filter w. $\frac{1}{2}$ cc. dil. alc. Dry 15 min. on porous tile at 100°. The product, diisobutyloxamide, forms fine ndl., m.p. 169° u.c. (172° c.). — B.HCl , v.s. aq., m.p. 160°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or. cryst., d. abt. 225°.
I079	76	87	Ethylisopropylamine, $\text{Et.NH.CH}_2\text{Me}_2$. — Misc. w. aq. — \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or. ndl. fr. aq., m.p. 180°.
I080	76–8	71	Methylisobutyramine, $\text{Me.NH.CH}_2\text{CH}_2\text{Me}_2$. — Odor fishy. Sp. gr. 0.722 (18°). — \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.*
I081	78c.	57	† Butylamine, $\text{Me}(\text{CH}_2)_3\text{NH}_2$. — Odor unpleasant, somewhat ammon. V.s. aq. Sp. gr. 0.740 (20°). — \textcircled{P} Gives Rimini's react. (T. 2.25). Prim. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § yel. lft. d.s. c. aq.
I082	79–80	83	N-Methylpyrrolidine, $\text{[NMe.CH:CH.CH}_2\text{CH}_2]$. — Odor unpleasant, ammon. Misc. w. aq. — B.HAuCl_4 , lft. fr. dil. alc., m.p. 190–1°.
I083	78c.	71	tert.-Amyl-amine, $\text{Me}_2\text{C}(\text{NH}_2)\text{Et}$. — \textcircled{P} Tert. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § v.s. aq., alc.
I084	82	71	Aminocyclobutane, $\text{[CH}_2\text{(CH}_2\text{)}_2\text{CH}(\text{NH}_2)]$. — Odor v. pungent. — \textcircled{P} Prim. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § deep-yel. octahedra fr. h. aq., blackens at 210–5°.
I085	81–3c.	85	N-Methylpyrrolidine, $\text{[NMe}(\text{CH}_2)_3\text{CH}_2]$. — Misc. aq. — \textcircled{P} Tert. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § (dried at 105°), m.p. 233°. — B.HAuCl_4 , yel. cryst., m.p. 218°.
I086	81–5	71	α -Crotylamine, $\text{Me.CH:CH.CH}_2\text{NH}_2$. — Odor penetrating, ammon. Mod. s. aq. — \textcircled{P} Prim. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § mic. tbl., m.p. 193° d.
I087	82–3	87	1-Amino-2,2-dimethylpropane, $\text{Me}_2\text{C}(\text{NH}_2)\text{CH}_2$. — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.* — B.HCl , m.p. abt. 275°.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I088	83	89	Ethoxyethylamine, EtO.NHET .—Odor fishy. E.s. aq. — \textcircled{P} Prim. amine.* — Reduces AgNO_3 on warming. — $\text{B.C}_2\text{H}_5\text{O}_4$, e.s. aq., m.p. 112 $^{\circ}$.
I089	84	85	Ethylallylamine, $\text{C}_2\text{H}_5\text{NHEt}$.—Misc. aq. — \textcircled{P} Sec. amine.* — $\text{B}_2\text{H}_5\text{PtCl}_6$, § monoclin. red pr., e.s. aq.; m.p. (after softening) 154—6 $^{\circ}$.
I090	84	87	3-Amino-2-methylbutane, $\text{Me}_2\text{CH.CH}(\text{NH}_2)\text{Me}$.—Unpleasant basic odor. — Sb. pr. 0.757(18.5 $^{\circ}$). — \textcircled{P} Should give Rimini's react. Prim. amine.*
I091	84	101	Diisopropylamine, $(\text{Me}_2\text{CH})_2\text{NH}$.—Sp. gr. 0.722 (22 $^{\circ}$). — \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.*
I092	84—6		s-Diethylhydrazine, EtNH.NHET .— B_2HCl , lft. fr. dil. HCl, m.p. 160 $^{\circ}$ d. — \textcircled{P} Reduces Fehling's sol., but only on heating!
I093	87.5c.		Methylhydrazine, MeNH.NH_2 .—Odor like methylamine. E.s. aq. Caustic. — \textcircled{P} Reduces c. Fehling's sol.! — $\text{B}_2\text{H}_5\text{SO}_4$, long ndl. e.s. aq., d.s. alc., m.p. 139.5 $^{\circ}$. — Picrate, † m.p. 162 $^{\circ}$ d.
I094	87.5—8.5	57	Pyrrolidine, Pentazane, $[\text{NH}.\text{CH}_2.\text{CH}_2.\text{CH}_2]^n$.—Odor basic, piperidine-like. Misc. aq. Sp. gr. 0.8520 (22.5 $^{\circ}$). — \textcircled{P} Sec. amine.* — Nitroso deriv., ** yel. oil, b.p. 214 $^{\circ}$ d. — B_2HAuCl_4 , yel. scales, v.s. h. aq., m.p. 206 $^{\circ}$ d.
I095	89c.	101	† Triethylamine, NET_3 .—Odor ammon. S. aq. Sp. gr. 0.7331 (15 $^{\circ}$). — \textcircled{P} Gives neither Rimini's nor Simon's react. (T. 2.25 & 2.28). Tert. amine.* \textcircled{P} Dis. 2 drops B in 1 cc. aq. Add 3 cc. c. sat. aq. picric ac. sol. Heat to boiling to dis. ppt. & cool slowly. Filter. Wash cryst. ppt. w. 1 cc. c. aq. Dry 15 min. on porous tile at 100 $^{\circ}$. The picrate (unanalyzed) forms yel. scales & ndl., softening fr. 168 $^{\circ}$, m.p. 170—2 $^{\circ}$ u.c. — B_2 . Picrolonate, † $\text{B}_2\text{C}_10\text{H}_8\text{O}_4\text{N}_4$, yel. cryst., slowly separating, s. in 63 pt. boiling, or in 536 pt. c. aq., d. 160 $^{\circ}$.
I096	90c. (748 mm.)	69	Pyrroline, $[\text{NH}.\text{CH}_2.\text{CH}:\text{CH}.\text{CH}_2]^n$.—Fumes & absorbs CO_2 in air. V.s. aq. — Sp. gr. 0.9097 (20/4). — B_2P_k , † yel. pr. fr. aq., m.p. 156 $^{\circ}$; e.s. aq., alc. — $\text{B}_2\text{H}_5\text{PtCl}_6$, § or-red ppt., d.s. c., e.s. h. aq., triclin. cryst., m.p. 182 $^{\circ}$ d.
I097	90	87	3-Aminopentane, $\text{Et.CH}(\text{NH}_2).\text{Et}$.—Odor ammon. Sp. gr. 0.749 (20/4). — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.* — B_2HCl , ndl., m.p. 216 $^{\circ}$.
I098	91	87	Methylbutylamine, $\text{MeNH.C}_2\text{H}_5$.— \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.* Sp. gr. 0.737 (15 $^{\circ}$). — B_2HCl , tbl., fr. acetone, m.p. 170—1 $^{\circ}$.
I099	92	87	2-Aminopentane, $\text{Me.CH}(\text{NH}_2).\text{CH}_2.\text{CH}_2.\text{Me}$.—Odor ammon. V.s. aq. — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.* — Sp. gr. 0.742 (18.5 $^{\circ}$). — $\text{B}_2\text{H}_5\text{O}_4$, m.p. 131 $^{\circ}$.
I100	95	87	† Isoamylamine, $\text{Me.CH.CH}_2.\text{CH}_2.\text{NH}_2$.—Odor strongly ammon., unpleasant, & slightly suggestive of amyl alc. V.s. aq. — \textcircled{P} Gives Rimini's react. (T. 2.25). Prim. amine.* Sp. gr. 0.7462 (17.5 $^{\circ}$). — \textcircled{P} Mix 3 drops B w. 4 drops ethyl oxalate. Dis. ppt. in 3 cc. h. 50% alc. Cool well. Recryst. fr. 2 cc. h. 50% alc. Dry 15 min. on tile at 100 $^{\circ}$. The product, diisoamyl oxamide, is obtained in white ndl. of slippery feel, m.p. 140 $^{\circ}$ u.c.
I101	95—6		Ethyldieneazine, $\text{Me.CH:CH.N:N:CH.Me}$.—Sp. gr. 0.832 (17 $^{\circ}$).
I102	95.5—6.5 (744 mm.)	85	2-Methylpyrrolidine, $[\text{NH}.\text{CHMe.CH}_2.\text{CH}_2.\text{CH}_2]^n$.—Odor penetrating & overpowering. Sp. gr. 0.84 (20/20). — $\text{B}_2\text{H}_5\text{O}_4$, softens 165 $^{\circ}$, m.p. 178—9 $^{\circ}$ d.

Explanation of typographical signs used in this Division: * = T. 2.35. † = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS. — Colorless and Liquid.
II103	96-6.5	99	1,2-Dimethylpyrrolidine, $[NMe.CHMe.CH_2.CH_2.CH_3]$. — Odor ammon. & piperidine-like. Misc. aq. Sp. gr. 0.83 (20°). — \textcircled{P} Tert. amine.* — $\text{B}_2\text{H}_4\text{PtCl}_6$, § or.-red pr., m.p. 223-4° d.
II104	96-9		uns-Diethylhydrazine, $Et_2N.NH_2$. — Odor ethereal, faintly ammon. E.s. aq. — \textcircled{P} Reduces Fehling's sol. (but only when warmed)! — Picrate,† fine yel. ndl.; evolves N when boiled w. aq.
II105	98	101	Ethylisobutylamine, $Et.NH.CH_2.CHMe_2$. — \textcircled{P} Should give Simon's react. (T. 2.28). Sec. amine.* — B_2HCl , m.p. 209°.
II106	101		Ethylhydrazine, $Et.NH.NH_2$. — Odor ethereal, faintly ammon. E.s. aq. Caustic. — \textcircled{P} Reduces Fehling's sol. in cold! Gives carbylamine react. (T. 2.12).
II107	100-3	101	4-Amino-2-methylpentane, $Me.CH(Me).CH_2.CH(NH_2).Me$. — Odor ammon. — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.* — $\text{B}_2\text{C}_2\text{H}_4\text{O}_4$, scales, m.p. 219°.
II108	103	101	3-Amino-2,2-dimethylbutane, $Me.CMe_2.CH(NH_2).Me$. — S. in 1 vol. c. aq.; less s. h. aq. Solidifies at -20°. — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.* — $\text{B}_2\text{H}_4\text{PtCl}_6$, § v.s. aq., alc.
II109	103-5	85	3-Methylpyrrolidine, $[NH.CH_2.CHMe.CH_3]$. — Odor piperidine-like. Fumes in air. Sp. gr. 0.8654 (0/4). — B.Pk ,† m.p. 105°, v.s. alc. — B_2HAuCl_4 , mic. tbl., e.s. aq., m.p. 170°.
II110	104	87	Amylamine, $Me.(CH_2)_5.NH_2$. — Sp. gr. 0.766 (19°). \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.*
II111	105	115	Methylethylisobutylamine, $Me.Et.BuN$. — Tert. amine.* — $\text{B}_2\text{H}_4\text{PtCl}_6$, § d.s. alc., m.p. 197°.
II112	106c.	85 $k.10^8 = 1.2$	† Piperidine, $[NH.(CH_2)_5]$. — Odor unpleasant, ammon., characteristic! Misc. w. aq., alc. — \textcircled{P} Gives good blue color (B) within 1 min. in Simon's react. (T. 2.28). Sp. gr. 0.8619 (19.6°). — \textcircled{D} Mix in v. small t.t. 4 drops B & 3 drops CS_2 . Vigorous react. ensues. Scrape down product fr. walls of tube. Add 3 or 4 more drops CS_2 to complete reaction. Dis. in 3 cc. alc. Concentrate sol. in v. small glass capsule by evapn. to abt. 1 cc. Transfer residue of lust. plates w. mother liquor to porous tile & allow to air-dry without heating. The product, piperyldihiocarbamate, when rapidly heated to 160° softens at 168° & melts at 172.2° u.c. (175.0° c.), subliming so quickly that it will all escape before melting unless the capillary is well filled & the heating is rapid. — B.Pk ,† ndl., e.s. h. aq., m.p. 145° d.
II113	106-8c.	85	Aminocyclopentane, $[CH(NH_2).(CH_2)_3.CH_3]$. — Odor strongly ammon. Misc. w. aq. — \textcircled{P} Prim. amine.* — $\text{B}_2\text{H}_4\text{PtCl}_6$, § red-yel. scaly ppt.
II114	107c.	97	2,5-Dimethylpyrrolidine, $[C_2H_5Me_2.NH]$. — Mobile liq. w. strong amine odor. Misc. w. little aq., sol. becoming turbid on further diln., & on still further diln. clear. Sp. gr. 0.8369 (20/4). $n_D^{20} = 1.4401$. — \textcircled{P} Sec. amine.* — B.Pk ,† yel. ndl. e.s. aq., alc., m.p. 105°.
II115	107	99	N-Methylpiperidine, $[NMe.(CH_2)_5]$. — Sp. gr. 0.8230 (10.2/4). — \textcircled{P} Tert. amine.* — $\text{B}_2\text{H}_4\text{PtCl}_6$, § or. cryst. fr. alc., m.p. 210-2° d.
II116	106-8c. (746 mm.)	99	2,5-Dimethylpyrrolidine, $[NH.CHMe.(CH_2)_3.CHMe]$. — Powerful odor. Misc. aq. Sp. gr. 0.8185 (12.3/4). — \textcircled{P} Sec. amine.* — B_2HCl , ndl., m.p. 188-90 .
II117	108-10	101	3-Amino-3-methylpentane, $Me.CH_2.CMe(NH_2).Et$. — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.*
II118	110c.	101	† Dipropylamine, $(Et.CH_2)_2NH$. — Odor strongly ammon. S. aq. Sp. gr. 0.736 (25°). — \textcircled{P} Gives Simon's react. (T. 2.28). Sec. amine.* — \textcircled{D} Dis. 2 drops B in 3 cc. pet.-eth. Add

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
			<i>3 drops CS₂. After 2 min. slender white ndl. begin to separate, & after 5 min. a compact mass of interlacing ndl. forms. Filter. Wash w. $\frac{1}{2}$ cc. pet.-eth. Recryst. fr. 2 cc. h. pet.-eth., allowing to stand w. occasional shaking for some time. Filter. Dry 15 min. at 60° on porous tile. The product, dipropylamine dipropylthiocarbamate, is obtained in matted silky ndl., m.p. 118.5° u.c.</i>
II19	111	97	Diallylamine, (C ₂ H ₅) ₂ NH. — Sec. amine.*
II20	110-4	109	Allylpropylamine, C ₂ H ₅ NH.Pr. — Sec. amine.* Sp. gr. 0.771 (18°).
II21	112	111	Methyldiallylamine, Me.N(C ₂ H ₅) ₂ . — Tert. amine.*
II22	111-3	113	1,2,4-Trimethylpyrrolidine, $^{\text{f}}\text{NMe.CHMe.CH}_2\text{CH}_2\text{CHMe}^{\text{l}}$. — Odor piperidine-like. Tert. amine.* — B ₂ H ₂ PtCl ₆ , § e.s. aq., m.p. 179-80°.
II23	113-4	101	1-Amino-2,2-dimethylbutane, NH ₂ CH ₂ CMe ₂ CH ₂ Me. — (P) Should give Rimini's react. (T. 2.25). Prim. amine.* — B.HCl, m.p. 225-8°.
II24	114	99	1-Amino-1-methylcyclopentane, $^{\text{f}}\text{CMe}(\text{NH}_2)\text{(CH}_2\text{)}_4^{\text{l}}$. — Odor ammon. Fumes in air. E.s. aq. Sp. gr. 0.820 (20/0). — (P) Prim. amine.* — B.HAuCl ₄ (110°), or. ndl., m.p. 173° d.
II25	115c.	79 k.10° = 2.3	† Pyridine, $^{\text{f}}\text{CH:N.CH:CH.CH:CH}^{\text{l}}$. — Odor strong, charac., unpleasant aromat., not distinctly ammon. when pure. Misc. aq., alc., eth., chlf. Hygroscopic. (Remove aq. by long standing over solid KOH.) Sp. gr. 0.9893 (15/4). Tert. amine.* — (P) Add alc. sol. of 2,4-dinitrochlorobenzene to alc. or aq. sol. containing not less than 0.1% B. Shake vigorously. Add NaOH sol. A red-violet color appears. (Von-gerichten, Ber., 32, 2571.) ① Add 3 cc. c. sat. alc. picric ac. sol. to sol. 1 drop B in 3 cc. alc. Filter the heavy ppt. of yel. scales which separate at once. Wash w. 3 cc. alc. Cryst. fr. 3 cc. h. alc. Air-dry on porous tile at room temperature. The product, B.Pk,† is obtained in Y scales or ndl., m.p. 163-5° u.c. (166-8° c.). — B ₂ H ₂ PtCl ₆ , § or.-yel. ndl., m.p. 240-2°.
II26	115-7	99	2,4-Dimethylpyrrolidine, $^{\text{f}}\text{NH.CHMe.CH}_2\text{CHMe.CH}^{\text{l}}$. — Sec. amine.* Sp. gr. 0.8297 (20/4). — B.Pk,† ndl. fr. aq. or alc., m.p. 116-7°. — B Picrolonate,† yel. ndl. fr. alc., m.p. 227°.
II27	116c.	101	2-Aminohexane, Me.CH(NH ₂).(CH ₂) ₄ Me. — Sp. gr. 0.764. — (P) Should give Rimini's react. (T. 2.25). Prim. amine.*
II28	116	101	1-Methylmorpholine, $^{\text{f}}\text{NMe.CH}_2\text{CH}_2\text{O.CH}_2\text{CH}_2^{\text{l}}$. — Odor strongly ammon. Misc. aq. Sp. gr. 0.9051 (20/4). — (P) Tert. amine.* — B.Pk,† hair-like ndl., alm. i. alc., eth., m.p. 225-6°.
II29	abt. 116c.	113	1,2,5-Trimethylpyrrolidine, $^{\text{f}}\text{NMe.CHMe}(\text{CH}_2)\text{CH}_2\text{CHMe}^{\text{l}}$. — D.s. aq. Sp. gr. 0.8149 (7.4/4). — (P) Tert. amine.* — B.Pk,† dendritic cryst. fr. alc., d.s. aq., m.p. abt. 163°. — B.C ₁₀ H ₈ O ₂ N ₄ ,† (picrolonate), ndl. fr. alc., e.s. aq., m.p. abt. 193°.
II30	117c.		† Ethylenediamine, NH ₂ CH ₂ CH ₂ NH ₂ . — Odor strongly ammon. Fumes in moist air. (Absorbs moisture, giving hydrate w. 1H ₂ O, b.p. 118°, m.p. +10°, fr. which aq. can be removed only by heating w. fused KOH at 100°.) S. aq. Not misc. w. bz. or eth. Sp. gr. 0.902 (15°); 0.970 (15°) for hydrate. Prim. amine.* — (P) Shake persistently in stoppered tube 1 drop B, 5 cc. NaOH sol. (1:10), & 8 drops benzoyl chloride, breaking lumps w. stirring rod until ppt. becomes dry & granular and odor of benzoyl chloride disappears. Filter. Wash thrice w. 2 cc. c. aq. Dis. in 8 cc. boiling alc. After

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No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
			<i>cooling & v. persistent shaking (!) the product, dibenzoyl-ethylenediamine, separates in minute pr., which after drying 15 min. at 100° melt at 244.5° u.c. (352° c.). — B.Pk., † lft., d.s. aq., m.p. 233–5° d.</i>
II131	117	115	Methyldipropylamine, Me.NPr₂. — \textcircled{P} Tert. amine.*
II132	117–8	101	5-Aminohexene(1), CH₂:CH(CH₂)₂.CH(NH₂).Me. — Odor piperidine-like. Misc. w. aq. Sp. gr. 0.779 (15°). — \textcircled{P} Prim. amine.*
II133	118	121	5-Dimethylamino-pentene(1), CH₂:CH(CH₂)₂.NMe₂. — Sp. gr. 0.7634 (14.9/4). — \textcircled{P} Tert. amine.*
II134	118–9	99	α-Pipecoline, 2-Methylpiperidine, $[\text{NH}.\text{CHMe}.\text{(CH}_2)_4]$. — Penetrating piperidine-like odor. E.s. aq. Sp. gr. 0.8622 (0°). — \textcircled{P} Sec. amine.* — B.Pk., † lust. yel. ndl., m.p. 134–5° (127–8°), e.s. alc., d.s. eth.
II135	119		1,2-Dimethylamino-ethane, NHMe.CH₂.CH₂.NHMe. — Odor ammon. Sp. gr. 0.828 (15/4). Sec. amine.* — B.Pk., † lft. fr. h. aq., m.p. 215–6°.
II136	119–20		1,2-Diaminopropane, NH₂.CH₂.CH(NH₂).CH₂. — [Absorbs moisture rapidly forming hydrate, B.H ₂ O, fr. which it is said aq. can be removed only by Na. Hydrate is misc. w. aq., b.p. 203–7°.] — Sp. gr. 0.878 (15°). Prim. amine.*
II137	121c.	97	2,4-Dimethylpyrrolidine, $[\text{C}_4\text{H}_8\text{Me}_2.\text{NH}]$. — Odor penetrating. Sp. gr. 0.8554 (20/4). Sec. amine.* — B.Pk., † yel. pr. fr. alc. + lgr., m.p. 102–4°. B ₁₀ H ₈ O ₂ N ₄ (picrolonate), † dark-yel. cryst. fr. alc., m.p. 225°.
II138	123–4c.	80	Pyrimidine, $[\text{N}:\text{CH}.\text{N}:\text{CH}.\text{CH}:\text{CH}]$. — Penetrating narcotic odor. Cryst. mass, m.p. 20–2°. Misc. aq. — B.Pk., † ndl., m.p. 156°.
II139	123	99	2-Amino-1-methylcyclopentane, $[\text{CHMe}.\text{CH}(\text{NH}_2).\text{(CH}_2)_3]$. — “V. little s. aq.” Sp. gr. 0.801 (20/0). — \textcircled{P} Prim. amine.* — B ₂ H ₂ PtCl ₆ , § dark-yel. ndl. fr. aq.; d.s. aq., d. 240°.
II140	123		Dimethylaminoacetone, Me₂N.CH₂.CO.Me. — Misc. w. aq., alc., eth. — B ₂ H ₂ PtCl ₆ , § cryst. d.s. aq., m.p. 176° d.
II141	123	113	Allylisobutylamine, C₃H₇.NH.CH₂.CHMe. — E.s. aq. — \textcircled{P} Sec. amine.* — B ₂ H ₂ PtCl ₆ , § red cryst., m.p. 182°.
II142	124	99	3-Amino-1-methylcyclopentane, $[\text{CHMe}.\text{CH}_2.\text{CH}(\text{NH}_2).\text{(CH}_2)_2]$. — Misc. w. aq. Sp. gr. 0.843 (20/20). — \textcircled{P} Prim. amine.* — Benzoyl deriv., §§ m.p. 115–7°.
II143	123–5	115	Propylisobutylamine, Pr.NH.CH₂.CHMe. — Odor basic & like fusel-oil. D.s. aq. — \textcircled{P} Sec. amine.* — Dioxalate, ndl., d.s. aq., m.p. 224°.
II144	125–6c.	99	d,l-3-(β)-Methylpiperidine, β-Pipecoline, $[\text{NH}.(\text{CH}_2)_3.\text{CHMe}.\text{CH}_2]$. — V.s. aq. Sp. gr. 0.8635 (0/4). — \textcircled{P} Sec. amine.* — B.Pk., † sulphur yel. pr., mod. s. aq., m.p. 136–8°. — B ₂ H ₂ PtCl ₆ , § chrome-red trimet. pr., m.p. 207° d. — B.HAuCl ₄ , mod. s. aq., m.p. 130–1°.
II145	125. 3c.	101	1-Amino-2-ethylbutane, Et.CH(Et).CH₂.NH₂. — Odor ammon. — \textcircled{P} Prim. amine.* — B.HCl, ndl. fr. alc., m.p. 187° d.
II146	126–8	113	2,3,5-Trimethylpyrrolidine, $[\text{NH}.\text{CHMe}.\text{CH}_2.\text{CHMe}.\text{CHMe}]$. — Odor piperidine-like. Misc. w. aq. Sp. gr. 0.816 (15°). — B ₂ H ₂ PtCl ₆ , § or.-red pr., m.p. 205–6° d.
II147	127	115	Ethylisoamylamine, Et.NH.C₄H₁₁. — D.s. aq. Sp. gr. 0.764. — \textcircled{P} Sec. amine.*
II148	126. 5–9c.	99	4-Methylpiperidine, γ-Pipecoline, $[\text{NH}.\text{(CH}_2)_3.\text{CHMe}.\text{(CH}_2)_2]$. — Fumes in air. E.s. aq. Sp. gr. 0.8674 (0°). — \textcircled{P} Sec. amine.* — B ₂ H ₂ PtCl ₆ , § lust. pr., m.p. 203° d. — Gold double salt, m.p. 125–7°.

No.	Boiling-point (C°).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
II49	126-30		Morpholine, $[\text{NH} \cdot (\text{CH}_2)_2 \cdot \text{O} \cdot (\text{CH}_2)_2]$. — Odor piperidine-like. V. volat. & hydroscopic. — \textcircled{P} Sec. amine.* — Nitroso deriv., ** odorous yel. cryst., e.s. aq., m.p. 29°, b.p. 225°. — $\text{B.Pk.} \ddagger$ pr. r.d.s. aq., m.p. 145-7°. — $\text{B.Pk.} \ddagger$ Picrolonate, † d.s. aq., d. 255°.
II50	128 ± 2	101	Hexylamine, $\text{Me} \cdot (\text{CH}_2)_5 \cdot \text{NH}_2$. — Odor distinctly ammon. Somewhat s. aq. Sp. gr. 0.768 (17°). — \textcircled{P} Should give Rimini's react. (T. 2.25). Prim. amine.*
II51	128c.	113	1,2-Dimethylpiperidine, N-Methyl- α -piperidine, $[\text{NMe} \cdot \text{CHMe} \cdot (\text{CH}_2)_4]$. — Odor strong, piperidine-like. S. in 10-12 vol. c. aq.; less s. h. aq. Sp. gr. 0.8345 (7.4/4). — \textcircled{P} Tert. amine.* — B.HCl , pr., m.p. 258-9°. — $\text{B.Pk.} \ddagger$ lust. yel. ndl., d.s. c. aq., m.p. 240-1°.
II52	128	113	2,6-Dimethylpiperidine, $[\text{NH} \cdot \text{CHMe} \cdot (\text{CH}_2)_5 \cdot \text{CHMe}]$. — Misc. w. aq. Sp. gr. 0.8492 (0°). — B.HCl , ndl., m.p. 279-81°. — $\text{B.Pk.} \ddagger$ m.p. 162-4°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or-red cryst., m.p. 212°.
II53	129c. k. $10^8 = 3.2$	93	† α -Picoline, 2-Methylpyridine, $\text{Me} \cdot \text{C}_6\text{H}_4\text{N}$. — Odor, strong, unpleasant, pyridine-like. V. s. aq. Sp. gr. 0.9497 (15/4). — \textcircled{P} Tert. amine.* — Boiled w. KMnO_4 sol. is oxidized to picolinic ac., No. 2.130. — \textcircled{P} Prepare the picrate by procedure described under pyridine (No. 2.1125). $\text{B.Pk.} \ddagger$ is obtained in yel. ndl., m.p. 164.3-5.3° u.c. d. (167.2-8.2° c.).
II54	129-30	(127)	1-Dimethylamino-2-methylpentene(4), $\text{NMe}_2 \cdot \text{CH}_2 \cdot \text{CHMe} \cdot \text{CH}_2 \cdot \text{CH}_2$. — Odor piperidine-like. Sp. gr. 0.767 (15°).
II55	abt. 130d.		Methyl Aminoacetate, $\text{NH}_2 \cdot \text{CH}_2 \cdot \text{CO} \cdot \text{Me}$. — Strong basic odor. Absorbs CO_2 . Polymerizes on keeping. — \textcircled{P} Should give N in T. 2.4. — Distn. w. dry Na_2CO_3 gives NH_3 & Et.NH_2 (No. 2.1062).
II56	abt. 130		2-Dimethylamino-ethanol(1), $\text{Me}_2\text{N} \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{OH}$. — B.HAuCl_4 , silky ndl. s. c. aq., m.p. 197°.
II57	130		Tetramethyltetrazone, $\text{Me}_2\text{N}_2 \cdot \text{N}_2\text{Me}_2$. — Yel. oil. Explodes violently above b.p.! D.s. aq. "Strong base." — \textcircled{P} Reduces Ag sol. in c. — Boiled w. dil. acids gives MeNH_2 , Me_2NH , etc.
II58	132-4	129	Ethyldipropylamine, Et.NPr. — \textcircled{P} Tert. amine.* — B.HAuCl_4 , ndl. s. aq., m.p. 96°.
II59	133-5		Pyrazolidone, $[\text{CO} \cdot \text{NH} \cdot \text{NH} \cdot \text{CH}_2 \cdot \text{CH}_2]$. — E.s. ac.; i. alk. — \textcircled{P} Reduces ammon. AgNO_3 sol. at once. Sol. colored violet-red by little FeCl_3 .
II60	134	99	Aminocyclohexane, $[\text{CH}(\text{NH}_2) \cdot (\text{CH}_2)_5]$. — Odor ammon. & coniine-like. Fumes slightly. Absorbs CO_2 . Sp. gr. 0.865 (20/0). — B.HCl , v.s. aq., alc.; i. eth., m.p. 206°.
II61	133-6	113	2-Amino-3-methylhexene(5), $\text{CH}_2 \cdot \text{CH}(\text{NH}_2) \cdot \text{CHMe} \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{CH}_2$. — Odor piperidine-like. E.s. aq. Sp. gr. 0.793 (15°). — \textcircled{P} Prim. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$ § (over H_2SO_4), or lft. e.s. aq., m.p. 157-8°.
II62	135c.		2-Methylpyrazine, $[\text{N} \cdot \text{CH} \cdot \text{CH} \cdot \text{N} \cdot \text{CH} \cdot \text{CMe}]$. — Misc. aq., alc., eth. Prob. v. weak base. — \textcircled{P} Sp. gr. 1.0441 (0/4). — $\text{B.Pk.} \ddagger$ yel. pr. fr. alc., d.s. c. aq., m.p. 133°. — B.HAuCl_4 , yel. lft. fr. HCl (deod. by aq.), m.p. 180°.
II63	136		1,3-Diaminopropane, $\text{NH}_2 \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{NH}_2$. — Fumes in moist air. Misc. w. alc., eth., bz. — $\text{B.Pk.} \ddagger$ yel. d.s. lft.
II64	138-9	115	1-Ethylmorpholine, $[\text{NEt} \cdot (\text{CH}_2)_2 \cdot \text{O} \cdot (\text{CH}_2)_2]$. — Mobile liq. w. ammon. odor. E.s. aq., alc. Sp. gr. 0.900 (20/4). — $\text{B.Pk.} \ddagger$ ndl., m.p. 189-90°.
II65	139-40	115	4-Aminoheptane, $\text{Me} \cdot (\text{CH}_2)_2 \cdot \text{CH}(\text{NH}_2) \cdot (\text{CH}_2)_2 \cdot \text{Me}$. — Odor ammon. Sp. gr. 0.767 (20/4). — \textcircled{P} Prim. amine.* — B.HCl , ndl., m.p. 241-2°.

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 ¶ = T. 2.1. ** = T. 2.36. ‡ = T. 2.37.

(ORDER II, SUBORDER I.)

No.	Boiling-point (C°).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
II66	139.5-40.3c.	129 $k \cdot 10^4 = 5$	† Diisobutylamine, $(Me_2CH.CH_2)_2NH$. — Odor unpleasant, ammon. V.d.s. aq. Sp. gr. 0.749 (15°). — ② Gives Simon's react. (T. 2.28). Sec. amine.* — ① Add 3 drops B to 3 cc. c. sat. aq. sol. oxalic ac. Heat to dis. ppt. Cool well. Filter. Wash w. 1 cc. alc. Dry on tile 15 min. at 100°. The product, $B_2C_6H_8O$, is obtained in ndl., m.p. 250.2-1.2° u.c. (268-9° c.). Rapid heating near m.p. is necessary to prevent complete volatilization.
II67	140-2	113	2,4-Dimethylpiperidine, $[NH.CHMe.CH_2.CHMe.CH_2.CH]$. — E.s. aq. Sp. gr. 0.8615 (0°). — ② Sec. amine.* — B.HCl, e.s. aq., m.p. 235°.
II68	141-3 (746 mm.)		3,5-Dimethylpyrazolidine, $[NH.NH.CHMe.CH_2.CHMe]$. — Odor faintly ammon. "Strong diacid base." — ② Reduces h. Fehling's sol. — B.2Pk,† ndl., m.p. 129-30° d.
II69	142.5c.	107	2,6-Dimethylpyridine, Lutidine, $Me_2C_6H_4N$. — [Fr. distn. of coal, shale, & animal oils.] Misc. w. c. aq., sol. becoming turbid on warming. Sp. gr. 0.9420 (0°). Tert. amine.* — B.Pk,† light-yel. ndl. fr. h. aq., m.p. 161°. — B.HCl.HgCl ₂ , ppt. crystg. fr. ac. sol., m.p. 186°, 188°, 191.5°.
II70	141-3	113	d,l- α -Ethylpiperidine, $[NH.CHEt(CH_2)]$. — S. in 20 vol. aq. Sp. gr. 0.867 (0/0). Sec. amine.* — B.HCl, m.p. 181°. — B.Pk,† pr., m.p. 133°.
II71	142	115	2-Aminoheptane, $Me.CH(NH_2).(CH_2)_4.Me$. — D.s. aq. E.s. alc. Sp. gr. 0.781 (0°). — ② Prim. amine.* — B.HCl, ndl., m.p. 133°. — B. ₂ H ₄ C ₂ O ₄ , tbl., m.p. 204-5° d.
II72	143.1c.	93	γ -Picoline, 4-Methylpyridine, $Me.C_6H_4N$. — [Fr. coal tar.] — Sp. gr. 0.957 (15/4). Tert. amine.* — B.Pk,† silky ndl., m.p. 167°, d. c. aq. — B.HCl.HgCl ₂ , m.p. 128°.
II73	143.5c.	93	β -Picoline, 3-Methylpyridine, $Me.C_6H_3N$. — [Fr. coal tar.] Misc. w. aq. Sp. gr. 0.9726 (0/4). $n_D^{20} = 1.50720$. Tert. amine.* — B.Pk,† lust. ndl. fr. alc., mod. s. aq., less s. alc., m.p. 149-50°. — B.HCl.2HgCl ₂ , ppt. of fine ndl., alm. i. c. aq., s. dil. HCl, m.p. 146°.
II74	143		6-Dimethylamino-hexene(1), $CH_2:CH.(CH_2)_4.NMe_2$. — Odor piperidine-like. Not misc. w. aq. — Sp. gr. 0.967 (15°).
II75	144		Pyrazoline, $[NH.N:CH.CH_2.CH]$. — Weak, basic cocoa-like odor. Oxidized in air. Misc. w. aq., alc. — B.HCl, pr. d.s. c. alc., m.p. 130°; dil. sol. colors pine splinter yel.! — B.Pk,† yel. ndl. fr. alc., m.p. 130°.
II76	145		Diethylethylenediamine, $NEt_2.CH_2.CH_2.NH_2$. — Fumes in moist air. Sp. gr. 0.827 (18.5°). — B.2Pk,† ndl. dec. at 211° w. frothing. — B. ₂ H ₄ PtCl ₆ , § pr. fr. alc., d. 230°.
II77	145-6	127	2,4,6-Trimethylpiperidine, Copellidine, $[NH.CHMe.CH_2.CHMe.CH_2.CHMe]$. — Sp. gr. 0.8475 (4°). Sec. amine.* — B. ₂ H ₄ PtCl ₆ , § or. lft., m.p. 205°, d. 242-4°.
II78	145.5	129	2-Amino-2,5-dimethylhexane, $Me.CMe(NH_2).(CH_2)_3.CHMe.Me$. — ② Prim. amine.* — B.HCl, m.p. 157-60°.
II79	146		Piperylhydrazine, $C_8H_{10}N.NH_2$. — Oil of penetrating ammon. odor. Misc. w. aq., alc., eth., bz., ligr. Sp. gr. 0.9283 (14.6°). — ② Reduces ammon. AgNO ₃ , sol. in cold. Gives carbonyl-amine react. (T. 2.12). — B.HCl, tbl. fr. alc., m.p. 162°, 163-4°.
II80	146	143	Ethylpropylisobutylamine, $N.Et.Pr.C_6H_5$. — Tert. amine.*
II81	147		Triacetone, $[NH.CMe_2.CH:CH.CH_2.CMe_2]$. — Odor piperidine-like. Sec. amine.* — Nitroso deriv., ** yellowish tbl. of camphorous odor. — B.HCl, e.s. aq., d.s. alc.
II82	145-50	141	Allyldipropylamine, $C_6H_5.NPr_2$. — Tert. amine.* — B. ₂ H ₄ PtCl ₆ , thick or.-red cryst., changing to B. ₂ H ₄ PtCl ₆ (yel. ndl. of m.p. 152-3°) when boiled some time w. aq.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
II83	148.6c.	107	2-Ethylpyridine, Et.C ₆ H ₅ N. — Tert. amine.* Sp. gr. 0.9371 (17 $^{\circ}$). — Oxidation w. KMnO ₄ gives picolinic ac. (No. 2.130). — B ₂ .H ₂ PtCl ₆ , § m.p. 165–7° d. — B.HCl.2HgCl ₂ , ndl. fr. aq., m.p. 103–6°.
II84	149–50		s-Diethylethylenediamine, EtNH.CH ₂ .CH ₂ .NHEt. — Exposed to moist air forms solid hydrate. — B.H ₂ PtCl ₆ .2H ₂ O, § or-yel. pr. fr. h. aq., m.p. 223–4°.
II85	149 sl. d.		Ethyl Aminoacetate, NH ₂ .CH ₂ .CO ₂ Et. — Odor basic & cocoa-like. Leaves residue in flask on distn. which cryst. fr. h. aq. in silky ndl. S. aq., alc., eth. Unstable & e. saponified. Sp. gr. 1.0358 (11.8/4). — ⊕ Should give N in T. 2.4.
II86	149–50		N-Propylpiperidine, C ₄ H ₁₀ N.Pr. — Tert. amine.* — B.Pk, † 121°; 108°.
II87	148–53		Allylisoamylamine, C ₆ H ₅ .NH.C ₆ H ₁₁ . — Sec. amine.* Sp. gr. 0.778 (18 $^{\circ}$).
II88	151–2	125	N-Allylpiperidine, C ₆ H ₅ .N.C ₆ H ₁₁ . — Tert. amine.* Sp. gr. 0.8445 (18.5 $^{\circ}$).
II89	151–3c.	113	Heptanaphtheneamine, C ₇ H ₁₂ .NH ₂ . — V.s. aq., alc. “Strong base.” — ⊕ Prim. amine.*
II90	152.6c. (154–5)	113	dL-3-(β)-Ethylpiperidine, [NEt.(CH ₂) ₃]. — Odor coniine-like. Fumes in air. D.s. aq. Sp. gr. 0.8658 (0 $^{\circ}$). — ⊕ Tert. amine.* — B.Pk, † ndl. d.s. c. aq., m.p. 63°. — B ₂ .H ₂ PtCl ₆ , § mod. s. cryst., m.p. 183–4°.
II91	154; 156.5	107	† α-Lutidine, Dimethylpyridine, Me ₂ C ₆ H ₄ N. — [Fr. bone oil.] Sp. gr. 0.947 (0 $^{\circ}$); 0.938 (0 $^{\circ}$). — B.HAuCl ₄ , lust. yel. lft. decd. by boiling w. aq. giving red lft.
II92	153–8		Diethylenedimethylamine, [NMe.CH ₂ .CH ₂ .NMe.CH ₂ .CH ₃]. — B.2HCl (100 $^{\circ}$), pr., m.p. 247–50° d.
II93	154–7c.	117	Trimethylamine Oxide, Et ₃ NO. — Thick oil. Somewhat s. aq. Sp. gr. 0.893 (0 $^{\circ}$). — ⊕ Reduces Ag, Cu, & Hg salts.
II94	155–6 sl. d.		Diethylaminoacetone, Et ₂ N.CH ₂ .CO.Me. — Misc. aq. — B ₂ .H ₂ PtCl ₆ , § cryst., m.p. 176°.
II95	155–6c.	137	Triallylamine, (C ₃ H ₇) ₃ N. — Odor v. unpleasant. Sp. gr. 0.809 (14 $^{\circ}$). — ⊕ Tert. amine.*
II96	155c.	143	Diethylisoamylamine, Et ₂ N.C ₆ H ₁₁ . — D.s. aq. — ⊕ Tert. amine.* — Picrate, † yel. ndl., m.p. 75°.
II96-1	155c.		2,5-Dimethylpyrazine, Ketine, Glykoline, [CH:CM ₂ .N:CH-CM ₂ :N]. — [In fusel-oil.] — M.p. 15°. Misc. w. aq., alc., eth. Sp. gr. 0.9896 (18/4). — B.Pk, † lust. yel. cryst., s. h. aq.; e.s. h. alc., m.p. 157°. — B.HAuCl ₄ .H ₂ O, lust. red-yel. ndl., m.p. when anhydrous 153°. — B.HCl, v.s. aq., sbl.
II97	156 ± 1c.	115	† Heptylamine, Me.(CH ₂) ₆ .CH ₂ .NH ₂ . — Odor, unpleasant, basic distinctly ammon. Alm. i. aq. Sp. gr. 0.78 (20 $^{\circ}$). — ⊕ Gives Rimini's react. (T. 2.25). Prim. amine.* — ⊕ Mix 3 drops B w. 3 drops ethyl oxalate. Dis. ppt. in 6–7 cc. 50% alc. Cool well. Filter. Wash w. 1 cc. dil. alc. Dry 15 min. on porous tile at 100°. The product, diheptyloxamide, is obtained in colorless cryst., m.p. 131–2° u.c. (132.8–3.8° c.).
II98	157c.	143	† Tripropylamine, NPr ₃ . — Odor ammon. V.d.s. aq. Sp. gr. 0.756 (18 $^{\circ}$). — ⊕ Tert. amine.* (T. 2.25 & 2.28, negative.) — ⊕ Mix 2 drops B w. 1 cc. aq. Add 3 cc. c. sal. aq. picric ac. sol. Filter. Wash w. 1 cc. aq. Dry 15 min. on porous tile at 100°. The product, B.Pk, † is obtained in soft yel. ndl., m.p. 115–6° u.c. (116.3–7.3° c.).

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(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I198-I	157; 159		2,4-Dimethylpyridine, 2,4-Lutidine, Me₂C₆H₃N. — [In coal tar.] — Sp. gr. 0.9380 (14 $^{\circ}$). 0.9493 (0/4). — B.Pk, No. 2.3515. — B.HCl.2HgCl ₂ . $\frac{1}{2}$ H ₂ O, silky ndl. (ppt. fr. h. acid sol. by HgCl ₂), m.p. 132 $^{\circ}$.
I199	158		α -Coniceine, C ₈ H ₁₁ N. — Odor coniine-like. D.s. aq. Sp. gr. 0.893 (15 $^{\circ}$). Tert. amine.* — B.Pk, \ddagger yel. ndl. fr. alc., alm. i. aq., d.s. c. alc., m.p. 225 $^{\circ}$ d.
I200	159c.	75	1-Methylamino-ethanol(1), MeNH.CH₂.CH₂.OH. — Odor fishy. Misc. aq., alc. Strong caustic base. Sp. gr. 0.937 (20 $^{\circ}$). — \oplus HgCl ₂ gives white ppt. fr. c., or or.-red fr. h. sol. — B.Pk, \ddagger matted ndl., m.p. 148–50 $^{\circ}$, e.s. alc.; alm. i. eth.
I201	158–60	k.10 ⁴ = 5.4	Putrescine, Tetramethylene diamine, NH₂.CH₂.(CH₂)₂.CH₂.NH₂. — [Non-toxic ptomaine.] — Odor piperidine-like. Fumes slightly in air & absorbs CO ₂ . E.s. aq.; d.s. eth. — B.2Pk, \ddagger greenish-yel. silky ndl., alm. i. c. aq., d. 250 $^{\circ}$. — B.2C ₁₀ H ₁₃ N ₂ O ₂ (picrolonate), \ddagger fine yel. ndl. s. in 13,157 pt. c., or 653 pt. boiling aq. — \oplus Prepare the dibenzoyl deriv., silky ndl. i. aq.; d.s. c. alc.; e.s. h. alc.; & (unlike cadaverine) pptd. fr. alc. sol. by eth., m.p. 175–6 $^{\circ}$. Cf. procedure given for preparation of corresp. deriv. of cadaverine, No. 2.1232.
I201-I	159–60c.		2,5-Dimethylpyridine, Me₂C₆H₃N. — [In coal-tar & shale oils.] — E.s. c., less s. warm aq.; misc. alc., eth. Vol. w. st. — Tert. amine.* — B.Pk, \ddagger ndl. d.s. alc., m.p. 165.5 $^{\circ}$; 151 $^{\circ}$, 156 $^{\circ}$.
I202	160	129	Dibutylamine, (C₄H₉)₂NH. — \oplus Should give Simon's react. Sec. amine.* — Picrate, \ddagger lft. s. in 157 pt. c. aq., m.p. 59.5 $^{\circ}$.
I203	161	117	2-Diethylamino-ethanol(1), Et₂N.CH₂.CH₂.OH. — Misc. w. aq. Tert. amine.*
I204	161.5		δ-Coniceine, 1-Piperolidine, C₈H₁₁N. — D.s. aq. Sp. gr. 0.9012 (15/4). Tert. amine.* — B.Pk, \ddagger m.p. 226 $^{\circ}$.
I205	162.3c.	123	Tropidine, (Tropene), C₈H₁₁N. — Strong coniine-like odor. V.e.s. c. aq., but d.s. h. aq., v.s. alc., eth. Sp. gr. 0.9467 (19/4). Mol. refraction, 61.73 $^{\circ}$. Tert. amine.* — B.Pk, \ddagger cryst. fr. h. aq. in light yel. pr., m.p. abt. 285 $^{\circ}$ w. frothing. — B.HAuHCl ₄ , \parallel yel. cryst. ppt., m.p. 205 $^{\circ}$.
I206	162–5	127	2-Methyl-5-ethylpiperidine Hexahydrocollidine, "NH.CHMe-(CH₂)₂CHEt.CH₂". — Odor ammon., penetrating. D.s. aq. Sp. gr. 0.8362 (18 $^{\circ}$). — Sec. amine.* — B ₂ H ₂ PtCl ₆ (110 $^{\circ}$) yel. ndl., v.s. aq.; i. alc., m.p. 145–7 $^{\circ}$.
I207	163	133	Aminoacetal, NH₂.CH₂.CH(OEt)₂. — Odor v. unpleasant. V.s. aq. alc. Combines w. CO ₂ . — Prim. amine.* — Picrate, \ddagger yel. d.s. ndl., m.p. 142–3 $^{\circ}$.
I208	163.5–4.5c.	107	3,4-Dimethylpyridine, (Lutidine), Me₂C₆H₃N. — [In coal tar.] — Tert. amine.* Oxidized by KMnO ₄ to cinchomeronic ac., No. 2.470. — B.HCl.2AuCl ₄ , \parallel fine yel. ndl., m.p. 160–2 $^{\circ}$ (giving dark oil).
I209	165c.	107	3-Ethylpyridine, β-Lutidine, Et.C₆H₃N. — Odor unpleasant. V.d.s. c. aq.; less s. h. aq. Sp. gr. 0.9585 (0/4). Oxidation by CrO ₃ , mixt. gives nicotinic ac., No. 2.410. — Tert. amine.* — B.Pk, \ddagger yel. ndl., d.s. aq., m.p. 128–30 $^{\circ}$. — B.HCl.HgCl ₂ , pr., m.p. 132 $^{\circ}$.
I210	164–6	107	4-(γ)-Ethylpyridine, Et.C₆H₃N. — Sp. gr. 0.9522 (0 $^{\circ}$). Tert. amine.* Oxidation by KMnO ₄ gives isonicotinic ac., No. 2.500. — B.Pk, \ddagger m.p. 168 $^{\circ}$. — Picrate, \ddagger m.p. 163 $^{\circ}$, 168 $^{\circ}$. — B.HCl.2HgCl ₂ , lft., m.p. 149–50 $^{\circ}$; 150–2 $^{\circ}$.
I211	165–8	121	2-(α)-Propylpyridine, Conyrine, Pr.C₆H₃N. — Lighter than aq. Oxidation by KMnO ₄ gives picolinic ac., No. 2.130. Tert. amine.* Salts v.s. — B ₂ H ₂ PtCl ₆ , \ddagger 4-sided or.-yel. monoclin. tbl., m.p. 159–60 $^{\circ}$; 172 $^{\circ}$.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I212	166		Triformalmethylamine, Trimethyltrimethylenetriamine, $[NMe_2CH_2NMeCH_2NMeCH_2]^3$. — [Fr. methylamine + formic aldehyde.] — Sp. gr. 0.921 (18.7°). — B.Pk, \dagger pr. fr. chlf., m.p. 127–8°. e.s. aq. — Mixed w. CS_2 forms dimethylformocarbostibaldine readily, fine ndl. fr. alc., m.p. 96°.
I213	165–8	135	2,4,5-Trimethylpyridine, $Me_2C_6H_3N$. — [In coal tar.] — Tert. amine.* — B.Pk, \dagger or. ndl., m.p. 128–31°.
I214	166–7c.	127 $k \cdot 10^3 = 1.3$	† d-Coniine, $[NH_2CH(n-Pr).(CH_2)_4]^2$. — [Narcotic alkaloid fr. Conium maculatum L., or "poison hemlock."] — Odor of natural base unpleasant, basic, like mouse urine! S. in 90 pt. aq.; less s. h.; e.s. alc., eth., chlf. Vol. w. st. Sp. gr. 0.8444 (20°). $[\alpha]_D^{20} = +15.7^\circ$. $n_D^2 = 1.4505$. <i>Moist HCl gas causes fuming.</i> Sec. amine* giving or.-red tint in T. 2.28. — (1) A clear sol. of 1 drop B in 2 cc. aq. becomes milky while when heated nearly to boiling, clearing again on cooling. (Some other liquid bases show similar behavior.) — (2) Charac. cryst. pptn. w. dinitroanthracrysonedisulphonic ac. described by Rosenthaler & Gerner, Z. anal. Chem., 49, 340 (1910). — For other reactions cf. Abderhalden, 5, 21, & Dilling, Pharm. J. [4], 29, 34 (1909).
			① (a) Mix 1 drop B w. 1 drop conc. HCl in 3-inch t.t. Add 1 cc. eth. Crush mass w. rod & decant eth. Wash w. eth. Transfer to porous tile & dry 15 min. at 100°. B. HCl is obtained as cryst. non-deliq. solid, m.p. 216.4–7.4° u.c. (222–3° c.). — (b) Heat sol. of 3 drops B in 5 cc. alc. w. 0.10 g. phthalic anhydride to obtain clear sol. Evap. on water-bath. Dis. sticky residue by stirring in 8 cc. h. 25% alc. containing 4 drops dil. HCl (sp. gr. 1.12). Cool well, & shake vigorously & persistently until the ppt., which slowly appears, no longer increases. Filter. Wash w. 3 cc. 25% alc. Recryst. fr. 3 cc. dil. alc. as before w. v. persistent cooling & shaking. Dry on porous tile 30 min. at 45°. The product, conylenephthalamic ac., $C_{10}H_{16}N.CO.C_6H_4.CO_2H$, is obtained in rosettes of microcryst. ndl., m.p. 154.2–5.2° u.c. (166.8–7.8° c.). (Yield small, but quite sufficient if time enough is allowed for each crystn.) — $B_2C_{10}H_8O_4N_2$ (picrolonate), \ddagger yel. rhombohedra, m.p. 195.5°. — $B_2H_2PtCl_6 \cdot H_2O$, § oily at first, then or.-yel. cryst.; deep red 4-sided pr. fr. abs. alc.; m.p. 78°; m.p. when anhydrous, 175°.
I215	166–7	143	4-Amino-2,6-dimethylheptane, $Me.CHMe.CH_2.CH(NH_2).CH_2.CHMe.Me$. — Sp. gr. 0.772 (20/4). — Prim. amine.* — B.HCl, ndl., m.p. 247–8°.
I216	167c.	125	Tropane, Hydrotropidine, $C_8H_{11}N$. — Odor coniine-like. D.s. c. aq.; less s. h. aq. Sp. gr. 0.934 (15/4). Tert. amine.* — $B_2H_2PtCl_6$, cryst. fr. h. conc. sol. in long light or red ndl. which suddenly change on cooling to small red tbl.; dimorphous; m.p. 220–1° d.! — Picrate, \ddagger gold-yel. pr. fr. alc., m.p. 280–1° d.
I217	167c.	129	1-Amino-2-propylpentane, $Me.CH_2.CH_2.CHPr.CH_2.NH_2$. — D.s. aq.; misc. eth. — ② Prim. amine.* — $B_2H_2PtCl_6$, § golden lft. fr. aq., m.p. 211° d.
I218	168		Tetraethylmethylenediamine, $(NEt_2)_2CH_2$. — Odor disagreeable. S. in 10 pt. aq.; misc. alc., eth. Sp. gr. 0.810 (18.7°).
I219	167.9c.	89	2-Ethylamino-ethanol(1), $EtNH.CH_2.CH_2.OH$. — Fumes absorbing aq. & CO_2 fr. air. Caustic. E.s. aq., alc., eth. Sp. gr. 0.914 (20/4).
I220	169c.	113	Aminocycloheptane, $[CH(NH_2).(CH_2)_6]^2$. — Absorbs CO_2 fr. air.
I221	169–70	107	3,5-Dimethylpyridine, $Me_2C_6H_3N$. — Mod. s. c. aq. Sp. gr. 0.9614 (0/4). Tert. amine.* — $B_2H_2PtCl_6$, § red cryst., d.s. aq., m.p. 255–6° d. — B_2HAuCl_4 , yel. ndl., m.p. 149°, d.s. aq. — $HgCl_2$ salt, m.p. 170°.

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(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I221-1	169-0.5		2,3,5-Trimethylpiperazine, $\text{NH} \cdot \text{CHMe} \cdot \text{CHMe} \cdot \text{NH} \cdot \text{CHMe} \cdot \text{CH}_3$. — Solid at 0 $^{\circ}$. E.s. aq., alc., chlf. — \textcircled{D} Sec. amine.* — $\text{B.2Pk, \dagger fine yel. ndl., v.d.s. aq., alc.; blackens abt. 250}^{\circ}$. — Dinitroso deriv.,** lt. fr. alc., lt. fr. alc., v.d.s. aq., m.p. 94-6 $^{\circ}$.
I222	171c.	61	2-Aminoethanol(1), $\text{NH}_2 \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{OH}$. — Viscous, strongly basic oil w. faint odor. Absorbs aq. & CO_2 . Misc. aq., alc.; s. in abt. 100 pt. eth. Sp. gr. 1.022 (20 $^{\circ}$). — $\text{B.Pk, \dagger 6-sided tbl. fr. alc., m.p. 159.5}^{\circ}$.
I223	171-2	121	2,4,6-Trimethylpyridine, γ-Collidine, $\text{Me}_3\text{C}_6\text{H}_3\text{N}$. — [In coal tar.] D.s. c. aq. Sat. sol. gives ppt. on warming. Sp. gr. 0.917 (15 $^{\circ}$). Gives cryst. ppt. w. AgNO_3 which dissolves on heating. — $\text{B.Pk, \dagger silky yel. ndl., d.s. aq., e.s. alc., m.p. 155-6}^{\circ}$. — $\text{B}_2\text{H}_2\text{PtCl}_6$,§ yel. cryst. ppt., m.p. 217 $^{\circ}$. — B.HAuCl_4 , woolly ndl. fr. h. aq., m.p. 112-3 $^{\circ}$, melting under boiling aq.
I224	172-3	102	1,4-Diamino-2-methylbutane, $\text{NH}_2 \cdot \text{CH}_3 \cdot \text{CHMe} \cdot \text{CH}_2 \cdot \text{CH}_3 \cdot \text{Me}$. — Fumes, absorbing aq. & CO_2 . Sp. gr. 0.884 (20/4). — $\text{B.2HCl.2AuCl}_4.2\text{H}_2\text{O}$, flat pr., m.p. 191 $^{\circ}$ after losing 2 aq. at 100 $^{\circ}$.
I225	173-4	121	2-Methyl-5-ethylpyridine, Aldehydecollidine, $\text{Me.EtC}_6\text{H}_3\text{N}$. — Strong aromatic odor. I. aq., dil. alc.; e.s. alc., eth., conc. H_2SO_4 . Fumes w. HCl. Sp. gr. 0.9184 (23 $^{\circ}$). — $\text{B.Pk, \dagger greenish yel. 4-cornered tbl. D.s. aq., m.p. 164}^{\circ}$! — Gives ppt. w. HgCl_2 . — $\text{B}_2\text{H}_2\text{PtCl}_6$,§ or-red ndl., alm. i. c. aq., m.p. 182 $^{\circ}$. Unstable in aq. — B.HAuCl_4 , yel. lt., mod. s. aq., m.p. 87 $^{\circ}$.
I226	173	125	γ-Coniceine, α-Propyltetrahydropyridine, $\text{Pr.C}_6\text{H}_3\text{N}$. — Odor coniine-like. V.d.s. aq. Sol. strongly alk. Sp. gr. 0.8724. — Sec. amine.* — $\text{B.Pk, \dagger m.p. 62}^{\circ}, 65^{\circ}, 78^{\circ}$.
I227	abt. 173	129	2-Aminooctane, $\text{Me.CH(NH)_2(CH_2)_6Me}$. — Sp. gr. 0.79 (0 $^{\circ}$). — Prim. amine.*
I228	174	127	3-(β)-Propylpiperidine, $\text{Pr.C}_6\text{H}_{10}\text{N}$. — Odor coniine-like. Fumes w. HCl. Browns in air. Sp. gr. 0.8475 (26/4). Sec. amine.* — $\text{B.HCl, ndl. fr. aq., i. eth., m.p. 127-9}^{\circ}$. — $\text{B.Pk, \dagger m.p. 121.5}^{\circ}$. — B.HAuCl_4 , lemon-yel. ndl. fr. aq., m.p. 95-6 $^{\circ}$, browning at 88 $^{\circ}$.
I229	174d.		Ethyl d,l-α-Aminoisovalerianate, $\text{Me}_2\text{CH} \cdot \text{CH}(\text{NH}_2) \cdot \text{CO} \cdot \text{Et}$. — E.s. aq. Unstable, yielding corresponding piperazine even by long keeping. Sp. gr. 0.9616 (15/4). — Saponify & isolate EtOH. (Procedure 2-B of Gen. T. V, Vol. I.) — Picrate,† yel. ndl., m.p. 135.5 $^{\circ}$ c., slowly decd. by boiling aq.
I230	175		1,4-Diamino-2-methylpentane, $\text{NH}_2 \cdot \text{CH}_3 \cdot \text{CHMeCH}_2 \cdot \text{CH}(\text{NH}_2) \cdot \text{Me}$. — $\text{B.H}_2\text{C}_2\text{O}_4$, fine ndl., d.s. abs. alc., m.p. 244 $^{\circ}$.
I231	abt. 176	129	Octylamine, $\text{Me} \cdot (\text{CH}_2)_6 \cdot \text{CH}_2 \cdot \text{NH}_2$. — Alm. i. aq. Prim. amine.* — Picrate, m.p. 112-4 $^{\circ}$.
I232	178-9		† Cadaverine, Pentamethylenediamine, $\text{NH}_2 \cdot \text{CH}_2 \cdot (\text{CH}_2)_4 \cdot \text{CH}_2 \cdot \text{NH}_2$. — [A non-toxic ptomaine.] — Syrupy liq. of peculiar piperidine-like odor, fuming, absorbing CO_2 , & solidifying in moist air. Solidified by freezing mixture, m.p. abt. +9 $^{\circ}$. E.s. aq.; d.s. alc.; v.d.s. eth. Sp. gr. 0.885 (15 $^{\circ}$). Vol. w. st. — \textcircled{D} Gives Rimini's react. (T. 2.25). — \textcircled{D} Dis. 1 drop B in 5 cc. 10% aq. NaOH sol. Add 8 drops benzoyl chloride. Shake persistently, breaking up lumps w. rod until ppt. becomes hard & granular, and odor of benzoyl chloride entirely disappears. Filter. Wash thrice w. 2 cc. c. aq. Dry on porous tile. Recryst. fr. 8 cc. h. bz. Filter hot if sol. is not clear. Dibenzoylcadaverine separates in small colorless scales & ndl. on cooling. Dried 10 min. at 100 $^{\circ}$, it melts at 129-30 $^{\circ}$ u.c. (130.8-1.8 $^{\circ}$ c.). — $\text{B.2Pk, \dagger v.d.s. aq., h. alc.; yel. ndl., m.p. 221}^{\circ}$. — $\text{B.2C}_{10}\text{H}_8\text{O}_4\text{N}_4$ (picrolonate),‡ or-yel. cryst. s.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
			in 7575 pt. c. or 357 pt. boiling aq., m.p. (after blackening) 250 $^{\circ}$. — $\text{B}_2\text{H}_2\text{PtCl}_6$, § or.-yel. pr. d.s. c. aq., d. 215 $^{\circ}$. — Phenylurea deriv., $\text{C}_8\text{H}_{10}(\text{NH}_2\text{CO}_2\text{NHPh})_2$, ndl. i. usual solvents, m.p. 207–9 $^{\circ}$. (Fr. suspension B in dry eth. + phenylisocyanate. Z. physiol. Chem. 43, 355.)
I233	183c.	135	Dimethylbenzylamine, $\text{Ph.CH}_2\text{NHMe}_2$. — Less s. h. than c. aq.; misc. alc., eth. — (P) Tert. amine.*
I234	183–5	141	5-Amino-1,1,3-trimethylcyclohexane, $^5\text{CMe}_2\text{CH}_2\text{CHMe}_2\text{CH}_2\text{CH}(\text{NH}_2)\text{CH}_2$. — Strongly basic smelling oil. — (P) Prim. amine.* (T. 2.4.)
I235	184.4c.	93 $k.10^{10} = 5$	† Aniline, PhNH_2 . — Odor, basic, peculiar. Taste (poisonous) burning. S. in 32 pt. aq. at 16 $^{\circ}$; misc. alc., eth. Sp. gr. 1.0219 (20/4). — (P) (1) Prim. amine.* The carbalamine react., T. 2.12, is convenient & v. delicate — (2) Dis. 1 drop B in 10 cc. aq. & add 4 drops c. sat. filtered aq. sol. bleaching powder. Within a minute an intense VRSI color (observed in t.t. by direct transmitted light fr. sky) appears! (Pure! methyl- or dimethyl-aniline does not give this color.) (D) Mix 2 drops B w. 4 drops acetic anhydride in small t.t. & heat to boiling. Cool. Add 2 cc. aq. Boil to secure clear sol. Cool well w. vigorous shaking. Wash ppt. on filter w. 2 cc. c. aq. Recryst. fr. 2 cc. h. aq. Strong cooling & vigorous shaking are necessary to start cryst.! Filter. Dry on porous tile at 90 $^{\circ}$ for 15 min. The product, acetanilide, is obtained in pearly scales, m.p. 111.7–2.1 $^{\circ}$ u.c. (112.7–3.1 $^{\circ}$ c.).
I236	184c.	107 $k.10^6 = 1.95$	† Benzylamine, PhCH_2NH_2 . — Odor strongly ammonial! Odor of warm aq. sol. w. 1 drop B per 10 cc. faintly aromatic, w. burning (pungency No. 2 in T. 2.29) taste. Sp. gr. 0.980 (20/0). — (P) Prim. amine giving RVT1 color in Rimini's react. (T. 2.25) within abt. 1 min. — A drop of B exposed to air on watch glass soon solidifies fr. absorption of CO_2 & moisture to frost-like incrustation effervescent w. HCl. (D) (a) Convert into corresponding picramide by procedure of T. 2.22. Product recryst. fr. gl. ac. ac. forms OY-YO ndl., m.p. 142 $^{\circ}$ u.c. — (b) Dis. 2 drops B in 3 cc. alc. Add 6 drops CS ₂ & heat to boiling. Evap. on water-bath in v. small glass capsule until cryst. begin to separate. Cool slowly. Filter. Wash w. few drops alc. Dry on porous tile at room temp. The product, benzylamine benzylidithiocarbamate, is obtained in pearly white scales, d. 127–8 $^{\circ}$ u.c. (128.7–9.7 $^{\circ}$ c.) — (c) Dis. 3 drops B in 2 cc. aq. Add sol. of 0.15 g. oxalic ac. in 3 cc. aq. Heat to boiling. Cool well. Dry ppt. of hexag. pr. 16 min. at 90 $^{\circ}$. The product, $\text{B}_2\text{H}_2\text{C}_2\text{O}_4$, melts w. decom. at 176 $^{\circ}$ u.c. (179.3 $^{\circ}$ c.).
I237	185	121	Methylbenzylamine, $\text{PhCH}_2\text{NHMe}_2$. — (P) Sec. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_6$, § (at 100 $^{\circ}$), m.p. abt. 193 $^{\circ}$ d.
I238	185c.	135 $k.10^6 = 3.1$	Dimethyl-o-toluidine, $\text{Me.C}_6\text{H}_4\text{NMe}_2$. — Sp. gr. 0.9286 (20/4). — (P) Tert. amine.*
I239	185–6	143	1-Amino-2-methyloctane, $\text{CH}_3(\text{CH}_2)_5\text{CHMe}_2\text{CH}_2\text{NH}_2$. — (P) Prim. amine.* — B_2HCl , ndl., m.p. 130 $^{\circ}$.
I240	184–6	143	Triisobutylamine, $\text{N}(\text{C}_2\text{H}_5)_3$. — Not misc. w. aq. Sp. gr. 0.785 (21%). — (P) Tert. amine.*
I241	185–8	121	2,3,4-Trimethylpyridine, $\text{Me}_3\text{C}_6\text{H}_3\text{N}$. — Strong pyridine odor. S. aq.; e.s. alc., eth. Sat. aq. sol. becomes turbid on heating. Sp. gr. 0.9127 (15%). — (P) Tert. amine.* — B_2HAuCl_4 , ndl. fr. h. aq., v.d.s. c. aq., m.p. abt. 100 $^{\circ}$.
I242	187.5c.	121	1'-Aminoethylbenzene, $\text{Ph.CH}(\text{NH}_2)\text{Me}$. — S. in 24 pt. aq. at 20 $^{\circ}$. Absorbs CO_2 . Sp. gr. 0.9395 (15%). — (P) Prim. amine.* — M.p. of hydrochloride, 158 $^{\circ}$. — $\text{B}_2\text{H}_2\text{C}_2\text{O}_4$, pr. fr. h. aq., m.p. 238 $^{\circ}$. — Acetyl deriv., ¶ m.p. 57 $^{\circ}$.

Explanation of typographical signs used in this Division: * = T. 2.35. † = T. 2.23. § = T. 2.14. || = T. 2.13. ¶ = T. 2.1. ** = T. 2.36. ¶¶ = T. 2.37.

(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I243	186; 188	155	N-Isoamylpiperidine, $C_9H_{16}N.C_6H_{11}$.—Lighter than aq. D.s. aq. — \textcircled{P} Tert. amine.* — Picrate,† m.p. 133°.
I244	189-90	119	2-Allylpyridine, $C_7H_7.C_6H_4N$.—Sp. gr. 0.9595 (0°).—Tert. amine.* Oxidation gives picolinic ac. (No. 2.130).— $\textcircled{B}_2.H_2PtCl_6$ § (at 100°), ndl. v.d.s. aq., m.p. 185-6° d.
I245	190c.	157 $k.10^{10} = 9.6$	† Diisoamylamine, $(C_6H_{11})_2NH$.—Odor unpleasant, faintly ammon., & a little like amyl alc. V.d.s. aq. — \textcircled{P} Sec. amine,* but too difficultly s. aq. to give Simon's react. (T. 2.28) distinctly. (W. large x.s. acetaldehyde & enough alc. to dissolve, only a v. faint blue color is obtained in this test.)—① Prepare the dioxalate by procedure given under No. 2.1166. Product melts at 255-4° u.c.
—	192-5		Formamide.—Cf. No. 2.2783. Sp. gr. 1.337 (14/4).
I247	194c.; 193.1c.	121 $k.10^{10} = 2.4$	† Dimethylaniline, Ph.NMe. — Odor unpleasant aromatic, char-ac., somewhat aniline-like. Old preparations dark colored, stoppers of containers showing bluish stains. Alm. i. aq. Sp. gr. 0.9575 (20/4). Tert. amine.* — \textcircled{P} (1) Heat 1 drop B + crushed fragment $ZnCl_2$ of size of small pea + 4 drops benzaldehyde to b.p. of liq. in small dry t.t. for 15 sec. by small flame. Cool. Add 3 cc. alc. + 1 drop HCl . Sol. viewed by direct light fr. sky will be intense green. (Aniline & methyl aniline, if pure, give colorless sol.)—(2) Unlike methylaniline, gives no oily ppt. when pure in the "preliminary test" of No. 2.1249 w. $NaNO_2$ & acid.
			① Dis. 0.1 g. B in 1 cc. HCl (sp. gr. 1.20) + 2 vol. H_2O in t.t. cooled to 0° by surrounding by ice water. Add 1 cc. 10% $NaNO_2$ sol. Filter off ppt. after 10 min. Wash w. 5 cc. dil. HCl (sp. gr. 1.12). Dis. the YT1 cryst. of <i>p</i> -nitrosodimethylaniline hydrochloride in 10 cc. aq. Add 5 cc. sat. K_2CO_3 sol., & shake out w. two 10 cc. portions of eth. As the eth. evaporates YG-G lf. of <i>p</i> -nitrosodimethylaniline, m.p. 85° u.c., separate.
I248	193-5		Hygrine, $C_8H_{11}ON$.—[In Cusko leaves.] Turns brown in air. Sp. gr. 0.935 (17/4).— \textcircled{B} .Pk,† yel. ndl., m.p. 148°.—Oxime, m.p. 116-20°.
I249	195.7c.; 195.5c.; 193.8c.	107 $k.10^{10} = 2.6$	† Methylaniline, Ph.NHMe. — Odor somewhat aniline like. Becomes yel. & brownish on keeping. V.d.s. aq. Sp. gr. 0.9865 (20/4). Sec. amine.— \textcircled{P} Dis. 2 drops B in 3 cc. aq. + 3 drops conc. HCl . Add 2 drops c. sat. aq. $NaNO_2$ sol. An oily! ppt. of nitrosomethylphenylamine separates at once. (Dif. fr. dimethylaniline.)—① Dis. 4 drops B in t.t. in 5 cc. conc. H_2SO_4 . Cool well & add w. constant shaking 10 drops HNO_3 (sp. gr. 1.42). After 1 min. cool w. ice water & then pour, drop by drop, constantly shaking, into 10 cc. ice water. After further vigorous shaking filter off the granular ppt. & wash w. 2 cc. c. aq. Dis. in 10 cc. boiling 50% alc. If sol. is not clear, filter rapidly while hot. Recryst. fr. 3 cc. h. 50% alc. Dry 15 min. on porous tile at 100°. The product, tetranitromethylaniline, is obtained in lust. YT1 scales, m.p. 126.8-7° u.c. (128.4-0.6° c.).
I250	abt. 195	121	4-Methyl-3-ethylpyridine, β -Collidine, Me.Et.C ₆ H ₄ N.—Sp. gr. 0.9656 (0°). Tert. amine.* — \textcircled{B} .Pk,† m.p. 148-50°.
I251	195d.	141	† Pelletierine, $C_9H_{11}ON$.—[Fr. pomegranate bark, <i>Punica granatum</i> L.] Odor conine-like. Darkens & resinifies in air. E.s. alc., eth., chlf. Aq. sol. reacts alk. Vol. w. st. Fumes w. HCl . Sp. gr. 0.988 (0°). Dextrorotatory.— $\textcircled{B}.HAuCl_4$, m.p. 62°.
I252	195-6		2-Dimethylamino-1,3-xylene, Me ₂ N.C ₆ H ₄ .Me ₂ .—Odor camphor-like. Tert. amine.*
I253	195	153	1-Fenethylamine, $C_{10}H_{17}NH_3$.—Absorbs CO_2 . Sp. gr. 0.9095 (22°). $[\alpha]_D = -24.63^\circ$. Prim. amine.*

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I254	195	153	Thujoneamine, $C_{10}H_{17}NH_2$. — Odor coniine-like. Sp. gr. 0.8712 (20/4). $[\alpha]_D = +101.00^{\circ}$. — Prim. amine.*
I255	196 sl.d.		Ethyl 1- α -Aminoisobutyacetate (Ester of Leucine), $Me_2CH-CH_2CH(NH_2)CO_2Et$. — B.p. 83.5° (12 mm.). $[\alpha]_D^{20} = +17.86^{\circ}$. Prim. amine.* — Picrate, \ddagger m.p. 128° u.c. — [The d,l-ester has same b.p. w. peculiar, not unpleasant odor; is s. in 23 pt. c. aq., misc. w. alc., eth. & gives picrate, \ddagger m.p. 134° u.c. (r.h.).]
I256	197-9	82 $k.10^7 = 2.1$	1-Methylglyoxaline, 1-Methylimideazole, Oxalmethyline, $[NMe.CH:CH.N:CH^2]$. — Misc. aq. M.p. -6° . Sp. gr. 1.0363 (10°). — \tilde{B} .Pk, \ddagger golden ndl. d.s. aq., m.p. 158° . — $\tilde{B}_2H_2PtCl_6$, \S or.-red ndl. s. in abt. 20 pt. c. aq., m.p. $190-1^{\circ}$ (frothing).
I257	197-8	121	ω -Phenylethylamine, $Ph.CH_2CH_2NH_2$. [Fr. putrefaction of gelatine.] — Odor peculiar. S. in 24 pt. aq.; v.s. alc., eth. Absorbs CO_2 . Sp. gr. 0.958 (24/4). Prim. amine.* — \tilde{B} .Pk, \ddagger tetrag. pr. fr. alc., m.p. $171-4^{\circ}$. — $\tilde{B}.HCl$, tbl. fr. alc., m.p. 217° .
I258	198c.	153	† Camphylamine, $C_{10}H_{17}NH_2$. — Odor basic. A drop exposed to air for some time on watch glass solidifies to crust efferv. w. HCl . — (1) Dis. 1 drop. \tilde{B} in 2 cc. alc. Add 3 cc. c. sal. aq. picric ac. sol. Shake. Filter cryst. ppt. & wash w. 3 cc. aq. Recryst. fr. 7 cc. boiling aq. Dry on porous tile 15 min. at 40° . The resulting picrate is obtained in YT1 scales or ndl., m.p. $195.8-6.5^{\circ}$ u.c. (blackening).
I259	199.5c.	121	1'-Amino-1,2-xylene, $Me.C_6H_4.CH_2NH_2$. — Prim. amine.* — \tilde{B} .Pk, \ddagger yel. ndl. d.w.m. a. 170° . — Acetyl deriv., \P ndl. fr. alc., m.p. 69° .
I260	199c.	135	Ethylbenzylamine, $Ph.CH_2.NHEt$. — I. aq.; e.s. alc., eth. Sec. amine.*
I261	199-200	149	3-Dimethylamino-1,2-xylene, $NMe_2C_6H_4Me_2$. — Tert. amine.*
I262	199.5c.; 200c.	107 $k.10^6 = 3.5$	† o-Toluidine, $Me.C_6H_4.NH_2$. — Odor aniline-like. Slowly darkens on exposure to air & light. D.s. c. aq. Sp. gr. 1.0031 (15/15). Prim. amine.* Color reactions for distinguishing the three toluidines & aniline are described by Lorenz, Ann., 172, 180 (1874), by Rosenstiehl, Ann. chim. phys., 26, 232 (1872), by Nietzski, Ber. 10, 1157 (1877), & No. 2.1264. — (1) Prepare the corresponding picramide by T. 2.22. This deriv. after recrystn. fr. alc. is readily obtained in YO-O ndl., m.p. 164° u.c. — Acetyl deriv., \P $C_6H_4NH.CO.Me$, ndl. s. in 116 pt. aq. at 19° , m.p. 110° ; b.p. 296° .
I263	201	133	Methylethylaniline, $Ph.NMe.Et$. — Tert. amine.*
I204	abt. 201c. d.		† Dimethylamino-dimethylethylcarbinol Benzoate, $Me_2N.CH_2-CMe_2Et(Ph.CO_2)$. — [$\tilde{B}.HCl$ is the anesthetic stovaine. B.p. of \tilde{B} is that of a preparation made fr. commercial stovaine by Poulen Frères, Paris.] — Odor faint, unpleasant. I. aq.; e.s. alc., eth. Extracted fr. alk. sol. by eth. — (2) (1) Dis. 0.002 g. \tilde{B} or $\tilde{B}.HCl$ in 2 drops conc. H_2SO_4 & add small cryst. KIO_3 . An OY color [strawberry red w. yel. & brown stripes (Gadamer)] develops in the cold, changing to VRT2-RT2 after heating for a very few sec. — (2) Dis. 1 or 2 mg. \tilde{B} or $\tilde{B}.HCl$ in 4 drops conc. H_2SO_4 . Add 5 drops alc. Heat 3 or 4 min. at temperature of boiling water-bath. The sweetish aromatic odor of ethyl benzoate will be noticed. [A 1% stovaine sol. on the eye causes slight dilation of the pupil & disturbance in power of accommodation, as well as local anesthesia. (Kobert).] — $\tilde{B}.HCl$ (stovaine), cryst. powd., e.s. aq., alc., alm. i. eth., m.p. 175° ; aq. sol. reacts sl. acid; taste sl. bitter; causes numbness of tongue.

Explanation of typographical signs used in this Division: * = T. 2.35. ‡ = T. 2.23. § = T. 2.14. ¶ = T. 2.13. \ddagger = T. 2.1. ** = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C.).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I265	203c.	107 $k.10^{10} = 5.5$	<p>① Dis. 0.05 g. stovaine in 7 cc. aq. Add 3 cc. sat. aq. picric ac. sol. Heat to boiling. Cool slowly. Filter. Wash cryst. w. 2 cc. c. aq. Recryst. fr. 12 cc. boiling aq. Allow to stand for some hours. Filter off the fine YT1→ cryst. of B.Pk‡ and wash w. 1 cc. aq. Dry 16 min. at 100°. Picrate ‡ melts to clear yel. liq. at 114.5–6.5° u.c.</p> <p>† m-Toluidine, Me.C₆H₄.NH₂. — Odor aniline-like. Darkens on exposure to air & light. D.s. aq. Sp. gr. 0.9891 (20/4). Prim. amine.* — ② Dis. 1 drop B in 10 cc. c. H₂SO₄ (2 vol. H₂SO₄ sp. gr. 1.84 + 1 vol. aq.). Add 2 drops HNO₃ (sp. gr. 1.42). At end of 3 min. a red-or. color (RO-ROT1) will develop. Heat gently for 30 sec. Allow to stand 30 sec. longer & then cool. The color by direct transmitted light fr. sky will then be intense red (R). (Directions must be closely followed.) — [In this preliminary test: aniline after 3 min. gives no color; o-toluidine gives deep red-brown at once, solution being entirely opaque after 3 min.; methylaniline & ethylaniline quickly give a red, changing to brown and opaque in 3 min.; dimethylaniline & diethylaniline behave much like B. Cf. No. 2.1262 for bibliography.]</p> <p>① (a) Prepare the corresponding picramide by procedure of T. 2.22. The product, consisting of small OY cryst., melts at 127° u.c.! — (b) Heat to boiling in 3-inch t.t. 2 drops B + 4 drops acetic anhydride. Dis. product in 3 cc. boiling aq. Cool to abt. 50° & filter quickly through v. small filter. Cool filtrate for 39 min. or longer w. ice water, shaking vigorously at short intervals. Wash ppt. w. 1 cc. c. aq. Dry on porous tile at 60°. The resulting m-acetotoluide forms lust. scales, m.p. 65.5–6° u.c.</p>
I266	203	135	1'-Aminopropylbenzene, Me.CH(NH ₂).CH ₂ .Ph. — Prim. amine.*
I267	203c.	149	4-Dimethylamino-1,3-xylene, Me ₂ N.C ₆ H ₃ .Me ₂ . — Tert. amine.*
I268	204		5-Methylpyrazole, ⁵ CH:CM ₂ :N.NH ² . — Misc. aq. Sp. gr. 1.0227 (23.7/4). — B.Pk,‡ cryst. ppt., e.s. aq., m.p. 144°. — B ₂ H ₂ PtCl ₆ .2H ₂ O, § cryst. powd., e.s. aq., m.p. 181° d.
I269	204	121	1'-Amino-1,4-xylene, Me.C ₆ H ₄ .CH ₂ .NH ₂ . — V.s. aq. Absorbs CO ₂ . Sp. gr. 0.9520 (20/0). Prim. amine.* — B.HCl, ndl., m.p. 235°. — B.Pk,‡ m.p. 194–9° d. — Acetyl deriv., ¶ cryst. fr. alc., m.p. 107°.
I270	205c.	121 $k.10^{10} = 4.2$	<p>† Ethylaniline, Ph.NHEt. — Odor aniline-like. Darkens on exposure. V.d.s. aq. Sp. gr. 0.963 (20/4). — ② Sec. amine.* — ① Mix 1 cc. conc. HNO₃, w. 3 cc. conc. H₂SO₄. Cool well & mix w. c. sol. of 4 drops B in 5 cc. conc. H₂SO₄. As soon as mixt. becomes brown, pour drop by drop into 15 cc. ice water & crushed ice. Filter. Wash ppt. w. 5 cc. aq. Recryst. twice, first fr. 5 cc. & then fr. 3 cc. alc., filtering h. if sol. are not clear. Dry on porous tile 16 min. at 60°. The product, tetranitroethylaniline, is obtained in YT3 lust. scales, m.p. 95.5–6.5° u.c. (96–7° c.).</p>
I271	206c.	(80)	Pyridazine, ⁵ CH:CH.CH:CH.N:N ² . — M.p. –8°. Misc. aq., e.s. alc., eth. Sp. gr. 1.1108 (18/16). — B.Pk,‡ lemon-yel. ndl. turning green at 160°, & black, melting at 170°.
I272	206–7	121	m-Methylamino-toluene, MeNH.C ₆ H ₄ .Me. — Sec. amine.*
I273	206–7	135	2-Methylamino-1,3-xylene, MeNH.C ₆ H ₃ .Me ₂ . — Odor camphor-like. — Sec. amine.*
I274	206–7	140	2-Aminotropane, Isotropylamine, NH ₂ .C ₆ H ₄ .N. — M.p. +8.5°. — B.Pk,‡ lust. pr., m.p. 236–7° d.
I275	205–8c.		Diethyldimethylenediamine, (CH ₃) ₂ N.Et ₂ . — Odor unpleasant. S. c. aq.; sol. separates in layers on warming. C. conc. HCl dec. to ethylamine & trioxymethylene.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid
I276	207-8		Triethyltrimethylenetriamine, $(\text{CH}_2:\text{N}=\text{Et})_3$. — Sp. gr. 0.892 (18.7°). W. CS, gives diethylformocarbothialdine readily, cryst. fr. abs. alc., m.p. 75°.
I277	207-8	121	Methyl-o-toluidine, $\text{NHMe.C}_6\text{H}_4.\text{Me}$. — I. aq. Sp. gr. 0.973 (15°). Sec. amine.*
I278	205-8	147	Allylbenzylamine, $\text{C}_6\text{H}_5.\text{NH.CH}_2.\text{Ph}$. — Odor basic. Sec. amine.*
I279	207.2-8.2c.; 206°c.; 209-10°		† 1-Menthylamine, $\text{C}_{10}\text{H}_{19}.\text{NH}_3$. — Odor, strong, characteristic, coniine-like; sl. suggestive of menthol when h. V.d.s. c. aq. Taste of c. sat. aq. sol. burning & bitter. Sp. gr. 0.860 (20°). $n_D = 1.46058$. $[\alpha]_D = -38.07^\circ$. — Strongly alk. to litmus. A drop exposed to moist air on watch glass soon becomes covered w. white incrustation wh. effervesces w. conc. HCl! — Conc. aq. sol. B.HCl + 1 mol. NaNO ₂ sol. boiled w. ac. ac. gives N gas & l-menthol! — ② Heat to boiling 3 drops B + 5 drops ac. anhydride. Boil w. 7 cc. aq. & just enough alc. to dis. White scales separate on cooling. Filter. Wash w. 2 cc. aq. Cryst. fr. 75% alc. Dry 15 min. at 100°. The product, acetylmenthylamine, melts at 142.2-3.2° u.c. (144.4-5.4° c.).
I280	208d.		Diethylenetriamine, $\text{NH}_2.\text{CH}_2.\text{CH}_2.\text{NH}.\text{CH}_2.\text{CH}_2.\text{NH}_3$. — Misc. aq., alc.; alm. i. eth. Strong. alk.
I281	208	121	Methyl-p-toluidine, $\text{NHMe.C}_6\text{H}_4.\text{NH}_3$. — Sec. amine.* — B.HCl, m.p. 119.5° (fr. eth. sol. + HCl). — Nitroso deriv., ¶ pr. fr. eth.-alc., m.p. 53°.
I282	208-9	133	Allylaniline, $\text{Ph.NH}(\text{C}_6\text{H}_5)$. — Sec. amine.* Sp. gr. 0.982 (25°).
I283	208-9	149	tert.-Butyl-aniline, Ph.NH.CMe_3 . — Sec. amine.*
I284	209-10		1-Ethylglyoxyline, $[\text{NEt}.\text{CH}:\text{CH.N}: \text{CH}]$. — Misc. aq. Sp. gr. 0.999.
I285	209.5	135	Dimethyl-p-toluidine, $\text{Me}_2\text{N.C}_6\text{H}_4.\text{Me}$. — Tert. amine.* Sp. gr. 0.9287 (20/4).
I286	210	135	2-Phenylpropylamine, $\text{Ph.CHMe.CH}_2.\text{NH}_3$. — Odor fishy. Oil v.d.s. aq.; misc. alc., eth. Prim. amine.* — B.Pk, † m.p. 182°. — B ₂ H ₂ PtCl ₆ , § yel. lft., d. abt. 140°.
I287	210	155	Campholamine, $\text{C}_9\text{H}_{17}.\text{CH}_2.\text{NH}_3$. — Odor ammon. & garlic-like. Somewhat s. aq. "Strong base." Prim. amine.* — B.HNO ₃ , silvery lft., m.p. abt. 210° d.
I288	210c.	163	Diethyl-o-toluidine, $\text{NEt}_2.\text{C}_6\text{H}_4.\text{Me}$. — Tert. amine.* Sol. of I in KI (T. 2.3) added to sol. of B in dil. acid gives steel blue ppt. of B.HI.I, m.p. 100°.
I289	211c.		3-Aminotropane, Tropylamine, $\text{C}_6\text{H}_{14}.\text{NH}_3$. — Prim. amine.* — B.Pk, † 4-sided lft. fr. h. aq., m.p. 235° d. — B ₂ H ₂ PtCl ₆ , § red pr., m.p. 257° d.
I290	211	149	Propylbenzylamine, $\text{Pr.NH.CH}_2.\text{Ph}$. — Sec. amine.*
I291	211-2c.	163	Diethylbenzylamine, $\text{Et}_2\text{N.CH}_2.\text{Ph}$. — Tert. amine.*
—	212	121	1,3-Dimethyl-2-aminobenzene. — Cf. No. 2.1307.
I293	212-3	133	β -Tetrahydroquinoline, $\text{C}_9\text{H}_{11}.\text{N}$. — I. aq.; v.s. alc., eth. Oxidation w. dil. KMnO ₄ gives quinoline, No. 2.1356.
I294	212-3	149	Methylisopropylaniline, $\text{Me}(\text{Me},\text{CH})\text{N.Ph}$. — Tert. amine.* — B ₂ H ₂ PtCl ₆ , § m.p. 196-7°.
I295	213c.	140	Ψ -3-Aminotropane, Ψ -Tropylamine, $\text{C}_6\text{H}_{14}.\text{N.NH}_3$. — Prim. amine.* — B.Pk, † cryst. fr. h. aq., m.p. abt. 236-8° d. — B ₂ H ₂ PtCl ₆ , 2H ₂ O, § m.p. 257° d., after losing aq. at 105°.
I296	214-5	121	m-Aminoethylbenzene, $\text{NH}_2.\text{C}_6\text{H}_4.\text{Et}$. — S. h. aq. Sp. gr. 0.990 (0°). — Prim. amine.* — Acetyl deriv., ¶ m.p. 24-5°; b.p. 312-3°.

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(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I297	214	135	1-Methyl-2-ethylaminobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{Me}$. — Still liquid at -15° . Sp. gr. 0.953 (15.5 $^{\circ}$). Sec. amine.*
I298	214c.	149	1'-Aminoisobutylbenzene, $\text{Ph}.\text{CH}(\text{NH}_2).\text{CHMe}_2$. — Sp. gr. 0.920 (20/0). Prim. amine.* — $\text{B}.\text{HCl}$, m.p. 275-7 $^{\circ}$ d. — Oxalate, m.p. 120.5-2 $^{\circ}$ d.
I299	215-6c.	121	o-Ethylaminobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{Et}$. — Alm. i. c. aq. Sp. gr. 0.983 (22 $^{\circ}$). Prim. amine.* — Acetyl deriv., ¶ m.p. 111-2 $^{\circ}$; b.p. 304-5 $^{\circ}$.
I300	215	135	Dimethyl-m-toluidine, $\text{NMe}_2\text{C}_6\text{H}_4\text{Me}$. — Tert. amine.* — Convert into nitroso deriv., $\text{NMe}_2\text{C}_6\text{H}_3(\text{NO})\text{Me}$, green lft., m.p. 92 $^{\circ}$. (Cf. Ber., 12, 1797.)
I301	215	135	o-Isopropylaminobenzene, o-Cumidine, $\text{NH}_2\text{C}_6\text{H}_4\text{CHMe}_2$. — Prim. amine.* — Acetyl deriv., ¶ m.p. 72 $^{\circ}$. — Urea deriv. (cf. Ber., 21, 1162), m.p. 133-4 $^{\circ}$.
I302	abt. 215c.	135	Isopropylaniline, $\text{Ph}.\text{NH}.\text{CHMe}_2$. — Sec. amine.* — Acetyl deriv., ¶ tbl. fr. lgr., m.p. 39 $^{\circ}$.
I303	215	135	β -m-Tolylethylamine, $\text{Me}.\text{C}_6\text{H}_4.\text{CH}_2.\text{CH}_2.\text{NH}_2$. — Prim. amine.* Absorbs CO_2 . — $\text{B}.\text{HCl}$, m.p. 159 $^{\circ}$. — $\text{B}.\text{Pk}$, ¶ lft. fr. alc., m.p. 173 $^{\circ}$. — Urea deriv. (cf. Ber., 33, 1080), m.p. 84 $^{\circ}$.
I304	abt. 215	147	Methylallylaniline, $\text{Ph}.\text{NMe}_2\text{C}_6\text{H}_4$. — Odor charac. Tert. amine.* — Picrate, ¶ m.p. 91-2 $^{\circ}$.
I305	215		Methylpelletierine, $\text{C}_8\text{H}_{17}\text{NO}$. — [Fr. pomegranate bark.] — S. aq.; e.s. alk., eth., chlf.
I306	216		Triethylenetriamine, $[\text{C}_2\text{H}_5\text{NH}.\text{C}_2\text{H}_5\text{NH}.\text{C}_2\text{H}_5\text{NH}]^3$. — $\text{B}.\text{PtCl}_6$, ¶ yel. ndl. mod. s. aq.
I307	216c.; 212-3c.	121	v-m-Xyldine, 1,3-Dimethyl-2-aminobenzene, 2-Amino-1,3-xylene, $\text{NH}_2\text{C}_6\text{H}_3\text{Me}_2$. — Sp. gr. 0.980 (15 $^{\circ}$). Prim. amine.* — $\text{B}.\text{H}_2\text{SO}_4.2\frac{1}{2}\text{H}_2\text{O}$, is much more s. in aq. than sulphate of xyldine No. 2.1308. (100 pt. aq. at 18 $^{\circ}$ dis. over 60 pt. of v-m-sulphate.) — Acetyl deriv., m.p. 176 $^{\circ}$. (Said to be less quickly formed by acetylation than corresponding deriv. of No. 2.1308, and to be not saponifiable by boiling 25% H_2SO_4 .) — Prepare formyl deriv. by boiling 1 g. B 30 min. w. 1.2 g. 50% formic ac. Fine ndl. separate on cooling. M.p., after recryst. fr. alc., 164.5 $^{\circ}$.
I308	216.5c.; 212	121	† (a)-m-Xyldine, 1,3-Dimethyl-4-aminobenzene, 4-Amino-1,3-xylene, $\text{NH}_2\text{C}_6\text{H}_3\text{Me}_2$. — V.d.s. aq. Sp. gr. 0.918 (15 $^{\circ}$). Prim. amine.* — (¶) T. 2.2-b w. conc. HNO_3 , gives intense VR color. — (¶) (a) Heat mixt. of 6 drops B w. 1 cc. acetic anhydride to boiling. Boil product w. 6 cc. aq. Cool sol. well & shake v. persistently. Filter. Wash ppt. w. 2 cc. c. aq. Recryst. fr. 4 cc. h. aq. The product, m-acetoxylide, is obtained in small white ndl., m.p. 127-7.5 $^{\circ}$ u.c. (128.7-9.2 $^{\circ}$ c.). — (b) The picramide is obtained by procedure of T. 2.22 in YO-O pr. , m.p. 158.5 $^{\circ}$.
I309	216-6.5c.	121	1-Ethyl-4-aminobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{Et}$. — M.p. —5 $^{\circ}$. Sp. gr. 0.975 (22 $^{\circ}$). Prim. amine.* — $\text{B}.\text{H}_2\text{SO}_4$, large lft. d.s. c. aq., more s. dil. H_2SO_4 , "Charac. salt."
I310	216c.	149	† Diethylaniline, $\text{PhN}(\text{Et})_2$. — Odor aniline-like, unpleasant. Alm. i. aq. Sp. gr. 0.9351 (20/4). Tert. amine.* — (¶) Heat together on oil-bath for 3 min. at 200 $^{\circ}$ a few cg. B & as much crushed ZnCl_2 . Fusion is blue or violet. On quickly cooling color changes to pink on edges. Dis. in 3 cc. c. alc. Sol. is bright pink, fading rapidly. — (¶) Dis. 3 drops B in 4 cc. gl. ac. ac. Add 5 drops conc. HNO_3 , drop by drop. Warm over small flame until sol. is light yel. Reaction will then usually proceed without further heating. When color is light brown, cool quickly & dilute w. 8 cc. aq. Shake vigorously. Filter. Wash w. 3 cc. c. aq. Recryst. fr. 5 cc. 50% alc. Filter h. if not clear. Dry ppt. which separates on cooling

No.	Boiling-point (C.).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
			<i>15 min. at 60° on porous tile. The product, p-nitrodiethylaniline, is obtained in small lust. Y-OY ndl., m.p. 77.5–8.5° u.c. (77.9–78.9° c.).</i>
I311	abt. 216	163	Ethylpropylaniline, Ph.NEt,Pr. — Sp. gr. 0.934 (15°). — Tert. amine.* — $\text{B}_2\text{H}_2\text{PtCl}_4$, § yel. cryst. powd., m.p. 199°, 186°. — Acetyl deriv., ¶ m.p. 129°.
I312	215–7c.		2,5-Dimethyl-3,6-diethylpyrazine, (1,4) Diethylketine, $\text{C}_{10}\text{H}_{14}\text{N}_2$. — Oil of faint narcotic odor. Forms cryst. hydrate, $\text{B}_2\text{xH}_2\text{O}$, m.p. 42.5°.
I313	217c.		1-Methyl-2,3-dihydroindole, $\text{C}^{\text{N}}\text{Me}.\text{CH}_2.\text{CH}_2.\text{C}_6\text{H}_4$? — D.s. aq.; e.s. alc., eth.; mod. vol. w. st. — $\text{B}.\text{Pk}$, † yel. tbl. fr. bz., m.p. 155°.
I314	217	135	Ethyl-p-toluidine, Ph.NHET. — Sp. gr. 0.939 (15°). Sec. amine.* Treatment of 1 pt. B disd. in 20 pt. conc. H_2SO_4 w. conc. HNO_3 gives nitro deriv., m.p. 48°.
I315	218c.; 217c.	121	† p-Xyldine, 1,4-Dimethyl-2-aminobenzene, $\text{NH}_2.\text{C}_6\text{H}_4.\text{Me}_2$. — M.p. +15.5°. V.d.s. aq. Prim. amine.* — $\text{B}.\text{H}_2\text{O}$, gives intense red-brown (ORS1) color. — $\text{B}.$ Heat to boiling 3 drops B + 1 cc. acetic anhydride. Boil product w. 10 cc. aq. Allow to cool slowly. Filter. Recryst. fr. 10 cc. aq. Dry 15 min. at 100°. The product, aceto-p-xyldine, is obtained in colorless ndl., m.p. 137.4–8°. u.c. (139.4–9.8° c.). Yield, good. — $\text{B}.\text{HCl}$, m.p. 228°. — Formyl deriv., m.p. 111–2°; 116–7°. — Benzenesulphonyl deriv., m.p. 138–9°.
I316	218–9	135	1'-Amino-1,3,4-trimethylbenzene, $\text{NH}_2.\text{CH}_2.\text{C}_6\text{H}_3.\text{Me}_3$. — D.s. aq.; e.s. alc., eth. Prim. amine.* — $\text{B}.\text{HCl}$, m.p. 210°. — $\text{B}.\text{Pk}$, † yel. lft., m.p. 223° d. — Urea deriv. (Ber., 22, 122), m.p. 184.5°.
I317	218	136	2-Aminodimethylaniline, $\text{NH}_2.\text{C}_6\text{H}_4.\text{NMe}_2$. — Odor menthol-like. Tert. amine.* — $\text{B}.\text{FeCl}_3$ colors aq. sol. brown, violet & finally deep blue, changed to red by boiling. — $\text{B}.\text{Pk}$, † yel. tbl. fr. alc., m.p. 138–40° d.
I318	218–20	153	Dihydrocarvylamine, $\text{C}_{10}\text{H}_{17}.\text{NH}_2$. — Sp. gr. 0.8875 (20°). Prim. amine.* $n_D = 1.48168$. Absorbs CO_2 .
I319	abt. 220		Arecoline, Methyl N-Methyl-4-tetrahydronicotinate, $\text{MeCO}_2-\text{C}_6\text{H}_4\text{N}$. — Misc. aq.! S. alc., eth. Tert. amine.* — $\text{B}.\text{HCl}$, deliq. ndl., m.p. 157–8° c. — $\text{B}.\text{MeI}$, ‡ by mixing components w. v. little MeOH, cooling, washing w. MeOH, & recryst. fr. same solvent; colorless pr., m.p. 173–4° c. (frothing). — $\text{B}_2\text{H}_2\text{PtCl}_4.5\text{H}_2\text{O}$, § or.-red cryst., m.p. 176° (frothing).
I320	218–21	163	Isobutylbenzylamine, $\text{Me}_2\text{CH}.\text{CH}_2.\text{NH}.\text{CH}_2.\text{Ph}$. — Sec. amine.* — $\text{B}.\text{HCl}$, ndl., m.p. 175°.
I321	220–1c.	121	s-m-Xyldine, 1-3-Dimethyl-5-aminobenzene, $\text{NH}_2.\text{C}_6\text{H}_3.\text{Me}_2$. — Sp. gr. 0.993 (0°). Prim. amine.* — Acetyl deriv., ¶ m.p. 140.5°, ndl. fr. alc. — Formyl deriv., pr. fr. dil. alc., m.p. 76.5°. — Nitroso deriv., ** (Ber., 18, 2679), m.p. 54°.
I322	219; 222–4	169	o-Propylaminobenzene, $\text{NH}_2.\text{C}_6\text{H}_4.\text{Pr}$. — Faint aniline-like odor. Prim. amine.* — $\text{B}.\text{HCl}$, m.p. 173°. $\text{B}.\text{Pk}$, m.p. 151°. — Acetyl deriv., ¶ m.p. 104–5°.
I323	220–1c.	135	1'-Amino-1,3,5-trimethylbenzene, $\text{NH}_2.\text{CH}_2.\text{C}_6\text{H}_3.\text{Me}_3$. — Sp. gr. 0.9500 (20/4). Prim. amine.* — $\text{B}.\text{HCl}$, m.p. 245° (brown). — $\text{B}.\text{Pk}$, † m.p. 225° d. — $\text{B}_2\text{H}_2\text{PtCl}_4$, § 6-sided yel. lft., d.s. c. aq., m.p. 204°.
I324	220	149	p-Methylisopropylaminobenzene, $\text{Me}.\text{C}_6\text{H}_4.\text{NH}.\text{CHMe}_2$. — Sp. gr. 0.923 (20°). Sec. amine.* — $\text{B}.\text{HCl}$, cryst. fr. alc., m.p. 170–1°. — Nitroso deriv., ** m.p. 58–9°. — $\text{B}_2\text{H}_2\text{C}_2\text{O}_4$, m.p. 129–30°.

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 1 = T. 2.1. ** = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS. — Colorless and Liquid.
I325	220.5 (th.i.)	149	1'-Aminobutylbenzene, Ph.CH(NH ₂).-(CH ₂) ₃ .Me. — Sp. gr. 0.937 (20/0). Prim. amine.* — B ₂ H ₅ PtCl ₆ , golden scales, m.p. 184–4.5° d., d.s.c. aq.
I326	221		Aminohydrindene, $\text{C}_6\text{H}_5\text{CH}(\text{NH}_2)\text{CH}_2\text{CH}_2$. — Absorbs CO ₂ . Prim. amine.* — B.HCl, cryst. v.s. aq., m.p. 208°.
I327	221	135	1,3-Dimethyl-4-methylaminobenzene, Me ₂ C ₆ H ₄ NHMe. — Sec. amine.* — Nitrosamine, oily. — Acetyl deriv., ¶ cryst. fr. lgr., m.p. 65°.
I328	221.5 (th.i.)	135	1-(γ)-Phenylpropylamine, Ph.CH ₂ .CH ₂ .CH ₂ .NH ₂ . — Absorbs CO ₂ . Somewhat s. aq. Misc. alc., eth. Sp. gr. 0.951 (15°). Prim. amine.* — B.HCl, lft. fr. alc. + eth., m.p. 218°. — B.Pk, ¶ ndl. fr. aq., m.p. 152–3°.
I329	222 (th.i.)	135	† Propylaniline, Pr.NH.Ph. — Sp. gr. 0.949 (18°). Sec. amine.* Acetyl deriv., ¶ tbl. fr. alc., m.p. abt. 49°. — B.HCl, m.p. 125°.
I330	220–5	122	ab-Methylphenylhydrazine, MeNH.NHPh. — Oil becoming yel. in air. — (¶) Reduces c. ammon. AgNO ₃ or Fehling's sol. — Sulphate e.s. aq., m.p. 180°.
I331	222	163	1,2,4-Trimethyl-5-dimethylaminobenzene, Dimethylpseudocumidine, Me ₂ C ₆ H ₄ NMe ₂ . — Tert. amine.*
I332	224c.	123	† o-Anisidine, MeO.C ₆ H ₄ .NH ₂ . — M.p. + 5.2° Odor mild aromatic. D.s. aq. Sp. gr. 1.098 (15/15). Prim. amine.* — (¶) Dis. v. small drop B in 5 cc. mixt. of 2 vol. conc. H ₂ SO ₄ + 1 vol. aq., & add 1 drop conc. HNO ₃ . Sol. becomes opaque brownish black at once. — (¶) Heat to boiling 3 drops B + 6 drops acetic anhydride. Dis. in 3 cc. boiling aq. Cool w. vigorous & persistent shaking. Recryst. ppt. fr. 3 cc. h. aq. The product, aceto-o-aniside, separates as oil fr. h. sol., changing to ndl. on further cooling, m.p. 84.5–85.0° u.c. (85.0–5.5° c.).
I333	224 (th.i.)	121	(v)-o-Xylidine, 1,2-Dimethyl-3-aminobenzene, Me ₂ C ₆ H ₃ .NH ₂ . — Sp. gr. 0.991 (15°). Prim. amine.* — B ₂ H ₅ SO ₄ , cryst. s. in 71 pt. c. aq. — Acetyl deriv., ¶ m.p. 131°, 134°.
—	225–6	130	Quinoxaline. — Cf. No. 2.536. (M.p. 27°.)
I334	225	135	p-Isopropylaminobenzene, Cumidine, Me ₂ CH.C ₆ H ₄ .NH ₂ . — Still liq. at –20°. Sp. gr. 0.953. Prim. amine.* — Acetyl deriv., ¶ lft. fr. h. aq., m.p. 102–2.5°. — Urea deriv. (Ber., 21, 1159), ndl. fr. aq., m.p. 152°.
I335	224–6	135	p-Propylaminobenzene, Pr.C ₆ H ₄ .NH ₂ . — Prim. amine.* — Acetyl deriv., ¶ lft. fr. dil. alc., m.p. 87°.
I336	227.5 (th.i.)	122	as-Methylphenylhydrazine, Me ₂ PhN.NH ₂ . — Boils w. sl. decon. & formation of NH ₃ . — (¶) Reduces Tollen's reagent (T. 2.30) r. slowly. — (¶) Prepare the corresponding hydrazone of benzaldehyde (Ber., 29, 814), yel. ndl. fr. lgr. softening at 102°, m.p. 104.5°.
I337	227	135	1,4-Dimethyl-2-methylaminobenzene, Me ₂ C ₆ H ₃ .NHMe. — Sp. gr. 0.962. Sec. amine.* — Nitrosamine, oily.
I338	226–8	149	1'-Aminomethyl-4-isopropylbenzene, Cumylamine, NH ₂ .CH ₂ -C ₆ H ₄ .CH.Me ₂ . — Alm. i. aq.; v.s. alc., eth. Absorbs CO ₂ . Prim. amine.* Acetyl deriv., ¶ pearly lft. fr. lgr., d.s. h. aq., m.p. 65°. — B.HNO ₃ , lft. e.s. aq., m.p. 155–7°.
I339	227–8	149	1,2-Dimethyl-3-ethylaminobenzene, NHEt.C ₆ H ₄ .Me ₂ . — Still liq. at –18°. Sec. amine.* — Nitroso deriv., ** green lft., m.p. 123–4°. — Acetyl deriv., ¶ liq. at –18°.
I340	227–8	163	o-Methyldiethylaminobenzene, Me.C ₆ H ₄ .NET ₂ . — Tert. amine.*
I341	abt. 228 (th.i.)		2-Methyl-1,3-dihydroindole, Me.C ₆ H ₄ N. — Pungent oil. "Strong base." Sp. gr. 1.0231 (20/4). — Picrate, ¶ m.p. 150–1°. — B.HCl + NaNO ₂ gives yel. nitroso deriv., pr. fr. lgr., m.p. 54–5°.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS. — Colorless and Liquid.
I342	228; 229	137	o-Phenetidine, EtO.C ₆ H ₄ .NH ₂ . — Liq. at -21 $^{\circ}$. Prim. amine.*
I343	227; 229-30	135	Mesidine, 1,3,5-Trimethyl-2-aminobenzene, Me ₃ C ₆ H ₃ .NH ₂ . — Sp. gr. 0.963. Prim. amine.*
I345	229	163	p-Methyldiethylaminobenzene, Me.C ₆ H ₄ .NEt ₂ . — Sp. gr. 0.924 (15 $^{\circ}$). Tert. amine.* — B.HCl, m.p. 157 $^{\circ}$.
I346	230	149	o-Methylpropylbenzene, Me.C ₆ H ₄ .NHPr. — Sec. amine.*
I347	230	149	p-Cymidine, 1-Methyl-4-isopropyl-3-aminobenzene, Me(Me-.CH)C ₆ H ₃ .NH ₂ . — Odor unpleasant. Alm. i. aq.; e.s. alc., eth. Prim. amine.* — Acetyl deriv., ¶ m.p. 112 $^{\circ}$.
I348	228-32	163	Butylbenzylamine, Bu.NH.C ₆ H ₄ .Ph. — Sec. amine.* Nitroso deriv.** oily. — B.HCl, m.p. 241 $^{\circ}$. — Chloroplatinate, § red-brown cryst. powd., m.p. 90 $^{\circ}$.
I349	231	171	2-Aminoundecane, Me.CH(NH ₂).(CH ₂) ₉ .Me. — Prim. amine.* — B.HCl, flat ndl. fr. lgr., m.p. 83-4 $^{\circ}$.
I349-I	231-2c.; (242)	149	Isobutylaniline, Me.CH.CH ₂ .NH.Ph. — Sp. gr. 0.940 (18/4 $^{\circ}$). — Sec. amine.* — S. in 12,500 pt. aq. at 15 $^{\circ}$; v.s. eth., bz.
I350	232 (th.i.)	133	3-Methyl-1,2-dihydroindole, Hydroksatole, Me.C ₆ H ₅ N. — D.s. aq.; e.s. alc., eth. — ② Alc. sol. colors pine splinter wet w. HCl or.-yel. — B.Pk, † m.p. 149-50 $^{\circ}$.
I351	232-3	133	Tetrahydroisoquinoline, C ₉ H ₁₁ N. — Liq. at -15 $^{\circ}$. Absorbs CO ₂ . Reduces Ag sol. — B.Pk, † glassy yel. ndl. fr. alc., m.p. 195 $^{\circ}$. — B.HCl, tbl., m.p. 195-7 $^{\circ}$. — B ₂ H ₂ PtCl ₆ , § red-yel. tbl., m.p. 231-2 $^{\circ}$. — Nitroso deriv.** flat ndl. fr. bz., m.p. 53 $^{\circ}$.
I352	232-4	135	2,3,4,5-Tetramethylpyridine, Parvoline, Me,C ₆ HN. — [In coal tar.] — B.Pk, † ndl. m.p. 170-2 $^{\circ}$. — B.HAuCl ₄ , yel. ndl., m.p. 216-8 $^{\circ}$. — B.HCl.2HgCl ₂ , ndl., e.s. aq., m.p. 156 $^{\circ}$.
I353	233c.	143	N-Methyl- α -pipecolylalkine, Hydrotropine, C ₉ H ₁₁ ON. — Odor faintly ammon. E.s. aq. — B.HAuCl ₄ , cryst. d.s. c. aq., m.p. 169-70 $^{\circ}$; 176 $^{\circ}$.
I354	235	149	p-Methylpropylaminobenzene, Me.C ₆ H ₄ .NHPr. — Odor caraway-like. Prim. amine.* — B.HCl, ndl., m.p. 150-1 $^{\circ}$. — B ₂ H ₂ O, m.p. 172-3 $^{\circ}$ d.
I354-I	231-2c.; 242	149	Isobutylaniline, Ph.NH.CH ₂ .CHMe ₂ . — Sp. gr. 0.940 (18/4 $^{\circ}$). S. in 12,500 pt. aq. at 15 $^{\circ}$; v.s. eth., bz. — ② Sec. amine.*
I355	234-6	163	Methylisobutylaniline, Me.(Me.CH.CH ₂).NPh. — Tert. amine.*
I356	236.2c.	129 k.10 ^a = 0.8	† Quinoline, C ₉ H ₇ N. — [In bone oil.] — <i>Odor strong, basic, unpleasant, charac!</i> V.d.s. aq.; misc. alc., eth., chlf. Not solidified by ice & salt. Tert. amine.* — ② Mix 4 drops B. w. 6 drops Mel. Set aside for 5 min. Crush lumps. Cryst. fr. 3 cc. abs. alc. Dry at room temp. on porous tile. B.Mel is obtained in fine Y-OYndl., m.p. 134.7-5.2 $^{\circ}$ u.c. (136.7-7.2 $^{\circ}$ c.). — B.Pk, † fine yel. ndl. fr. bz., m.p. 203 $^{\circ}$.
I357	235-7	133	Styrylamine, Ph.CH:CH.CH ₂ .NH ₂ . — Absorbs CO ₂ . Prim. amine.* — B.Pk, cryst. ppt., m.p. 173 $^{\circ}$. — B.HCl, m.p. 210 $^{\circ}$. — B ₂ H ₂ PtCl ₆ , § egg-yel. ppt., m.p. 205-7 $^{\circ}$.
I358	236	149	Butylaniline, Me.(CH ₂) ₂ .CH ₂ .NH.Ph. — Sec. amine.*
I359	236 (th.i.)	150	Isopropylphenylhydrazine, (Me.CH) ₂ PhN.NH ₂ . — Sp. gr. 0.9588 (15 $^{\circ}$). Prob. reduces Tollen's reagt. slowly. — B.HCl, cryst. fr. bz., m.p. 135 $^{\circ}$.
I360	237 (th.i.)	136	as.-Ethylphenylhydrazine, Et.Ph.N.NH ₂ . — Sp. gr. 1.018 (15 $^{\circ}$). Reduces Fehling's sol. but only on heating. — B.HCl, lft. e.s. bz., eth.!, m.p. 137 $^{\circ}$.
I361	237 (th.i.)	149	1,4-Dimethyl-3-ethyl-2-aminobenzene, Me ₂ Et.C ₆ H ₃ .NH ₂ . — Sp. gr. 0.963 (15 $^{\circ}$). — Prim. amine.* — Acetyl deriv., ¶ m.p. 142 $^{\circ}$. — Formyl deriv., m.p. 104-5 $^{\circ}$.

Explanation of typographical signs used in this Division: * = T. 2.35. † = T. 2.23. § = T. 2.14. || = T. 2.13.
¶ = T. 2.1. ** = T. 2.36. ¶¶ = T. 2.37.

(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I362	236; 240	135	1,2,4-Trimethyl-3-aminobenzene, Me ₃ C ₆ H ₃ NH ₂ . — Prim. amine.* — Acetyl deriv., ¶ m.p. 186°.
I363	238	163	1-Methyl-3,5-diethyl-4-aminobenzene, Me,Et ₂ C ₆ H ₃ NH ₂ . — Prim. amine.* — Acetyl deriv., ¶ m.p. 167°.
I364	240	177	Isoamylbenzylamine, C ₁₁ H ₂₁ NH ₂ CH ₂ Ph. — Sec. amine.* — B.HCl, d.s. aq., m.p. 253°. — B ₂ H ₂ PtCl ₆ , § cryst. or. powd., m.p. 203°.
I365	240.8c.	129	† Isoquinoline, C ₉ H ₇ N. — Solidified by ice & salt. M.p. 24.6°. V.d.s. aq. Sp. gr. 1.099 (21.4/4). Tert. amine.* — ① Warm 4 drops B w. 6-8 drops MeI in small t.t. Set aside 10 min., occasionally shaking. Recryst. ppt. fr. 3 cc. h. abs. alc. Dry at room temp. on porous tile. B.MeI, ‡ the product, is obtained in pale yel. ndl., m.p. 157.3-9.5° u.c. (159.9-61.9° c.). — B.Pk, ‡ yel. ndl., d.s. aq., alc., m.p. 223°.
I366	240-2; (235-40)	170	Menthylhydrazine, C ₁₀ H ₁₁ NH.NH ₂ . — Reduces ammon. AgNO ₃ sol. E.s. aq., alc., eth. [α] _D = 46.05 (for sol. of 9.62 g. B in 100 cc. aq.).
I367	240-4	122	m-Tolylhydrazine, Me,C ₆ H ₄ NH.NH ₂ . — Opt. i.
I368	241-2	149	Carvacrylamine, 1-Methyl-4-isopropyl-2-aminobenzene, Me,-(Me ₂ CH) ₂ C ₆ H ₃ NH ₂ . — Sp. gr. 0.944 (20°). Prim. amine.* — B.HCl, lft., m.p. 207°. — Acetyl deriv., ¶ tbl. fr. dil. alc., m.p. 71°.
I369	243.5 (th.i.) sl. d.	108 k.10 ⁹ = 1.6	† Phenylhydrazine, Ph.NH.NH ₂ . — Soon becomes yel. & brown on exposure. Disd. in 2 vol. anhydrous eth. & cooled to -10°, white monolin. tbl., m.p. +17.5°, separate. A hydrate, B ₄ H ₈ O, melts at 24.1°. D.s. aq. Misc. w. alc., eth., chlf., bz. Sp. gr. 1.097 (22.7/4). — ② Reduces Tollen's reagt. (T. 2.30) instantly! Reduces Fehling's sol. in cold w. efferv. & escape of N ¹⁺ — ③ (a) Convert 1 drop into benzaldehyde-phenylhydrazone by procedure of T. 1.113. — (b) Prepare the charac. blood-red 2,4-dinitrobenzaldehydophenylhydrazone, m.p. 216°, as described by Sachs & Kempf (Ber., 1902, 1230).
I370	243	163	1-Methyl-3-tert.-butyl-6-aminobenzene, Me,(CMe ₃).C ₆ H ₄ NH ₂ . — Prim. amine.* — Acetyl deriv., ¶ lft. fr. dil. alc., m.p. 162°.
I371	243-4	163	1-Methyl-3-isobutyl-2-aminobenzene, Me,(NH ₂).C ₆ H ₃ CH ₂ Me. — Prim. amine.* — Acetyl deriv., ¶ silky ndl., m.p. 141-2°.
I372	243-6		Phenylethylenediamine, Ph.CH(NH ₂).CH ₂ (NH ₂). — Not solidified by cold. V.s. aq.! E.s. alc. — Picrate, ‡ yel. cryst. granules i. bz., m.p. 160°. — Diacetyl deriv., ¶ lft. fr. bz., m.p. 152°.
I373	245	164	as.-Isobutylphenylhydrazine, (C ₆ H ₅) ₂ PhN.NH ₂ . — Sp. gr. 0.9633 (15°). Prob. reduces Tollen's reagt.
I374	245	175	N-Benzylpiperidine, Ph.CH ₂ NC ₆ H ₅ . — Alm. i. aq. Tert. amine.* — B ₂ H ₂ PtCl ₆ , § d.s., m.p. 191-3°.
I375	245-8		1',3'-Diaminoxylene, (NH ₂ CH ₂) ₂ C ₆ H ₄ . — Misc. alc., eth. — B.Pk, ‡ yel. lft. d.w.m. 185-90°. — Diacetyl deriv., ¶ cryst. e.s. h. aq., m.p. 118-9°.
I376	246-7c.	143 k.10 ⁹ = 4	† Quinaldine, 2-Methylquinoline, Me,C ₆ H ₄ N. — V.d.s. c. aq. Tert. amine.* — ① Heat 6 drops B w. 0.2 g. phthalic anhydride in t.t. immersed in oil at 170° for 5 min. Dis. in 2 cc. warm H ₂ SO ₄ (sp. gr. 1.84). Cool. Add aq. cautiously until ppt. no longer increases. Filter. Wash the yel. to dark brown ppt. w. 3 cc. aq. Drain thoroughly. Dis. in 1 cc. gl. ac. ac. Ppt. again w. aq. Filter the light yel. ppt. Wash w. aq. Recryst. fr. h. alc. The product, quinophthalon or "quinoline yellow," C ₁₄ H ₁₁ N.CH ₂ O ₄ C ₆ H ₄ , is obtained in fine yel. (Y) ndl., m.p. 232.7-3.7° u.c. (239.4-40.4° c.). — B.Pk, ‡ yel. ndl., d.s. aq., alc., m.p. 191°. — B ₂ H ₂ PtCl ₆ , § or.-red pr. fr. h. aq., m.p. 226°.

No.	Boiling-point (C°).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I377	246.5	147	ac.-Tetrahydro- α -naphthylamine, $C_{10}H_{13}N$. — Thick oil. Odor ammon. S.c. aq.; e.s. alc. — Acetyl deriv., \parallel m.p. 148–9°.
I378	246–9		Methyl- α -phenylenediamine, $MeNH.C_6H_4.NH_2$. — Vol. w. st.
I379	247 (th.i.)	150	Propylphenylhydrazine, $Pr.PH.N.NH_2$. — Sp. gr. 0.9471 (15°). Prob. reduces Tollen's reagt. — $B.HCl$, m.p. 135°.
I380	247.3c.; 247.7–9.7c.	k.10 ⁴ = abt. 3	† 1-Nicotine, 1-Methyl-2- β -pyridylpyrrolidine, $Me.NC_6H_7.C_6H_4.N$. — [Toxic alkaloid fr. tobacco.] — Odor strong & unpleasant, recalling piperidine & tobacco, but quite different from that of either! Distillate in hydrogen atmosphere colorless, becoming distinctly yel. on contact w. air within 24 hr. Dist. w. v.s.t. deca. in air. Misc. aq., alc., eth. Vol. w. st. Sp. gr. 1.0097 (20/4). $[\alpha]D^{20} = -166.39^\circ$. Salts dextrorotatory. $nD^{20} = 1.5280$. — A few drops of sol. of 1 drop B in 5 cc. aq. held on tongue for few seconds & then ejected fr. mouth causes burning taste (No. 2 in T. 2.29) followed by disagreeable irritation that extends to the throat and persists for some minutes! (This test, if used, should be made only w. great caution, the fatal dose of pure B being probably as little as 2 or 3 drops.) — Characteristic color reactions are lacking.
			② Add to ethereal sol. containing 1 mol. B an equal vol. ethereal I sol., containing 2 atoms I. Employing 1% to 0.2% B sol., colored cryst. ppt. of v. charac. appearance (Roussin's crystals) are obtained within a few minutes or hours. Cf. Kippenberger, Z. anal. Chem., 42, 232–76, for full discussion of this reaction.
			③ Dis. 1 drop B in 5 cc. alc. Add 10 cc. c. sat. aq. sol. picric ac. Heat to dis. heavy ppt. Cool & shake. Wash cryst. w. 3 cc. 33% alc. Recryst. fr. 10 cc. h. 33% alc. by cooling & shaking. Wash w. 3 cc. dil. alc. Dry 15 min. on porous tile at 100°. The product, $B.Pk_2$, † is obtained in distinct Y/T1 ndl., which may be more than 1 cm. in length if sol. is allowed to cool slowly, m.p. 219.2–20.2° u.c. (224.8–5.8° c.) — $B.2C_6H_4O_4N_4$ (picrolonate), ndl., m.p. 213°. — $B.H_2PtCl_6$, § monolin. cryst. (fr. dil. sol.), m.p. 275° d. (darkening fr. 250°).
I381	245–50	205	Diisobutylaniline, $(C_6H_5)_2NPh$. — Tert. amine.*
I382	248c.	143	8-Methylquinoline, $Me.C_6H_4.C_6H_3N$. — Odor unpleasant, quinoline-like. Sp. gr. 1.0730 (20/4). Tert. amine.* — $B.Pk$, † yel. lft. d.s. alc., m.p. 200°.
I383	248	143	1-Methylisoquinoline, $^TC_6H_4.CH:CH.N:CMe^3$. — Sp. gr. 1.0768 (20/4). Tert. amine.* — $B.H_2PtCl_6.4H_2O$, § light red pr., v.d.s. aq., m.p. 200° (frothing).
I384	247–50c.	147	N-Methyltetrahydroquinoline, "Kairolin," $C_6H_{10}N.Me$. — [Sulphate is commercial.] Sp. gr. 1.022 (20/4). Tert. amine.* — ② The v. dil. acid sol. gives charac. yel.-red color w. $NaNO_2$. — Is colored intense blue by treatment w. $COCl_2 + AlCl_3$. — Picrate, † yel. ndl., m.p. 125°. — $B.C_6H_5O_4N_4$ (picrolonate), † pr. fr. alc., m.p. 192–5° w. efferv.
I385	251c.; (260c.)	133	† Tetrahydroquinoline, $C_6H_{11}N$. — Darkens on exposure. V.d.s. c. aq. Vol. w. st. Sp. gr. 1.063 (15/15). — ③ Dis. 3 drops B in 3 cc. dil. HCl (1 vol. conc. HCl + 4 vol. aq.). Add 1 cc. c. sat. aq. sol. of $NaNO_2$. Separate ppt. of brownish oil & shake it vigorously w. 3 cc. dil. HNO_3 (1 vol. conc. HNO_3 + 2 vol. aq.). Oil gradually solidifies without dissolving. Crush lumps. Filter. Wash w. 5 cc. c. aq. Recryst. twice, first fr. 3 cc., & then fr. 2 cc. h. alc. The product, nitronitrosotetrahydroquinoline, is obtained in O-OY ndl., m.p. 143.7–6.7° u.c. (145.9–7.9° c.).
I386	251–2 (th.i.)	133	Ethyldiethylolamine, $EtN(CH_2.CH_2.OH)_2$. — Odor faintly ammon. Fumes, absorbing CO_2 & aq. fr. air. Caustic. Sp. gr. 1.013 (20/4).

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 ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37.

No.	Boiling-point (C.).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
1387	251-2 (th.i.)	147	ac.-Tetrahydro- β -naphthylamine, $C_{10}H_{11}.NH_2$. — Odor intensely ammon. & piperidine-like. Absorbs CO_2 . Sp. gr. 1.0344 (15/15). Prim. amine.* — Acetyl deriv., pr. fr. bz., e.s. h. aq., m.p. 107.5°. — Phenylurea deriv. (fr. eth. sol. $B +$ phenyl isocyanate), m.p. 165.5°.
1388	252	143	3-Methylquinoline, $Me.C_8H_4N$. — M.p. + 10 to 14°. — Tert. amine.* — $B.Pk$, † yel. ndl., m.p. 187°. — $B.HAuCl_4$, ndl. d.s. c. aq., m.p. 145°.
1389	252c.	157	† 2,8-Dimethylquinoline, 8-Methylquinaldine, $Me_2.C_8H_4N$. — D.s. aq.; e.s. alc., eth. Tert. amine.* — ① Heat 6 drops B w. 0.2 g. phthalic anhydride 10 min. at 170°. Boil brown mass w. 5 cc. HCl (sp. gr. 1.12), breaking up lumps. Filter. Wash w. 5 cc. HCl. Cryst. fr. smallest possible quantity h. alc. (10 to 20 cc.). Dry on porous tile 15 min. at 100°. The product, 8-methylquinophthalone, $C_{11}H_9N.C_6O_2.C_6H_4$, forms fine yel. ndl., m.p. 271.8-2.6° u.c. (280.7-1.7° c.). $B.Pk$, † yel. pr. fr. chlf. & alc., m.p. 180°.
1390	251-4 (th.i.)	147	4-Methyl-1,2,3,4-tetrahydroquinoline, $Me.C_8H_{10}N$. — Oil of penetrating odor. Tert. amine.*
1391	250-5		Nicotimine, $C_{10}H_8N_2$. — [Fr. tobacco in small quantity.] Odor sharper & more unpleasant than for nicotine. Misc. aq. Sec. amine.* — Picrate, † pr., m.p. 163°.
1392	254.2-0.7c.	137	p-Phenetidine, $EtO.C_6H_4.NH_2$. — M.p. + 2.4°. Soon reddens & darkens on exposure. Prim. amine.* Sp. gr. 1.0613 (15°). — ② Conc. HNO_3 (T. 2.2-6) gives blue ($B-BT1$) color. — ③ Heat to boiling 2 drops B + 5 drops acetic anhydride. Dis. product in 7 cc. boiling aq. Cool. Filter. Wash ppt. w. 3 cc. c. aq. Recryst. fr. 3 cc. boiling aq. Dry 15 min. on tile at 100°. The product, phenacetine, is obtained in pearly white scales, softening fr. 133° u.c., m.p. 134° u.c. (135.9° c.).
1393	abt. 254c.	163	Isoamylaniline, $C_8H_{11}.NH.Ph$. — Sp. gr. 0.928 (15/4). Sec. amine.*
—	255	149	Isouridine. — Cf. No. 2.531. (M.p. 23-4°.)
1394	255	171	2-Isopropylquinoline, $Pr.C_8H_4N$. — Oil of quinoline-like odor. Tert. amine.* — $B.Pk$, † yel. lt. fr. alc., m.p. 150°; 155-7°.
1395	256	143	4-Methylisoquinoline, $Me.C_8H_4N$. — Liq. at -75°. Tert. amine.* — Picrate, † ndl., m.p. 194-5°. — $B_2H_2PtCl_6$, § (100°) brown-red cryst., m.p. 253.5°.
1396	abt. 256	147	8-Methyltetrahydroquinoline, $Me.C_8H_{10}N$. — R.d.s. aq.; e.s. alc. — $B.Pk$, † m.p. 212°. — Nitroso deriv., ** tbl. fr. eth., m.p. 51°.
1397	257	177	Methylisoamylaniline, $Me.(C_8H_{11}).NPh$. — Sp. gr. 0.906 (20°). Tert. amine.*
1398	256.6-8.6c.	157	2-Ethylquinoline, $Et.C_8H_4N$. — D.s. aq.; e.s. alc., eth. Tert. amine.* — $B.Pk$, † yel. ndl., d.s. aq., m.p. 146-7°; 148°. — $B_2H_2PtCl_6$, § (at 100°), tbl. d.s. aq., m.p. 190° d. — $B.HCl-HgCl_2$, ndl., m.p. 118°.
1399	257-9.5		Methyl-p-phenylenediamine, $NH_2.C_6H_4.NHMe$. — E.s. aq.! Dil. neutral sol. of salts colored intensely red by trace $FeCl_3$, becoming colorless on addition of HCl , & then blue on passage of H_2S .
1400	258	143	6-Methylquinoline, $Me.C_8H_4N$. — Odor unpleasant, quinoline-like. Alm. i. aq. Sp. gr. 1.0664 (20/4). Tert. amine.*
1401	258	143	8-Methylisoquinoline, $Me.C_8H_4N$. — Tert. amine.* — Picrate, † m.p. 204-5°. — $B_2H_2PtCl_6.2H_2O$, § light red ndl.
1402	260c.	143	7-Methylquinoline, $Me.C_8H_4N$. — Sp. gr. 1.0722 (20°). Tert. amine.* — Picrate, † yel. pr. v.d.s. h. alc., m.p. 237°.

No.	Boiling-point (C. $^{\circ}$).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I403	260 sl. d.		1-Phenylpyrazolidine, $[\text{NPh.NH}(\text{CH}_2)_3]$. — I. aq.; s. alc. Oxidizes in air. Sp. gr. 1.20 (15 $^{\circ}$). Sol. in conc. HCl is purple-red. — B.Pk, \ddagger yel. cryst. powd., m.p. 102 $^{\circ}$ d.
I404	260	269	Trihexylamine, $(\text{C}_6\text{H}_{11})_3\text{N}$. — Tert. amine.* Alm. i. aq.
—	262c.		Dimethyl-p-phenylenediamine. — Cf. No. 2.560. (M.p. 41 $^{\circ}$.)
I405	262	178	Isoamylphenylhydrazine, $(\text{C}_6\text{H}_{11})\text{Ph.N.NH}_2$. — Prob. reduces Tollen's reagt.
I406	262	191	Ethylisoamylaniline, Et, $(\text{C}_6\text{H}_{11}).\text{N.Ph}$. — Oil. Tert. amine.*
I407	263		\ddagger Tetramethyl-m-phenylenediamine, $\text{C}_6\text{H}_4(\text{NMe}_2)_2$. — Sp. gr. 0.9879 (15.8 $^{\circ}$). M.p. — 2°.
I408	262-4		Phenylethylenediamine, Ph.NH.CH ₂ .CH ₂ .NH ₂ . — Thick oil misc. w. aq. to strongly alk. sol. — Picrate, \ddagger flat ndl. fr. gl. ac. ac., m.p. 142-3 $^{\circ}$.
I409	264-5	157	2,4-Dimethylquinoline, Me ₂ C ₆ H ₄ N. — Sp. gr. 1.0611 (15 $^{\circ}$). Tert. amine.* — B.Pk, \ddagger tbl., d.s. h. alc., m.p. 193.5 $^{\circ}$ (r.h.). — B ₂ H ₂ PtCl ₆ , \ddagger flesh-colored ndl. fr. dil. HCl, m.p. 229 $^{\circ}$; also red ndl. w. 2H ₂ O.
I410	265d.	91	1-Aminopropanediol(2,3), NH ₂ .CH ₂ .CH(OH).CH ₂ .OH. — Odor, weak basic. E.s. aq. (!), alc.; i. eth. Sp. gr. 1.175 (20/4). Prim. amine.* — B ₂ H ₂ PtCl ₆ , \ddagger ndl. e.s. aq., alc., m.p. 185 $^{\circ}$.
I411	260-70	241	Diocytamine, $(\text{C}_6\text{H}_{17})_2\text{NH}$. — Sec. amine.*
I412	266-7		Triethylenetetramine, NH ₂ .C ₂ H ₄ .NH.C ₂ H ₄ .NH.C ₂ H ₄ .NH ₂ . — M.p. + 12°. S. aq. (!), alc. Sp. gr. 0.982 (15 $^{\circ}$). — B ₂ H ₂ Cl ₂ . \parallel i. aq.
I413	266 (th.i.); 261-3 (th.i.)	129	Py-4-(γ)-Methylquinoline, Lepidine, Me.C ₆ H ₄ N. — [In coal tar.] Odor unpleasant, basic, quinoline-like. Solidified by freezing mixt. Alm. i. aq.; misc. alc., eth., bz., lgr. Sp. gr. 1.0862 (20 $^{\circ}$). — Tert. amine.* Oxidn. by CrO ₃ mixt. gives cinchonic ac., No. 2.302. — B.Pk, \ddagger yel. cryst. ppt. fr. alc. sol., m.p. 208 $^{\circ}$; 212-3 $^{\circ}$.
I414	266-7		Nicotine, C ₁₀ H ₁₁ N ₂ . — [Fr. tobacco.] — Odor suggestive of parsley & pyrrole. Brown on keeping. Diacid tert. base.* Sp. gr. 1.0778 (12.5/4). $[\alpha]_D^{27} = -46.41^{\circ}$. Misc. aq.! — Picrate, \ddagger yel. pr., m.p. 165 $^{\circ}$.
I415	266	157	5,8-Dimethylquinoline, Me ₂ C ₆ H ₄ N. — M.p. + 4-5 $^{\circ}$. Sp. gr. 1.070 (21 $^{\circ}$). Tert. amine.* — B ₂ H ₂ Cr ₂ O ₇ , or. ndl. fr. aq., m.p. 149 $^{\circ}$.
I416	267	157	3-Ethylquinoline, Et.C ₆ H ₄ N. — Tert. amine.* Oxid. by CrO ₃ mixt. to quinoline-3-carbonic ac., No. 2.484. — Picrate, \ddagger yel. ndl., m.p. 163 $^{\circ}$.
I417	265-70		Methyl-m-phenylenediamine, MeNH.C ₆ H ₄ .NH ₂ .
I418	268-9c.		6,8-Dimethylquinoline, Me ₂ C ₆ H ₄ N. — Sp. gr. 1.0665 (4 $^{\circ}$). Tert. amine.*
I419	268-70		p-Aminobenzylamine, NH ₂ .C ₆ H ₄ .CH ₂ .NH ₂ . — Mod. s. aq.; more s. alc.; i. eth. Absorbs CO ₂ fr. air. Sp. gr. 1.08 (20 $^{\circ}$).
I420	268-70; 258		Dimethyl-m-phenylenediamine, Me ₂ N.C ₆ H ₄ .NH ₂ .
I421	269.5-70.5		3-(β)-Phenylpyridine, Ph.C ₆ H ₄ N. — I. aq.; e.s. alc., eth. Odor like diphenylamine. Sp. gr. 1+. Oxid. by KMnO ₄ + H ₂ SO ₄ to benzoic ac. & pyridine-3-carbonic ac. No. 2.410. — Picrate, \ddagger yel. ndl. fr. alc., m.p. 161-3.5 $^{\circ}$.
I422	269-71		2-(α)-Phenylpyridine, Ph.C ₆ H ₄ N. — I. aq. Sp. gr. 1+. Oxid. by CrO ₃ mixt. to pyridine-2-carbonic ac. No. 2.130. — Picrate, \ddagger yel. ndl., s. h. alc., m.p. 175 $^{\circ}$. — B ₂ H ₂ PtCl ₆ , \ddagger (dried at 100 $^{\circ}$), i. aq., alc., m.p. 204 $^{\circ}$.

Explanation of typographical signs used in this Division: * = T. 2.35. \ddagger = T. 2.23. $\ddot{\delta}$ = T. 2.14. \parallel = T. 2.13. $\overline{\eta}$ = T. 2.1. $\overline{\alpha}$ = T. 2.36. $\overline{\beta}$ = T. 2.37. $\overline{\gamma}$ = T. 2.38.

No.	Boiling-point (C.).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
I423	270c.	161	α -Tetrahydronaphthobenzylamine, $C_{10}H_{11}.CH_2.NH_2$. — Odor, sweetish ammon. Absorbs CO_2 . Prim. amine.* Oxidn. by alk. $KMnO_4$ gives oxalic & phthalic ac. — Acetyl deriv., pearly lft., m.p. 88.5. — Phenylurea deriv. (fr. phenyl isocyanate) e.s. h. alc., m.p. 88.5°.
I424	271c.	161	β -Tetrahydronaphthobenzylamine, $C_{10}H_{11}.CH_2.NH_2$. — Absorbs CO_2 . Prim. amine.* Oxidn. by alk. $KMnO_4$ gives phthalic ac., etc. — Acetyl deriv., ndl. fr. dil. alc., m.p. 64–5°. — Phenylurea deriv. (fr. phenyl isocyanate), ndl. fr. alc., m.p. 141°.
I425	271–4c.	157	4-Ethylquinoline, $Et.C_6H_5N$. — Tert. amine.* — $B.Pk$, † yel. ndl., m.p. 178–80°. — $B_2H_2PtCl_6$, § m.p. 204° d.
I426	273–4 (th.i.)	157	4,8-Dimethylquinoline, $Me_2C_6H_4N$. — Pungent oil. Alm. i. aq.; e.s. alc., eth. Tert. amine.* — $B_2H_2PtCl_6.H_2O$, § m.p. 220°. — $B_2H_2AuCl_6$, ndl. fr. HCl , m.p. 181§.
I427	273–4		6,5-(or 4)-Dimethylquinoline, $Me_2C_6H_4N$. — Tert. amine.*
I428	275–8		Metanicotine, $C_{10}H_{14}N_2$. — Opt. i. Misc. aq.; d.s. eth. Sp. gr. 1.006 (16/4). — $B_2P_2H_2O$, † m.p. 114°; (anhydrous) 163°. — $B_2H_2PtCl_6$, § yel.-brown pr., d.s. aq., m.p. abt. 255° d.
I429	276		1-Phenylimidazole, $[NPh.CH:CH.N:CH^2]$. — M.p. +13°. I. aq. Misc. alc. — $B.Pk$, † yel. ndl. fr. alc., m.p. 152°.
I430	276.7	169	2-Benzylpyridine, $Ph.CH_2.C_6H_4N$. — Odor, lemon-like. I. aq.; v.s. alc., eth. Sp. gr. 1.0536 (20/0). — $B.Pk$, † cryst. fr. alc., m.p. 140°. — $B_2H_2PtCl_6$, § powd. s. h. aq., m.p. 183°.
I431	abt. 276 (th.i.)	171	Dimethyl- α -naphthylamine, $NMe_2C_{10}H_7$. — Odor petroleum-like. Shows intense green fluor. Sp. gr. 1.0446 (15/15).
I432	277	147	† ar- α -Tetrahydronaphthylamine, $C_{10}H_{11}.NH_2$. — Viscous oil of aromatic odor. Reduces Au, Ag, & Pt salts, but not Fehling's sol. Prim. amine. — Acetyl deriv., (acetylated by 1 pt. NaA + 2 pt. acetic anhydride), silky ndl., s. h. aq., m.p. 158°.
I433	277–8	171	2,4,6-Trimethylquinoline, $Me_3C_6H_3N$. — Cryst. w. H_2O fr. aq., m.p. 62°. R.d. vol. w. st. — $B.Pk$, † green-yel. ndl. fr. acetone, m.p. 200–1°.
I434	275–80	233	Diisoamylaniline, $(C_6H_{11})_2NPh$. — Oil. Sec. amine.*
I435	280 (th.i.); 273–4	157	4,6-Dimethylquinoline, $Me_2C_6H_4N$. — Solid at 0°. Alm. i. aq.; e.s. alc., eth. — $B.Pk$, † ndl. fr. h. alc., d. abt. 230°. — $B_2H_2PtCl_6.2H_2O$, § ndl. fr. conc. HCl , d. 231°.
I436	281–2c.		Phenyltrimethylenediamine, $Ph.NH.CH_2.CH_2.CH_2.NH_2$. — Sp. gr. 1.025 (15°). S. in 300 pt. c. aq. — B_2P_2 , † greenish cryst., d. 195°. — B_2HCl , ndl. v.s. aq.
I437	280–2		Bipyridyl, $NC_6H_4.C_6H_4N$. — D. vol. w. st. Salts usually sol. — B_2HCl . — Picrate, † yel. ndl., m.p. 208°.
I438	283–5		3,5-Diaminotoluene, $(NH_2)_2C_6H_4Me$. — Syrup. V.s. aq.! — B_2HCl , ndl., v.s. aq., m.p. 255–60° d. — Diacetyl deriv., pr., m.p. 235–6°.
I438-I	287–9		3,2'-(α)-Bipyridyl, $C_{10}H_8N_2$. — Odor pyridine-like. Alm. i. aq.; e.s. eth. Little vol. w. st. — $B.Pk$, † dull yel. ndl. M.p. 149.5°, v.d.s. c. alc.
I439	287–9	211	† Ethylbenzylaniline, $Et.(Ph.CH_2).NPh$. — I. aq.
I440	288–9	183	Benzhydrylamine, Aminodiphenylmethane, $Ph_2CH.NH_2$. — Absorbs CO_2 . Prim. amine.* — B_2HCl , ndl., m.p. 270°.
I441	288–90		m-Nitrodiethylaniline, $NO_2.C_6H_4.NEt_2$. — Yel. oil.
I442	290; 283–5		Diethyl- α -naphthylamine, $NEt_2.C_10H_7$. — Browns on keeping. Misc. alc., eth., bz. Sp. gr. 1.005. — Convert into red nitroso deriv., m.p. 165° (Cf. Soc., 41, 180).

No.	Boiling-point (C°).	Neut. Equiv., etc.	BASIC COMPOUNDS.—Colorless and Liquid.
1443	293	157	Methyl- α -naphthylamine, $\text{Me.NH.C}_{10}\text{H}_7$. — Reddish oil soon becoming opaque in air. I. aq.; e.s. alc., eth. — Sec. amine.* — \textcircled{P} FeCl_3 colors alc. sol. dark violet. — Acetyl deriv., m.p. 95°. — Nitroso deriv., ** golden cryst. fr. bz., m.p. 157° d.
1444	295.5–6.5c.		3,3'-Bipyridyl, $\text{NC}_6\text{H}_4\text{C}_6\text{H}_4\text{N}$. — Deliq. ndl., m.p. 68°, but usually obtained as oil. Misc. alc., d.s. eth. Sp. gr. 1.1635 (20°). Oxidn. by $\text{KMnO}_4 + \text{H}_2\text{SO}_4$ gives pyridine-3-carbonic ac. No. 2.410. — $\text{B.Pk.} \ddagger$ mic. pr., v.d.s. alc.
1445	298	157	Methyl- β -naphthylamine, $\text{MeNH.C}_{10}\text{H}_7$. — Turns dark in air. Sec. amine.* — Nitroso deriv., ** pearly lft. fr. dil. alc., m.p. 90°.
1446	297–300	185	2,5,6,8-Tetramethylquinoline, $\text{Me}_4\text{C}_8\text{H}_4\text{N}$. — Cryst. mass. M.p. 20°. I. aq.; e.s. alc., eth.
—	298–300	183	Benzylaniline. — Cf. No. 2.541. (M.p. 32°.)
1447	306c.		Ethyldibenzylamine, $\text{NEt}(\text{Ph.CH}_3)_2$. — Tert. amine.*
1448	309 ± 6 sl. d.	197	† Dibenzylamine, $(\text{Ph.CH}_3)_2\text{NH}$. — Odor aromatic like brom-toluene. I. aq.; e.s. alc., eth. Sp. gr. 1.034 (15/15). Sec. amine.* — \textcircled{D} Prepare the picramide by procedure of T. 2.22. Recrystd. fr. gl. ac. ac. gives OY-YO ndl., m.p. 173°. — B.HCl , m.p. 255.1–6.1° u.c. (263.8–4.6° c.).
1449	310–1	197	Diphenylethylamine, $\text{Ph.CH}_3\text{CH}(\text{NH}_2)\text{Ph}$. — Sp. gr. 1.031 (15°). Prim. amine. V.s. alc., eth. — Acetyl deriv., long. ndl. fr. alc., m.p. 148°.
1450	312–3	197	p-Benzyltoluidine, $\text{Ph.CH}_3\text{NH.C}_6\text{H}_4\text{Me}$. — Yel. oil. Solidifies slowly. Sec. amine.* E.s. alc., eth.
1451	315–6; 191 (25 mm.)		Ethyl- β -naphthylamine, $\text{C}_{10}\text{H}_7\text{NHEt}$. — Sec. amine.* Does not solidify. — B.HCl , lft. d.s. aq.; m.p. 235°.
1452	318 (th.i.)		Diethyl- β -naphthylamine, $\text{C}_{10}\text{H}_7\text{NET}_2$. — Viscous oil. Tert. amine.* — B.HCl , ndl. fr. aq., m.p. 175°. — $\text{B}_2\text{H}_2\text{PtCl}_6$, decd. by boiling aq., m.p. 95°.
1453	abt. 317		Propyl- α -naphthylamine, $\text{C}_{10}\text{H}_7\text{NH.Pr}$. — Oil. I. aq. Sec. amine.* — Acetyl deriv., m.p. 93–4°.
1454	322–4		Propyl- β -naphthylamine, $\text{C}_{10}\text{H}_7\text{NH.Pr}$. — Sec. amine.*
1455	327–8 (th.i.)		Benzyl-phenylethyl-amine, $\text{Ph.CH}_3\text{NH.CH}_2\text{CH}_2\text{Ph}$. — Sec. amine.* Misc. alc., eth. — B.HCl , plates, m.p. 264–6°.
1456	abt. 328c. sl. d.; 311–0.5, (723 mm.); 180–1 (20 mm.)	k.10 ⁴ = 1+	† Sparteine, Lupinidine, $\text{C}_{18}\text{H}_{28}\text{N}_2$. — [Alkaloid fr. Spartium scoparium (broom)]. Colorless oil w. pronounced basic, piperidine-like odor. Taste v. bitter. V.d.s. aq.; e.s. alc., eth., chlf.; i. bz., lgr. Sp. gr. 1.0199 (20°). $[\alpha]_D = -14.6$ (in alc. sol.). $n_D = 1.5291$. — \textcircled{D} (1) Mix 1 drop B , or cryst. of a salt, w. 2 drops ammon. polysulphide. Heat 2 or 3 min. on water-bath. An OR-RO color develops. — (2) Dis. 1 drop B in 5 cc. aq. Add 2 cc. T.S. of I in K1. An OS1 ppt., becoming cryst. & greenish black within 2 hr., forms. — (3) T. 2.2-(a,f,g) give no colorations.
			— \textcircled{D} Dis. 0.1 g. B in 5 cc. aq. + 2 drops HCl (sp. gr. 1.12). Add 4 cc. c. sat. aq. picric ac. sol. Wash curdy ppt. w. 2 cc. aq. Dis. in 4 cc. h. alc. Add 4 cc. aq. & heat to boiling. Allow to stand until crystn. is complete. Filter. Wash w. 1 cc. alc. Dry 15 min. at 100°. The product, B.2Pk , forms YT1 cryst., m.p. 178–80° u.c., sl. d. — $\text{B}_2\text{H}_2\text{PtCl}_6.2\text{H}_2\text{O}$, § rhomb. pr. fr. HCl , v.d.s. aq., alc., m.p. 243° (frothing), darkening fr. 239°.
1457	344–8		Dicarvacrylamine, $(\text{C}_{10}\text{H}_{12})_2\text{NH}$. — Sec. amine.* Sol. in conc. H_2SO_4 colored blue by HNO_3 . — Acetyl deriv., ¶ scales fr. bz., m.p. 78°.
1458	370		Trioctylamine, $(\text{C}_8\text{H}_{17})_3\text{N}$.

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 ¶ = T. 2.1. ** = T. 2.36. §§ = T. 2.37. §§ = T. 2.38.

SUBORDER I OF ORDER II.
COLORLESS COMPOUNDS CONTAINING C, N, H, AND O.

GENUS III, NEUTRAL COMPOUNDS.

DIVISION A, SOLID SPECIES.

No.	Melting-point (C. $^{\circ}$).	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
1459	-5	m-Nitrostyrene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{CH}_2$. — \oplus Nitro comp.‡ — Odor cinnamon-like. E.s. abe. alc., eth., lgr.
1460	+4	Lauronitrile, $\text{C}_{11}\text{H}_{22}\text{CN}$. — \oplus Sapn. T.* products: NH_3 ; lauric ac. (Vol. I.). — B.p. 198° (100 mm.). Sp. gr. 0.827 (15°).
1461	11	Diocetylketoxime, $(\text{C}_2\text{H}_5)_2\text{C}:\text{NOH}$. — \oplus Oxime.§ Waxy tbl.
1462	12-3.5	o-Nitrostyrene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{CH}_2$. — \oplus Nitro comp.‡ — Vol. w. st. S. in conc. H_2SO_4 w. blue color!
1462-1	13	Nitroglycerine, Glycerol Trinitrate, $\text{NO}_2\text{CH}_2\text{CH}(\text{NO}_2)\text{CH}_2\text{NO}_2$. — \oplus (1) Gives blue color in T. 2.15. — (2) A small drop explodes violently when heated by flame in thin capillary, or struck sharply w. hammer on anvil. — Pale yel. or colorless oily liq. at ordinary temperature. Sp. gr. 1.6009 (15°). S. in 800 pt. aq., in 4 pt. abs. alc.; misc. eth., chlf., gl. ac. ac. Poisonous. — Sapn. w. aq.-KOH sol. gives glycerine (T. 1.816), etc.
—	15.9	† m-Nitrotoluene. — Cf. No. 2.2815.
1463	15-16	Propylsuccinimide, $[\text{NPr.CO.CH}_2\text{CH}_2\text{CO}]$. — \oplus \ominus Sapn. T.* products: propylamine (No. 2.1067); succinic ac. (Vol. I.). — B.p. 247-8°.
1464	19	Myristonitrile, $\text{C}_{18}\text{H}_{37}\text{CN}$. — Sapn. T.* products: NH_3 ; myristic ac. (Vol. I.). — B.p. 226° (100 mm.). Sp. gr. 0.828 (19/4).
1465	20	6-Nitropseudocumene, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}_2[\text{Me}_2 = 1,2,4]$. — \oplus Nitrocomp.‡ — Thick pr. alm. i. aq.
1466	20	Diheptylketoxime, $(\text{C}_7\text{H}_{15})_2\text{C}:\text{NOH}$. — \oplus Oxime.§ — Tbl. fr. c. dil. MeOH; v.s. MeOH, eth., lgr.
1467	23	Oximinocycloheptane, Suberoxime, $[\text{CH}_2\text{C}(:\text{NOH})(\text{CH}_2)_5]$. — \oplus Oxime.§ Odor camphorous! B.p. 230°. Sp. gr. 1.023 (20°). 4-sided pr. i. aq.; e.s. alc., eth., lgr. Dec. in moist air to NH_3 & suberone.
1467-1	24	tert.-Nitrobutane, $\text{Me}_2\text{C.NO}_2$. — \oplus Prob. gives T. 2.15. — Misc. alc., eth. I. KOH sol.
1468	24.5	† Methyl o-Aminobenzoate, $\text{NH}_2\text{C}_6\text{H}_4\text{CO}_2\text{Me}$. — \oplus Synthetic perfume of agreeable odor. Occurs in orange blossom oil. Alc. sol. shows bluish violet fluor. Cryst. mod. s. aq.; e.s. alc., eth.; d.s. mineral ac. Vol. w. st. B.p. 135.5° (15 mm.). Sp. gr. 1.168 (15°). Gives YO color in pine splinter T. 2.24-b. — B.Pk (T. 2.23) ndl., m.p. 105-6°. — [Cf. Ber., 24, 296; 24, 2335 for discussion of detection & determination in essential oils.]
1469	25	γ -Aminobutyric-acid-anhydride, Pyrrolidone, $[\text{NH}_2\text{CO.CH}_2\text{CH}_2\text{CH}_2]$. — \oplus V.s. aq. Deliq. in air, finally solidifying to 6-sided tbl. w. 1 mol. aq., m.p. 35°. — Cryst. mass fr. pet.-eth. V.s. alc., eth., chlf., bz.; d.s. pet.-eth.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		"Aq. sol. reacts neut." Odor like acetamide when warmed. — \textcircled{D} Conc. aq. sol. dis. yel. HgO . The sol. on evapn. yields colorless ndl. of $\text{Hg}(\text{C}_4\text{H}_9\text{NO})_2 \cdot \text{H}_2\text{O}$ (losing H_2O at 100°), m.p. abt. 218° d., browning fr. 180°.
1470	26	Ethylsuccinimide , $[\text{N}=\text{C}(\text{OCH}_2\text{CH}_2\text{CO})_2]$. — \textcircled{P} \textcircled{D} Sapon. T.* products: EtNH_2 (No. 2.1062); succinic ac. (Vol. I). E.s. aq., alc., eth. B.p. 234°.
1471	26.2	Isosuccinonitrile , $\text{Me.CH}(\text{CN})_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_2 ; iso-succinic ac. ($\text{C}_4\text{H}_6\text{O}$, Vol. I). — Long ndl. S. aq. B.p. 197–8°.
1472	27	m-Nitrobenzyl Alc. , $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{OH}$. — \textcircled{P} Nitro comp.† — Rhomb. cryst. B.p. 175–80° (3 mm.).
1473	28	Methylacetamide , Me.CO.NH.Me . — \textcircled{P} \textcircled{D} Sapon. T.* products: MeNH_2 (No. 2.1059); acetic ac. (Vol. I). — Ndl. B.p. 206°.
1474	28	Propylpentadecylketoxime , $\text{Pr.C}(:\text{NOH})\text{C}_{11}\text{H}_{24}$. — \textcircled{P} Oxime.§
1475	29	p-Nitro-o-xylene , $\text{NO}_2\text{C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Nitro comp.† — B.p. 258° (th.i.) sl. d. Sp. gr. 1.139 (30°). Pale yellowish pr. fr. alc. Misc. alc. in all proportions at 30°.
1476	29	p-Nitrostyrene , $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{CH}_2$. — \textcircled{P} Nitro comp.† — Pr. fr. lgr. E.s. warm lgr., but d.s. c. Changes to i. comp. on heating or long keeping.
—	29.5(?)	p-Tolunitrile . — Cf. No. 2.1498.
1477	29–30	Malononitrile , $\text{CH}_2(\text{CN})_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_2 ; malonic ac. (Vol. I). — B.p. 218–9°. Cryst. s. in 7.5 pt. aq., 2.5 pt. alc., or 5 pt. eth.
1478	29	Palmitonitrile , $\text{C}_{15}\text{H}_{31}\text{CN}$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_2 ; palmitic ac. (Vol. I). — B.p. 196° (15 mm.). Sp. gr. 0.822 (31/4).
1479	29	Triethyl n-Cyanurate , $(\text{EtOCN})_3$. — \textcircled{P} Ethyl ester.* 100 pt. c. aq. dis. 0.7 pt., sol. becoming turbid at 29°. E.s. alc., eth. E. vol. w. st.
1480	30	3-Nitrocumene , $\text{NO}_2\text{C}_6\text{H}_5\text{Me}$; (Me. = 1,2,4). — \textcircled{P} Nitro comp.† — Alm. i. c. aq.
1481	30	Ethyl o-Nitrobenzoate , $\text{NO}_2\text{C}_6\text{H}_4\text{CO}_2\text{Et}$. — \textcircled{P} Nitro comp.† & ethyl ester.* — Triclin. cryst.
1482	30	Methylformo-p-toluidine , $\text{Me.C}_6\text{H}_4\text{NMe}(\text{CHO})$. — \textcircled{P} \textcircled{D} Sapon. T.* products: methyl-p-toluidine (No. 2.1281); formic ac. (Vol. I). — B.p. 273–7°.
1483	30.5	Methylphenylcyanamide , Me.PhN.CN . — Lft. fr. chlf. B.p. 135° (10 mm.).
1484	31	p-Nitrodiphenylmethane , $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{CH}_2\text{Ph}$. — \textcircled{P} Nitro comp.† Cryst. fr. c. lgr. E.s. alc., eth.; d.s. c. lgr.
1485	31–2	Dimethylmalononitrile , $\text{Me}_2\text{C}(\text{CN})_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_2 ; dimethylmalonic ac. ($\text{C}_4\text{H}_6\text{O}_4$, Vol. I). — Cryst. fr. lgr. w. camphorous odor. Sbl. in ndl. B.p. 169.5° (th.i.). D.s aq.; e.s. alc.
1486	32	5-Nitro-3-tert.-butyltoluene , $\text{NO}_2\text{C}_6\text{H}_4\text{Me}(\text{Me}_2\text{C})$. — \textcircled{P} Nitro comp.† B.p. 120° (15 mm.). — Fuming HNO_3 gives dinitro deriv., m.p. 175°.
1487	32–3	Benzoyl Cyanide , Ph.CO.CN . — \textcircled{P} Odor, pungent, aromatic, v. irritating! — Slow cooling solidifies to tabular cryst. B.p. 206–8°. — \textcircled{D} Sapon. by alk. (T. 2.26–c) to benzoic ac., m.p. 121° (Vol. I), formic ac., & NH_3 .
1488	32.5–3	1-(p)-Tolylpyrazole , $[\text{N}(\text{C}_6\text{H}_5\text{Me})\text{N}:\text{CH}.\text{CH}:\text{CH}]$. — B.p. 258–9° (th.i.). — Lst. of greasy feel fr. alc. — $\text{B}_2\text{H}_6\text{PtCl}_6 \cdot 2\text{H}_2\text{O}$, or. ndl. fr. aq.; (aq. lost at 110°), m.p. then 183–4° d. (T. 2.14).
1489	33; 35	† α-anti-Benzaldoxime , $\text{Ph.CH}:\text{NOH}$. — \textcircled{P} Oxime.§ giving good reduction & hydroxylamine reactions & benzaldehyde odor in T. 2.17-(a & b). — Odor, aromatic. B.p. 118–9° (10 mm.). D.s. aq.; e.s. alc., eth., bz.
1490	34	m-Nitrophenol-ethyl-ether , $\text{EtO.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Nitro comp.† — B.p. 264° sl. d.
1491	34.5	† aa-Diphenylhydrazine , $\text{Ph}_2\text{N.NH}_2$. — \textcircled{P} Dis. in pure conc. H_2SO_4 w. the intense blue color of T. 2.15! — Reduces Tollen's reagt. in T. 2.30 after shaking 20–30 sec. — B.p. 220° (40–50 mm.). Cryst. fr. lgr. after distn. in vacuo in monoclin. tbd.; but usually a slightly colored oil. V.d.s. aq.; e.s. eth., alc. — \textcircled{D} Dis. 0.05 g. substance in 3 cc. boiling dil. (2:1) alc.

Explanation of typographical signs used in this Division * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		Add 2 drops benzaldehyde & boil 30 sec. Shake. Cool & filter. Wash w. 5 cc. dil. alc. Recryst. fr. 30 cc. boiling dil. (2:1) alc. Wash w. 4 cc. dil. alc. Dry at 100°. The product, benzaldehydediphenylhydrazone, is obtained in fine colorless or v. faintly yellowish ndl., m.p. 123° u.c.
1492	35-6	Hexylpentadecylketoxime, $C_9H_{12} \cdot C(=NOH) \cdot C_{15}H_{32}$. — Oxime. § — Flat ndl. fr. dil. wood spirit.
1493	37	o-Nitrobiphenyl, $Ph \cdot C_6H_4 \cdot NO_2$. — ⊖ Nitro comp. † — B.p. abt. 320°. Lst. fr. alc. I. aq.
1494	37	3-Methyl-1-phenylpyrazole, $[NPh \cdot N : CMe \cdot CH : CH^2]$. — ⊖ Odor quinoline-like. Vapors provoke sneezing. — B.p. 255° (th.i.). Vol. w. st. Thick ndl. fr. dil. alc. E.s. alc., eth., chlf., bz., lgr. "Weak base." — Gives fuchsine-red color in "Knorr's pyrazole react." (Ann., 238, 200; Ber., 26, 100!) — $B_2H_2 \cdot PtCl_6 \cdot 3H_2O$ (fr. conc. HCl), m.p. 153°.
1495	37.5 u.c.	† α-Naphthonitrile, $C_{10}H_8 \cdot CN$. — ⊖ Gives NH_3 in T. 2.7. — Broad lust. ndl. fr. c. lgr. Odor aromatic, recalling brombenzene & naphthalene. I. aq.; e.s. alc.; v.s. eth. — ⊖ Boil 0.3 g. nitrile w. 1 g. KOH + 10 cc. alc. for 30 min. under reflux. Evap. to dryness. Treat residue w. 10 cc. c. aq. Filter. Wash w. 3 cc. aq. Recryst. fr. 15 cc. h. 50% alc. Wash w. 2 cc. 50% alc. Dry 15 min. at 100°. The product, α-naphthionamide, $C_{10}H_7 \cdot CO \cdot NH_2$, is obtained in lust. white ndl., m.p. 202° u.c. (206.7° c.).
1496	37.5	1,2-Dinitroethane, $NO_2 \cdot CH_2 \cdot CH_2 \cdot NO_2$. — ⊖ Should give blue color in T. 2.15. — 4-sided pr. I. aq.; e.s. alc., eth.
1497	38	m-Nitrophenolmethylether, $MeO \cdot C_6H_4 \cdot NO_2$. — ⊖ Nitro comp. † — Flat ndl. fr. alc. E. vol. w. st. B.p. 258° (th.i.).
1496	38; 29.5	† p-Tolunitrile, $Me \cdot C_6H_4 \cdot CN$. — ⊖ D. Sapd. by procedure of T. 2.26-c to NH_3 & p-toluic ac., using 35 cc. aq. in the final crystn. & bone blacking. The p-toluic ac. is obtained in alm. quantitative yield in ndl., m.p. 176-7° u.c. — Cryst. mass of strong aromatic, bitter almond-like odor. B.p. 217.3° c. [A preparation fr. Kahlbaum of Berlin melted at 29-30°. Kröher (Ber., 23, 1030) found the m.p. 29.5°. The m.p. of 38° was obtained by Pinner & Caro (Ber., 27, 3275) under circumstances which seem to give it greater weight.]
1499	38	s-Diisoamylurea, $(C_6H_{11} \cdot NH)_2 \cdot CO$. — ⊖ ⊖ Sapn. T.* products: $C_6H_{11} \cdot NH_2$ (No. 2.1100); CO_2 . — B.p. 270°. I. c. aq.; e.s. alc., eth.
1500	38	Diacetanilide, $Ph \cdot N(CO \cdot Me)_2$. — ⊖ Should give powerful carbylaniline odor in T. 2.12 (!); & aniline & ac. ac. in Sapn. T.* — [One acetyl group is so easily removed by alk. that it may influence the Generic T.]
1501	38.5	3-Methyl-1-phenyl-5-oxyprazolemethylether, $[N : CMe \cdot CH : C(OH) \cdot NPh]$. — B.p. 300-2°. Tbl. fr. lgr. I. aq., alk.; e.s. alc., eth. — Heated w. conc. HCl at 160° gives 3-methyl-1-phenylpyrazolone(5), No. 2.106.
1502	38-9	α-Dimethylaminobenzophenone, $Me_2N \cdot C_6H_4 \cdot CO \cdot Ph$. — B.p. 330-40°. Ndl. fr. lgr. E.s. alc.
1503	39	Nitrosoformanilide, $NO \cdot N(CHO) \cdot Ph$. — ⊖ Should give blue color in T. 2.15. — Yellowish white e.s. ndl. V. unstable.
1504	39-40	β-Naphthindole, $[C_{10}H_8 \cdot NH \cdot CH : CH^2]$. — ⊖ Gives intense blue-violet color in pine splinter indole react., T. 2.24-b. — E.s. alc., eth., chlf.; r.d.s. lgr.; d.s. aq.
1505	39-40	Lauroneoxime, $(C_{11}H_{22})_2 \cdot C : NOH$. — ⊖ Oxime. § — Ndl. fr. dil. alc.; e.s. bz., lgr.
1506	39	Isopropylacetanilide, $Ph \cdot NPr \cdot (Me \cdot CO)$. — ⊖ ⊖ Sapn. T.* products: isopropylaniline (No. 2.1302); acetic ac. (Vol. I). — B.p. 262-3° (th.i., 712 mm.).
1507	39-40	Piperidone, δ-Aminovalerianic Ac. Anhydride, $[CH_2 \cdot NH \cdot CO \cdot (CH_2)_3]^2$. — B.p. 256°. — V.s. aq., alc., eth. — Boiling w. conc. NaOH or HCl gives 5-(δ)-aminovalerianic ac. (No. 2.2238).
1508	40	3-Nitro-2,2-dimethylbutane, $Me \cdot C(Me)_2 \cdot CH(NO_2) \cdot CH_2$. — ⊖ Nitro comp. † — Cryst. mass of camphorous odor. E. volatile.
1509	41	Stearonitrile, $C_{18}H_{36} \cdot CN$. — ⊖ ⊖ Sapn. T.* products: NH_3 ; stearic ac., (Vol. I). — B.p. 214° (13 mm.).

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
—	41-2	† Cyanamide. — Cf. No. 2.1525.
I511	41-2	Dimethylbenzamide, Ph.CO.NMe ₂ . — ⊕ ⊖ Sapn. T.* products: dimethylamine (No. 2.1061); benzoic ac. (Vol. I). — B.p. 255-7°. Cryst. e.s. aq.
I512	41	Phenyl-o-toluidine, Ph.NH.C ₆ H ₄ .Me. — I. aq. B.p. 306° (th.i.). Gives blue-violet coloration w. HNO ₃ !
I513	42	† tert-Amyl Phenylcarbamate, Ph.NH.CO.CEt ₂ .Me. — ⊕ Should give carbylamine odor in T. 2.12. — ⊖ Sapn. T.* products: aniline; dimethyl-ethylcarbinol (C ₆ H ₁₂ O, Vol. I); CO ₂ .
I514	42; 44	† Ethyl o-Nitrocinnamate, NO ₂ .C ₆ H ₄ .CH:CH.CO ₂ .Et. — ⊕ Nitro comp.‡ — Rhomb. cryst. e.s. c. alc., eth. — ⊖ Sapn. T.* products: ethyl alc.; o-nitrocinnamic ac. (No. 2.437).
I515	43	Benzoylacetoxime, Me ₂ C:N.O(CO.Ph). — ⊕ Gives hydroxylamine in T. 2.17. — ⊖ Products by alk. sapn.* acetoxime (No. 2.1597); benzoic ac. (Vol. I). — Cryst. fr. lgr. or eth. D.s. c. aq.; v.s. alc., eth.
I516	43	Acet-m-ditolylamine, Me.CO.N.(C ₆ H ₄ .Me) ₂ . — ⊕ ⊖ Sapn. T.* products: di-m-tolylamine (No. 2.2868), & acetic ac. (Vol. I).
I517	abt. 44	Nitromesitylene, NO ₂ .C ₆ H ₂ .Me ₂ [Me ₂ = 1,3,5]. — ⊕ Nitro comp.‡ — B.p. 255°. Pr. fr. alc. Alm. i. c. aq.
—	44	† o-Nitrobenzaldehyde. — Cf. No. 2.2889 (GYT3 color).
I518	44	Dipropionanilide, (Et.CO) ₂ .N.Ph. — ⊕ Should give carbylamine odor in T. 2.12. — Cryst. fr. lgr. B.p. 179° (30 mm.). — ⊖ Sapn. T.* products: aniline; propionic ac. (Vol. I).
I519	44-5	Diethylmalononitrile, Et ₂ C(CN) ₂ . — ⊕ ⊖ Sapn. T.* products: NH ₃ ; diethylmalonic ac. (C ₇ H ₁₂ O ₄ , Vol. I).
I520	44-5	Benzoyl- α -pipecoline, Ph.CO.NC ₆ H ₅ . — ⊕ ⊖ Sapn. T.* products: α -pipecoline (No. 2.1134); benzoic ac. (Vol. I). — Cryst. mass. E.s. alc.
I521	44	Ethyloctadecylketoxime, Et.C(:NOH).(CH ₂) ₁₇ .Me. — Oxime. § — V.s. alc., eth.
I522	45	Methylnonylketoxime, Me.C(:NOH).C ₆ H ₁₁ . — ⊕ Oxime. § — V.s. alc.
I523	45-6	o-Tolylurethane, Me.C ₆ H ₄ .NH.CO.Et. — ⊕ ⊖ Sapn. T.* products: o-toluidine (No. 2.1262); ethyl alc. (Vol. I); CO ₂ . — Cryst. Vol. w. st. S.c. alc., bz., lgr.
I524	46	Nitrosodiisopropylamine, (Me ₂ .CH) ₂ N.NO. — ⊕ Should give blue color in T. 2.15. — B.p. 194.5° c. V.d.s. aq.; e.s. alc.
I525	46; 41-2	† Cyanamide, CN.NH ₂ . — [According to G. Henschel, Diss. Univ. of Leipzig, the m.p. of pure comp. is 46°; but as usually obtained, 41-2°.] — ⊕ Gives NH ₃ in T. 2.7. Sapn. T.* products: NH ₃ ; CO ₂ . — Polymerizes slowly above m.p. & when impure, v. quickly on keeping. E.s. aq., alc., eth.; less s. chlf., bz. Vol. w. st. Has alkaline "feel." HNO ₃ added to eth. sol. gives ppt. of urea nitrate. CuSO ₄ sol. gives brown-black ppt. w. aq. sol. Ammon. lead acetate gives ppt., pale yel. at first, finally lemon-yel. & cryst.
I526	46	† Formanilide, NHPh(CHO). — ⊕ T. 2.12 gives strong carbylamine odor. — 4-sided pr. fr. slow evapn. aq. sol. S. aq.; e.s. alc.; alm. i. pet.-eth. Taste burning. — ⊖ Sapd. by T. 2.26-a to aniline, No. 2.1235.
I527	46.8	1'-Nitromesitylene, Me ₂ .C ₆ H ₂ .CH ₂ .NO ₂ [Me ₂ = 3,5]. — ⊕ Nitro comp.‡ — Ndl. e.s. alc., eth., bz.; d.s. lgr. — ⊖ Oxidized by boiling dil. HNO ₃ (Cf. T. 1.905-3) to mesitylenic ac. (C ₆ H ₁₀ O ₂ , Vol. I).
I528	47; 42	Ethyl m-Nitrobenzoate, NO ₂ .C ₆ H ₄ .CO ₂ .Et. — ⊕ ⊖ Sapn. T.* products: ethyl alc.; m-nitrobenzoic ac. (No. 2.139). — B.p. 296°; 298°.
I529	47	† Acetaldoxime, Me.CH:NOH. — ⊕ Gives the oxime reactions of T. 2.17. — Ndl. Misc. aq., alc., eth.
I530	47	Methyl Phenylcarbamate, Ph.NH.CO.Me. — ⊕ Should give carbylamine odor in T. 2.12. — Pr. alm. i. c. aq.; e.s. alc., eth. — ⊖ Sapn. T.* products: methyl alc.; aniline; CO ₂ .

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I531	48	Diacet-p-toluide, $\text{Me.C}_6\text{H}_4.\text{N}(\text{C}_2\text{H}_5\text{O})_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); acetic ac.
I532	48	Methyl Anilinoacetate, $\text{Ph.NH.CH}_2.\text{CO}.\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: anilinoacetic ac. (No. 2.99); methyl alc. — Ndl. alm. i. aq.; e.s. alc., eth., HCl. Vol. w. st.
I533	48	N-Benzoylpiperidine, $\text{Ph.CO.NC}_6\text{H}_{10}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: piperidine (No. 2.1112); benzoic ac. (Vol. I).
I534	49	(stable) β -Mesityloxime, $\text{C}_6\text{H}_{10}.\text{NOH}$. — \textcircled{P} Oxime. § — Lft. I. aq.; e.s. alc., eth.; s. NaOH sol. E. vol. w. st. B.p. 102° (13 mm.).
I535	49	Propylacetanilide, $\text{Pr}_3(\text{C}_2\text{H}_5\text{O})\text{N}.\text{Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: propylaniline (No. 2.1329); acetic ac. — B.p. 267° c. Monoclin. tbl. fr. alc. Alm. i. aq.; e.s. alc.
I536	49-50	Acetophenonemethylphenylhydrazone, $\text{Me}.\text{PhC}:\text{N.NPh}.\text{Me}$. — \textcircled{P} Hydrazone. § — Cryst. fr. lgr. I. aq.; e.s. alc., eth., bz., chlf. Dist. undec.
I537	50	Acetoacetesterphenylhydrazone, $\text{Me.C}(:\text{N.NH.Ph}).\text{CH}_2.\text{CO}.\text{Et}$. — Long ndl. Unstable, oxidizing in air. V.s. alc. Warmed w. dil. HCl or alc. KOH gives 3-methyl-1-phenyl-pyrazalone (No. 2.106)!
I538	abt. 50	Myristoneoxime, $(\text{C}_{18}\text{H}_{37})_2\text{C}:\text{NOH}$. — Oxime. §
I539	49-50	Diethyl Iminodicarbonate, $\text{NH}.\text{(CO.Et)}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH; CO ₂ ; ethyl alc.
I540	49-50	† Urethane, Ethyl Carbamate, $\text{NH}_2.\text{CO.Et}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH; EtOH; CO ₂ . — B.p. 180°. Clear, slender ndl. fr. lgr. sol. V.s. c. aq.; alc., eth. Taste of aq. sol. bitter & burning. — Dis. 0.2 g. in 2 cc. warm 10% NaOH. Add abt. 0.1 g. I in small portions — enough to give faint permanent yel. color. Iodoform is produced. (Flückiger.)
I541	50.5-1	Nitrosoacetanilide, $(\text{NO})(\text{MeCO}).\text{N}.\text{Ph}$. — \textcircled{P} Gives Liebermann's reagt. (T. 2.18). — Yel.-white ndl. fr. pet.-eth. V. unstable. I. aq.; e.s. eth.; s. alc.
I542	50-1	b-Formyl-aa-methylphenylhydrazine, Ph.NMe.NH(CHO) . — \textcircled{P} Should reduce Tollen's reagt. after hydrolysis by T. 2.17. — Cryst. fr. eth. E.s. alc., eth. — B.p. 183° (11 mm.).
I543	51-2d.	Nitrosourethane, $\text{NH}(\text{NO}).\text{CO.Et}$. — \textcircled{P} Gives odor of acetaldehyde on fusion. — Ndl. fr. lgr. Decd. by alk. or even by h. aq. to CO ₂ , alc., & N!
I544	51.5-2u.c.	† Phenylurethane, Ethyl Phenylcarbamate, Ph.NH.CO.Et . — \textcircled{P} Gives powerful carbylamine odor in T. 2.12! — Long ndl. fr. h. aq. Odor, faint aromatic. Taste, pungent. Alm. i. c. aq.; e.s. alc., eth. B.p. 237-8° sl. d. — \textcircled{D} Boil 0.1 g. substance w. 1 cc. aniline for 15 min. in reflux t.t., as in T. 1.317-3. Cool. Add 10 cc. dil. HCl (sp. gr. 1.12). Mix well. Filter. Wash residue w. 5 cc. aq. Dis. in 6 cc. boiling 66% alc. Cool, shake, & filter. Recryst. again fr. 4 cc. boiling 66% alc. Wash w. 3 cc. dil. alc. Dry 15 min. at 100°. The product, carbanilide, is obtained in fine white ndl., m.p. 236° u.c.
I545	51-2; 54.5	† Succinonitrile Ethylene Cyanide, $\text{CN}.\text{CH}_2.\text{CH}_2.\text{CN}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; succinic ac. (Vol. I). — Amorph. E.s. aq., chlf., alc.; less s. eth. B.p. 265-7° d.
I546	52u.c.	† Indole, Benzopyrrole, $\text{C}_9\text{H}_7\text{N}$. — Odor, fecal, persistent, like α -naphthylamine!! Lust. lft. fr. lgr. R.s. h. aq.; e.s. alc., eth., hydrocarbons, chlf. E. vol. w. st. \textcircled{P} (1) Gives deep red stain within 1 or 2 min. in the cold in pine splinter pyrrole T. 2.24-b, or more quickly on heating. — (2) [Ehrlich's T.] — To sol. of 0.5 mg. indole in 10 cc. aq., add 5 cc. 2% alc. sol. of p-dimethylaminobenzaldehyde. Then add, drop by drop, 0.5 cc. dil. HCl (sp. gr. 1.12). A VRT2 color, which gradually deepens, appears. — (3) [Legal's T.] — To sol. of 1 mg. indole in 1 cc. aq., add abt. 1 mg. solid sodium nitroprusside & 1 drop 10% NaOH sol. A VRS1 color (by transmitted light), changing to VB on addition of 2 drops conc. HCl, appears. — (4) Add 3 drops HNO ₃ (sp. gr. 1.42) & 1 drop 6.1% aq. KNO ₃ sol. to sol. of 1 mg. indole in 5 cc. aq. An OR color quickly develops, & after standing for some time an unstable RT1 ppt. of nitrosoindole nitrate separates. [Cf. Abderhalden, 4, 851, for bibliography & additional qualitative & quantitative methods.]

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		<p style="text-align: center;">① Add 0.03 g. picric ac. diss. in 1 cc. bz. to sol. of 0.05 g. B in 1 cc. bz. Allow to stand several hr. Filter off ppt. Wash w. 2 cc. c. bz. Recryst. fr. 3 cc. bz. Dry on porous tile 15 min. at 100°. The product, BPk, is obtained in long lust. R ndl., which darken at 160° & melt to black liq. at 176–7° u.c. (frothing at 180°).</p>
1547	52 <u>c.</u>	<p>† p-Nitrotoluene, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}$. — ② Gives nitro group T. 2.21. — B.p. 237.7° c. Colorless ndl. of strong aromatic odor fr. alc. Alm. i. aq.; e.s. alc.; v.s. eth. Taste of sat. aq. sol. at first sweet (No. 2-3 in T. 2.29), then burning! — ③ Convert into 2,4-dinitrotoluene by procedure used in corresponding test for o-nitrotoluene (No. 2.2804), except that 0.1 g. substance is used in place of 3 drops. In crystallizing the product, the sol. is at first milky, but within about 20 min. it will clear w. separation of cryst. of the dinitro deriv. of m.p. 70.4° u.c. (70.7° c.).</p>
1548	52	Dinitro-p-ethyltoluene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}$, Et. — ② Nitro comp.‡ — Tbl. e.s. h. alc.
1549	52.5	2,5-Dinitrotoluene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}$. — ② Nitro comp.‡ — Ndl. fr. alc. E.s. alc., CS ₂ , bz. — Reduced by boiling alc. ammon. sulphide to 6-nitrom-toluidine (No. 2.3261).
1550	52-3	m-Dinitrobenzene-naphthalene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{C}_{10}\text{H}_8$. — ② Nitro comp.‡ — Thick ndl. w. odor of naphthalene fr. bz. — Decd. to m-dinitrobenzene & naphthalene by h. aq.
1551	52-3	Ethylphenylketoxime, Et.C(:NOH)Ph. — Oxime § — Tbl. fr. lgr. B.p. 245–6° d. Corresponding ketone, b.p. 215.5°; m.p. +18.5°.
1552	52	Methyl Carbamate, Urethan, $\text{NH}_2\text{CO}_2\text{Me}$. — ② ③ Sapn. T.* products: NH ₂ ; MeOH; CO ₂ . — B.p. 177°. Tbl. 100 pt. aq. at 11° dis. 217 pt.; 100 pt. alc. at 15° dis. 73 pt.
1553	52	Ethyl p-Tolylcarbamate, $\text{Me.C}_6\text{H}_4\text{NH.COOC}_2\text{Et}$. — Sapn. T.* products: p-toluidine (No. 2.566); ethyl alc.; CO ₂ . — I. aq.; monoclin. pr. fr. alc.
1554	52	4-Formylamino-1,2-xylene, $(\text{CHO})\text{NH.C}_6\text{H}_4\text{Me}_2$. — Sapn. T.* products: 4-amino-1,2-xylene (No. 2.588); formic ac. (Vol. I).
1555	52-3	Diacetonitrile, $\text{Me.C}(\text{NH}).\text{CH}_2\text{CN}$. — ② Gives NH ₂ in T. 2.7. — Ndl. fr. lgr. S. aq.; v.s. alc., eth. Boiling w. aq. gives NH ₂ , HCN, etc. Alc. sol. w. Na gives ethylamine, No. 2.1062.
1556	52-3	Methylphenyl-β-naphthylamine, $\text{MePhN.C}_{10}\text{H}_7$. — Cryst. s. alc.
1557	53	2,2-Dinitropropane, $\text{Me}_2\text{C}(\text{NO}_2)_2$. — ② Nitro comp.‡ — Camphor-like cryst. V.d. s. aq. Sbl. at room temp. B.p. 185.5° c.
1558	abt. 53d.	α-Nitroisobutyronitrile, $\text{Me}_2\text{C}(\text{NO}).\text{CN}$. — ② Melts to deep blue liq. Abt. 80° becomes colorless w. vigorous gas evolution. Deliq. in moist air. E.s. alc.; d.s. bz. E. vol. w. st. or eth.
1559	53; 60	Propyl Carbamate, $\text{NH}_2\text{CO}_2\text{Pr}$. — ② ③ Sapn. T.* products: NH ₂ ; CO ₂ ; propyl alc. (Vol. I). — Pr. V.s. aq., alc. B.p. 194–5°.
1560	53	Cetyl Cyanide, $\text{C}_{16}\text{H}_{33}\text{CN}$. — ② ③ Sapn. T.* products: NH ₂ ; margaric ac. (Vol. I).
1561	53	Formo-p-toluide, $(\text{CHO})\text{NH.C}_6\text{H}_4\text{Me}$. — Sapn. T.* products: p-toluidine (No. 2.566); formic ac. — Ndl.; s. aq., alc.
1562	54 <u>c.</u>	<p>† p-Nitroanisole, $\text{NO}_2\text{C}_6\text{H}_4\text{OMe}$. — ② Nitro comp.‡ — Colorless rhomb. plates w. strong aromatic odor. Taste of sat. aq. sol. sweet (No. 2 in T. 2.29). Alm. i. c. aq.; e.s. alc.; v.s. eth.; sl. s. 10% NaOH to YT1 sol. B.p. 258–60°.</p> <p>③ Mix in 6-inch t.t. 0.5 g. substance, 1.5 g. granulated Sn., & 3 cc. conc. HCl, adding latter in 1 cc. portions to prevent too violent action. Allow reduction to continue until all oily drops disappear. Then add 15 cc. KOH sol. (1 : 2). Warm. Filter through "hardened" paper. Wash residue w. 5 cc. aq. Dis. in 10 cc. boiling aq. Filter hot. Cool filtrate w. ice water. Filter. Air dry on tile. Dis. in 3 to 5 cc. boiling lgr. Filter hot. Cool filtrate w. ice water. Filter, & dry on porous tile. The product, o-anisidine, $\text{NH}_2\text{C}_6\text{H}_4\text{OMe}$, is obtained in nearly colorless rhomb. pr., m.p. 57–7.5° u.c.</p>

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I563	54	2,4-Dinitrocymene, $(NO_2)_2.C_6H.Me.(Me_2CH)$, [Me:C ₆ H = 1:4]. — \oplus Nitro comp.† — Rhomb. tbl.
I564	54	Diethylidiphenylurea, $NET_2CO.NPh_2$. — \oplus \ominus Sapon. T.* products: diethylamine (No. 2.1068-1); diphenylamine (No. 2.1568); CO ₂ . — [Like diphenylamine, below, may give blue color in test w. H ₂ SO ₄ + HNO ₃ .] — Lft. i. aq.; e.s. alc.
I565	54.5	Ethylacetanilide, $Et.(C_6H_5O)NPh$. — \oplus \ominus Sapon. T.* products: ethylaniline (No. 2.1270); acetic ac. — Cryst. v.s. eth. B.p. 258° (th.i., 731 mm.).
I566	53; 55	1,3,5-Trimethylbenzonitrile(2), $Me_3C_6H_3CN$. — \oplus Odor strong & cinnamon-like. B.p. 225–30°.
I567	54–5	Thujoneoxime, $C_{10}H_{14}:NOH$. — \oplus $[\alpha]_D = +108.46^{\circ}$ (in alc., c = 21.79). — Pr. B.p. 135.6° (20 mm.). — May, perhaps, give oxime T. 2.17-a.
I568	54.0 u.c.	† Diphenylamine, Ph_2NH . — Lust. colorless scales w. faint aromat. odor. Tasteless. Alm. i. aq.; v.s. c. alc., eth.; alm. i. dil. min. acids. B.p. 302.0° c. \oplus Dis. 1 or 2 mg. substance in 5 cc. of a mixt. of 2 vol. pure H ₂ SO ₄ (sp. gr. 1.84) + 1 vol. aq. Add trace HNO ₃ on stirring rod. A pure blue color so intense as to render sol. alm. opaque develops at once! Dilution w. conc. H ₂ SO ₄ gives B color. Diln. w. aq. gives violet color. \ominus Dis. 0.1 g. substance in 5 cc. alc. Add 20 drops conc. HCl, warming to dis. if necessary. Cool. Add, drop by drop, 10 drops c. sat. aq. NaNO ₂ sol. Sol. becomes blue, dirty green, & finally yellow, a pale yel. gran. ppt. separating. Add 2 cc. aq., shaking. Chill w. ice water for 5 min., w. occasional shaking. Filter. Wash w. 1 cc. ice water. Dry on porous tile. Recryst. fr. 1 cc. warm lgr. Chill w. ice. Filter. Dry at room temp. on porous tile. The product, diphenylnitrosamine, is obtained in sl. yellowish cryst., m.p. 66.8–6.4° u.c. (65.1–6.7° c.). — Diphenylaminetetrabromide may be prepared by adding sl. x.s. Br to alc. sol. of diphenylamine & recryst. fr. bz. The product melts at 183° (Ann., 132, 166).
I569	53.5–5	β -Methylpiperidone, $^{\beta}NH.(CH_2)_3CHMe.CO$. — Odor coniine-like. B.p. 249–50°. Small cubes fr. lgr. D.s. lgr.; e.s. alc.
I570	55	Erucianilide, $(C_{12}H_{24}O_2)NHPh$. — \oplus \ominus Sapon. T.* products: aniline; erucic ac. (Vol. I). — I. aq.; d.s. alc.; e.s. eth., bz.
I571	55–6	† Methyl-o-acettouide, $Me.(C_6H_5O)N.C_6H_4.Me$. — Sapon. T.* products: methyl-o-toluidine (No. 2.1278); acetic ac. — B.p. abt. 250°. Nitro deriv. (Ann., 304, 98), m.p. 119°.
I572	55–6	3,4,4-Trimethyl-1-phenylpyrazolone(5), $^{\beta}NPh.N:CMe.CMe.CO$. — B.p. 309° c. I. aq., alk., dil. ac.; e.s. alc., eth., chlf., bz. — Alc. sol. + Na gives product whose sol. in dil. acid is colored fuchsine-red by drop FeCl ₃ sol.
I573	56	Ethyl- β -naphthylnitrosamine, $Et.(NO)N.C_{10}H_7$. — \oplus Should give blue color in T. 2.15. — Pure white cryst. I. aq. — \ominus Convert into nitrosoethyl-naphthylamine by Fischer & Hepp rearrangement (Ber., 20, 2471).
I573-I	56	Trinitroethane, $Me.C(NO_2)_3$. — \oplus Odor like nitrous ac. & v. irritating, inducing flow of tears & headache. Persistently shaken w. c. conc. (2:3) KOH sol. gives yel. & then red sol. of the explosive K salt of dinitroethane & KNO ₃ . — Cubical cryst. d.s. aq., lgr.; e.s. other solvents. V. volatile. (Only slowly attacked by weak alk.)
I574	56	Pr-1,2-Dimethylindole, $Me.C_6H_3N.Me$. — \oplus Gives pine splinter coloration in T. 2.24-b. — Ndl. fr. lgr. V.d.s. h. aq.; e.s. alc., eth., bz., h. lgr., conc. HCl. Vol. w. st. — Picrate, fine dark red ndl. fr. bz.
I575	abt. 56	† α Enantholoxime, $Me.(CH_2)_5CH:NOH$. — \oplus (1) Gives oxime react., T. 2.17. — (2) To 1 cc. of a 1% aq. sol. of the oxime acidified w. 1 drop HCl (sp. gr. 1.12), add 1 drop 10% aq. FeCl ₃ sol. A VRTI-RTI color slowly develops. — B.p. 195°. Tbl. fr. alc. D.s. c. aq.; e.s. alc., eth.
I576	56	Ethylheptadecylketoxime, $Et.C:(NOH).C_{17}H_{35}$. — \oplus Oxime. § — Ndl. fr. alc. D.s. alc., pet.-eth.
I577	56.5	Cyclopentanoneoxime, $^{\beta}CH_2.C:(NOH).(CH_2)^2$. — B.p. 196° (th.i.). Glassy pr. fr. lgr. Mod. s. aq.; v.s. alc., eth.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1578	57-8	Nitrodimethylamine, $\text{Me}_2\text{N}.\text{NO}_2$. — \textcircled{P} Nitro comp.‡ — Cryst. e.s. aq., alc., eth., bz. Vol. w. st. B.p. 187°. Reduction w. Zn dust & ac. ac. gives dimethylhydrazine, No. 2.1072.
1579	57	Ethyl p-Nitrobenzoate, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.Et}$. — \textcircled{P} Nitro comp.‡ — Sapn. T.* products: p-nitrobenzoic ac. (No. 2.425); ethyl alc.
1580	57-8; 60	† p-Nitrophenol-ethyl-ether, $\text{NO}_2\text{C}_6\text{H}_4\text{OEt}$. — \textcircled{P} Nitro comp.‡ B.p. 283°.
1581	57	Isobutylbenzamide, $(\text{C}_3\text{H}_7)(\text{Ph.CO}).\text{NH}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: isobutylamine (No. 2.1078); benzoic ac. (Vol. I). — Ndl. fr. bz. B.p. 308-13° sl. d.
1582	57.5	1,2,4-Trimethylbenzonitrile(5), $\text{Me}_3\text{C}_6\text{H}_3\text{CN}$. — \textcircled{P} Sapn. T.* products: NH_3 ; 1,2,4-trimethylbenzene-5-carbonic ac., m.p. 149-50°. — Ndl. fr. alc. B.p. 250°. E.s. alc., eth.
1583	57	1'-Acetaminoethylbenzene, $\text{Ph.CH}(\text{NH.C}_2\text{H}_5\text{O}).\text{Me}$. — \textcircled{P} Sapn. T.* products: 1'-aminoethylbenzene (No. 2.1242); acetic ac. — B.p. 292-3° c. E.s. alc.
1584	57-8	Ethyl Anilinoacetate, $\text{Ph.NH.CH}_2\text{CO.Et}$. — \textcircled{P} Sapn. T.* products: anilino-acetic ac. (No. 2.99); ethyl alc. — B.p. 273-4°. Lft. d.s. h. aq. E.s. eth., HCl, h. alc. Vol. w. st.
1585	58	Pseudobutylnitrol, $\text{Me.C(NO}_2\text{)(NO)Et}$. — \textcircled{P} Melts to blue liq. — Pr. fr. chlf. I. aq.
—	58u.c.	α -Nitronaphthalene. — Cf. No. 2.2917 [GYT2 colored.]
—	58u.c.	† m-Nitrobenzaldehyde. — Cf. No. 2.2918 [GY-Y, T3 colored.]
1586	58	Phenylnitroethylene, Ph.CH:CH.NO_2 . — \textcircled{P} Nitro comp.‡ Odor, cinnamon-like, irritating, provoking flow of tears. Blisters the skin. — Yellowish rhomb. pr. fr. alc. I. c. aq.; d.s. h. aq.; v.s. eth., bz. Polymerizes in moist air & sunlight. Warmed w. KOH sol. gives benzaldehyde! — \textcircled{D} Dibromide (fr. CS_2 sol. + Br_2), ndl. fr. lgr., m.p. 86°.
1587	58	α -Nitrophenyl Benzoate, $\text{Ph.CO.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Nitro comp.‡ Lust. triclin. pr. fr. lgr. I. aq. E.s. eth., bz., chlf.
1588	58	p-Nitrophenyllactic acidketone, $\text{NO}_2\text{C}_6\text{H}_4\text{CH(OH).CH}_2\text{CO.CMe}$. — \textcircled{P} Nitro comp.‡ Cryst. I. c. aq., lgr.; e.s. alc., eth.
1589	58.5	Bz-3-Methylindole, $\text{Me.C}_6\text{H}_4\text{N}$. — \textcircled{P} Gives pine splinter react. (T. 2.24-b). Odor fecal. — Ndl. fr. h. aq. Mod. s. h. aq.; e.s. alc., eth., bz., lgr. Vol. w. st. — \textcircled{D} Picrate (T. 2.23), red ndl. fr. h. aq., m.p. 151°.
1590	58	Phenylacetaldehydephenylhydrazone, $\text{Ph.CH}_2\text{CH:N.NH.Ph}$. — \textcircled{P} Prob. gives hydrazone T. 2.17-a. — Pr. fr. lgr. I. aq.
1591	58	Lactanilide, $\text{Me.CH(OH).CO.NH.Ph}$. — \textcircled{P} Should give carbylamine odor in T. 2.12. — Plates. D.s. c. aq.; v.s. alc., eth.; i. lgr.
1592	57-9	Propyl Phenylcarbamate, Ph.NH.CO.Pr . — \textcircled{P} Should give carbylamine odor in T. 2.12. Ndl. I. c. aq.; s. alc., eth. — \textcircled{D} Sapn. T.* products: aniline (No. 2.1235); propyl alc.; CO_2 .
1593	58	Cerotonitrile, $\text{C}_{12}\text{H}_{11}\text{CN}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: NH_3 ; cerotic ac. (Vol. I). — Fine ndl. fr. alc. I. aq.; e.s. alc., eth.
1594	58	Dipropionyl Dicyanide, $(\text{Et.CO.CN})_2$. — \textcircled{P} \textcircled{O} Sapn. T.* (w. KOH) products: NH_3 ; propionic ac.; HCN. — B.p. abt. 220°. D.s. aq.; v.s. alc., eth., bz.
1595	58	Ethylphenyl- β -naphthylamine, $\text{Et.Ph.N.C}_1\text{H}_7$. — Lft. I. aq.
1596	58-9; 60	α -Nitrobenzylideneacetone, $\text{Me.CO.CH:CH.C}_6\text{H}_4\text{(NO}_2\text{)}$. — \textcircled{P} Nitro comp.‡ — Ndl. V.s. alc., eth., chlf., bz. — If KOH is added to alc. sol., and then, after some time, HCl, and the sol. boiled, indigo separates.
1597	59u.c.	† Acetoneoxime, $\text{Me}_2\text{C:NOH}$. — \textcircled{P} Gives the oxime reactions of T. 2.17. — Odor, faint chloral-like. B.p. 134.8° (th.i.). Pr. V.s. aq., alc., eth., lgr. — $[\text{k}_A \times 10^{13}] = 6.0$; $\text{k}_B \times 10^{13} = 6.5$]

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1598	59	Methylphenylketoxime , Me.C:(NOH).Ph. — \textcircled{P} Oxime. § — Silky ndl. v.s. alc., eth., chlf., bz., lgr. Dist. w. decn. After several days contact w. ac. ac. + HCl changes to acetanilide.
1599	59	Palmitoneoxime , $(\text{C}_{16}\text{H}_{31})_2\text{C:NOH.}$ — Oxime. § — Ndl. fr. gl. ac. ac.
1600	59–60	Pr-2-Methyl-3-phenylindole , $\text{Me.Ph.C}_6\text{H}_4\text{N.}$ — Pr. fr. h. lgr. Odorless! Does not give pine splinter indole react. Not vol. w. st. I. aq.; e.s. alc. eth., bz. — \textcircled{P} Picrate (T. 2.23), dark red ndl. fr. alc., e.s. bz., m.p. 141–2°, after sintering abt. 125°.
1601	60	† Pr-2-(α-Methylindole , Methylketole , $\text{Me.C}_6\text{H}_4\text{N.}$ — \textcircled{P} Odor fecal. Gives red coloration in pyrrole splinter T. 2.24-b. — Cryst. fr. lgr. Vol. w. st. B.p. 272° (th.i., 750 mm.). Somewhat s. h. aq.; v.s. alc., eth.; e.s. HCl. — Gives orange picrate. — W. benzaldehyde at 100° gives benzylidenemethylketole, m.p. 246–7°, lft. fr. acetone (Ann., 242, 272–377).
1602	60–1	2,4-Dinitrotoluene-naphthalene , $(\text{NO}_2)_2\text{C}_7\text{H}_4\text{C}_{10}\text{H}_8.$ — Nitro comp.‡
1603	59–61	m-Methylhydrazobenzene , $\text{Me.C}_6\text{H}_4\text{NH.NH.Ph.}$ — \textcircled{P} Should reduce Tollen's reagt. (T. 2.30). — Pr. fr. lgr.; v.s. alc.
1604	60–1	Acetaldehydediphenylhydrazone , Me.CH:N.NPh. — \textcircled{P} Prob. gives Ag reduction slowly after hydrolysis in T. 2.17. Prob. gives deep blue color to conc. H_2SO_4 . — Pr. fr. lgr.
1605	60	† Isoamyl Carbamate , $\text{NH}_2\text{CO.C}_6\text{H}_{11}.$ — \textcircled{P} \textcircled{D} Sapon. (w. NaOH) T.* products: NH_3 ; isoamyl alc.; $\text{CO}_2.$ — Odor aromat. Taste burning. — Cryst. s. alc., eth., h. aq. — B.p. 220°.
1606	60	Cyanoformanilide , $\text{CN.CO.NH}_2.$ — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; oxalic ac. (Vol. I). — Tbl. E.s. aq., alc., eth. — Dec. at 120° to HCN & cyanuric ac.
1607	60–1	Benzylacetamide , $\text{Me.CO.NH(Ph.CH}_3).$ — \textcircled{P} \textcircled{D} Sapon. (w. alc. KOH) T.* products: benzylamine (No. 2.1236); acetic ac. — Lft. fr. eth.; e.s. alc., eth. — B.p. a. 300°.
1608	60.5	Ethyl Hippurate , $\text{Ph.CO.NH.CH}_2\text{CO.Et.}$ — \textcircled{P} \textcircled{D} Final sapon. T.* products: glycine (No. 2.2568), benzoic ac. (Vol. I), ethyl alc.
1609	59–60; 57; 61–2	Anisonitrile , $p\text{-MeO.C}_6\text{H}_4\text{CN.}$ — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; anisic ac. (Vol. I). — Ndl. fr. h. aq. Mod. s. h. aq.; v.s. alc., eth. B.p. 253–4°; 256–7°.
1610	60	Pimiliimid , $[\text{CH}(\text{Me},\text{CH}).\text{CH}_2\text{CO.NH.CO}]_2.$ — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; pimiliac ac. — Dist. Ndl. fr. eth. + lgr. E.s. alc., eth.
1611	60–1	p-Cyanoacetophenone , $\text{CN.C}_6\text{H}_4\text{CO.Me.}$ — \textcircled{P} \textcircled{D} Sapon. (w. alc. KOH) T.* products: NH_3 ; p-acetylbenzoic ac. — Ndl. fr. dil. alc. E.s. alc., eth. — Gives oxime, m.p. 160°.
1612	61	Nitroerythrite , $\text{NO}_2\text{CH}_2[\text{CH}(\text{NO}_2)]_2\text{CH}_2\text{NO}_2.$ — \textcircled{P} Should give blue color in T. 2.15. — Lft. fr. alc. I. c. aq. Explodes w. violence on percussion!
1613	61	3,4-Dinitrotoluene , $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me.}$ — \textcircled{P} Nitro comp.‡ — Ndl. fr. CS_2 . 100 pt. c. CS_2 dis. 2.19 pt.
1614	61	v-Nitrotetramethylbenzene , Nitroprennitole , $\text{NO}_2\text{C}_6\text{H}_4\text{Me}_4.$ — \textcircled{P} Nitro comp.‡ Cryst. e.s. alc., eth., lgr. B.p. 295° d (th.i.).
1615	61; 55; 67	Isobutyl Carbamate , $\text{NH}_2\text{CO.C}_6\text{H}_5.$ — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; isobutyl alc.; $\text{CO}_2.$ — Cryst. alm. i. aq. Rotates on aq. like camphor. B.p. 206°.
1616	59; 63	Methylbenzoylanilide , $\text{Me}(\text{PhCO})\text{N.Ph.}$ — \textcircled{P} \textcircled{D} Sapon. T.* products: methylaniline (No. 2.1249); benzoic ac. (Vol. I). — Pr. fr. lgr. I. aq.; e.s. alc. — B.p. 315–30°.
1617	61	Diethyl Acetophenylglycine-o-carbonate , $\text{Ph}(\text{CO.Et})_2(\text{C}_6\text{H}_4\text{O})\text{N.CH}_2\text{CO.Et.}$ — \textcircled{P} Heated w. fuming H_2SO_4 at 100° gives blue sol. (indigosulphonic ac.)
1617-I	60–2	Phenyl-α-naphthylamine , $\text{C}_{10}\text{H}_7\text{NHPH.}$ — \textcircled{P} Chlf. sol. fluor. blue. Sol. in conc. H_2SO_4 colored blue by nitrous ac. — Pr. or lft., i. dil. ac.; e.s. alc., eth., bz., chlf. — Nitroso deriv. yel.-red, m.p. 92° (Ber., 20, 1247).
1618	62	1-Phenylpyrrole , $\text{C}_6\text{H}_5\text{N.Ph.}$ — \textcircled{P} Odor camphorous. — Pearly scales. Sbl. B.p. 234°. Vol. w. st. I. aq.; s. alc., eth., chlf., bz.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1619	62	aa-Ethylphenylurea, $\text{NH}_2\text{CO.NEtPh}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: NH_2 ; ethylaniline (No. 2.1270); CO_2 . — Silvery lft. V.s. alc.
1620	62	Undecyl Phenylcarbamate, $\text{Ph.NH.CO}_2\text{C}_{11}\text{H}_{22}$. — \textcircled{P} Prob. gives carbylamine odor in T. 2.12. — Ndl. I. aq. — \textcircled{O} Sapn. T.* products: aniline; undecylic alc. (Vol. I); CO_2 .
1621	62	o-Formotoluide, $\text{Me.C}_6\text{H}_4\text{NH(CHO)}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: o-toluidine (No. 2.1262); formic ac. (Vol. I). — B.p. 288°. Tbl. V.s. alc.
1622	62-3	Stearoneoxime, $(\text{C}_{17}\text{H}_{33})_2\text{C:NOH}$. — Oxime. §
—	63d.	1-Nitromesitylene (iso form). — Cf. No. 2.23-1.
1624	63	2,3-Dinitrotoluene, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Me}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. lgr.
1625	63	Amylpseudonitrol, $\text{Et}_2\text{C}(\text{NO})(\text{NO}_2)$. — \textcircled{P} Sol. in chlf. blue! — Tbl. fr. eth.
1626	62-4	n-Nonyl Phenylcarbamate, $\text{Ph.NH.CO}_2\text{C}_9\text{H}_{19}$. — \textcircled{P} Should give carbylamine odor in T. 2.12. — Cryst. fr. alc. I. aq. — \textcircled{O} Sapn. T.* products: nonyl alc. (Vol. I); CO_2 .
1627	63	ab-Phenylnonylurea, $\text{Ph.NH.CO.NH.C}_9\text{H}_{19}$. — \textcircled{P} Should give carbylamine odor in T. 2.12. — Pr. fr. dil. alc. I. aq.; e.s. alc., eth., bz. — \textcircled{O} Sapn. T.* products: aniline; nonylamine; CO_2 .
1628	64 u.c.	Butyl 3,5-Dinitrobenzoate, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{CO}_2[\text{Me}(\text{CH}_2)_3]$. — \textcircled{P} Nitro comp.‡ — Pearly cryst. fr. h. dil. alc.
1629	64	Methyl-3-nitro-p-acet-toluide, $\text{Me.C}_6\text{H}_4(\text{NO}_2)\text{NMe}(\text{C}_2\text{H}_5\text{O})$. — \textcircled{P} \textcircled{O} Sapn. T.* products: methyl-3-nitro-p-toluidine; acetic ac. — B.Pk (T. 2.23), yel. cryst. ppt., m.p. 210-2° d.
1630	64-5	Pr-1-Methyl-3-phenylindole, $[\text{C}_6\text{H}_4\text{NMe.CH:CPh}]$. — \textcircled{P} Pine splinter pyrrole T. 2.24 gives red-violet color. — Cryst. fr. lgr. E.s. alc., eth., bz.; r.d.s. c. lgr. — B.Pk, m.p. 90°.
—	63; 65	Ethylenediphenyldiamine. — Cf. No. 2.644.
1632	64.5; 66	1,3,4-Xylylhydroxylamine, $\text{Me}_2\text{C}_6\text{H}_3\text{NH(OH)}$. — \textcircled{P} Prob. reduces Tollen's reagt. (T. 2.30). — Lft. fr. bz. + lgr. Air bubbled through aq. suspension gives azoxyxylene.
1633	65	Dinitro-p-dipropylbenzene, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Pr}_2$. — \textcircled{P} Nitro comp.‡ — I. aq. Tbl. fr. alc.
1634	65	Dypnoneoxime, $\text{Me.CPh:CH.C(:NOH).Ph}$. — Oxime. § — V.s. alc., eth., bz.
1635	65.5	Acetone- β -naphthylhydrazone, $\text{Me}_2\text{C:N.NH.C}_{10}\text{H}_7$. — Pale yellowish pr. fr. lgr. E.s. alc., eth.
1636	65.5	m-Acettoluide, $\text{Me.C}_6\text{H}_4\text{NH(C}_2\text{H}_5\text{O)}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: m-toluidine (No. 2.1265); acetic ac. — Cryst. 100 pt. aq. at 13° dis. 0.44 pt. B.p. 303°.
1637	65	Butylphthalimide, $[\text{NBu.CO.C}_6\text{H}_4\text{CO}]$. — \textcircled{P} \textcircled{O} Sapn. T.* products: butylamine (No. 2.1081); phthalic ac. (Vol. I). — Thin lft. fr. dil. alc. B.p. 312° c.
1638	65	Ethyl-glycolate Phenylcarbamate, $\text{Ph.NH.CO.CH}_2\text{CO}_2\text{Et}$. — \textcircled{P} \textcircled{O} Sapn. T.* products: aniline; glycollic ac. (Vol. I); ethyl alc. — Pr. fr. alc. + pet. eth. S. alc., eth., h. aq.
1639	66	2,6-Dinitrotoluene, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Me}$. — \textcircled{P} Nitro comp.‡ — Ndl. s. alc. Reduction by ammon. sulphide gives 6-nitro-o-toluidine (No. 2.3018).
1640	66 u.c.	† β -Naphthonitrile, $\text{C}_{10}\text{H}_7\text{CN}$. — \textcircled{P} \textcircled{O} Sapn. T. 2.26-c, w. alc. KOH, using 0.5 g. nitrile & boiling for 1 hr., gives β -naphthoic ac. in colorless cryst. (if boneblackened), m.p. 182° u.c. (185.7° c.). — Lft. fr. c. lgr. I. aq.; s. alc., eth. B.p. 304-5° c.; 306.5° (th.i.). Odor strongly aromatic.
1641	66	Propylphthalimide, $[\text{NPr.CO.C}_6\text{H}_4\text{CO}]$. — \textcircled{P} \textcircled{O} Sapn. T.* products: propylamine (No. 2.1067); phthalic ac. (Vol. I). — Cryst. fr. alc. B.p. 297° c.
1642	66.5	Methylsuccinimide, $[\text{NMe.CO.CH}_2\text{CH}_2\text{CO}]$. — \textcircled{P} \textcircled{O} Sapn. T.* products: methylamine; succinic ac. (Vol. I). — Tbl. fr. acetone. B.p. 234°.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I643	66	Pyrotartarimide, $[NH.CO.CH_2.CHMe.CO]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; pyrotartaric ac. (C ₄ H ₄ O ₄ , Vol. I). — 6-sided tbl. S. aq. alc., eth.
I644	66	2-Ethyl-3-methylindole, Et,Me,C ₈ H ₇ N. — B.p. 185° (35 mm.). — Picrate (T. 2.23), m.p. 150–1°.
I645	67.5–8	Nitrosobenzene, C ₆ H ₅ NO. — \textcircled{P} Melts to emerald-green liq.! Sol. in bz. grass-green! Odor pungent! — Colorless monoclin. cryst. fr. eth. Mod. s. organic solvents. V. volatile. — Heated w. aniline + ac. ac. gives azobenzene (No. 2.2935).
I646	67–8	N-Ethylcarbazole, C ₁₁ H ₈ N.Et. — Lft. fr. eth. V.s. eth., h. alc. — \textcircled{D} B.Pk (T. 2.23), red ndl., e.s. alc., m.p. 97°.
I647	68	ab-Dimethyl-b-acetylphenylhydrazine, Me,(C ₂ H ₅ O)N.NMePh. — Cryst. fr. lgr. Dist. undecd. “V. stable towards alk.” — Protracted boiling w. 20% HCl gives methylaniline (No. 2.1249).
I648	69	o-Nitrophenacetole, NO ₂ .C ₆ H ₄ OCH ₂ CO.Me. — [D.R.P. 97,242.] — \textcircled{P} Nitro comp.‡ — Nearly colorless ndl. fr. aq. S. h. aq., alc.; e.s. eth., bz. Not vol. w. st. — Oxime, m.p. 102° (Ber., 30, 1635). — Semicarbazone, m.p. 178°.
I649	69	syn-o-Tolylphenylketoxime, Me.C ₆ H ₄ C(:NOH).Ph. — Oxime. §
I650	69	N-Acetylcarbazole, C ₁₁ H ₈ N.C ₂ H ₅ O. — Fine ndl. D.s. h. aq.; v.s. alc., eth., bz. — Gives a yellowish red picrate (T. 2.23).
I650-I	69–5.70	Pyrazole, $[NH.N:CH.CH:CH]$. — B.p. 186–8°. Ndl. fr. eth. E.s. c. aq., alc., eth., bz. Odor, faint, pyridine-like. Taste, unpleasant. $k_B \times 10^4 = 3$. Aq. sol. reacts neutral.” — Ammon. AgNO ₃ gives ppt. of AgC ₆ H ₅ N ₂ fr. aq. sol. HgCl ₂ gives white ppt. fr. aq. sol. — B.Pk, ppt. of fine yell. ndl. fr. eth. sol., m.p. 159–60°.
I651	70.7 u.c.	† 2,4-Dinitrotoluene, (NO ₂) ₂ C ₆ H ₄ Me. — \textcircled{P} Nitro comp.‡ — Odorless ndl. fr. alc. Alm. i. c. aq.; d.s. c. alc.; e.s. eth. Taste of c. sat. aq. sol., bitter (No. 2 of T. 2.29). — \textcircled{D} Place in 6" t.t. 0.2 g. substance, 2 cc. HCl (sp. gr. 1.20), & 1.5 g. granulated tin. Heat sufficiently to liquefy nitro comp. & maintain moderate action, adding more tin, if necessary, until all oil drops disappear. Cool. Add 10 cc. KOH sol. (1:2). Shake & cool. Shake w. 10 cc. eth. Pipe off eth. sol. & evap. on water-bath. Dis. residue in 15 cc. h. lgr. Filter hot. Allow filtrate to cool slowly. Wash separated cryst. w. 5 cc. c. lgr. Dry 15 min. on tile at abt. 75°. The product, p-toluylenediamine, is obtained in long sl. colored ndl., m.p. 99° u.c.
I652	70–1	p-Aacetocumeneoxime, Me.C(:NOH).C ₆ H ₄ CHMe ₂ . — \textcircled{P} Oxime. § — 4-sided tbl. fr. lgr.
I653	70	p-Phenylaminophenol, Ph.NH.C ₆ H ₄ OH. — Lft. V.d.s. aq.; e.s. dil. alk. or mineral ac.! E.s. alc., eth., chlf. — B.p. 330°. — \textcircled{D} Add 1 mol. NaNO ₂ sol. (1:200), cooling w. ice, to sol. of 1 mol. substance in HCl. The nitroso deriv. obtained cryst. in yell. lft. fr. dil. HCl, m.p. 95° d.
I654	70	Melissonitrile, C ₁₁ H ₈ CN. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; melissic ac. (Vol. I).
I655	70–1	Allylphthalimide, $[N(C_6H_5).CO.C_6H_4.CO]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: allylamine (No. 2.1068); phthalic ac. (Vol. I.). — Tbl. B.p. 295°.
I656	70	ab-Pseudohexylphenylurea, Et,CH.CH ₂ NH.CO.NH.Ph. — \textcircled{P} Should give strong carbylamine odor in T. 2.12. — Ndl. fr. alc. I. aq. — \textcircled{D} Sapn. T.* products: aniline; pseudohexyl alc. (b.p. 139–43°).
I657	70	Phenyl o-Aminobenzoate, NH ₂ .C ₆ H ₄ CO.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: o-aminobenzoic ac. (No. 2.148); phenol (Vol. I).
I658	70–1	CEnanthanilide, C ₁₁ H ₈ .CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; enanthic ac. (Vol. I.). — Lust. ndl. fr. lgr. E.s. alc., eth.
I659	71	Dinitro-o-xylene, (NO ₂) ₂ C ₆ H ₄ Me. — \textcircled{P} Nitro comp.‡ — Lust. ndl. D.s. c. alc.
I660	71	Acetone- α -naphthylhydrazone, Me,C:N.NH.C ₁₀ H ₇ . — \textcircled{P} Prob. gives hydrazone T. 2.17-a. — Cryst. fr. lgr. V.s. alc., eth., bz.
I661	71	Ethylbenzamide, EtNH.CO.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylamine (No. 2.1062); benzoic ac. (Vol. I.). — Lust. ndl. d.s. h. aq. B.p. abt. 290°.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1662	72-3d.	s-Diethylpropylpseudonitrole, Et.CH ₂ (NO)(NO ₂).Et. — ⊕ Melts to blue liq. — Rhombohedra d.s. eth.
1663	72-2.5	o-Nitrosotoluene, Me.C ₆ H ₄ .NO. — ⊕ Melts to green liq. & gives green sol. in chlf. — Lust. cryst. E.s. eth., alc.; v.s. chlf. V. vol. w. st.
1664	72-3	Methyl o-Nitrocinnamate, NO ₂ .C ₆ H ₄ .CH:CH.CO.Me. — ⊕ Nitro comp.‡ — Ndl. fr. h. alc. — ⊖ Sapn. T.* products: o-nitrocinnamic ac. (No. 2.437); methyl alc.
1665	72-3	Trinitro-1-methyl-3-isopropylbenzene, (NO ₂) ₂ .C ₆ H.Me.(C ₂ H ₅). — Yellowish white lft. fr. lgr. V.s. alc., eth.
1666	72	5-Nitroquinoline, NO ₂ .C ₆ H ₄ N. — ⊕ Nitro comp.‡ — Lust. ndl. fr. h. aq. Loses cryst. aq. over H ₂ SO ₄ . Sbl. in ndl. D.s. h. aq. — B.MeI (T. 2.37), large dark red pr. fr. aq., m.p. 215° d.
1667	72	Ethyl Diphenylcarbamate, Ph.N.CO.Et. — ⊕ ⊖ B.p. a. 360°. — ⊖ 6 Br ₂ added to ac. ac. sol. gives hexabromo deriv., brownish ndl. fr. ac. ac., m.p. 184°.
1668	72-3	Camphonitrite, C ₁₀ H ₁₇ .CN. — ⊕ ⊖ Sapn. T.* products: NH ₃ ; campholic ac., (C ₁₀ H ₁₆ O ₂ , Vol. I). — E.s. alc. B.p. 217-9°.
1669	72	Pentadecyl Phenylcarbamate, Ph.NH.CO.C ₁₅ H ₃₁ . — ⊕ ⊖ Sapn. T.* products: aniline; pentadecyl alc. (C ₁₅ H ₃₁ O, Vol I); CO ₂ . — Lft. i. aq.
1670	72	d-Carvoneoxime, C ₁₀ H ₁₄ :NOH. — ⊕ May give T. 2.17-a. — Monoclin. pr. fr. alc. [α] _D = +40.9° in abs. alc. — Benzoate (fr. benzoyl chloride), ndl. fr. pet.-eth., m.p. 96°.
1671	72-3	Homoantipyrine, ¹ NPh.NEt.CMe:CH.CO ² . — B ₂ H ₆ PtCl ₆ .2H ₂ O (T. 2.14) or-red ndl., ppt., m.p. 180°.
1672	73-5 <i>u.c.</i>	Propyl 3,5-Dinitrobenzoate, (NO ₂) ₂ .C ₆ H ₄ .CO ₂ .CH ₂ .CH ₂ .Me. — ⊕ Nitro comp.‡
1673	73-4	o-Nitrodesoxybenzoin, C ₁₀ H ₁₁ (NO ₂)O. — ⊕ Nitro comp.‡ — Ndl. fr. bz. Mod. s. alc., eth., bz. — Alc. sol. colored blue by KOH!
1674	73.5	o-Nitromethylcinnamarylvinylketone, Me.CO.C ₆ H ₄ .C ₆ H ₄ (NO ₂). — ⊕ Nitro comp.‡ — Ndl. e.s. alc.
1675	73	N-Ethyl-β-naphthindole, ¹ C ₁₀ H ₈ .NET.CH:CH ² . — ⊕ Colors pine splinter violet in T. 2.24-b. — Ndl. fr. alc. V.d.s. aq.; e.s. alc., eth.
1676	73	Behenolanilide, Ph.NH(C ₂₁ H ₃₈ .CO). — ⊕ Prob. gives carbylamine odor in T. 2.12. I. aq.
1677	73	Butyro-p-toluidine, C ₈ H ₇ .CO.NH.C ₆ H ₅ .Me. — ⊕ ⊖ Sapn. T.* products: p-toluidine (No. 2.566); butyric ac. (Vol. I).
1678	73-4	Formyldiphenylamine, Ph.N(CHO).Ph. — ⊕ ⊖ Sapn. T.* products: diphenylamine (No. 2.1568); formic ac. (Vol. I). — Orthorhomb. cryst. fr. alc.
1679	74	† o-Nitrobenzyl Alc., NO ₂ .C ₆ H ₄ .CH ₂ OH. — ⊕ Nitro comp.‡ — Ndl. D.s. c. aq.; e.s. alc., eth. Deflagrates when rapidly heated.
1680	74-5	5-Nitro-o-xylene, NO ₂ .C ₆ H ₄ .Me ₂ . — ⊕ Nitro comp.‡ — B.p. 274° (th.i.).
1681	74-5	Isodinitrobenzyl, C ₆ H ₁₂ (NO ₂) ₂ . — ⊕ Nitro comp.‡ — Ndl. s. h. alc. Oxid. by CrO ₃ in gl. ac. ac. sol. gives easily some p-nitrobenzoic ac. (No. 2.425).
1682	74	3,4'-Dimethylhydrazobenzene, Me.C ₆ H ₄ .NH.NH.C ₆ H ₄ .Me. — ⊕ Prob. reduces Tollen's reagt. directly in T. 2.30. — Alm. i. aq. Tbl. fr. lgr.
1683	74	† Lactamide, Me.CH(OH).CO.NH ₂ . — ⊕ ⊖ Sapn. T.* products: NH ₃ ; lactic ac. (Vol. I). — Cryst. mass. Luster pearly. V.s. c. aq.! E.s. alc.
1684	74	as-Diethylurea, NH ₂ .CO.NEt ₂ . — ⊕ ⊖ Sapn. T.* products: diethylamine, (No. 2.1068-1); NH ₃ ; CO ₂ . — Deliq. cryst. V.s. aq., alc., bz.; s. eth.
1685	74; 5	Cinnamicaldehydecyanhydrine, Ph.CH:CH.CO(OH).CN. — Lft. fr. h. CS ₂ . E.s. alc., eth., bz., chh.; v.d.s. lgr. — ⊖ Mix in freezing mixt. 1 g. sub-

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No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		stance & 15 cc. eth. w. enough conc. HCl to form homogenous liq. & allow to stand 2 to 3 days at ordinary temperature. The product, α -hydroxy- β -benzalpropionic ac., cryst. fr. h. aq. in ndl., m.p. 137 $^{\circ}$.
1686	74	N-Methyl- α -quinolone, $^{[C_8H_7NMe.CO.CH:CH]}.$ — Ndl. fr. lgr. B.p. 324 $^{\circ}$ (728 mm.). E.s. alc., chlf.; less s. aq., lgr. "Reacts neutral." — $B.HgCl$, d.s. lust. pyramids, m.p. 189 $^{\circ}$.
1687	75-6	β -Nitrosoisopropylacetone, $Me.CO.CH_2.C(NO).Me.$ — $\textcircled{P} \textcircled{D}$ Melts to blue liq. Gives Liebermann's react. (T. No. 2.18). — Pr. fr. pet.-eth. Odor pungent.
1688	75	Myricyl Cyanide, $C_{10}H_{21}CN.$ — Amorph.
1689	75	Iminoacetonitrile, $NH.(CH_2CN)_2.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* (w. alk.) products: NH ₂ ; iminoacetic ac. (No. 2.401). — Lft. fr. eth. Less s. eth. than alc. or aq.
1690	75.5	Trimethylurea, $NHMe.CO.NMe_2.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: methylamine; dimethylamine (No. 2.1059, 2.1061); CO ₂ . — Cryst. fr. eth. V.s. aq., alc. B.p. 232.5 $^{\circ}$ c.
1691	75	α -Tolylsuccinamide, $^{[CO.NC_6H_5.CO.(CH_2)_2]}.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: α -toluidine (No. 2.1262); succinic ac. (Vol. I). — Cryst. i. aq. B.p. 340 $^{\circ}$.
1692	75	Lactyl- α -toluide, $Me.CH(OH).CO.NH(C_6H_5).$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: α -toluidine (No. 2.1262); lactic ac. (Vol. I). — Cryst. powd. fr. bz. + lgr. S. alc., eth., chlf.; i. lgr.
1693	75	3-Acetmethyl- α -xylene, $Me(C_2H_5O)N.C_6H_4.Me.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products (?): v- α -xylidene (No. 2.1333); acetic ac. "Not saponifiable." — Cryst. fr. lgr. Vol. w. st.
1694	75	3-Methylpentanoyl-p-toluide, $(Et.CHMe.CH_2CO).NH(C_6H_5).$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: β -methylethyl-propionic ac. (C ₆ H ₁₂ O ₂ , Vol. I); p-toluidine (No. 2.566). — Ndl. i. c. aq.
1695	75-6	3-Methyl-1-phenyl-5-benzoylpyrazolone, $^{[NPh.N:CMc.CH:C(PhCO_2)]}.$ — Ndl. fr. lgr. V.s. alc. — Not sapd. by aq. KOH; but gives sodium benzoate & methylphenylpyrazolone w. sodium ethylate.
1696	76	Pseudopropynitrole, $Me.C(NO)(NO_2)Me.$ — \textcircled{P} Melts to blue liq.! — Monoclin. cryst. I. aq.
1697	76	as-Dipropylnitro, $NH_2.CO.NPr_2.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: dipropylamine (No. 2.1118); NH ₂ ; CO ₂ . — V.s. aq.! — Picrate, fine yel. ndl., m.p. 135 $^{\circ}$.
1698	abt. 76	Oleamide, $C_{17}H_{33}.CO.NH_2.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: NH ₂ ; oleic ac. (Vol. I).
1699	76.5	5-Formylamino-1,3-xylene, $NH(CHO).C_6H_4.Me.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: s-m-xylidine (No. 2.1321); formic ac. (Vol. I). — Pr. fr. dil. alc.
1700	77-8	2,5-Dinitro-p-cymene, $(NO_2)_2.C_6H_3.Me.(Me_2CH).$ — \textcircled{P} Nitro comp.† — Colorless cryst. I. c. aq.
1701	77-8	Pinacolineoxime, $Me.C(:NOH).CMe_2.$ — \textcircled{P} Oxime.‡ — Ndl. v.d.s. c. aq.; e.s. h. aq. E. vol. w. st.
1702	77.5	Ethyl Carbanil- α -hydroxyisobutyrate, $Ph.NH.CO_2.CMe_2.CO_2Et.$ — \textcircled{P} Should give carbylamine odor in T. 2.12. — \textcircled{D} Short boiling w. 2 mol. NaOH sol. gives cryst. of α -hydroxyisobutyranilide (No. 2.2114).
1703	77-8	Ethyl Acetylcarbamate, $NH(C_2H_5O)CO_2Et.$ — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: NH ₂ ; ethyl alc.; acetic ac., CO ₂ . — Ndl. s. aq., alc., eth.
1704	77-9	Mannite Pentanitrate, $C_6H_9O.(NO_3)_5.$ — \textcircled{P} Should give blue color in diphenylamine react. (T. 2.15). — Detonates w. violence on percussion or heating! Sol. in 500 pt. aq. at 60 $^{\circ}$; v.s. c. alc., eth. Dextrorotatory.
1705	77-9	2,4-Dimethylhydrazobenzene, $Me_2.C_6H_4.NH.NH.Ph.$ — \textcircled{P} Sol. in alc. prob. reduces Tollen's reagt. directly. — I. c. aq.; e.s. alc., eth. Ndl. fr. lgr.
1706	78.5	Trinitrophenetole, Picricacidethylether, $EtO.C_6H_3.(NO_2)_3.$ — \textcircled{P} Nitro comp.‡ — Alm. colorless ndl.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I707	78.5; (70)	Methyl m-Nitrobenzoate, $\text{NO}_2\text{C}_6\text{H}_4\text{CO}_2\text{Me}$. — \textcircled{P} Nitro comp.† — Odorless cryst. D.s. methyl alc. B.p. 279°. — \textcircled{D} Sapn. T.* products: methyl alc.; m-nitrobenzoic ac. (No. 2.139).
I708	78	Brassidanilide, $(\text{C}_6\text{H}_4\text{CO})\text{NHPh}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline (T. 2.1235); brassidiac ac. (Vol. I). — I. aq.
I709	78-9	Cenanth-p-toluide, $(\text{C}_6\text{H}_4\text{CO})\text{NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); cenanthic ac. (Vol. I). — Tbl. fr. dil. alc. Alm. i. c. aq.; e.s. alc.; s. lgr.
I710	78	Diacetamide, $(\text{C}_6\text{H}_4\text{O})_2\text{NH}$. — \textcircled{P} \textcircled{D} Sapn. T.* (KOH sol.) gives NH_3 & acetic ac. easily. — Ndl. fr. eth. V.s. aq.; s. eth. B.p. 223°. Boiling w. aq. gives ac. ac., acetamide (No. 2.1737), & NH_3 . "Reacts neut."
I711	78-9	Triacetamide, $(\text{C}_6\text{H}_4\text{O})_3\text{N}$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. KOH sol.) gives NH_3 & ac. ac. v. easily. — Sm. ndl. fr. eth. Decd. easily to NH_3 & ac. ac. "Reacts neut." May belong to Gen. I.
I712	78-9	Ethylphthalimide, $[\text{C}_6\text{H}_4\text{CO.NEt.CO}]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylamine (No. 2.1062); phthalic ac. (Vol. I). — Ndl. B.p. 285° c.
I713	79	† Propionamide, Et.CO.NH_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; propionic ac. (Vol. I). — Lft. fr. chlf. S. aq. B.p. 213°.
I714	79-80	Campholamide, $\text{C}_6\text{H}_{17}\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; campholic ac. ($\text{C}_{10}\text{H}_{16}\text{O}_2$, Vol. I). — Pr. fr. lgr. V.s. alc.
I715	79	Diethylcarbanilide, $(\text{Et}_2\text{PhN})_2\text{CO}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylaniline (No. 2.1270); CO_2 . — Cryst. fr. alc. I. aq.
I716	79	Ethyl α -Naphthylcarbamate, $\text{C}_{10}\text{H}_7\text{NH.CO.Et}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: α -naphthylamine (No. 2.589); ethyl alc.; CO_2 . — Ndl.
I717	79	α -Acetphenetidide, $\text{EtO.C}_6\text{H}_5\text{NH}(\text{C}_6\text{H}_4\text{O})$. — \textcircled{P} \textcircled{D} Sapn. T.* products: α -phenetidine (No. 2.1342); acetic ac. — Lft. fr. dil. alc. B.p. above 250°.
I718	79	Di-p-tolylamine, $(\text{Me.C}_6\text{H}_4)_2\text{NH}$. — B.p. 330.5°. Ndl. i. aq. Sol. in HCl pptd. by aq. Colored yel. by HNO_2 . — \textcircled{D} Add 1 g. HCl (sp. gr. 1.21) to sol. 1 g. B in alc., then, gradually, 1 g. NaNO_2 in conc. aq. sol. Wash ppt. w. aq. Dry. Recryst. fr. lgr. The nitrosamine formed is obtained in yel. rhomb. cryst., m.p. 101°; 103°. — Acetyl deriv., m.p. 85°; benzoyl deriv., m.p. 125°.
I719	80d.	Nitroso-p-acettoluide, $(\text{NO})(\text{C}_6\text{H}_4\text{O})\text{N.C}_6\text{H}_4\text{Me}$. — \textcircled{P} Prob. gives blue color in T. 2.15. — Ndl. fr. lgr. V.s. alc., eth.; chlf.
I720	80	Tetramethyloxamide, $\text{NM}_2\text{CO.CO.NM}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: dimethylamine (No. 2.1061); oxalic ac. — Ndl. fr. eth. V.s. chlf., bz.; d.s. eth.
I721	80	Isobutyl Phenylcarbamate, $\text{Ph.NH.CO.C}_6\text{H}_5$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; isobutyl alc.; CO_2 . — B.p. 216° sl. d. D.s. aq.; e.s. alc., eth.
I722	80	Ethytriphenylurea, EtPhN.CO.NPr_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylaniline (No. 2.1270); diphenylamine (No. 2.1568); CO_2 . — Ndl. i. aq. Sol. in conc. H_2SO_4 prob. gives blue color w. trace HNO_2 as described for No. 2.1568.
I723	80-1	Methylbenzamide, MeNH.CO.Ph . — \textcircled{P} \textcircled{D} Sapn. T.* products: methylamine (No. 2.1059); benzoic ac. (Vol. I). — Cryst. fr. alc. B.p. 291°.
I724	79-81	m-Cyanobenzaldehyde, $\text{CN.C}_6\text{H}_4\text{CHO}$. — \textcircled{P} \textcircled{D} Sapn. T.* products (w. alk.) prob.: NH_3 ; isophthalic ac. (Vol. I). — B.p. 210°. Ndl. fr. eth. Vol. w. st. E.s. aq., alc., eth.
I725	80.5	1 ² -Cyanoacetophenone, $\text{Ph.CO.CH}_2\text{CN}$. — \textcircled{P} \textcircled{D} Sapn. T. ("w. conc. KOH") products: NH_3 ; acetic & benzoic ac. — Ndl. v.d.s. c. aq., lgr.; e.s. alc., eth.
I726	81	Propion-m-toluide, $\text{Et.CO.NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: m-toluidine (No. 2.1265); propionic ac. — Ndl. fr. eth. V.s. alc.
I727	81-2	† β -Phenylhydroxylamine, Ph.NH.OH . — \textcircled{P} Dis. in conc. H_2SO_4 w. deep blue color! — Unstable, turning brown on exposure. Cryst. s. in 50 pt. c., or 10 pt. h. aq. V.s. alc., eth., chlf.; v.d.s. lgr. — \textcircled{D} Add to sol. of 1 mol. substance in bz., 1 mol. phenyl isocyanate, drop by drop, w. cooling.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. ^o).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		Recryst. ppt. that separates on standing fr. pet.-eth. The product, carbaniplphenylhydroxylamine, is obtained in colorless ndl., m.p. 125°. Its alc. sol. is colored red by FeCl ₃ . (J. prakt. Chem. [2], 56, 84.)
I728	81.5-2	Phenylaminophenol , m-Hydroxydiphenylamine, PhNH.C ₆ H ₄ .OH. — ⊖ Distillate fr. Zn dust gives diphenylamine. Use prelim. T. of No. 2.1568. — Pearly lft. fr. aq. B.p. 340°. D.s. h. aq.; e.s. alc., eth.; less s. lgr.; sol. dil. NaOH or mineral ac., & repptd. fr. ac. sol. by sodium acetate.
I729	81-2	† o-Nitrophenylacetylene, NO ₂ .C ₆ H ₄ .C:CH. — ⊖ Give yel.-white ppt. w. ammon. AgNO ₃ sol., & red ppt. w. ammon. CuCl ₂ sol. (T. No. 1.906), which when dried deflagrate or detonate on heating. — Colorless ndl. fr. dil. alc., yellowing in the air. E.s. h. aq. Vol. w. st. Odor pungent.
I730	78; 84	o-Acetaniside, MeO.C ₆ H ₄ .NH(C ₂ H ₅ O). — ⊖ ⊖ Sapn. T.* products: o-anisidine (No. 2.1332); acetic ac. — B.p. 303-5°. Pearly cryst. E.s. h. aq., gl. ac. ac. 100 pt. alc. dis. 55.28 pt. at 21°.
I731	81-2	† 2,4-(α)-Diphenylpyridine, Ph ₂ C ₆ H ₃ N. — Lust. ndl. fr. dil. alc. B.p. 396-8° c., sl. d. I. aq.; e.s. alc., eth. — B.Pk, dark yel. ndl.; d.s. c. aq., m.p. 169°. — B ₂ H ₆ PtCl ₆ (T. 2.14), fr. alc. in or. ndl. w. alc. of crystn., m.p. 205° (after drying).
I732	79-84	Diacetonitrile (labile form), C ₆ H ₄ N ₂ . — 100 g. bz. dis. 1.22 g. at 16.5°. Boiling bz. sol. finally changes to stable form, m.p. 52°.
I733	82 u.c.	† 2,4,6-Trinitrotoluene, (NO ₂) ₂ .C ₆ H ₃ .Me. — ⊖ S. in c. 10% aq. NaOH sol. w. RO color. — Colorless, odorless cryst. fr. h. alc. Alm. i. c. aq.; v.d.s. c. alc. S. eth. Taste of sat. aq. sol., bitter (No. 3 of T. 2.29). — ⊖ Dis. 0.05 g. substance in 15 cc. boiling lgr. Add 5 drops aniline & gently boil. Filter hot. Wash cryst. w. 2 cc. c. lgr. Dry 5 min. on porous tile at 55°. The aniline salt formed melts at 84.1° u.c.
I734	82	1,3-Dinitro-2,4-xylene, (NO ₂) ₂ .C ₆ H ₃ .Me ₂ . — Nitro comp.† Scaly lft. Ammon. sulphide reduction gives 2-nitro-4-amino-1,3-dimethylbenzene.
I735	82-3	2,3,4-Trimethyl-1-phenylpyrazolone(5), ^[(NMe.CMe:CMe.CO.NPh)] . — ⊖ Neutral FeCl ₃ sol. colors aq. sol. red-violet! — Cryst. e.s. aq., alc., chlf.; d.s. eth., lgr. B.p. 362° c. — Picrate (T. 2.23), ndl., m.p. 94°.
I736	82	m-Hydroxybenzonitrile, HO.C ₆ H ₄ .CN. — ⊖ Taste intensely sweet, but also pungent! Lft. fr. aq. E.s. h. aq.; v.s. alc., eth. — ⊖ Sapn. T.* (w. HCl) products: NH ₃ ; m-hydroxybenzoic ac. (Vol. I).
I737	82-3	† Acetamide, Me.CONH ₂ . — ⊖ ⊖ Sapn. products* by T. 2.26-c: NH ₃ ; acetic ac. — B.p. 222° c. Cryst. fr. h. chlf. in clear odorless ndl. on cooling. Taste faintly bitter. Odor of ordinary st. impure preparation like mouse urine, and m.p. sl. lower than stated. V.s. c. aq., alc.; d.s. eth. — [k _B .10 ⁴ = 0.31.]
I738	82	Methylphenylurea, Me.PhN.CO.NH ₂ . — ⊖ ⊖ Sapn. T.* products: NH ₃ ; methylaniline (No. 2.1249); CO ₂ . — Thin flat striped rhombs of fatty luster fr. bz. & lgr. 100 cc. aq. sol. at 45° contains 74 g.
I739	82	Citronellamide, C ₁₀ H ₁₇ .CO.NH ₂ . — ⊖ ⊖ Sapn. T.* products: NH ₃ ; citronellic ac. (C ₁₀ H ₁₆ O ₂ , Vol. I). — Lft. i. aq.; e.s. alc., eth. Eth. sol. dextro-rotatory.
I740	82-3	Vinylphenylurea, C ₆ H ₅ .NH.CO.NHPh. — ⊖ ⊖ Sapn. T.* products: vinylamine (No. 2.1070); aniline; CO ₂ . — Ndl. of adamantic lust. fr. eth. I. c. aq.
I741	83-3.5 u.c.	Isobutyl 3,5-Dinitrobenzoate, (NO ₂) ₂ .C ₆ H ₄ .CO ₂ C ₆ H ₅ . — Nitro comp.†
I742	83	Methylbenzylketonephenylhydrazone, Me.C(:N.NH.Ph).CH ₂ .Ph. — ⊖ Prob. gives hydrazone react. in T. 2.17-a. — Lft. fr. lgr. E.s. alc., eth., bz.
I743	83	Methyl-p-acettoluide, Me.(C ₆ H ₅ O)N.C ₆ H ₄ .Me. — ⊖ ⊖ Sapn. T.* products: p-toluidine (No. 2.566); acetic ac. — B.p. 283°. Lft. fr. eth.-alc. V.s. h. lgr.
I744	83.5	Ethyllactanilide, Me.CH(OH).CO.NEtPh. — ⊖ ⊖ Sapn. T.* products: ethylaniline (No. 2.1270); lactic ac. (Vol. I). — Pr. fr. aq. E.s. aq., bz.
I745	84	4,6-Dinitro-1,3-dimethyl-5-tert.-butylbenzene, (NO ₂) ₂ .C ₆ H ₃ .Me ₂ (CMe ₃). — ⊖ Nitro comp.† Rhomb. tbl. fr. lgr.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS. — Colorless and Solid.
I746	84	Myristanilide, $(C_{12}H_{27}.CO)NH.Ph.$ — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; myristic ac. (Vol. I). — Ndl. i. aq.; e.s. eth., chlf., bz.
I747	84	Erucamide, $C_{21}H_{44}.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; erucic ac. (Vol. I). — Ndl. fr. alc. D.s. alc.; e.s. eth.
I748	85	2-Nitro-1,3-dimethyl-5-tert.-butylbenzene, $NO_2.C_6H_3.(CMe_3)$. — Nitro comp.‡
I749	85	m-Nitrophenylditolylmethane, $NO_2.C_6H_4.CH(C_6H_4.Me)_2$. — Nitro comp.‡ — Cryst. fr. lgr.
I750	85	Trinitro-1,4-dimethyl-2-propylbenzene, $(NO_2)_3.C_6H_3.Me_2.Pr.$ — \textcircled{P} Nitro comp.‡ — Ndl. i. aq.
I751	85	Aceto-acetanilide, $Me.CO.CH_2.CO.NH.Ph.$ — \textcircled{P} Aq. sol. colored dark violet by $FeCl_3$. — Cryst. fr. lgr. D.s. aq.; e.s. alk. or acid; s. alc., eth. — Reduces ammon. $AgNO_3$. — \textcircled{D} Sapn. T.* products: aniline; ac. ac.; acetone; CO ₂ .
I752	85	Cyclooctanonesemicarbazone, $[CH_2.C(:N.NH.CO.NH_2).(CH_2)]^7$. — Semicarbazone. § — Pr. fr. 60% alc.
I753	85.5	Limonenoneoxime, $C_{10}H_{14}.NOH$. — Oxime. § — After melting & solidifying remelts at 72° & has properties of l-carvoxime.
I754	85	Allylurea, $NH_2.CO.NH.C_2H_5$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; allylamine (No. 2.1068); CO ₂ . — Ndl. V.s. aq., alc.; alm. i. eth.
I755	85	Diethylphenylurea, $NEt_2.CO.NH.Ph.$ — \textcircled{P} \textcircled{D} Sapn. T.* products: diethylamine (No. 2.1068-1); aniline; CO ₂ . — Ndl. E.s. alc.
I756	85	Isopropylphthalimide, $C_6H_5.CO.N(C_3H_7).CO^2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: isopropylamine (No. 2.1063); phthalic ac. (Vol. I). — Ndl. fr. alc. B.p. 286° c.
I757	84.5-5.5	Undecylenamide, $C_{10}H_{19}.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; undecylenic ac. ($C_{11}H_{20}O_2$, Vol. I). — Sol. alc.
I758	85	Phenylanilinoacetonitrile, $(Ph.NH_2)Ph.CH.CN$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. alc. NaOH) products: NH_2 ; aniline; mandelic ac. (Vol. I). — Ndl. fr. dil. alc. I. aq. Gives off HCN on ignition.
I759	85	Acet-p-ditolyamine, $(C_6H_4O).N.(Me.C_6H_4)_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: di-p-tolyamine (No. 2.1718); acetic ac. — I. aq.
I760	86-7	p-Methylhydrazobenzene, $Me.C_6H_4.NH.NH_2.Ph$. — \textcircled{P} Prob. reduces Tolten's reagt. directly on shaking. — Scales. E.s. alc., bz.
I761	86	Dinitromesitylene, $(NO_2)_2.C_6H_3.Me_2$, [Me ₂ = 1:3:5]. — Nitro comp.‡ — Pr. s. h. alc.
I762	86	Benzyl Carbamate, $NH_2.CO_2.CH_2.Ph$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; benzyl alc. (Vol. I); CO ₂ . — Lft. fr. h. aq. D.s. h. aq.; v.s. alc.; mod. s. eth.
—	86-7	Benzoyleccgonine, $C_{18}H_{21}O_4.N_2H_2O$. — Cf. No. 2.2433.
I763	86	2,4-Dimethylpentanoyl-p-toluide, $(Me_2CH.CH_2.CHMe.CO)NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); methylisobutylacetic ac., b.p. 204.5°. — Ndl. fr. lgr. V.s. alc., eth.
—	86	Tetramethylpyrazine. — Cf. No. 2.706.
I764	87	N-Methylcarbazole, $C_{11}H_9N.Me$. — \textcircled{P} Sol. in conc. H ₂ SO ₄ becomes deep green w. drop HNO ₃ . — Lft. fr. alc. D.s. c. alc.; v.s. eth. — \textcircled{D} B.Pk, dark redndl., m.p. 141°, e.s. alc. (T. 2.23).
I765	87	Benzylacetophenoneoxime, $Ph.CH_2.CH_2.C(:NOH).Ph$. — Oxime. § — Ndl. fr. dil. alc. V.s. alc.; e.s. chlf., bz.
I766	87	Iminoformyl Cyanide, $NH:CH.CN$. — \textcircled{P} \textcircled{D} E. sapd. by dil. HCl to NH ₂ & formic ac. — Sbl. B.p. 120-5°. E.s. aq.; d.s. alc., eth.
I767	87	Propion-o-toluide, $(Et.CO).NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: o-toluidine (No. 2.1262); propionic ac. — E.s. alc., eth. — B.p. 298-9° c. (730 mm.).

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I768	87	Phenyl-p-toluidine, Ph.NH.C ₆ H ₄ .Me. — \textcircled{P} Dis. in HNO ₃ w. blue color. — B.p. 334.5° c. — Cryst. i. aq. Less s. aq. than diphenylamine.
I769	87u.c.	† Diphenylbenzylamine, Ph.N.CH ₂ .Ph. — D.s. c. alc. — Heated w. HCl & arsenic ac. gives bronzy mic. cryst. of green dyestuff, "Viridin."
I770	88-9	γ -Trinitrotoluene-naphthalene, (NO ₂) ₃ C ₆ H ₂ (Me).C ₁₀ H ₈ . — \textcircled{P} Nitro comp.‡ — Yellowish white cryst.
I771	88-9	8-Nitroquinoline, NO ₂ .C ₆ H ₄ N. — \textcircled{P} Nitro comp.‡ — Cryst. fr. alc. D.s. c. aq.; mod. s. alc., eth.; e.s. bz. "Salts decd. by aq." — B ₂ H ₆ PtCl ₆ , yel. ppt., cryst. fr. dil. HCl in yel.-red ndl.
I772	88-9	Pr-3-Phenylindole, $^{\text{r}}\text{C}_6\text{H}_4.\text{NH.CH:CHPh}^{\text{l}}$. — \textcircled{P} Gives yel. & then blue-violet color in T. 2.24-b w. pine splinter! — Lft. fr. lgr. E.s. alc., eth., bz.; r.d.s. lgr.
I773	88	Tetramethylsuccinanil, $^{\text{r}}\text{CO.NPh.CO.CMe}_2\text{.CMe}_2^{\text{l}}$. — \textcircled{P} \textcircled{D} Prob. sapn. T.* products: aniline; tetramethylsuccinic ac. (Vol. I). — Ndl. fr. dil. alc. I. aq.; e.s. alc.; d.s. lgr.
I774	89	Amylene Nitrosate, Me ₂ C(NO ₂).C(:NOH).Me. — \textcircled{P} Prob. gives blue color in T. 2.15. — Ndl. fr. bz. i. aq.; d.s. alc. Also cryst. fr. bz. in cubes, m.p. 96-7°.
I775	89	Cyclohexanoneoxime, $^{\text{r}}\text{CH}_2\text{.C(:NOH).}(\text{CH}_2)_4^{\text{l}}$. — \textcircled{P} Oxime.§ — 6-sided pr. fr. lgr. S. aq., alc., eth. — B.p. 206-10°.
I776	89	syn-Furfuraldoxime, C ₄ H ₆ O.CH:NOH. — \textcircled{P} Oxime.§ — Ndl. fr. lgr. D.s. c. aq.; e.s. alc., eth., bz. B.p. 201-8°.
I777	89-90	p-Propionyltolueneoxime, Et.C(:NOH).C ₆ H ₄ .Me. — \textcircled{P} Oxime.§ — Cryst. fr. alc.
I778	89	o-Hydrazophenetole, EtO.C ₆ H ₄ .NH.NH.C ₆ H ₄ .OEt. — \textcircled{P} Prob. reduces Tollen's reagt. directly on shaking. — Air bubbled through h. sol. oxidizes to azophenetole (No. 2.3225).
—	90.0u.c.	† m-Dinitrobenzene. — Cf. No. 2.3016. [Nitro comp.‡ GY-Y, T3.]
I779	90	m-Nitrotriphenylmethane, NO ₂ .C ₆ H ₄ .CHPh ₃ . — \textcircled{P} Nitro comp.‡ — Cryst. fr. lgr.
I780	90	Acenaphthenonephenylhydrazone, C ₁₂ H ₈ :N.NH.Ph. — Hydrazone.§ — Cryst. fr. alc. D.s. c. alc.; e.s. eth., chlf.
I781	90	Furfuraldiphenylhydrazone, C ₆ H ₆ O.CH:N.NPh ₃ . — Hydrazone.§ — Flat ndl. fr. dil. alc. I. aq.; e.s. abs. alc., eth.
I782	90	Brassidamide, C ₁₂ H ₁₄ .CONH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; brassidic ac. (Vol. I). — D.s. alc.
I783	90	Methylisobutylacetamide, Me ₂ C ₆ H ₅ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; methylisobutylacetic ac. (Vol I). — Ndl. fr. lgr. V.s. alc.
I784	90	Butyranilide, (Pr.CO)NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline (T. 2.1235); butyric ac. (Vol. I). — Lft. fr. dil. alc. I. aq.; e.s. alc., eth.
I785	90	α -Hydroxybutyranilide, Me.CH ₂ .CH(OH).CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; α -hydroxybutyric ac. (C ₄ H ₆ O ₃ , Vol. I). — Ndl. fr. eth. + lgr. V.s. alc., eth., chlf.; v.d.s lgr.
I786	90.5	Palmitanilide, (C ₁₆ H ₃₂ .CO)NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; palmitic ac. (Vol. I). — Silky ndl. fr. alc. E.s. alc., eth., bz.
I787	89.5-90u.c.	† Anæsthesine, Ethyl p-Aminobenzoate, NH ₂ .C ₆ H ₄ .CO ₂ Et. — Rhomb. cryst. fr. eth. Alm. i. aq.; s. alc., bz., chlf., dil. acids. S. in 7 pt. eth. Odorless. Taste, sl. bitter. — \textcircled{P} Rub together on crucible cover 1 mg. substance & 2 drops conc. H ₂ SO ₄ . Touch w. rod moistened w. conc. HNO ₃ . A Y-GY color appears. Mix w. 0.3 cc. aq. Add 0.5 cc. 10% NaOH sol. Color changes to O. \textcircled{D} Dis. 0.05 g. substance in 3 cc. aq. + 1 drop HCl (sp. gr. 1.12). Add 3 cc. c. sat. aq. picric ac. sol. Filter off slender ndl. which slowly separate. Wash w. 1 cc. aq. Recryst. fr. 8 cc. boiling aq. Wash w. $\frac{1}{2}$ cc. aq. Dry on tile 15 min. at 100°. The resulting picrate is obtained in hair-like GY-Y, T2 ndl., m.p. 128.5-9.5° u.c. (130.2-131.2° c.).

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1788	90	<i>o</i> -Tolyl-p-aminophenol, $\text{Me.C}_6\text{H}_4.\text{NH.C}_6\text{H}_4.\text{OH}$. — Lft. fr. lgr. + bz. B.p. 366–8° c. E.s. alc., eth., dil. KOH(!); d.s. lgr. — \textcircled{P} Ignition w. Zn dust gives acridine (No. 2.3102).
1789	90; 91; 92	p-Dimethylaminobenzophenone, $\text{NMe}_2.\text{C}_6\text{H}_4.\text{CO.Ph}$. — Lft. fr. alc. I. aq.; s.c. alc.; v.s. h. alc., eth.
1790	90	Methylpyrrolketone, $[\text{NH.C}(\text{CO.Me}): \text{CH.CH}: \text{CH}]$. — Ndl. B.p. 220°. — $\text{AgC}_6\text{H}_4\text{NO}$, cryst. ppt. fr. aq. sol. w. AgNO_3 , & little ammon.
1791	91	Trinitroisobutyl, $\text{C}_6\text{H}_{14}(\text{NO}_2)_3$. — \textcircled{P} Nitro comp.‡ — Tbl. fr. bz. + pet-eth. Not s. in dil. NaOH.
1792	88; 95	Dinitro-p-tert.butyltoluene, $(\text{NO}_2)_2.\text{C}_6\text{H}_4.\text{Me}(\text{CMe}_3)$. — \textcircled{P} Nitro comp.‡ w. v. faintly musky odor. — Cryst. fr. dil. alc.
1793	91	4-Methylbenzalacetophenoneoxime, $\text{Me.C}_6\text{H}_4.\text{CH}: \text{CH.C}(\text{:NOH})\text{Ph}$. — \textcircled{P} Oxime. § — Ndl. fr. alc.
1794	91	Campholanilide, $(\text{C}_9\text{H}_{17}.\text{CO})\text{NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; campholic ac. ($\text{C}_{10}\text{H}_{18}\text{O}_2$, Vol. I). — Ndl. fr. alc.
1795	91	Ethylcarbanilide, $\text{Et}.\text{PhN.CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; ethylaniline (No. 2.1270); CO_2 . — Clear pr. fr. alc. I. aq. Dec. to ethylaniline & phenylisocyanate on distn.
1796	91	1-Methylcyclohexyl Phenylcarbamate, $\text{Ph.NH.CO}_2.\text{C}_7\text{H}_{15}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; cyclohexanol ($\text{C}_6\text{H}_{12}\text{O}$, Vol. I). — Alm. i. aq., lgr.; e.s. alc.
1797	91	p-Tolyl-m-aminophenol, $\text{Me.C}_6\text{H}_4.\text{NH.C}_6\text{H}_4.\text{OH}$. — \textcircled{P} Sol. in conc. H_2SO_4 colored blue by trace HNO_2 . — B.p. 350° c. Ndl. fr. bz. + lgr. D.s. h. aq.; e.s. alc., eth., bz. — Nitroso deriv. (T. 2.36), yel. ndl. fr. dil. alc., m.p. 105° d.
1798	92–3	3,5-Dinitrotoluene, $(\text{NO}_2)_2.\text{C}_6\text{H}_4.\text{Me}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. h. aq. D.s. aq.; s.c. alc.; e.s. eth.; v.s. bz. Vol. w. st.
1799	92	Trinitro-p-ethyltoluene, $(\text{NO}_2)_3.\text{C}_6\text{H}_4.\text{Me.Et}$. — \textcircled{P} Nitro comp.‡ — Short pr. fr. alc. D.s. c. alc.
1800	92–3 u.c.	Ethyl 3,5-Dinitrobenzoate, $(\text{NO}_2)_2.\text{C}_6\text{H}_3.\text{CO}_2\text{Et}$. — \textcircled{P} Nitro comp.‡ — Colorless ndl. fr. h. dil. alc. 100 pt. 90% alc. dis. 0.592 pt. at 13°.
1801	92	Dinitro-1-methyl-3-tert.butylbenzene, $(\text{NO}_2)_2.\text{C}_6\text{H}_4.\text{Me}(\text{CMe}_3)$. — Nitro comp.‡ — Ndl. i. c. aq.
1802	92	Methyl 4-Nitro-3-hydroxybenzoate, $(\text{NO}_2)(\text{HO}).\text{C}_6\text{H}_4.\text{CO}_2\text{Me}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. — \textcircled{D} Sapn. T.* products: methyl alc.; m-hydroxybenzoic ac. (Vol. I).
—	92 u.c.	Acet-o-nitroanilide. — Cf. No. 2.3021. (Yellowish.)
1803	92–3	d,L-Carboxime, $\text{C}_{10}\text{H}_{14}:\text{NOH}$. — Oxime. § — Monoclin. cryst.
1804	92–3	Methylacetylphenylhydrazine, $\text{Me}(\text{C}_6\text{H}_5\text{O})\text{N.NMe.Ph}$. — S. h. aq.; e.s. alc., bz.; d.s. eth. Does not reduce boiling Fehling's sol. directly, but is saponifiable to methylphenylhydrazine, No. 2.1336, which reduces Tollen's reagt. slowly.
1805	91; 87–8	m-Tolualdehydephenylhydrazone, $\text{Me.C}_6\text{H}_4.\text{CH}: \text{N.NH.Ph}$. — \textcircled{P} Hydrzone. § — Pr. fr. lgr. E.s. alc., eth., chlf.
1806	92	Ethylurea, EtNH.CO.NH_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; ethylamine; CO_2 . — Pr. V.s. aq., c. alc.
1807	89–91; 92–3.5	Isoamylurea, $(\text{C}_6\text{H}_{11})\text{NH.CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; isoamylamine (No. 2.1100); CO_2 . — D.s. aq.
1808	91–3	Ricinelauidamide, $\text{C}_{17}\text{H}_{34}.\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; ricinelidic ac.
1809	93	Hydrazineacetic Ac., Glycollic-acid-hydrazide, $\text{NH}_2.\text{NH.CO.CH}_2.\text{OH}$. — \textcircled{P} Reduces Tollen's reagt. in T. 2.30. — Glassy tbl. fr. h. alc. E.s. c. aq., h. alc.; i. eth. Tastes sweet. "Reacts neutral." Warmed w. alk. or ac. gives hydrazine & glycollic ac. (Vol. I).

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
I810	93-4	Methylphenylhydroxyurea, Ph.NH.CO.N.Me(OH) . — \textcircled{P} FeCl_3 , colors alc. sol. deep blue-violet. Tbl. fr. lgr. E.s. alc. — Boiled w. NaOH sol. gives aniline, β -methylhydroxylamine & carbanilide.
I811	93	4,6-Dinitro-m-xylene, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} Nitro comp.‡ — Pr. Alm. i. c. aq.
I812	93	(β)-2,3-Dinitro-p-xylene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Nitro comp.‡ — Monoclin. cryst. fr. toluene. Alm. i. c. aq.
I813	93	p-Nitrotriphenylmethane, $\text{NO}_2\text{C}_6\text{H}_4\text{CH.Ph}_3$. — \textcircled{P} Nitro comp.‡ Lft. fr. alc.
I814	93	p-Nitrobenzyl Alc., $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{OH}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. aq. D.s. c. aq.; e.s. h. aq.
I815	93.5	o-Nitro-p'-nitrobiphenyl, $\text{NO}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NO}_2$. — Nitro comp.‡ — Monoclin. cryst. e.s. h. alc.
I816	93	Methyl Acetylicarbamate, $\text{NH}(\text{C}_2\text{H}_5\text{O})\text{CO}_2\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; methyl alc.; ac. ac.; CO_2 . — Tbl. fr. chlf.
I817	93	Isobutylphthalimide, $^{\text{C}}\text{C}_6\text{H}_4\text{CO.N(C}_6\text{H}_5\text{)}\text{CO}^{\text{C}}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: isobutylamine (No. 2.1078); phthalic ac. (Vol. I). — Lft. fr. CS_2 . B.p. 293-5°.
I818	93-4	Elaidamide, $\text{C}_{17}\text{H}_{34}\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; elaidic ac., (Vol. I). — D.s. alc.
I819	93.6	Stearanilide, $(\text{C}_{17}\text{H}_{34}\text{CO})\text{NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; stearic ac. (Vol. I). — Fine ndl. fr. alc. I. aq.
I820	94	p-Tolylhydroxylamine, $\text{Me.C}_6\text{H}_4\text{NH(OH)}$. — \textcircled{P} Reduces Tollen's reagt. (T. 2.30). — Lft. fr. bz. S. in abt. 100 pt. c., or 2 pt. h. aq.; alm. i. c. lgr.; e.s. alc., eth., chlf. Dec. slowly in aq. sol.
I821	94	Ethyl α -Oximinopropionate, Me.C(:NOH).CO.Et . — \textcircled{P} Oxime.§ — Ndl. E.s. alc., eth.
—	94-5	3-Nitro-p-acettoluide. — Cf. No. 2.3028 (slightly colored).
I823	94-5	m-Nitrobenzophenone, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.Ph}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc.
I824	94-5	Acetyl methyl-m-nitroaniline, $(\text{C}_2\text{H}_5\text{O})\text{MeN.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Nitro comp.‡ — Ndl. — \textcircled{D} Sapn. T.* products: m-nitromethylaniline (No. 2.2933); acetic ac.
I825	94	ab-Pentadecylphenylurea, $(\text{C}_{15}\text{H}_{30})\text{NH.CO.NHPh}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; pentadecylamine; CO_2 . — I. aq.
I826	94c.	m-Toluamide, $\text{Me.C}_6\text{H}_4\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; m-toluic ac. ($\text{C}_6\text{H}_5\text{O}_2$, Vol. I).
I827	94-5	Piperonylonitrile, $\text{m,p-CH}_2\text{:O}_2\text{:C}_6\text{H}_4\text{CN}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; piperonylic ac. or its decn. products (Vol. I). — Ndl. fr. aq. V.s. alc., eth., chlf., bz.
I828	94-5	o-Tolyl- α -naphthylamine, $\text{Me.C}_6\text{H}_4\text{NH.C}_10\text{H}_7$. — Flat ndl. fr. lgr. E.s. alc., eth., bz.
I829	94	Diphenylfurazan, $^{\text{C}}\text{C}_6\text{H}_4\text{N.O.N:CPh}^{\text{C}}$. — Feathery cryst. fr. alc. S. c. alc.; e.s. eth. Not easily attacked by boiling w. alk. or ac. sol. — Dinitro deriv., cryst. fr. gl. ac. ac., m.p. 218-20° (Ann., 264, 182).
I830	95	† Skatole, 3-Methylindole, $^{\text{C}}\text{C}_6\text{H}_4\text{NH.CH:CM}^{\text{C}}$. — [Fr. excrement, various plants, & protein putrefaction.] — Shining lft. fr. lgr. B.p. 265-6° (th.i.). Vol. w. st. 1000 pt. c. aq. dis. 0.45 g. E.s. alc., eth., chlf., bz. Odor, disagreeable, fecal, persistent!
		† (1) Gives red coloration in pine splinter T. 2.24. — (2) [Legal's T.] Dis. abt. 1 mg. sodium nitroprusside in sol. of 1 mg. skatole in 1 cc. aq. Add 1 drop 10% aq. NaOH sol. Dilute the OY sol. w. 3 cc. aq. Add 2 cc. gl. ac. ac. Boil 1 to 2 min., & set aside for 5 min. A R-VR color gradually develops. — (3) 1 mg. skatole heated w. 1 cc. conc. H_2SO_4 , gives R-VR coloration. — Does not give the nitrosoindole reaction numbered "(4)" among prelim. tests under No. 2.1546. — [Cf. Herter, J. Biol. Chem., 1908, 267-271 & 1908, 101-109 for separation & determination in presence of indole; and Rosenthaler, p. 580, for bibliography of other tests.]

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		<p>① Suspend 0.05 g. skatole in 5 cc. boiling aq. Add 4 cc. c. sat. picric ac. sol. & heat to boiling. Set aside until cold. Filter. Wash w. 2 cc. c. aq. Dis. in 10 cc. boiling aq. Allow to stand over night. Wash ppt. w. 1 cc. aq. Dry on tile 15 min. at 100°. The picrate obtained, B.Pk, cryst. in lust. OS2 ndl., melts at 168–70° u.c. (171–8° c.) to dark brown liq.</p>
1831	95	m-Nitrophenyl Benzoate, Ph.CO ₂ .C ₆ H ₄ .NO ₂ —② Nitro comp.‡ Cryst. v.s. alc., gl. ac. ac., h. lgr.
1832	95	2,3,3-Trinitro-2-methylpentane, Me.CMe(NO ₂).C(NO ₂).CH ₂ .Me. — ② Nitro comp.‡ — Plates & ndl. fr. lgr. & h. alc.
1833	95	Caproanilide, (C ₆ H ₁₁ .CO)NH.Ph. — ② Sapn. T.* products: aniline; caproic ac., (Vol. I). — Lust. ndl.
1834	95	Methylallactanilide, Me.CH(OH).CO.N.Me.Ph. — ② ③ Sapn. T.* products: methylaniline; lactic ac. (Vol. I). — Tbl. fr. aq. E.s. alc., bz.; v.d.s. lgr.
1835	95	Cenanthamide, Me.(CH ₂) ₆ .CO.NH ₂ . — ② ③ Sapn. T.* products: NH ₂ ; cenanthic ac., (Vol. I). — Mod. s. c. aq. B.p. 250–8°.
1836	95	Methyl- α -acetnaphthalide, Me,(C ₆ H ₅ O).N.C ₁₀ H ₇ . — ② ③ Sapn. T.* products: methyl- α -naphthylamine (No. 2.1443); acetic ac. — Pr. D.s. aq.; e.s. alc., eth.
1837	95	Triethyl(Iso-)Cyanurate, Triethylcarbonimide, C ₃ O ₂ N.Et ₃ . — ② ③ Sapn. T.* (by alk.) products: ethylamine (No. 2.1062); CO ₂ . — B.p. 276°. Pr. Vol. w. st. S. h. aq.; e.s. alc., acids.
1838	95	Diphenyl-m-phenylenediamine, Ph.NH.C ₆ H ₄ .NH.Ph. — Sol. in conc. H ₂ SO ₄ is colored yel.-green & then blue-violet by trace HNO ₃ ! — Flat ndl. I. aq., dil. ac. or alk.; e.s. eth., h. bz.; v.d.s. c. alc., warm lgr. — Gives brown-red nitroso deriv., pr. w. bluish reflections fr. bz., m.p. 153°.
1839	95–6	α -Tolyl- β -naphthylamine, Me.C ₆ H ₄ .NH.C ₁₀ H ₇ . — Lft. fr. lgr. I. aq.; e.s. alc., eth., bz. — Picrate (T. 2.23), red-brown ndl. fr. eth., m.p. 110°.
1840	95	Tetraethyldiaminobenzophenone, CO.[C ₆ H ₄ .NET ₂] ₂ . — Lft. fr. alc.
1841	96	Methyl p-Nitrobenzoate, NO ₂ .C ₆ H ₄ .CO.Me. — ② Nitro comp.‡ — Lft. — ③ Sapn. T.* products: methyl alc., p-nitrobenzoic ac. (No. 2.425).
1842	96	Ethyl 5-Nitrosalicylate, NO ₂ .C ₆ H ₅ (OH)(CO ₂ Et). — ② Nitro comp.‡ — Ndl. fr. alc. E.s. alc., eth. — ③ Sapn. T.* products: ethyl alc.; 5-nitrosalicylic ac. (No. 2.416).
1843	96–7	† 2,4,6-Trinitro- Ψ -butyltoluene, artificial or Baur Musk, (NO ₂) ₃ .C ₆ H ₄ .Me-(Me,C). — ② Odor like musk! Nitro comp.‡ — Sl. yellowish ndl. fr. alc. E.s. alc.
1844	96	Arachidanilide, (C ₁₁ H ₂₂ .CO).NH.Ph. — ② ③ Sapn. T.* products: aniline; arachidic ac. (Vol. I). — I. aq. Ndl. fr. alc.
1845	96	α -Hydroxybutyro- α -naphthalene, Et.CH(OH).CO.NH.C ₁₀ H ₇ . — ② ③ Sapn. T.* products: α -naphthylamine (No. 2.589); α -hydroxybutyric ac. (C ₄ H ₆ O ₃ , Vol. I). — Ndl. fr. alc.
1846	97–8	α -Trinitrotoluene-naphthalene, (NO ₂) ₃ .C ₆ H ₄ .Me.C ₁₀ H ₇ . — ② Nitro comp.‡ — Ndl. d.s. alc.
1847	97	Dimethyl Phenylglycine- α -carbonate, (CO ₂ Me).C ₆ H ₄ .NH.CH ₂ .CO ₂ Me. — ② Sol. in bz. or eth. fluor. blue-violet! — Vol. w. st.
1848	97 u.c.	Furfurolphenylhydrazone, C ₆ H ₅ O.CH:N.NH.Ph. — ② Hydrzone. § — Pearly yellowish lft. fr. h. 50% alc. E.s. alc., eth.; v.d.s lgr.
1849	97–8	Acetophenonediphenylhydrazone, Ph.MeC:N.NPh ₂ . — ② Hydrzone. § — Warty masses fr. alc. I. aq.; e.s. eth., h. alc.
1850	97	Oxyhydrastinine, C ₁₁ H ₁₁ O ₂ N. — [Fr. action of h. conc. KOH on hydrastinine.] Ndl. fr. pet.-eth. V.s. alc., bz. B.p. a. 350°. — B ₂ H ₅ PtCl ₆ (T. 2.14), yel. cryst. e.s. alc., m.p. 160°. [“V. weak base; s. in HCl, but pptd. by aq.”]

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1851	98	a-Ethyl-a-hydroxy-b-phenylurea, Et(OH)N.CO.NHPh. — \textcircled{P} FeCl ₃ colors alc. sol. deep blue-violet! — Cryst. e.s. except in lgr. — Sapn. w. alk. gives methylhydroxylamine (No. 2.564); aniline; CO ₂ .
1852	98-9	(β)-Benzoinoxime, Ph.C(:NOH).CH(OH).Ph. — \textcircled{P} Oxime. § — Ndl. — Eth. sol. satd. w. HCl gas gives in 2 or 3 hr. α -deriv., pr. fr. bz., m.p. 151-2°.
1853	98	Isocarvoneoxime, C ₁₀ H ₁₄ :NOH. — Oxime. § — Cryst. fr. alc. V.s. alc., eth.
1854	98	Citraconanil, C ₈ H ₆ O ₂ .NPh. — \textcircled{P} \textcircled{D} Sapn. T.* (w. alk.) products: aniline; mesaconic ac. (Vol. I). — Ndl. fr. aq. D.s. aq.; e.s. alc., eth., chlf., bz.
1855	98; 108	Capramide, C ₈ H ₁₆ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; capric ac. (Vol. I). — Lft. fr. eth.
1856	98.5	Tridecylamide, C ₁₂ H ₂₅ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; tridecyl ac. (C ₁₂ H ₂₅ O ₂ , Vol. I). — Silky lft. fr. alc. E.s. alc., eth.
1857	98	α -Hydroxyisobutyramide, Me ₂ C(OH).CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; α -hydroxyisobutyric ac. (C ₄ H ₆ O ₂ , Vol. I). — E.s. aq., alc. B.p. 260°.
1858	98-9	m-Cyanoacetophenone, CN.C ₆ H ₅ .CO.Me. — \textcircled{P} \textcircled{D} Sapn. T.* (w. 20% KOH) products: NH ₃ ; m-acetophenonecarboxylic ac. (ndl. fr. h. aq., m.p. 172°). — Ndl. fr. alc.
1859	99; (92-3)	Pelargonamide, C ₈ H ₁₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; pelargonic ac. (Vol. I). — Pearly cryst. mass. Alm. i. c. aq.
1860	99-100	Phenylpropiolamide, Ph.C:CCO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; phenylpropionic ac. (Vol. I).
1861	99	s-Ethylphenylurea, Et.NH.CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; ethylamine (No. 2.1062); CO ₂ . — Ndl. fr. dil. alc.
1862	99	Phenylseptadecylurea, Ph.NH.CO.NH(C ₁₇ H ₃₄). — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; septadecylamine. — Lust. ndl. fr. alc. I. aq.
1863	99	4-Acetamino-o-xylene, (C ₈ H ₈ O)NH.C ₈ H ₈ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: 4-amino-1,2-xylene (No. 2.588); acetic ac. — Pr. fr. dil. alc. V.s. alc.
—	99	Carbazolin. — Cf. No. 2.746-1.
1864	100; 96-8; (92)	p-Cyanobenzaldehyde, CN.C ₆ H ₄ .CHO. — \textcircled{P} Alc. sol. may reduce Tollen's reagt. directly. — Pr. E.s. alc., eth., chlf.
1865	100	Dinitro-o-benzyltoluene, (NO ₂) ₂ .C ₆ H ₁₂ . — \textcircled{P} Nitro comp. ‡ — Ndl. I. aq.
—	100	β -Trinitrotoluene-naphthalene. — Cf. No. 2.3064. — (Yellowish.)
1867	100	Methylpropylketonesemicarbazone, Me.C:(N.NH.CO.NH ₂).CH ₂ .CH ₂ .Me. — \textcircled{P} Semicarbazone. § — Cryst. fr. aq. E.s. alc., h. aq.
1868	100	Caproamide, Me.(CH ₂) ₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; caproic ac. (Vol. I). — Lft. D.s. c. aq.; e.s. alc. B.p. 255°.
1869	100	s-Dimethylurea, MeNH.CO.NHMe. — \textcircled{P} \textcircled{D} Sapn. T.* products: methylamine; CO ₂ . — B.p. 268-73° c.
1870	100	s-Diallylurea, Sinapolin, (C ₆ H ₅)NH.CO.NH(C ₆ H ₅). — \textcircled{P} \textcircled{D} Sapn. T.* products: allylamine (No. 2.1068); CO ₂ . — Lft. E.s. alc., eth., h. aq. Vol. w. st.
1871	100	Succin-o-toluide, C ₈ H ₇ .NH.CO.CH ₂ .CH ₂ .CO.NH.C ₆ H ₅ . — \textcircled{P} \textcircled{D} Sapn. T.* products: o-toluidine (No. 2.1262); succinic ac. (Vol. I). — V.d.s. aq.; s. alc.
1872	100-1	Pr-1-Methyl-2-phenylindole, [C ₆ H ₄ .NMe.CPh:CH]. — \textcircled{P} Gives dark red color to splinter in T. 2.24-b! — Pointed pr. fr. alc. S. eth., chlf., bz., h. aq., lgr. Dist. undecd.
—	100	Triphenylrosaniline. — Cf. No. 2.748-1. (Becomes bluish when washed & dried.)
1873	101	2,5-Methylphenylpyrrole, [MeC:CH.CH:CPH]. — \textcircled{P} Vapors color pine splinter red in T. 2.24-a! — Lust. lft. E.s. alc., eth., chlf., bz. Sbl. w. partial carbonization. Sol. in gl. ac. ac. warmed w. conc. H ₂ SO ₄ , & little isatin gives purple-red sol. Gives dark red picrate decd. by aq.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1874	101-2	$\text{o-Methylhydrazobenzene, } \text{Me.C}_6\text{H}_4.\text{NH.NH.Ph. — } \textcircled{P} \text{ Prob. reduces Tollen's reagt. slowly. — Lft. fr. alc. I. aq.}$
1875	101-2	$\text{o-Nitro-m-acettoluide, } (\text{NO}_2)(\text{Me.CO.NH).C}_6\text{H}_4.\text{Me. — } \textcircled{P} \text{ Nitro comp.‡ — Cryst. fr. alc. — Savn. T.* products: o-nitro-m-toluidine (No. 2.2901); acetic ac.}$
1876	101	$\text{Acetyl diphenylamine, } \text{Ph}_2\text{N(C}_6\text{H}_4\text{O). — } \textcircled{P} \text{ Savn. T.* products: diphenylamine (No. 2.1568); acetic ac. — Tbl. fr. lgr. S. eth. Sbl. in fine ndl.}$
1877	102	$\text{o-Hydrazoanisole, } \text{MeO.C}_6\text{H}_4.\text{NH.NH.C}_6\text{H}_4.\text{OMe. — } \textcircled{P} \text{ Prob. reduces Tollen's reagt. slowly. — Colorless ndl.}$
1878	102	$\text{Dinitrodiisoamyl, } \text{Me}_2\text{C}(\text{NO}_2)_2(\text{CH}_2)_4\text{C}(\text{NO}_2)_2\text{Me}_2. — \textcircled{P} \text{ Nitro comp.‡ — Pr. fr. bz. Sbl. at } 100^\circ. \text{ R.d.s. eth.; e.s. bz.}$
1879	102-3	$\text{Piperonalphenylhydrazone, } \text{CH}_2:\text{O}_2:\text{C}_6\text{H}_5\text{CH: N.NH.Ph. — } \textcircled{P} \text{ Hydrazone.§ — Ndl. fr. dil. alc. E.s. eth., chlf.}$
1880	102	$\text{† Methylacetanilide, } \text{Me}(\text{C}_6\text{H}_5\text{O})\text{N.Ph. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: methylaniline (No. 2.1249); acetic ac. — Pr. fr. alc. B.p. } 245^\circ. \text{ — Convert into 2,4-dinitroaniline (No. 2.3539-1, yel. cryst. fr. dil. alc., m.p. } 174^\circ \text{ u.c.) by boiling 2 hr. w. 100 pt. dil. HNO}_3 \text{ (sp. gr. 1.029).}$
1881	102-2.5	$\text{p-Acetaminoisopropylbenzene, } (\text{C}_6\text{H}_5\text{O})\text{NH.C}_6\text{H}_4.\text{CH.Me}. — \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: p-isopropylaminobenzene (No. 2.1334); acetic ac.}$
1882	102-3	$\text{p-Lactotoluide, } \text{Me.CH(OH).CO.C}_6\text{H}_4.\text{Me. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: p-toluidine (No. 2.566); lactic ac. (Vol. I). — Ndl. fr. h. aq. V.d.s. c. aq.; l. lgr.}$
1883	102	$\text{Myristamide, } \text{C}_{12}\text{H}_{25}.\text{CO.NH}_2. — \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: NH}_2; \text{ myristic ac. (Vol. I). — Scales. E.s. alc. B.p. } 217^\circ \text{ (12 mm.).}$
1884	102	$\text{† Methylurea, } \text{NH}_2.\text{CO.NHMe}. — \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: NH}_2; \text{ MeNH}_2; \text{ CO}_2. — \text{Pr. V.s. aq., alc.}$
1885	103	$\text{Undecylamide, } \text{C}_{11}\text{H}_{23}.\text{CO.NH}_2. — \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: NH}_2; \text{ undecylic ac. (C}_{11}\text{H}_{23}\text{O, Vol. I).}$
1886	103-4	$\text{2,3-Dimethylbutanoyl-p-toluide, } \text{Me.CHMe.CHMe.CO.NH.C}_6\text{H}_4.\text{Me. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: p-toluidine (No. 2.566); methylisopropyl acetic ac. (b.p. } 189-91^\circ).$
1887	abt. 103	$\text{3,4'-Dinitrodiphenylmethane, } (\text{NO}_2\text{C}_6\text{H}_4)_2\text{CH}_2. — \textcircled{P} \text{ Nitro comp.‡ — Silky ndl. fr. alc.}$
1888	103-4	$\text{Hydraltalphenylhydrazone, } \text{CH}_2:\text{O}_2:\text{C}_6\text{H}_5\text{(C}_6\text{H}_4\text{).CH: N.NHPH. — } \textcircled{P} \text{ Hydrazone.§ Ndl.}$
1889	103; (97-9)	$\text{Phenylacetaldoxime, } \text{Ph.CH}_2\text{CH: NOH. — } \textcircled{P} \text{ Oxime.§ — Cryst. fr. eth. E.s. alc., eth.}$
1890	104	$\text{Acetyl-o-tolyldihydrazine, } \text{Me.C}_6\text{H}_4.\text{NH.NH.CO.Me. — Lft. fr. aq. — o-Tolylhydrazine (the product of hydrolysis) reduces Tollen's reagt. easily.}$
—	104	$\text{2,4,5-Trinitrotoluene. — Cf. No. 2.3081 ("yellowish").}$
1891	104-5	$\text{Benzoylmethyl-m-nitroanilide, } (\text{Ph.CO})\text{Me.N.C}_6\text{H}_4.\text{NO}_2. — \textcircled{P} \text{ Nitro comp.‡ — Tbl. Savn. T.* products: m-nitromethylaniline (No. 2.2933); benzoic ac.}$
1892	104	$\text{† Propionanilide, } (\text{Et.CO})\text{NH.Ph. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: aniline; propionic ac. (Vol. I). — Lft. 100 pt. aq. at } 24^\circ \text{ dis. 0.42 pt.; e.s. alc., eth.}$
1893	102.5; 105	$\text{Isobutyranilide, } (\text{Me}_2\text{CH.CO})\text{NH.Ph. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: aniline; isobutyric ac. (Vol. I). — Pr. e.s. h. aq.; e.s. alc., eth.}$
1894	104	$\text{Pyruvanilide, } \text{Me.CO.CO.NH.Ph. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: aniline; pyruvic ac. (Vol. I). — Ndl. Alm. i. c. aq.; d.s. alc.; e.s. chlf.}$
1895	104-5	$\text{Acrylanilide, } \text{CH}_2:\text{CH.CO.NH.Ph. — } \textcircled{P} \text{ } \textcircled{D} \text{ Savn. T.* products: aniline; acrylic ac. (Vol. I).}$

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1896	104	Methylcarbanilide, MePhN.CO.NHPh . — $\text{P} \text{ D}$ Sapon. T.* products: methylaniline (No. 2.1249); aniline; CO_2 ; — Ndl. fr. alc. I. c. aq., lgr.; d.s. c. alc.; v.s. eth., bz. — Boils at 204° decg. to methylaniline & phenyl isocyanate (of irritating odor).
1897	104	α -Hydroxyvaleramide, $\text{Me}_2\text{CH.CH(OH).CO.NH}_2$. — $\text{P} \text{ D}$ Sapon. T.* products: NH_3 ; α -hydroxyvaleric ac. ($\text{C}_6\text{H}_{10}\text{O}_2$, Vol. I). — Cryst. s. aq., alc.
1898	104	α -Dimethyloxamide, $\text{Me}_2\text{N.CO.CO.NH}_2$. — $\text{P} \text{ D}$ Sapon. T.* (w. alk.) products: NH_3 ; dimethylamine (No. 2.1061); oxalic ac. (Vol. I). — Tbl. fr. bz. E.s. aq., alc.; d.s. eth.
1899	105-6	Ethylsemicarbazide, $\text{NH}_2\text{CO.NH.NHET}$. — P Semicarbazide. (Reduces Fehling's sol. only on heating.) Lft. V.s. aq. (!); alc.; d.s. eth.
1900	105	Acetophenonephenylhydrazone, Me.C(=O.NHPh).Ph . — P Hydrazone. § — Fine ndl. D.s. aq., c. alc.; e.s. eth.
1901	105	† Vanillinphenylhydrazone, $(\text{HO})(\text{MeO})\text{C}_6\text{H}_4\text{CH(=O)NHC}_6\text{H}_4\text{Ph}$. — P Hydrazone. § Gives Bülow's react. (T. 2.11). — Silvery lft. V.d.s. aq., lgr.; e.s. alc., eth., bz. Becomes colored on keeping.
1902	105	anti- α -Tolylphenylketoxime, $\text{Me.C}_6\text{H}_4\text{C(=NOH)Ph}$. — Oxime. §
1903	105	α -Nitrobenzophenone, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.Ph}$. — Nitro comp. † — Cryst. R.d.s. abs. alc.
1904	105	s-Dipropylurea, PrNH.CO.NH.Pr . — $\text{P} \text{ D}$ Sapon. T.* products: propylamine (No. 2.1067); CO_2 . — Ndl. fr. h. aq. B.p. 255°.
1905	105-6	Piperidylurea, $\text{NH}_2\text{CO.C}_6\text{H}_{10}$. — $\text{P} \text{ D}$ Sapon. T.* products: NH_3 ; piperidine (No. 2.1112); CO_2 . — Ndl. fr. alc. Sol. in v. conc. HNO_3 , giving nitro-piperidine. — B.HNO_3 , m.p. 67°.
1906	105	Diethylacetamide, $\text{Et}_2\text{CH.CO.NH}_2$. — $\text{P} \text{ D}$ Sapon. T.* products: NH_3 ; diethylacetic ac. ($\text{C}_6\text{H}_{10}\text{O}_2$, Vol. I). — Ndl. fr. alc. E.s. aq. (!), alc., eth. B.p. 230-5°.
1907	105-6	Butyroformamide, $\text{Et.CH}_2\text{CO.CO.NH}_2$. — $\text{P} \text{ D}$ Sapon. T.* products: NH_3 ; butyroformic ac. (b.p. 180-5° d.).
1908	105 u.c.	Isobutyro-p-toluide, $\text{Me}_2\text{CH.CO.NH.C}_6\text{H}_4\text{Me}$. — $\text{P} \text{ D}$ Sapon. T.* products: p-toluidine (No. 2.566); isobutyric ac. (Vol. I).
1909	105-6	Benzoylbenzylamine, $\text{Ph.CO.NH.CH}_2\text{Ph}$. — $\text{P} \text{ D}$ Sapon. T. ("by HCl at 175°") products: benzylamine (No. 2.1236); benzoic ac. (Vol. I).
1910	105	Diacetylhydrazobenzene, $\text{Ph.N(C}_6\text{H}_4\text{O)}\text{N(C}_6\text{H}_4\text{O).Ph}$. — $\text{P} \text{ D}$ Protracted boiling w. conc. HCl gives benzidine (No. 2.840). — Rhomb. cryst. fr. alc. D.s. aq.; e.s. alc., eth.
1911	106; (102-3)	Pr-2,3-Dimethylindole, $\text{Me}_2\text{C}_6\text{H}_3\text{N}$. — P Odor disagreeable, fecal. Gives coloration in pyrrole splinter react. (T. 2.24-b). — Lust. lft. fr. lgr. V.d.s. h. aq.; d.s. c. lgr.; v.s. alc., eth. S. conc. HCl, but repptd. by aq. Vol. w. st. — B.p. 285° (th.i., 750 mm.). — D B.Pk (T. 2.23), lust. brown ndl. fr. alc., m.p. 157°.
1912	106.5 (frothing)	Methylene-N,N-bisphenylhydroxylamine, $\text{Ph.N(OH).CH}_2\text{N(OH).Ph}$. — P FeCl_3 colors sol. transiently brownish violet. Pungent odor of nitroso-benzene then appears, color becoming violet on diln. w. aq. — Decd. by boiling in presence of air. Ndl. fr. chlf. Alm. i. aq.; e.s. eth., chlf.; less s. c. alc., lgr. — Al amalgam reduces to aniline & methylaniline.
1913	106 u.c.	p-Nitrobenzaldehyde, $\text{NO}_2\text{C}_6\text{H}_4\text{CHO}$. — P Reduces Tollen's reag. quickly in T. 2.30. — Thin alm. colorless pr. Odor, faint, bitter almond-like. D.s. h. aq., alc.; alm. i. c. aq.; s. eth. Taste of c. sat. aq. sol. sl. bitter—not burning. D Convert into p-nitrobenzaldehydophenylhydrazone, closely following procedure given under No. 2.2898, for the corresponding ortho aldehyde. The product cryst. in v. thin pr. of RS1 color, when dry, w. m.p. 156.5° u.c.
1914	106	(β)-Benzoinphenylhydrazone, $\text{C}_2\text{H}_5\text{C}_6\text{H}_4\text{N}_2$. — P Hydrazone. § — Monoclin. pr. fr. lgr. Abs. alc. at 20° dis. 8.8%.
1915	106-7	Levulinicacidphenylhydrazoneanhydride, $\text{C}_11\text{H}_{12}\text{ON}_2$. — P Gives hydrazone reduction T. § — Tbl. fr. alc. I. aq., c. alk.; r.d.s. alc., v.s. chlf. B.p. 340-50° (th.i.), w. sl. deen.

No.	Melting-point (C. ^o). ³	NEUTRAL COMPOUNDS.—Colorless and Solid.
I916	106-7	α -Pyridone, 2-(α)-Oxypyridine, $\text{HO.C}_6\text{H}_4\text{N}$. — B.p. 280-1°. Ndl. fr. bz. V.s. aq. (!), alc., chlf.; mod. s. eth., bz. Aq. sol. dis. HgO , & is reddened by FeCl_3 ("Aq. sol. reacts neut. & has lower electrical conductivity than that of phenol.")
I917	106	anti-Dimethylsuccinimide, $^{\text{t}}\text{NH.CO.CHMe.CHMe.CO}^{\text{t}}$. — \textcircled{P} \textcircled{D} Sappn. T.* (w. KOH) products: NH_2 ; anti-dimethylsuccinic ac. ($\text{C}_8\text{H}_{10}\text{O}_4$, Vol. I). — E.s. aq., alc.; d.s. eth. B.p. 300°.
I918	107	Propylurea, $\text{NH}_2\text{CO.NHPr}$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; propylamine (No. 2.1067); CO_2 . — S. aq.
I919	106-7; 104-5	Palmitamide, $\text{C}_{16}\text{H}_{31}\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; palmitic ac. (Vol. I). — S. alc. B.p. 235-6° (12 mm.).
I920	107-8d.	Levulinamide, $\text{Me.CO.CH}_2\text{CH}_2\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; levulinic ac. (Vol. I). — 6-sided tbl. fr. alc. + chlf. E.s. aq. (!), alc.; d.s. chlf.
I921	107; (104)	Pyrotartaranil, $^{\text{t}}\text{PhN.CO.CH}_2\text{CHMe.CO}^{\text{t}}$. — \textcircled{P} \textcircled{D} Prob. final sappn. T.* products: aniline; pyrotartaric ac. ($\text{C}_8\text{H}_8\text{O}_4$, Vol. I). [Pyrotartaranilic ac., No. 2.169, may be prepared by warming w. alk.] — Cryst. powd. d.s. h. aq.; v.s. alc.; e.s. eth.
I922	107-8	p-Cyanobenzophenone, $\text{CN.C}_6\text{H}_4\text{CO.Ph}$. — \textcircled{P} \textcircled{D} Sappn. T.* (w. alc. KOH) products: NH_2 ; p-benzoylbenzoic ac. ($\text{C}_{14}\text{H}_{10}\text{O}_3$, Vol. I). Cryst. fr. dil. alc.
I923	107	Carbonyldiurethane, $(\text{EtCO}_2\text{NH})_2\text{CO}$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; ethyl alc.; CO_2 . — V. stable. Tbl. e.s. aq! Gives d.s. Ag salt w. alk. AgNO_3 sol.
I924	107-8	Diacetylphenylhydrazine, $\text{Ph}(\text{C}_2\text{H}_5\text{O})\text{N.NH}(\text{C}_2\text{H}_5\text{O})$. — \textcircled{P} "Reduces Fehling's sol." — Tbl. fr. alc. + bz. E.s. alc., h. aq.; v.d.s. eth. — Sappn. T.* products: phenylhydrazine (No. 2.1369); ac. ac.
I925	107-8	Hippuricaldehydephenylhydrazone, $\text{Ph.CO.NH.CH}_2\text{CH}_2\text{N.NH.Ph}$. — \textcircled{P} Hydrazone § — Pr. fr. bz. E.s. alc.; less s. eth., bz.
I926	107.5 <u>c</u> .	Methyl 3,5-Dinitrobenzoate, $[(\text{NO}_2)_2\text{C}_6\text{H}_3\text{CO}_2\text{Me}]$. — \textcircled{P} Nitro comp.† — Pr. & lft. D.s. c. alc.; e.s. h. alc.
I927	107.5-8	Phenyl- β -naphthylamine, $\text{Ph.NH.C}_{10}\text{H}_7$. — \textcircled{P} Sol. show blue fluor. — B.p. 395-5.5° c. Ndl. fr. MeOH. E.s. h. org. solvents. W. conc. HCl at 240° gives β -naphthol & aniline. Br. in gl. ac. ac. sol. gives dibromo deriv., ndl. fr. bz., m.p. 140°.
I928	108	Arachidamide, $\text{C}_{18}\text{H}_{38}\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; arachidic ac. (Vol. I). — Lust. ndl. fr. alc. I. eth.
I929	108; 108.5-9; 107	Stearamide, $\text{C}_{17}\text{H}_{36}\text{CO.NH}_2$. — [Used as an emulsifier.] — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; stearic ac. (Vol. I). — Sol. alc.
I930	108	α -Aminobenzamide, $\text{NH}_2\text{C}_6\text{H}_4\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sappn. T.* (using HCl) products: NH_2 ; α -aminobenzoic ac. (No. 2.148). — Lft. fr. chlf. E.s. alc., h. aq.; d.s. eth.; alm. i. bz.
I931	108	Lacto- α -naphthalide, $\text{Me.CH(OH).CO.NH(C}_{10}\text{H}_7)$. — \textcircled{P} \textcircled{D} Sappn. T.* products: α -naphthylamine (No. 2.589); lactic ac. (Vol. I). — Pr. fr. dil. alc.
I932	109	Cerotamide, $\text{C}_{22}\text{H}_{41}\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sappn. T.* products: NH_2 ; cerotic ac. (Vol. I). — Cryst. gran. I. c. alc.; e.s. h. alc.
I933	109	α -Nitrobenzonitrile, $\text{NO}_2\text{C}_6\text{H}_4\text{CN}$. — \textcircled{P} Nitro comp.† — Silky ndl. E.s. h. aq., alc., chlf.; less s. aq., lgr. — Sappn. T.* products: NH_2 ; α -nitrobenzoic ac. (No. 2.139).
I934	109	Trinitrocumene, $(\text{NO}_2)_2\text{C}_6\text{H}_2\text{CH}_2\text{Me}$. — \textcircled{P} Nitro comp.† — Colorless ndl. Alm. i. aq. E.s. h. alc.
I935	110	† 8, or 5-Nitroisoquinoline, $\text{NO}_2\text{C}_6\text{H}_4\text{N.H}_2\text{O}$. — \textcircled{P} Nitro comp.† — Ndl. fr. aq. E.s. alc. — B.HCl , tbl., m.p. 245°; 240°. — B.Pk , yel. lft., m.p. 220°.
I936	110-1	Trinitro-m-diisopropylbenzene, $(\text{NO}_2)_2\text{C}_6\text{H}(\text{CH}_2\text{Me})_2$. — \textcircled{P} Nitro comp.† — Ndl. i. c. aq.

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(ORDER II, SUBORDER I.)

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1937	110	Trinitro-1,3-dimethyl-4-propylbenzene, $(NO_2)_3.C_6H_3.Me_2$. — \textcircled{P} Nitro comp.‡ — Ndl. i. aq.
1938	110	2,4,6-Trinitro-1,3-dimethyl-5-(tert.)butylbenzene, $(NO_2)_3.C_6H_3.(CMe_2)_2.C$. — \textcircled{P} Nitro comp.‡ — Yellowish white ndl. fr. alc. Odor strongly musky!
1939	110	Methylisopropylketonesemicarbazone, $Me.C(:N.NH.CO.NH_2).CH_2.Me_2$. — \textcircled{P} Gives semicarbazone react. § Lft. w. fatty luster. S. aq. E.s. alc., eth.
1940	110 u.c.	† Benzaldehydephenylbenzylhydrazone, $Ph.CH_2:N.N(Ph.CH_2)Ph$. — \textcircled{P} Gives hydrazone react. § — YT4 lust. ndl. fr. h. dil. alc. I. aq.; e.s. alc., eth., bz.
1941	110; 102; 97	Lauramide, $C_{11}H_{22}.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; lauric ac. (Vol. I). Lft. B.p. 200° (12.5 mm.).
1942	109–11	Isovaleranilide, $(Me_2.CH.CH_2.CO)NH.Ph$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; isovaleric acid, (Vol. I). — Lft. d.s. h. aq.; e.s. alc., eth.
1943	110	† o-Acetoluide, $Me.C_6H_4.NH(C_2H_5O)$. — \textcircled{P} \textcircled{D} Saponify 1 g. w. HCl by T. 2.26-a & identify the o-toluidine (No. 2.1262) formed by Sivobof's method (Vol. I), after drying over solid KOH. — B.p. 296°. — Cryst. s. in 116 pt. aq. at 19°. Gives powerful carbamine odor in T. 2.12. — \textcircled{D} Dis. 2 pt. in 13 pt. gl. ac. ac. & pass current of air charged w. Br vapors through sol. until a mass of white cryst. forms. Drain. Press. Recryst. fr. alc. The product, m-bromoacetoluide, melts at 156–7°.
1944	110–1	$\alpha\beta$ -Dinaphthylamine, $(C_{10}H_7)_2.NH$. — Pr. E.s. h. alc., eth., bz.; d.s. lgr. — \textcircled{D} B.Pk., brown-black ndl. fr. eth., m.p. 172–3°.
1945	110	β -Methyl- γ -phenylhydantoin, $[NPh.CO.NMe.CH_2.CO]^+$. — Pr. D.s. c. aq.; e.s. alc.; v.s. chlf.
1946	111 u.c.; 111–2; 113	Antipyrine, Analgesin, 2,3-Dimethyl-1-phenylpyrazolone (5), $[NPh.CO.CH_2:CMe.NMe]^+$. — Colorless, alm. odorless cryst. w. faint bitter taste. S. at 25° in less than 1 pt. aq., 1 pt. alc., 1 pt. chlf., 30 pt. eth. B.p. 319° c. (174 mm.). — \textcircled{P} (1) 1 drop 10% aq. FeCl ₃ sol. added to 2 cc. $\frac{1}{2}\%$ aq. antipyrine sol. gives OR color, changing to OY T2 on addition of 2 drops dil. H ₂ SO ₄ . — (2) 2 drops fuming HNO ₃ added to 2 cc. $\frac{1}{2}\%$ aq. sol. gives G color, changing to reddish brown on addition of 4 to 5 more drops acid & heating to boiling. — (3) Tannic ac. sol. gives heavy white ppt. in aq. sol. \textcircled{D} Add 0.5 cc. 10% NaNO ₂ sol. & 1 drop conc. H ₂ SO ₄ to sol. of 0.1 g. substance in 2 cc. aq. Shake. Wash cryst. green ppt. on filter w. 5 cc. aq. Recryst. fr. 10 cc. boiling acetone. Wash w. 2 cc. acetone. Dry 20 min. on tile at 100°. The product, nitrosoantipyrine, forms bril. B-BG ndl. deg. w. efferv. at abt. 195° u.c. (199.5° c.), & becoming dark brown.
1947	111	Behenamide, $C_{21}H_{42}.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; behenic ac. (Vol. I). — Tbl. fr. alc.
1948	111	Tetramethylenecarbonanilide, $C_6H_4.CO.NHPh$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; tetramethylenecarbonic ac. ($C_6H_4O_2$, Vol. I). — Long ndl. fr. alc.
1949	111–2	o-Acetylaminooethylbenzene, $(C_6H_5O)NH.C_6H_4.Et$. — \textcircled{P} \textcircled{D} Sapn. T.* products: o-aminoethylbenzene (No. 2.1299); acetic ac. — S. h. aq. B.p. 304–5°.
1950	112	s-Diethylurea, $CO.(NHEt)_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylamine (No. 2.1062); CO ₂ . — E.s. alc., eth. B.p. 263° c.
1951	112; (102)	† Methyl p-Aminobenzoate, $NH_2.C_6H_4.CO_2Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-aminobenzoic ac. (No. 2.290); methyl alc. — Lft. fr. dil. MeOH.
1952	112	Mannite Hexanitrate, Nitromannite, $CH_2(NO_3)_2.[CH_2(NO_3)_4].CH_2(NO_3)_2$. — \textcircled{P} Should give blue color in T. 2.15! — Explodes violently on percussion, & deflagrates when thrown on hot surface. Ndl. s. in 34.4 pt. alc. at 12.8°.
1953	112.5	Benzoylhydrazine, Benzhydrazide, $Ph.CO.NH.NH_2$. — \textcircled{P} Reduces Fehling's sol. or Tollen's reagt. in the cold. ["Reacts neutral."] Silvery tbl. Mod. s. aq.; less s. eth., chlf., bz. — Boiled w. acid or alk. gives hydrazine & benzoic ac.
1954	112	β - or 2,3,4-Trinitrotoluene, $(NO_2)_3.C_6H_3.Me$. — \textcircled{P} Nitro comp.‡ — Lft. fr. alc. D.s. c. alc.; e.s. eth., acetone, bz. Heated w. alc. ammon. gives β -dinitrotoluidine, m.p. 94°. — Heated w. alk. gives dinitrocresol.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1955	112-3	1,2,5-Trinitronaphthalene, $(NO_2)_2.C_{10}H_4$. — Nitro comp.‡ — Colorless ndl. fr. alc.
1956	111; 113	Di- α -naphthylamine, $(C_{10}H_7)_2NH$. — B.p. 310-5° (15 mm.). Quadrat. lft. S. alc.; v.s. eth., chlf., bz. — ⊕ B.2Pk (T. 2.23), small black ndl. fr. eth., m.p. 168-9°.
1957	113	Methylsemicarbazide, $NH_2.CO.NH.NHMe$. — ⊕ Semicarbazide.§ — Ndl. fr. bz. E.s. aq. (!), alc.; d.s. eth.
1958	113-4	γ -Benziloxime, $Ph.CO.CPh:NHOH$. — ⊕ Oxime§ [split at 100° by conc. HCl to benzil (Vol. I) & hydroxylamine.] — Cryst. w. $\frac{1}{2}C_6H_6$ in ndl. fr. bz., m.p. 70°. I. aq.; d.s. lgr.; v.s. alc.
1959	113	Benzophenonephenylimide, $Ph.C:NPh$. — ⊕ ⊖ Sapn. T.* w. HCl splits quickly to aniline & benzophenone (Vol. I). — Lft. fr. abs. alc. E.s. h. alc., eth., bz. B.p. a. 360°.
1960	113	Anilinoacetanilide, $Ph.NH.CH_3.CO.NH.Ph$. — ⊕ ⊖ Sapn. T.* products: aniline; anilinoacetic ac. (No. 2.99). — Ndl. D.s. c. aq.; s. alc., eth., h. bz.
1961	113-4	4-Formylamino-1,3-xylene, $(CHO)NH.C_6H_4.Me$. — ⊕ ⊖ Sapn. T.* products: 4-amino-1,3-xylene (No. 2.1308); formic ac. (Vol. I). — Cryst. fr. aq.; e.s. alc., eth.
1962	114	Methyloxanilide, $Ph.NH.CO.CO_2Me$. — ⊕ ⊖ Sapn. T.* products: aniline; methyl alc.; oxalic ac. — Tbl. E.s. alc.
1963	114-5	Ethyl Oxamate, $NH_2.CO.CO_2Et$. — ⊕ ⊖ Sapn. T.* products: NH_3 ; ethyl alc.; oxalic ac.
1964	114	p-Nitrobiphenyl, $Ph.C_6H_4.NO_2$. — ⊕ Nitro comp.‡ — B.p. 340° (th.i.). Ndl. D.s. c. alc.; e.s. chlf., eth.
1965	114-5	2,5-Dimethylindole, $Me_2.C_6H_3N$. — Dist. unded. Alm. i. h. aq.; e.s. h. alc., eth., bz. — ⊖ B.Pk (T. 2.23), dark red ndl. fr. bz., m.p. 155°.
1966	115-6	† Benzylideneacetoneoxime, $Ph.CH:CH.C(:NOH).Me$. — ⊕ Gives oxime react.§ — Odor somewhat like cumarine. Ndl. D.s. c. aq.; e.s. alc., eth. Boils w. decn. & evolution of NH_3 . Heated w. P_2O_5 gives isoquinoline (No. 2.1365).
1967	115-6	(β)-p-Phenyltolylketoxime, $Ph.C(:NOH).C_6H_4Me$. — ⊕ Oxime.§ — Ndl.
1968	115-6	Benzaldehydebenzoylphenylhydrazone, $Ph.CH:N.N(Ph.CO)Ph$. — ⊕ Hydrazone.§ — Silky ndl. Alm. i. aq.; e.s. bz.; v.s. chlf.
1969	115.5	Acetonebenzoylphenylhydrazone, $Me_2C:N.N(Ph.CO)Ph$. — ⊕ Hydrazone.§ — Lust. pr. V.s. alc., acetone.
1970	115	Butyramide, $Pr.CO.NH_2$. — ⊕ ⊖ Sapn. T.* products: NH_3 ; n-butyric ac. (Vol. I). Tbl. V.s. aq. B.p. 216°.
1971	114-6	Valeramide, $Me.(CH_2)_2.CO.NH_2$. — ⊕ ⊖ Sapn. T.* products: NH_3 ; n-valerianic ac. (Vol. I). — E.s. aq., alc., eth. Odor perspiration-like.
1972	115-6	p-Ethylbenzamide, $Et.C_6H_4.CO.NH_2$. — ⊕ ⊖ Sapn. T.* products: NH_3 ; p-ethylbenzoic ac. ($C_6H_5O_2$, Vol. I).
1973	115-6	Benzylphthalimide, $[C_6H_5.CO.N(Ph.CH_2).CO]^+$. — ⊕ ⊖ Sapn. T.* products: benzylamine (No. 2.1236); phthalic ac. (Vol. I). — Ndl. fr. alc.
1974	115	ab-Allylphenylurea, $(C_6H_5)NH.CO.NHPh$. — ⊕ ⊖ Sapn. T.* products: aniline; allylamine (No. 2.1068); CO_2 . — Thick ndl. fr. bz.
1975	114.2c.; 115-6	† Acetanilide, $Me.CO.NHPh$. — Colorless lft. fr. h. aq. [M.p. usually given as abt. 113°, but preparations recryst. fr. 95% alc. appear to melt 2° higher (Ber., 31, 661)]. S. at 25° in 179 pt. aq., 2.5 pt. alc., 12 pt. eth., or 5 pt. chlf.; in 18 pt. boiling aq., or 0.4 boiling alc. B.p. 303.8° (th.i., 760 mm.); 305°. Odorless. Taste, faintly burning. ⊕ ⊖ Sapn. T.* products; aniline; acetic ac. — Gives carbylamine odor directly in T. 2.12. — ⊖ Dis. 0.1 g. in 10 cc. h. aq. Filter, when cold, & add Br aq., drop by drop, as long as ppt. continues to form. (Antipyrine, No. 2.1948, gives no ppt.). Filter. Wash w. 5 cc. c. aq. Cryst. fr. 3 cc. h.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

(ORDER II, SUBORDER I.)

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		<i>25% alc. Wash w. 1 cc. 25% alc. Recryst. fr. 2 cc. h. 25% alc. Wash w. ½ cc. dil. alc. Dry at 100°. The product, p-bromoacetanilide, is obtained in colorless ndl., m.p. 170° u.c. (173° c.).</i>
1976	115	Acetylcarbonanilide , $(C_6H_5O)PhN.CO.NHPh$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline (T. 2.1235); acetic ac.; CO_2 . — Lft. fr. aq.
1977	115	Dehydراacetanilide , $C_6H_5O.NHPh$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. NaOH) prob. gives: aniline; acetone; ac. ac.; CO_2 ; etc. — Ndl. E.s. alc., eth. Vol. w. st.
1978	115–6	Isobutyro-o-toluidine , $Me.CH.CO.NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: o-toluidine (No. 2.1262); isobutyric ac. (Vol. I). — Ndl. fr. bz. D.s. lgr.
1979	115	Phenylacet-α-naphthalide , $Ph(C_6H_5O)N.C_{10}H_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: phenyl- α -naphthylamine (No. 2.1617–1); acetic ac. — Cryst. I. aq.; e.s. alc., bz.; d.s. eth.
1980	116	Melissamide , $C_6H_5CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; melissic ac. (Vol. I). — I. aq.; s. h. alc.
1981	116–7	Propionylformamide , $Et.CO.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; propionylformic ac. [oil of peculiar odor, b.p. 74–8° (25 mm.)]. — Lft. fr. eth. E.s. aq., alc.; less s. eth.
1982	116–7	2-Formylamino-1,4-xylene , $(CHO)NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-xylidine (No. 2.1315); formic ac. — Ndl. fr. aq.
1983	116.5	Formyl-aa-diphenylhydrazine , $(CHO)NH.NPh$. — \textcircled{P} May reduce Tollen's reagt. — \textcircled{D} Sapn. T.* products: diphenylhydrazine (No. 2.1491); formic ac. — Ndl. fr. dil. alc. E.s. alc. d.s.; lgr.
1984	117	Phenylacetanilide , $Ph.CH_2.CO.NHPh$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; phenylacetic ac. (Vol. I). — Pr. fr. alc. E.s. alc., eth.
1985	117.5–8	Lactophenine , Lacto-p-phenetidine , $Me.CH(OH).CO.NH.C_6H_4.OEt$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-phenetidine (No. 2.1392); lactic ac. (Vol. I). — Ndl. fr. aq. E.s. alc., bz., h. aq.; d.s. eth., lgr.
1986	117–8; (115)	† m-Nitrobenzonitrile , $NO_2.C_6H_4.CN$. — \textcircled{P} \textcircled{D} Sapn. T.* (prob. w. HCl) products: NH ₃ ; m-nitrobenzoic ac. (No. 2.139). — Ndl. D.s. aq.; e.s. alc., eth., chlf.
1987	117	Camphoranol , $C_6H_{14}.CO.NPh.CO^+$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; mixt. of camphoric acids. — Ndl. fr. eth. I. c. aq.; e.s. alc., eth.
1988	116–7; 116; 120–1	Ethyl Indoxylate , $(C_6H_5ON).CO.Et$. — \textcircled{P} Heated w. conc. H_2SO_4 forms (blue) indigosulphonic ac. — Thick colorless pr. s. without decn. in alk. & pptd. unchanged fr. sol. by CO_2 .
1989	117	o-Dinitrobenzene , $(NO_2)_2.C_6H_4$. — \textcircled{P} Nitro comp.† — B.p. 319° (773 mm.). Tbl. fr. alc. 100 pt. aq. at 100° dis. 0.38 pt.; 100 pt. methyl alc. at 20.5°, 3.3 pt.; v.s. chlf.; e.s. c. bz. — \textcircled{D} Convert into o-nitrophenol (Cf. Ber., 9, 1829), or reduce to o-phenylenediamine, No. 2.751.
1990	116; 118	2,6-Dinitro-1-methoxybenzene , $(NO_2)_2.C_6H_4.OMe$. — \textcircled{P} Nitro comp.† — Colorless ndl. S. in 110 pt. 95% alc.
—	118	2,4'-Dinitrodiphenylmethane . — Cf. No. 2.3157. (Yellowish.)
1992	118	p-Ethylacetnitroanilide , $Et(C_6H_5O)N.C_6H_4.NO_2$. — \textcircled{P} Nitro comp.† — White or sl. yellowish cryst., d.s. aq., eth.; i. lgr.; e.s. alc., bz. — \textcircled{P} Sapn. T.* (w. 1 mol. KOH) products: p-nitroethylaniline; ac. ac.
1993	118–9	p-Dinitrobenzene-naphthalene , $(NO_2)_2.C_6H_4.C_{10}H_8$. — \textcircled{P} Nitro comp.† — Colorless ndl. fr. alc. V.d.s. alc. — Decd. by boiling w. aq. to naphthalene, (Vol. I) & p-dinitrobenzene (No. 2.2319).
1994	118–9	p-Hydrazophenetole , $EtO.C_6H_4.NH.NH.C_6H_4.OEt$. — \textcircled{P} Prob. reduces Tollen's reagt. directly (T. 2.30). — Ndl.
1995	119.5	Dibenzylketoxime , $(Ph.CH_2)_2C(:NOH)$. — \textcircled{P} Oxime.‡ — Cryst.
1996	119.5	† Camphoroxime , $C_{10}H_{16}.NOH$. — \textcircled{P} Oxime.‡ <i>The hydrolytic splitting in T. 2.17-a is so slight that the Tollen's reagt. is merely darkened without giving a distinct ppt. Odor strongly camphorous!</i> — Ndl. fr. dil. alc. I. aq.; e.s. alc., eth., alk. acids.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
1997	119-20	Isothujoneoxime, $C_{10}H_{16}NOH$. — \textcircled{P} Oxime. § — Ndl. fr. methyl alc. E. vol. w. st. Opt. i. D.s. lgr.; more s. aq.
1990	119	Trinitrocymene, $(NO_2)_2C_6HMe(Me_2CH)$. — \textcircled{P} Nitro comp. † — Thin lft. S. h. alc.
1999	120	Acet-2,4-dinitroanilide, $(C_6H_4O)NH.C_6H_4.(NO_2)_2$. — \textcircled{P} Nitro comp. † — Ndl. fr. alc. I. c. aq.; e.s. h. alc. — \textcircled{D} Sapn. T.* products: 2,4-dinitroaniline (No. 2.3539-1); acetic ac.
2000	120	Acetonylacetoneosazone, $Me.C(N:NHPh).CH_2.CH_2.C(:N.NHPh).Me$. — \textcircled{P} Osazone. § — Unstable, resinifying. Lust. white lft. fr. dil. alc. E.s. alc., eth., bz.; i. lgr.; s. acids w. decn.
2001	120; 128-9	Dibenzylketonephenylhydrazone, $(Ph.CH_2)_2C:N.NH.Ph$. — \textcircled{P} Hydrazone. § — Lft. fr. alc. E.s. eth., bz.; h. alc.
2002	120-1	Anisicaldehydophenylhydrazone, $MeO.C_6H_4.CH:N.NH.Ph$. — \textcircled{P} Hydrazone. § — Cryst. fr. alc. I. aq.; e.s. eth., h. alc.
2003	120	† Cyanacetamide, $CN.CH_2.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ & probably malonic ac. (Vol. I). — Hexag. tbl. s. in 6.5 pt. c. aq., or in 55 pt. c. aq.
2004	120	Isobutylacetamide, $Me_2CH.CH_2.CH_2.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; isobutylacetic ac. ($C_6H_{12}O_2$, Vol. I). — S. aq.
2005	120-1	Diisobutylacetamide, $(Me_2CH.CH_2)_2CH.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; diisobutylacetic ac. — Ndl. Alm. i. aq.; v.s. alc.
2005	120	Glycollamide, $HO.CH_2.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; glycolic ac., (Vol. I). — Cryst. E.s. aq.; d.s. alc.
2007	120	Ethylacetylurea, $EtNH.CO.NH(C_2H_5O)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; ethylamine (No. 2.1062); CO ₂ . — Thick pr. fr. eth. E.s. aq., alc., eth. Sbl.
2008	120	Methacrylanilide, $CH_2:CMe.CO.NH.Ph$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; methacrylic ac. ($C_6H_6O_2$, Vol. I). — Pr. w. adamantine luster fr. dil. alc.
2005	120-1	Dimethylcarbanilide, $(MePhN)_2CO$. — \textcircled{P} \textcircled{D} Sapn. T.* products: methyl-aniline (No. 2.1249); CO ₂ . — B.p. 350°. Monoclin. tbl. fr. alc. I. aq.; e.s. alc., eth., bz.
2010	120	Oxindole, $[CO.NH.C_6H_4.CH_3]$. — \textcircled{P} Continued boiling w. ammon. AgNO ₃ sol. gives Ag mirror. — Long ndl. fr. aq. E.s. h. aq.; s. alc., eth.; more s. in aq. alk. than in aq., but extracted fr. alk. sol. by eth. — Ag \ddot{A} floc. ppt. — B.HCl, cryst. s. aq.
2011	120-2	4-Hydrazo-1,3-xylene, $Me_2C_6H_3.NH.NH.C_6H_4.Me$. — \textcircled{P} Prob. reduces Tollen's reagt. — Ndl. fr. alc.
2012	121-5	Ironoxime, $(C_{11}H_{17})C(:NOH)Me$. — \textcircled{P} Oxime. § — Lft. fr. lgr.
2013	121	Trinitro-1,2-dimethyl-4-ethylbenzene, $(NO_2)_2C_6Me_2Et$. — \textcircled{P} Nitro comp. † — Ndl. I. c. aq.
2014	121	2,5-Dinitroacetanilide, $(NO_2)_2C_6H_3.NH(C_2H_5O)$. — \textcircled{P} Nitro comp. † — Warming w. conc. H ₂ SO ₄ gives 2,5-dinitroaniline (No. 2.3254) & acetic ac. — Nearly colorless ndl. fr. alc. E.s. alc. Boiling KOH sol. gives NH ₃ .
2015	120-2	Glycero-p-toluide, $CH_2(OH).CH(OH).CONH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); glyceric ac. (Vol. I). — Lft. fr. aq. E.s. alc., eth.; d.s. bz., lgr.
—	122	s-Trinitrobenzene & 1,3,5-Trinitronaphthalene. — Cf. Nos. 2.3177, 2.3174. (Yellowish.)
2016	122	p-Tolyl-p-aminophenol, $HO.C_6H_4.NH.C_6H_4.Me$. — Lft. B.p. 350-60°. E.s. alc., bz. — Diacetyl deriv., tbl. fr. alc., m.p. 101°.
2017	120; 124	Anilinoantipyrine, $C_{17}H_{14}N_2$. — Cryst. fr. alc. "Not deod. by HCl or alc. KOH." Mixed w. conc. HNO ₃ gives a nitroazo deriv., $C_{12}H_8O_2N_4$, lemon-yel. lft., m.p. 164°, s. in alk. w. intense fuchsin-red color.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2018	123-4	Pr-2,3-Diphenylindole, $\text{Ph}_2\text{C}_6\text{H}_3\text{N}$. — \textcircled{P} Solutions fluor. blue. — Cryst. fr. bz. + lgr. B.p. 290-6° (10 mm.). I. aq.; e.s. alc., eth., bz.; r.d.s. lgr. "Does not give indole splinter react." — \textcircled{D} B.Pk (T. 2.23), red ndl. fr. bz., m.p. 154° d.
2019	123 u.c.	† Benzaldehydediphenylhydrazone, $\text{Ph}.\text{CH}:\text{N}.\text{NPh}_2$. — \textcircled{P} Gives good Ag reduction after hydrolysis in hydrazone T. 2.17. — YT4 ndl. fr. h. dil. alc. Alm. i. c. aq.; e.s. eth., bz. Sol. in conc. H_2SO_4 w. OYS1-2 color.
2020	123-4d.	Nitrosomethylurea, $\text{NH}_2\text{CO.N(NO)Me}$. — \textcircled{P} Prob. gives blue color in T. 2.15 w. diphenylamine. — Yellowish tbl. fr. eth. E.s. h. aq., alc., eth.
2021	123.5	2,6-Dinitro-1,4-xylene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Nitro comp.‡ — Hair-like ndl. fr. toluene. Alm. i. aq.
2022	123	4,6-Dinitroethylmesitylene, $(\text{NO}_2)_2\text{C}_6\text{Et}_2\text{Me}_2$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. I. aq.
2023	123-4	m-Nitrooctylbenzene, $\text{NO}_2\text{C}_6\text{H}_4\text{C}_8\text{H}_{17}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. I. aq., eth.; d.s. c. alc. E. vol. w. st.
2024	123.5	Pyromuconilide, $\text{C}_6\text{H}_4\text{O.CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; pyromucic ac. (Vol. I). — Silvery lft. & ndl. fr. alc. E.s. h. alc.
2025	123	Ethylisopropylacet-p-toluide, $\text{Et}(\text{Me}_2\text{CH})\text{CH.CO.NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); ethylisopropylacetic ac. ($\text{C}_6\text{H}_4\text{O}_2$, Vol. I). — Ndl. fr. pet.-eth.; i. aq.; e.s. organ. solvents.
2026	123	Piperovatin, $\text{C}_{15}\text{H}_{21}\text{O}_2\text{N}$. — [Toxic comp. of strychnine-like action fr. Piper ovatum of Trinidad.] — Ndl. fr. eth.-alc. I. aq., dil. ac. or alk.; v.d.s. lgr., abs. eth.; e.s. alc., chlf. ["Without basic properties."] W. aq. at 160° gives a vol. base, an acid, & an oil w. odor like anisole.
2027	124	Benzylidene-o-nitroacetophenone, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.CH:CH.Ph}$. — \textcircled{P} Dec. in sunlight to indigo, benzaldehyde & benzoic ac. — Silky colorless ndl. fr. alc.
2028	124-5	Hydrazopseudocumene, $\text{Me}_2\text{C}_6\text{H}_4\text{NH.NH.C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Prob. reduces Tollen's reagt. (T. 2.30).
2029	124-5	5-Hydrazo-1,3-xylene, $\text{Me}_2\text{C}_6\text{H}_4\text{NH.NH.C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} Prob. reduces Tollen's reagt. (T. 2.30). — Ndl. fr. alc.
2030	121-8	β -Protocatechuicaldehydophenylhydrazone, $(\text{HO})_2\text{C}_6\text{H}_4\text{CH:CH.NH.Ph}$. — \textcircled{P} Hydrazone.§ — Tbl. Changes on recrystn. or keeping to α -isomer, m.p. 175-6°.
2031	122; 127	o-Dinitrobibenzyl, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{C}_6\text{H}_4(\text{NO}_2)_2$. — \textcircled{P} Nitro comp.‡ — Flat lust. pr. fr. ac. ac. D.s. alc.; e.s. eth., bz.
2032	124-5	2,5-Dinitro-2,5-dimethylhexane, $\text{Me.CMe}(\text{NO}_2)_2(\text{CH}_2)_2\text{CMe}(\text{NO}_2)_2\text{Me}$. — \textcircled{P} Nitro comp.‡ — Lft. fr. bz. D.s. eth.; i. alk.
2033	124	2,4,6-Trinitro-1-methyl-3-isobutylbenzene, $(\text{NO}_2)_3\text{C}_6\text{H}_2\text{Me}(\text{C}_6\text{H}_5)$. — \textcircled{P} Nitro comp.‡ of musk-like odor! — Cryst. fr. dil. alc.
2034	124-5	as-Dibenzylurea, $\text{NH}_2\text{CO.N(Ph.CH}_2)_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; dibenzylamine (No. 2.1448); CO ₂ . — Pr. d.s. c. aq.; e.s. h. aq., alc.
2035	124	sec.-Butylacetamide, $\text{Et.CH(Me).CH}_2\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; sec.-butylacetic ac. — Ndl. fr. aq. E.s. alc., eth.
2036	124-5	Pyruvamide, Me.CO.CO.NH_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; pyruvic ac. (Vol. I). — Tbl. fr. alc. E.s. aq.; s. alc.
2037	124 u.c.	† Succinimide, $[\text{CO.NH.CO.CH}_2\text{CH}_2]^2$. — \textcircled{P} Ignite abt. 0.05 g. substance sharply in small t.t. w. 1 g. Zn dust. When destructive distn. begins insert pine splinter that has been soaked 30 sec. in conc. HCl in vapors. The wood becomes bright red. Cf. T. 2.24-b (2). — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; succinic ac. (Vol. I). — Mic. double octahedral pyramids w. 1 H ₂ O fr. sat. h. alc. sol. V.s. c. aq.; e.s. c. alc.; d.s. to v.d.s. eth. Odorless. Taste of conc. aq. sol. bitter.
2038	124	Diethylacetanilide, $(\text{Et}_2\text{CH.CO})\text{NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; diethyl acetic ac. ($\text{C}_6\text{H}_{12}\text{O}_2$, Vol. I). — Ndl. fr. alc.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2039	124 u.c.	Propion-p-toluide, Et.CO.NH.C ₆ H ₄ .Me. — ⊕ ⊖ Sapn. T.* products: p-toluidine (No. 2.566); propionic ac. (Vol. I). — Cryst. d.s. h. aq.
2040	125	α-Campholenamide, C ₁₀ H ₁₆ .CO.NH ₂ . — ⊕ ⊖ Sapn. T.* products: NH ₃ ; campholenic ac. (C ₁₀ H ₁₆ O ₂ , Vol. I). — Lft. d.s. aq.; e.s. alc., eth. Conc. HCl changes to β-isomer, m.p. 86°. [α] _D = -4.4° (alc. sol.).
2041	125	m-Benzotoluide, Ph.CO.NH.C ₆ H ₄ .Me. — ⊕ ⊖ Sapn. T.* products: m-toluidine (No. 2.1265); benzoic ac. — Cryst. fr. dil. alc.
2042	125-6	Isovaleryl-α-naphthalide, C ₁₀ H ₁₆ .CO.NH.C ₁₀ H ₇ . — ⊕ ⊖ Sapn. T.* products: α-naphthylamine (No. 2.589); isovaleric acid (Vol. I). — Silky ndl. fr. bz.
2043	125	Ethyldeneurethane, Me.CH(NH.CO.Et) ₂ . — ⊕ Warmed w. v. dil. H ₂ SO ₄ evolves acetaldehyde (pungent odor, Vol. I). — Minute ndl. fr. h. aq. E.s. h. aq.; alm. i. c. aq.; e.s. alc., eth. Odorless.
2044	125d	Hydroanthracene Nitrite, C ₁₄ H ₁₀ O ₄ N ₂ . — ⊕ Melts w. violent evolution NO. Hot dil. NaOH gives deep or. colored sol. — Lgr. ppts. rosettes of clear cryst. fr. bz. sol. — ⊖ Oxid. in ac. ac. sol. by CrO ₃ to anthraquinone (Vol. I).
2045	125	Carbanilphenylhydroxylamine, Ph.CH.CO.N(OH).Ph. — ⊕ FeCl ₃ colors alc. sol. red. — Ndl. fr. pet.-eth.
2046	125	Benzene-α-hydrazonaphthalene, Ph.NH.NH.C ₁₀ H ₇ . — ⊕ Prob. reduces Tollen's reag. slowly.
2047	125	anti-p-Hydroxybenzophenoneoxime, HO.C ₆ H ₄ C(:NOH).Ph. — ⊕ Oxime. § — Pr. fr. dil. ac. ac. E.s. alc., eth.; less s. gl. ac. ac., HCl, bz. — Boiling w. KOH sol. gives syn. deriv., m.p. 81°.
2048	126	anti-2,4-Dimethylbenzophenoneoxime, Me ₂ .C ₆ H ₄ C(:NOH)Ph. — Oxime. § Cryst. I. aq.; v.s. alc., eth.; less s. bz. Beckmann's rearrangement gives 2,4-xylic ac. anilide.
2049	126	2,3'-Dinitrobenzophenone, [(NO ₂) ₂ .C ₆ H ₄] ₂ .CO. — Nitro comp.‡ — Lust. cryst. fr. toluene.
2050	126u.c.	† Hydrazobenzene, Ph.NH.NH.Ph. — ⊕ Gives black ppt. in T. 2.30 w. Tollen's reag. on shaking. — Lust. colorless lft. fr. h. 50% alc. on cooling. Becomes or. colored by oxidn., especially in alk. sol. on contact w. air. V.d.s. aq. 100 pt. satd. abs. alc. sol. contain 5 pt. at 16°. — ⊖ (1) Dis. 0.01 g. in 2 cc. conc. HCl + 5 cc. aq. by heating. Add 2 drops conc. H ₂ SO ₄ . Sol. soon becomes turbid fr. separation of shimmering ppt. of minute cryst. of benzidine sulphate. — (2) Add 2 cc. 10% FeCl ₃ sol. to sol. of 0.1 g. substance in 11 cc. boiling 50% alc. Wash the YO ppt. w. 5 cc. dil. alc. Recryst. fr. 5 cc. h. 50% alc. Wash w. 4 cc. c. dil. alc. The product, azobenzene, forms O-YO scales, m.p. 68°.
2051	126u.c.; 128	† Benzamide, Ph.CO.NH ₂ . — ⊕ ⊖ Sapn. by T. 2.26-a w. HCl, boiling 30 min., gives NH ₃ & benzoic ac. (pearly scales fr. h. aq., m.p. 121-1.5°). — Odorless, tasteless monoclin. lbl. fr. h. aq. D.s. h. aq.; e.s. alc., eth., h. bz. Dist. w. sl. decn. giving some benzonitrile, recognizable by its strong bitter almond-like odor! — Evolves NH ₃ freely in T. 2.7.
2052	126	Phenyl Phenylcarbamate, Ph.NH.CO.Ph. — ⊕ ⊖ Sapn. T.* products: aniline; phenol; CO ₂ . — Lft. i. c. aq.; e.s. alc., eth.
2053	126; (131)	Anthranilanilide, o-NH ₂ .C ₆ H ₄ .CO.NH.Ph. — ⊕ ⊖ Sapn. T.* products: aniline (No. 2.1235); anthranilic ac. (No. 2.148). — Ndl. fr. bz. Mod. s. aq.; v.s. eth., chlf.; d.s. bz.
2054	126-7	α-Diethyloxamide, NH ₂ .CO.CO.NET ₂ . — ⊕ ⊖ Sapn. T.* products: diethylamine (No. 2.1068-1); NH ₃ ; oxalic ac. — B.p. 266-8° c.
2055	126	Nitroacetonitrile, N(CH ₃).CN ₃ . — Sapn. w. Ba(OH) ₂ gives NH ₃ & nitriloacetic ac. — Ndl. fr. alc.
2056	126-7c.d.	Ethyl-γ-phenylhydantoin, "NPh.CO.NH.CEt.CO". — D.s. h. aq.; e.s. h. alc.
2057	127	p-Methoxyacetaminophenol, "Methacetin," MeO.C ₆ H ₄ .NH(C ₆ H ₅ O). — ⊕ ⊖ Sapn. T.* products: p-anisidine (No. 2.614); ac. ac. Gives carbonylamine odor readily in T. 2.12! — Scales or ndl. s. in 50 pt. aq. at 15°, or 12 pt. on boiling; e.s. alc., chlf.; 100 pt. abs. eth. dis. 0.7 pt. at 20°.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2058	127	Triphenylamine, Ph_3N . — \textcircled{P} Sol. in gl. ac. ac. is colored violet & then blue by conc. H_2SO_4 , or green by trace HNO_3 . — Monoclin. cryst. fr. eth. D.s. alc.; e.s. bz. Forms no salts even w. picric ac.
2059	127-8	Rhamnoseoxime, $\text{C}_6\text{H}_{12}\text{O}_4(\text{NOH})$. — \textcircled{P} Oxime. § Tbl. fr. MeOH. V.s. aq.; i. eth.
2060	128	Cuminicaldehydephenylhydrazone, $p\text{-Me}_2\text{CH.C}_6\text{H}_4\text{CH: N.NHPh}$. — \textcircled{P} Hydrazone. § Ndl. fr. lgr. S. h. alc., eth., lgr.
2061	128	Isovaleramide, $\text{Me}_2\text{CH.CH.CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; isovaleric ac. (Vol. I). — S. aq. B.p. 230-2°.
2062	128-9	Isobutyramide, $\text{Me}_2\text{CH.CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; isobutyric ac. (Vol. I). — E.s. aq. B.p. 216-20°.
2063	128	α -Methylcinnamamide, Ph.CH: CMe.CONH_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; α -methyl cinnamic ac. ($\text{C}_{10}\text{H}_{10}\text{O}_2$, Vol. I). — Lst. fr. aq. D.s. c. alc., eth.
2064	128-9	Diisoamyloxamide, $(\text{C}_5\text{H}_{11})\text{NH.CO.CO.NH}(\text{C}_5\text{H}_{11})$. — \textcircled{P} \textcircled{D} Sapn. T.* products: isoamylamine (No. 2.1100); oxalic ac.
2065	128-8.5 u.c.	† 4-Acetamino-1,3-xylene, Acet-a-m-xylide, $(\text{C}_6\text{H}_4\text{O})\text{NH.C}_6\text{H}_4\text{Me}_2$. — \textcircled{P} \textcircled{D} Sapn. products in T. 2.26-a w. HCl ; 4-amino-1,3-xylene (No. 2.1308), identified by b.p. by Siwoloboff's method after drying over solid KOH; acetic ac. — Flat ndl. fr. h. alc. D.s. aq. Odorless. Tasteless. Gives strong carbyleamine odor in T. 2.12.
2066	128	Glycollic- α -naphthalimide, $\text{HO.CH}_2\text{CO.NH.C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: α -naphthylamine (No. 2.589); glycollic ac. (Vol. I). — Tbl. fr. acetone. E.s. h. aq.; d.s. chlf., c. bz.
2067	128.5	† ab-Acetylphenylhydrazine, $\text{Ph.NH.NH}(\text{C}_6\text{H}_4\text{O})$. — \textcircled{P} Reduces Tollen's reagt. immediately (T. 2.30). — Cryst. d.s. c. aq., eth.; e.s. h. aq., alc. — Sapn.* gives phenylhydrazine (No. 2.1369) & acetic ac.
2068	128-30(r.h.)	syn-(β)-Benzaldoxime, Ph.CH: NOH . — \textcircled{P} Oxime. § — Thin lust. trimet. tbl. or ndl. fr. eth. D.s. bz. (unlike anti isomer). Is transformed to anti isomer by contact w. dil. H_2SO_4 or prolonged heating.
2069	129d.	Ethylacetooacetatesemicarbazone, $\text{Me.C(: NH.CO.NH}_2\text{).CH}_2\text{CO.Et}$. — \textcircled{P} Semicarbazone. § — Ndl. fr. eth. E.s. h. aq. Heating at 120° or boiling w. aq. gives 3-methylpyrazolone (No. 2.378).
2070	129d.	l-Xylonicacidphenylhydrazone, $\text{CH}_2\text{(OH).(CHOH).CO.NH.NH.Ph}$. — Unstable ndl. — E.s. except in lgr., bz.
2072	128-30	Hydroxyurea, $\text{NH}_2\text{CO.NH(OH)}$. — \textcircled{P} Gives intense blue-violet color w. FeCl_3 . Reduces h. ammon. AgNO_3 , sol.! — Ndl. fr. alc. V.s. aq.
2073	129	3-(β)-Oxypyridine, $\text{HO.C}_6\text{H}_4\text{N}$. — \textcircled{P} FeCl_3 colors aq. sol. red! — Ndl. E.s. aq., alc. Dist. undecd. Ignition w. Zn. dust. gives pyridine, No. 2.1125. — Oxalate, d.s. abs. alc., m.p. 175°.
2074	129	Trinitro-1,4-dimethyl-2-ethylbenzene, $(\text{NO}_2)_3\text{C}_6\text{Me}_2\text{Et}$. — \textcircled{P} Nitro comp. † Pr. fr. alc. I. c. aq.
2075	129-30	Pr-2- α -Naphthylindole, $[\text{C}_8\text{H}_7\text{NH.C}(\text{C}_{10}\text{H}_7):\text{CH}]^+$. — \textcircled{P} \textcircled{D} B.Pk (T. 2.23), purple-red scales, m.p. 179°, d.s. lgr., e.s. bz. — Ndl. fr. alc.
2076	129	Formyl- β -naphthalide, $(\text{CHO})\text{NH.C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: β -naphthylamine (No. 2.781); formic ac. — Lst. e.s. alc., bz., chlf.; less s. eth.; d.s. h. aq.
2077	129-30u.c.	Dibenzoylpentamethylenediamine, $\text{Ph.CO.NH(CH}_2)_5\text{NH.CO.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: pentamethylenediamine (No. 2.1232); benzoic ac. (Vol. I). — Ndl. & scales e.s. alc. I. aq.
2078	130-1	1,2-Dimethylbenzamide(4), $\text{Me}_2\text{C}_6\text{H}_4\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; 1,2-dimethylbenzoic(4) ac. ($\text{C}_8\text{H}_{10}\text{O}_2$, Vol. I). — Lust. ndl. fr. h. aq.
2079	130	d-Glucosemethylphenylhydrazone, $\text{CH}_2\text{OH.(CHOH).CH: N.NMePh}$. — \textcircled{P} hydrazone. § — Tbl. fr. 98% alc.
2080	130-1d.	α -Hydrindonephenylhydrazone, $\text{C}_9\text{H}_8\text{: N.NH.Ph}$. — \textcircled{P} Hydrazone. § — Pr. fr. alc. D.s. lgr.; e.s. h. alc.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2001	129–30u.c.	† Piperine, Piperonylpiperidine, $C_4H_{10}N.CO.C_4H_4.C_4H_3 : O_2 : CH_2$. — [Fr. pepper.] Monoclin. cryst. Alm. i. c. aq.; s. in 15 pt. alc., 36 pt. eth., or 1.7 pt. chlf. at 25° ; s. in 4.4 alc. at 60° . Odorless. Tasteless at first, later pungent. ① (1) 1 mg. piperine w. 1 drop mixt. of 1 vol. 40% formalin + 2 vol. conc. H_2SO_4 gives permanent GS2 color. — (2) T. 22-a w. conc. H_2SO_4 gives OS1 color entirely disappearing on diln. w. aq. — (3) T. 22-b w. HNO_4 gives GY T1 color quickly changing to O, w. undissolved cryst. RS1. If 2 cc. 10% NaOH be added to the O sol. in a t.t. the turbid YT1 mixt. changes to RO → when heated to boiling. — Sappn.* w. conc. KOH sol. gives piperidine, (No. 2.1112) & piperic ac. (Vol. I).
2002	131	2,4,6-Trinitro-3-hexyltoluene, $(NO_2)_3.(C_6H_{11}).C_6H.Me$. — ② Nitro comp.‡ of faintly musky odor.
2033	131	Dinitro-1-methyl-5-tert. butyl-2-acetylbenzene, $(NO_2)_2.Me.(Me_2C).(C_6H_4O).-C_6H$. — ② Nitro comp.‡ of strong musk-like odor. — Ndl. fr. alc.
2004	131	Dinitrobenzil, $(NO_2)_2.C_6H_4O_2$. — ② Nitro comp.‡ — Octahedral cryst. s. in 41 pt. boiling, or 137 pt. c. alc. Separates fr. h. alc. in moss-like cryst. mixt., which after standing some weeks under bz. + alc. consists of octahedra & leafy cryst., the latter melting at 147° & s. in 290 pt. c. alc. [D.R.P. 44,269.]
2065	131	Benzyl-p-tolylketoxime, $Ph.CH_2.C(:NOH).C_6H_4.Me$. — ② Oxime.§ — Lft. fr. alc.
2086	131	Coumaroxime, $[O.C_6H_4.CH:CH.C(:NOH)]_2$. — ② Oxime.§ — Ndl. Alm. i. c. aq.; e.s. alc., eth., bz., alk. Prolonged heating w. HCl gives coumarin & hydroxylamine.
2067	131	Oxal-m-toluidine, $Me.C_6H_4.CO.CO.C_6H_4.Me$. — ② ③ Sappn. T.* products: m-toluidine (No. 2.1265); oxalic ac.
2008	131–2	Mandelamide, $Ph.CH(OH).CO.NH_2$. — ② ③ Sappn. T.* products: NH ₂ ; mandelic ac. (Vol. I). — Tbl. S. in 33.7 pt. aq. at 24° ; d.s. eth.
2089	131	2-Methylfurancarbonamide, $Me.C_6H_4O.CO.NH_2$. — ② ③ Sappn. T.* products: NH ₂ ; 2-methylpyromucic(5) ac. (Vol. I). — Pr. e.s. alc.
2000	131	Diacetyltriphenylguanidine, $[(C_6H_5O)PhN]_3.C.NPh$. — ② ③ Sappn. T.* (w. HCl) products: triphenylguanidine (No. 2.882); acetic ac.
—	131	uns.-Tetraphenylguanidine. — Cf. No. 2.855.
2091	131–2	Diphenylethaneguanidine, $Me.CH(:NPh)(NH.Ph)$. — ② ③ Sappn. T.* products: aniline; acetic ac. — Ndl. d.s. c. aq.; e.s. h. aq.; e.s. eth., acids. "Reacts neut." — $B_2.H_2PtCl_6$, d.s. c. aq., m.p. 210° .
2092	132u.c.	† Urea, Carbamide, $NH_2.CO.NH_2$. — ② ③ Sappn. T.* (w. NaOH) products: NH ₃ ; CO ₂ . — 4-sided pr. w. rough longitudinal striations fr. aq.; or smooth delicate ndl. when crysd. rapidly. Begins to soften at 130° . V.s. c. aq.; s. in 20 pt. c. alc.; alm. i. c. chlf. Odorless. Taste, cooling, saline, sl. bitter. — Heat 0.06 g. substance in t.t. over small flame. Comp. melts & boils giving white sublimate & odor of NH ₃ . When liquid disappears leaving white residue at bottom of t.t. remove flame & cool. Dis. residue in 2 cc. aq. + 10 drops 10% NaOH sol. Drop in CuSO ₄ component of Fehling's sol. (Cf. Vol. I), shaking after each drop. The first drop gives a VRT $\frac{1}{2}$ coloration, the second a RV, the third a V, and the fourth a BV. ("Biuret reaction") ③ Dis. 10 mg. in 2 drops aq. on small watch glass. Mix w. 3 drops conc. HNO_3 . Examine cryst. ppt. of urea nitrate under low power microscope. It is obtained in charac. rhombic, quadrangular or hexag. plates, separate or stratified-like shingles on a roof. The acute angles on the rhombs are about 82° . — [For the isolation of small quantities of urea fr. mixtures in dil. solutions (e.g. blood) & identification, see Schroeder, Z. anal. Chem., 22, 138.]
2093	132	Methylphthalimide, $[CO.C_6H_4.CO.NMe]^2$. — ② ③ Sappn. T.* products: methylamine (No. 2.1059); phthalic ac. (Vol. I). — Silky ndl. Alm. i. aq.; e.s. alc. Sbl. in lft. B.p. 285° c.
—	132.5	1-Canadine. — Cf. No. 2.858-1.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2094	132.5	Cyclopentyl Phenylcarbamate, $\text{Ph.NH.CO}_2(\text{C}_5\text{H}_5)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; cyclopentanol.
2095	132	Acet- β -naphthalide, $(\text{C}_2\text{H}_4\text{O})\text{NH.C}_10\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. HCl) products: β -naphthylamine (No. 2.781); ac. ac. — Lft. fr. alc.
2096	132	Pr-2-Methyl- α -naphthindole, $[\text{C}_{10}\text{H}_8.\text{NH.CMe}:\text{CH}]$. — \textcircled{P} Gives blue-violet color in pine splinter T. 2.24! — Ndl. fr. aq. E.s. alc., eth., bz.; d.s. c. lgr. D. vol. w. st. FeCl_3 colors sol. in gl. ac. ac. cherry-red. — \textcircled{D} B.Pk, dark red ndl. fr. bz., m.p. 167–8°.
2097	132–3	d,l-Glucosediphenylhydrazone, $\text{C}_6\text{H}_{12}\text{O}_6:\text{N.NPh}_2$. — \textcircled{P} Hydrazone. §— Colorless cryst.
2098	132–3	7-Nitroquinoline, $\text{NO}_2\text{C}_6\text{H}_4\text{N}$. — \textcircled{P} Nitro comp.† — Silvery cryst. fr. dil. alc. V.d.s. c. alc.; e.s. eth. — B.HCl, ndl., m.p. 225° d.
2099	132	p-Nitromethylcinnamylvinylketone, $\text{Me.CO.CH:CH.CH:CH.C}_6\text{H}_4(\text{NO}_2)$. — \textcircled{P} Nitro comp.† — Ndl. fr. dil. alc. E.s. alc.
2100	133–4; (126)	p-Hydrazotoluene, $\text{Me.C}_6\text{H}_4.\text{NH.NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} Prob. reduces Tollen's reag. directly on shaking. — Cryst. e.s. alc., eth. Oxid. by air in alc. sol. to orange colored p-azotoluene (No. 2.3303). Contact w. conc. HCl causes rearrangement to 3-toluidino-4-amino-toluene.
2101	133	Mesitylenamide, 1,3-Dimethylbenzenecarbonamide(5), $\text{Me}_2\text{C}_6\text{H}_4.\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; mesitylenic ac. (Vol. I). — Ndl. V.d.s. c. aq.; v.s. alc., eth.
2102	133	Anilinoacetamide, $\text{Ph.NH.CH}_2.\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; anilinoacetic ac. (No. 2.99). — Mic. ndl. V.s. alc., eth., h. aq.
2103	133	α -Hydroxyisovaleranilide, $\text{Me}_2\text{CH.CH(OH).CO.NHPH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; α -hydroxyisovaleric ac. ($\text{C}_6\text{H}_10\text{O}_3$, Vol. I). — Ndl. V.d.s. aq., lgr.; e.s. alc., eth., chlf.
2104	134; 131	3-Acet-1,2-xylide, $(\text{C}_2\text{H}_4\text{O})\text{NH.C}_6\text{H}_4.\text{Me}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: (v)-o-xylidine (No. 2.1333); ac. ac. — Ndl. E.s. alc., eth.; d.s. c. bz.
2105	134–4.5	Phenacetine, p-Acetaminoethoxybenzene, $(\text{C}_2\text{H}_4\text{O})\text{NH.C}_6\text{H}_4.\text{OEt}$. — [An antipyretic.] <i>Lust., odorless, tasteless lf. fr. h. aq.</i> S. at 25° in 900 pt. aq., 16 pt. alc., 64 pt. eth., 21 pt. chlf.; s. at 30° in 153 pt. bz., 9.3 pt. acetone; s. in 70 pt. boiling aq. \textcircled{D} (1) <i>Dis. abt. 0.05 g. in 2 cc. HCl (sp. gr. 1.20) & boil for 1 min. Dil. w. 15 cc. c. aq. Filter. Add 1 drop CrO_3; mixt. (Vol. I, p. 147). Sol. becomes VRS1. — (2) Add 3 drops HNO_3 (sp. gr. 1.42) to a few mg. of the dry poud. & warm. A YO sol. forms fr. which nitrophenacetine separates on cooling. \textcircled{D} <i>Boil 0.1 g. substance for 30 sec. w. 2 cc. dil. HNO_3 (1 vol. HNO_3, sp. gr. 1.20 + 5 vol. aq.). Cool. Wash ppt. on filter w. 5 cc. c. aq. Recryst. fr. 8 cc. boiling aq. Wash w. 3 cc. c. aq. Dry 15 min. on tile at 60°, & 20 min. in 100° oven. The product, nitrophenacetine, is obtained in YO-YO ndl., m.p. 103° u.c. (104° c.). [Arch. Pharm., 220 (1891), 208.]</i></i>
2106	134–5; (120)	1,2-Naphthocarbazole, $\text{C}_{10}\text{H}_{11}\text{N}$. — \textcircled{P} Sol. fluor. blue! — E.s. alc., eth., bz. Colors pine splinter violet in T. 2.24. — Ndl. fr. lgr. — B.Pk, ruby-red ndl. fr. bz., m.p. 174–5° d.
2107	134–5	Acetonylacetoxime, $\text{Me.C:(NOH).CH}_2.\text{CH}_2.\text{C:(NOH)Me}$. — \textcircled{P} Oxime. §— Lft. fr. bz. V.d.s. h. bz.; e.s. alc., eth., ac., alk., h. aq.
2108	134–5	† Salicylanilide, o-HO.C ₆ H ₄ .CO.NH.Ph. — \textcircled{P} FeCl_3 gives violet color w. alc. sol.! — Lft. fr. h. aq. V.d.s. h. aq.; e.s. alc., eth., chlf., bz. Dist. alm. unded. — \textcircled{D} Slowly sapd. by boiling w. dil. alk. (aniline & salicylic ac.).
—	134.5	Quinolinic Acid Anhydride. — Cf. No. 2.122–1.
2109	135–6	Methylethylketonesemicarbazone, $\text{Me.C:(N.NH.CO.NH_2).Et}$. — \textcircled{P} Gives semicarbazone reaction, T. 2.17. — Lft. E.s. alc., h. aq.
2110	135	Acetophenine, $\text{C}_{10}\text{H}_7\text{N}$. — [Fr. acetophenone, NH ₂ & P ₂ O ₅ .] — Ndl. fr. alc. V. weak base. Not attacked by CrO_3 .
2111	135–6	s-Diisobutylurea, $(\text{C}_4\text{H}_9)_2\text{NH.CO.NH(C}_4\text{H}_9)_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: isobutylamine (No. 2.1078); CO ₂ . — Ndl. i. aq.; v.s. alc.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2112	136	tert. Butyl Phenylcarbamate, $\text{Ph.NH.CO}_2\text{C.Me}_3$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; tert.-butyl-amine (No. 2.1066); CO_2 . — Ndl. Alm. i. aq.; d.s. pet.-eth.; s. alc., eth., bz.
2113	136	Triphenylurea, Ph.NH.CO.NPh_3 . — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; diphenylamine (No. 2.1568); CO_2 . Gives the products just named on dry distn.!
2114	136	α -Hydroxyisobutyranilide, $\text{Me}_2\text{C(OH).CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; α -hydroxyisobutyric ac. ($\text{C}_4\text{H}_6\text{O}_3$, Vol. I). — Rhomb.tbl. fr. aq. I. bz.
2115	136	p-Toluidino-p-acettoluide, $\text{Me.C}_6\text{H}_4\text{CH}_2\text{CO.NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); p-toluidinoacetic ac. (No. 2.85). — Lit. V.d.s. h. aq.; e.s. eth., h. alc.
2116	abt. 136	Diacetyl piperazine, $[\text{N}(\text{C}_2\text{H}_5\text{O}).\text{CH}_2.\text{CH}_2.\text{N}(\text{C}_2\text{H}_5\text{O}).\text{CH}_2.\text{CH}_2]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: piperazine (No. 2.765); ac. ac. — Tbl. E.s. aq., alc.
2117	136	Triphenylmethanehydrazobenzene, $\text{Ph}_3\text{C.NH.NH.Ph}$. — \textcircled{P} Prob. reduces Tollen's reagt. directly (being v. easily oxidized to triphenylmethaneazo-benzene). — Cryst. fr. eth.
2118	136	β -Benzaldehydophenylhydrazone, Ph.CH:N.NHPh . — \textcircled{P} Hydrazone. § — Labile isomer of No. 2.2220 to which it is quite similar in reactions, but 3 times more s. in alc. Ndl. fr. gl. ac. ac. Changes to α -form slowly on keeping or warming, or quickly in presence of little KOH in alc. sol.
2119	136	6-Nitro-1,3-acettoluide, $\text{NO}_2\text{C}_6\text{H}_3\text{Me}(\text{C}_2\text{H}_5\text{O})$. — \textcircled{P} Nitro comp.‡ — I. c. aq.; e.s. alc. — \textcircled{D} Sapn. T.* products: 6-nitro-1,3-toluidine (No. 2.3261) ac. ac.
2120	136	p-Nitrotriphenylcarbinol, $\text{Ph}_3\text{C(OH)(C}_6\text{H}_4\text{NO}_2)$. — \textcircled{P} Nitro comp.‡ — Cryst. mass fr. dil. ac. ac. V.d.s. lgr.
2121	137	8-Nitroquinaldine, $\text{NO}_2\text{C}_9\text{H}_7\text{N}$. — \textcircled{P} Nitro comp.‡ — Pale yellowish ndl. fr. dil. alc. D.s. c. aq.; e.s. alc., eth., bz. — B.HCl , glassy ndl. fr. alc. + HCl, losing HCl w. aq.
2122	137	Dinitrobenzyltoluene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{C}_6\text{H}_3(\text{NO}_2)\text{Me}$. — \textcircled{P} Nitro comp.‡ — Ndl. d.s. c. alc., eth.; e.s. chlf., bz.
2123	137	Benzophenonephenylhydrazone, $\text{Ph}_2\text{C:N.NH.Ph}$. — \textcircled{P} Hydrazone. § — Lust. ndl. fr. alc. I. aq.; r.d.s. h. alc.; v.d.s. c. alc.
2124	137-8	α -Benziloxime, Ph.C:(NOH).CO.Ph . — Oxime. § — Pearly lft. fr. h. 30% alc. E.s. c. alc., eth., chlf., less s. bz.; v.d.s. lgr.; e.s. dil. alk. — Heated 2 h. at 135° gives γ -deriv., cryst. fr. bz., m.p. 113-4° after losing $\frac{1}{2}$ mol. cryst. aq. & first melting at 70°.
2125	137	Hydrocyanocarbodiphenylimide, $(\text{Ph.NH})(\text{Ph.NH})\text{C.CN}$. — \textcircled{P} Pour few drops of sol. in conc. H_2SO_4 into aq. & add NaOH sol. An intense but transient blue color appears! — Cryst. fr. alc. or bz. I. aq.; e.s. alc., eth., bz. — Sapn.* w. HCl gives aniline, NH_3 & oxalic ac.
2126	137	1,3-Di-p-tolylphenylenediamine, $\text{C}_8\text{H}_4(\text{NH.C}_6\text{H}_4\text{Me})_2$. — Ndl. D.s. c. alc., eth. bz. — Nitroso deriv.‡ (fr. gl. ac. ac. + NaNO_2), yel. ndl. fr. lgr. + alc., d. 150°.
2127	137.5	Lacto- β -naphthalide, $\text{Me.CH(OH).CO.NH.C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: β -naphthylamine (No. 2.781); benzoic ac. (Vol. I). — Ndl. fr. dil. alc.
2128	138.5	Formyl- α -naphthalide, $(\text{CHO})\text{NH.C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: α -naphthylamine (No. 2.589); formic ac. (Vol. I). Long ndl. fr. aq.; e.s. h. aq.
2129	138.5	† Ethyl p-Nitrocinnamate, $\text{NO}_2\text{C}_6\text{H}_4\text{CH:CH.CO.Et}$. — \textcircled{P} Nitro comp.‡ — \textcircled{D} Sapn. T.* products: p-nitrocinnamic ac. (No. 2.488); ethyl alc. — Fine ndl. Alm. i. c. alc., eth.
2130	138-9	2-Acetamino-1,4-Xylene, $(\text{C}_6\text{H}_5\text{O})\text{NH.C}_6\text{H}_4\text{Me}$. — \textcircled{P} Nitro comp.‡ — \textcircled{D} Sapn. T.* products: 2-amino-1,4-xylene (No. 2.1315); ac. ac.
2131	138	p-Nitrobenzophenone, $\text{Ph.CO.C}_6\text{H}_4(\text{NO}_2)_2$. — \textcircled{P} Nitro comp.‡ — Lft. fr. alc.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2132	138c.	Perseite Heptanitrate, $C_7H_9(NO_3)_7$. — \textcircled{P} Nitrate.‡ — Ndl. fr. alc.; s. h. alc. (Prob. explosive.)
2133	abt. 139	2,3,6-Trinitro-1,4-xylene, $(NO_2)_3C_6H_4Me$. — Nitro comp.‡ — Monoclin. ndl. Alm. i. aq. Boiled 3 days w. alc. ammon. gives a dinitroxylidine, cryst. fr. gl. ac. ac., m.p. 203°.
2134	abt. 139u.c.	o-Toluamide, $Me.C_6H_4CONH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; o-toluidic ac. (Vol. I). — Ndl. E.s. alc., warm eth., conc. HCl; d.s. c. aq.; e.s. h. aq.
2135	140d.	Nitroanthrone, $[C_6H_4.CH(NO_2)_2]_2CO$. — \textcircled{P} Sol. in alc. KOH w. deep or-red color! Becomes violet at 100° & in melting evolves nitrous fumes yielding anthraquinone (Vol. I). — Lust. ndl. fr. bz. E.s. h. bz.; less s. h. alc.
2136	140	† Ethoxycaffeine, 1,3,7-Trimethyl-2,6-oxy-8-ethoxypurine, $C_{10}H_{14}O_2N_6$. — \textcircled{P} Gives murexide react. (T. 2.20)! — Ndl. D.s. aq., eth., c. alc.; e.s. h. alc.; i. alk. Dist. alm. undec.
2137	139-40	Colchicine, Acetotrimethylcolchicinic Ac., $(MeO)_3C_{18}H_9(NH.CO.Me)CO_2H$. — [Fr. Colchicum autumnale]. — \textcircled{P} Sol. in alk., ammon. & alk. carbonates w. intense yellow color! — Lust. ndl. fr. aq. Loses aq. at 140°, but not at 100°; then softens at 161° & melts at 172°. V.d.s. c. aq.; v.s. alc., chlf.; alm. i. abs. eth., bz. — Gives same color reactions as colchicine (No. 2.2152), fr. which it differs in being extracted fr. ac. sol. by bz., solubility in aq., & in being cryst. ["Reacts neut."]
2138	139-40	Nitroso- β -dinaphthylamine, $(C_{18}H_{17})_2N.NO$. — \textcircled{P} Prob. gives deep blue or green-blue color in T. 2.15. — Colorless ndl. fr. bz. R.d.s. alc., eth.; e.s. bz.
2139	140	Tetranitrobiphenyl, $(NO_2)_4C_{12}H_6$. — \textcircled{P} Nitro comp.‡ — Amorph. I. aq.; d.s. alc.; s. or r.d.s. eth.
2140	137-9.5; 141-2	o-Oxycarbanil, Carbonylaminophenol, $[NH.CO.O.C_6H_4]^+$. — Lust. plates fr. h. dil. HCl. B.p. above 360°. D.s. c. aq.; e.s. alc.; less s. eth.; v.s. alk. Decd. by HCl above 150° to CO ₂ & o-aminophenol.
2141	140-1	Diisobutylacet-p-toluide, $(C_6H_5)_2CH.CO.C_6H_4Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); diisobutylacetic ac. — D.s. c. pet.-eth.; i. aq.
2142	141	Isobutylurea, $NH_2.CO.NH(C_6H_5)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; isobutylamine (No. 2.1078); CO ₂ . — Ndl. fr. acetone. D.s. acetone.
2143	141-2	Pyromucamide, $C_6H_5O.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; pyromucic ac. (Vol. I). — Cryst. Subl. fr. 100° in ndl. — Treated w. Br-aq., & then w. NaOH sol., gives charac. dark blue coloration! — W. 3.6 pt. Br at 0° gives a tetrabromide, cryst. fr. ac.-eth., m.p. 121° d.
2144	141	Phenyl Carbamate, Ph.CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; phenol (Vol. I), CO ₂ . — Lft. D.s. c. aq.; more s. h. aq.; e.s. alc., eth.
2145	141	Acrylo-p-toluide, $CH_2:CH.CO.C_6H_4Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); acrylic ac. (Vol. I).
2146	141-2u.c.	† Benzophenoneoxime, Ph.C(:NOH).Ph. — \textcircled{P} Oxime.§ — Silky ndl. fr. h. dil. alc. V.d.s. c. aq.; v.s. eth.; s. alk. & pptd. by alk. fr. sol.; s. conc. HCl & pptd. by aq.; s. bz., chlf., lgr.
2147	140-2	Phenyl- α -naphthylketoxime, Ph.C(:NOH).C ₁₀ H ₇ . — \textcircled{P} Oxime.§ — Ndl. fr. dil. alc.
2147-I	141-2	α -Anilinopalmitic Ac., $C_{16}H_{29}CH(NH.Ph)CO_2H$. — Powd. I. aq.; d.s. alc.; more s. eth., bz.
2148	141	5-Nitrophthalide, $[CO.C_6H_4(NO_2)_2]CH_2O$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. I. c. aq.; Na ₂ CO ₃ ; r.d.s. alc., eth.; v.s. h. chlf., bz. — Sn + HCl reduces to aminophthalide, pr. fr. chlf., m.p. 178°.
2149	140-2; 145	p-Nitrodesoxybenzoin, $NO_2C_6H_4CO.CH_2Ph$. — \textcircled{P} Nitro comp.‡ — Pr. S. in 597 pt. c. or 22.5 pt. boiling 95% alc.; v.d.s. h. eth. Gives violet coloration w. alc. KOH!
2150	141	6-Phenylindanone(7), $[C_6H_5CO.CHPh.CH_2]^+$. — Ndl. fr. alc.
2151	142	Diphenyl- α -naphthylamine, $Ph_2N.C_10H_7$. — Silky ndl. fr. h. dil. alc. E.s. eth., bz. B.p. 335-40° (80-85 mm.).

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2152	142.5; 143-7	† Colchicine, $C_{22}H_{22}O_6N$. — [Highly toxic alkaloid fr. <i>Colchicum autumnale</i> L., or meadow saffron.] Amorph. powd., or lft. containing chlf. of cryst. in case of "cryst." commercial product. Houdé (Bull. soc. chim., 42, 298) claims to have obtained it colorless; but commercial preparations are of a light broken yellow, becoming darker on exposure to light. Odor like moist straw. Taste v. bitter! (Dose, 0.0005 g.) — S. at 25° in 22 pt. aq., 155 pt. eth., 87 pt. bz.; s. at 80° in 20 pt. aq.; v. s. alc., chlf.; i. pet.-eth.; s. in strong min. ac., or dil. alk. w. intense yel. color! — Levorotatory. — [Is best extracted fr. ac. aq. sol. by chlf. after removal of impurities by pet.-eth., & remains after evapn. of solvents as sticky varnish-like residue.] ② (1) Dis. 1 mg. in 3 drops H_2SO_4 (sp. gr. 1.84) on crucible cover. To the yel. sol. add 1 small drop HNO_3 (sp. gr. 1.42). The color changes more or less rapidly according to manner of mixing, going within 15 min. through green, blue, red-violet, & or-yel. to pale yel. Addition of conc. $NaOH$ in sl. x.s. then gives a strong or.-red. — (2) T. 2.2-b w. HNO_3 gives VRS ₂ color, changing after 5 min. to ROS ₂ , and then slowly to OYS ₁ . Addition of 6 drops $NaOH$ sol. then gives brilliant OR. — (3) Tannic ac. sol. gives floc. white ppt., e.s. ac. ac., fr. dil. aq. sol.
2153	142-3	Methyl- β -naphthylketoxime, $Me.C.(NOH)C_{10}H_7$. — ② Oxime. §
2154	142	p-Nitrophenyl Benzoate, $NO_2.C_6H_4.CO_2Ph$. — ② Nitro comp. † — Cryst. Alm. i. aq.; e.s. h. alc.
2155	142	m-Tolylurea, $NH_2.CO.NH.C_6H_4.Me$. — ② ③ Sapn. T.* products: NH ₂ ; m-toluidine (No. 2.1265); CO ₂ . — Lft. fr. aq.
2156	142-3 u.c.	† o-Benzotoluide, $Ph.CO.NH.C_6H_4.Me$. — ② ③ Sapn. T. 2.26-b w. sulfuric ac. gives benzoic ac. & o-toluidine (No. 2.1262). — Slender ndl. fr. dil. alc. Odorless. I. c. aq.; s. alc., conc. HCl. — Gives strong carbarylamine odor in T. 2.12.
2157	140.5; 144.5	5-Acetamino-1,3-Xylene, $(C_2H_5O)NH.C_6H_4.Me$. — ② ③ Sapn. T.* products: 5-amino-1,3-xylene (No. 2.1321); ac. ac.
2158	143	Glycolyl-p-toluide, $CH_2(OH).CO.NH.C_6H_4.Me$. — ② ③ Sapn. T.* products: p-toluidine (No. 2.566); glycollic ac. (Vol. I). — Ndl. fr. h. aq. E.s. aq., chlf., bz.; v.s. alc.; d.s. eth., lgr.
2159	143	Acet- α -naphthylhydrazine, $(C_2H_5O)NH.NH(C_10H_7)$. — ② ③ Sapn. T.* products: α -naphthylhydrazine (No. 2.804); ac. ac. — Ndl. fr. dil. alc. E.s. alc., eth., bz. [Possibly gives Ag reduction in T. 2.17-a.]
2160	143	Methyldibenzoylhydrazine, $Me(Ph.CO).N.NH(Ph.CO)$. — ② ③ Sapn. T.* products: methylhydrazine (No. 2.1093); benzoic ac. — Ndl. fr. alc. E.s. alc., dil. alk.; less s. aq.; alm. i. eth. — [Possibly gives Ag reduction in T. 2.17-a.]
2161	143-4 u.c.	† Salicycaldehydophenylhydrazone, $\alpha-OH.C_6H_4.CH:N.NH.Ph$. — ② Reduces Tollen's reagt. in T. 2.30 (giving heavy ppt.) after shaking for 1 min.! Gives yel. coloration to conc. H_2SO_4 , & yel. sol. in h. 10% NaOH! — Fine GYT ₃₋₄ ndl. fr. h. 67% alc. Exposed 15 min. to direct sunlight becomes YOT ₁₋₂ , color disappearing after 5 min. heating at 100°! (Phototropy.) — I. c. aq.; s. h. aq., alc.; s. eth.; e.s. chlf. — Odorless. Tasteless.
2161-I	143	Anhydroformaldehydianiline, $(Ph.N.CH_2)_2$. — [May reduce Tollen's reagt. before or after treatment of T. 2.17, since it is said to give trioxymethylene & aniline w. aq. in tube at 100°.] — Silky lft. I. aq.; v.d.s. alc.; d.s. eth.; e.s. chlf., bz. Sbl., giving cryst., m.p. 177-8°. Reduction w. Zn dust & HCl gives methylaniline (No. 2.1249). Sol. in HCl soon reddens.
2162	143	Tanacetonesemicarbazone, $C_9H_{16}:N.NH.CO.NH_2$. — ② Semicarbazone. §
2163	142-4	Dibenzalacetoneoxime, $(Ph.CH:CH)_2.C:NOH$. — ② Oxime. § — Treatment w. ac. anhydride for several hr. at 100° gives acetyl deriv., ndl. fr. alc., m.p. 93-4°.
2164	143.5	2,5-Diphenylpyrrole, $[NH.CPh:CH.CH:CPh]$. — ② Vapors color pine splinter dark red in T. 2.24-a! — Lust. lft. fr. gl. ac. ac. I. aq., alk.; e.s. alc., eth. — Sol. in conc. H_2SO_4 becomes red-violet & then fluor. blue on warming!

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2165	144	3,6-Dinitro-1,2,4,5-tetraethylbenzene, $(NO_2)_2C_6H_4Et_4$. — \textcircled{P} Nitro comp.‡ Silky ndl. fr. alc. I. aq.
—	144	3,4-Dinitroacetanilide. — Cf. No. 2.3309. (Yellowish.)
2166	144	6,8-Dinitroquinoline, $(NO_2)_2C_6H_3N$. — \textcircled{P} Nitro comp.‡ — Mic. ndl. D.s. h. aq.; v.s. h. alc. — Sol. in h. alkalies w. dark red color!
2166-1	144d.	a-Phenyl-b-hydroxyurea, $Ph.NH.CO.NH(OH)$. — \textcircled{P} Reduces Fehling's sol. (& prob. Tollen's reag.)! — $FeCl_3$ colors alc. sol. deep blue-violet. — Powd. V.d.s. "ordinary solvents." — \textcircled{D} Boiled w. alc. gives d.s. ndl. of diphenyloxybiuret, m.p. 178°.
2167	144	Di-p-tolylparabanic Ac., $[CO.N(C_6H_5Me).CO.CO.N(C_6H_5Me)]^2$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. alc. KOH) products: p-toluidine (No. 2.566); oxalic ac.; CO_2 . — Lft. Alm. i. aq.; e.s. alc.
2168	144-5	Glutaranil, $[CO.CH_2CH_2CH_2CO.NPh]^2$. — \textcircled{P} \textcircled{D} Prob. Sapn. T.* products: aniline; glutaric ac. (Vol. I). — Ndl. fr. alc. E.s. alc.; alm. i. eth.
2169	145	Formylphenylhydrazine, $(CHO)NH.NPhH$. — \textcircled{P} Should reduce Tollen's reag. in T. 2.17, if not in T. 2.30. — Lust. lft. fr. alc. R.d.s. c. aq.; v.s. h. alc. — Sapn. T.* products: phenylhydrazine (No. 2.1369); acetic ac.
2170	146	Indanone-(1)-oxime, $[C_6H_4.C(:NOH).CH_2CH_3]^2$. — \textcircled{P} Oxime.§ — Lust. ndl. fr. alc.
2171	146. 2u.c. (148. 5c.)	4-(γ)-Oxypyridine, Pyridone, $[NH.CH:CH.CO.CH:CH^2]$. — \textcircled{P} Cryst. w. H_2O (m.p. 62°; 66-7°), lost over H_2SO_4 in <i>vacuo</i> . — S. in 1 pt. c. aq.; e.s. alc.; alm. i. eth. — Dist. a. 350°. — $B.HgCl_2$, cryst. ppt., d.s. aq.
2172	146	Diphenyl-p-phenylenediamine, $(Ph.NH)_2C_6H_4$. — \textcircled{P} Sol. in conc. H_2SO_4 colored cherry-red by little KNO_2 . — Silvery lft. Alm. i. dil. acids; e.s. h. bz., eth., chlf.; less s. h. alc. — Diacetyl deriv., cryst., e.s. h. bz., m.p. 191.7°.
2173	146u.c.	† Acet-p-toluide, $(Me.CO)NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. by T. 2.26-a (w. HCl) gives p-toluidine (No. 2.566), & ac. ac. (Recryst. the toluidine by spontaneous evapn. of lgr. sol. before taking m.p.) — Ndl. fr. alc. B.p. 307°. 1000 pt. aq. at 22° dis. 0.886 pt.; 100 cc. 95% alc. dis. 10.77 g. at 25°. Odorless. Tasteless. Gives carbylamine odor (but less readily than acetanilide) in T. 2.12.
2174	147	Tetraphenylhydrazine, $Ph_4N.NPh_3$. — \textcircled{P} Sol. w. deep blue color in conc. H_2SO_4 . — Trimet. pr. fr. 1 vol. chlf. + 5 vol. alc.
2175	146. 5-7u.c.	† Phenylurea, $Ph.NH.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; aniline (No. 2.1235); CO_2 . — Odorless, tasteless, monoclin. cryst. D.s. c. aq.; e.s. h. aq.; e.s. alc., eth. Gives powerful carbylamine odor in T. 2.12. — \textcircled{D} Is easily converted into carbanilide by boiling w. aniline by procedure described for phenylurethane (No. 2.1544).
2176	147	α-Triphenylbiuret, $Ph.NH.CO.NPh.CO.NH.Ph$. — \textcircled{P} \textcircled{D} Sapn. T.* products & other derivatives same as described for No. 2.2175 above. — Pr. i. aq. — Strongly heated, dec. to phenylisocyanate (v. irritating odor) & s-diphenylurea.
2177	147; (141. 5)	Cinnamamide, $Ph.CH:CH.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; cinnamic ac. (Vol. I). — Cryst. D.s. c. aq.; e.s. h. alc.
2178	147	Methyl 2,6-Dinitrobenzoate, $(NO_2)_2C_6H_3CO.Me$. — \textcircled{P} Nitro comp.‡ — Plates fr. alc. E.s. h. alc. — Sapn. T.* products prob.: 2,6-dinitrobenzoic ac. (No. 2.340); methyl alc. (Vol. I).
—	147	Dinitrobenzil. — Cf. No. 2.3326.
2179	147	p-Nitrobenzonitrile, $NO_2C_6H_4CN$. — \textcircled{P} Nitro comp.‡ — Lft. fr. alc. D.s. c. aq. alc.; e.s. h. alc., chlf.
2180	147c.	m-Hydroxybenzaldehydophenylhydrazone, $HO.C_6H_4.CH:NH.Ph$. — \textcircled{P} Hydrazone.§ — Colorless pr. fr. toluene quickly becoming brownish yel. in light! — E.s. chlf., h. toluene, KOH; i. lgr.
2181	147-8	3-Methyl-2-acetylindole, Acetyliskatole, $Me(C_2H_5O).C_6H_4N$. — Long hair-like ndl. fr. dil. alc. I. c. aq.; e.s. alc. R. vol. w. st. Boiled w. conc. HCl gives skatole (No. 2.1830). — \textcircled{D} Picrate, cryst. fr. bz. in long or. yel. ndl., m.p. 156-7°.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2182	147-9	d-(& l)-Gulonic acidphenylhydrazide, $\text{CH}_2(\text{OH}).(\text{CH}.\text{OH})_2\text{CO.NH.NH.Ph.}$ — \textcircled{P} Hydrazide. § — Dec. at 195°.
2183	148-9	3,3'-(β)-Dinitrobenzophenone, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{CO.C}_6\text{H}_4(\text{NO}_2)_2$. — \textcircled{P} Nitro comp. † — Cryst.
2184	147-9	Ethylene Carbamate, $\text{NH}_2\text{CO}_2\text{CH}_2\text{CH}_2\text{CO}_2\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; ethylene glycol (Vol. I). — Cryst. flocks. E.s. h. aq., alc.; d.s. eth.
2185	148-9	m-Acetaminophenol, $(\text{C}_6\text{H}_5\text{O})\text{NH.C}_6\text{H}_4\text{OH}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: m-aminophenol (No. 2.830); ac. ac. — Ndl. fr. aq. E.s. aq., alc.; d.s. eth., chlf.
—	149	a-Diethylsemicarbazide. — Cf. No. 2.890-1.
2187	149; 152	p-Nitrophenylacetylene, $\text{NO}_2\text{C}_6\text{H}_4\text{C}\text{:CH.}$ — \textcircled{P} Gives brick red ppt. w. ammon. CuCl sol. which explodes when carefully dried & heated! Odor strongly cinnamon-like! — Fine ndl. fr. h. aq. Vol. w. st. V.d.s. c. aq.; s. h. aq.; e.s. alc., eth.
2188	149-50	† 6-Nitroquinoline, $\text{NO}_2\text{C}_6\text{H}_4\text{N}$. — \textcircled{P} Nitro comp. † — Cryst. w. zH_2O (lost in the air). Sbl. undecd. D.s. c. aq., alc., eth., lgr.; v.s. bz. — $\text{B}_2\text{H}_6\text{PtCl}_6$, cryst. ppt.; yel. ndl. fr. h. dil. HCl.
2189	149-50	Sarcosine Anhydride, $[\text{CO.NMe.CH}_2\text{CO.NMe.CH}_2]$. — B.p. 350°. V.s. aq.; e.s. alc., eth. Taste bitter. Boiled w. HCl gives sarcosine (No. 2.2456-1).
2190	150 u.c.	Acet-m-nitroanilide, $(\text{C}_6\text{H}_5\text{O})\text{NH.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Nitro comp. † — Colorless lft. fr. alc. S. h. but not c. aq. — Sapn. T.* products: p-nitroaniline (No. 2.3319); ac. ac.
2191	150-1	4-Nitroacet-o-toluidine, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}(\text{Me.CO.NH})$. — \textcircled{P} Nitro comp. † — Yellowish white ndl. Sapn. T.* products: 4-nitro-o-toluidine (No. 2.3098-1); ac. ac.
2192	150	ab-Methylphenylurea, MeNH.CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: methylamine; aniline; CO_2 . [“8 hr. heating w. conc. HCl gives only partial splitting.”] — Lust. lft. fr. h. aq.
2193	150-1	† Cinnamylanilide, $\text{Ph.CH:CH.CO.NH.Ph.}$ — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; cinnamic ac. (Vol. I).
2194	150	p-Tolylsuccinimide, $[\text{CO.N(C}_6\text{H}_5\text{)}_2\text{CO.CH}_2\text{CH}_2]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); succinic ac. (Vol. I). — Ndl. fr. h. aq. I. c. aq. E.s. h. alc., eth., chlf. — B.p. 344-5° (733 mm.).
2195	150	Phthalimidine, $[\text{NH.CH}_2\text{C}_6\text{H}_4\text{CO}]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH; o-oxymethylbenzoic ac. ($\text{C}_6\text{H}_5\text{CO}_2$, Vol. I), or its anhydride, phthalide, (Vol. I). — Cryst. E.s. alc., eth., chlf. Oxid. by alk. KMnO_4 to phthalimide (No. 2.2555). — B.Pk, light yel. ndl. fr. alc., d.s. c. aq., m.p. 140°.
2196	151-2	† Glutarimide, $[\text{NH.CO.(CH}_2)_2\text{CO}]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH; glutaric ac. (Vol. I). — Lust. scales fr. alc. Alm. i. eth. Sbl.
2197	151	Acetacetetherphenylsemicarbazone, $\text{Me.C(:N.H.CO.NHPh).CH}_2\text{CO}_2\text{Et}$. — \textcircled{P} Semicarbazone. § — Ndl. E.s. alc., eth.
2198	151-1.5	Fluorenonephenylhydrazone, $[\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{C(:N.NHPh)}]$. — \textcircled{P} Hydrasone. § — Lust. pr. fr. alc.
2199	150-1; 153	l-Arabinosephenylhydrazone, $\text{CH}_2(\text{OH}).(\text{CH}.\text{OH})_2\text{CH:N.NH.Ph.}$ — \textcircled{P} Hydrazone. § — Colorless ndl. S. at 25° in 85 pt. aq. or 30 pt. 90% alc. $[\alpha]_D = +2.5^\circ$ in sl. supersat. sol. in 80% alc.
2200	151-2	(α)-Benzoinoxime, $\text{Ph.C(:NOH).CH(OH).Me.}$ — \textcircled{P} Oxime. § — Mic. pr. fr. bz.
2201	152	Isorhamnonic acidphenylhydrazide, $\text{C}_6\text{H}_5\text{O}_5\text{N}_2\text{H}_2\text{Ph.}$ — \textcircled{P} Hydrazide. § — Ndl. fr. h. alc. E.s. aq.; d.s. acetone.
2202	152	Trinitrobenzene-naphthalene, $(\text{NO}_2)_3\text{C}_6\text{H}_4\text{C}_{10}\text{H}_8$. — \textcircled{P} Nitro comp. † — Fine white ndl. w. naphthalene odor. — On steam distn., naphthalene (Vol. I), passes into distillate. S. h. alc. w. some loss of naphthalene.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2203	152	Succinanil, $[CO.CH_2.CH_2.CO.NHPh]$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. alk.) products: aniline & succinic ac. w. the intermediate product, succinanilic ac. (No. 2.162). — Ndl. D.s. h. aq.; e.s. h. alc. B.p. abt. 400°.
2204	152-3	s-Diacetylurea, $(C_2H_4O)NH.CO.NH(C_2H_4O)$. — \textcircled{P} \textcircled{D} Sapn. T.* ("long boiling w. aq. NaOH") products: NH ₂ ; ac. ac.; CO ₂ . — Ndl. fr. alc. Sbl. undec. D.s. aq., alc.
2205	152-3	Cyclobutanecarbonamide, $[CH_2.(CH_2)_2.CH(CO.NH)]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; tetramethylenecarbonic ac. (C ₆ H ₁₀ O ₂ , Vol. I). — Sbl. Tbl. fr. eth. D.s. eth.
2206	153-4	Dipropionamide, $(Et.CO)_2.NH$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; propionic ac. (Vol. I). — E.s. alc.; i. lgr. B.p. 210-20°. — Silky ndl. fr. aq.
2207	153.5	Cuminamide, p-Me ₂ CH.C ₆ H ₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; cuminic ac. (Vol. I). — Tbl. i. c. aq.; d.s. h. aq.; misc. alc., eth.; e.s. h. bz.
2208	153-4	Trimethylacetamide, Me ₃ C.CONH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; trimethylacetic ac. (C ₄ H ₁₀ O, Vol. I). — Fine ndl. fr. aq. B.p. 212°. V. conc. HNO ₃ gives N ₂ O in the cold.
2209	153	Dibutylloxamide, $(C_4H_9)NH.CO.CO.NH(C_4H_9)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: butylamine (No. 2.1081); oxalic ac. — Ndl. fr. alc. V.d.s. aq.
2210	153-4	(α -p-Phenyltolylketoxime, Ph.C(:NOH).C ₆ H ₄ .Me. — \textcircled{P} Oxime. § — Ndl. fr. dil. alc. E.s. alc., eth., bz.
2211	152-5	Indanone(θ)-oxime, $[C_6H_5CH_2.C(:NOH).CH_2]$. — \textcircled{P} Oxime. § — Ndl. fr. dil. alc. E.s. alc., eth., chlf.
2212	153	b-Benzoyl-a-methylphenylhydrazine, Ph.CO.NH.NMePh. — \textcircled{P} Prob. gives Ag reduction in T. 2.17, being split to methylphenylhydrazine (No. 2.1336) & benzoic ac. by boiling w. conc. HCl. — Fine ndl. fr. alc. V.s. h. alc. Alm. i. aq., alk.; e.s. conc. HCl.
2213	153	2,2'-Dimethyl-5,5'-dinitrodiphenylmethane, Me(NO ₂).C ₆ H ₄ .CH ₂ .C ₆ H ₄ .(NO ₂).Me. — \textcircled{P} Nitro comp.‡ — I. aq.
2214	153-4	m-Nitrobenzanilide, NO ₂ .C ₆ H ₄ .CO.NH.Ph. — \textcircled{P} Nitro comp.‡ — Lft. fr. h. aq. V.d.s. c. aq.; e.s. alc., eth. Sbl. — Sapn. T.* products: m-nitroaniline (No. 2.3125); benzoic ac.
2215	153	Methylacet-p-nitroanilide, Me(C ₆ H ₄ O)N.C ₆ H ₄ .NO ₂ . — \textcircled{P} Nitro comp.‡ — Lft. fr. h. aq. — Sapn. T.* products: nitro-p-methylaniline (No. 2.3343); ac. ac.
2216	154	Nitrofluorene, C ₁₃ H ₈ .NO ₂ . — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. I. aq. — Oxid. in gl. ac. ac. sol. by CrO ₃ to p-nitrodiphenyleneketone, m.p. 217-8°.
2217	153-5d.	d,L-Gulonicacidphenylhydrazide, CH ₂ (OH).(CH.OH) ₄ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. § — Ndl. fr. abs. alc. D.s. c. aq.; e.s. h. aq.
2218	154	s-Diallyloxamide, $(C_4H_9)_2NH.CO.CO.NH(C_4H_9)_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: allylamine (No. 2.1068); oxalic ac. — B.p. 274° d.
2219	154	Ethyldeneurea, $[CO.NH.CHMe.NH]$. — \textcircled{P} \textcircled{D} (Conc. HCl splits to acetaldehyde & urea.) Boil w. HCl. Pass vapors into 10% NaOH & boil sol. (T. 1.111-1 for acetaldehyde.)
2220	154.5-5.5c.; 156u.c.	† Benzaldehydophenylhydrazone, Ph.CH ₂ :N.NH.Ph. — \textcircled{P} Hydrazone. § — YT4 cryst. fr. h. 50% alc. Exposure of dry comp. to direct sunlight for 15 min. changes color to OT1-2 (phototropy), the color discharging to OT3 on heating 15 min. at 100°.
2221	154-5	† Phenylacetamide, Ph.CH ₂ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; phenylacetic ac. (Vol. I). — Lft. E.s. h. aq.; v.d.s. c. aq.; e.s. alc.; v.d.s. eth. B.p. 281-4°. Aq. sol. dis. HgO giving comp. crystg. in fine ndl., m.p. 208°.
2222	155	Isoamylphenylurea, (C ₆ H ₁₁)NH.CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; isoamylamine (No. 2.1100); CO ₂ . — Ndl. fr. alc. I. aq.
2223	155-6	1-Methylcyclohexane-3-carbonamide, $[CHMe.CH_2CH(CO.NH_2).(CH_2)_2]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 1-methylcyclohexanecarbonic ac. (b.p. 245° c.). — Lust. ndl. fr. aq.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2224	abt. 155	p-Ethoxyphenylsuccinimide, "Pyrantin," $[CO.N(EtO.C6H4).CO.CH2.CH2}]$. — [An antipyretic.] — \textcircled{P} \textcircled{D} Sapn. T.* products: p-phenetidine (No. 2.1392); succinic ac. (Vol. I). — Pr. fr. alc. S. in 1317 pt. aq. at 17°, or 83.6 pt. at 100°; e.s. h. alc.; i. eth. — Dis. 0.05 g. in 2–3 cc. h. HCl. Dil. w. aq. Add 1 drop 3% CrO_3 sol. A ruby-red color appears. — Fuse w. KOH. Add bleaching powd. sol. to sol. of fusion. A red color, which increases on standing, develops. — \textcircled{D} Dis. 80 pt. I & 50 pt. KI in 50 pt. aq. Run into sol. of 60 pt. compound in 200 pt. warm gl. ac. ac. A ruby-red cryst. ppt. of $(C_{12}H_{14}O_3N)_2I_2KI$, m.p. 175°, forms on cooling.
2225	155	Pimelanilide, Ph.NH.CO.(CH ₂) ₄ .CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline (T. 2.1235); pimelic ac. (Vol. I). — Ndl. fr. dil. alc.
2226	155–5.5u.c.	† p-Benzotoluide, Ph.CO.NH.C ₆ H ₄ .Me. — \textcircled{P} \textcircled{D} Sap. by procedure prescribed for No. 2.2156 to benzoic ac. & o-toluidine (No. 2.1262)! — Scales or pr. fr. dil. alc. E.s. alc., conc. HCl; i. aq., alk. Odorless. Tasteless. Gives carbylamine odor in T. 2.12.
2227	155–6	Salicyl-p-toluide, o-HO.C ₆ H ₄ .CO.NH.C ₆ H ₄ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); salicylic ac. — Pr. fr. alc.
2228	155d.	d-Talonic acid phenylhydrazone, CH ₂ (OH).(CH.OH) ₄ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. § — Pr. E.s. aq.
2229	155–6	Acetonephenylsemicarbazone, Me ₂ C:N.NH.CO.NH.Ph. — \textcircled{P} Semicarbazone. § — Ndl. D.s. aq.; e.s. alc., eth.
2230	156	4,6-Dinitro-1,2,3,5-tetramethylbenzene, (NO ₂) ₂ C ₆ Me ₄ . — \textcircled{P} Nitro comp.‡ — Pr. fr. alc. D.s. c. alc.; e.s. h. alc.
2231	155–7	(β)-Methylhydantoin, $[CO.NMe.CH_2.CO.NH]$. — Pr. E.s. aq., alc. Sbl. Sol. dis. AgO giving sol. w. alk. reaction.
2232	157	Pinacoline semicarbazone, Me.C(:N.NH.CO.NH ₂).CMe ₂ . — \textcircled{P} Semicarbazone. § — Cryst. fr. aq. S. eth.
2233	157	d-Galactosediphenylhydrazone, CH ₂ (OH).(CH.OH) ₄ .CH:N.NPh. — \textcircled{P} Hydrazone. § — Pr. fr. aq.
2234	157.5–8	6-Nitro-acet-o-toluide, NO ₂ C ₆ H ₄ .Me.(Me.CO.NH). — \textcircled{P} Nitro comp.‡ — Pr. e.s. alc., eth. — Sapn. T.* products: 6-nitro-o-toluidine (No. 2.3018); ac. ac.
2235	157.5	Ethylene Phenylcarbamate, Ph.NH.CO.CH ₂ .CH ₂ .NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; ethylene glycol (Vol. I). — Pr. fr. alc.
2236	157	Benzoyl- β -naphthalide, Ph.CO.NH.C ₁₀ H ₇ . — \textcircled{P} \textcircled{D} Sapn. T.* products: β -naphthylamine (No. 2.781); benzoic ac. — Ndl. fr. bz. E.s. eth., chlf., h. alc.
2237	157	Benzyl Cyanurate, (Ph.CH ₂ .NCO) ₃ . — Ndl. fr. alc. I. aq.; d.s. eth. B.p. a. 320°. — Fusion w. KOH gives benzylamine (No. 2.1236) & CO ₂ .
2238	157–8	δ -Aminovaleric Ac., Homopiperidinic Ac., NH ₂ (CH ₂) ₄ .CO ₂ H. — Pearly lft. V.s. aq.; alm. i. abs. alc.; i. eth. — In melting forms anhydride, piperidone, m.p. 39–40°. — Heated w. solid NaOH gives NH ₂ butylamine (No. 2.1081); etc. — Benzoyl deriv. (by Schotten-Baumann method, T. 2.27), m.p. 94°.
2239	158	Dinitrosopiperazine, $[N(NO).CH2.CH2.N(NO).CH2.CH2}]$. — \textcircled{P} Prob. gives blue coloration in T. 2.15. — Lust. lft. fr. h. aq.; d.s. c. aq., eth.
2240	158d.	(α)-Styrene Nitrite, (C ₆ H ₅ O ₂ N ₂) ₂ . — \textcircled{P} May give blue coloration in T. 2.15. Dec. on fusion giving NO, benzonitrile, etc. — Cryst. fr. ac.-eth. — Heated in sealed tube w. aq. gives CO ₂ , NH ₃ , benzonitrile & benzoic ac.
2241	157–8; 160	ab-Propionylphenylhydrazine, Ph.NH.NH(Et.CO). — \textcircled{P} Prob. reduces Tollen's reagt. directly. — Lft. fr. chlf.
2242	158	Succinylphenylhydrazine, $[CO.CH2.CH2.CO.N(Ph.NH)]$. — \textcircled{P} Prob. reduces Tollen's reagt. after hydrolytic treatment. — Lft. e.s. h. aq., alc.
2243	158–9; 155	(α)-Benzoinphenylhydrazone, Ph.CH(OH).C(:N.NH.Ph).Ph. — \textcircled{P} Hydrazone. § — Abs. alc. at 20° dis. 2.2%; e.s. eth., chlf.; d.s. lgr.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2244	158; 160-2	d-Galactosephenylhydrazone, $\text{CH}_2(\text{OH}).(\text{CH}.\text{OH})_4.\text{CH}:\text{N}.\text{NH}.\text{Ph}$. — \textcircled{P} Hydrazone. § — Ndl. fr. alc. S. in 50 pt. c. aq.; e.s. h. aq.; s. in 10 pt. h. alc.; i. eth.
2245	158	3-Nitroacet-o-toluide, $\text{NO}_2.\text{C}_6\text{H}_4.\text{Me},(\text{C}_6\text{H}_4\text{O.NH})$. — \textcircled{P} Nitro comp. † — V.d.s. qq.; e.s. alc., bz., ac. ac.; d.s. eth. — Sapn. T.* products: 3-nitro-o-toluidine (No. 2.3041); ac. ac.
2246	158-9	p-Toluanide, $\text{Me}.\text{C}_6\text{H}_4.\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; p-tolue ac. (Vol. I). Cryst. fr. h. aq. D.s. c. aq., eth., bz.; e.s. h. aq., alc.
2247	158-9; 156; 160-1	Isophthalonitrile, m- $\text{C}_6\text{H}_4(\text{CN})_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; isophthalic ac. (Vol. I). — Ndl. D.s. h. aq.; r. more s. h. alc. Sbl.
2248	abt. 158	γ -Phenylhydantoin, $[\text{NPh}.\text{CO}.\text{NH}.\text{CH}_2.\text{CO}]$. — [Dehydration product fr. phenylureidoacetic ac. (No. 2.318); by boiling w. 25% HCl.] — S. in 105 pt. aq. at 10°; e.s. alc., bz.; d.s. eth.
2249	159	Acetylhydrazobenzene, $\text{Ph}.\text{NH}.\text{N}(\text{C}_6\text{H}_5\text{O})\text{Ph}$. — \textcircled{P} \textcircled{D} Decd. by heat to acetanilide (No. 2.1975); ac. ac. — Cryst. s. h. aq.; e.s. alc.
2250	159	Rhamnosephenylhydrazone, $\text{Me}(\text{CH}.\text{OH})_4.\text{CH}:\text{N}.\text{NH}.\text{Ph}$. — \textcircled{P} Hydrazone. § — Colorless lft. S. in 80 pt. c. aq.; s. alc.; i. eth. $[\alpha]_D^{20} = +54.2^{\circ}$ (in 1% aq. sol.; l = 2 dm.).
2251	159	Acet- α -naphthalide, $(\text{C}_8\text{H}_7\text{O})\text{NH}.\text{C}_8\text{H}_7$. — \textcircled{P} \textcircled{D} Sapn. T.* products: α -naphthylamine (No. 2.589); ac. ac. — Cryst. s. h. aq.; e.s. alc.
2252	159-60	Cyclopentanecarbonanilide, $[\text{CH}_2.\text{CH}(\text{CO}.\text{NH}.\text{Ph}).(\text{CH}_2)]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; pentamethylenecarbonic ac. ($\text{C}_6\text{H}_{10}\text{O}_2$, Vol. I). — Lust. pr. fr. alc. E.s. alc., chlf., bz.
2253	160d.	Arabinosecarbonamide, Amide of l-Mannonic Ac., $\text{CH}_2(\text{OH}).(\text{CH}.\text{OH})_3.\text{CO}.\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; l-mannonic ac. — Mic. ndl. V.d.s. c. aq.; e.s. h. aq.; i. alc., eth.
2254	160-1	Ethylenephenylurea, $[\text{CH}_2.\text{CH}.\text{NH}.\text{CO}.\text{NPh}]$. — \textcircled{P} \textcircled{D} Prob. sapn. T.* products: aniline; ethyleneglycol (Vol. I); CO_2 . — Lft. fr. h. aq. I. c. aq., lgr.; e.s. alc., bz.; d.s. eth.
—	160	"Dulcin." — Cf. No. 2.2313. (Very sweet.)
2255	160	Polymeric Carbodiphenylimide, $(\text{C}_6\text{H}_{10}\text{N}_2)_n$. — [Partially changed to carbani-lide (which is saponifiable to aniline & CO_2) by boiling w. alc. HCl.] Lft. w. bz. of cryst. fr. bz.; m.p. 154°. I. aq., abs. eth., lgr.; s. abs. alc. B.p. 235-6° (65 mm.).
2256	160-1u.c.	† Benzanilide, $\text{Ph}.\text{CO}.\text{NH}.\text{Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products by procedure (b.) w. H_2SO_4 gives: aniline (T. 2.1235); benzoic ac. (Vol. I). (Recryst. the benzoic ac. fr. 20 cc. boiling aq.). — Odorless, tasteless lft. fr. h. alc. Dist. undec. Gives carbylanine odor directly in T. 2.12.
2257	160	Phthalidanol, $[\text{CO}.\text{C}_6\text{H}_4.\text{CH}_2.\text{NPh}]$. — Lust. lft. fr. alc. V.d.s. h. aq.; d.s. eth.; e.s. chlf., bz. Is said not to be decd. by boiling ac. or alk.
2257-I	abt. 160d.	p-Acetaminophenol-phenacylether, "Hypnoacetin," $\text{Ph}.\text{CO}.\text{CH}_2.\text{O}.\text{C}_6\text{H}_4.\text{NH}.\text{CO}.\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* may give: p-aminophenol, No. 2.963); acetophenone (Vol. I). — Lft. fr. alc. Alm. i. aq., eth. S. in 353.8 pt. 85% alc. at 23.3°; d.s. chlf., bz.
2258	160-1	Furoinoxime, $\text{C}_6\text{H}_5\text{O.C(:NOH).CH(OH).C}_6\text{H}_5\text{O}$. — \textcircled{P} Oxime. § — Pr. d.s. c. alc.
2259	160	Tetranitrobenzyltoluene, $(\text{NO}_2)_4.\text{C}_6\text{H}_{10}$. — \textcircled{D} Nitro comp. † — Pr. fr. bz. I. aq.; d.s. c. alc., eth.
—	155-65	Formopyrine. — Cf. No. 2.2357. (Loses 1 H_2O at 130°.)
2260	161	Methyl p-Nitrocinnamate, $\text{NO}_2.\text{C}_6\text{H}_4.\text{CH}:\text{CH}.\text{CO}.\text{Me}$. — \textcircled{P} Nitro comp. † — Ndl. B.p. 281-6°. — Sapn. T.* products: methyl alc.; p-nitrocinnamic ac. (No. 2.488).
2261	161-2	d-Glucosedi phenylhydrazone, $\text{CH}(\text{OH}).(\text{CH}.\text{OH})_4.\text{CH}:\text{N}.\text{NPh}_2$. — \textcircled{P} "Reduces Fehling's sol." — Pr. fr. h. aq. V.s. h. aq., alc.; i. eth., chlf., bz.
2262	161	Xanthoneoxime, $[\text{O}.\text{C}_6\text{H}_4.\text{C}(:\text{NOH}).\text{C}_6\text{H}_4]$. — \textcircled{P} The yel. sol. in conc. H_2SO_4 fluor. blue! — Oxime. § Cryst. fr. alc.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2263	161-2; 159-60	Benzoyl- α -naphthalide, Ph.CO.NH.C ₁₀ H ₇ . — \textcircled{P} \textcircled{D} Sapn. T.* products: α -naphthylamine (No. 2.589); benzoic ac. (Vol. I). — Cryst. fr. alc.
2264	162	s-Dipropylloxamide, Pr.NH.CO.CO.NH.Pr. — \textcircled{P} \textcircled{D} Sapn. T.* products: propylamine (No. 2.1067); oxalic ac. — D.s. h. aq.; e.s. chlf.
2265	162	† Acetaldehydesemicarbazone, Me.CH:N.NH.CO.NH. — \textcircled{P} Semicarbazone. § — Ndl. or plates fr. aq. or alc. — \textcircled{D} Boil w. dil. H ₂ SO ₄ & identify acetaldehyde in dist. by T. 1.111.
2266	162	Undecanonedioxime (2.3), Me.C(:NOH).C(:NOH).C ₈ H ₁₇ . — \textcircled{P} Oxime. § — Tbl. fr. alc. I. lgr.
2267	162c.	4-Nitropyrazole, $^{\text{C}}$ NH.N:CH.C(NO ₂):CH ² . — B.p. 323° (th.i.). Flat ndl. fr. bz.
2268	162-3u.c.	Methylbiuret, Me.NH.CO.NH.CO.NH. — \textcircled{P} Gives the biuret react., T. 2.10. — Cryst. or spheroidal cryst. aggregates fr. alc. S. in 8 pt. h. aq.
2269	abt. 163d.	Mesityloxidesemicarbazone, Me ₃ C:CH.C(:N.NH.CO.NH).Me. — \textcircled{P} Semicarbazones § ("Split by acids" to its components). — Lft. E.s. alc.
2270	163-4	Suberonesemicarbazone, $^{\text{C}}\text{C}(:\text{N.NH.CO.NH}).(\text{CH}_2)_6\text{C}^{\text{2}}$. — \textcircled{P} Semicarbazone. § —
2271	163-4d.	1-Arabinosesemicarbazone, CH ₂ (OH).(CH.OH) ₅ .CH:N.NH.CO.NH. — \textcircled{P} Semicarbazone. § — Cryst. E.s. c. aq.; i. eth.; slowly s. h. alc.
2272	162-4	1-Ribonicacidphenylhydrazide, CH ₂ (OH).(CH.OH) ₅ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. § — Ndl. fr. abs. alc. S. aq.
2273	162-3u.c.	d-Lyxonicacidphenylhydrazide, CH ₂ (OH).(CHOH) ₅ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. § — Cryst. fr. alc. E.s. aq.; d.s. alc. (Cryst. w. 2H ₂ O, lost at 105°, m.p. 148-9°).
2274	163	4,4'-Dimethylbenzophenoneoxime, Me.C ₆ H ₄ .C(:NOH).C ₆ H ₄ .Me. — \textcircled{P} Oxime. § — Lust. pr. fr. alc.
—	163-4	3-Nitrophthalic Anhydride. — Cf. No. 2.226-2.
2277	162-4	2,2'-Hydrazonaphthalene, C ₁₀ H ₈ .NH.NH.C ₁₀ H ₈ . — \textcircled{P} May reduce Tollen's reagt. directly. Prob. e. oxidized to colored azo comp. — Flocks. I. aq.; d.s. alc.; e.s. eth.; bz.
2278	163d.	β -Benzylhydroxylaminecarbanilide, Ph.CH ₂ .N(OH).CO.NH.Ph. — \textcircled{P} FeCl ₃ gives blue ppt. in alc. sol. — Benzoylation w. benzoyl chloride & alk. (Cf. T. 2.27) gives deriv. crystg. fr. methyl alc., m.p. 120°.
2279	163.5	† Diphenylpiperazine, $^{\text{C}}\text{NPh.CH}_2\text{CH}_2\text{NPh.CH}_2\text{CH}_2\text{C}^{\text{2}}$. — Ndl. fr. alc. + acetone. I. aq.; d.s. c. alc. B.p. 300° d. — Gives charac. cryst. black dinitroso deriv. (Cf. Ber., 12, 1795).
2280	163	Malonphenylamide, NH ₂ .CO.CH ₂ .CO.NHPh. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; aniline; malonic ac. (Vol. I). — Fine ndl. fr. aq. or alc.
2281	163-4	5-Aacetamino-1,2,3-trimethylbenzene, (C ₆ H ₅ O)NH.C ₆ H ₃ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: 5-amino-1,2,3-trimethylbenzene (No. 2.669); ac. ac.
2282	163	Hydrocarbostyrene, o-C ₆ H ₅ .NH.CO.CH ₂ .CH ₂ C ² . — Pr. fr. alc. Alm. i. aq.; e.s. alc., eth., h. conc. HCl. V. stable. Dist. undecd. — B ₂ H ₂ PtCl ₆ .2H ₂ O, yel. tbl., m.p. 172°.
2283	164	Bis-Phenyldimethylpyrazolone, C ₂ H ₂ O ₂ N ₄ . — Pr. fr. gl. ac. ac. I. aq., dil. ac. or alk. Is pptd. unchanged by aq. fr. sol. in conc. H ₂ SO ₄ .
2284	164	Dinitroditolylmethane, (NO ₂) ₂ .C ₁₂ H ₁₄ . — \textcircled{P} Nitro comp.‡ — E.s. abs. alc., bz.; s. eth.
2285	164	5-Aacetamino-1,2,4-trimethylbenzene, Acetpseudocumidine, (C ₆ H ₅ O)NH.C ₆ H ₃ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: s-pseudocumidine (No. 2.646); ac. ac. — Ndl. E.s. h. aq.; v.s. chlf.
2286	164-5; 176-7	2-Formylamino-1,3-xylene, (CHO)NH.C ₆ H ₄ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: 2-amino-1,3-xylene (No. 2.1307); formic ac. (Vol. I). — Ndl. s. alc.
2287	165	ab-Phenyl-m-tolylurea, Ph.NH.CO.NH.C ₆ H ₄ .Me. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; m-toluidine (No. 2.1265); CO ₂ . — Ndl. fr. alc.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2268	165	ab-sec.-Butylphenylurea, Et.CHMe.NH.CO.NH.Ph. — ⊕ ⊖ Sapn. T.* products: aniline; sec.-butylamine (No. 2.1073). — Pr. fr. alc. I. aq.
2280	165	β-Diphenylbiuret, Ph.NH.CO.NPh.CO.NH. — ⊕ ⊖ Sapn. T.* products: aniline (T. 2.1235); NH ₃ ; CO ₂ . — Pr. E.s. alc.; i. aq.
2260	165	o-Hydrazotoluene, Me.C ₆ H ₄ .NH.NH.C ₆ H ₄ .Me. — ⊕ Reduces Tollen's reag. directly. — Lft. becoming yellowish in air by oxidn. to azotoluene. Alm. i. aq.; s. alc.
2281	165	Acetylsemicarbazide, Me.CO.NH.NH.CO.NH ₂ . — ⊕ Semicarbazide. § ("Sapd. by boiling aq., " and may reduce Tollen's reag. directly.) Cryst. e.s. aq., warm alc.; i. eth.
2292	165	tert.Butylphenylketoxime, Me.C ₆ C(:NOH).Ph. — ⊕ Oxime. § — (Readily split by heating w. conc. HCl. The corresponding ketone is an aromatic oil, b.p. 219–21°.) — Ndl. fr. alc. I. aq.; v.d.s. c. alc.
2253	165	Dibenzoylinethaneoxime, Ph.CO.CH ₂ .C(:NOH)Ph. — Glassy pr. fr. eth. D.s. c. alc. Transformed by action of dil. HCl or alk. to v. stable diphenylisoxazole, tbd. fr. alc., m.p. 140.5–141°!
2294	167	Cyclohexanonesemicarbazone, ¹² CH ₂ .C(:N.NH.CO.NH ₂).CH ₂ . — ⊕ Semicarbazone. §
2295	167	Benzoylcumidine, (Ph.CO.NH) ⁴ .C ₆ H ₄ .(CH.Me ₂) ² . — ⊕ ⊖ Sapn. T.* products: cumidine (No. 2.1334); benzoic ac. (Vol. I). — Ndl. fr. alc.
2295	167	s-Dibenzylurea, Ph.CH ₂ .CO.CH ₂ .Ph. — ⊕ ⊖ Sapn. T.* products: benzylamine (No. 2.1236); CO ₂ . — Ndl. i. aq.; e.s. alc.
2207	167	m-Hydroxybenzamide, HO.C ₆ H ₄ .CO.NH ₂ . — ⊕ ⊖ Sapn. T.* products: NH ₃ ; m-hydroxybenzoic ac. (Vol. I). — Lft. fr. h. aq. D.s. c. aq.; e.s. alc., eth., h. aq.; i. chlf., bz. — Taste bitter.
2298	167	o-Benzoylaminophenol, Ph.CO.NH.C ₆ H ₄ .OH. — ⊕ ⊖ Sapn. T.* products: o-aminophenol; benzoic ac. — Lft. Loses aq. in melting, giving benzylaminophenol, m.p. 103°. Lft. D.s. c. aq.; e.s. h. aq., alc., bz.; s. c. alk. & repnd. by ac.
2299	167	Benzimide, C ₂₂ H ₁₈ O ₂ N ₂ . — [Fr. benzaldehyde & alc. HCN.] — ⊕ Boiled w. HCl gives benzaldehyde & NH ₄ Cl! — Floc. mass. I. aq., KOH, HCl; mod. s. alc., eth.
2300	164; 169–70	N-Methylpyrrolidine-α-carbonic Ac., Hygric Ac., ¹² NMe.CH ₂ .CH ₂ .CH ₂ -CH(CO ₂ H) ² . — ⊕ Heated above m.p. decd., superheated vapors giving pyrrole react. to pine splinter soaked in HCl (Cf. T. 2.24)! Ndl. w. 1H ₂ O fr. moist chlf. V.s. aq., alc., h. chlf.; i. eth., bz. ["Reacts neut. in aq. sol."] Sol. dis. Ag ₂ O & gives Ag mirror on warming. — B.HCl, lft. fr. alc. + eth., v.s. aq. & h. alc., m.p. 187–8°.
2301	168	† ab-Benzoylphenylhydrazine, Ph.CO.NH.NH.Ph. — ⊕ Gives immediate heavy ppt. in T. 2.30 w. Tollen's reag.! — Cryst. D.s. h. aq., eth.; s. h. alc.
2302	167–9	p-Hydrazobiphenyl, Ph.C ₆ H ₄ .NH.NH.C ₆ H ₄ .Ph. — ⊕ Alc. sol. prob. reduces Tollen's reag. Soon oxid. by air to the azo comp. — Colorless lft. I. aq.; d.s. alc.
2303	168–9	Formylurea, (CHO)NH.CO.NH ₂ . — ⊕ ⊖ Sapn. T.* products: NH ₃ ; formic ac. (Vol. I). — Cryst. s. aq.; v.d.s. c. alc. Dec. in boiling aq. sol. to formic ac. & urea.
2304	168(r.h.)	Aminopropylmethylacetic Ac., (NH ₂ .CH ₂ .CH ₂ .CH ₂).CHMe.CO ₂ H. — ⊕ Melts w. efferv. giving anhydride (cubes fr. lgr., m.p. 53.5–55°) of coniine-like odor. — V.s. aq.; i. abs. alc., eth. — B ₂ H ₂ PtCl ₆ , lft. e.s. aq., dec. at 190°.
2305	165; 171	Phenylglycine-p-toluide, Ph.NH.CH ₂ .CO.NH.C ₆ H ₄ .Me. — ⊕ ⊖ Sapn. T.* products: p-toluidine (No. 2.566); phenylglycine (No. 2.99). — Cryst. i. aq.; e.s. alc., eth.
2306	169	Ethylideneacetamide, Me.CH(C ₂ H ₅ O.NH) ₂ . — ⊕ Boil w. dil. H ₂ SO ₄ & test distillate for acetaldehyde (T. 1.111). — Pr.
2307	169	Acetylphenylsemicarbazide, Me.CO.NH.NH.CO.NH.Ph. — ⊕ Should reduce Tollen's reag. in T. 2.17 & perhaps directly in T. 2.30. — Cryst. V.d.s. c. aq., eth.; e.s. h. aq., alc.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2308	169	s-Tetramethylethylene Cyanide, $(\text{CN})\text{Me}_2\text{C}(\text{CN})\text{Me}_2$. — \oplus Prob. gives NH_2 in T. 2.7. — Lust. lft. fr. dil. alc. V. volat.
2309	169	Dehydroacridine, $\text{C}_8\text{H}_4\text{CH}_2 : \text{C}_8\text{H}_4\text{NH}^+$. — $\oplus \ominus$ CrO_3 , mixt. (Vol. I) oxid. at once to acridine (No. 2.3102). — Cryst. fr. alc. I. aq.; d.s. c. alc.; e.s. h. alc., eth. Sbl. ["Does not unite w. acids."]
2310	169–70	Diphenyloxamide, $\text{Ph}_2\text{N.CO.CO.NH}_2$. — $\oplus \ominus$ Sapn. T.* products: NH_2 ; diphenylamine (No. 2.1568); oxalic ac. — Cryst. fr. h. aq. V.d.s. eth.
2311	170	† Malonamide, $\text{NH}_2\text{CO.CH}_2\text{CO.NH}_2$. — $\oplus \ominus$ Sapn. T.* products: NH_2 ; malonic ac. (Vol. I). Ndl. S. in 12 pt. aq. at 8°. I. abs. alc.
2312	170	p-Isopropyltoluylamide, $\text{Me}_2\text{CH.C}_6\text{H}_4\text{CH}_2\text{CONH}_2$. — $\oplus \ominus$ Sapn. T.* products: NH_2 ; p-isopropyltoluyl ac. (ndl. fr. aq., m.p. 52°). — Tbl. fr. bz. V.d.s. eth.; i. lgr.
2313	170–1; 173; (160)	"Dulcin," "Sucrol," p-Ethoxyphenylurea, $\text{EtO.C}_6\text{H}_4\text{NH.CO.NH}_2$. — \oplus Taste v. sweet!! — Cryst. fr. dil. alc. S. in 800 pt. c., or 55 pt. boiling aq.; s. in 25 pt. alc. — A suspension of comp. in 5 cc. aq. heated on boiling water-bath w. 2–4 drops mercuric nitrate sol. gives violet-blue color, changing to violet w. PbO_2 . (Delicacy, 2 mg. in 100 cc.) — Sapn. T.* products: NH_2 ; p-phenetidine (No. 2.1392).
2314	abt. 170d. (r.h.)	β -Rhamnohexanicacidphenylhydrazide, $\text{C}_9\text{H}_{11}\text{O}_4\text{NH.NH.Ph}$. — \oplus Hydrazide. § — Cryst. E.s. aq.; s. in abt. 200 pt. h. acetone; d.s. alc.
2315	170u.c.	† Narceine, $\text{C}_2\text{H}_7\text{NO}_3\text{SH}_2\text{O}$. — [Opium alkaloid. Alm. non-toxic.] — Silky ndl. fr. h. 50% alc. Separates fr. h. sat. aq. sol. as sticky mass. M.p. data conflicting. Said to cryst. w. $3\text{H}_2\text{O}$, & when dried 4 hr. at 100° to melt at 140–5°, the m.p. rising rapidly as aq. is taken up fr. air. (Cf. Freund, Ann., 277, 32.) — S. at 13° in 1285 pt. aq., or 945 pt. alc.; e.s. h. aq., alc.; d.s. chlf., amyl alc., KOH; i. bz. Odorless. Taste faintly bitter. Opt. i. Neut. to litmus. [Best extracted fr. aq. sol. containing alk. carbonate w. chlf. containing 10% alc.; partly extracted fr. acid sol. by chlf.] \oplus (1) Add 0.0005 g. B to sol. of 2 mg. resorcinol in 5 drops H_2SO_4 (sp. gr. 1.84). An OYS1 color develops. Heat 1 min. on steam bath w. constant stirring. Color becomes VRT1→. In 1 hr. after cooling color is R-OR; after 3 hr. OT1. — (2) Aq. I sol. gives BV-VBT1 color, gradually changing to light broken VR tones. — (3) Husemann's T. (Cf. No. 2.952) gives the colors described for narcotine. — (4) Heat together on steam-bath, constantly stirring, 1 mg. B, 2 mg. tannic ac. & 5 drops H_2SO_4 (sp. gr. 1.84). The color at first OYS1, soon changes to G, after some min. to BGS2, & finally to GYS3. (Narcotine & hydrastine give similar colors.) — (5) H_2SO_4 gives at first OYS1 color, changing after heating some sec. on steam-bath to OR-ORS1. (According to Dragendorff, as little as 0.01 mg. gives a blood-red color.) \ominus (a) Dis. 0.05 g. B in 3 cc. aq. + 2 drops HCl (sp. gr. 1.12). Add 0.5 cc. chloroplatinic ac. T. sol. A cryst. ppt. separates slowly on shaking & standing. Wash on filter w. 1 cc. dil. HCl. Dry at 100°. The product, $\text{B}_2\text{H}_2\text{PtCl}_6$, forms mic. OYT1 ndl., melting to dark brown liq. at 194–5° u.c. (198.3–9.5° c.) after beginning to darken abt. 190°. — (b) The picrate (un-analyzed) is easily prepared by ppn. of the acidified B sol. of the foregoing paragraph by 3 cc. c. sat. aq. picric ac. sol., washing w. 2 cc. aq. & recrystn. fr. 3 cc. 10% ac. ac. It forms fine Y ndl. which melt, after drying at 100°, at 127° u.c. (128.5° c.) to a clear yell. liq.
2316	170.5	$\beta\beta$ -Dinaphthylamine, $(\text{C}_{10}\text{H}_7)_2\text{NH}$. — \oplus Solutions fluor. blue. — B.p. 471° c. — Silvery lft. S. in 93.2 pt. bz. at 14.5°; d.s. h. alc.; e.s. h. gl. ac. ac. — \ominus B.Pk, red-brown hair-like ndl. fr. bz., m.p. 164–5°.
2317	171.5u.c. (174c.)	† 4,4'-Tetramethylaminobenzophenone, "Michler's Ketone," $\text{Me}_2\text{N.C}_6\text{H}_4\text{CO.C}_6\text{H}_4\text{NMe}_2$. — [Important intermediate in color industry.] — \oplus Fused w. NH_4Cl & ZnCl_2 at 150° gives the yellow dyestuff auramine (No. 3.1085). — Silvery lft. E.s. alc., eth. Dist. w. sl. decn. a.360°. — B.Pk (Cf. T. 2.23) purple-red pr. fr. alc., m.p. 156–7°.
2318	170–2	2-Methylhexanedioxime(4,5), $\text{Me.CHMe.CH}_2\text{C}(:\text{NOH}).\text{C}(:\text{NOH}).\text{Me}$. — \oplus Oxime. §

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2319	171-2	p-Dinitrobenzene, $(NO_2)_2.C_6H_4$. — \textcircled{P} Nitro comp.† — B.p. 298.4°. Sbl. 100 pt. aq. at 100° dis. 0.18 pt.; 100 pt. c. alc. dis. 0.4 pt.; s. c. bz.; s. to d.s. c. chlf. — Boiled for some hr. w. 5% NaOH sol. gives $NaNO_3$ & p-nitrophenol. — Gives a naphthalene deriv., $C_{10}H_8.C_6H_4(NO_2)_2$, which separates in fine white ndl., v.d.s. c. alc., when naphthalene is dissolved w. comp. in warm alc. & sol. allowed to cool.
2320	171-2	4,4',4"-Trinitrotriphenylcarbinol, $(NO_2C_6H_4)_3.C.OH$. — \textcircled{P} Nitro comp.‡ S. in alc. KOH w. intense violet-blue color. — "Nearly colorless" cryst. fr. bz. D.s. h. alc., eth.; more s. bz., gl. ac. ac. — <i>The ac. ac. sol. treated w. Zn dust becomes red.</i>
2321	171	p-Isobutylbenzamide, $(C_3H_7)_2.C_6H_4.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; p-isobutylbenzoic ac. ($C_{11}H_{14}O_2$, Vol. I). — Hair-like ndl. fr. h. aq.
2322	171	Gluconanilide, $CH_3(OH).(CH.OH)_4.CO.NHPh$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; gluconic ac. (Vol. I). — E.s. c. aq.
2323	169.5; 172	5-Acetamino-1,2,3,4-tetramethylbenzene, $(C_2H_5O)NH.C_6H_4Me_4$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 5-amino-1,2,3,4-tetramethylbenzene (No. 2.659); ac. ac. — Ndl. E.s. alc.
2324	172	Ethylenediacetamide, $(C_2H_5O)NH.CH_2.CH_2.NH(C_2H_5O)$. — \textcircled{P} \textcircled{D} Sapn. T. products: ethylenediamine (No. 2.1130); ac. ac. — Ndl. V.s. aq., alc., d.s. eth.
2325	172	1-Phenylsemicarbazide, $Ph.NH.NH.CO.NH_2$. — \textcircled{P} Should reduce Tollen's reagt. after splitting in T. 2.17 (possibly directly). — Lft. fr. dil. alc. E.s. h. aq., alc. D.s. c. aq. — Split by fuming HCl to CO_2 , NH ₃ & phenylhydrazine. HCl sol. w. NaNO ₂ gives nitroso deriv., pale yel. ndl. fr. aq., m.p. 126-7°.
2325	172u.c.	Dinitroso-2,5-(α)-dimethylpiperazine, $[N(NO).CHMe.CH_2.N(NO).CHMe-CH_2]$. — \textcircled{P} Gives blue color in T. 2.15! — Broad faintly yellowish ndl. fr. h. aq. D.s. h. aq., c. alc.; e.s. chlf.
2327	167; 177	Dinitrodi-(tert.)butylbenzene, $(NO_2)_2.C_6H_3.(CMe_3)_2$. — \textcircled{P} Odor musk-like. — Ndl. fr. alc.
2320	172-3c.	Methyl- γ -phenylhydantoin, $[CHMe.CO.NPh.CO.NH^2]$. — Ndl. fr. dil. alc.
—	172	Colchicine. — Cf. No. 2.2137. (M.p. 139-40° when dried at 100°.)
2329	172; 175	3,4-Dinitrobenzophenone, $NO_2.C_6H_4.CO.C_6H_4.NO_2$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. gl. ac. ac.
2330	170; 173; 176	1,4-(ac)-Diphenylsemicarbazide, $Ph.NH.CO.NH.NH.Ph$. — Cryst. fr. alc. D.s. h. aq.; e.s. alc. — Gradual addition of FeCl ₃ to c. alc. sol. (Ber., 29, 1690) gives phenylazocarbanilide, lust. or.-red lft. fr. dil. alc., m.p. 121-2°.
2331	173-5; 182	Propiophenonesemicarbazone, $Ph.C(:N.NH.CO.NH_2)Et$. — \textcircled{P} Semicarbazone.§ — Ndl. fr. alc.
2332	abt. 174-5 (r.h.)	Benzilmonosemicarbazone, $Ph.C(:N.NH.CO.NH_2).CO.Ph$. — \textcircled{P} \textcircled{D} Dis. slowly on warming in conc. HCl giving benzil (Vol. I), which suddenly separates out. — 8-cornered tbl. fr. alc. E.s. alc., chlf.; v.d.s. eth., lgr.
2333	174	Tetramethyldiaminobenzophenonephenylhydrazone, $(Me_2N.C_6H_4)_2C:N.NH-Ph$. — \textcircled{P} Hydrazone, becoming red, and then giving green sol. when covered w. conc. HNO ₃ ! — Ndl. fr. bz. + alc. I. aq.; s. h. alc.; e.s. eth.; v.s. chlf. bz.
2334	174-5	α -Naphthindole, $[C_{10}H_8.NH.CH:CH^2]$. — \textcircled{P} Gives deep blue-violet color to pine splinter in T. 2.24-b. — Lft. fr. lgr. Alm. i. h. aq.; d.s. lgr.; v.s. alc., eth., bz. D. vol. w. st.
2335	174	Benzoyl-m-aminophenol, $Ph.CO.NH.C_6H_4.OH$. — \textcircled{P} \textcircled{D} Sapn. T.* products: m-aminophenol (No. 2.830); benzoic ac. — Silky ndl. fr. toluene. V.s. alc., eth.; d.s. bz.
2335	174	Diisobutyramide, $(Me_2CH.CO)_2.NH$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; isobutyric ac. (Vol. I). — Ndl. fr. alc. I. aq. Rapidly decd. by distn. to isobutyric ac. (of rancid odor) & isobutyronitrile. — Sbl. below 100°.
2337	175	† s-Diethyloxamide, $EtNH.CO.CO.NHET$. — \textcircled{P} \textcircled{D} Sapn. T.* products: ethylamine (No. 2.1062); oxalic ac.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2330	175	Pyrotartaramide, $\text{Me}.\text{CH}(\text{CO.NH}_2).\text{CH}_2.\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; pyrotartaric ac. (Vol. I). — E.s. c. aq.
2339	abt. 175	Maleanilide, $\text{Ph}.\text{NH}.\text{CO}.\text{CH}:\text{CH}.\text{CO}.\text{NH}.\text{Ph}$. — \textcircled{P} \textcircled{D} Sapon. T.* products: aniline; maleic ac. (Vol. I). — E.s. alc.; d.s. dry eth., bz.
2340	175.5	Citraconanilide, $\text{Me}.\text{C}(\text{CO.NH.Ph}):\text{CH}.\text{CO}.\text{NH}.\text{Ph}$. — \textcircled{P} \textcircled{D} Sapon. T.* products: aniline; citraconic ac. (Vol. I). — Flat ndl. fr. alc. D.s. h. aq.; e.s. alc., eth.
2341	175	Acetonylurea, $[\text{CMe}_2.\text{NH}.\text{CO}.\text{NH}.\text{CO}]$. — [Slowly attacked by boiling baryta. Conc. HCl in sealed tube at 150–60° gives NH_3 , CO_2 & α -amino-isobutyric ac.] — Pr. E.s. aq., alc.; d.s. eth. — Evapd. on water-bath w. 5 pt. v. conc. HNO_3 gives nitro deriv., ndl. fr. bz., d.s. aq., m.p. 140–1°.
2342	175	Acenaphthenoneoxime, $\text{C}_{12}\text{H}_8.\text{NOH}$. — \textcircled{P} Oxime. — Lft. fr. alc.
2343	175d.	d-Glucosemicarbazone, $\text{CH}_2(\text{OH}).(\text{CH}.\text{OH})_4.\text{CH}:\text{N}.\text{NH}.\text{CO}.\text{NH}_2$. — \textcircled{P} Semicarbazone. § — Ndl. fr. abs. alc.
2344	175–6d.	(α)-Protocatechuicaldehydephenylhydrazone, $(\text{HO})_2\text{C}_6\text{H}_4.\text{CH}:\text{N}.\text{NH}.\text{Ph}$. — \textcircled{P} Hydrazone. § — Mic. tbl. fr. aq. E.s. alc., eth.; i. lgr.
2345	174–6	α -Nitrobenzamide, $\text{NO}_2.\text{C}_6\text{H}_4.\text{CO}.\text{NH}_2$. — \textcircled{P} Nitro comp.† — Short ndl. Mod. s. h. aq. — Sapon. T.* products: NH_3 ; α -nitrobenzoic ac. (No. 2.164).
2345	175–6	Trimethylcyanurate, $\text{Me}_3\text{C}_3\text{N}_3\text{O}_3$. — Odorless pr. D.s. h. aq.; s. alc. Normal KOH at 40–45° gives trimethylbiuret & CO_2 .
2347	175	Diphenyldiisocyanate, $[\text{NPh}.\text{CO}.\text{NPh}.\text{CO}]$. — \textcircled{P} Strongly heated gives phenylisocyanate (No. 2.2750), of powerfully irritating odor! — Tbl. fr. alc. I. aq.; d.s. h. alc.; alm. i. eth. — Alc. NH_3 converts at once to β -diphenylbiuret, pr. i. aq., m.p. 165°.
2348	176	α -Biquinolyline, $2,3'$ -Diquinolyl, $\text{C}_{12}\text{H}_{12}\text{N}_2$. — [Fr. quinoline & Na.] Silvery lft. I. h. aq.; alm. i. c. alc.; e.s. h. alc. Sbl. B.p. a. 400° sl. d. — [“Alc. sol. reacts neut.” $\text{B}_2\text{HCl}.\text{4H}_2\text{O}$, asbestos-like ndl., losing all aq. & iHCl at 100°, & decd. to components by aq.] — $\text{B}_2\text{HAuCl}_4.\text{2H}_2\text{O}$, yel. ndl., v.d.s. aq., m.p. 248°.
2349	abt. 176 sl. d.	† Cerebrin, Phrenosin, $\text{C}_{10}\text{H}_{16}\text{O}_{12}\text{N}_2$. — [A cerebroside fr. brain or protagon.] — \textcircled{P} Stirred w. conc. H_2SO_4 gives yellowish color, the undissolved portion after some time becoming purple. The purple (VR) is more quickly & satisfactorily developed when a little cane sugar is dissolved in the acid! (This reaction is also given by other cerebrosides.) — Spheroidal cryst. fr. alc., or light sl. hydroscopic powd. Swells up w. aq. E.s. h. alc., bz., chlf., acetone, gl. ac. aq.; i. h. eth.
2350	176	Glutaramide, $\text{NH}_2.\text{CO}.\text{(CH}_2)_2.\text{CO}.\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; glutaric ac. (Vol. I). — S. in 14 pt. aq. at 10.4°; i. eth.
2351	176.5	Mesaconamide, $\text{NH}_2.\text{CO.CMe}:\text{CH}.\text{CO}.\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; mesaconic ac. (Vol. I). — Plates fr. aq.
2352	176.5–7; (180)	Diphenylbenzamide, $\text{Ph}.\text{CO}.\text{NPh}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: diphenylamine (No. 2.1568); benzoic ac. — Rhomb. pr. fr. alc. D.s. alc., eth.
2353	176.5c.	† Pyrrole-2-carbonamide, $\text{C}_5\text{H}_5\text{NH}.\text{CO}.\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; pyrrole-2-carbonic ac. (No. 2.356). Lft. fr. alc. D.s. aq.; e.s. alc., eth. Tastes sweet!
2354	177	α -Biphenylcarbonamide, $\text{Ph}.\text{C}_6\text{H}_4.\text{CO}.\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH_3 ; α -phenylbenzoic ac. ($\text{C}_10\text{H}_{10}\text{O}_2$, Vol. I). — Ndl. Dist. alm. undecd. Alm. i. eth.
2355	177–8	p-Hydroxybenzaldehydephenylhydrazone, $\text{HO}.\text{C}_6\text{H}_4.\text{CH}:\text{N}.\text{NH}.\text{Ph}$. — \textcircled{P} Hydrazone. § — Colorless ndl. fr. alc. E.s. eth.; less s. alc.; s. KOH.
2356	177–8	Thujonesemicarbazone, $\text{C}_{10}\text{H}_{16}.\text{N}.\text{NH}.\text{CO}.\text{NH}_2$. — \textcircled{P} Semicarbazone. § — Ndl.
2357	177	† Methylene-bis-antipyrine, Formopyrine, $(\text{CO}.\text{NPh}.\text{NMe}.\text{CMe}:\text{C}^2).\text{CH}_2$. — Cryst. fr. alc. w. $1\text{H}_2\text{O}$ (lost at 130°), m.p. 155–65° w. frothing. I. aq., eth., alk.; e.s. bz., acids; 100 pt. 90% alc. dis. 12 pt. at 8–9°. — $\text{B}.\text{Pk}$

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2358	178	(T. 2.23), ppt., yel. ndl. fr. alc., m.p. 185 $^{\circ}$. — Tetraiodide, $C_{22}H_{24}O_2N_4I_4$, ndl. of I color, m.p. 135 $^{\circ}$ (fr. equal parts I & formopyrine in alc. sol.)
2359	178	Trinitro-o-xylene, $(NO_2)_3C_6H_2Me_2$. — \textcircled{P} Nitro comp.‡ — Lust. scales. Alm. i. c. alc.
2360	178d.	1,2-Dinitroprehnitol, $(NO_2)_2C_6Me_4$, [Me ₄ = 3,4,5,6]. — \textcircled{P} Nitro comp.‡ Glassy pr. D.s. c. alc.
2361	178	Diphenyloxobiuret, Ph.NH.CO.N(OH).CO.NH.Ph. — \textcircled{P} FeCl ₃ colors alc. sol. cherry-red. — Ndl. I. aq. — Sapn.* w. NaOH sol. gives aniline, diphenylurea, hydroxyamine, CO ₂ . — Does not reduce Fehling's sol.
2362	179	6,6'-(γ)-Biquinolyl, $C_{14}H_{12}N_2$. — Tbl. fr. alc. Dist. undecd. V.d.s. h. aq.; e.s. bz.; less s. alc., eth.
2363	abt. 180	Anisoylphenylhydrazide, p-MeO.C ₆ H ₄ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. § Phthalylphenylhydrazine, $[N(NHPh)CO.C_6H_4.CO]$. — \textcircled{P} Reduces Tollen's reagt. S. in warm conc. H ₂ SO ₄ w. violet color. — Ndl. fr. alc. I. aq., alk., acids, lgr.; d.s. c. alc.; e.s. h. alc.
2364	180	Benzoyl-o-tolylhydrazine, Ph.CO.NH.NH.C ₆ H ₄ .Me. — \textcircled{P} Should reduce Tollen's reagt. in T. 2.17 after sapn., if not directly. — Long lust. ndl.
2365	180	6-Pseudolutidostyryl, Lutidone, $[CO.CH:CM\bar{e}.CH:CM\bar{e}.NH]$. — Lust. ndl. fr. alc. B.p. 303–5 $^{\circ}$. E.s. aq., alc.; v.d.s. eth., bz. ["Reacts neut. Combines w. bases & acids."] Ignition w. Zn dust gives di & trimethylpyridines.
2366	180 (to violet liq.)	Dioxindole, $[NH.C_6H_4.CH(OH).CO]$. — \textcircled{P} Ammon. colors alc. sol. violet, & gives violet ppt. on boiling. — Colorless rhomb. pr. fr. alc.; yellowish fr. aq. S. in 12 pt. c., or 6 pt. boiling aq.; in 15 pt. c., or 10 pt. boiling abs. alc. Taste, bitter. Sinters fr. abt. 160 $^{\circ}$. Dec. at 190 $^{\circ}$, giving aniline (No. 2.1235). — Ag \ddot{A} , white cryst. ppt. (fr. aq. sol. Na \ddot{A} w. AgNO ₃ & ammon.); evolves aldehyde at 60 $^{\circ}$.
2367	180–1	5,6-Dinitropseudocumene, $(NO_2)_2C_6H_2Me_2$, [Me ₂ = 1:2:4]. — \textcircled{P} Nitro comp.‡ — Ndl. fr. h. alc. Alm. i. c. alc.
2368	180d. (r.h.)	Polymeric Hydrocyanic Ac., (HCN) _n . — \textcircled{P} Boiling w. aq. dec. to HCN, NH ₃ & brown matters. — Cryst. fr. aq. or alc. Brown at 140 $^{\circ}$. 100 pt. aq. at 24° dis. 0.55 pt., or 9–10 pt. at 100 $^{\circ}$; e.s. h. alc.; d.s. eth. — Sapn. w. baryta sol. gives NH ₃ , CO ₂ & glycine.
2369	180	as-Dimethylurea, Me ₂ N.CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; methylamine (No. 2.1059); CO ₂ . — Cryst. D.s. c. alc.; v.d.s. eth. Taste v. sweet!
2370	180	† ab-Methylacetylurea, MeNH.CO.NH(C ₂ H ₅ O). — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; methylamine; CO ₂ ; ac. ac. — Monoclin. pr. S.c. aq.; e.s. h. aq.; d.s. alc., eth.
2371	179–81	1,3-Dimethylbenzamide(4), Me ₂ C ₆ H ₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products (?) : NH ₃ ; 1,3-dimethylbenzoic(4) ac. (C ₉ H ₁₀ O ₂ , Vol. I). "Not decd. by boiling NaOH sol." — Ndl. Alm. i. c. aq.; v.s. alc.
2372	179; 182	o-Tolylphthalimide, $[CO.N(C_6H_5.Me).CO]$. — \textcircled{P} \textcircled{D} Sapn. T.* products: o-toluidine (No. 2.1262); phthalic ac. — Ndl. fr. gl. ac. ac. D.s. alc., eth.
2373	181	p-Tolylurea, Me.C ₆ H ₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; p-toluidine (No. 2.566); CO ₂ . — E.s. h. aq., alc., eth.; alm. i. c. aq., eth.
2374	181	Succinphenylamide, Ph.NH.CO.CH ₂ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; aniline; succinic ac. (Vol. I). — Ndl. s. h. aq.; d.s. h. alc.
2375	181	o-Nitrophenylurea, NO ₂ .C ₆ H ₄ .NH.CO.NH ₂ . — \textcircled{P} Nitro comp.‡ — Sapn. T.* products: NH ₃ ; o-nitroaniline (No. 2.2945); CO ₂ .
2376	182	2,4,6-Trinitro-1,3-xylene, $(NO_2)_3C_6H_2Me_2$. — \textcircled{P} Nitro comp.‡ — Ndl. alm. i. c. alc.
2377	182	Trinitro-1,3-dimethyl-4-tert.butylbenzene, $(NO_2)_3Me_2(Me_2CH).C_6$. — \textcircled{P} Nitro comp.‡ — Ndl. I. aq.
2378	182–3	5,8-Dinitroquinoline, $(NO_2)_2C_6H_2N$. — \textcircled{P} Nitro comp.‡ s. in warm NaOH w. red color! — Lust. ndl. fr. aq. V.d.s. h. aq., dil. ac., eth., bz.; more s. h. alc.; v.s. conc. acids.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2379	182	σ -Hydrazobiphenyl, $\text{Ph.C}_6\text{H}_4.\text{NH.NH.C}_6\text{H}_4.\text{Ph}$. — \textcircled{P} Becomes pink in air & prob. reduces Tollen's reagt. directly. — Ndl. fr. alc. D.s. c. alc. W. warm conc. HCl gives diphenylbenzidine.
2380	182	(α)-Methylhydantoin, $[\text{CO.NMe.CO.CH}_2.\text{NH}]$. — Pr. fr. aq. V.s. alc., h. aq.; v.d.s. eth.
2381	183-4	Phthalazone, $[\text{CO.C}_6\text{H}_4.\text{CH}_2.\text{N.NH}]$. — Long glassy ndl. fr. aq. E.s. aq.; v.s. h. alc., bz. B.p. 337°.
2382	183-4	γ -Aminobutyric Ac., Piperidinic Ac., $\text{NH}_2.\text{CH}_2.\text{CH}_2.\text{CH}_2.\text{CO}_2\text{H}$. — [$k_A \cdot 10^{11} = 3.7$; $k_B \cdot 10^{10} = 1.7$.] Lft. fr. MeOH + eth. V.s. aq. Sol. in 4 pt. aq. + 25 pt. alc. after long standing gives ndl., m.p. 202°. Heated above m.p. loses aq. readily, giving anhydride (pyrrolidone), m.p. 245°. — B_3HAuCl_4 (T. 2.13), monoclinc. tbl., m.p. 138°.
2383	183	4,4'-Dinitrodiphenylmethane, $(\text{NO}_2\text{C}_6\text{H}_4)_2\text{CH}_2$. — \textcircled{P} Nitro comp. — Ndl. I. c. aq., alc.; d.s. eth.; e.s. h. bz.
2384	183	Tetraphenylurea, $\text{Ph}_4\text{N.CO.NPh}_3$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. HCl, sp. gr. 1.12 at 250°) products: diphenylamine (No. 2.1568); CO_2 . — Cryst. I. aq.
2385	183	ab-Acetylphenylurea, $(\text{C}_6\text{H}_5\text{O})\text{NH.CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; aniline; ac. ac.; CO_2 . — Ndl. fr. aq.
2386	183	Hippuramide, $\text{Ph.CO.NH.CH}_2.\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; benzoic ac. (Vol. I); glycine (No. 2.2568). — Cryst. i. c. aq., alc., eth.; s. h. aq., alc.
2387	183	Suberanilide, $\text{Ph.NH.CO.(CH}_2)_4\text{CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; suberic ac. (Vol. I). — Scales. I. aq.; d.s. c., e.s. h. aq.; e.s. eth. — ["Not sapd. by h. KOH sol."]
2388	184	p-Tolylacetamide, $\text{Me.C}_6\text{H}_4.\text{CH}_2.\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; p-tolylacetic ac. ($\text{C}_9\text{H}_{10}\text{O}_2$, Vol. I). — Lft. fr. h. aq. D.s. c. aq., eth.; e.s. h. alc. Sbl.
2389	184	Acetyl diphenylhydrazine, $(\text{C}_6\text{H}_5\text{O})\text{NH.NPh}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: diphenylhydrazine (No. 2.1491); ac. ac. — Lust. ndl. I. aq.; e.s. h. alc.; d.s. eth. Dist. undec.
2390	184	β -Aminobutyric Ac., $\text{Me.CH}(\text{NH}_2).\text{CH}_2.\text{CO}_2\text{H}$. — Ndl. i. abs. alc., eth. — $\text{B}_3\text{H}_2\text{PtCl}_6$ (T. 2.14), e.s. or.-red cryst.
2391	185	Polymeric p-Tolylcarbonimide, $(\text{Me.C}_6\text{H}_4\text{NCO})_n$. — After fusion does solidify on cooling. I. aq. Ndl. fr. eth.
2382	185	3,5,6-Trinitropseudocumene, $(\text{NO}_2)_3\text{C}_6\text{H}_3\text{Me}$; [Me, = 1,2,4]. — \textcircled{P} Nitro comp. — Short quad. pr. fr. bz. V.d.s. h. alc.; v.e.s. bz.
2393	185	Azelaanilide, $\text{Ph.NH.CO.(CH}_2)_7\text{CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; azelaic ac. (Vol. I). — Ndl. Alm. i. aq., bz., eth.
2364	185	Pseudoitaconanilide, $\text{C}_6\text{H}_5\text{O}_2(\text{NH.Ph})_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; itaconic ac. (Vol. I). — Scales fr. h. aq. Alm. i. c. aq.; e.s. eth., h. alc.
2395	185	Hexahydrobenzamide, $\text{C}_6\text{H}_{11}.\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; hexahydrobenzoic ac. ($\text{C}_7\text{H}_{12}\text{O}_2$, Vol. I). — Flat pr. fr. aq. V.s. aq., eth.
2395	186	1,4-Dimethylbenzamide(2), $\text{Me}_2\text{C}_6\text{H}_4.\text{CONH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; 1,4-dimethylbenzoic(2) ac. ($\text{C}_7\text{H}_{10}\text{O}_2$, Vol. I). — Ndl. Mod. s. h. aq.; v.s. alc.
2397	185-7d.	Citraconamide, $\text{NH}_2.\text{CO.CMe}_2:\text{CH.CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 & citraconic ac. or its isomers. — E.s. c. aq.; d.s. alc.; i. eth.
2398	186	Mesaconanilide, $\text{Ph.NH.CO.CMe}_2:\text{CH.CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; mesaconic ac. or its transformation products. — Flat silky ndl. D.s. h. aq.; v.s. alc., eth.
2399	186-7	Hydrochelidonanilide, $\text{CO}(\text{CH}_2.\text{CH}_2.\text{CONH.Ph})_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; hydrochelidonic ac. ($\text{C}_7\text{H}_{10}\text{O}_2$, Vol. I). — Ndl. fr. alc.
2400	186	Acet-2,3-dinitroanilide, $(\text{C}_6\text{H}_5\text{O})\text{NH.C}_6\text{H}_4(\text{NO}_2)_2$. — \textcircled{P} Nitro comp. — Boiled w. KOH gives NH_3 . — Colorless ndl. D.s. c. alc., eth., bz.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS. — Colorless and Solid.
2401	186	Benzoyl-3,5-dinitro-p-toluide, $(NO_2)_2C_6H_3Me(Ph.CO.NH)$. — \textcircled{P} Nitro comp.† — Colorless ndl. fr. alc. — Mod. s. h. alc.; e.s. bz., h. Na_2CO_3 sol., alc. KOH.
2402	187 u.c.	p-Tolylsemicarbazide, $Me.C_6H_4.NH.NH.CO.NH$. — \textcircled{P} Reduces Tollen's reag. directly (T. 2.30)! — Cryst. E.s. h. aq., alc.
2403	187d.	† Acetonesemicarbazone, $Me_2C:N.NH.CO.NH$. — \textcircled{P} Reduces Tollen's reag. slowly in T. 2.30, but quickly & strongly after the splitting by HCl of T. 2.17. — Ndl. fr. acetone. S.c. aq., alc.; i. eth. — Distil w. dil. H_2SO_4 & test distillate for acetone (T. 1.711). [$k_b, 10^4 = 3.29$.]
2404	187	Helicinephenylhydrazone, $C_8H_{11}O_4.O.C_6H_4.CH:N.NH.Ph$. — \textcircled{P} Hydrzone. § — Indistinctly cryst. mass. Alm. i. c. aq., bz.; r.s. alc., eth., h. aq.
2405	187	Tetramethylsuccinimide, $[CO.CMe_2.CMe_2.CO.NH]^2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; tetramethylsuccinic ac. ($C_8H_{14}O_4$, Vol. I). — Ndl. fr. bz. + lgr. E.s. alc.
2405-I	187-8	2,4-Dibenzoylamino phenol, $(Ph.CO.NH)_2C_6H_3.OH$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 2,4-diaminophenol (No. 2.686); benzoic ac. — Lft. E.s. chlf., alc.; d.s. bz., eth.
2406	187	4-Acetaminonaphthol(1), Naphthacetol, $(C_8H_9O)NH.C_10H_8.OH$. — [A component in azo color manufacture.] Ndl. fr. alc. E.s. alc.; s. Na_2CO_3 , ammon. — Gives nitroso deriv., yel. ndl. fr. gl. ac. ac., s. in alk. w. yel. color, m.p. 203°.
2407	187; 186	Pr-2-Phenylindole, $[C_8H_4.NH.CPh:CH]^2$. — \textcircled{P} Gives intense violet color to pine splinter in T. 2.24-b! — Lft. fr. CS. B.p. a. 360°; 240-50° (10 mm.). I. aq.; s. in 38.5 pt. 94% alc., 47.4 pt. bz.; v.s. eth., chlf., gl. ac. ac.; less s. CS. — B.Pk (T. 2.23), red pr. fr. alc., decd. by aq., m.p. 127°.
2408	188-9	3-Acetylindole, $[C_8H_4.NH.CH:C(CO.CH)]^2$. — \textcircled{P} Boiling w. conc. HCl gives indole (No. 2.1546). (Cf. T. 2.24.) — Ndl. fr. bz. Sbl. in lft. D.s. c. aq., vz. — Picrate (T. 2.23), m.p. 183°.
2409	186-90d. (r.h.)	Rhammonic acidphenylhydrazide, $C_8H_{11}O_4.NH.NH.Ph$. — \textcircled{P} Hydrazide. § Lft. d.s. c. aq., alc.
2410	188	2,2'-(γ)-Dinitrobenzophenone, $NO_2.C_6H_4.CO.C_6H_4(NO_2)$. — \textcircled{P} Nitro comp.† — Ndl. fr. toluene.
2411	188.5 u.c.	† Biuret, $NH_3.CO.NH.CO.NH_2.H_2O$. — \textcircled{P} Gives the color reaction described in T. 2.10! — Long slender ndl. fr. slowly cooling h. half saturated aq. sol. containing a little NH_3 . Spontaneous evapn. of alc. sol. on microscope slide also gives slender ndl. Melts w. sl. efferv. Odorless. Taste faintly bitter. S. in 65 pt. aq. at 15°, or in 2.2 pt. at 106°; e.s. alc.; alm. i. eth. — Boiled w. aq. NaOH gives NH_3 & CO_2 .
2412	188-90	d,L-Gluconic acidphenylhydrazide, $CH_2.OH(CH.OH)_4.CO.NH.NHPh$. — \textcircled{P} Hydrazide. § — Warts. I. aq.
2413	189-90	4,4'-(α)-Dinitrobenzophenone, $NO_2.C_6H_4.CO.C_6H_4.NO_2$. — \textcircled{P} Nitro comp.† Ndl. fr. gl. ac. ac.
2414	abt. 189d.	1-Mannosaccharamide, $NH_2.CO.(CH.OH)_4.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; saccharic ac. (Vol. I).
2415	189	aa-Diphenylurea, $Ph_2N.CO.NH_2$. — \textcircled{P} Gently warmed w. conc. H_2SO_4 containing trace of HNO_3 gives blue sol.! — Long ndl. I. aq. — Sapn. T.* (w. alc. KOH) products: NH_3 ; diphenylamine (No. 2.1568); CO_2 .
2416	190-1	o-Tolylurea, $NH_2.CO.NH.C_6H_4.Me$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; o-toluidine (No. 2.1262); CO_2 . — Alm. i. c. aq., bz., eth.; d.s. c. acetone. — Acetylation w. acetyl chloride in pyridine sol. gives acetyl deriv., m.p. 168-9°.
2417	190	† Ethyl Allophanate, $NH_2.CO.NH.CO.Et$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; CO_2 ; ethyl alc. (T. 1.814). — Mic. ndl. fr. h. aq. V.d.s. c. aq.; e.s. h. aq.; d.s. c. alc.; v.d.s. eth. Heated in dry t.t. gives woolly sublimate. Distn. gives EtOH & cyanuric ac. — Odorless. Tasteless.
2418	190	Acet-4-nitro- α -naphthalide, $(C_8H_9O)NH.C_{10}H_8.NO_2$. — \textcircled{P} Nitro comp.† giving 4-nitro- α -naphthylamine (No. 2.3598), on sapn. by boiling 4 pt. w. 1- $\frac{1}{2}$ pt. KOH in alc. sol.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2419	191; 192.5	7,2'- β -Biquinolyl, $C_{16}H_{12}N_2$. — [Fr. quinoline vapor in red hot tube.] — \textcircled{P} Nitro comp. — Silky tbl. fr. alc. Sbl. in tbl. I. aq.; d.s. eth.; s. h. alc.; alm. i. c. alc.; e.s. chlf., h. bz. ["Feeble base, completely pptsd. fr. sol. of salts by aq."]
2420	191	Isovalerylurea, $NH_2CO.NH(C_6H_5CO)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; isovalerianic ac. (Vol. I). — Pr. Alm. i. c. aq., alc.
2421	191-2	α -Cumaramide, $HO.C_6H_5CH:CH.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; cumaric ac. ($C_9H_8O_3$, Vol. I). — Ndl. fr. alc.
2422	191	Diacetyl-m-phenylenediamine, $(C_6H_5O)NH.C_6H_4.NH(C_6H_5O)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: m-phenylenediamine (No. 2.634); ac. ac. — Pr. fr. dil. alc. D.s. c. aq.; more s. h. aq., alc.
2423	192	β -Naphthamide, $C_{10}H_8CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; β -naphthoic ac. ($C_{11}H_8O_2$, Vol. I). — Tbl. D.s. aq., alc. Dist. undec.
2424	192-3	Camphoramide, $C_8H_{14}(CONH_2)$. — \textcircled{P} \textcircled{D} Sapn. T.* (boiling w. KOH sol.) products: NH_2 ; camphorimide (cryst., s. in KOH sol., pptsd. by CO_2 , m.p. 244-5° c.). — Glassy pr. fr. alc. V.s. aq.; e.s. alc.; i. eth.
2425	192	4-Benzoylamino-1,3-xylene, $Ph.CO.NH.C_6H_4Me_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 4-amino-1,3-xylene (No. 2.1308); benzoic ac. (Vol. I). — Ndl. E.s. aq. Boiled w. conc. HNO_2 gives mononitro deriv., ndl. fr. alc., m.p. 184.5°.
2426	192	Benzoyldiphenylhydrazine, $Ph.CO.NH.NPh_2$. — \textcircled{P} \textcircled{D} Sapn. T.* ("slowly by conc. acids") products: diphenylhydrazine (No. 2.1491); benzoic ac. — Ndl. fr. acetone. E.s. h. acetone; less s. alc., eth.
2427	192	Di- β -naphthyl-m-phenylenediamine, $(C_{10}H_7)NH.C_6H_4.NH(C_{10}H_7)$. — \textcircled{P} Sol. fluor. blue. — Long ndl. fr. alc. + bz. B.p. a. 460° d. (45 mm.). Alm. i. alc., c. bz., aq.; s. acetone. — Diacetyl deriv., tbd. fr. alc., m.p. 175°.
2428	193	γ -Aminovalerianic Ac., $Me.CH(NH_2).CH_2.CH_2.CO_2H$. — Cryst. V.s. aq.; alm. i. alc.; i. eth., lgr. — $B.HCl$ (fr. alc. + eth.), m.p. 154°.
2429	193	α -Phenylhydantoin, $[NH.CO.NPh.CH_2.CO]^+$. — Ndl. fr. h. aq. E.s. alk. & pptsd. unchanged fr. sol. by acids! D.s. c. alc. Ammon. sol. gives ppt. w. $AgNO_3$ or $BaCl_2$.
2430	193	Acetylbiuret, $(C_2H_5O)NH.CO.NH.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; CO_2 ; ac. ac. — Ndl. fr. aq. V.s. aq.; v.d.s. eth.
2431	193-4 u.c.; 199	† Benzoyl-p-nitroanilide, $Ph.CO.NH.C_6H_4.NO_2$. — \textcircled{P} \textcircled{D} Sapn. of 0.5 g. w. sulphuric ac. by T. 2.26-b gives benzoic ac. & p-nitroaniline (No. 2.3319). The last-named product is obtained from the acid filtrate fr. which the benzoic ac. has been separated by making it strongly alk. w. conc. $NaOH$, cooling, shaking, washing w. aq., recrystg. twice fr. 50% alc., & drying at 50°. Lust., odorless, nearly colorless ndl. fr. h. alc. I. h. aq.; alm. i. c. alc., chlf. Gives good nitro† & carblyamine reactions (T. 2.21).
2432	194d.	Galactosecarbamide, $CH_2OH.(CH.OH)_2.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; galactosecarboxic ac. ($C_7H_{14}O_8$, Vol. I). — Mic. ndl. fr. ac. ac. Alm. i. alc.
2433	195-5.5	† Benzoylecgonine, $Ph.CO.O.C_6H_4O_2N$. — [In leaves of Erythroxylon coca.] \textcircled{P} \textcircled{D} Sapn. T.* (has been made w. conc. HCl in tube at 100°) products: ecgonine (No. 2.2249); benzoic ac. — Cryst. w. $4H_2O$ (m.p. 86-7°). D.s. c. aq.; more s. alc.; e.s. dil. ac., alk., h. aq.; i. eth.
2434	195-5.5	Benzoylpiperlyhydrazine, $(Ph.CO)NH.N.C_6H_4H_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: piperlyhydrazine (No. 2.1179), which reduces Tollen's reagt. directly; benzoic ac. — Pearly scales. Alm. i. c. aq., alc.; e.s. eth., bz., h. lgr., h. alc.
2435	195	Glycovanillinephenylhydrazone, $C_8H_7O_3.O.C_6H_4(OMe).CH:N.NH.Ph$. — \textcircled{P} Hydrazone, § — Cryst. mass fr. aq. V.d.s. c. aq.; alm. i. eth.; i. bz.; e.s. alc.
2436	195; 193-4	Diphenyleneketoxime, $[C_6H_4C(:NOH).C_6H_4]^+$. — \textcircled{P} Oxime, § — Ndl. fr. chlf. + lgr. HCl gas passed into eth. sol. gives yel. ppt. of $B.HCl$.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

(ORDER II, SUBORDER I.)

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2437	195-6	2-Methyl-3-acetylindole, $^{\text{r}}\text{C}_8\text{H}_7\text{NH.CMe: C(Me.CO)}^{\text{r}}$. — P After being boiled several min. w. conc. HCl gives pyrrole splinter react. (T. 2.24). ["Not sapd. by boiling NaOH sol." "V. weak base."] — Ndl. fr. bz. D.s. aq.; e.s. alc., h. bz.
2438	196	Diacetylmethylenediamine, $(\text{Me.CO.NH})_2\text{CH}_2$. — P D Sapn. T.* (w. HCl) products: NH ₃ ; formaldehyde (T. 1.114); ac. ac. — 4-sided pr. fr. alc. V.s. aq.; i. eth. B.p. 288°.
2439	196d.	Tartronamide, HO.CH(CO.NH) ₂ . — P D Sapn. T.* products: NH ₃ ; tartronic ac. (Vol. I). — Silky ndl. fr. dil. alc. D.s. c. aq.; alc.
2440	abt. 196u.c.	Ricinine, $\text{C}_3\text{H}_5\text{O}_2\text{N}_2$. — [Fr. seeds of Ricinus communis.] — P Sol. in conc. H ₂ SO ₄ colorless; but becomes yel.-green & then brilliant green on adding little K ₂ Cr ₂ O ₇ . (Delicate)! — Pr. fr. alc., aq. Sbl. undecd. Opt. i. Taste bitter. S. h. aq.; d.s. c. aq.; e.s. alc., eth., chlf., bz. ["Aq. sol. reacts neut."] — Sapn. w. alc. KOH gives ricinic ac. & methyl alc.
2441	196-7	2,4'-(δ)-Dinitrobenzophenone, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.C}_6\text{H}_4\text{NO}_2$. — P Nitro comp.‡ Cryst. V.d.s. alc., bz.
2442	197	Acet-2,6-dinitroanilide, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.CO.Me}$. — Nitro comp.‡ — Colorless ndl. fr. gl. ac. ac.
2443	197-8; 201.4c.	† p-Nitrobenzamide, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.NH}_2$. — P Nitro comp.‡ — Ndl. alm. i. aq. — Sapn. T.* products: NH ₃ ; p-nitrobenzoic ac. (No. 2.425).
2445	198-200; 195-8	Acetophenonesemicarbazone, Ph.C(:N.NH.CO.NH ₂)Me. — P Semicarbazone. § — Lft. fr. 90% alc.
2445	198	Malanilide, Ph.NH.CO.CH(OH).CH ₂ .CO.NH.Ph. — P D Sapn. T.* products: aniline; malic ac. (Vol. I). — Thin scales. R.d.s. alc., eth.
2447	198	Sebacanilide, Ph.NH.CO.(CH ₂) ₈ .CO.NH.Ph. — P D Sapn. T.* products: aniline; sebacic ac. (Vol. I). — Silky scales fr. alc. I. aq., lgr.; d.s. h. eth. B.p. a. 360°.
2448	198	Pr-3-Methyl- α -naphthindole, $^{\text{r}}\text{C}_{10}\text{H}_7\text{CMe: CH.NH}^{\text{r}}$. — P Colors splinter blue in T. 2.24-b! — Silvery lft. fr. dil. ac. ac. E.s. bz., h. abs. alc., gl. ac. ac.; d.s. lgr.
2449	198d. u.c.	† 1-Egonine, 3-Hydroxytropane-2-carbonic Ac., Me.NC ₆ H ₁₀ -(OH)(CO ₂ H)-H ₂ O. — [Fr. sapn. of coca alkaloids by HCl.] — Monoclin. hemimorph. pr. fr. alc. containing 1 mol. H ₂ O. After dehydration by heating 2 hr. at 185-40°, melts at 205.2° d. u.c. Browns & efferv. in melting. Odorless. Taste bitter-sweet. 1 g. B dis. at 17° in 4.6 cc. aq., or 67 cc. 95% alc.; v.d.s. eth.; s. ethyl acetate; alm. i. chlf., CS ₂ . [$k_B \cdot 10^{11}$ lies between 0.2 & 8.] — P Evapd. to dryness w. Cl aq. gives green color w. conc. H ₂ SO ₄ . — B ₂ H ₂ PtCl ₆ , prepared by T. 2.14, is obtained in broad orange ndl., m.p. 226° (recryst. fr. dil. alc. & dried at 140°), containing 24.93% Pt. — B.HAuCl ₄ (T. 2.13), cryst. w. 2H ₂ O fr. alc., m.p. 202°, or 71° (anhydrous).
2450	199-200	† 2-Hydroxyquinoline, Carbostyril, $\text{HO.C}_8\text{H}_7\text{N}$. — Pr. fr. alc. Cryst. fr. h. 1% aq. sol. w. 1H ₂ O in long threads. Sbl. Alm. i. c., e.s. h. aq.; e.s. alc., eth.; i. ammon.; d.s. HCl. Na salts e.s. aq.; deod. by CO ₂ . Fusion w. NaOH gives some indole (No. 2.1546).
2451	abt. 200d.	d- & l-Gluconic acidphenylhydrazide, $\text{CH}_2(\text{OH}).(\text{CH.OH})_4\text{CO.NH.NHPh}$. — P Hydrazide. § — Gives violet-red color w. conc. H ₂ SO ₄ + 1 drop FeCl ₃ . — Pr. fr. h. aq. 100 pt. boiling aq. dis. 15.1 pt. D.s. h. alc.; i. eth.
2452	196; 206-7; not melted at 220	β -Alanine, β -Aminopropionic Ac., NH ₂ .CH ₂ .CH ₂ .CO ₂ H. — [$k_A = 7.1 \times 10^{11}$; $k_B = 5.1 \times 10^{11}$.] Oblique rhomb. pr. fr. aq. E.s. aq.; alm. i. abs. alc.; i. eth. Tastes faintly sweet. — Cu ₂ .6H ₂ O, deep blue cryst. s. w. deep blue color in alc. — Benzoyl deriv. (fr. benzoyl chloride & NaOH, T. 2.27), pr. d.s. c. aq., e.s. alc., m.p. 120°. — Phenylureidoacid deriv. (T. 2.31) ndl. or tbd. fr. aq., m.p. 171-2° (w. efferv.). — Et ₂ A, b.p. 58° (14 mm.).
2453	abt. 200	† 3,5-Diphenylpyrazole, $^{\text{r}}\text{CH}_2\text{CPh:N.N.CPh}^{\text{r}}$. — Cryst. B.p. 347° (th.i., 175 mm.). Sbl. 202°. — B.HCl, ndl. fr. alc. + HCl, m.p. 232-3°.
2454	200-1	1,2,4-Trimethylbenzamide(5), Me ₃ C ₆ H ₃ .CO.NH ₂ . — P D Sapn. T.* products: NH ₃ ; 1,2,4-trimethylbenzoic ac. (C ₁₀ H ₁₂ O ₂ , Vol. I). — Lust. ndl. fr. dil. alc.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2455	201	o-Acetaminophenol, $(C_6H_5O)NH.C_6H_4.OH$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. conc. HCl at 130°) products: o-aminophenol (No. 2.938); ac. ac. — Lft. fr. dil. alc. E.s. alc., h. aq.; s. KOH sol.
2456	201	4-Hydroxyquinoline, Kyanurin, $HO.C_6H_4N$. — [“Reacts feebly alk.”] — Cryst. w. $3H_2O$ (lost at 115°), m.p. abt. 60°. 100 pt. aq. at 15° dis. 0.477 pt.; more s. c. alc.; d.s. abs. eth., lgr., bz. Taste bitter. Colored faintly carmine-red by $FeCl_3$. Ignition w. Zn dust gives quinoline (No. 2.1356).
2456-I	abt. 210 <u>c.</u> d.	† Sarcosine, Methylglycine, $MeNH.CH_2.CO_2H$. — Rhomb. pr. Taste faintly sweetish. V.s. aq.; d.s. alc. On fusion gives sarcosine anhydride, CO_2 , & dimethylamine (No. 2.1061). — Picrate, ndl. fr. aq.; m.p. 149.5° c.
2457	202	α -Benzalaminophenylacetonitrile, Benzoylazotide, $Ph.CH:N.C(=O)Ph.CN$. — Cryst. gran. I. aq.; s. in 300–400 pt. h. alc. Softens at 198°. Not decd. by dil. acids, but gives benzaldehyde & phenylaminoacetic ac. (No. 2.465) when long boiled w. conc. HCl.
2458	202; 202.5; 203.5; 204.5; 206	Benzalbenzoylhydrazine, $Ph.CH:N.NH.CO.Ph$. — Ndl. fr. alc. E.s. alc., chlf. — Reduced to benzamide (No. 2.2051) & benzylamine (No. 2.1236) by Zn dust & ac. ac.
2459	202	α -Naphthoamide, $C_{10}H_7.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; α -naphthoic ac. (Vol. I). — Cryst. fr. alc. V.d.s. aq., alc.
2460	201–2; 204	p-Tolylphthalimide, $[CO.C_6H_4.CO.NC_6H_4]^2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-toluidine (No. 2.566); phthalic ac. (Vol. I). — Ndl. I. h. aq.; d.s. h. alc. Sbl.
2461	202	Diacetyldiphenylin, $(C_6H_5O)NH.C_6H_4.C_6H_4.NH(C_6H_5O)$. — \textcircled{P} \textcircled{D} Sapn. T.* products: diphenylin (No. 2.573); ac. ac. — Cryst.
2462	202–3	Ethyloxamide, $EtNH.CO.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; ethylamine (No. 2.1062); oxalic ac. — Cryst. Sbl. E.s. h. aq.
2463	202	Tribenzamide, $(Ph.CO)_3N$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; benzoic ac. (Vol. I). — Silky ndl. fr. alc. Sbl. I. c. alc.; v.d.s. eth.
2464	202–3	ϵ -Aminocaproic Ac., $NH_2.(CH_2)_5.CO_2H$. — Lft. fr. MeOH + eth. V.s. aq.; d.s. MeOH. Begins to soften at 190°.
2465	200–5	d- & L-Galactonicacidphenylhydrazide, $CH_2(OH).(CH.OH)_4.CO.NH.Ph$. — Hydrazide. § — Cryst. fr. h. aq. D.s. c. aq.; more s. alc.; h. aq. [The d,l-compound also melts at 205° on rapid heating.]
2466	203	1,3,6,8-Tetranitronaphthalene, $(NO_2)_4C_{10}H_4$. — \textcircled{P} Nitro comp.‡ which explodes violently when rapidly heated. — Long thin ndl. fr. alc. Gives red color w. alc. NH ₃ .
2467	204	p-Nitrooctylbenzene, $NO_2.C_6H_4.C_8H_{17}$. — \textcircled{P} Nitro comp.‡ — Lust. ndl. Sbl. S. h. alc. — Oxid. to p-nitrobenzoic ac. (No. 2.425) by KMnO ₄ (Cf. T. 1.905–1).
2468	204–5	Methyl-p-tolylketonesemicarbazone, $Me.C(:N.NH.CO.NH_2).C_6H_4.Me$. — \textcircled{P} Semi-carbazone. § — Mic. pr. fr. alc. R.d.s. alc.
2469	204; 204.5; 216–8(r.h.)	† L-Arabinosediphenylhydrazone, $CH_2.OH(CH.OH)_2.CH:N.NPh_2$. — \textcircled{P} Gives immediate deep blue color in Bülow's react. T. 2.21! Hydrazone. — Colorless or faintly yellowish ndl. fr. dil. alc. D.s. aq., c. alc.; e.s. pyridine, h. alc.; v.d.s. bz., lgr. $[\alpha]_D = +0.42^\circ$ (for sol. 0.2 g. hydrazone in 4 cc. pyridine + 6 cc. abs. alc.).
2470	204	Oxalydiethylhydrazide, $Et.NH.NH.CO.CO.NH.NH.Et$. — \textcircled{P} “In alk. sol. dec. CuO, Ag ₂ O or HgO w. evol. of gas.” — Fine ndl. S. h. aq., alc. E.s. dil. min. acids: pptd. on exact neutralization by alk., ppt. rediss. in v. sl. x.s. alk. — Nitroso deriv. (fr. NaNO ₂ & dil. H ₂ SO ₄), ndl. e.s. h. aq., m.p. 144–5°.
2471	203–5	Phenylphthalimide, $[CO.NPh.CO.C_6H_4]^2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; phthalic ac. (T. 1.318). — Ndl. fr. alc. I. aq. Sbl.
2472	204	† Benzoylmesidine, $Ph.CO.NH.C_6H_4.Me$, [Me ₂ = 1:3:5]. — \textcircled{P} \textcircled{D} Sapn. T.* products: mesidine (No. 2.1343); benzoic ac. — Ndl.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2473	204	Oxalylcarbanilide, (Diphenylparabanic Ac.), $[(CO.NPh.CO.CO.NPh)]_2$. — \textcircled{P} Sapon. T.* (w. KOH) products: aniline; oxalic ac.; CO ₂ . — Ndl. I. aq.; e.s. alc., eth.
2475	205 u.c.	† Dicyandiamide, C ₂ H ₄ N ₂ . — \textcircled{P} Evolves NH ₃ freely in T. 227. — Pearly lft. fr. h. sat. aq. sol. on slow cooling. E.s. c. aq.; s. 95% alc.; alm. i. c. eth. Odorless. Taste of c. conc. aq. sol. sl. bitter. — For delicate color react., cf. Ber., 26, 1527. — Sol. w. AgNO ₃ gives ppt. of C ₂ H ₄ N ₂ .AgNO ₃ , lust. ndl., d.s. c., more s. h. aq.; alm. i. HNO ₃ .
2477	205	3,6-Dinitro-1,2,4,5-tetramethylbenzene, (Dinitrodurene), (NO ₂) ₂ C ₆ Me ₄ . — \textcircled{P} Nitro comp.‡ — Rhomb. pr. Sbl. undecd. in ndl. V.d.s. c. alc.; e.s. eth.
2478	205 u.c.; 208. 5c.; 210	3-Nitrocarbazole, $[(NH.C6H5.(NO2).C6H5)]_2$. — \textcircled{P} Nitro comp.‡ — Lft. fr. h. alc. Alm. i. eth., lgr.; d.s. h. chlf., bz.
2479	206	Dinitro- β -methylnaphthalene, (NO ₂) ₂ C ₁₀ H ₈ . — \textcircled{P} Nitro comp.‡ — Nearly colorless ndl. D.s. alc.
2480	206-7; 203	† p-Trinitrotriphenylmethane, (NO ₂ C ₆ H ₅) ₂ CH. — \textcircled{P} Nitro comp.‡ — Dissolving w. intense violet-blue color in alc. KOH! — Small scales fr. bz. V.d.s. c. gl. ac. bz., eth.
2481	abt. 206	d,L-Arabinosediphenylhydrazone, CH ₂ (OH).(CH.OH) ₂ CH.N.NPh ₂ . — \textcircled{P} Should give blue color in Bülow's react., T. 211. — Colorless ndl. fr. pyridine + alc. D.s. aq., alc., chlf., bz.
2482	206	Iosuccinimide, Me.CH(CO.NH ₂) ₂ . — \textcircled{P} \textcircled{D} Sapon. T.* products: NH ₃ ; iosuccinic ac. (C ₄ H ₆ O ₄ , Vol. I). — Pr. D.s. h. alc.; i. eth.
2483	207 u.c.	† Acet-p-nitroanilide, (C ₆ H ₅ O)NH.C ₆ H ₄ .NH ₂ . — \textcircled{P} \textcircled{D} S. in c. aq.-KOH (1:2) w. intense red-brown or orange color. After abt. 12 hr. pr. of p-nitraniline (No. 2.3319) of m.p. 141° separate fr. this sol. — Colorless ndl. fr. h. aq. V.d.s. aq., chlf., eth., lgr. Sol. in 10% NaOH, Y; in conc. H ₂ SO ₄ , colorless. Gives good nitro comp. react.‡
—	abt. 207(r.h.)	β -Alanine. — Cf. No. 2.2452.
2484	208	Sebacamide, NH ₂ CO.(CH ₂) ₈ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapon. T.* products: NH ₃ ; sebacic ac. (Vol. I). — Cryst. I. c. aq.; s. h. aq.; d.s. c. alc.
2485	208d.	Methyl Allophanate, NH ₂ CO.NH.CO.Me. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH ₃ ; CO ₂ ; methyl alc. (Vol. I). — Ndl. fr. h. aq.; alm. i. c. aq.
2486	208	1,3,5-(s)-Triacetaminobenzene, (Me.CO.NH) ₂ C ₆ H ₄ . — \textcircled{P} \textcircled{D} Sapon. T.* products: 1,3,5-triaminobenzene (unknown in free state); ac. ac. — Lft. fr. alc. E.s. h. aq.; alm. i. bz., eth., c. aq.
2486-I	208-9	1-Hydroxyisoquinoline, Isocarbostyryl, $[(C6H5.CH:CH.NH.CO)]_2$. — Cryst. fr. bz. D.s. aq., eth., bz., e.s. alc., chlf.; alm. i. lgr. Distrn. w. Zn dust gives isoquinoline (No. 2.1365).
2487	209	Trinitro-1,2,3-trimethylbenzene, (NO ₂) ₃ C ₆ Me ₃ . — \textcircled{P} Nitro comp.‡ — Glassy pr. fr. alc.; alm. i. alc.
2488	209-10	6-Nitro-4-acetamino-1,2-xylene, (NO ₂).Me.CO.NH.C ₆ H ₄ .Me. — \textcircled{P} Nitro comp.‡ — Colorless ndl. fr. alc.
2488	209. 5d. u.c.	Dinitrosopentamethylenetetramine, C ₁₀ H ₁₀ O ₂ N ₄ . — Boiled w. dil. HCl gives NH ₃ ; formaldehyde (Vol. I), & N. — Ndl. fr. alc. S. in 100 pt. h. alc.
2490	abt. 210w. efferv.	d-Saccharic acid bisphenylhydrazide, Ph.NH.CO.(CH.OH) ₄ .CO.NH.Ph. — \textcircled{P} Hydrazide giving red coloration w. conc. H ₂ SO ₄ & FeCl ₃ (T. 211). — Yellowish white tbl. I. aq., alc., eth.; s. alc.-NaOH.
2491	210d.	α -Rhamnohexonic acid diphenylhydrazide, Me.(CH.OH) ₄ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide, prob. giving coloration in T. 211. — Hexag. lft. S. in 72 pt. aq., at 17°.
2492	210; 195	Dibenzoylurea, Ph.CO.NH.CO.NH.CO.Ph. — \textcircled{P} \textcircled{D} Sapon. T.* (w. conc. HCl) products: NH ₃ ; benzoic ac.; CO ₂ . — Ndl. fr. h. gl. ac. ac. Dec. at 205-10° to benzonitrile (No. 2.2781), benzamide (No. 2.2051), & CO ₂ .
2493	210	3,4-Diacetaminotoluene, (Me.CO.NH) ₂ C ₆ H ₄ .Me. — \textcircled{P} \textcircled{D} Sapon. T.* products: 3,4-toluylenediamine (No. 2.707); ac. ac. — Long fine ndl. fr. h. aq.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2494	210	Oxal-o-toluide, $\text{Me.C}_6\text{H}_4.\text{NH.CO.CO.NH.C}_6\text{H}_4.\text{Me}$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: o-toluidine (No. 2.1262); oxalic ac. — Lft. i. aq.; d.s. alc. Ignition w. Zn dust gives indole (No. 2.1546).
2495	210	α -Diphenylbiuret, $\text{Ph.NH.CO.NH.CO.NH.Ph}$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: aniline; NH_3 ; CO_2 . — Cryst. I. aq.; d.s. alc.
2496	211	Phenylasparticphenylimide, Anilino-succinimide, $[\text{CH}(\text{NHPH}).\text{CO.NPh.CO.-CH}_2]$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* (w. alc. KOH or conc. HCl at 100°) products: aniline; phenylaspartic ac. — Small ndl. I. aq.; v.d.s. c. alc.; e.s. gl. ac. ac.
2497	211	Phenyl-s-pseudocumeylurea, $\text{Ph.NH.CO.NH.C}_6\text{H}_4.\text{Me}$, [NH : Me, = 5 : 1,-2,4]. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: aniline; s-pseudocumidine (No. 2.646); CO_2 . — Fine ndl. fr. alc.
2498	211-2; 209	\dagger 4-(δ)-Oxyquinazoline, $[\text{C}_6\text{H}_4.\text{CO.NH.CH:N}^{\ddagger}]$. — \textcircled{P} Sol. fluor. blue! — Ndl. fr. aq. E.s. aq., alc.; less s. eth.; i. lgr. — $\text{B}_2\text{H}_6\text{PtCl}_6\text{H}_2\text{O}$, or.-red cryst. ppt.
2499	216-7c. = 210.5-11.5u.c.	Purine, $\text{C}_5\text{H}_4\text{N}_4$. — Mic. ndl. fr. toluene or alc. Sinters before melting. Carbonizes w. partial volatilization above m.p. V.s. c. aq.; e.s. warm alc.; v.d.s. eth., chlf. [\dagger Sol. reacts neutral to litmus & turmeric.] Tannic ac. gives floc. ppt. — B.Pk (Cf. T. 2.23), lust. lft., d.s. c. aq., s. in abt. 20 pt. h. aq., m.p. abt. 208°.
2500	211	1,5-(α)-Dinitronaphthalene, $(\text{NO}_2)_2\text{C}_10\text{H}_4$. — \textcircled{P} Nitro comp.‡ Boiled w. 10 pt. alc. & 2 pt. KCN gives blue-green sol. — 6-sided colorless ndl. fr. gl. ac. ac. V.d.s. ordinary solvents; d.s. c. bz. — Reduction w. ammon. sulfide (Ann., 169, 87) gives 1,5-nitronaphthylamine, small red ndl. fr. aq., m.p. 118-9°.
2501	212d. (r.h.)	d-Mannosaccharicacidbisphenylhydrazide, $\text{Ph.NH.NH.CO.(CH.OH)}_2\text{CO.-NH.NH.Ph}$. — \textcircled{P} Hydrazide, prob. giving coloration in Bülow's react., T. 2.11. — Sl. yellowish lft. I. h. aq.
2502	212	Phenyl-o-tolylurea, $\text{Ph.NH.CO.NH.C}_6\text{H}_4.\text{Me}$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: aniline; o-toluidine (No. 2.1262); CO_2 . — Fine ndl.
2503	210-5d.	Citramide, $\text{NH}_3\text{CO.CH}_2\text{C(OH)(CO.NH}_2\text{).CH}_2\text{CO.NH}_3$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: NH_3 ; citric ac. (Vol. I). — Melts to black liquid. S. c. aq.; v.s. h. aq.; i. alc., eth.
2504	210-5d.	Anthranil, o-Aminobenzoicacidanhydride, $[\text{C}_6\text{H}_4.\text{NH.CO}]$. — \textcircled{P} Reduces Au & Ag sol. on warming. — Odor basic & bitter almond-like. Resinifies in the air. E. vol. w. st. — Sol. in h. dil. NaOH, giving salt of anthranilic ac. (No. 2.148).
2505	212-4 w. efferv.	α -Aminosaradic Ac., $\text{C}_{12}\text{H}_{20}(\text{NH}_2).\text{CO.H}$. — Lft. fr. gl. ac. ac. V.d.s. h. alc.; i. eth. [\dagger Does not combine w. acids or bases.]
2506	213 (frothing)	Areccaine, $[\text{NMe.CH}_2\text{CHMe.CO.CO.CH}_2]$. — [Alkaloid fr. the areca nut, Areca Catechu.] — Cryst. w. H_2O (lost at 100°). E.s. aq., dil. alc.; alm. i. abs. alc.; i. eth., chlf., bz. [\dagger Aq. sol. reacts neut.] — KBi iodide gives red ppt., soon becoming cryst., fr. sol. acidified w. H_2SO_4 . — $\text{B}_2\text{H}_6\text{PtCl}_6$ (T. 2.14), or.-yel. octahedra fr. aq., m.p. 213-4° (frothing). — B.HAuCl_4 (T. 2.13), pr. fr. h. dil. HCl, m.p. 186-7°.
2507	210-6	β -p-Dinitrostilbene, $\text{NO}_2\text{C}_6\text{H}_4.\text{CH:CH.C}_6\text{H}_4.\text{NO}_2$. — \textcircled{P} Nitro comp.‡
2508	213-4	β -Naphthylurea, $\text{C}_{10}\text{H}_7\text{NH.CO.NH}_3$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: NH_3 ; β -naphthylamine (No. 2.781); CO_2 . — After fusion & solidification remelts above 300° giving dinaphthylurea. — Ndl. fr. alc.
2509	214	Tartaricaciddi- α -naphthalide, $\text{C}_{10}\text{H}_7\text{NH.CO.(CH.OH)}_2\text{CO.NH.C}_{10}\text{H}_7$. — \textcircled{P} Sapn. T.* products: α -naphthylamine (No. 2.589); tartaric ac. (Vol. I). — Ndl. fr. alc. E.s. eth.; d.s. bz.
2510	213-5	Ethylmalonanilide, $\text{Ph.NH.CO.CHEt.CO.NH.Ph}$. — $\textcircled{P} \textcircled{D}$ Sapn. T.* products: aniline; ethylmalonic ac. ($\text{C}_4\text{H}_8\text{O}_4$, Vol. I). — Ndl. fr. alc. I. aq.; e.s. alc., ac. ac.
2511	214d.	\dagger Benzalsemicarbazide, $\text{Ph.CH:CH.NH.CO.NH}_3$. — \textcircled{P} Semicarbazide.§ — Cryst. fr. aq. V.d.s. h. aq.; e.s. alc.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2512	214d.; abt. 215	d- & l-Arabonic acidphenylhydrazide, $\text{CH}_3\text{OH}(\text{CH}_3\text{OH})_2\text{CO.NH.NH.Ph.}$ — \textcircled{P} Hydrazide. § — Colorless cryst. fr. warm aq. D.s. c. aq.
2513	214-6d.	d- & l-Mannonic acidphenylhydrazide, $\text{CH}_3\text{OH}(\text{CH}_3\text{OH})_2\text{CO.NH.NH.Ph.}$ — \textcircled{P} Hydrazide. § — Cryst. S. alc., warm aq.
2514	abt. 215d.	α -Aminocerotic Ac., $\text{C}_{14}\text{H}_{26}\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$. — Cryst. powd. Alm. i. alc., etc.
2515	214-6; 200	† Amygdalin, $\text{C}_{20}\text{H}_{22}\text{O}_5\text{N}_2$. — [Glucoside.] — \textcircled{P} Boiled w. dil. HCl gives odor of bitter almonds! — Rhomb. cryst. w. $3\text{H}_2\text{O}$ fr. aq. (Aq. lost at 110-20°.) After fusion solidifies to glassy mass remelting at 125-30°. Cryst. w. $2\text{H}_2\text{O}$ in lust. scales fr. 80% alc. S. at 8-10° in 12 pt. aq., or 904 pt. alc. (sp. gr. 0.819); i. eth. Taste of aq. sol. bitter. $[\alpha]_D = -41.1^{\circ}$ (in 3.883% aq. sol.).
2516	215	Inosin, $\text{C}_{10}\text{H}_{12}\text{O}_4\text{N}_4$. — [Glucoside giving hypoxanthin & a pentose on hydrolysis.] E.s. h. aq.; d.s. other solvents. $[\alpha]_D = -49.2^{\circ}$ (in 9% aq. sol.).
2517	215	Hydantoin, Glycolylurea, $[\text{CO.NH.CH}_2\text{CO.NH}]$. — \textcircled{P} W. NH ₃ & AgNO ₃ gives ppt. $\text{AgC}_2\text{H}_4\text{O}_2\text{N}_2\text{H}_2\text{O}$, s. in x.s. NH ₃ . — Ndl. S. c. aq.; e.s. h. aq. Boiling w. baryta aq. gives hydantoic ac. (No. 2.187).
2518	215	Benzoylurea, Ph.CO.NH.CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* (w. NaOH sol.) products: NH ₃ ; benzoic ac.; CO ₂ . — Lft. fr. alc. S. in 24 pt. c. alc.; s. h. aq.; e.s. KOH sol.; i. eth.
2519	215	β -Naphthoylurea, $\text{C}_{10}\text{H}_8\text{CO.NH.CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; β -naphthoic ac. (Vol. I). — Ndl. R.d. s. alc.; v.d.s. chlf., bz.
2520	216	Dibenzylloxamide, Ph.CH ₂ .NH.CO.CO.NH.CH ₂ .Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: benzyl amine (No. 2.1236); oxalic ac. — Scales fr. alc. I. aq.; d.s. h. alc.
2521	216-7	Suberamide, $\text{NH}_2\text{CO}(\text{CH}_2)_4\text{CO.NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; suberic ac. (Vol. I). — Cryst. S. in abt. 1170 pt. aq. at 18°; v.d.s. c. alc; i. eth.
2522	216-7	Acetmesidine, $(\text{C}_2\text{H}_5\text{O})\text{NH.C}_6\text{H}_5\text{Me}$; [NH:Me, = 2:1,3,5]. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; mesidine (No. 2.1343); ac. ac. — Pr. fr. alc. Sbl. undec. E.s. h. alc.
2523	216-7; 207-9	Santonin oxide, $\text{C}_{10}\text{H}_{10}\text{O}_3\text{N.H}_2\text{O}$. — \textcircled{P} Oxime. § — Ndl. I. c., v.d.s. h. aq.; i. alk. $[\alpha]_D = -80.83^{\circ}$ (2.3364 g. in 100 cc. gl. ac. ac.).
2524	217	β -Aminoisovaleric Ac., Me ₂ C(NH ₂).CH ₂ .CO ₂ H.H ₂ O. — Glassy cryst. fr. eth.-alc. V.s. aq.; d.s. abs. alc.; i. eth. — $2\text{AgA}.\text{AgNO}_3.\text{H}_2\text{O}$ (fr. conc. aq. sol. w. AgNO ₃ & few drops ammon.), ndl. e.s. h. aq., less s. c.
2525	217.4u.c.	2-Oxylepidine, Lepidone, γ -Methylcarbostyryl, $[\text{NH.C}_6\text{H}_4\text{CMe}:\text{CH.CO}]$. — Ndl. fr. aq. V.d.s. c. aq., eth., chlf., bz., lgr.; e.s. h. alc. — B.p. 270° (17 mm.).
2526	218d.	Trigoniellin, Methylbetaine of Nicotinic Ac., $\text{C}_7\text{H}_9\text{O}_2\text{N}$. — \textcircled{P} Boiled w. KOH sol. gives off methylamine (No. 2.1059). — Cryst. w. $1\text{H}_2\text{O}$ fr. 96% alc. in flat pr. of saline taste & neut. react. Melts in cryst. aq. abt. 130°. Darkens at 200°. I. eth., chlf., bz.; v.s. aq.
2527	218	1,3,8-Trinitronaphthalene (NO_2) ₃ .C ₁₀ H ₆ . — \textcircled{P} Nitro comp.‡ — Cryst. fr. gl. ac. ac. 100 cc. 88% alc. dis. 0.046 g. at 23°; v.d.s. eth., chlf.
2528	215; 222	Terephthalonitrile, p-CN.C ₆ H ₄ .CN. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; terephthalic ac. (Vol. I). — D.s. c. alc., h. eth.
2529	219-20	† Phthalamide, o-NH ₂ .CO.C ₆ H ₄ .CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₃ ; phthalic ac. (Vol. I). — Mic. rhombohedra. I.c. aq., alc., eth. (Boiling w. aq. or alc. gives NH ₃ & phthalimide.)
2530	219	3,6-Dimethylcarbazole, $[\text{C}_6\text{H}_5(\text{Me}).\text{C}_6\text{H}_4(\text{Me}).\text{NH}]$. — \textcircled{P} Shaken w. sol. of benzoquinone in gl. ac. ac. + drop conc. H ₂ SO ₄ gives indigo-blue sol. — Ndl. fr. bz. E.s. alc., bz.; mod. s. eth. — B.Pk (T. 2.23), m.p. 192°.
2531	219	α -Dimethyluracil, $[\text{CO.NMe.CMe}:\text{CH.CO.NH}]$. — Glassy lft. fr. abs. alc. S. aq.; d.s. alc.; i. eth.
2532	220-1d.	† Semioxamazide, $\text{NH}_2\text{CO.CO.NH.NH}_2$. — \textcircled{P} Reduces Tollen's reagt. immediately in T. 2.30, & gives NH ₃ in T. 2.27. — Lft. fr. h. aq. S. in 400 pt. aq. at 19°; e.s. h. aq.; i. alc., eth.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2533	220	Adipamide, $\text{NH}_2\text{CO}(\text{CH}_2)_6\text{CO}\text{NH}_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; adipic ac. (Vol. I). — S. in 227 pt. aq. at 12.2° .
2534	221-2	α -Aminostearic Ac., $\text{C}_{18}\text{H}_{32}\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$. — Cryst. powd. fr. h. ac. ac.; e.s. h. gl. ac. ac.; i. aq., alc., eth.
2535	221; 225	β -Naphthylsemicarbazide, $\text{C}_{10}\text{H}_7\text{NH.NH.CO.NH}_2$. — \textcircled{P} Semicarbazide (T. 2.17). — Lft. alm. i. c. aq.; d.s. c. alc., eth., bz.
2536	220-5d. (r.h.)	d,L-Mannosaccharicacidbisphenylhydrazide, $\text{Ph.NH.NH.CO}(\text{CH}_2\text{OH})_4\text{CO-NH.NH.Ph}$. — Hydrazide prob. giving coloration in T. 2.11. — Colorless lft. Alm. i. aq.
2537	222u.c.; 228c.	Tetramethyluric Ac., 1,3,7,9-Tetramethyl-2,6,8-trioxypurine, $\text{C}_6\text{H}_12\text{O}_4\text{N}_4$. — \textcircled{P} Gives the murexide react., T. 2.20! — Ndl. fr. alc. S. in 39 pt. aq. at 20° , or 3 pt. h. aq., or 27 pt. h. alc. Taste v. bitter. Without acid properties. Boiling w. KOH sol. gives methylamine (No. 2.1059).
2537-1	abt. 223u.c., d. (r.h.); 228c.	L-Serine, α -Amino- β -hydroxypropionic Ac., $\text{CH}_3(\text{OH}).\text{CH}(\text{NH}_2).\text{CO}_2\text{H}$. — [Proteolytic product.] — \textcircled{P} \textcircled{D} Yields 13.33% α -amino N in T. 2.5. — 6-sided tbl. or pr. fr. c. aq. S. in 3 pt. aq. Tastes sweet. Easily racemized. $[\alpha]_D^{20} = -6.83^\circ$ (in 10% sol.) — Begins to darken abt. 207° u.c. Melts w. efferv.
2538	223	Malonanilide, $\text{Ph.NH.CO.CH}_2\text{CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; malonic ac. (Vol. I). — Ndl. fr. alc. i. aq., eth.
2539	223-4	Glutaranilide, $\text{Ph.NH.CO}(\text{CH}_2)_3\text{CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; glutaric ac. (Vol. I). — Ndl. fr. alc.
2540	222; 226	d-Camphoranilide, $\text{C}_9\text{H}_{14}(\text{CO.NH.Ph})_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: aniline; camphoric ac. (Vol. I). — Ndl. fr. h. gl. ac. ac. V.d.s. alc., etc.
2541	224	Phenylloxamide, Ph.NH.CO.CO.NH_2 . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; aniline; oxalic ac.
2542	221; 224	Diacetyl-2,4-toluylenediamine, $(\text{Me.CO.NH})_2\text{C}_8\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 2,4-toluylenediamine (No. 2.744); ac. ac. — Ndl. fr. h. aq. HNO_2 (sp. gr. 1.47) gives nitro deriv., ndl. fr. acetone, m.p. 253° .
2543	224	Dibenzoyl-2,4-toluylenediamine, $(\text{Ph.CO.NH})_2\text{C}_8\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 2,4-toluylenediamine (No. 2.744); benzoic ac. — Tbl. fr. gl. ac. ac. D.s. alc.
2544	224-5	4-Acetaminocresol(2), $\text{Me.CO.NH.C}_6\text{H}_4\text{Me.OH}$. — \textcircled{P} \textcircled{D} Sapn. T.* products: 4-aminocresol(2), (No. 2.910); ac. ac. — Sbl. in lft. D.s. c. aq.; v.s. alc., NaOH sol.
2545	224	Triphenyl Cyanurate, $(\text{PhCON})_3$. — Ndl. fr. alc. Alm. i. aq., eth.; s. bz. ["Scarcely attacked by boiling acid or alk."] In sealed tube at 180° gives phenol & cyanuric ac.
2546	225	Phenyl- α -naphthylcarbazole, $[\text{C}_6\text{H}_5\text{C}_10\text{H}_8\text{NH}]$. — Cryst. fr. alc., bz., lgr. D.s. c. alc.; e.s. h. bz.
2547	225	2,6-Dimethylpyridone, Lutidone, $[\text{CO.CH:CM}_2\text{NH.CMe:CH}]$. — Lust. ndl. or pointed monoclin. pyramids fr. aq. E.s. aq., alc.; alm. i. eth., chlf., bz. B.p. $349-51^\circ$. — B.Pk (T. 2.23), ppt. of fine yel. ndl., m.p. $219-20^\circ$.
2548	225u.c. (232c.)	Arecaidin, N-Methyl- Δ^1 -tetrahydronicotinic Ac., $\text{C}_8\text{H}_7\text{NMe.CO}_2\text{H}$. — [Fr. seeds of betel nut, Areca Catechu.] — Tbl. fr. alc. Cryst. w. $1\text{H}_2\text{O}$ & then melts after drying at $222-3^\circ$ (c. d. E.s. aq., dil. alc.; i. eth., bz., chlf. ["Aq. sol. reacts neut., or when conc., feebly ac."]) — $\text{B}_2\text{H}_6\text{PtCl}_6$ (T. 2.14), yel. octahedra, m.p. 225° c. (r.h.) w. efferv.
2549	225-6	Cyanphenylhydrazine, Diamidrazon, $\text{Ph.NH.N:C}(\text{NH}_2).\text{C}(\text{NH}_2):\text{N.NH.Ph}$. — \textcircled{P} Reduces Fehling's sol. Dis. in conc. H_2SO_4 w. indigo-blue color! — Lft. fr. alc. i. aq. D.s. h. alc., chlf.
2550	225	Tetranitrobenzophenone, $(\text{NO}_2)_4\text{C}_6\text{H}_2\text{O}$. — \textcircled{P} Nitro comp.† — Small tbl. fr. gl. ac. ac. V.d.s. gl. ac. ac.; i. bz.
2551	226	Dinitrooctylbenzene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{C}_8\text{H}_{17}$. — \textcircled{P} Nitro comp.† — S. h. alc.
2552	227	Succinanilide, $\text{Ph.NH.CO.CH}_2\text{CH}_2\text{CO.NH.Ph}$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. HCl in sealed tube at 100°) products: aniline; succinic ac. (Vol. I). —

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17. —

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		Ndl. fr. alc. I. aq.; e.s. alc., eth.; s. undecd. in conc. H_2SO_4 or HNO_3 . "Not attacked by alc. KOH." C. fuming HNO_3 gives dinitro deriv., m.p. 260°.
2553	227	Benzoyl-p-aminophenol, $Ph.CO.NH.C_6H_4OH$. — \textcircled{P} \textcircled{D} Sapn. T.* products: p-aminophenol (No. 2.963); benzoic ac. — Hair-like ndl. fr. gl. ac. ac. Alm. i. aq., bz., chlf., lgr.
2554	228(w. sbl.)	Methyloxamide, $MeNH.CO.CO.NH_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_2 ; methylamine (No. 2.1059); oxalic ac.
2555	228.5(u.c.)	† Phthalimide, $O^{\prime}-CO.C_6H_4.CO.NH^{\prime}$. — \textcircled{P} (1) Gives NH_2 in T. 2.7. — (2) Heat 2 to 3 min. at 100–110° 0.05 g. substance, 0.5 g. resorcinol, & 10 cc. conc. H_2SO_4 . Sol. after 30–40 sec. becomes YOSI, & finally alm. opaque, w. color appearing OR by strong direct light fr. sky, or BGS2 by reflected light! ("Phthalimide blue.") D.R.P. 44,268. — Flat ndl. fr. slowly cooling sol. Odorless. Tasteless. Alm. i. aq., bz., lgr.; s. strong KOH sol. [$k_A 10^6$ = 5] Sbl. — \textcircled{D} Heat to boiling for some sec. 1 cc. conc. ammon. (sp. gr. 0.90) & 0.05 g. substance. Set aside 30 min. Filter off the white powd. which separates & wash w. 0.5 cc. ammon. Dry on tile 10 min. at 100°. The product, phthalamide, is obtained in mic. rhombohedra, which melt w. loss of NH_2 at 218–20° u.c. & then solidifies, through formation of phthalimide, to melt again at 227–8° u.c.
2556	228–30	Hemipinicimide, $[NH.CO.C_6H_4(MeO)_2.CO]$. [$(CO)_2:(MeO)_2 = 1:2:3:4$]. — \textcircled{P} Alc. or dil. aq. sol. fluor. blue! — Sapn. T.* (w. alk.) products: NH_2 & hemipinic ac. ($C_{10}H_{10}O_6$, Vol. I). — Fine ndl. fr. alc. S. h. aq.; s. NaOH (w. yel. color which disappears after some min.); not s. Na_2CO_3 sol. — AgA, ppt. i. alc., eth.
—	229 ± 2d.	Allantoin. — Cf. No. 2.413.
2557	abt. 230 (r.h.)	d,l-Mannonicacidphenylhydrazide, $[CH_2.OH.(CH.OH)_4.CO.NH.NH.Ph]$. — \textcircled{P} Phenylhydrazide, prob. giving coloration in T. 2.11. — D.s. aq., alc.
2558	230	† Nitroguanidine, $NO_2.NH.C(:NH).NH_2$. — \textcircled{P} Loses NH_2 in melting. — Odorless ndl. fr. aq. V.d.s. c. aq., alc.; i. eth.; e.s. KOH sol. [$k_B 10^{14}$ 2.1].
2559	229; 233	pp'-Dinitrobiphenyl, $NO_2.C_6H_4.C_6H_4.NO_2$. — \textcircled{P} Nitro comp.† — Fine ndl. S. h. alc. — Sn & HCl reduces to benzidine, No. 2.840.
2560	230–2	† 2,4,6-Trinitromesitylene, $(NO_2)_3.C_6Me_2$. — \textcircled{P} Nitro comp.† — Ndl. alm. i. c. alc.; d.s. h. alc.
2561	229.5–30.5 u.c.	† Caffeine, 1,3,7-Trimethyl-2,6-dioxypurine, $C_8H_{10}O_2N_4$. — Cryst. fr. h. aq. w. 1 H_2O in hair-like ndl. 100 pt. of each of following solvents dis.: aq. at 16°, 1.35 pt., or 45.5 pt. at 65°; 85% alc. at 16°, 2.3 pt.; eth. at 16°, 0.044 pt.; chlf., 12.97 pt. — Odorless. Taste of powder slightly but distinctly bitter. — \textcircled{P} Gives the murexide react., T. 2.20! — \textcircled{D} Dis. 0.05 g. substance in 7 cc. warm aq. Cool. Add 2 cc. sol. $HgCl_2$ sol. After a few min. long white ndl. separate, completely filling the sol. Filter. Wash w. 3 cc. c. aq. Recryst. fr. 3 cc. h. aq. The product, $C_8H_{10}O_2N_4.HgCl_2$, after drying, sinters w. sl. blackening at 241° & melts at 243.5° u.c. (251° c.).
2562	230	Oxal-s-pseudocumadide, $Me_2.C_6H_5.NH.CO.CO.C_6H_5.Me_2$. — \textcircled{P} \textcircled{D} Sapn. T.* products: s-pseudocumidine (No. 2.646); oxalic ac. — Ndl. fr. gl. ac. ac.
2563	230–1d. u.c.	† Anhydroecgonine, Tropenecarbonic Ac.(2), $C_8H_{10}O_2N$. — \textcircled{P} 1 cg. gives unsaturation T. No. 1.304 w. $KMnO_4$ & 2.901 w. Br. — Cryst. fr. MeOH + eth. V.s. aq.; alm. i. eth., chlf., lgr., bz. — $B_2H_6PtCl_6$, compact yel.-red pr., m.p. 223° d.
—	sbl. 230	d-Isoleucine. — Cf. No. 2.2629.
2564	231	α -Naphthylsemicarbazide, $C_{10}H_7.NH.NH.CO.NH_2$. — \textcircled{P} Semicarbazide. § — Lust. lft. I. aq., eth.; v.s. c. alc., dil. alk.
2565	231–2; 233–4	Carboxyldiurea, $(NH_2.CO.NH_2.CO)$. — \textcircled{P} \textcircled{D} Sapn. T.* (w. NaOH) products: NH_2 ; cyanuric ac. (Heat alone gives same products.) — Micaceous scales fr. h. aq. D.s. c. aq.; alm. i. alc.; i. eth.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2566	231	4,4'-Diacetamino-3,3'-dimethoxybiphenyl, Diacetyl-o-anisidine, ($\text{Me.CO.NH}(\text{MeO})\text{C}_6\text{H}_4\text{C}_6\text{H}_4(\text{MeO})(\text{Me.CO.NH})$). — \textcircled{P} \textcircled{D} Sapon. T.* products: o-anisidine (No. 2.1332); ac. ac. — Pr. fr. gl. ac. ac. I. aq.; d.s. alc., bz.; s. chlf., gl. ac. ac.
2567	232	Ethylenediphthalimide, $[\text{CO.C}_6\text{H}_4\text{CO.N}]^{\text{—}}\text{CH}_2\text{CH}_2\text{N.CO.C}_6\text{H}_4\text{CO}^{\text{—}}$. — \textcircled{P} \textcircled{D} Sapon. T. (w. conc. HCl at 200°) products: ethylenediamine (No. 2.1130); phthalic ac. (Vol. I). — Lust. ndl. fr. gl. ac. ac.
2567-I	233; 231	Kyaphenin, (PhCN). — Trimet. pr. Sbl. Dist. a. 350° undecd. I. aq., dil. ac.; alm. i. c. alc., eth.; v.d.s. h. alc.; e.s. h. toluene. Unchanged by boiling KOH sol. in aq. or alc. — Heated 4 or 5 hr. w. HNO_3 (sp. gr. 1.5) gives trinitro deriv., ndl. fr. HNO_3 , m.p. 250–60° d.
2568	$d.233 \pm 2u.c.$	† Glycocol, Glycine, Aminoacetic Ac., $\text{NH}_2\text{CH}_2\text{CO}_2\text{H}$. — Begins to blacken at 225° u.c., decomposing w. efferv. abt. 235°. Monoclin. cryst. s. in 4.3 pt. c. aq.; alm. i. alc.; i. eth. Odorless. Taste of powder distinctly sweet!! — $[k_A]10^{10} = 1.8$; $k_B10^{12} = 2.8$. The interesting behavior of this amphoteric species in the generic titration tests is described in a foot-note of Chapter 2.] \textcircled{P} (1) Gives N gas in T. 2.4. Quantitative determin. of α -amino N by T. 2.5 gives, according to Van Slyke, 19.98%, instead of the theoretical 18.67%. — (2) Boil sol. of 0.2 g. substance in 1 cc. aq. for nearly 1 min. w. powd. CuO . Filter the deep blue sol. hot. Silky sky-blue ndl. of $\text{CuA}_2\text{H}_2\text{O}$ separate after cooling. — (3) If a drop of phenol be added to a glycine sol. & then sodium hypochlorite, a fine blue color soon develops. (Ber., 8, 699.) [The color phenomena in the two foregoing tests are not specific, being also given by other amino acids.] \textcircled{D} (a) Dis. 0.1 g. substance in 1 cc. aq. Add 3 cc. c. sat. alc. sol. picric ac. After v. vigorous shaking (and then only) an abundant pptn. of yell. cryst. appears. Filter. Wash w. 2 cc. 50% alc. Dry on tile 15 min. at 100°. The resulting picrate, B.Pk, blackens & decomposes w. efferv. at 198–201° u.c. (202.5–5.5° c.). — (b) Benzoylate w. benzoyl chloride & alk. to hippuric ac. (No. 2.304), by Schollen-Baumann reaction, T. 2.27-c or d. — (c) T. 2.31 gives phenylureidoacetic ac. (No. 2.318), & γ -phenylhydantoin (No. 2.2248).
2568-I	233	Nor-l-ecgonine, $\text{C}_7\text{H}_{10}(\text{OH})(\text{CO}_2\text{H})(:\text{NH})$. — Ndl. fr. MeOH sol. (upon addition of eth.). — B.HAuCl ₄ .H ₂ O (T. 2.13), yell. ndl. fr. aq., m.p. 211°.
2569	234. 5u.c.	† Dimethylglyoxime, Diacetylloxime, $\text{Me.C}(:\text{NOH}).\text{C}(:\text{NOH}).\text{Me}$. — Colorless cryst. fr. h. 50% alc. I. aq.; e.s. alc., eth. — \textcircled{P} Add 0.5 cc. 10% NiCl_2 sol. to sol. of 0.01 g. substance in 5 cc. 66% alc. A floc. ppt. of nickel-dimethylglyoxime, $\text{NiC}_6\text{H}_5\text{O}_2\text{N}_2$, of ORT1 color, while in suspension, or R-RT1 when examined moist on filter paper, is obtained! The ppt. is s. in abs. alc. & mineral acids, but v.d.s. 50% alc. When dry it is said to sublime at 250°.
2570	234	Tetraphenylsuccinamide, $[\text{NPh}_2\text{CO.CH}_2\text{CH}_2\text{CO.NPh}_2]^{\text{—}}$. — \textcircled{P} \textcircled{D} Sapon. T.* ("w. v. conc. KOH") products: diphenylamine (No. 2.1568); succinic ac. (Vol. I). — Lust. ndl. fr. alc. E.s. chlf.; d.s. c. alc., bz., eth., lgr.
2571	234	1,2-Diacetaminonaphthalene, ($\text{Me.CO.NH}.\text{C}_10\text{H}_8$). — \textcircled{P} \textcircled{D} Sapon. T.* products: 1,2-diaminonaphthalene (No. 2.730); ac. ac. — Ndl. fr. alc.
2572	234	Oxalyl- α -naphthalide, $\text{C}_{10}\text{H}_7\text{NH.CO.CO.NH.C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Sapon. T.* products: α -naphthylamine (No. 2.589); oxalic ac. — Ndl. fr. gl. ac. S. in NaOH sol.
2573	235-5	Benzoyladenine, $\text{Ph.CO.C}_6\text{H}_4\text{N}_2$. — \textcircled{P} \textcircled{D} Sapon. T.* products: adenine (No. 2.1057), benzoic ac. — Thin lust. ndl. fr. aq. E.s. h. alc., ammon., dil. acids.
2574	235d.	1-Acetaminonaphthol(2), $\text{Me.CO.NH.C}_{10}\text{H}_8\text{OH}$. — Sapon. T.* products: 1-aminonaphthol(2); ac. ac. — Lft. fr. v. dil. alc.
2575	235	Stachydrin, Dimethylbetaine of α -Proline, $\text{C}_7\text{H}_{11}\text{O}_2\text{N}$. — [In tubers of <i>Stachys tuberifera</i> & leaves of <i>Citrus aurantium</i> .] — \textcircled{P} Heated w. con. KOH evolves dimethylamine (No. 2.1061). Vapors evolved on heating give pyrrole splinter react. (Cf. T. 2.24). — Clear deliq. cryst. w. 1H ₂ O (partly lost over H ₂ SO ₄). Sol. reacts neutral & has unpleasant sweetish taste. V.s. aq., alc.; i. chlf., eth. — B.Pk, ndl., m.p. 195–6°.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. ‡ = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS. — Colorless and Solid.
2576	231-2; 239-40	Dibenzylidenebenzidine, Ph.CH:N.C ₆ H ₄ .C ₆ H ₄ .N:CH.Ph. — Lft. fr. bz. D.s. alc. eth.; e.s. chlf., bz.; i. lgr.
2577	d.235.5-236.5 u.c. (=243-4c.)	Benzildisemicarbazone, Ph.C(:N.NH.CO.NH).C(:N.NH.CO.NH)Ph. — \textcircled{P} Semicarbazone, giving T. 2.17. — Lft. fr. alc. D.s. c. alc.; v.d.s. chlf., bz.
2578	abt. 236	Trinitro-1-ethyl-3,5-dimethylbenzene, (NO ₂) ₃ C ₆ H ₃ EtMe ₂ . — \textcircled{P} Nitro comp. \ddagger — Alm. i. alc., aq.
2579	236-7	p-Tolyloxamide, Me.C ₆ H ₄ .NH.CO.CO.NH ₂ . — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; p-toluidine (No. 2.566); oxalic ac. — Ndl. I. aq.; e.s. alc.
2580	236-7 u.c.	† Carbanilide, Diphenylurea, Ph.NH.CO.NH.Ph. — Odorless, tasteless ndl. fr. boiling 95% alc. V.d.s. aq.; e.s. alc., eth.; d.s. lgr.; s. bz.; s. acetone. \textcircled{P} Gives powerful odor of carbalamine in T. 2.12. — Sapn. T.* products: aniline; CO ₂ . — \textcircled{D} Dis. 0.05 g. substance in 10 cc. boiling alc. Cool. Add 1 drop liquid Br. Shake. Wash ppt. on filter w. 10 cc. c. alc. The ppt., consisting of micro-cryst. ndl. of silky luster, is a bromine substitution deriv. or mixture which sublimes without melting above 290° when dried & heated in t.t. by small flame.
2581	237	ab-Dibenzoylhrazine, Ph.CO.NH.NH.CO.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* (by persistent boiling w. ac. or alk.) products: hydrazine; benzoic ac. (Has only feeble Ag. reducing power before hydrolysis.) — Silky ndl. fr. alc. Alm. i. c. aq., alc., chlf.; s. alk.
2582	237d.	† α -Benzildioxime, Ph.C(:NOH).C(:NOH)Ph. — \textcircled{P} Oxime. § — Micro-cryst. powd. fr. alc. I. aq. 100 pt. alc. at 17° dis. 0.05 pt. Alm. i. eth., gl. ac. ac.; s. conc. NaOH & repptd. by ac. — Heated w. aq. at 200° gives β -deriv., lft., s. alc., m.p. 104-5°.
2583	238	ab-n-Amylphenylurea, C ₉ H ₁₁ .NH.CO.NH.Ph. — \textcircled{P} \textcircled{D} Sapn. T.* products: amyramine; aniline; CO ₂ . — Pr. i. aq.
2584	238.5 u.c.	† Carbazole, [C ₆ H ₄ .NH.C ₆ H ₄]. — Cryst. fr. h. alc. or toluene in lft. w. pearly luster, sl. violet fluor. & faint indole-like odor. Tasteless. B.p. 351.5° c. 100 pt. alc., at 14°, dis. 0.92 pt.; at b.p., 3.88 pt. 100 pt. toluene, at 16.5° dis. 0.55 pt.; at 100°, 5.46 pt. I. aq.; d.s. c. bz., eth., CS ₂ , chlf., gl. ac. ac. \textcircled{P} (1) Gives deep red (R) color in pyrrole pine splinter react. T. 2.24! — (2) 0.01 g. substance fused w. 0.1 g. oxalic ac. gives blue mass ("carbazole blue"), s. w. VB color in 10 cc. h. gl. ac. ac. — (3) 0.001 dis. in conc. H ₂ SO ₄ w. YGT2 color, becoming deep BG-G w. 1 drop HNO ₃ ! \textcircled{D} Dis. 0.1 g. carbazole in 4 cc. boiling toluene. Add 0.15 g. picric ac. disd. in 2 cc. h. toluene. Wash cryst. which separate on cooling w. 5 cc. c. toluene. Recryst. fr. 2 cc. boiling toluene, shaking & cooling. Wash w. 1 cc. toluene. Dry 15 min. at 100°. The resulting picrate, B.Pk, is obtained in fine RO — ndl., m.p. 181-2° u.c.
2585	237; 240	† Guanosin, Vernin, C ₁₀ H ₁₃ O ₄ N ₅ .2H ₂ O. — [Fr. hydrolysis of vegetable nucleic acids & in pancreas] — \textcircled{P} Gives purple ring in first part of Molisch react. (Vol. I, Chap. IV), but no ppt. w. aq. or NH ₃ in second part of test. — Thin, lust., tyrosine-like ndl. or flat pr. D.s. c. aq.; e.s. h. aq.; s. alk. or dil. min. acids; i. alc. Does not reduce Fehling's sol. $[\alpha]_D^{20} = -60^\circ$ (3% sol. in N/10 NaOH). — Pptd. fr. acid sol. by phosphotungstic ac. AgNO ₃ gives gelat. ppt. s. in ammon. — Picrate, spheroidal cryst. aggregates, m.p. abt. 185° d.
2586	238.5	Dinitroisoquinoline, (NO ₂) ₂ C ₆ H ₃ N. — \textcircled{P} Nitro comp. \ddagger — Lust. cryst. fr. gl. ac. ac. D.s. alc. — B ₂ H ₂ PtCl ₆ , reddish ppt.
2587	abt. 240d.	Mucicacidbisphenylhydrazide, Ph.NH.NH.CO.(CH.OH) ₄ .CO.NH.NH.Ph. — \textcircled{P} Hydrazide. — Lft. fr. phenylhydrazine. V.d.s. common solvents.
2588	abt. 240d.	Anthraniilcarbonic Ac., Isatoic Anhydride, [C ₆ H ₅ .NH.CO.O.CO]. — \textcircled{P} Sol. in c. 1% NaOH w. transient blue fluor. — Monoclin. tbl. fr. acetone. S. in 23 pt. h. acetone; d.s. common solvents. Evolves CO ₂ in melting. ["Does not redden blue litmus paper at once, & is only v. slowly sol. in Na ₂ CO ₃ sol."] Long boiling w. aq. gives CO ₂ & anthranilic ac. (No. 2.148).
—	d. abt. 240	p-Benzquinonedioxime. — Cf. No. 2.3821.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2589	abt. 240d.	d,L-Serine, α -Amino- β -hydroxypropionic Ac., $\text{CH}_2(\text{OH})\cdot\text{CH}(\text{NH}_2)\cdot\text{CO}_2\text{H}$.—[Usually obtained fr. proteolysis in this racemic form.]—Thin irregular lft. S. in 3–4 pt. h. aq., or 23.1 pt. at 20°. Taste, sweet. Begins to brown fr. 207° u.c. — P_2O_5 gives N in T. 2.4. Percentage of α -amino N by T. 2.5 is 13.33! — (1) Convert into β -naphthalenesulphonyl deriv., $\text{C}_{10}\text{H}_7\text{SO}_2\text{NH}\cdot\text{CH}(\text{CH}_2\text{OH})\cdot\text{CO}_2\text{H}$, ndl. fr. alc., s. in 70–80 pt. boiling aq., or 7 pt. h. alc., m.p. 210° u.c. (Ber., 35, 3784–3785). — (2) Convert into corresponding phenylisocyanate deriv. of m.p. 165–6° u.c. & its hydantoin. (Cf. T. 2.31 & Ber., 25, 3787–3805.)
2590	240	Dibenzoyl-m-phenylenediamine, $\text{Ph}\cdot\text{CO}\cdot\text{NH}\cdot\text{C}_6\text{H}_4\cdot\text{NH}\cdot\text{CO}\cdot\text{Ph}$.— P_2O_5 Sappn. T.* products: m-phenylenediamine (No. 2.634); benzoic ac.—Matted ndl. fr. gl. ac. ac. D.s. alc.; more s. gl. ac. ac.
2591	242–3	s-Succinamide, $\text{NH}_2\cdot\text{CO}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{CO}\cdot\text{NH}_2$.— P_2O_5 Sappn. T.* products: NH_2 ; succinic ac. (Vol. I).—Ndl. S. in 220 pt. aq. at 15°, or in 9 pt. at 100°; i. eth.
2592	d.240–5	p-Nitro-d,L-phenylalanine, $\text{NO}_2\cdot\text{C}_6\text{H}_4\cdot\text{CH}_2\cdot\text{CH}(\text{NH}_2)\cdot\text{CO}_2\text{H}$.— P_2O_5 Nitro comp.‡—Browns abt. 220°. Cryst. w. $\frac{1}{2}\text{H}_2\text{O}$ (lost over H_2SO_4). Anhydrous pr. fr. alc. ["Reacts neut."] Taste, bitter-sweet. R.d.s. c. aq.; d.s. c. alc.; e.s. ammon.
2593	242d. (r.h.)	β -Amino- α -hydroxypropionic Ac., Isoserine, $\text{NH}_2\cdot\text{CH}_2\cdot\text{CH}(\text{OH})\cdot\text{CO}_2\text{H}$.—Long thin monoclin. pr. S. in abt. 65.35 pt. aq. at 20°; e.s. h. aq.—Benzoyl deriv. (T. 2.27 w. benzoyl chloride + NaOH at 0°), pr. fr. h. aq., m.p. 151° c.—Phenylureido deriv. (T. 2.31), tbd., m.p. 183–4° d. (c.).
2594	244.5u.c.	Dibenzoylethylenediamine, $\text{Ph}\cdot\text{CO}\cdot\text{NH}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{NH}\cdot\text{CO}\cdot\text{Ph}$.— P_2O_5 Sappn. T.* (w. dil. NaOH) products: ethylenediamine (No. 2.1130); benzoic ac.—Pr. fr. h. alc.
2595	244–5c.	Camphorimide, $\text{C}_{10}\text{H}_{14}\text{O}_2\text{NH}$.— P_2O_5 Sappn. T.* products: NH_2 ; camphoric ac. (Vol. I).—B.p. 300°, undecd. S. in 10% NaOH & reproto. by CO_2 ! S. in 150 pt. aq. at 15°; e.s. h. alc., eth. $[\alpha]_D = -10.6^\circ$.—AgA, ppt., d.s. boiling aq., stable in light.
2596	245u.c.	† Oxanilide, $\text{Ph}\cdot\text{NH}\cdot\text{CO}\cdot\text{CO}\cdot\text{NH}\cdot\text{Ph}$.— P_2O_5 Sappn. T.* products: aniline; oxalic ac.—Lust. scales fr. h. xylene. D.s. h. alc.; s. h. bz.; i. h. aq., eth. B.p. 320°; a. 360°.—(Gives powerful carbylamine odor in T. 2.12!)!
2597	245–6	Tetraphenylpyrazine, Amaron, $\text{C}_{22}\text{H}_{20}\text{N}_2$.— P_2O_5 Sol. in conc. H_2SO_4 , blood-red!—Lust. ndl. fr. gl. ac. ac. Sbl. D.s. h. alc., eth.; e.s. chlf., h. bz.; i. dil. ac., alk.—Fuming HNO_3 gives tetranitro deriv., s. bz.; i. eth., m.p. 130–40°.
2598	245	Bisantipyrine, $\text{C}_{22}\text{H}_{22}\text{O}_4\text{N}_4$.—Cryst. fr. wood spirits. Alm. i. aq., eth.; d.s. alc., toluene; e.s. chlf.—Picrate (T. 2.23), ndl. fr. alc., m.p. 161°.
2599	246–7d.	† Benzalbismethylketole, $\text{C}_{12}\text{H}_{14}\text{N}_2$.—Lft. fr. acetone. I. aq.; v.d.s. h. alc., eth.— FeCl_3 in gl. ac. ac. sol. oxid. to dimethylrosindole.
2600	249–50	α -Hydroxy- β -phenyl- α -aminopropionic Ac., α -Tyrosine, $\text{HO}\cdot\text{C}_6\text{H}_4\cdot\text{CH}_2\cdot\text{CH}(\text{NH}_2)\cdot\text{CO}_2\text{H}$.— P_2O_5 Gives Millon's react. (T. 2.19). Gives violet coloration w. FeCl_3 . T. 2.4 w. nitrous ac. should yield N.—S. in 500 pt. aq.; i. alc.
2600-I	251d.	rac-Ecgonine, Ψ -Tropine-C-carbonic Ac., $\text{C}_{10}\text{H}_{15}\text{O}_3\text{N}$.—[Synthetic product.]—Cryst. w. $2\text{H}_2\text{O}$ (lost in vacuo over H_2SO_4) in monoclin. 6-sided tbd. V.s. aq.; d.s. h. alc. Reacts neutral. Stable in KMnO_4 sol., acidified w. H_2SO_4 .— BAuCl_4 (T. 2.13), ndl. without cryst. aq., m.p. 213°.
2601	250; 243; 256	Di-o-tolylurea, $(\text{Me}\cdot\text{C}_6\text{H}_4\cdot\text{NH})_2\text{CO}$.— P_2O_5 Sappn. T.* products: o-toluidine, (No. 2.1262); CO_2 .—Ndl. i. aq.; d.s. h. alc., bz. Sbl.
2602	253	Aminomyristic Ac., $\text{NH}_2\cdot\text{C}_{11}\text{H}_{22}\cdot\text{CO}_2\text{H}$.—Cryst. powd. fr. h. gl. ac. ac. I. aq., alc., eth.
2603	255u.c.	Succin-p-toluide, $\text{Me}\cdot\text{C}_6\text{H}_4\cdot\text{NH}\cdot\text{CO}(\text{CH}_2)_2\cdot\text{CO}\cdot\text{NH}\cdot\text{C}_6\text{H}_4\cdot\text{Me}$.— P_2O_5 Sappn. T.* products: p-toluidine (No. 2.566); succinic ac. (Vol. I).—Cryst. i. h. aq., c. alc.
2603-II	257d.	d- Ψ -Ecgonine, $\text{C}_{10}\text{H}_{15}\text{O}_3\text{N}$.—[Fr. rearrangement of ordinary L-ecgonine by h. conc. KOH, but not optical antipode of the latter compound.] Lust.

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.17.

No.	Melting-point (C.).	NEUTRAL COMPOUNDS.—Colorless and Solid.
		oblique pr. & t-bl. fr. alc. without cryst. aq. Less s. in abs. alc. than ordinary ecgonine.—B.HAuCl ₄ (T. 2.13), lemon-yel. ndl. or lft. without cryst. aq., m.p. 220°.
2504	259	α -Tetranitronaphthalene, (NO ₂) ₄ .C ₁₀ H ₄ . — (P) Nitro comp.† Detonates violently when heated! — Rhomb. cryst. fr. chlf. Alm. i. c. alc. & common solvents.
2605	abt. 260 ± 10 u.c. d.	† 1-Tryptophane, β -Indole- α -aminopropionic Ac., C ₁₁ H ₁₂ O ₂ N ₂ . — [Proteolytic product.] Begins to yellow abt. 30° below m.p. & does not melt sharply. The m.p. varies greatly according to rate of heating. The m.p. here given approximates the value that will be found for the temperature of complete fusion when heating at the usual rate (Cf. Vol. I, p. 220). With rapid heating, according to Abderhalden, fusion occurs abt. 280° c. — 6-sided & rhomb. silky lft. D.s. c. aq.; e.s. h. aq.; d.s. abs. alc. [α] _D = abt. -30° (for aq. sol.). (Easily racemized.) Odorless. Taste faintly bitter. (P) (1) 5 mg. ignited in t.t. gives strong fecal odor of skatole! — (2) Reddens pine splinter in T. 2.24-b. — (3) Add to aq. tryptophane sol. Br-water, carefully avoiding any x.s. A $\leftarrow RT^2$ color, or in more conc. sol. a ppt. of dull VRT^2 color (as viewed on filter), appears. Or, dis. 3 mg. tryptophane in 1 cc. warm aq. Cool. Add 3 or 4 drops fresh saturated Br-aq. Shake out w. 10 cc. ethyl acetate. A deep VR sol., showing greenish fluor. & charac. absorption spectrum (Biochem. Z., 24, 423) is obtained!! — (4) Gives the xanthoproteic & Adamkiewicz reactions (T. 2.32 & 2.33!). (5) Gives 6.86% amino-N in T. 2.5.
		① B.Pk (T. 2.23), carmine-red cryst., d.s. c. aq., m.p. 195-6°. — B.C ₁₀ H ₈ O ₂ N ₄ (picrolonate), or.-red ndl. v.d.s. aq.; m.p. 203-4° d. — Benzenesulphonyl deriv., C ₁₁ H ₁₁ O ₂ N ₂ .SO ₃ Ph (Z. physiol. Chem., 55, 22), ndl. fr. dil. alc., m.p. 185° d. — Phenylureido deriv., C ₁₁ H ₁₁ O ₂ N ₂ .CO.NH.Ph (Cf. T. 2.31), ndl. fr. methyl alc. reddening in sunlight, m.p. 166°. [d,l]-Tryptophane differs little fr. the l-comp. in m.p. or chemical reactions, but is opt. i. & has a sweetish taste.]
2606	260d.	α -Methylaminopropionic Ac., Me.CH(NH.Me).CO ₂ H. — (P) Tastes sweet. — Rhomb. pr. S. aq.; alm. i. c. abs. alc.
2607	260	β -Dimethyluracil, $^5\text{CO}.\text{NMe}.\text{CO}.\text{CH}:\text{CM}\text{e}.\text{NH}^6$. — Thick glassy pr. E.s. h. aq.; d.s. c. aq., alc.; alm. i. eth. — [k ₄ .10 ¹¹ = 7.4.]
2600	260	ab-Di-p-tolylurea, (Me.C ₆ H ₄ .NH) ₂ CO. — (P) (D) Sapn. T.* products: p-toluidine (No. 2.566); CO ₂ . — Ndl. I. aq.; d.s. c. alc.
2609	259; 265	Dibenzoyl-o-tolidine, Ph.CO.NH.C ₆ H ₄ .Me.C ₆ H ₄ .Me.NH.CO.Ph. — (P) (D) Sapn. T.* products: o-tolidine (No. 2.849); benzoic ac. — Ndl. fr. quinoline + alc. I. common solvents.
2610	263	Oxal-p-toluide, Me.C ₆ H ₄ .NH.CO.CO.NH.C ₆ H ₄ .Me. — (P) (D) Sapn. T.* products: p-toluidine (No. 2.566); oxalic ac. — Sbl. D.s. h. alc., bz., chlf.; e.s. h. gl. ac. ac.
2611	263-4	3,4-Dibenzoylaminotoluene, (Ph.CO.NH) ₂ .C ₆ H ₄ .Me. — (P) (D) Sapn. T.* products: 3,4-toluylenediamine (No. 2.707); benzoic ac. — Ndl. fr. gl. ac. ac. I. aq.; s. h. alc.
2612	263	Diphenyldiacipiperazine, $^5\text{CO}.\text{NPh}.\text{CH}_2.\text{CO}.\text{NPh}.\text{CH}_2^6$. — Ndl. I. aq., eth., chlf.; e.s. h. gl. ac. ac. Sbl.
2612-I	264-6d.	2,4,2',4'-Tetranitrostilbene, (NO ₂) ₂ .C ₆ H ₄ .CH:CH.C ₆ H ₄ .(NO ₂) ₂ . — V. fine ndl. fr. gl. ac. ac. I. eth.
2613	266	Succin- β -naphthalide, C ₁₀ H ₈ .NH.CO.CH ₂ .CH ₂ .CO.NH.C ₁₀ H ₇ . — (P) (D) — Sapn. T.* products: β -naphthylamine (No. 2.781); succinic ac. (Vol. I). — Mic. cryst. fr. nitrobenzene. I. common solvents.
—	abt. 268d.	d,l-Phenylalanine. — Cf. No. 2.478-1. [Weak acid.]
—	268-9(in s.t.)	Pyrocoll. — Cf. No. 2.3890. [Yellowish. Fr. dry distn. of gelatine.]
2616	271-2d.	Guvaccine, C ₁₀ H ₁₂ O ₂ N. — [Alkaloid fr. Areca Catechu.] — Lust. cryst. S. aq., dil. alc.; i. other solvents. Sol. neut. Darkens abt. 265°. — B ₂ H ₅ PtCl ₆ .4H ₂ O, 6-sided overlapping t-bl., darkening at 210° & melting w. frothing a few degrees higher. — B.HAuCl ₄ (T. 2.13), flat pr. fr. dil. HCl, m.p. abt. 194-5°.

No.	Melting-point (C°).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2617	271d. u.c.; 290-2d.u.c. (r.h.)	† d-Alanine, α -Aminopropionic Ac., $\text{Me}.\text{CH}(\text{NH}_2).\text{CO}_2\text{H}$.—[Proteolytic product.] — Large cryst. of rhombic system on spontan. evapn. of aq. sol. Melts w. efferv. to yel. liquid. S. in 4-5 pt. c. aq.; alm. i. abs. alc. † Tastes sweet. $[\alpha]_D^{20} = +2.7^\circ$ (0.80 g. disd. in aq. to 8.0014 g.), or $+10.4 \pm 0.2$ (0.4452 g. disd. in aq. w. 1 mol. HCl to weigh 6.2513 g.). ② Evolves N in T. 2.4, giving 15.73% in T. 2.5! — $\text{Cu}\bar{\text{A}}$, (prepared by disg. CuO in aq. sol. & concentrating), 6-sided lft. or pr. mod. s. aq., containing 26.52% Cu. — Benzoyl deriv. (No. 2.165), prepared by T. 2.27-d by shaking 3 g. disd. in 30 cc. aq. w. 22 g. NaHCO_3 , adding 14.5 g. benzoyl chloride, acidifying sol., & removing benzoic ac. by boiling out w. lgr.; m.p. $150-1^\circ$ c., s. in 85 pt. c. aq. — β -Naphthalenesulphonyl deriv., small ndl. fr. aq., sintering at 117° , m.p. $122-3^\circ$. (Cf. Ber., 32, 2458; Monatsh., 23, 1351, 26, 1219). — $\text{Et}\bar{\text{A}}$, b.p. 48° (11 mm.). — Picrolonate (T. 2.23), pr. s. in 62 pt. aq. at 20° , m.p. 214° d.
2618	273	Nitroantipyrine, $^{\text{r}}\text{NPh.NMe.CMe:C(NO}_2\text{).CO}^{\text{l}}$. — Ndl. I. aq., alk.; d.s. conc. mineral ac.
2619	$d.273 \pm 5$ u.c.	† Betaine, Trimethylglycocol, $^{\text{r}}\text{CO.CH}_2\text{NMe}_2\text{O}^{\text{l}}$. — [Abundant in plants.] V. deliq. cryst. mass. V.s. aq. Ppid. by eth. fr. warm alc. sol. Odorless. $[k_A \cdot 10^4 = 1$; $k_B \cdot 10^4 = 8$.] Should not give N w. HNO_3 in T. 2.4. — ② Cryst. on porcelain gives or.-red color w. drop FeCl_3 sol. — ③ Dis. 0.1 g. betaine (or its hydrochloride) in 2 cc. aq. Add 5 cc. sat. alc. picric ac. sol. Filter ppt. which slowly separates. Recryst. fr. 1 cc. h. alc. Dry 15 min. at 100° . The product, B.Pk , forms Y-Y T1 ndl., m.p. $180-1^\circ$ u.c. ($183.5-4.5^\circ$ c.), after softening at lower temperature. — $\text{B}_2\text{C}_6\text{H}_5\text{O}_4\text{N}_4$ (picrolonate), pale yel. ndl. e.s. aq., alc., d. abt. 200° . — $\text{B}_2\text{HAuCl}_4\cdot\frac{1}{2}\text{H}_2\text{O}$ (when recrystd. fr. aq.), thin ndl. or lft., d.s. c. aq. — B_2HCl ("Acidol"), monoclin. tbl., e.s. aq., i. c. abs. alc., d. $227-8^\circ$.
2620	275d.	Glycine Anhydride, $^{\text{r}}\text{CO.NH.CH}_2\text{CO.NH.CH}_2^{\text{l}}$. — Tbl. E.s. h. aq., alc. Browns at 245° . Sbl. in ndl. "Reacts neut."
2621	275	Lactimide, α -Aminopropionic Anhydride, $^{\text{r}}\text{CO.NH.CHMe.CO.NH.CHMe}^{\text{l}}$. — Ndl. or lft. E.s. aq. Sbl. "Indifferent."
2622	d.w.m. 270-80	Methyluracil, $^{\text{r}}\text{CO.NH.CMe:CH.CO.NH}^{\text{l}}$. — Ndl. fr. alc. 100 pt. aq. at 22° dis. 0.74 pt.; alm. i. eth.; s. undecd. in conc. H_2SO_4 .
2623	abt. 275d.	Asparaginimide, $^{\text{r}}\text{NH.CO.CH}(\text{NH}_2)\text{CH}_2\text{CO}^{\text{l}}$. — ② ③ Sapn. T.* products: NH ₄ ; aspartic ac. (No. 2.414). — Ndl. V.d.s. c. aq.; i. alc., eth.
2624	274-5	Triphenyl Isocyanurate, $(\text{PhCON})_3$. — Pr. I. h. aq.; s. h. alc.
2625	275	Tetrinitrotetraphenylmethane, $(\text{NO}_2\text{C}_6\text{H}_4)_4\text{C}$. — ② Nitro comp. † White ndl. S. bz., h. chlf.
2626	275	1,1'-Hydrazonaphthalene, $\text{C}_{10}\text{H}_7\text{NH.NH.C}_{10}\text{H}_7$. — ② May reduce Tollen's reagt. on shaking. — Lft. fr. bz. I. aq.; e.s. alc., eth., bz.
2627	274; 277-8	Oxalylphenylhydrazine, $\text{Ph.NH.NH.CO.CO.NH.NH.Ph}$. — ② ③ Sapn. T.* (w. conc. NaOH) products: phenylhydrazine (No. 2.1369); oxalic ac. — Leafy cryst. D.s. org. solvents; e.s. dil. NaOH; s. conc. H_2SO_4 w. red-violet color. Sbl. alm. undecd.
2628	277	Dinitro-p-diphenylbenzene, $(\text{NO}_2)_2\text{C}_{12}\text{H}_{12}$. — ② Nitro comp. † — Monoclin. pr. fr. nitrobenzene. V.d.s. alc., eth.
—	abt. 273u.c., d.	1-Phenylalanine. — Cf. No. 2.478-1.
2629	280d. (r.h. in s.t.)	d-Isoleucine, α -Amino-sec.-butylacetic Ac., $\text{Et.CHMe.CH}(\text{NH}_2).\text{CO}_2\text{H}$. — [Proteolytic product, in plants & beet molasses.] Lust. lft. similar to leucine fr. cooling dil. alc. sol. Sbl. in woolly flocks at 230° giving odor like freshly crushed green beans. Dec. at 200° to d-amylamine & CO_2 . S. in 26 pt. aq. at 15.5° . I. c. abs. alc.; s. h. alc.; i. most organic solvents. $[\alpha]_D^{20} = \text{abt. } +10^\circ$ (in 3% aq. sol.). Taste of sol., sl. astringent, bitter & chalky. ② Evolves N in T. 2.4 w. HNO_3 , giving 10.69% in T. 2.5. — $\text{Cu}\bar{\text{A}}$, rosettes of blue lft., s. at 18° in 278 pt. aq. or 476 pt. alc. & containing 19.64% Cu. (Prepared by disg. Cu(OH) ₂ or CuCO_3 in h. aq. sol. of the acid.) — $\text{B}_2\text{C}_{10}\text{H}_8\text{O}_4\text{N}_4$ (picrolonate), cryst. s. in 176 pt. aq. at 20° . — Benzoyl deriv. (preparation fully described in T. 2.27-d), lust. ndl. V.d.s. c. aq.,

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. ‡ = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2630	280-1d. (r.h. in s.t.)	m.p. 116-7°, sintering at 114°; $[\alpha]_D^{20} = +26.36^{\circ}$ (fr. 1.4612 g. disd. in 9 cc. normal NaOH & diluted to weight 19.6544 g.). — The phenylureido acid of T. 2.31 cryst. fr. dil. alc. in lft., m.p. 119-20° w. frothing. It is i. c. aq.; e.s. h. aq.; v.s. alc. The corresponding phenylhydantoin (Ber., 37, 1829) forms long silky ndl., d.s. c. aq., m.p. 78-9°. — Et \ddot{A} , b.p. 90-2° (15 mm.).
2631	280-1	d'Alloisoleucine, Et.CHMe.CH(NH ₂).CO ₂ H. — [V. similar in most respects to No. 2.2629, of which it is a synthetic isomer.] Taste sweet. Separated fr. d-isoleucine by treating sol. w. sugar & fermenting. This destroys d-isoleucine, but not d'alloleucine. (Ber., 40, 2538-2562.) Taste sweet. Aq. sol. levorotatory. — Should give N in T. 2.4.
2632	280	m-Tyrosine, m-Hydroxyphenyl- α -aminopropionic Ac., HO.C ₆ H ₄ .CH ₂ .CH(NH ₂).CO ₂ H. — [Synthetic compound.] — Lst. s. 12 pt. c. or 22 pt. h. aq.; d.s. alc. — Should give N in T. 2.4.
2633	280-5	Dinitro-5-acetamino-1,2,4-trimethylbenzene, (NO ₂) ₂ (Me.CO.NH).C.Me ₂ . — (P) Nitro comp. — Sbl. D.s. alc. ("Sapd. by H ₂ SO ₄ , but not by KOH or HCl.")
2634	abt. 283-5u.c. d. (r.h. in s.t.) (= 293-5c.)	(α)p-Dinitrostilbene, NO ₂ .C ₆ H ₄ .CH:CH.C ₆ H ₄ .NO ₂ . — (P) Nitro comp. — Ndl. fr. nitrobenzene. I. aq. † l-Leucine, α -Aminoisobutylic Ac., Me ₂ CH.CH ₂ .CH(NH ₂).CO ₂ H. — [In plants, beet molasses, & proteolytic products.] — Thin lust. lft. of greasy feel moistened w. difficulty by aq. Sbl. in woolly flocks w. amylamine odor. S. in 41 pt. aq. at 22°; in 1385 pt. 99% alc. at 17°, or in 826 pt. h. alc. 100 pt. gl. ac. ac. dis. 10.87-10.95 pt. at 16°. $[\alpha]_D^{20} = +15.64^{\circ}$ in 20% HCl. Taste insipid, faintly bitter. $[k_A \cdot 10^{10}] = 2$. $k_B \cdot 10^{12} = 2.5$. (P) (1) Vapors give strong pyrrole react. when substance is ignited w. Zn dust in T. 2.24-(2). — (2) Evolves N in T. 2.4 w. nitrous ac., giving 10.69% in T. 2.5. Cu \ddot{A} , pale blue scales or mic. rhomb. tbl. s. in 3045 pt. c., or 1460 pt. boiling aq. — Benzoyl deriv. (prepared by T. 2.27-d), m.p. 105-7° c. — B.C ₆ H ₅ O ₂ N ₄ , cryst. s. in 181 pt. aq. at 20°, m.p. 145-50°. — Phenylureido-acid deriv. (T. 2.31), m.p. 200-10° d.; corresponding phenylhydantoin, ndl. fr. bz., m.p. 119-20°, $[\alpha]_D^{20} = -39.0^{\circ}$ (1.085 g. in 4 cc. normal KOH). — Et \ddot{A} , b.p. 83.5° (12 mm.); 196° (761 mm.).
2635	abt. 283-5u.c., d. (r.h. in s.t.)	d,l-Leucine, α -Aminoisobutylic Ac., Me ₂ CH.CH ₂ .CH(NH ₂).CO ₂ H. — [Product of proteolysis.] Cryst. S. in abt. 106 pt. aq. at 15°. Taste faintly sweet. — (P) Should give results as described above under (1) & (2) for l-deriv. — Benzoyl deriv. (prepared by T. 2.27-d), plates or pr., m.p. 135-9° u.c., s. in 200 pt. h. aq. (Ber., 23, 2373). — Phenylureido deriv. (Cf. T. 2.21) s. in 300 pt. boiling aq., m.p. 165° c. The phenylhydantoin corresponding (formed by dehydration w. boiling 25% H ₂ SO ₄) melts at 125°. — Et \ddot{A} , b.p. 83.5° (12 mm.), 196° (761 mm.), giving picrate, d.s. h. aq., m.p. 134° u.c.
2636	285d.	Succin- α -naphthalide, C ₁₀ H ₈ .NH.CO.CH ₂ .CH ₂ .CO.NH.C ₁₀ H ₇ . — (P) (D) Sapn. T.* products: α -naphthylamine (No. 2.589); succinic ac. (Vol. I). — Ndl. fr. gl. ac. ac. Alm. i. aq.; v.d.s. alc.; d.s. gl. ac. ac.
2637	abt. 288u.c., d. (r.h. in s.t.) (= 298c.)	d,l-Valine, α -Aminoisovaleric Ac., Me ₂ CH.CH(NH ₂).CO ₂ H. — [Synthetic, or by racemization of opt. act. acids.] — S. in 11.7 pt. aq. at 15°. Taste, sweet. — (P) Should evolve N in T. 2.4 w. nitrous ac., & give 11.96% N in T. 2.5. — Benzoyl deriv. (prepared by T. 2.27-d, Ber., 35, 403), ndl. fr. eth. + lgr., m.p. 132.5° c. — Phenylureido deriv. (T. 2.31), lft. s. in 130 pt. h. aq., m.p. 163.5° c. The corresponding phenylhydantoin melts at 124.5°. — Et \ddot{A} , b.p. 63.5° (8 mm.).
2638	288u.c. (= 298-9c.)	Paraxanthine, 1,7-Dimethylxanthine, C ₈ H ₁₀ O ₂ N ₄ . — [In urine.] — (P) Gives murexide react., T. 2.20, but not the "xanthine test (1)" described under No. 2.517. — Glassy, usually 6-sided tbl. D.s. c. aq.; s. 24 pt. h. aq.; i. alc., eth. ["Sol. reacts neut."] V. conc. sol. slowly solidifies to mass of silky interlacing ndl. — Picric ac. gives cryst. ppt. in HCl sol. decd. by aq. — In conc. sol. NaOH ppts. Na deriv. in lust. tbl. — B.HAuCl ₄ .H ₂ O, (T. 2.13), m.p. 227-8°.
2639	290-1	Phenylactimide, 3,6-Dibenzyl-2,5-diacipiperazine, (d,l-Phenylalanineanhydride) $^2\text{CO.NH.CH(CH}_2\text{.Ph).CO.NH.CH}^2$. — Silky ndl. fr. h. alc. After melting in t.t. sbl. in woolly ndl. I. h. aq.; mod. s. h. alc.; d.s. c. alc.; i. eth., ac., alk.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2640	abt. 292 u.c., d. (r.h. in s.t.) (=303c.)	d- α -Aminobutyric Ac., Et.CH(NH ₂).CO ₂ H. — Colorless lft. fr. dil. alc. $[\alpha]_D^{20} = +8.0^{\circ}$ (0.738 g. in 13.6546 g. aq. sol.). Properties similar to d,l-compound, No. 2.2642. — \oplus Should give N in T. 2.4.
—	abt. 290-2 c. (r.h. in s.t.)	d-Alanine. — Cf. No. 2.2617. M.p. abt. 297° d. (r.h. in s.t.) is also given.
2641	293d. (r.h. in s.t.?)	d,l-Alanine, rac.- α -Aminopropionic Ac., Me.CH(NH ₂).CO ₂ H. — Cryst. e.s. aq. 100 g. 80% alc. at 25° dis. 0.37 g. Closely resembles d-alanine, No. 2.2617. — \oplus (1) Gives strong pyrrole react. in T. 2.24-d when ignited w. Zn dust. — (2) Should give 15.73% N in T. 2.5. — Benzoyl deriv. (prepared by T. 2.27-d, Ber., 32, 2451), e.s. h. aq., m.p. 162-3° u.c. — Phenylureido deriv. T. 2.31), lust. lft. fr. h. aq., m.p. 168°; corresponding phenylhydantoin, ndl. fr. h. dil. alc., m.p. 172-3°.
—	abt. 293-5c.d. (r.h. in s.t.)	l-Leucine. — Cf. No. 2.2634.
2642	abt. 295d.w. efferv.; abt. 304u.c., d. (r.h.) (=316c.)	l-Tyrosine, p-Hydroxyphenyl- α -aminopropionic Ac., HO.C ₆ H ₄ .CH ₂ .CH(NH ₂).CO ₂ H. — [In germinating plants & proteolytic products. M.p. given are both Fischer's (Ber., 32, 3640). The first was obtained by slow heating, the rate being v. important. Much lower melting points (e.g. 272° u.c. in open tube, 265° in s.t., Z. physiol. Chem., 37, 18) have been recorded.] — Small silky ndl. often united in tufts. Odorless. Tasteless. Gives odor of burned horn on ignition. S. in 2491 pt. aq. at 17°, or 154 pt. at 100°. 100 pt. 95% alc. dis. 0.01 pt. at 17°. I. eth., abs. alc.; e.s. alk., alk. carbonates, ammon., dil. mineral ac.; d.s. ac. ac. $[\alpha]_D^{20} = -8.64^{\circ}$ (for 4% sol. in 21% HCl, or -13.2° (for 4.6% sol. in 4% HCl). — \oplus (1) Evolves N w. nitrous ac. in T. 2.4, giving 7.73% N in T. 2.5. — (2) Vapors on ignition w. Zn dust. (T. 2.24-(2)) give pyrrole splinter react. — (3) Gives Millon's react., T. 2.19! — (4) Warm 30 min. w. little conc. H ₂ SO ₄ on water-bath. Dil. cooled sol. Neutralize w. BaCO ₃ . Filter & concentrate. Sol. becomes violet when a little FeCl ₃ is added. (Piria's react., Ann., 82, 252.) — (5) Add a little tyrosine to 2-3 cc. of sol. of 1 cc. formaldehyde in 50 cc. conc. H ₂ SO ₄ . A wine-red color soon appears. Add 2 vol. gl. ac. ac. & boil. Sol. becomes green. — (6) A boiling aq. tyrosine sol. acidified w. ac. ac. becomes violet-red when 1% NaNO ₂ sol. is dropped in with continued heating. (Wurster's test, Zentr. Physiol., I, 193.) — (7) Cf. Z. physiol. Chem., 42, 517, for description of Pauly's test w. diazobenzenesulphonic ac., which is said to be especially specific. — Cu \ddot{A} , (by boiling aq. sol. w. Cu(OH) ₂ , blue pr. s. in 1230 pt. c., or in 240 pt. boiling aq.). — Dibenzoyl deriv. (T. 2.27), ndl. fr. gl. ac. ac., i. c. aq., e.s. alc., m.p. 211-2°. — B.C ₁₀ H ₈ O ₄ (picrolonate) (T. 2.23), pr. s. in 345 pt. aq., d. 260°. — Et \ddot{A} , flat pr., d.s. c. aq., e.s. alc., m.p. 108-9°.
2643	abt. 296u.c., d. (r.h. in s.t.) (=307c.)	d- α -Aminobutyric Ac., Et.CH(NH ₂).CO ₂ H. — Lft. s. in 3.5 pt. c. aq., or in 550 pt. boiling alc. Slowly heated in open capillary melts abt. 285° & subl. above 300°. Taste, sweet. — \oplus Should evolve N in T. 2.4 w. nitrous ac. — FeCl ₃ gives dark brown-red color w. aq. sol. — Aq. sol. w. Cu acetate gives violet-blue ppt., Cu \ddot{A} , v.d.s. h. aq. — Benzoyl deriv. (prepared by T. 2.27-d, Ber., 33, 2388), cryst. fr. 25 pt. h. aq., m.p. 143-4° u.c. — Phenylureido deriv. (T. 2.31; Ber., 33, 2395), ndl. fr. 50 pt. h. aq., m.p. 170° c., d.; corresponding phenylhydantoin, cryst. fr. h. dil. alc., m.p. 126-7° c., d.
2644	abt. 297d.	d,l-Tyrosine. — Opt. i., but otherwise closely resembling the l-deriv. No. 2.2642.
2645	300	Naphthalimide, $[C_{10}H_8.CO.NH.CO]^{\pm}$. — Ndl. fr. alc. D.s. h. alc.; v.d.s. eth., bz. E.s. warm dil. KOH. — E.s. warm dil. NaOH. Normal sapon. products (T. 2.26); NH ₃ & naphthalic ac. (C ₁₂ H ₈ O ₄ , Vol. I).
2646	abt. 303u.c., d. (in s.t.) (=315c.)	d-Valine, l-Aminoisovaleric Ac., Me.CH.CH(NH ₂).CO ₂ H. — [Proteolytic product & in plants.] — Silvery, usually 6-cornered, mic. lft. S. in 11 pt. c. aq. Taste, feebly sweet & bitter. $[\alpha]_D^{20} = +28.2^{\circ}$ (5% sol. in 20% HCl). — \oplus Evolves N w. nitrous ac. in T. 2.4, giving 11.96% in T. 2.5. — Cu \ddot{A} , blue lft. s. in 52 pt. methyl alc. at 18°. — B.C ₁₀ H ₈ O ₄ (picrolonate), cryst., s. in 83 pt. aq. at 20°, m.p. 170-80°. — Phenylureido deriv. (T. 2.31; Ber., 39, 2320), cryst. s. in 130 pt. h. aq., m.p. 154°; 147° c. M.p. of corresponding phenylhydantoin, cryst. fr. eth. + pet.-eth., 131-3° c.; 124°.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.17.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2647	306; 314	Diacetyl-(o)3,4-tolidine, 4,4'-Diacetamino-3,3'-dimethylbiphenyl, (Me.CO.NH)Me.C ₆ H ₄ .C ₆ H ₄ .Me(NH.CO.Me). — Lust. ndl. Alm. i. alc., eth., bz.; s. h. gl. ac. ac. — Normal sapn. products (T. 2.26): o-tolidine (No. 2.849); ac. ac.
2647-I	313-4d.	Fumaranilide, Ph.NH.CO.CH:CH.CO.NH.Ph. — (P) (D) Sapn. T.* products: aniline; fumaric ac. (Vol. I). — Mic. ndl. fr. gl. ac. ac.
2640	abt. 317-20; 330-1c.	Diacetylbenzidine, (Me.CO.NH).C ₆ H ₄ .C ₆ H ₄ .(NH.CO.Me). — V.d.s. common solvents. Normal sapn. products (T. 2.26): benzidine (No. 2.840); ac. ac.
—	abt. 314-8c., d. (r.h. in s.t.)	L-Tyrosine. — Cf. No. 2.2643.
2643	abt. 321d. (r.h.), w. efferv.; 325-335	Thymine, 5-Methyluracil, $^{\text{f}}\text{NH.CO.NH.CH:CM}\text{e.CO}^{\text{f}}$. — [Fr. hydrolysis of nucleic ac.] — Stellate or dendritic clusters of small lft., or rarely, short ndl. fr. aq. Luster fatty. Sbl. on cautious heating in lft. 100 pt. aq. at 25° dis. 0.404 pt. E.s. h. aq.; d.s. alc.; v.d.s. eth. [“Aq. sol. reacts neut.”] Taste, bitter. Decolorizes Br. aq. Sol. in NaOH is colored red by diazobenzenesulphonic ac. AgNO ₃ gives no ppt. alone in thymine sol., but voluminous ppt. appears, s. in x.s. of precipitant, on cautious addition of ammonia.
2650	330	Phenyl- β -naphthylcarbazole, C ₁₀ H ₁₁ N. — [In crude anthracene.] Lft. B.p. 440-50°. S. in 400 pt. boiling alc.; i. aq., c. alc., c. toluene; s. in 200 pt. boiling toluene. — (P) Sol. shows blue fluor. — Nitroso deriv. (Ann., 201, 1), m.p. 240°.
2651	337u.c.	† Theobromine, 3,7-Dimethyl-2,6-dioxypyurine, C ₇ H ₈ O ₂ N ₄ . — [In the cocoa bean.] Separates fr. h. sat. aq. sol. in minute micro-cryst. of rhombic syst. Sbl. abt. 290°. [k _A .10 ⁴ = 0.01; k _B .10 ⁴ = 1.5.] S. in 3282 pt. aq. at 18°, or 148.5 pt. boiling; in 4284 pt. abs. alc. at 17°; s. in 105 pt. h. chlf., v.d.s. c.; 100 pt. c. bz. dis. 0.0015 g.; alm. i. c. eth.; s. dil. ac. or alk. Odorless. Taste of sat. aq. sol., bitter. (P) (1) Gives the murexide react. (T. 2.20). — (2) Dis. 0.05 g. in 2 cc. aq. + 1 cc. conc. HCl. Add 5 cc. sat. Br-aq. Boil to expel x.s. Br. Dilute w. 5 cc. aq. Add 1 drop FeSO ₄ sol., & then ammon., drop by drop, shaking after each addition until the blue color formed is permanent. The intense color (BS1) fades on long standing & is destroyed by x.s. of ammon. [François (Chem. Zentr., 1898, II, 66) states that this reaction is also given by caffeine and describes other tests for theobromine.] (D) Ppt. the gran. cryst. Ag. deriv., C ₇ H ₈ O ₂ N ₄ .Ag.1½H ₂ O, by adding N/10 AgNO ₃ sol. in x.s. to boiling ammon. aq. sol. of substance, & filter. Determine percentage of Ag in deriv. It loses its aq. at 120-30°.
2651-I	338c.	s-Tetra-p-nitrotetraphenylethane, (NO ₂ .C ₆ H ₄) ₂ .CH.CH.(C ₆ H ₄ .NO ₂) ₂ . — Ndl. fr. nitrobenzene. I. alc. Stable toward conc. H ₂ SO ₄ & fuming HNO ₃ .
2652	338 (r.h.) w. efferv.	Uracil, 2,6-Dioxypyrimidine, $^{\text{f}}\text{NH.CO.NH.CH:CH.CO}^{\text{f}}$. — [Proteolytic product.] Cryst. powd. (Ndl. in rosettes.) D.s. c. aq.; e.s. h. aq.; alm. i. alc., eth.; e.s. ammon. Not decd. by boiling NaOH. — Ammon. sol. gives amorph. ppt. of Ag salt w. AgNO ₃ . — Is ptd. by HgN ₂ O, but not by picric ac. or phosphotungstic ac.
2653	344	† 2,4-Diketotetrahydroquinazoline, Uramidobenzoyl, $^{\text{f}}\text{C}_6\text{H}_4.\text{CO.NH.CO.NH}^{\text{f}}$. — Ndl. s. in 300 pt. boiling aq. V.d.s. alc., eth.; s. NaOH & ptd. by CO ₂ . Sbl. Gives white ppt. w. AgNO ₃ sol.
2654	352	Dibenzoylbenzidine, Ph.CO.NH.C ₆ H ₄ .C ₆ H ₄ .NH.CO.Ph. — Pr. fr. boiling phenol. I. usual solvents. Sbl.
—	360-5(r.h.) d.	Adenine. — Cf. No. 2.1057.
2655	abt. 380(r.h.), d. w. efferv.	Heteroxanthine, 7-Methylxanthine, 7-Methyl-2,6-dioxypyurine, C ₆ H ₈ O ₂ N ₄ . — [In urine.] — White powd., or rosettes of ndl. Sinters & colors above 360° before melting. S. w. “neut. react.” in 142 pt. boiling aq.; d.s. c. aq.; i. alc., eth.; e.s. ammon. or h. HCl. — (P) Gives the murexide react. (T. 2.20), but not the “xanthin tests” (Cf. No. 2.517-(1),(2)). — Picric ac. gives no ppt. — Is ptd. by copper acetate or phosphotungstic ac.

No.	Melting-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Colorless and Solid.
2656	384d.	9-Methyl-2,6-oxypurine , $C_6H_5O_2N_4$. — \textcircled{P} Gives the murexide react. (T. 2.20). — Ndl. s. in 280 pt. h. aq.
2657	d.w.m. a. 360	3-Methylxanthine , 3-Methyl-2,6-dioxypurine , $C_6H_5O_2N_4$. — \textcircled{P} Gives the murexide react. T. 2.20. — Fine lust. ndl. or pr. fr. h. aq. S. in 350 pt. boiling aq.; less s. alc., chlf.; e.s. dil. alk.; mod. s. ammon., separating when NH ₃ is boiled off. — Ba salt d.s. h. aq., cryst. on cooling in fine ndl.
2658	d.w.m. a. 390	Epiguanine , 7-Methylguanine , 7-Methyl-2-amino-6-oxypurine , $C_6H_5ON_5$. — [In urine.] — \textcircled{P} Gives the murexide react. T. 2.20 & the "xanthine test" (1) & (2) (Cf. No. 2.517). — Fine ndl. fr. aq. S. in 900 pt. boiling aq.; alm. i. c. aq., alc., ammon.; e.s. dil. NaOH, HCl or H ₂ SO ₄ ; d.s. dil. HNO ₃ . — \textcircled{D} B.Pk. (T. 2.23), s. in 2740 pt. aq. at 18°, d. w. efferv. at 257°!
2659	unchanged at 360 <u>c.</u>	† Guanine , 2-Amino-6-oxypurine , $C_6H_5ON_4$. — [Fr. animal secretions & tissues.] — \textcircled{P} Gives the murexide react. & "xanthine tests" (1) & (2). (Cf. T. 2.20 & No. 2.517.) In test last named the color w. NaOH is at first O, then deep red, brown-red, & after protracted heating on water-bath, lightening to ORT2. — <i>Usually amorph. powd. Odorless. I. aq., alc., eth.; d.s. ammon.; e.s. alk. or dil. min. acids. Conc. K₂CrO₄ or K₄Fe(CN)₆ sol. added to dil. sol. of guanine in acidulated aq. gives ppt. which does not appear w. xanthine or hypoxanthine (Z. physiol. Chem., 4, 235).</i> † <i>Dis. 0.05 g. in boiling mixture of 1 cc. conc. HCl + 9 cc. aq. Add 4 cc. sat. aq. picric ac. sol. Allow to cool v. slowly. The picrate, B.Pk.H₂O separates in clusters of long slender OY ndl. Wash w. 2 cc. c. dil. HCl (1:10) sol. Dry 15 min. on porous tile at 100°. Picrate darkens at 170° & dec. abt. 190° u.c. — B.2H₂SO₄.2H₂O ndl. fr. h. dil. H₂SO₄. (Fischer recommends determination of H₂O, lost at 120°, in characterization, & to distinguish fr. the very similar 6-amino-2-oxypurine). — Unlike adenine, xanthine, & hypoxanthine, is completely pptd. fr. sol. by metaphosphoric ac.</i>
2660	417-9d. (in s.t.)	† Oxamide , $NH_2.CO.CO.NH_2$. — Cryst., tasteless, odorless powd. I. aq., alc., eth. — \textcircled{P} \textcircled{D} Sapn. T.* (w. NaOH) products: NH ₃ ; oxalic ac.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.15.

SUBORDER I OF ORDER II.
COLORLESS (OR YELLOWISH) COMPOUNDS CONTAINING C, N, H, AND O.

GENUS III, NEUTRAL COMPOUNDS.

DIVISION B, LIQUID SPECIES.

No.	Boiling-point (C°).	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
2661	-20.7	Cyanogen, $(CN)_2$. — M.p. -34.4°. Odor sl. pungent (poisonous). — S. in $\frac{1}{2}$ vol. c. aq., more s. alc. — (1) Burns w. charac. peach blossom colored flame! — (2) Absorbed by KOH sol. giving KCN & KCNO! (Cf. No. 2.519). — C. conc. HCl saturated w. the gas gives ppt. of solid oxamide (No. 2.2660) after 12 hr.
2662	-12	Methyl Nitrite, $MeNO_2$. — Sp. gr. 0.991 (15°). — (2) Nitrite. § — (1) Sapn. T.* (w. NaOH) products: methyl alc. (Vol. I); nitrous ac.
2663	+17	Ethyl Nitrite, $Et.NO_2$. — Sp. gr. 0.900 (15.5°). D.s. aq. Odor, fruity. — (2) Nitrite. § — Sapn.* gives ethyl alc. (Vol. I).
2664	abt. 39-9.5; 45	Isopropyl Nitrite, $Me_2CH.NO_2$. — Sp. gr. 0.856 (0°); 0.844 (25°). — (2) Nitrite. § — (1) Sapn. T.* (w. NaOH) products: isopropyl alc. (Vol. I); nitrous ac.
2665	43-5	Methyl Isocyanate, Methylcarbonimide, $Me.N:C:O$. — (2) Odor, powerful & disagreeable! — (1) Sapn. T.* (w. NaOH) products: methylamine (No. 2.1059); CO_2 .
2666	44	Allyl Nitrite, $C_3H_5NO_2$. — Sp. gr. 0.955 (0°). — (2) Nitrite. § — (1) Sapn. T.* products: allyl alc. (Vol. I); nitrous ac. — V. unstable. Vapors explode at 100°. I. aq. "Decd. by shaking w. aq."
2667	57; 43-6	Propyl Nitrite, $Pr.NO_2$. — Sp. gr. 0.998 (0°). — (2) Nitrite. § — (1) Sapn. T.* products: propyl alc. (Vol. I); nitrous ac.
2668	59.6	Methyl Isocyanide, Methylcarbylamine, $MeNC$. — Sp. gr. 0.7557 (4°). S. in 10 pt. aq. at 15°. — (2) Odor, powerful, charac., nauseating!
2669	60	Ethyl Isocyanate, Ethylcarbonimide, $Et.N:C:O$. — Sp. gr. 0.898. — (2) Odor, powerful & disagreeable! — (1) Sapn. T.* (w. NaOH) products: ethylamine (No. 2.1062); CO_2 .
2670	66	Methyl Nitrate, $MeNO_3$. — Sp. gr. 1.182 (22°)! Odor, ethereal. I. aq. Percussion or heat may cause violent explosion. — (2) Nitrite. § — (1) Sapn. T.* (w. NaOH) products: methyl alc. & nitric ac.
2671	67	Isopropyl Isocyanate, Isopropylcarbonimide, $Me_2CH.N:C:O$. — (2) Odor, strong & unpleasant! — (1) Sapn. T.* (w. NaOH) products: isopropylamine (No. 2.1063); CO_2 .
2672	67	Isobutyl Nitrite, $Me_2CH_2.NO_2$. — Sp. gr. 0.908 (0°). — (2) Nitrite. § — (1) Sapn. T.* (w. NaOH) products: isobutyl alc. (Vol. I); nitrous ac.
2673	68	sec.-Butyl Nitrite, $Et.CHMe.NO_2$. — Sp. gr. 0.898 (0°). — (2) Nitrite. § — (1) Sapn. T.* products: sec.-butyl alc. ($C_4H_{10}O$, Vol. I); nitrous ac.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.15.

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Liquids.
2674	75	Butyl Nitrite, Et.CH ₂ .CH ₂ .NO ₂ . — Sp. gr. 0.911 (0 $^{\circ}$). — \textcircled{P} Nitrite. \textcircled{S} — \textcircled{D} Sapon. T.* products: n-butyl alc. (T. 1.813); nitrous ac.
2675	78-9	† Ethyl Isocyanide, Ethylcarbylamine, EtNC. — Sp. gr. 0.7591 (4 $^{\circ}$). — \textcircled{P} Odor, powerful, charac., nauseating! — E.s. aq. Conc. HCl reacts w. hissing sound, destroying odor & giving ethylamine & formic ac. Reduces Tollen's reagent.
2676	81.5	† Acetonitrile, Methyl Cyanide, Me.CN. — Sp. gr. 0.7906 (14.1 $^{\circ}$ /4 $^{\circ}$). M.p. -41 $^{\circ}$. Misc. aq. Odor ethereal. Burns w. pink edged flame. — \textcircled{P} Gives NH ₃ in T. 2.7. — \textcircled{D} Sapon. 0.3 g. w. alc. KOH to NH ₃ ; & ac. ac.
2677	82	Allyl Isocyanate, Allylcarbonimide, C ₃ H ₅ N:C:O. — \textcircled{P} \textcircled{D} Sapon. T.* (w. NaOH) products: allylamine (No. 2.1068); CO ₂ .
2678	84	Formaldoxime, CH ₂ :NOH. — \textcircled{P} Reduces AgNO ₃ , HgCl ₂ , etc., directly. — Conc. HCl splits to hydroxylamine & formaldehyde. Metallic deriv. are explosive.
2679	85.5c.	tert.-Butyl Isocyanate, tert.-Butylcarbonimide, Me ₃ C.N:CO. — Sp. gr. 0.868 (0 $^{\circ}$). Odor, aromatic, then pungent. — \textcircled{P} \textcircled{D} Sapon. T.* (w. NaOH) products: tert.-butylamine (No. 2.1066); CO ₂ .
2680	87.6	† Ethyl Nitrate, EtNO ₂ . — Sp. gr. 1.116 (15 $^{\circ}$). — Odor v. similar to chlf. Taste of sol. v. sweet, & sl. burning. — \textcircled{P} Nitrate. \textcircled{S} — \textcircled{D} Sapon. T.* (w. NaOH) gives ethyl alc. & nitric ac.
2681	87	Isopropyl Isocyanide, Isopropylcarbylamine, Me ₂ CH.NC. — Sp. gr. 0.7596 (0 $^{\circ}$). — \textcircled{P} Odor, intolerable, though at first rather ethereal! — I. aq.; s. alc., eth. Reacts vigorously w. conc. HCl giving isopropylamine & formic ac.
2682	91	tert.-Butyl Isocyanide, Me ₃ C.NC. — \textcircled{P} Odor exceedingly disagreeable! — E. decd. by conc. HCl to tert.-butylamine (No. 2.1066) & formic ac. (Vol. I).
2683	93	Acetyl Cyanide, Me.CO.CN. — [Generic position doubtful.] — Odor like ac. ac. & HCN, which are formed by moisture. Contact w. solid KOH, or on long keeping, gives solid polymer. Careful treatment w. HCl gives pyruvic ac. (Vol. I).
2684	94-5	† Isoamyl Nitrite, C ₆ H ₁₁ NO ₂ . — Sp. gr. 0.880 (15 $^{\circ}$) GYT ² liquid of charac. odor. D.s. to v.d.s. aq. — \textcircled{P} Nitrite. \textcircled{S} — \textcircled{D} Sapon. T.* (w. NaOH) products: isoamyl alc. (Vol. I); nitrous ac.
2685	96-8	Ethylene Nitrite, C ₂ H ₄ (NO ₂) ₂ . — Sp. gr. 1.216 (0 $^{\circ}$). — \textcircled{P} Nitrite. \textcircled{S} — Unstable. I. aq. E. sapd. Forms ethyl nitrite in alc. sol.
2686	97.1c.	† Propionitrile, Ethyl Cyanide, EtCN. — Sp. gr. 0.8010 (0 $^{\circ}$). Odor ethereal. Mod. s. aq.; salted out by CaCl ₂ . — Sapon. T.* (w. NaOH) products: NH ₃ ; propionic ac. (Vol. I).
2687	100-1	Methyl Cyanoformate, CN.CO.Me. — Lighter than aq. Odor, ethereal, irritating. I. aq. & decd. by it, especially on heating. — \textcircled{P} \textcircled{D} Sapon. T.* products: HCN, CO ₂ , MeOH.
2688	101	† Nitromethane, MeNO ₂ . — Sp. gr. 1.144 (15 $^{\circ}$). — Mobile liq. of sweetish ethereal odor. Sol. of 1 drop in 2 cc. aq. tastes sweet like chlf. E.s. aq. — \textcircled{P} Gives T. 2.15 & 2.17-b. — 3 drops mixed w. 3 cc. 10% NaOH sol. & a pinch Zn dust give on boiling vapors of ammon. odor which blue litmus. — Dis. bit of Na of size of grain of wheat in 1 cc. alc. in small t.t. Cool. Add 3 drops nitro comp. Stir. Transfer pearly ppt. to porous tile & wash w. 0.5 cc. alc. Place moist salt in t.t. & dis. in 1 cc. aq. Add 1 drop c. sat. NaNO ₂ . Acidify (using litmus) w. dil. H ₂ SO ₄ . Make strongly alk. w. NaOH. The alk. sol. shows an O-YO color (sodium methylnitrolate) which is discharged by ac. & restored by alk.
2685	101-2	Isopropyl Nitrate, Me ₂ CH.NO ₂ . — Sp. gr. 1.036 (19 $^{\circ}$). — \textcircled{P} Nitrate. \textcircled{S} — \textcircled{D} Sapon. T.* (w. NaOH) products: isopropyl alc. (T. 1.818); nitric ac.
2680	96-106	Allyl Isocyanide, C ₃ H ₅ N.C. — Sp. gr. 0.794 (17 $^{\circ}$). — \textcircled{P} Odor, penetrating, persistent, & highly disagreeable! — D.s. aq.; misc. eth., ac. ac.
2691	105-6	Trimethylacetonitrile, Me ₃ C.CN. — Cryst. mass of m.p. 15-16 $^{\circ}$. — \textcircled{P} \textcircled{D} Sapon. T.* products: NH ₃ ; trimethylacetic ac. (C ₆ H ₁₀ O ₂ , Vol. I).

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.15.

No.	Boiling-point (C°).	NEUTRAL COMPOUNDS.—Liquids.
2692	106	Allyl Nitrate, $C_3H_5NO_2$. — Sp. gr. 1.09 (10°). — \oplus Nitrate. § — \ominus Sapon. T.* (w. NaOH) products: allyl alc. (Vol. I); nitric ac.
2693	107-8	Isopropyl Cyanide, $Me_2CH.CN$. — \oplus \ominus Sapon. T.* products: NH_3 ; isobutyric ac. (Vol. I).
2684	108-10	Propionyl Cyanide, $Et.CO.CN$. — [Unstable. Generic position doubtful.]
2695	110	Isobutyl Isocyanate, Isobutylcarbonimide, $Me_2CH.CH_2.N:CO$. — Odor disagreeable! — \oplus \ominus Sapon. T.* (w. NaOH) products: isobutylamine (No. 2.1078); CO_2 .
2696	110.5	Propyl Nitrate, $Pr.NO_2$. — Sp. gr. 1.063 (15°). — \oplus Nitrate. § — \ominus Sapon. T.* (w. NaOH) products: propyl alc. (Vol. I); nitric ac.
2697	112-3; 114-5 (th.i. 747.5 mm.)	1-Methylpyrrole, $C_4H_5N.Me$. — Sp. gr. 0.9203 (10°). — Odor somewhat like pyrrole.
2698	114.5	† Nitroethane, $Et.NO_2$. — Sp. gr. 1.101. — Mobile liq. of sweet, ethereal, chlf.-like odor. Taste of aq. sol. sweet & chlf.-like. E.s. to s. aq. — \oplus Gives results described in prelim. tests for No. 2.2688. — [Isomerization by alk. is a little too slow in generic titration to give species position w. acids.]
2699	115-6	Ethyl Cyaniformate, $CN.CO.Et$. — Sp. gr. 1.101. — \oplus Odor ethereal, pungent, irritating. — I. aq., by which it is slowly decd. to CO_2 , HCN & EtOH, the decn. being more rapid in presence of NaOH. — Treatment w. ammon. gives urethane (No. 2.1540).
2700	114-7	Isobutyl Isocyanide, $Me_2CH.CH_2.NC$. — Sp. gr. 0.7873 (4°). — \oplus Insufferable carbylamine odor! — Alm. i. aq.; s. alc., eth.
2701	118	2-Nitropropane, $Me.CH(NO_2).Me$. — Sp. gr. 1.024 (0°). — \oplus Gives T. · 2.21, T. 2.17-b. — Dropped into conc. aq. NaOH sol., solid Na deriv. ppts. — Convert into charac. propylpseudonitrole (Cf. Ann., 175, 120).
2702	118	† Butyronitrile, Propyl Cyanide, $Pr.CN$. — Sp. gr. 0.795 (12.5°). — Sapon. T.* products: NH_3 ; butyric ac. (Vol. I).
2703	119c.	Crotononitrile ("Allyl Cyanide"), $Me.CH:CH.CN$. — Sp. gr. 0.835 (15°). Odor alliaceous. — \oplus \ominus Sapon. T.* (w. NaOH) products: NH_3 ; α -crotonic ac. (Vol. I).
2704	120	α -Hydroxyisobutryonitrile (Acetonecyanhydrine), $Me_2C(OH).CN$. — Completely decd. by $AgNO_3$ sol. to AgCN & acetone (Vol. I).
2705	120-5	Cyanoacetone, $Me.CO.CH_2.CN$.
2706	123	Isobutyl Nitrate, $Me_2CH.CH_2.NO_2$. — Sp. gr. 1.021 (15°). — \oplus Nitrate. § — \ominus Sapon. T.* (w. NaOH) products: isobutyl alc. (Vol. I); nitric ac.
2707	124	sec.-Butyl Nitrate, $Et_2Me.CH.NO_2$. — Sp. gr. 1.038 (0°). — \oplus Nitrate. § — \ominus Sapon. T.* (w. NaOH) products: sec.-butyl alc. (Vol. I); nitric ac.
2708	125	Methylethylacetonitrile, sec.-Butyl Cyanide, $Me.CHEt.CN$. — Sp. gr. 0.806 (0°). — \oplus \ominus Sapon. T.* products: NH_3 ; methylethylacetic ac. ($C_6H_{10}O_2$, Vol. I).
2709	126-7	1-Methylpyrazole, $[NMe.N:CH.CH:CH^2]$. — Oil of pyridine-like odor. [$k_B \cdot 10^{12} = 1.1$. "Aq. sol. reacts neut."] — $B_2H_2PtCl_6$ (T. 2.14), or.-yel. pr., m.p. 196-8° d.
—	127	Tert.-Nitrobutane. — Cf. No. 2.1467-1. (M.p. 24°.)
2710	129	Isovaleronitrile, Isobutyl Cyanide, $Me_2CH.CH_2.CN$. — Sp. gr. 0.807 (20°). — \oplus \ominus Sapon. T.* products: NH_3 ; isovalerianic ac. (Vol. I).
2711	128-30	Dimethylethylacetonitrile, $Me_2CEt.CN$. — \oplus \ominus Sapon. T.* products: NH_3 ; dimethylethylacetic ac. ($C_6H_{12}O_2$).
2712	130c.	† Pyrrole, $[NH.CH:CH.CH:CH^2]$. — Sp. gr. 0.9669 (21/4). — Odor sl. empyreumatic & chlf.-like. Soon turns yel. in air, & after a few days, brown. I. aq., dil. alk.; e.s. alc., eth.; dis. slowly in dil. ac. — \oplus (1) A splinter of soft pine which has been soaked in HCl (sp. gr. 1.12) becomes carmine red (R) at once when held in vapors rising from mixture of 1 drop pyrrole & 1 cc. aq. boiling in a t.t.! — (2) 1 drop pyrrole mixed w. 1 drop HCl (sp. gr. 1.20)

No.	Boiling-point (C.).	NEUTRAL COMPOUNDS. — Liquids.
		<i>reacts vigorously giving a red-brown resin, becoming hard on cooling. — (3) Mix 1 drop sat. aq. isatine sol. w. 1 drop pyrrole & 1 drop dil. H₂SO₄. An indigo-blue ppt., v.s. gl. ac. ac. w. dark blue color, d.s. alc., eth., appears!</i>
2713	131	1-Ethylpyrrole, "NET.CH:CH.CH:CH". — Sp. gr. 0.8881 (16°). — I. aq.; misc. alc., eth. — (P) Vapors color pine splinter moistened w. HCl bright red as described for pyrrole above. — Unlike pyrrole does not react w. K.
2714	131c.	† Nitropropane, Et.CH ₂ .NO ₂ . — Sp. gr. 1.011 (15°). — Odor sweet-ethereal like chlf. Taste of sat. aq. sol. sweet-pungent like chlf. sol. D.s. aq. — [Under the titration conditions of Gen. T. 2.I, abt. 0.7 cc. N/10 alk. is neutralized.] — (P) Gives the three "preliminary tests" described for nitromethane, No. 2.2688.
2715	131-5c.	Propionaldoxime, Et.CH:NOH. — Sp. gr. 0.9258 (20/4). — (P) Oxime (T. 2.17). — M.p. +21.5°; abt. 40°. n _D = 1.4287.
2716	134-5	Isoamyl Isocyanate, C ₅ H ₁₁ .N:C:O. — Lighter than aq. — (P) Odor v. unpleasant. — (P) Sapn. T.* (w. NaOH) products: isoamylamine (No. 2.1100); CO ₂ .
2717	135	Cyclopropanecarbononitrile, "CH(CN).CH ₂ .CH ₂ ". — Sp. gr. 0.911 (16°). — Sapn. T.* (w. NaOH) products: NH ₃ ; cyclopropanecarbonic ac. (C ₄ H ₆ O ₂ , Vol. I).
2718	abt. 135d.	α-Hydroxyisovaleronitrile, Me ₂ CH.CH(OH).CN. — Sp. gr. 0.956 (0°). — S. in 5 vol. aq.; v.s. alc., eth. — (P) Dis. easily in fuming HCl giving corresponding amide (No. 2.1897).
2719	136	Butyl Nitrate, C ₅ H ₁₁ .NO ₂ . — Sp. gr. 1.048 (0°). — (P) Nitrate. § — (P) Sapn. T.* (w. NaOH) products: butyl alc. (Vol. I); nitric ac.
2720	137	Isoamyl Isocyanide, C ₅ H ₁₁ .NC. — Lighter than aq. — (P) Odor, powerful, unpleasant, aromatic, suggesting HCN. — I. aq., alc., eth. Reacts explosively w. conc. HCl giving isoamylamine & formic ac.
2721	139	1-Nitro-2-methylpropane, Me ₂ CH.CH ₂ .NO ₂ . — Sp. gr. 0.987 (7.5°). — (P) Should give T. 2.21 & T. 2.17-b. — Gives no ppt. w. alc. NaOH.
2722	139	Isobutyraldoxime, Me ₂ CH.CH:NOH. — Sp. gr. 0.8943 (20/4). — (P) Oxime (T. 2.17). — Mod. s. aq.
2723	140	Allylacetonitrile, C ₅ H ₁₁ .CH ₂ .CN. — Sp. gr. 1.180 (13°). — Odor agreeable. I. aq. — (P) (P) Sapn. T.* products: NH ₃ ; allylactic ac. (C ₅ H ₈ O ₂ , Vol. I). [The isomeric nitriles of β-ethylacrylic ac. & ββ-dimethylacrylic ac. are similar in b.p., but have sp. gr. abt. 0.82.]
2724	141c.	Valeronitrile, Butyl Cyanide, Me ₂ (CH ₂) ₃ .CN. — Sp. gr. 0.816 (0°). — (P) (P) Sapn. T.* products: NH ₃ ; valerenic ac. (Vol. I).
2725	143	3-Methylpyrrole, "CMe:CH.NH.CH:CH". — Unstable in air. Less easily resinated by HCl than pyrrole (No. 2.2712).
2726	144-6	Diethylacetonitrile, Et ₂ CH.CN. — (P) (P) Sapn. T.* products: NH ₃ ; diethylacetic ac. (C ₄ H ₈ O ₂ , Vol. I).
2727	147	Isoamyl Nitrate, C ₅ H ₁₁ .NO ₂ . — Heavier than aq. — (P) Nitrate. § — (P) Sapn. T.* products: isoamyl alc. (Vol. I); nitric ac.
2728	145-50	Isovaleryl Cyanide, Me ₂ CH.CH ₂ .CO.CN. — [Generic position doubtful.] Decd. by Zn + HCl to HCN (No. 2.519) & isovalerenic ac. (Vol. I).
2729	148-9	Tetramethylene carbononitrile, "CH(CN).(CH ₂) ₃ ". — (P) (P) Sapn. T.* products: NH ₃ ; tetramethylenecarbonic ac. (C ₄ H ₈ O ₂ , Vol. I).
2730	abt. 150d.	Methaneazobenzene, Me.N:N.Ph. — Yel. oil. V. vol. w. st.
2731	151-2c.	1-Nitrobutane, Et.CH ₂ .CH ₂ (NO ₂). — Taste of sat. aq. sol. sweetish-pungent. Alm. i. aq. — (P) Should give T. 2.21, T. 2.17-b, & nitrolie ac. color reaction described for No. 2.2688.
2732	152.5(th.i.); 148	† Nitrosodimethylamine, Me ₂ N.NO. — (P) Should give nitrosomine T. 2.15. Yellowish oil. Odor peculiar & sl. mint-like on deep inhalation. V.s. aq. Taste of sol. burning & sweetish. Boiling HCl dec. to dimethylamine & nitrous ac.

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.15.

(ORDER II, SUBORDER I.)

No.	Boiling-point (C.).	NEUTRAL COMPOUNDS.—Liquids.
2733	152-3	† Methylmethylethylketoxime, $\text{MeC}(\text{:NOH})\text{Et}$. — Sp. gr. 0.919 (24°). — (P) Oxime giving T. 2.17-a, b. — S. in 10 vol. aq.; misc. alc., eth. Odor, aromatic.
2734	153-4	Butyraldoxime, Pr.CH:NOH . — (P) Oxime (T. 2.17). — Remains liq. at —80°.
2735	155	n-Heptyl Nitrite, $\text{C}_7\text{H}_{15}\text{NO}_2$. — Sp. gr. 0.894 (0°). — (P) Nitrite. § — (P) Sapn. T.* (w. NaOH) products: heptyl alc. ($\text{C}_7\text{H}_{15}\text{O}$, Vol. I); nitrous ac.
2736	155	Dimethylformamide, HCO.NMe_2 . — Sp. gr. 0.968 (20°). — (P) (P) Sapn. T.* products: dimethylamine (No. 2.1060); formic ac. (Vol. I).
2737	155.5	† Capronitrile, Isoamyl Cyanide, $\text{C}_6\text{H}_{13}\text{CN}$. — Sp. gr. 0.806 (20°). — (P) (P) Sapn. T.* products: isocaproic ac. ($\text{C}_6\text{H}_{13}\text{O}_2$, Vol. I).
2738	157-8	Methylisopropylketoxime, $\text{Me.C}(\text{:NOH}).\text{CHMe}_2$. — Lighter than aq. — (P) Oxime (T. 2.17).
2739	158	Methyl Methylcarbamate, MeNH.CO.Me . — Sp. gr. 1.065 (15°). — (P) (P) Sapn. T.* (w. NaOH) products: methylamine (No. 2.1059); methyl alc. (Vol. I); CO_2 .
2740	abt. 163	Isovaleraldoxime, $\text{Me}_2\text{CH.CH}_2.\text{CH:NOH}$. — Sp. gr. 0.8934 (20/4). — (P) Oxime (T. 2.17). — Solidified by freezing mixt.; m.p. 48.5°. Cryst. liquefy after a few days.
2741	163.5	Dimethylcyanamide, $\text{Me}_2\text{N.CN}$.
2742	164	† 4-Nitro-2-methylbutane, $\text{Me.CHMe.CH}_2.\text{CH}_2.\text{NO}_2$. — Sp. gr. 0.960 (20/4). — (P) Should give T. 2.21 & T. 2.17-b. — Gives nitric ac. color react. described for No. 2.2688. — Odor, peculiar, sweetish. Alm. i. aq. Taste of sat. aq. sol. v. pungent & sweetish.
2743	163-5	3-Ethylpyrrole, $^{\text{C}}\text{Et:CH.NH.CH:CH}^{\text{C}}$. — Turns brown in air.
2744	165	2,5-Dimethylpyrrole, $^{\text{C}}\text{Me:CH.CH}_2:\text{CMe.NH}^{\text{C}}$. — Sp. gr. 0.9353 (20/4). — (P) Vapors color pine splinter bright red in T. 2.24. — Unpleasant pungent odor; not chloroform-like. I. aq., alk.; v.s. alc., eth. Not easily attacked by acids.
2745	165	2,3-Dimethylpyrrole, $^{\text{C}}\text{Me:CM:CH:CH.NH}^{\text{C}}$.
2746	165-6	Phenyl Isocyanide, Phenylcarbylamine, PhNC . — Sp. gr. 0.9775 (15°). — (P) Odor, intolerable, nauseating, persistent! — Vol. w. st. Dist. w. partial polymerization leaving blue residue. Fresh distillate colorless, but becomes deep blue within an hour!
2747	165	Diethylketoxime, $\text{Et}_2\text{C:NOH}$. — Sp. gr. 0.914 (20/4). — (P) Oxime (T. 2.17).
2748	165	Methyl Ethylcarbamate, EtNH.CO.Me . — Sp. gr. 1.019 (15°). — (P) (P) Sapn. T.* products: ethylamine (No. 2.1062); methyl alc.; CO_2 .
2749	165.7	Dimethylacetamide, Me.CO.NMe_2 . — Sp. gr. 0.940 (20°). — (P) (P) Sapn. T.* products: dimethylamine (No. 2.1061); ac. ac.
2750	166	† Phenyl Isocyanate, Phenylcarbonimide, Carbanil, Ph.N:CO . — Sp. gr. 1.092 (15°). — (P) Vapors of extraordinarily irritating odor, provoking flow of tears! — (P) Treatment w. h. aq. changes to solid carbanilide, No. 2.2580, w. evolution of CO_2 . Recryst. product fr. h. dil. alc. — [May neutralize enough NaOH in Gen. T. 2.1 titration to be taken for "acidic species" if stirring is vigorous & long continued.]
2751	167	Acetilacetonitrile, Me.CH(O.OC.Me).CN . — Sp. gr. 1.032 (14°). — S. in 25 pt. aq.; e.s. alc., eth. — Heated w. alk., or w. aq. at 140° gives acetaldehyde, ac. ac., & HCN.
2752	168	† Methylpropylketoxime, Me.C(:NOH).Pr . — Sp. gr. 0.907 (20/0). — (P) Oxime (T. 2.17).
2753	169.5	1,2,5-Trimethylpyrrole, $^{\text{NMe:CM:CH.CH:CM}}\text{Me}$. — E. vol. w. st. w. pine resin odor! E.s. alc., eth., bz. Is colored cherry-red when boiled w. FeCl_3 sol.
2754	170	Ethyl Methylcarbamate, MeNH.CO.Et . — Sp. gr. abt. 1. — (P) (P) Sapn. T.* products: methylamine (No. 2.1059); ethyl alc.; CO_2 .

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Liquids.
2755	abt. 170d.	Nitrosoparaldimine, $C_6H_{12}O_2N.NO$. — \textcircled{P} Sol. in conc. H_2SO_4 blood-red, becoming deep blue on addition of KOH! — Lemon-yel. liq. I. aq.; misc. alc., eth. — Boiled w. HCl gives paraldehyde (Vol. I).
2756	170-1	Cyclopentanecarbononitrile, $^rCH(CN).(CH_2)_4^r$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; cyclopentanecarbonic ac. ($C_6H_{10}O_2$, Vol. I).
2757	171	3-Nitro-3-methylpentane, $Me.CH_2.CMe(NO_2).Et$. — Sp. gr. 0.977 (0 $^{\circ}$). — \textcircled{P} Nitro comp. (T. 2.21).
2758	170-4	2-Nitro-2,3-dimethylbutane, $Me.CMe(NO_2).CHMe.Me$. — Sp. gr. 0.961 (20/4). — \textcircled{P} Nitro comp. (T. 2.21). — Scales. M.p. 5-7 $^{\circ}$.
2759	171c.	2,4-Dimethylpyrrole, $^rNH.CMe:CH.CMe:CH^r$. — Odor pungent & chloroform-like. V. vol. w. st. D.s. aq.; e.s. alc., eth., bz. — Aq. sol. colored cherry-red on warming w. FeCl ₃ .
2760	174-5	Ethyl Ethylcarbamate, $EtNH.CO.Et$. — Sp. gr. 0.986 (21 $^{\circ}$). — \textcircled{P} \textcircled{D} Sapn. T.* (w. NaOH) products: ethylamine (No. 2.1062); ethyl alc.; CO ₂ .
2761	175.4c.	† Nitrosodiethylamine, $Et_2N.NO$. — Sp. gr. 0.951 (17.5 $^{\circ}$). — \textcircled{P} Should give blue color w. diphenylamine reagt. — <i>Y T'2 out of pungent mint-like odor. Taste v. pungent & mint-like.</i> — Boiling w. conc. HCl gives diethylamine (No. 2.1068-1), & nitrous ac.
2762	175-7	n-Octyl Nitrite, $C_8H_{17}NO_2$. — Sp. gr. 0.862 (17 $^{\circ}$). — \textcircled{P} Nitrite. § — \textcircled{D} Sapn. T.* (w. NaOH) products: n-octyl alc. ($C_8H_{16}O$, Vol. I); nitrous ac.
2763	176c.	2-Nitrohexane, $Me.CH(NO_2).(CH_2)_4.Me$. — Sp. gr. 0.936 (20/0). — \textcircled{P} Nitro comp. (T. 2.21).
2764	175-8(th.i.)	Hexyl Cyanide, $C_6H_{13}N.CN$. — Sp. gr. 0.895 (22 $^{\circ}$). — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; oenanthrylic ac. (Vol. I).
2765	177(th.i.)	Tetramethylurea, $Me_2N.CO.NMe_2$. — Sp. gr. 0.972 (15 $^{\circ}$). E.s. alc., eth. — \textcircled{P} \textcircled{D} Sapn. T.* products: dimethylamine (No. 2.1061); CO ₂ .
2766	178(th.i.)	Diethylformamide, $Et_2N(CHO)$. — Sp. gr. 0.908 (19 $^{\circ}$). Misc. aq. Salted out fr. sol. by K ₂ CO ₃ . — \textcircled{P} \textcircled{D} Sapn. T.* products: diethylamine (No. 2.1068-1); formic ac.
2767	175-85(sl.d.)	† Ethaneazobenzene, $Et.N:NPh$. — Pale yel. pungent smelling oil. Vol. w. st. V.d.s. aq.; e.s. alc., eth., bz.; d.s. v. dil. ac.; e.s. conc. ac. Mixed w. 60% H ₂ SO ₄ gradually changes to acetaldehydophenylhydrazone.
2768	180-1	1-Nitrohexane, $Me.(CH_2)_4.CH_2.NO_2$. — Sp. gr. 0.960 (17 $^{\circ}$). — \textcircled{P} Nitro comp. (T. 2.21). — Pale yellowish oil of ethereal odor. I. aq.; s. alk. — Gives color in nitrolic ac. test described for nitromethane (No. 2.2688).
2769	181-2(th.i.)	1-Acetylpyrrole, $^rN(C_2H_5O).CH:CH.CH:CH^r$. — \textcircled{P} Hot vapors redden pine splinter moistened w. HCl in T. 2.24! — Resinified by HCl. — Alm. i. aq. Aq. sol. pptd. by HgCl ₂ . Reduces AgNO ₃ .
2770	182-4(sl.d.)	† α -Ethylenelactonitrile, $Me.CH(OH).CN$. — Misc. alc., aq. — \textcircled{P} \textcircled{D} Sapn. T.* w. KOH gives HCN & aldehyde resin; w. conc. HCl, lactic ac. (Vol. I) & NH ₄ Cl.
2771	183d.	Glycollonitrile, $HO.CH_2.CN$. — Sp. gr. 1.100 (12 $^{\circ}$).
2772	181-5	Diisopropylketoxime, $Me_2.CH.C(:NOH).CH.Me_2$. — \textcircled{P} Oxime (T. 2.17). — M.p. +6-8 $^{\circ}$.
2773	183-4	o-Tolyl Isocyanide, $Me.C_6H_4.N:C$. — Sp. gr. 0.968 (24 $^{\circ}$). — \textcircled{P} Powerful, nauseating & persistent, carbylamine odor! — Turns greenish yellow, & later dark yellow, on keeping, but is more stable than No. 2.2746.
2774	184	† 3-Cyanopentanol(3), $Et_2C(OH)(CN)$. — Sp. gr. 0.933 (22 $^{\circ}$). — \textcircled{P} Odor agreeable. — I. aq.; s. alc., eth.
2775	185-6	Diethylacetamide, $Me.CO.NEt_2$. — Sp. gr. 0.925 (8.5 $^{\circ}$). — \textcircled{P} Sapn. T.* products: diethylamine (No. 2.1068-1); ac. ac.
2776	186	o-Tolyl Isocyanate, o-Tolylcarbonimide, $Me.C_6H_4.N:C:O$. — \textcircled{P} Vapors exceedingly irritating to eyes & nose! — \textcircled{D} Decd. by aq. to o-tolylurea, No. 2.2416, & CO ₂ . [Hence prob. neutralizing some alk. in titration of Gen. T. 2.I.]

Explanation of typographical signs used in this Division: * = T. 2.26. † = T. 2.21. § = T. 2.15.

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Liquids.
2777	187	p-Tolyl Isocyanate, p-Tolylcarbonimide, Me.C ₆ H ₄ .N:C:O. — (P) Action of vapors as for No. 2.2776. — (D) Deen. by aq. gives p-tolylurea, No. 2.2373, & CO ₂ .
2778	185–90	3-Nitro-3-ethylpentane, Me.CH ₂ .CEt(NO ₂). — Sp. gr. 0.955 (0°). — (P) Nitro comp. (T. 2.2 & prob. T. 2.17-b).
2779	abt. 188	Diethylcyanamide, Et ₂ N.CN. — (P) (D) Sapn. T.* (w. HCl) products: diethylamine (No. 2.1068–1); NH ₃ ; CO ₂ .
2780	abt. 189	3-Ethyl-4-oximinopentane, Et.CHEt.C(:NOH).Me. — Sp. gr. less than 1. — (P) Oxime (T. 2.17).
2781	191c.	† Benzonitrile, Phenyl Cyanide, Ph.CN. — Sp. gr. 1.0005 (24.8/4). Strong bitter almond odor & taste! S. in 100 pt. boiling aq.; misc. alc., eth. — (P) (D) Sapn. T.* 2.26-c gives NH ₃ & benzoic ac. (Vol. I). (Do not dry the benzoic ac. at 100° for more than 10 min., or there will be much loss by sublimation.)
2782	abt. 193	Dipropylketoxime, Pr ₂ C:NOH. — Oxime (T. 2.17).
2783	192–5d.	† Formamide, H.CO.NH ₂ . — Sp. gr. 1.337 (14/4). — Boils w. partial deen. to NH ₃ & CO. — (P) (D) Sapn. T.* (w. NaOH) products: NH ₃ ; formic ac. (Vol. I). Conc. KOH gives NH ₃ , even in the cold.
2784	193–5	1-Nitroheptane, Me.(CH ₂) ₆ .CH ₂ .NO ₂ . — (P) Nitro comp. (T. 2.21 & prob. 2.17-b). — Pale yellowish oil. I. aq.; e.s. alc., eth. Gives nitrolie ac. color react. (Cf. No. 2.2688).
2785	195–6d.	Methylisoamylketoxime, Me.C(:NOH).C ₆ H ₁₁ . — Sp. gr. 0.888 (20/4). — (P) Oxime (T. 2.17).
2786	194–8(sl.d.)	2-Nitroheptane, Me.CH(NO ₂).(CH ₂) ₅ .Me. — Sp. gr. 0.947 (0°). — (P) Nitro comp. (T. 2.21). — S. in warm conc. KOH sol.
2787	198–200	Caprylonitrile, C ₇ H ₁₅ .CN. — Sp. gr. 0.820 (13°). — (P) (D) Sapn. T.* products: NH ₃ ; caprylic ac.
2788	199	Ethylformamide, EtNH.(CHO). — Sp. gr. 0.952 (21°). — (P) (D) Sapn. T.* products: ethylamine (No. 2.1062); formic ac. (Vol. I).
2789	199c.	2,2-Dinitrobutane, Me.CH ₂ .C(NO ₂) ₂ .Me. — “Neutral oil.” (P) Nitro comp. (T. 2.21).
2790	201–2	2-Nitro-2,5-dimethylhexane, Me.CMe(NO ₂).CH ₂ .CHMe ₂ . — Sp. gr. 0.920 (20/0). — (P) Nitro comp. (T. 2.21). — Reduction gives diisobutylamine.
—	204	5-Methylpyrazole. — Described as No. 2.1268. — [k _B .10 ¹¹ = 3.6.]
2791	205	Ethylacetamide, Me.CO.NHET. — Sp. gr. 0.942 (4.5°). — (P) (D) Sapn. T.* products: ethylamine (No. 2.1062); ac. ac.
2792	205.2(th.i.)	† o-Tolunitrile, Me.C ₆ H ₄ .CN. — Sp. gr. 0.9975 (15°). — Odor like nitrobenzene! I. aq.; misc. alc. — (P) Gives NH ₃ in T. 2.7. — (D) Sapn. T.* 2.26-c (w. alc. KOH, boiling for 1 hr.) gives o-toluamide (No. 2.2134), which is recrystd. first fr. 8 & then fr. 5 cc. boiling aq.
2793	206–10(sl.d.)	1-Nitrooctane, Me.(CH ₂) ₆ .CH ₂ .NO ₂ . — Sp. gr. 0.935 (20°). Pale yellowish oil of agreeable odor. — (P) Nitro comp.†
2794	208–10	m-Tolunitrile, Me.C ₆ H ₄ .CN. — (P) (D) Sapn. T.* products: NH ₃ ; m-toluic ac. (C ₆ H ₅ O ₂ , Vol. I). — S. in 60 pt. boiling, or 1170 pt. c. aq.
2795	210 ± 5	Tetraethylurea, Et ₂ N.CO.NEt ₂ . — (P) Odor peppermint-like. — S. acids; repptd. by alk. — (D) Sapn. T.* products: diethylamine (No. 2.1068–1); CO ₂ .
2796	210.0c.	† Nitrobenzene, Ph.NO ₂ . — Sp. gr. 1.2116 (13/4). M.p. +5.4°. Color (freshly distd.), GYT ² . Odor, bitter almond-like! Taste of c. sat. filtered aq. sol., sweet (No. 9 on scale of T. 2.29). V.d.s. aq.; v.s.c. alc., eth.; misc. bz.
		(P) (1) Gives nitro comp. test (T. 2.21). — (2) Place in a 30-cc. dislg. flask granulated tin (bulk of a pea), 1 drop nitro comp., & 10 drops conc. HCl. Boil v. gently for 3 min. Add 5 cc. of 10% NaOH sol., cooling, & drop in ebullator tube. Dist. off slowly into well-cooled t.t., without use of condenser, abt. 5 drops of liquid. Dil. w. 5 cc. aq. Add 5 drops satd. bleaching powd.

No.	Boiling-point (C°).	NEUTRAL COMPOUNDS.—Liquids.
		<i>sol. Shake & filter. On further dilution w. aq. a VR sol. (react. fr. aniline) will be obtained.</i> ① Convert 5 drops to <i>m</i> -dinitrobenzene, following procedure given for benzene in T. 1.913.
2797	210-2d.	2-Nitrooctane, $\text{Me}(\text{CH}_2)_7\text{CH}(\text{NO}_2)\text{Me}$. — Sp. gr. 0.936 (0°). — ② Nitro comp.‡
2798	213-6	Nitrosodiisobutylamine, $(\text{Me}_2\text{CHCH}_2)_2\text{NNO}$. — Lighter than aq. Odor unpleasant. Solidifies in freezing mixt. — ② Prob. gives blue color in T. 2.15.
2799	214-6	Pelargononitrile, $\text{Me}(\text{CH}_2)_7\text{CN}$. — Sp. gr. 0.786 (16°). — ② ③ Sapn. T.* products: NH_3 ; pelargonic ac. (Vol. I).
2800	215-6	Acetonemethylphenylhydrazone, $\text{Me}_2\text{C}:\text{N.NMePh}$. — ② Hydrzone (T. 2.17). [“Reduces h. ammon. AgNO_3 , but not Fehling’s sol.”] — D.s. h. aq.; e.s. alc., eth., lgr.
2801	216.5	Nitrosopiperidine, $\text{C}_4\text{H}_{10}\text{NNO}$. — Sp. gr. 1.0659 (16.5°). — ② Prob. gives blue color in T. 2.15. — Mod. s. aq.; v.s. conc. HCl. — Reduction w. Zn dust & HCl gives NH_3 & piperidine (No. 2.1112).
2802	215-8d.	1-Nitrononane, $\text{Me}(\text{CH}_2)_7\text{CH}_2\text{NO}_2$. — Sp. gr. 0.923 (17°). Pale yel. liq. — ② Nitro comp. (T. 2.21 & prob. T. 2.17-b).
2803	abt. 217	Methylhexylketoxime, $\text{Me.C}(:\text{NOH})\text{C}_6\text{H}_{12}$. — Sp. gr. 0.886 (20/4). — ② Oxime (T. 2.17).
—	217.3c.	p-Tolunitrile. — Cf. No. 2.1498. (M.p. 29-30°)
2804	220.4c.	† o-Nitrotoluene, $\text{Me.C}_6\text{H}_4\text{NO}_2$. — Sp. gr. 1.168 (15°). M.p. -10.5°. Yel. liq. of bitter almond odor. Taste of c. sat. aq. sol. burning & sweet (No. 1 on sweetness scale, T. 2.29). Alm. i. c. aq.; misc. alc., eth., bz. ② (1) Nitro comp.‡ — (2) Warm together for 1 min. 5 drops substance, 5 cc. alc., a bit of Na half as large as a pea, & 0.5 g. Zn dust. Cool & add 2 cc. conc. HCl. Heat 30 sec. & allow to stand 5 min. Filter. Filtrate shows permanent RT1 color. ③ Place 3 drops substance, 1 cc. fuming HNO_3 , & 1 cc. H_2SO_4 (sp. gr. 1.84) in 6-inch t.t. Boil v. gently for 2 min. Cool. Pour into 5 cc. c. aq. Shake vigorously! Filter. Wash w. 3 cc. aq. Dis. in 10 cc. boiling 50% alc. Filter hot. After 15 min. wash cryst. w. 2 cc. 50% alc. Recryst. fr. 5 cc. boiling 50% alc. Filter hot. After 10 min. turbid sol. clears, yielding fine colorless ndl. Wash w. 2 cc. 50% alc. Dry 15 min. on tile at abt. 50°. The product, 2,4-dinitrotoluene, has bitter almond odor, sweet taste, & melts at 70.7° u.c.
2805	221-3	Ethylenecyanhydrine, $\text{HO.CH}_2\text{CH}_2\text{CN}$. — Sp. gr. 1.059 (0°). Misc. aq., alc.; s. eth. — Sapn. T.* (w. HCl) gives NH_3 , acrylic ac. & hydrylic ac.
2805	222	N-Formylpiperidine, $(\text{CHO})\text{NC}_6\text{H}_{10}$. — Sp. gr. 1.0193 (23/4). Misc. aq., alc. — ② ③ Sapn. T.* products: piperidine (No. 2.1112); formic ac. — B.HgCl_2 , fine ndl., m.p. 148-9°.
2807	222	1,3-Dimethylbenzonitrile(4), $\text{Me}_2\text{C}_6\text{H}_3\text{CN}$. — Sp. gr. 0.9871 (19°). M.p. 23-5°. Odor bitter almond like. — ② ③ Sapn. T.* products: 1,3-dimethylbenzoic ac.(4), ($\text{C}_9\text{H}_{10}\text{O}_2$, Vol. I).
2808	226(th.i.)	† 2-Nitro-1,3-xylene, $\text{NO}_2\text{C}_6\text{H}_3\text{Me}_2$. — Sp. gr. 1.112 (15°). — ② Nitro comp.‡
2809	225 ± 2c., d.	† Nitrosomonomethylaniline, Methylphenylnitrosamine, $(\text{NO})\text{Me.NPh}$. — YO oil of aromatic odor, after distn. w. st. Boils at atmospheric pressure w. formation of brown fumes & much tar. I. c. aq.; v.s. alc., eth., bz., chlf. — ② Gives deep blue color w. diphenylamine reag. (T. 2.15)! — ③ Dis. 6 drops substance in 0.5 cc. dry eth. Add 2 cc. abs. alc. said. w. dry HCl gas. Allow to stand over night in stoppered t.t. Filter off the yel. cryst. of <i>p</i> -nitrosomonomethylaniline hydrochloride which separates fr. the dark orange sol. Wash w. 2 cc. alc. + 2 cc. eth. Dis. in 2 cc. c. aq. Add 0.5 cc. ammon. (sp. gr. 0.90). Filter off the shimmering dark green lft. Wash w. 2 cc. ammon. Dry at 50°. Recryst. fr. 3 cc. h. bz., allowing sol. to concentrate by spontaneous evapn. Filter off the dark steel-blue ndl. of <i>p</i> -nitrosomonomethylaniline & wash w. 1 cc. c. bz. Dry 15 min. at 100°. The product melts at 114.5-5° u.c. (r.h.).

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.15.

(ORDER II, SUBORDER I.)

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Liquids.
2810	226-7	† α -Campholenonitrile, $C_9H_{14}CN$. — Sp. gr. 0.915 (23°). $n_D^{20} = 1.46653$. I. aq. acids; s. alc., eth. — \textcircled{P} \textcircled{D} Sapn. T.* (w. alc. KOH — protracted boiling) products: NH_3 ; campholenic ac. ($C_{10}H_{16}O_2$, Vol. I).
2811	226-7	N-Acetylpyridine, $(C_6H_5O)NC_6H_5$. — Sp. gr. 1.0111 (9°). Misc aq. — \textcircled{P} \textcircled{D} Sapn. T.* products: pyridine (No. 2.1112); ac. ac. — $B_2H_6PtCl_6$ (T. 2.14), red cryst., m.p. 107-9°, e.s. aq.
2812	227	p-Propylbenzonitrile, $Pr.C_6H_4CN$. — \textcircled{P} \textcircled{D} Sapn. T.* ("w. conc. HCl at 100°") products: NH_3 ; p-propylbenzoic ac. ($C_{10}H_{16}O_2$, Vol. I).
2813	227-8	σ -Nitroethylbenzene, $NO_2.C_6H_5Et$. — Sp. gr. 1.126 (24.5°). — \textcircled{P} Nitro comp.† — "Not oxidized by CrO_3 mixt."
2814	230-2	1,2-Dimethylbenzonitrile(4), $Me_2C_6H_3CN$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; 1,2-dimethylbenzoic ac.(4), ($C_8H_{10}O_2$, Vol. I). — Misc. alc., eth.
2815	232c.	† m-Nitrotoluene, $Me.C_6H_4.NO_2$. — Sp. gr. 1.168 (22°). M.p. +15.9° u.c. ($Y-GY$) T_2 oil of faint bitter-almond-like odor. Taste of c. sat. aq. sol., sweet & burning (Nos. 1 & 2, T. 2.29). Alm. i. c. aq.; misc. alc., bz., eth. — \textcircled{P} Nitro comp.† — \textcircled{D} Reduce to m-nitrotoluidine by treatment w. granulated Sn & conc. HCl. Add x.s. NaOH & dist. w. steam. Separate a few drops of oily toluidine w. pipette & acetylate by heating to boiling w. 2 vol. ac. anhydride. Recryst. resulting m-acettoluide fr. a little boiling aq., & dry at 50°. (The solution has a tendency to remain supersaturated.) The product melts at 65° u.c.
2816	233.5(th.i.)	† Benzyl Cyanide, $Ph.CH_2CN$. — Sp. gr. 1.0214 (15/15). M.p. -24.6°. — \textcircled{P} \textcircled{D} Sapn. T.* ("w. 3 vol. conc. H_2SO_4 + 2 vol. aq.") products: NH_3 ; phenylacetic ac. (Vol. I).
2817	234-7c.	Nitrosodibutylamine, $(C_6H_5)_2N.NO$. — Yellowish oil. — \textcircled{P} Prob. gives blue color w. diphenylamine reag. (T. 2.15).
2818	236d.	γ -Nitrobutyronitrile, $NO_2.C_4H_7CH_2CN$. — Sp. gr. 1.138 (12°). — \textcircled{P} Nitro comp.† — Has faint odor & sweetish pungent taste. I. aq.; e.s. alc., eth.
2819	235-7d.	2-Nitro-2,7-dimethyloctane, $Me.CMe(NO_2).(CH_2)_6CHMe.Me$. — Sp. gr. 0.909 (20/4). Not solidified by freezing mixt. — \textcircled{P} Nitro comp.†
2820	235-7	Caprinitrile, $Me.(CH_2)_8CN$. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; capric ac. (Vol. I).
2821	237	† Methyl Dimethyloxamate, $Me_2N.CO.CO_2Me$. — Sp. gr. 1.105 (15°). — \textcircled{P} \textcircled{D} Sapn. T.* products: dimethylamine (No. 2.1061); methyl alc.; oxalic ac.
2822	239(th.i.)	2-Nitro-1,4-xylene, $NO_2.C_6H_3Me_2$. — Sp. gr. 1.132 (15°). Vol. w. st. Alm. i. aq. — \textcircled{P} Nitro comp.†
2823	241.5-2(th.i.)	N-Methylindole, $Me.NC_6H_5$. — Sp. gr. 1.0707 (0°). — Yel. oil. Alm. i. aq.; v.s. alc., eth., bz. — \textcircled{P} Gives red-violet color in pyrrole splinter react., T. 2.24. — \textcircled{D} B_2Pk (T. 2.23), long dark red pr. fr. eth., m.p. 150°.
2824	242	m-Nitroethylbenzene, $NO_2.C_6H_5Et$. — Sp. gr. 1.134 (0°). — \textcircled{P} Nitro comp.†
2823	243-4c.	4-Nitro-1,3-xylene, $NO_2.C_6H_4Me_2$. — Sp. gr. 1.126 (17.5°). — \textcircled{P} Nitro comp.†
2826	243-4	Ethyl Methylphenylcarbamate, $Me.PhN.CO.Et$. — \textcircled{P} \textcircled{D} Sapn. T.* products: methylaniline (No. 2.1249); ethyl alc.; CO_2 .
2827	244(th.i.)	Cuminonitrile, $p-(Me_2CH)_2C_6H_3CN$. — Sp. gr. 0.765 (14°). D.s. aq.; misc. alc., eth. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH_3 ; cuminic ac. ($C_{10}H_{12}O_2$, Vol. I). Treatment w. c. alc. KOH sol. gives cuminamide (No. 2.2207).
2828	244d.	1'-Nitroisobutylbenzene, $Me_2CH.CH(NO_2).Ph$. — \textcircled{P} Nitro comp.† — Slowly disd. by aq. NaOH sol. Dil. H_2SO_4 ppts. the unstable "iso" form fr. this sol., m.p. 54° d. — Gives Konowalow's react. (T. 2.41).
2829	245-6	p-Nitroethylbenzene, $NO_2.C_6H_5Et$. — Sp. gr. 1.124 (25°). — \textcircled{P} Nitro comp.† — Oxidation by CrO_3 mixt. (Cf. T. 1.905-2) gives p-nitrobenzoic ac. (No. 2.425).

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS.—Liquids.
2830	244-6	Ethyl Ethyloxamate, EtNH.CO.CO.Et. — S. aq., alc., eth. — $\oplus \ominus$ Sapn. T.* products: ethylamine (No. 2.1062); ethyl alc., & oxalic ac. (Vol. I).
2831	245	Allylsuccinimide, $[N(C_2H_5).CO.CH_2.CH_2.CO]$. — Sp. gr. 1.154 (0 $^{\circ}$). — $\oplus \ominus$ Sapn. T.* products: allylamine (No. 2.1068); succinic ac. (Vol. I).
2832	246.5c.	1-Phenylpyrazole, $[NPh.N:CH.CH_2:CH_2:CH]$. — Sp. gr. 1.1125 (16 $^{\circ}$). Golden yel. oil. Solidified by freezing mixt.; m.p. 11-11.5 $^{\circ}$. I. aq.; s. alc., eth.; s. conc. HCl, but repprd. by aq. — \oplus Gives Knor's pyrazole react. (Ann., 238, 200; Ber., 26, 100.) — $B_2H_2PtCl_6.2H_2O$, red-yel. ndl., m.p. 171-2 $^{\circ}$ d.
2833	246.5c.	1-o-Tolylpyrazole, $[N(C_6H_5.Me).N:CH.CH_2:CH_2:CH]$. — Sp. gr. 1.0868 (0 $^{\circ}$). Remains liquid at -10 $^{\circ}$. — $B_2H_2PtCl_6$, yel.-red pr., m.p. 200-1 $^{\circ}$ d.
2834	249-51	Dimethyloxanilide, Me ₂ PhN.CO.CO.NPh.Me. — $\oplus \ominus$ Sapn. T.* products: methylaniline (No. 2.1249); oxalic ac.
2835	251d. (th.i.)	3-Nitro-1,2-xylene, NO ₂ .C ₆ H ₄ .Me ₂ . — Sp. gr. 1.147 (15 $^{\circ}$). Vol. w. st. — \oplus Nitro comp.‡
2836	250-4	† Ethyl Diethyloxamate, Et ₂ N.CO.CO.Et. — $\oplus \ominus$ Sapn. T.* products: diethylamine (No. 2.1068-1); ethyl alc.; oxalic ac. — I. aq.
2837	252-3c.	N-Ethylindole, C ₈ H ₇ N.Et. — Sp. gr. 1.256 (15 $^{\circ}$). Oil of not unpleasant odor. — B_2P_2 , red ndl. fr. lgr., m.p. 105 $^{\circ}$.
2838	253	m-Nitroisobutylbenzene, NO ₂ .C ₆ H ₄ .CH ₂ .CHMe ₂ . — \oplus Nitro comp.‡ — Yellowish oil of aromatic odor.
2833	250-6d.	1' Nitrobutylbenzene, Me.(CH ₂) ₃ .CH(NO ₂).Ph. — Sp. gr. 1.059 (2010). — \oplus Nitro comp.‡
2840	253-4	Undecylonitrile, Me.(CH ₂) ₁₀ CN. — $\oplus \ominus$ Sapn. T.* products: NH ₃ ; undecylic ac. (C ₁₁ H ₂₂ O ₂ , Vol. I).
2841	254-5	Cinnamonic nitrile, Ph.CH:CH.CN. — Sp. gr. 1.037 (0 $^{\circ}$). M.p. +11 $^{\circ}$. E.s. alc. — $\oplus \ominus$ Sapn. T.* products: NH ₃ ; cinnamic ac. (Vol. I).
2842	257-9	Triethyloxamide, EtNH.CO.CO.NEt ₃ . — Misc. aq. — $\oplus \ominus$ Sapn. T.* products: ethylamine & diethylamine (Nos. 2.1062 & 2.1068-1); oxalic ac.
2843	258	Ethyl-p-acettoluide, Et.(C ₆ H ₅ O)N.C ₆ H ₄ .Me. — $\oplus \ominus$ Sapn. T.* products: ethyl-p-toluidine (No. 2.1314); ac. ac.
2844	259c.	Ethyformanilide, Et.(CHO)N.Ph. — Sp. gr. 1.063 (16/4). — $\oplus \ominus$ Sapn. T.* products: ethylaniline (No. 2.1270); formic ac. (Vol. I).
2845	260-2	p-Isoamylbenzonitrile, C ₈ H ₁₁ .C ₆ H ₅ CN. — $\oplus \ominus$ Sapn. T.* products: NH ₃ ; p-isoamylbenzoic ac. (C ₁₂ H ₂₁ O ₂ , Vol. I).
2845	266-8	Ethyl o-Aminobenzoate, NH ₂ .C ₆ H ₄ .CO.Et. — Heavier than aq. M.p. 13 $^{\circ}$. D.s. c. alc.; i. eth. Odor faint. — $\oplus \ominus$ Sapn. T.* products: o-aminobenzoic ac. (No. 2.148); ethyl alc. (Vol. I).
2847	268(th.i.)	Propylformanilide, Pr.(CHO)N.Ph. — Sp. gr. 1.044 (16/4). — $\oplus \ominus$ Sapn. T.* products: propylaniline (No. 2.1329); formic ac.
2840	abt. 268; 258	o-Nitrophenetole, NO ₂ .C ₆ H ₄ .OEt. — Yel. oil of mild aromatic odor. — \oplus Nitro comp.‡
2845	268-71	Ethyl-p-propiontoluicide, Et ₂ N(C ₆ H ₅ O).C ₆ H ₄ .Me. — $\oplus \ominus$ Sapn. T.* products: ethyl-p-toluidine (No. 2.1314); propionic ac. (Vol. I).
2850	269-70	α -Naphthyl Isocyanate, C ₁₀ H ₇ .N:C:O. — \oplus Odor v. irritating. \ominus Sapn. T.* products: α -naphthylamine (No. 2.589); CO ₂ .
2851	271d.	6-Nitro-1,3-dimethyl-5-ethylbenzene, (NO ₂).Me ₂ .Et.C ₆ H ₃ . — \oplus Nitro comp.‡
2852	265; 277	† o-Nitroanisole, NO ₂ .C ₆ H ₄ .OMe. — Sp. gr. 1.249 (26 $^{\circ}$); 1.268 (20 $^{\circ}$). M.p. +9 $^{\circ}$. Yellowish oil of mild aromatic odor.
2853	274(th.i.)	Isobutylacetanilide, (C ₃ H ₇).(C ₆ H ₅ O)N.Ph. — $\oplus \ominus$ Sapn. T.* products: isobutylaniline (No. 2.1354-1); ac. ac.
2854	275	Tridecyronitrile, Me.(CH ₂) ₁₁ CN. — E.s. alc., eth. — $\oplus \ominus$ Sapn. T.* products NH ₃ ; tridecyclic ac. (C ₁₅ H ₃₀ O ₂ , Vol. I).

Explanation of typographical signs used in this Division: * = T. 2.26. ‡ = T. 2.21. § = T. 2.15.

No.	Boiling-point (C. $^{\circ}$).	NEUTRAL COMPOUNDS. — Liquids.
2855	275	Butylacetanilide, $(C_6H_5)(C_2H_5O)NPh$. — \textcircled{P} \textcircled{D} Sapn. T.* products: butylaniline (No. 2.1358); ac. ac.
2856	275	Methyl o-Nitrobenzoate, $NO_2C_6H_4CO_2Me$. — Sp. gr. 1.2855 (20 $^{\circ}$). M.p. —13 $^{\circ}$. Odor strawberry-like. — \textcircled{P} Nitro comp.† — \textcircled{D} Sapn. T.* products: o-nitrobenzoic ac. (No. 2.164); methyl alc.
2857	280	Pr-1,2,3-Trimethylindole, $Me_3C_6H_3N$. — M.p. 18 $^{\circ}$. D.s. h. aq.; e.s. alc., eth., bz. — \textcircled{P} \textcircled{D} B.Pk (T. 2.23), dark red ndl. fr. bz., m.p. 150 $^{\circ}$.
2858	281c.	Diethylbenzamide, $Et_2N.CO.Ph$. — Sp. gr. 1.019 (15 $^{\circ}$). S. in HCl, but repprd. by aq. — \textcircled{P} \textcircled{D} Sapn. T.* products: diethylamine (No. 2.1068-1); benzoic ac.
2859	283-5c.	Pr-3-Ethylindole, β -Ethylindole, $Et.C_6H_4N$. — Pale yel. oil of unpleasant fecal odor! Vol. w. st. V.d.s. aq.; e.s. alc., eth.; i. dil. ac. or alk. — \textcircled{P} Gives indole pine splinter react. in T. 2.24-b. — Picrate (T. 2.23) pptd. fr. bz. sol. by lgr. in red flocks, m.p. 143 $^{\circ}$.
2860	285-7. 4c.	Trimethylene Cyanide, $CN.(CH_2)_2CN$. — Sp. gr. 0.995 (15 $^{\circ}$). I. eth. — \textcircled{P} \textcircled{D} Sapn. T.* products: NH ₂ ; glutaric ac. (Vol. I).
2861	288(th.i.)	Isoamylacetanilide, $(C_{11}H_{11})(C_2H_5O)NPh$. — E.s. alc., eth. — \textcircled{P} \textcircled{D} Sapn. T.* products: isoamylaniline (No. 2.1393); ac. ac.
—	288-90	m-Nitrodiethylaniline. — Cf. No. 2.1441. (Yel. oil.)
2862	291-3(th.i., 750 mm.)	Pr-2,3-Methylethylindole, $Me.Et.C_6H_3N$. — V.d.s. aq.; v.e.s. alc., eth.
2863	abt. 295	Ethyldiphenylamine, $EtN.Ph_2$. — I. aq. — \textcircled{P} HNO ₃ gives violet-red color. (Cf. T. 2.15.)
2864	296	Nitrosodipropylamine, $Pr_2N.NO$. — Sp. gr. 0.924 (14 $^{\circ}$). Yel. liq. of aromat. odor. I. aq. — \textcircled{P} Should give blue color in T. 2.15.
	296c.	† Methyldiphenylamine, $MeN.Ph_2$. — Sp. gr. 1.052 (15/15). I. aq. — \textcircled{P} HNO ₃ gives violet coloration. — \textcircled{D} Nitroso deriv., $Me_2PhN.C_6H_4NO$ (Cf. Compt. rend., 124, 898 for preparation), green lft., m.p. 44 $^{\circ}$.
2866	300-5	Phenyl-m-toluidine, $PhNH.C_6H_4Me$. — Reddish oil. — \textcircled{P} Sol. in conc. H ₂ SO ₄ , is colored intensely green by little HNO ₃ !
2867	313-4d. (th.i.)	Di-o-tolylamine, $(Me.C_6H_4)_2NH$.
2868	319-20	Di-m-tolylamine, $(Me.C_6H_4)_2NH$. — Still liq. at -12 $^{\circ}$. Vol. w. st. E.s. alc., eth.; i. aq.
2859	326-7(th.i.)	N-Phenylindole, $C_6H_5N.Ph$. — Yellowish oil. I. aq.; e.s. alc., eth., bz. — \textcircled{P} Gives blue-violet color in pine splinter react., T. 2.24-b!
2870	330-1c.	† Carbodiphenylimide, $PhN:C:NPh$. — Syrup slowly stiffening to glassy mass. Polymerizes alone or in sol. to cryst. polymer, m.p. 160-1 $^{\circ}$. — Reacts vigorously w. aniline, giving α -triphenylguanidine (No. 2.882). — Boiled w. alc. HCl gives carbanilide (No. 2.2580).
2871	330-40	Isoamylidiphenylamine, $(C_{11}H_{11})N.Ph_2$. — I. aq.
2872	314-20 (223 mm.)	Pr-2-Methyl- β -naphthindole, $[C_{10}H_8.NH.CMe:CH]^2$. — V.d.s. aq.; e.s. alc., eth., bz. — \textcircled{P} Colors pine splinter like No. 2.2873 below. — B.Pk (T. 2.23), fine red-brown ndl. fr. bz., m.p. 176 $^{\circ}$.
2873	a. 360; 222 (th.i., 18 mm.)	$\beta\beta$ -Naphthindole, $[C_{10}H_8.NH.CH:CH]^2$. — E.s. alc., eth., bz.; d.s. lgr.; sl. s. aq. Sol. fluor. green-blue! — \textcircled{P} Pine splinter soaked in alc. sol. is colored intense violet when immersed in conc. HCl (T. 2.24-b). — B.Pk, fine dark red ndl. fr. bz.; i. aq.
2874	370-5c.	o-Tolyl-m-aminophenol, $(Me.C_6H_4.NH).C_6H_4.OH$. — Brownish oil. E.s. alc., eth., bz. — Ignition w. Zn dust gives acridine (No. 2.3102). — Boiled w. formic ac. gives formyl deriv., tbl. fr. alc., m.p. 169 $^{\circ}$.

SUBORDER II, DIVISION A.
(ORDER II.)

COLORED COMPOUNDS CONTAINING C, N, H, AND OFTEN O.

DIVISION A, SOLID* SPECIES.

No.	Melting-point (C.).	Color.	SPECIFIC NAMES.—Tests and Miscellaneous Properties.
2875	18-9	Dark red	Benzene-azo-m-toluene, Ph.N ₂ C ₆ H ₄ .Me. — Cryst. mass. Sp. gr. 1.065 (20/4). B.p. 179° (19 mm.). — ⊕ Azo comp.
—	29	Light yel.	4-Nitro-1,2-xylene. — Cf. No. 2.1475.
2876	30	Yellow	Nitroso-o-nitroethylaniline, (NO)EtN.C ₆ H ₄ .NO ₂ . — Long ndl. fr. gl. ac. ac. — ⊕ Prob. gives T. 2.15 w. diphenylamine reagt.
2877	30	Yellow	Benzooquinoneoximeethylether, O:C ₆ H ₄ :NOEt. — Tbl. fr. lgr.
2878	33.5	Yellow	3-Nitrocresol(4), NO ₂ .C ₆ H ₄ (Me)(OH). — Flat ndl. Vol. w. st. V.s. alc., eth. — ⊕ Nitro comp.† giving colored sol. in alk. — NaA, dark red ndl. — AgA, brick-red ppt.
2879	35-6	Red	o-Nitromethylaniline, NO ₂ .C ₆ H ₄ .NHMe. — Ndl. w. bluish-violet reflections fr. lgr. D.s. c. aq.; e.s. alc. — ⊕ Nitro comp.†
2880	35	Or.-red	2-Nitrodimethyl-p-toluidine, NO ₂ .C ₆ H ₄ (Me)(NMe ₂). — Pr. fr. alc.; i. aq. — ⊕ Nitro comp.†
2881	36	Yellowish	Nitroso-o-nitromethylaniline, NO ₂ .C ₆ H ₄ .NMe(NO). — Nitrosamine. §
2882	36.2	YT2	Azoxybenzene, Ph.N ₂ O.Ph. — Ndl. fr. h. alc. I. aq.; v.s. eth., chlf., bz. 100 pt. sat. abs. alc. sol. at 16° contain 17.5 pt. 100 pt. lgr. (b.p. 70-80°) dis. 43.5 pt. at 15°. T. 2.15 gives no coloration. — ⊕ ⊖ Mix 0.1 g. w. 0.2 g. powdered PCl ₅ in 3-inch t.t. Heat gently over small flame until mixt. fuses to dark red liquid. Cool. Dec. x.s. PCl ₅ w. 10 cc. aq. Filter. Wash residue w. 20 cc. boiling aq. Cryst. fr. 5 cc. 66% h. alc. Wash cryst. w. 8 drops dil. alc. Recryst. fr. 4 cc. boiling 66% alc. Wash w. 8 drops dil. alc. Dry on tile 20 min. at 50°. The product, azobenzene, is obtained in small lust. YO scales, m.p. 68° u.c.
2883	37-9	Light yel.	m-Azoxytoluene, Me.C ₆ H ₄ N ₂ O.C ₆ H ₄ .Me. — Ndl. fr. eth. E.s. alc.
2884	38-9	Intense yel.	p-Diazophenol, C ₆ H ₄ .N ₂ O + 4H ₂ O (air dried). — Ndl. v.s. aq., alc.; d.s. eth., bz. Deflagrates violently at 75° when anhydrous!
2885	40	Yellow	2,4-Dinitrodipropylaniline, (NO) ₂ .C ₆ H ₄ .NPr ₂ . — ⊕ Nitro comp.†
2886	42	Emerald-green	p-Nitrosodipropylaniline, NO.C ₆ H ₄ .NPr ₂ . — Triclin. cryst. fr. lgr. — Boiling w. aq. KOH gives dipropylamine (No. 2.1118) & nitrosophenol.

* For reasons elsewhere stated the few colored liquids of this suborder are described in the tables with the colorless liquids of Suborder I.

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No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
2887	43-4	Orange	5-Nitroeugenol, $\text{NO}_2\text{C}_6\text{H}_3(\text{MeO})^2(\text{C}_6\text{H}_5)(\text{OH})^2$. — Triclin. cryst. fr. lgr., resembling $\text{K}_2\text{Cr}_2\text{O}_7$. V.d.s. aq.; e.s. alc., eth. Vol. w. st. — \textcircled{P} Nitro comp. — K salt, s. aq.; cryst. of or.-red color w. metallic luster.
2888	44	Green	Nitrosomethyldiphenylamine, $(\text{NO})\text{C}_6\text{H}_4\text{NMePh}$. — Lft. [Gives oxazine dyestuff w. diethyl-m-aminophenol (D.R.P., 75,127).]
2889	44.5; 44.2 u.c.; 46	GYT3	o-Nitrobenzaldehyde, $\text{NO}_2\text{C}_6\text{H}_4\text{CHO}$. — Long ndl. fr. h. aq. V.d.s. c.; d.s. h. aq.; v.s. eth. Odor, when cold, pungent & sl. aromatic. When hot, vapors provoke violent sneezing! Taste of c. sat. aq. sol. powerfully pungent! (No. 4+, scale of T. 2.29). \textcircled{P} Dis. 0.02 g. in 5 cc. h. aq. After cooling, add 0.5 cc. acetone, or enough to remove turbidity, and 1 drop 10% NaOH sol. Sol. becomes yel., then green, & after heating over small flame for 1 min., blue. Cool & filter. Wash w. 5 cc. aq. & dry filter at 100°. The dry ppt. shows VB color of indigo, & coppery metallic streak when rubbed w. spatula. \textcircled{P} Heat to boiling in t.l. 0.05 g. substance disd. w. 4 drops pure phenylhydrazine in 2 cc. alc. Filter hot. Add 2 cc. alc. + 3 cc. aq. to filtrate. Heat to boiling. Set aside 15 min. Cool well & shake. Wash ppt. w. 2 cc. 50% alc. Dry on tile at 100°. The resulting phenylhydrazone is obtained in red (R) ndl., m.p. 155° u.c. (This phenylhydrazone when disd. by heating w. conc. H_2SO_4 & sol. then diluted w. aq. gives brown sol. Corresponding treatment of the phenylhydrazone fr. m-nitrobenzaldehyde yields green sol.)
2890	45.2 u.c.	YT1	o-Nitrophenol, $\text{NO}_2\text{C}_6\text{H}_4\text{OH}$. — B.p. 214°. Pr. of strong phenolic odor, v. little s. c aq.; e.s. alc.; v.s. eth., bz. Taste of c. sat. aq. sol. sweet (No. 4 on scale of T. 2.29), & sl. sour. H. sat. aq. sol. efferv. sl. w. CaCO_3 . — \textcircled{P} (1) Sol. of 0.03 g. in 10 cc. 10% NaOH sol. shows YO color! — (2) Nitro comp. — \textcircled{P} Place 0.2 g. substance, 0.5 g. Zn dust, 5 cc. aq., & 2 cc. 10% CaCl_2 sol., in a 25-cc. flask fitted w. reflux tube. Boil (abt. 5 min.) until the sol. becomes colorless. Filter hot. Cool w. ice water to cause separation of cryst. fr. filtrate. Wash on filter w. 2 cc. c. aq. Partially dry on porous tile. Recryst. fr. 15 cc. boiling bz., separating cryst. by cooling & shaking. Filter. Dry on tile at abt. 70° for 15 min. The product, o-aminophenol, is obtained in sl. colorless rhombic scales, m.p. 170.8° u.c., w. sl. decn. fr. 165°.
2891	46-7	Light red	3,5,2',4'-Tetramethylazobenzene, $\text{Me}_2\text{C}_6\text{H}_3\text{N}_2\text{C}_6\text{H}_3\text{Me}_2$. — Cryst. e.s. alc.; i. aq. — \textcircled{P} Azo comp.
2892	46-7	Light red	6-Nitro-5-amino-1,2,4-trimethylbenzene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}_2$. — Ndl. fr. dil. alc. V.s. eth. — \textcircled{P} Nitro comp. ‡
2893	45-7	Yel.-red	3-Nitro-4-aminoethylbenzene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{Et}$. — \textcircled{P} Nitro comp. ‡ — Pr. fr. lgr. V.s. alc., eth., chlf., bz.; less s. lgr.
2894	45-7	Light yel.	3-Nitro-4-acetaminoethylbenzene, $(\text{NO}_2)(\text{NH.CO.Me})\text{C}_6\text{H}_4\text{Et}$. — \textcircled{P} Nitro comp. ‡ — Silky ndl. fr. lgr.; v.s. alc., eth., chlf., bz.; less s. lgr. — Sapn. w. conc. HCl gives No. 2.2892.
2895	47-8	Gold-yel.	4-Nitro-1-isopropylphenol (3), $\text{NO}_2\text{C}_6\text{H}_3(\text{C}_2\text{H}_5)(\text{OH})$. — \textcircled{P} Nitro comp. I — Ndl. fr. alc. B.p. 260-2° d. Vol. w. st. — Reduced to amino deriv., m.p. 122°, by Sn + HCl. — NaA . $2\text{H}_2\text{O}$, vermillion-red lt.
2896	48	Red-yel.	6-Nitro-1,2,4-trimethylphenol (5), Nitropseudocumeneol, $\text{NO}_2\text{C}_6\text{H}_3(\text{Me})_2(\text{OH})$. — \textcircled{P} Nitro comp. ‡ — Ndl. w. fatty luster fr. alc. Mod. s. h. aq.; v.s. alc. — B.HNO_3 , lust. cryst. fr. eth., decd. by h. aq., m.p. 84°; 81.5°.
2897	51	Or.-yel.	2,2'-Dimethyldiazoaminobenzene, $\text{Me.C}_6\text{H}_4\text{N}=\text{NH.C}_6\text{H}_4\text{Me}$. — Cryst. powd.

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No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
2898	50-2	Yellow	3,3'-Dimethylazobaminobenzene, Me.C ₆ H ₄ .N.NH.C ₆ H ₄ Me. — Ndl. fr. pet.-eth.; e.s. org. solvents.
—	50.5-1.0	Yel.-white	Nitrosoacetanilide. — Cf. No. 2.1541.
2898	52	Red	2-Methylbenzene-azo- α -naphthalene, Me.C ₆ H ₄ .N.C ₁₀ H ₇ . — Lft. fr. alc. — (P) Azo comp.
2900	52	Gold-yel.	Nitroso-2,4-dinitroethylaniline, (NO)EtN.C ₆ H ₄ .(NO ₂) ₂ . — (P) Prob. gives T. 2.15 w. diphenylamine reagt.
2901	53	Yellow	2-Nitro-m-toluidine, NO ₂ .C ₆ H ₄ .Me.NH ₂ . — (P) Nitro comp.‡ & prim. amine (T. 2.4). — D.s. c. aq.; e.s. alc.
2902	53	Green-yel.	Methyl o-Nitrocarbanilate, NO ₂ .C ₆ H ₄ .NH.CO.Me. — (P) Nitro comp.‡ — Should give methyl alc. as a sapon. product (T. 2.26).
2903	54-5	Or.-red	3,3'-Azotoluene, Me.C ₆ H ₄ .N ₂ .C ₆ H ₄ .Me. — (P) Azo comp. — Cryst. I. aq.; e.s. alc., eth.
2904	54	Yel.-brown	Nitroso-p-methyltoluidine, (NO)MeN.C ₆ H ₄ .Me. — (P) Nitrosamine. § — Pr. fr. eth.-alc. I. aq.; e.s. alc., eth.
2905	54	Yellow	4-Nitro-5-amino-1,3-xylene, (NO ₂)(NH ₂).C ₆ H ₄ .Me ₂ . — (P) Nitro comp.‡ & prim. amine (T. 2.4). — Ndl. E. vol. w. st.
2905	55	Red	2,2'-Dimethylazobenzene, o-Azotoluene, Me.C ₆ H ₄ .N ₂ .C ₆ H ₄ .Me. — (P) Azo comp. — Pr. fr. eth. E. vol. w. st. I. aq.; 100 pt. alc. at 14.5° dis. 6.027 pt.; 100 pt. eth. at 16.5° dis. 147.7 pt.
2907	55	Yellow	Nitromethyl-p-toluidinenitrosamine, (NO ₂).Me.C ₆ H ₄ .NMe-(NO). — (P) Nitrosamine. § — Ndl. fr. alc. E.s. eth., chlf., warm alc.; d.s. lgr.
2908	55	Yellow	2,6-Dinitrothymol, (NO ₂) ₂ .C ₆ H ₄ .Me ¹ ,(C ₆ H ₅) ⁴ (OH) ² . — (P) Nitro comp.‡ — Cryst. V.d.s. aq.; v.s. alc., eth. — KÄ, yel. ndl. d.s. aq. — (P) Gently heated w. nitro-sulphuric ac. gives trinitro deriv., yel. ndl. fr. aq., m.p. 111°.
2909	55; 56-8	Brown-red	3,4-Dimethylazobenzene, Me.C ₆ H ₄ .N ₂ .C ₆ H ₄ .Me. — (P) Azo comp. — Lit. fr. dil. alc. E.s. alc.
2910	56-7	Orange	m-Aminoazobenzene, NH ₂ .C ₆ H ₄ .N ₂ .Ph. — (P) Prim. amine (T. 2.4). — Silky ndl. fr. lgr. E.s. alc., eth., chlf., bz. — (P) Reduction T. products: aniline (No. 2.1235); m-phenylenediamine (No. 2.634).
2911	56; 51-2	Yellow	4-Nitrocresol(3), NO ₂ .C ₆ H ₃ .Me(OH). — (P) Nitro comp.‡ S. in alk. — Monoclin. cryst. fr. bz. D.s. aq.; e.s. bz., alc., eth. — K salt, red lft., e.s. aq.
2912	57	Red	2-Nitromethyl-p-toluidine, NO ₂ .C ₆ H ₄ .Me ¹ ,(NHMe) ⁴ . — (P) Nitro comp.‡ — Cryst. fr. alc. — NaNO ₂ & HCl give nitrosamine (T. 2.36), yel. ndl. fr. alc., m.p. 55°. (Ber., 28, 3039.)
2913	57	Brown	Phenyl-o-nitrobenzylamine, NO ₂ .C ₆ H ₄ .CH ₂ .NHPH. — (P) Nitro comp.‡ — Lust. tbl. fr. alc. (Sometimes, lust. or. ndl. of m.p. 44°, which become opaque on rubbing & change to 57° m.p. form.) — E.s. alc., eth., chlf., bz.; d.s. lgr.
2914	57.5	Yellow	1,2,4-Trinitrobenzene, (NO ₂) ₃ .C ₆ H ₃ . — (P) Nitro comp.‡ — Cryst. d.s. aq.
2915	58	Carmine-red	5-Nitro-4-methylamino-1,3-xylene, (NO ₂)(NHMe).C ₆ H ₄ .Me ₂ . — (P) Nitro comp.‡ — Plates showing greenish reflections. R.d.s. c. alc., lgr.
—	58	Yellowish	Phenylnitroethylene. — Cf. No. 2.1586. (Odor irritating & cinnamon-like!)
2916	58	Yellowish	Nitrosophenylbenzylamine, (NO)PhN.CH ₂ .Ph. — (P) Prob. gives T. 2.15 w. diphenylamine reagt. — Ndl. e.s. alc., eth., chlf., lgr.
2917	58u.c.	GYT2	α -Nitronaphthalene, C ₁₀ H ₇ .NO ₂ . — B.p. 304°. Long fine pr. fr. alc. I. aq. 100 pt. 88% alc. dis. 2.8 pt.; e.s. eth. — (P) Nitro

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
2918	58	(GY-Y)T3	<i>comp.† — ① Reduce by heating & shaking w. x.s. Sn & conc. HCl, as in preparation of aniline fr. nitrobenzene, & identify the α-naphthylamine (No. 2.589) produced.</i> m-Nitrobenzaldehyde, $\text{NO}_2\text{C}_6\text{H}_4\text{CHO}$. — ② Odor, faintly bitter almond-like, sl. pungent, provoking sneezing. — Ndl. Alm. i. c. aq.; d.s. h. aq.; s. alc.; v.s. eth. — Taste of sat. aq. sol. bitter & burning (abt. No. 2 on scale of bitterness & pungency, T. 2.29). ③ Convert into <i>m-nitrobenzaldehydophenylhydrazone</i> (No. 2.3164), employing the procedure described for preparing corresponding <i>ortho</i> deriv. under (No. 2.2889).
2919	59-60	Red-yel.	m-Nitroethylaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NHET}$. — ② Nitro comp.‡ — Ndl. e.s. alc., eth., lgr. Vol. w. st.
2920	59	Dark yel.	Propylpicramide, $\text{PrNH.C}_6\text{H}_5\text{(NO}_2)_2\text{C}_6\text{H}_4\text{Me}$. — Ndl.
2921	59-60	Yellowish	o-Azoxytoluene, $\text{Me.C}_6\text{H}_4\text{N.O.C}_6\text{H}_4\text{Me}$. — ② Azoxy comp. — Monoclin. tbl. fr. lgr. — Heated w. conc. H_2SO_4 at 116° gives much o-azo-toluene (No. 2.2906). — Reduction w. Sn & HCl should give o-toluidine (No. 2.1262).
2922	59	Steel-blue	p-Nitrosopropylaniline, $\text{NO.C}_6\text{H}_4\text{NHPr}$. — Ndl. fr. alc. E.s. alc., eth., bz., w. green color! — Boiled w. NaOH sol. yields propylamine (No. 2.1067), & p-nitrosophenol (No. 2.3191).
2923	60	Carmine-red	Anethole Picrate, $\text{C}_{10}\text{H}_{12}\text{O.C}_6\text{H}_4\text{O}_2\text{N}_3$. — Picrate of anethole (Cf. Vol. I). — Ndl. fr. alc.
2924	60-1	Red	m-Nitrodimethylaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NMe}_2$. — ② Nitro comp.‡ — B.p. 280-5° d. Pr. fr. eth. Vol. w. st.
2925	61-2	Yellow	Dinitro-tert.-butylbenzene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{CMe}_3$. — ② Nitro comp.‡ — Pr. e.s. alc., eth.
2926	61.8c.	Light yellow	2,6-Dinitrophenol, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{OH}$. — Nitro comp.‡ — V.d.s. c. aq.; e.s. h. aq.; v.s. bz., chlf., eth., h. alc. — Nitration w. x.s. conc. HNO_3 gives picric ac. (No. 2.3168). — $\text{BaAl}_2\text{H}_2\text{O}$, gold-yel. ndl., s. in 555 pt. c. aq.; alm. i. h. 90% alc. — MeA, m.p. 118°; 116°.
2927	62	Yellow	Trinitro-m-diethylbenzene, $(\text{NO}_2)_3\text{C}_6\text{H}_2\text{Et}_2$. — ② Nitro comp. — Short pr. fr. lgr.
2928	63..	Or.-yel.	4-Nitro-2-aminodimethylaniline, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{NMe}_2$. — Nitro comp.‡ & prim. amine (T. 2.4). — Ndl. fr. alc. E.s. alc., eth.; v.s. chlf., bz.
2928	63	Lemon-yellow	Propion-o-nitroaniline, $(\text{Et.CO})\text{NH.C}_6\text{H}_4\text{NO}_2$. — ② Nitro comp.‡ — Cryst. — Sapn. T. products (T. 2.26): o-nitroaniline (No. 2.2945), & propionic ac. (Vol. I).
2930	64	Yellow	Phenylnitropropylene, $\text{Ph.CH:C}(\text{NO}_2)\text{Me}$. — ② Odor like nutmegs. — Lust. ndl. fr. lgr. Vol. w. st. — Adds Br_2 giving dibromide, pr. fr. lgr., m.p. 78°.
2931	64	Lemon-yellow	2,4,6-Trinitroanisole, Methyl Picrate, $(\text{NO}_2)_3\text{C}_6\text{H}_2\text{OMe}$. — ② Nitro comp.‡ — Monoclin. cryst. I. aq.; s. alc., eth.; e.s. acetone. — Boiled w. conc. KOH sol. gives K picrate (No. 2.3168) & methyl alc.
2932	65-6	Scarlet	3-Nitro-4-amino-1,2-xylene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}_2$. — ② Nitro comp.‡ & prim. amine (T. 2.4). — Vol. w. st. — Acetyl deriv., ** white ndl. m.p. 115-6°.
2933	65-6	Red-yel.	m-Nitromethylaniline, $(\text{NO}_2)\text{C}_6\text{H}_4\text{NHMe}$. — ② Nitro comp.‡ — Ndl. E.s. alc., eth.
2934	66.5	Pale yel.	Nitrosodiphenylamine, Ph.N.NO. — ② Dis. in conc. H_2SO_4 w. the intense blue color of T. 2.15! — 4-sided tbl. fr. bz. + alc. D.s. c. alc.; e.s. h. alc., bz.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. ‡ = T. 2.36. || = T. 2.34.
¶ = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
2935	68 u.c.	O or YO	Azobenzene, Ph.N:N.Ph.—Color of large cryst., O; of powder or small cryst., YO. Plates fr. 90% alc. B.p. 295–7° c. (749 mm.). 100 pt. sat. alc. sol. contain 8.5 pt. at 16°. — (D) Gives silver reduction in T. 2.21. — (D) Dis. 0.3 g. in 15 cc. 66% alc. w. 0.3 g. KOH. Add 3 g. Zn dust & boil 5 min. under reflux (or until color is discharged). Filter hot. Wash colorless scales of hydrazobenzene (No. 2.2050), which separate on cooling, w. 5 cc. c. 50% alc. Boil moist product 1 min. w. conc. HCl. Shake w. 10 cc. c. aq. & filter if not clear. Add 4 drops conc. H ₂ SO ₄ to sol. After standing for a min. & shaking, shimmering silvery scales of benzidine (No. 2.840), will be seen to separate.
2936	69.5	Yellow	3-Nitro-o-cresol, NO ₂ .C ₆ H ₄ .Me.(OH). — (D) Nitro comp.‡ — Pr. fr. dil. alc. — I. aq.; v.s. alc., eth.
2937	68–70	Yellow	Nitroso-m-nitromethylaniline, NO ₂ .C ₆ H ₄ .NMe(NO). — (D) Nitrosamine. § — Cryst. e.s. alc.
2938	69.5–70.5; 71–2	Or.-red	p-Benzeneazotoluene, Ph.N ₂ .C ₆ H ₄ .Me. — (D) Azo comp. — Lft. fr. alc. I. aq.; e.s. eth., chlf., lgr. — Reduced by Zn & HCl to aniline & p-toluidine (Nos. 2.1235 & 2.566).
2939	70	Pale yel.	p-(α)-Azoxytoluene, Me.C ₆ H ₄ .N.O.C ₆ H ₄ .Me. — (D) Dec. on distn. to p-azotoluene (No. 2.3303), & p-toluidine (No. 2.566). — Ndl. fr. alc. E.s. alc., eth. — Reduction w. Sn & HCl gives p-toluidine!
2940	70	Gold-yel.	3,4-Dinitroanisole, (NO ₂) ₂ .C ₆ H ₄ .OMe. — (D) Nitro comp.‡ — Fine ndl.
2941	70	Yellow	β -Ethynaphthalene Picrate, C ₁₁ H ₁₂ .Pk. — (D) (D) Picrate of hydrocarbon of b.p. 251° (Vol. I). — Fine ndl.
2942	70	Yellow	Hexylpicramide, (C ₆ H ₅)NH.C ₆ H ₅ .(NO ₂) ₂ .
2943	70.5–1.5	Red	4-Hydroxy-3-methoxyazobenzene, (HO)(MeO).C ₆ H ₄ .N ₂ .Ph. — (D) Azo comp. — Lust. pr. fr. lgr. E.s. alc.; v.s. eth., bz.; d.s. c. lgr.
2944	71	Or.-red	2,4'-Azotoluene, Me.C ₆ H ₄ .N ₂ .C ₆ H ₄ .Me. — (D) Azo comp. — Ndl. fr. alc. I. aq. — Reduction w. Sn & HCl should give o- & p-toluidine (Nos. 2.1262 & 2.566).
2945	71.4 u.c.	RO	o-Nitroaniline, NO ₂ .C ₆ H ₄ .NH ₂ . — (D) Nitro comp.‡ & prim. amine (T. 2.4). — Ndl. fr. alc. V.d.s. aq.; e.s. alc., eth. — (D) Mix 0.1 g. substance, 0.4 cc. ac. anhydride, & 1 drop conc. H ₂ SO ₄ . Heat to boiling. Cool & shake. Filter off ppt. Wash w. 1 cc. c. aq. Recryst., first fr. 5 cc., & then fr. 3 cc. h. aq. Dry on tile at 60°. The product, o-nitroacetanilide, is obtained in v. pale yel. ndl., m.p. 91–1.5° u.c.
2946	72	Orange	p-Nitrobenzylaniline, Phenylnitrobenzylamine, NO ₂ .C ₆ H ₄ .CH ₂ -NHPh. — (D) Nitro comp.‡ — Pr. fr. eth.; d.s. lgr.; e.s. h. alc., eth., bz. — HNO ₂ gives straw-yel. nitrosamine, m.p. 76°.
2947	72–3	Wine-yel.	Furfural-p-phenetidine, EtO.C ₆ H ₄ .N:CH.C ₆ H ₅ O. — [D.R.P., 96,658.] (D) Alc. sol. w. FeCl ₃ becomes blood-red w. green surface reflections. — Tbl. fr. eth. Alm. i. aq.; otherwise e.s.
2948	72; 68–9	Yellow	2,4-Dinitrobenzaldehyde, (NO ₂) ₂ .C ₆ H ₄ .CHO. — (D) Reduces ammon. AgNO ₃ or Fehling's sol. — Pr. fr. alc. w. 1 mol. alc. (lost at 80–90°). D.s. aq.; v.s. alc., eth., bz.; d.s. pet.-eth. — Decd. by alk. w. brown color. Oxidation gives 2,4-dinitrobenzoic ac. (No. 2.274).
2949	74; 71	Straw-yel.	p-Nitrodiazobenzeneimide, NO ₂ .C ₆ H ₄ .N ₂ . — Lust. lft. fr. lgr. E.s. h. alc., eth. — Boiled w. alc. KOH gives NH ₃ , p-nitroaniline (No. 2.3319) & p-azophenetole. — (D) Reduction by Sn & HCl gives p-phenylenediamine (No. 2.877).
2950	72	Yellow	5-Nitro-1,3-xylenol(4), (NO ₂).C ₆ H ₄ .Me ₂ (OH). — (D) Nitro comp.‡ — Ndl. — K ₂ 3H ₂ O, dark red lft.
2951	72–3	Yellow	3,4,6-Trinitrophenol-naphthalene, (NO ₂) ₃ .C ₆ H ₄ .(OH).C ₁₀ H ₈ . — (D) Nitro comp.‡ — Ndl.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
2952	73-4	Gold-yel.	Nitromesidine, Nitro-2-amino-1,3,5-trimethylbenzene, $(NO_2)_2.C_6H_3.Me_3$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Ndl. E.s. alc., eth. — Acetyl deriv.,** m.p. abt. 190°.
2953	73-5	Straw-yel.	α -Nitrophenanthrene, $NO_2.C_14H_8$. — \textcircled{P} Nitro comp.‡ — Oxidation by CrO_3 in gl. ac. ac. gives α -nitrophenanthrenequinone, or.-yel. lft., d.s. alc., m.p. 215-20°.
2954	74-5	Deep yel.	α -Phentriazine, $C_7H_4N_3$. — B.p. 235-40°. Lust. ndl. fr. bz. E. vol. w. st. E.s. h. aq., warm eth., alc.
2955	74-5	Orange	6-Nitro-4-amino-1,2-xylene, $(NO_2)(NH_2).C_8H_6.Me_2$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Lft. fr. alc. — Acetyl deriv.,** white, m.p. 209-10°.
2956	75	Red	Azomesitylene, 2,4,6,2',4',6'-Hexamethylazobenzene, $Me_2.C_6H_2.N_2.C_6H_2.Me_2$. — \textcircled{P} Azo comp. — Ndl. fr. alc. R.d.s. c. alc.
2957	74-6	Red-yel.	α -Nitrophenylbenzylamine, $NO_2.C_6H_4.NH.CH_2.Ph$. — \textcircled{P} Nitro comp.‡ — Pr. e.s. alc.
2958	75	Or.-yel.	p-(β)-Azoxytoluene, $Me.C_6H_4.N.O.C_6H_4.Me$. — Pr. fr. lgr. — \textcircled{P} Reduction by Sn & HCl gives p-toluidine (No. 2.586).
2959	75.5	Gold-yel.	α -o-Hydroxyazoxybenzene, $HO.C_6H_4.N.O.Ph$. — \textcircled{P} S. in NaOH sol. w. or.-red color. — Conc. H_2SO_4 gives o-hydroxy-azobenzene. — Reduction gives aniline & o-aminophenol (Nos. 2.1235 & 2.938).
2960	75	Yel.-red	α -Nitrodiphenylamine, $NO_2.C_6H_4.NHPH$. — Nitro comp.‡ — Trimet. tbl.
2961	76	Or.-red	5-Nitro-4-amino-1,3-xylene, $(NO_2)(NH_2).C_8H_6.Me_2$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Ndl. fr. alc. Vol. w. st. — Acetyl deriv.,** yellowish ndl. fr. h. aq., m.p. 172-3°.
2962	76	Light yel.	α -Nitrobenzalhydrazine, $NO_2.C_6H_4.CH:N.NH_2$. — \textcircled{P} Sol. colored intense red by HgO . — Pr. fr. alc. V.e.s. — Boiled w. dil. alc. gives o,o'-dinitrobenzalzin, m.p. 182° (Ber., 33, 2463).
2963	76-7	Pale yel.	α -Nitrophenylnitropropylene, $NO_2.C_6H_4.CH:C(NO_2).Me$. — \textcircled{P} Nitro comp.‡ — Lft. I. c. aq. — Oxidized by $KMnO_4$ to o-nitrobenzoic ac. (No. 2.164).
2964	77	Brown-red	(α)-Dinitrophenyl- α -naphthylamine, $(NO_2)_2.C_10H_7.NH_2$. — \textcircled{P} Nitro comp.‡ E.s. alk. w. orange color. — D.s. org. solvents.
2965	77	Yellow	2-Nitrocresol(4), $NO_2.C_6H_5.Me(OH)$. — \textcircled{P} Nitro comp.‡ prob. giving yel. sol. in alk. — Pr. fr. c. eth. V.s. alc., eth.; d.s. c. aq., lgr.; e.s. bz.
2965	77.5	Yellow	2-Nitro-p-toluidine, $NO_2.C_6H_5.Me(NH_2)$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Broad monoclin. ndl. fr. aq. D.s. CS ₂ .
2967	77-8	Sulphur-yellow	p-Nitrodiethylaniline, $NO_2.C_6H_4.NEt_2$. — \textcircled{P} Nitro comp.‡ — Monoclin. ndl. E.s. h. alc. D.s. lgr.
2968	78	Or.-yel.	2-Amino-3,5,2',4'-tetramethylazobenzene, $(NH_2)Me_2.C_6H_3.N_2.C_6H_2.Me_2$. — \textcircled{P} Azo comp. & prim. amine. — Lft. fr. alc. or bz. D.s. c. alc.; e.s. bz. — Reduction by Sn & HCl gives 1,3,4-xylidine (No. 2.1308), & 4,5-diamino-m-xylene (No. 2.676).
2969	78	Green	p-Nitrosoethylaniline, $NO.C_6H_4.NHEt$. — Lft. fr. bz. E.s. alc., eth. — \textcircled{P} Boiled w. NaOH sol. evolves ethylamine (No. 2.1062).
2970	79-80	Red-brown	Fluorene Picrate, $C_{10}H_{10}.Pk$. — \textcircled{P} \textcircled{P} Picrate of h.c., m.p. 109-10° of Vol. I. Pr. Deed by boiling w. aq. or alc.
2971	79	Yellow	β -Nitronaphthalene, $NO_2.C_{10}H_7$. — \textcircled{P} Nitro comp.‡ of cinnamon-like odor. — Small ndl. fr. dil. ammon. Vol. w. st. E.s. alc., eth., chlf.

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(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
2972	79-81	Blood-red	γ -Methylenebiphenyl Picrate, $C_{12}H_{10} \cdot Pk$. — \textcircled{P} Picrate of h.c., m.p. 116° (Vol. I).
2973	80	Yellow	Allylpicramide, $(C_2H_5)NH \cdot C_6H_5 \cdot (NO_2)_2$.
2974	80	Yellow	2,4-Dinitrodiethylaniline, $(NO_2)_2 \cdot C_6H_4 \cdot NEt_2$. — \textcircled{P} Nitro comp.‡ — Long ndl. V.s. h. alc.; d.s. lgr.
2975	80	Yellow	Isobutyl-2,4-dinitroaniline, $(Me_2CH \cdot CH_2)NH \cdot C_6H_4 \cdot (NO_2)_2$. — \textcircled{P} Nitro comp.‡ Ndl.
2976	78; 81-2	Gold-yel.	2-Nitro-4-amino-1,3-xylene, $(NO_2)(NH_2) \cdot C_6H_4 \cdot Me_2$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Ndl. s. alc. — Acetyl deriv.,** m.p. 149°.
2977	81-2	Light red	4-Nitro-2-ethyltoluidine, $(NO_2)Me \cdot C_6H_4 \cdot NHET$. — \textcircled{P} Nitro comp.‡ — 1 pt. Br. added to 1 pt. comp. in 11 pt. gl. ac. ac. gives Br. deriv., ndl. fr. alc., m.p. 170°.
2978	81	Or.-yel.	α -Dihydroxyazoxybenzene-dimethylether, $MeO \cdot C_6H_4 \cdot N_2O \cdot C_6H_4 \cdot OMe$. — Pr. fr. methyl alc. I. aq.; e.s. alc., eth., bz.
2979	81-2	Sulphur-yellow	4-Nitro-2-amino-1,3-xylene, $(NO_2)(NH_2) \cdot C_6H_4 \cdot Me_2$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Ndl. fr. dil. alc. — Acetyl deriv.,** m.p. 170°.
2980	82	Gold-yel.	Nitrosocarbazole, $C_{12}H_8 \cdot N \cdot NO$. — \textcircled{P} Nitrosamine.§ — Long flat lust. ndl. fr. alc. E.s. alc., bz., gl. ac. ac., chlf., eth. — \textcircled{P} Boiled w. alc. & ac. gives carbazole (No. 2.2584!).
2981	82	Gold-yel.	Phenylaminoazobenzene, $Ph \cdot NH \cdot C_6H_4 \cdot N_2 \cdot Ph$. — \textcircled{P} Alc. sol. colors yel. w. ac., salt afterwards separating in gray cryst. — Lft. s. alc., eth., lgr.
2982	82	Pale yel.	4,6-Dinitro-1,2-xylanol(3), $(NO_2)_2 \cdot C_6H_4 \cdot Me_2(OH)$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc.
2983	83-4	Red	α -Trinitrotoluene-aniline, $[(NO_2)_3 \cdot C_6H_3 \cdot Me] \cdot PhNH$.
2984	82.5-3	Or.-red	2-Hydroxyazobenzene, $HO \cdot C_6H_4 \cdot N_2 \cdot Ph$. — \textcircled{P} Sol. in alk. or.-red. — Ndl. w. bluish reflections fr. eth. — Vol. w. st. D.s. aq.; e.s. org. solvents.
2985	83	Yellow	Quinoneoxime-methylether, $O \cdot C_6H_4 \cdot NOMe$. — Flat ndl. fr. lgr. V. vol. w. st. V.s. alc. Odor ethereal.
2986	84-5	Red	3-Nitromethyl-p-toluidine, $(NO_2)Me \cdot C_6H_4 \cdot NHMe$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. I. aq.
2987	84	Yellow	Nitrodimethyl-m-toluidine, $(NO_2)Me \cdot C_6H_4 \cdot NMe_2$. — \textcircled{P} Nitro comp.‡ — Ndl. I. aq.
2988	84.3; 85-90	Light yellow	Benzene Picrate, $C_6H_6 \cdot Pk$. — \textcircled{P} Picrate* of h.c. of Vol. I. — Rhomb. cryst. soon losing benzene in air at ordinary temperature.
2989	84u.c.	Dark green	p-Nitrosodiethylaniline, $NO \cdot C_6H_4 \cdot NEt_2$. — Ndl. w. bluish reflections & metallic lustre! D.s. aq.; e.s. alc., eth. — \textcircled{P} Treat 0.1 g. w. 2.5 cc. HNO_3 (sp. gr. 1.42) + 2.5 cc. aq. After 30 sec. add 15 cc. c. aq. & shake vigorously. (Too long continued nitration gives oily product.) Filter. Wash ppt. w. 3 cc. aq. Recryst., first fr. 4 cc., & then fr. 2 cc. h. 50% alc. Dry on tile 15 min. at 60°. The product, dinitrodiethylaniline, is obtained in OY ndl., m.p. 79.6-80.2° u.c.
2990	85	Or.-red	2-Nitroresorcinol, $NO_2 \cdot C_6H_3 \cdot (OH)_2$. — \textcircled{P} Nitro comp.‡ — Pr. fr. dil. alc. Dist. Strong odor like o-nitrophenol!
2991	85	Yellowish	Nitroso-2,4-dinitromethylaniline, $(NO)MeN \cdot C_6H_4 \cdot (NO_2)_2$. — \textcircled{P} Nitrosamine.§ — Ndl. fr. alc. Unstable.
2993	85-6	Yellow	Trinitro-5-pseudobutylcresol(2), $(NO_2)_3 \cdot (Me_2C) \cdot Me \cdot C_6 \cdot OH$. — Ndl. fr. alc.
2994	85	Gold-yel.	(β)-Nitromethaneazobenzene, Nitroformaldehydophenylhydrazone, $NO_2 \cdot CH \cdot N \cdot NH \cdot Ph$. — \textcircled{P} S. in NaOH sol. w. red color. — Lust. ndl. fr. bz. + lgr. D.s. lgr.; e.s. alc.

(ORDER II.)

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
2995	85	Yellow	Nitrophenyl-β-naphthylamine, $\text{NO}_2\text{C}_{14}\text{H}_{12}\text{N}$. — (P) Nitro comp.‡ — Cryst. mass fr. dil. alc.
2996	85 u.c.	Green	p-Nitrosodimethylaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NMe}_2$. — Scales w. yellowish reflections. D.s. aq., giving deep brown sol. which stains skin bright yel. — (P) Dis. 0.05 g. in 1 cc. alc. Add 2.5 cc. HNO_3 (sp. gr. 1.42) + 2.5 cc. aq. If a ppt. forms, shake to redissolve. After 2 min. dilute w. 10 cc. c. aq. Shake vigorously. Filter. Wash w. 3 cc. c. aq. Recrys. fr. 5 cc. h. 50% alc., filtering h. if not clear. Cool. Filter. Dry cryst. on tile 15 min. at 60°. Dinitrodimethylaniline, the product, is obtained in fine OY ndl., m.p. 86.3-9° u.c.
2997	86	O-YO	Guaiacol Picrate, [$\text{o-MeO.C}_6\text{H}_4\text{OH}$] ₂ Pk. — (P) Picrate* of species of Vol. I. — Cryst. ppt.
2998	86	Yellow	Heptadecylpicramide, $(\text{C}_{17}\text{H}_{35})\text{NH.C}_6\text{H}_5\text{(NO}_2)_2$. — Tbl. D.s. alc.
2999	86	Yellow	3,5-Dinitrocresol(2), $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Me(OH)}$. — (P) Nitro comp.‡ dyeing (T. 2.16-a) deep yel. — Pr. fr. alc. Somewhat vol. w. st. D.s. aq., lgr.; s. in 13 pt. alc. at 15°. — $\text{KA.xH}_2\text{O}$, long lust. yel. ndl. fr. aq.
3000	86	Yellow	Nitrodiphenylethylene, $\text{NO}_2\text{C}_6\text{H}_4\text{CPh:CH}_2$. — (P) Nitro comp.‡ — Cryst. fr. eth.
3001	86-7	Yellow	Dinitrophenetole, 1-Ethoxy-2,4-dinitrobenzene, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{OEt}$. — (P) Nitro comp.‡ — Ndl. S. in 64 pt. 90% alc. at 21°.
3002	86-7	Yellow	2,4,6-Trinitro-1-methyl-3,5-diethylbenzene, $(\text{NO}_2)_3\text{C}_6\text{H}_3\text{MeEt}_2$. — (P) Nitro comp.‡ Lft. fr. alc.
3003	86	Yellow	3-Nitropyrocatechol, $\text{NO}_2\text{C}_6\text{H}_3\text{(OH)}_2$. — (P) Nitro comp.‡ colored purple-red by alk. — Ndl. fr. dil. alc. Mod. s. aq.
3004	87	Yellow	2,4-Dinitrodimethylaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NMe}_2$. — (P) Boiled w. KOH sol. evolves dimethylamine (No. 2.1061), & gives 2,4-dinitrophenol (No. 2.3126). — Pr. fr. CS ₂ . D.s. h. aq., eth.; e.s. chlf., less s. eth. R.d.s. alc.
3005	85-7	Lemon-yellow	α-Isoamylnaphthalene Picrate, $\text{C}_{15}\text{H}_{15}$.Pk. — (P) Picrate* of h.c., b.p. 288-92°.
3005	88-9	Or.-red	m-Tetramethyldiaminoazoxybenzene, $\text{NMe}_2\text{C}_6\text{H}_4\text{N.O.C}_6\text{H}_4\text{-NMe}_2$. — Cryst. fr. alc.
3007	88	Or.-yel.	o-Cresol Picrate, [$\text{Me.C}_6\text{H}_4\text{OH}$] ₂ Pk. — (P) Picrate* of species of Vol. I. — Ndl.
3008	88-9	Yellowish	2,4,5-Trinitrotoluene-naphthalene, $[(\text{NO}_2)_3\text{C}_6\text{H}_3\text{Me}]\text{C}_{10}\text{H}_8$.
3009	88	Pale yellow	Dinitroanisole, 1-Methoxy-2,4-dinitrobenzene, $\text{MeO.C}_6\text{H}_3\text{-(NO}_2)_2$. — (P) Nitro comp.‡ — Flat ndl. S. in 64.2 pt. 95% alc. at 21°. — Boiled w. alc. KOH gives NH ₃ (T. 2.6). — W. ammon. (sp. gr. 0.93) at 200° gives 2,4-dinitroaniline (No. 2.3539-1).
3010	88-9	Light yel.	Ethylacet-m-nitroanilide, $\text{Et}(\text{C}_2\text{H}_5\text{O})\text{N.C}_6\text{H}_4\text{NO}_2$. — (P) Nitro comp.‡
3011	89	Sulphur-yellow	4-Nitro-2-amino-1,3-dimethyl-5-tert.-butylbenzene, $(\text{NO}_2)_2\text{(NH}_2\text{)(Me}_2\text{)}(\text{C}_6\text{H}_3)\text{C}_6\text{H}_5$. — (P) Nitro comp.‡ & prim. amine.¶
3012	89-90	Lemon-yellow	β-Propynaphthalene, Picrate, $\text{C}_{11}\text{H}_{14}$.Pk. — (P) (P) Picrate* of h.c. of Vol. I.
3013	90	Brick-red	o-Nitrophenylhydrazine, $\text{NO}_2\text{C}_6\text{H}_4\text{NH.NH}_2$. — (P) Prob. reduces Tollen's reagt. directly in T. 2.30. — Silky ndl. fr. bz. E.s. h. aq.; d.s. c. alc., eth., bz. — Warming w. 25% NaOH & pptn. w. conc. HCl gives azimidol, m.p. 157° (Ber., 27, 3381).

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 1 = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
3014	90.5	Or.-yel.	3-Nitrohydrocumaric Ac., $(NO_2)_2C_6H_3(OH)^2\cdot(C_6H_4CO_2H)^1$. — \textcircled{D} Strong acid. — Ndl. fr. aq. D.s. h. aq.; v.s. c. alc. — MeA, yel. ndl., m.p. 64°.
3015	90-1	Yellow	4-Methylidazoaminobenzene, $Me.C_6H_4.N_2NHPh$. — Lft. Boiled w. dil. H_2SO_4 gives aniline, p-toluidine, phenol, & p-cresol!
3015-I	90-1	Light yellow	5-Nitrocresol(3), $NO_2C_6H_3Me(OH)$. — \textcircled{D} Nitro comp.‡ — Cryst. w. $1H_2O$ (lost at 100°), m.p. 60-1°. V.s. alc., eth.; less s. bz.
3016	90.0 u.c.	(GY-Y)T3	m-Dinitrobenzene, $(NO_2)_2C_6H_4$. — B.p. 297° c. Thin rhomb. bl. or fine ndl. fr. h. dil. alc. Odorless, cold. On heating, vapors have st. pungent aromatic odor & bitter taste. Taste of sat. aq. sol. v. bitter (No. 2-3, scale of T. 2.29). S. in 10,000 pt. c. aq.; more s. h. aq.; s. in c. 96% alc.; v.s. h. alc.; e.s. c. eth.; v.s. bz. \textcircled{D} Dis. 0.1 g. in 10 cc. 96% alc. Add 1 cc. ammon. (sp. gr. 0.90). Pass H_2S into mixt. heated nearly to boiling in t.t. for 5 min. Dist. in small flask until reduced to 1 cc. Acidify w. HCl. Add 1 cc. aq. Filter & make filtrate alk. w. conc. Na_2CO_3 sol. Filter, & wash ppt. w. c. aq. Dis. in 2 cc. boiling aq. Filter hot. Cool & shake persistently. Filter. Recryst. fr. 2 cc. h. aq., washing w. not more than 2 cc. c. aq. Dry on tile at 100° for 10 min. The product, m-nitroaniline, is obtained crystg. in OY ndl., m.p. abt. 115° u.c.
3017	91	Leather-yellow	6-Nitro-1,4-xylenol(2), $NO_2C_6H_3Me_2(OH)$. — \textcircled{D} Nitro comp.‡ Lft. fr. lgr.
3018	91.5	Pale yellow	6-Nitro-o-toluidine, $(NO_2)(NH_2)_2C_6H_4Me$. — \textcircled{D} Nitro comp.‡ & prim. amine. — Ndl. S. in abt. 75 pt. h. aq.; e.s. alc., eth., bz. Tasteless.
3019	91-2	Steel-blue	p-Nitrosophenylbibenzylaniline, $NO_2C_6H_4N(CH_3)_2Ph$. — Thin plates fr. CS ₂ . E.s. CS ₂ , eth.; less s. alc.
3020	92	Pale yel.-red	Nitrosophenyl- α -naphthylamine, $Ph.N(NO).C_{10}H_7$. — \textcircled{D} Nitrosamine.§ — Isomerized by Fischer & Hepp react. (Ber., 29, 1247) to comp. of m.p. 150°.
3021	92 u.c.	V. pale yellow	Acet-o-nitroanilide, $(C_6H_5O.NH)_2C_6H_4NO_2$. — \textcircled{D} Nitro comp. — Lft. or ndl. fr. h. aq. V.s. alc., chlf.; e.s. in aq. KOH w. yel. color! — \textcircled{D} E. saponified by KOH sol. (T. 2.26) to o-nitroaniline (No. 2.2945), & ac. ac!
3022	92	Green	Nitroso-m-dimethyltoluidine, $NO_2C_6H_3Me(NMe_2)$. — \textcircled{D} Sol. in eth. or bz. intensely green. — Does not give the Liebermann react. (T. 2.18). — $\textcircled{B}HCl$, yel. ndl., d.s. c. aq. — Aniline deriv., deep steel-blue.
3023	93-4	Brown	Benzeneazopseudocumenol, $Ph.N_2C_6H_4Me_2(OH)$. — \textcircled{D} Azo comp. — Lust. pr. fr. alc. Dist. alm. undecd. — Acetate, red-yel. pr. fr. alc., m.p. 73-4°.
3024	93	Yellow	Nitrosophenyl- β -naphthylamine, $Ph.N(NO).C_{10}H_7$. — \textcircled{D} Nitrosamine.§ — Pr. fr. bz. D.s. h. alc.; e.s. bz.
3025	93	Canary-yellow	m-Nitrophenylhydrazine, $NO_2C_6H_4.NH.NH_2$. — \textcircled{D} Should reduce Tollen's reagt. directly (T. 2.30). — Ndl. fr. alc. D.s. h. aq., c. alc. or bz. — \textcircled{D} Prepare the corresponding hydrazone of benzaldehyde (No. 2.2220), (Ber., 22, 2813).
3026	93	Light yellow	Benzalzin, $C_{14}H_{12}N_2$. — \textcircled{D} Boiled w. dil. ac. gives hydrazine & benzaldehyde (cf. T. 2.17). — Lust. pr. I. c. aq.; mod. s. h. alc., eth., chlf., bz. — Dec. to N ₂ & stilbene (Vol. I) when boiled.
3027	93-4	YT1→	Benzoyl-o-nitroanilide, $(Ph.CO)NH_2C_6H_4NO_2$. — \textcircled{D} Nitro comp.‡ giving heavy black ppt. in T. 2.21. — Silky hair-like ndl. fr. h. 50% alc. Alm. i. c. aq.; v.d.s. h. aq.; e.s. alc.; v.s. eth. Odorless. — \textcircled{D} Sap. 0.5 g. w. H_2SO_4 (T. 2.26-b) to

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
			benzoic ac. (Vol. I) & o-nitroaniline (No. 2.2945). To obtain the o-nitroaniline, make filtrate fr. benzoic ac. strongly alk. w. conc. NaOH; cool; shake; filter; wash w. c. aq.; recryst. fr. 8 cc. boiling 50% alc.; wash w. 3 cc. 50% alc., & dry at 50-60°.
3028	93.5	Yellow	3-Nitro-p-acetoluide, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}(\text{NH.CO.Me})$. — ② Nitro comp.‡ — Fine ndl. fr. aq. or conc. alc. sol. Also cryst. in long colorless ndl. fr. dil. alc., m.p. 94-5°. By fusion & inoculation either modification may be converted into the other. — S. in c. conc. aq. KOH & rapidly saponified to 3-nitrotoluidine (No. 2.3129), & ac. ac.
3029	94	Gold-yellow	β -Dinitro-p-toluidine, $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_4\text{Me}$. — ② Nitro comp.‡ & prim. amine.¶ — Short ndl. fr. ac. ac. S. h. alc., eth.; d.s. c. HCl.
3030	95	Blood-red	N-Ethylisatine, $\text{Et.C}_6\text{H}_4\text{O}_2\text{N}$. — ② Gives indophene react. w. H_2SO_4 + benzene containing thiophene. — Plates fr. eth. E.s. alc., h. aq.; s. alk. w. yel. color.
3031	95d.	Pale yellow	Phenylnitrosourea, $\text{Ph}(\text{NO})\text{N.CO.NH}_2$. — ② Nitrosamine.§ Ndl. fr. eth. + pet.-eth. V. unstable. V.s. alc.; i. pet.-eth. Heated w. aq. emits odor of phenol.
3032	95	V. light yellow	5-Nitrocresol(2), $\text{NO}_2\text{C}_6\text{H}_4\text{Me(OH)}$. — ② Nitro comp.‡ Silky ndl. fr. aq. D.s. aq.; v.s. alc., eth.; v.d.s. lgr. Dec. Na_2CO_3 giving yel. sol. — Reduction w. Sn & HCl gives 5-amino-cresol(2), lft. fr. bz., m.p. 174°.
3033	95	Yellow	6-Nitro-1,3-xylenol(4), $\text{NO}_2\text{C}_6\text{H}_4\text{Me(OH)}$. — ② Nitro comp.‡ — Ndl. — KÄ.2H ₂ O, red cryst., e.s. aq., alc.
3034	95	Yellow	Propyl-2,4-dinitroaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NHPr}$. — ② Nitro comp.‡ — Ndl.
3035	95	Yellow	Isobutylpicramide, $(\text{Me}_2\text{CH.CH}_2\text{NH.C}_6\text{H}_4\text{NO}_2)_2$. — ② Nitro comp.‡
3036	95	Sulphur-yellow	p-Nitroethylaniline, $(\text{NO}_2)\text{C}_6\text{H}_4\text{NHET}$. — ② Nitro comp.‡ — Pr. fr. alc. w. blue-violet reflections. E.s. alc., eth.; d.s. CS ₂ , lgr.
3037	95	Green	β -Nitroso- α -ethylnaphthylamine, $\text{NO.C}_6\text{H}_4\text{NHEt}$. — Lft. fr. dil. alc.
3038	96-7	Red	8-Nitro- α -naphthylamine, $\text{NO}_2\text{C}_{10}\text{H}_8\text{NH}_2$. — ② Nitro comp.‡ & prim. amine.¶ — Scales. — Acetyl deriv.,** m.p. 187-8°.
3039	96-7	Red	p-Nitrosophenylacetanilide, $\text{NO.C}_6\text{H}_4\text{N}(\text{C}_6\text{H}_5\text{O})\text{Ph}$. — Pr. fr. lgr. E.s. alc., eth., bz.; less s. lgr.
3040	96	Or.-yel.	3,6-Dinitro-1,2,4-trimethylbenzene, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}_3$. — ② Nitro comp.‡ — Cryst. fr. alc. Alm. i. c. aq.
3041	96	Pale yel. or or.-yel.	3-Nitro-o-toluidine, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)$. — ② Nitro comp.‡ & prim. amine.¶ — Pr. fr. 45% alc. E.s. alc., eth., chlf., bz. — Acetyl deriv.,** e.s. alc., m.p. 158°.
3042	96-7	Lemon-yellow	2,4,6-Triaminophenol Picrate, $[(\text{NH}_2)_3\text{C}_6\text{H}_3\text{OH}]\text{Pk}$. — ② Picrate* of a v. unstable phenol whose hydrochloride in dil. sol. is colored deep blue by FeCl ₃ . — Ndl. s. in 500 pt. c. aq.
3043	96	Yellow	Kreosol Picrate, $[\text{Me}^1\text{C}_6\text{H}_4\text{(OH)}^4(\text{OMe})^4]\text{Pk}$. — ② Picrate* of species of Vol. I. — Ndl.
3044	96	Yellow	β -Isobutylnaphthalene, Picrate, $\text{C}_{14}\text{H}_{16}\text{Pk}$. — ② Picrate* of h.c. of b.p. 280°.
—	96	Yellowish	3,4,6-Trinitrophenol. — Cf. No. 2.56.
3045	96u.c.	YT2	m-Nitrophenol, $\text{NO}_2\text{C}_6\text{H}_4\text{OH}$. — ② Nitro comp.‡ giving O colored sol. when 0.03 g. is diss. in 10 cc. 10% NaOH. — 5 drops 3% FeCl ₃ added to 5 cc. sat. aq. sol. gives VRS1-RS1

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(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
			<i>sol.</i> — Cryst. of aromat. odor. Sat. aq. sol. tastes at first sweet (No. 3, scale of T. 2.9) & then pungent (No. 2). V.d.s. c. aq.; e.s. h. aq.; v.s. alc., eth. — B.p. 194°. — \textcircled{D} Heat 0.2 g. w. 5 drops benzoyl chloride to boiling in t.t. for 30 sec. Wash w. 5 cc. aq. Decant. Add 5 cc. 15% Na_2CO_3 sol. Warm. Filter. Wash residue w. 2 cc. aq. Dry on porous tile. Dis. in 10 cc. boiling lgr. Filter hot. Cool well. Allow to stand 5 min. Shake. Filter. Recryst. fr. 5 cc. boiling lgr. Filter. Dry on tile 15 min. at 76°. The product, 3-nitrophenylbenzoate, cryst. in thin white pr., m.p. 95° u.c.
3046	96	Yellow	2'-Amino-4-methylbenzophenone, $\text{NH}_2\text{C}_6\text{H}_4\text{CO.C}_6\text{H}_4\text{Me}$. — \textcircled{D} Prim. amine. — Cryst. fr. alc. — Picrate*, pr. fr. alc., browns at 140°, m.p. 146°.
3047	96	Yellow	p-Nitrodimethylaminobenzhydrol, $\text{NO}_2\text{C}_6\text{H}_4\text{CH(OH).C}_6\text{H}_4\text{-NMe}_2$. — \textcircled{D} Nitro comp.‡ — Ndl. fr. dil. alc. I. aq., lgr.; v.s. alc.
3048	96-7	Light yellow	Nitrophenetidine, $(\text{NO}_2)_2\text{C}_6\text{H}_4(\text{EtO})^2(\text{NH}_2)^2$. — \textcircled{D} Nitro comp.‡ & prim. amine. — Ndl. fr. dil. alc. V.d.s. aq.; v.s. alc. — Acetyl deriv.** yel. ndl. fr. alc., m.p. 196°.
3049	96	Light yellow	Tetranitroethylaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_2\text{NET}(\text{NO}_2)_2$. — \textcircled{D} Boiled w. KOH sol. gives ethylamine (No. 2.1062) & picric ac. (No. 2.3168). Lft. fr. alc.
3050	96	Yellow	Diazoaminobenzene, Ph.N. ₂ NH.Ph. — \textcircled{D} Heated w. conc. HCl efferv. briskly w. evolution of N gas & formation of phenol! Deflagrates when rapidly heated on Pt foil! — Lft. fr. alc. Flat pr. fr. bz. I. aq., dil. ac.; d.s. c. alc.; mod. s. h. alc.; e.s. eth., bz.
3051	97	Light red	N-Ethylcarbazole Picrate, $\text{C}_{12}\text{H}_{13}\text{N.Pk}$. — \textcircled{D} Picrate* of No. 2.1646. — Ndl. E.s. alc.
3052	97-8	Yellow	Desylanilide, Anilbenzoin, Ph.CO.CH(NH.Ph).Ph. — \textcircled{D} Warming w. alc. KOH gives purple-violet sol.! — Ndl. fr. alc. D.s. c. alc., eth.; e.s. bz., chlf. — Acetyl deriv.** (by boiling w. ac. anhydride), ndl. fr. bz., m.p. 153°.
3053	96-8	Light yellow	Petrocine Picrate, $\text{C}_{12}\text{H}_8\text{Pk}$. — Picrate* of h.c. of m.p. 101-2°.
—	97	Pale green	Furfurolphenylhydrazone. — Cf. No. 2.1847.
3055	98	Yel.-or.; brown-red	5-Nitro-m-toluidine, $\text{NO}_2\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)$. — \textcircled{D} Nitro comp.‡ & prim. amine. — Ndl. V.d.s. c. aq.; e.s. alc., bz.; v.s. eth.
3056	98	Dark yellow	5-Nitro-2-ethyltoluidine, $\text{NO}_2\text{C}_6\text{H}_4\text{NHEt}$. — \textcircled{D} Nitro comp.‡ — Lft. fr. alc. R.d.s. c. alc.
3057	98	Yellow	o-Nitrobenzil, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.CO.Ph}$. — \textcircled{D} Nitro comp.‡ — Ndl. w. pale green reflections fr. alc. D.s. alc.; e.s. eth., bz.
3058	98	Lemon-yellow	α -Ethynaphthalene Picrate, $\text{C}_{12}\text{H}_{11}\text{Pk}$. — Picrate* of h.c. of Vol. I. — Fine ndl.
3059	98	Yellow	Acetaldehyde-m-nitrophenylhydrazone, $\text{Me.CH: N.NH.C}_6\text{H}_4\text{-NO}_2$. — \textcircled{D} Hydrazone. ‡ — Cryst. fr. h. alc. D.s. h. aq.; e.s. alc., eth., bz.
3060	98d.	Yel.-green	Nitroso-p-nitrosophenylaniline, $(\text{NO})\text{NPh.C}_6\text{H}_4\text{NO}$. — \textcircled{D} Nitrosamine. § — Lft. fr. eth. E.s. alc., eth., bz.
3061	98.5; 95.5	Pale green	N-Benzoylcarbazole, $\text{C}_{12}\text{H}_9\text{N.CO.Ph}$. — Silky pr. fr. alc. D.s. c. alc., bz.; s. eth.
3062	100-1	Yellow	α -Benzylnaphthalene Picrate, $[\text{Ph.CH}_2\text{C}_{10}\text{H}_7]\text{Pk}$. — \textcircled{D} Picrate* of h.c. of Vol. I. — E.s. c. alc.
3063	100	Gold-yellow	2,3,6-Trinitrophenol-naphthalene, $(\text{NO}_2)_3\text{C}_6\text{H}_2(\text{OH}).\text{C}_{10}\text{H}_8$. — Ndl. fr. alc.
3064	100	Yellow-white	β -Trinitrotoluene-naphthalene, $[(\text{NO}_2)_3\text{C}_6\text{H}_3\text{Me}].\text{C}_{10}\text{H}_8$. — Ndl. fr. alc.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3065	101-2	Blood-red	Isatine-o-methylether, $C_8H_4ON.OMe$. — Rhomb. pr. fr. bz. E.s. eth.; less s. alc.; i. lgr. V. unstable. Slowly s. dil. KOH, acids pptg. isatine (No. 2.3633) fr. sol..
3066	101	Yellowish	p-Nitrosomethylphenylnitrosamine, $NO.C_6H_4.N(NO)Me$. — \textcircled{P} Nitrosamine. \textcircled{S} — Cryst. giving greenish sol. in alc.
3067	101-2	Yellow	Nitroacenaphthene, $NO_2.C_{12}H_8$. — Nitro comp. \ddagger — Ndl. fr. lgr. I. aq.
3068	102-3	Red or or.-yel.	4'-Hydroxy-2-methylazobenzene, $HO.C_6H_4.N_2.C_6H_5.Me$. — \textcircled{P} Azo comp. \parallel — Cryst. fr. lgr. + bz. V.s. alc., eth.; d.s. lgr. V.s. NaOH sol.! (Sometimes cryst. w. H_2O , m.p. 76°; 66°.)
3069	102	Or.-yel.	3,4,5(?) -Trinitrocresol(2), $(NO_2)_3.C_8H_6.Me(OH)$. — \textcircled{P} Nitro comp. \ddagger — Thick pr. fr. acetone. D.s. c. aq.; e.s. alc., eth., chlf. — Gives naphthalene deriv., yel. ndl. fr. acetone, m.p. 106°.
3070	102-3	Yellow	3-Nitro-4-dimethylaminobenzaldehyde, $NO_2.C_6H_4(NMe_2).CHO$. — Ndl. fr. alc. E.s. bz., eth. [D.R.P. 89,244.]
3071	102	Yellow	Nitroso-p-nitromethylaniline, $(NO)MeN.C_6H_4.NO_2$. — \textcircled{P} Nitrosamine. \textcircled{S} — Ndl.
3072	101; 103	Gold-yel.	Di-p-tolylnitrosamine, $(Me.C_6H_4)_2.N.NO$. — \textcircled{P} Nitrosamine. \textcircled{S} — E.s. eth., bz., lgr.; d.s. alc.
3073	103.5	Red	8-Nitro- β -naphthylamine, $NO_2.C_{10}H_8.(NH_2)$. — \textcircled{P} Nitro comp. \ddagger & prim. amine. \textcircled{I} — Lust. ndl. E.s. alc. I. lgr. — Benzoyl deriv., m.p. 162°.
3074	102-4	Red	Benzeneazo- β -naphthylamine, $Ph.N_2.C_{10}H_8.NH_2$. — \textcircled{P} Azo comp. \parallel Sol. in conc. H_2SO_4 w. blue color! — Rhomb. tbl. fr. abs. alc. I. aq.; mod. s. alc., gl. ac. ac. — Acetyl deriv., ** red ndl. fr. alc., m.p. 152-3°; i. aq.; e.s. alc.
3075	103-4	Yellow	p-Nitroguaiacol, $NO_2.C_6H_4.(OMe)^2(OH)^1$. — \textcircled{P} Sol. in alk. purple-red! Odor, faint & vanilla-like. — Ndl. fr. aq. E.s. alc., eth., h. aq.
3076	103	Yellow	1-Nitronaphthol(2), $NO_2.C_{10}H_8.OH$. — \textcircled{P} Nitro comp. \ddagger — Cryst. fr. alc. D.s. c. alc.; e.s. eth. — NaÅ cryst. in red ndl., aq. sol. giving red ppt. w. Pb or Ba salts. — $C_{10}H_8(NH_2)-(OH)$, the reduction product w. Sn & HCl, forms a yel. picrate of m.p. 109-10°.
3077	103	Yellow	3-Nitro-4-acetaminophenoletylether, $(NO_2)(NH.CO.Me)-C_6H_4.OEt$. — \textcircled{P} Nitro comp. \ddagger — Silky ndl. fr. aq. E.s. alc., eth., chlf.
3078	104.5	Or.-yel. or red- brown	p-Nitrodiphenylbutylene, $NO_2.C_6H_4.N_2.NH.NH.Ph$. — \textcircled{P} Sol. in conc. H_2SO_4 w. yel.-green color. — Ndl. fr. alc. E.s. chlf., bz.
3079	104.5-5	Or.-yel.	o-Nitro diazoaminobenzene, $NO_2.C_6H_4.NH.N_2.Ph$. — Ndl. fr. alc. E.s. chlf., bz.
3080	104	Light yellow	2,5-Dinitrophenol, $(NO_2)_2.C_6H_3.OH$. — \textcircled{P} Nitro comp. \ddagger — Ndl. fr. aq. D.s. aq., c. alc.; e.s. eth. E. vol. w. st.! — Methyl ether, m.p. 96°.
3081	104	Yellowish	(γ) 2,4,5-Trinitrotoluene, $(NO_2)_3.C_6H_3.Me$. — [An explosive.] — \textcircled{P} Nitro comp. giving blue color w. acetone + ammon. — 6-sided tbl. fr. acetone. D.s. c. alc.; e.s. eth., bz., acetone. — Treated w. KOH gives dinitrocresol (Ber., 37, 704). — Naphthalene addition product, m.p. 88-9°.
3082	104.5	Yellow	Benzaldehydemethylphenylhydrazone, $Ph.CH:N.NMe.Ph$. — \textcircled{P} Hydrazone (T. 2.17). — Ndl. fr. lgr. I. aq.
3083	105	Dark red	Pr-3-Phenylindole Picrate, $C_{11}H_11N.Pk$. — \textcircled{P} \textcircled{I} Picrate* of No. 2.1772. — Ndl. v.s. alc.; v.d.s. lgr.

Explanation of typographical signs used in this Division: * = T. 2.39. \ddagger = T. 2.21. \textcircled{S} = T. 2.36. \parallel = T. 2.34.
 \textcircled{P} = T. 2.35. \textcircled{I} = T. 2.1. \textcircled{II} = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3064	105d.	Yellow	Nitrosophenylglycine, $(NO)NPh.CH_2CO_2H$. — \textcircled{P} Nitrosoamine. § — Long ndl. E.s. h. aq.; v.s. alc., eth. Dec. carbonates. Boiling w. aq. give CO_2 & methylphenylnitrosoamine (No. 2.2809).
3085	105	Yellow	Anilbenzil, $Ph.CO.C(NPh).Ph$. — \textcircled{P} S. in conc. H_2SO_4 w. blood-red color. — Trichin. pr. fr. alc. D.s. c. alc.; more s. eth., bz.
3086	105-6	Yellow	2,4,6-Trinitrocresol(3), $(NO_2)_3C_6H_3Me(OH)$. — \textcircled{P} Nitro comp. ‡ giving yel. sol. in alk. — Ndl. fr. aq. S. in 449 pt. aq. at 20°, or in 123 pt. at 100°; e.s. alc., eth.
3087	105-6	Light yellow	o-Aminobenzophenone, $Ph.CO.C_6H_4.NH_2$. — \textcircled{P} Prim. amine. ¶ — Cryst. fr. alc.
3088	104-6	Light yellow	Picrylpiperidine, $(NO_2)_3C_6H_3NC_6H_{10}$. — Pr. fr. alc. Alm. i. methyl alc., chlf., lgr.; e.s. bz.
3089	106	Garnet-red	m-Nitrocinnamicaldehydephenylhydrazone, $(NO_2).C_6H_4.CH:CH.CH:N.HPh$. — \textcircled{P} Hydrazone (T. 2.17). — Tbl. fr. alc. I. aq.
3090	106.5	Pale yellow	Nitroso-2,4,6-trinitromethylaniline, $(NO)MeN.C_6H_4(NO_2)_3$. — \textcircled{P} Nitrosamine. § — Cryst. fr. alc. Alc. sol. colored blood-red by KOH.
3091	106	Yellowish	Trinitro-o-cresol-naphthalene, $[(NO_2)_3C_6HMe(OH)].C_{10}H_8$. — Ndl. fr. acetone. Softens at 102°.
3092	106	Yellow	2,6-Dinitromethylaniline, $(NO_2)_3C_6H_3.NHMe$. — \textcircled{P} Nitro comp. ‡
3093	106-7	Yellow	2,1'-Dinitrostyrene, $NO_2.C_6H_4.CH:CH(NO_2)$. — \textcircled{P} Nitro comp. ‡ — Ndl. fr. alc. Vol. w. st. S. in NaOH when freshly pptd. fr. alc. by aq. — Oxid. by $KMnO_4$ gives o-nitrobenzoic ac. (No. 2.164).
3094	107.5	Red or yellow	4-Nitromethyl-o-toluidine, $NO_2.C_6H_4.Me(NH.Me)$. — \textcircled{P} Nitro comp. ‡ — Red lft. or yel. ndl. fr. alc. or lgr. E.s. eth., chlf.; d.s. alc., lgr. — Nitrosamine, § m.p. 95°. — B.Pk, compact red pr.
3095	107	Yellow	Dinitrodimethyl-m-toluidine, $(NO_2)_3C_6H_2.Me(NMe_2)$. — \textcircled{P} Nitro comp. ‡ — Ndl. i. c. aq.
3096	107	Pale yellow	5-Nitro-4-acetamino-1,2-xylene, $(NO_2)(Me.CO.NH).C_6H_4.Me_2$. — \textcircled{P} Nitro comp. ‡ — Ndl. fr. alc.
3097	107	Gold-yel.	m-Nitrophenylbenzylamine, $NO_2.C_6H_4.NH(CH_2Ph)$. — \textcircled{P} Nitro comp. ‡ — Lamella fr. alc.
3098	107	Light yellow	m-Nitrobenzalhydrazine, $NO_2.C_6H_4.CH:N.NH_2$. — \textcircled{P} HgO colors sol. intense red. — Tbl. fr. alc. — In alc. sol. is changed by acid to m,m'-dinitrobenzalzin, ndl. fr. alc., m.p. 194°.
3098-I	107; 109; 104-5		4-Nitro-o-toluidine, $(NO_2).C_6H_4.Me(NH_2)$. — \textcircled{P} Taste, v. sweet. — Monoclin. pr. S. alc., ether.
3099	107	Golden	m-Nitrobenzenediazoamino-p-toluene, $NO_2.C_6H_4.NH.N_2C_6H_4.Me$. — Lust. ndl. fr. bz.
3100	106-8	Dark violet	1,3,5-Trinitrobenzenedimethylaniline, $[(NO_2)_3C_6H_2]Ph.NMe_2$. — V.d. s. alc.
3101	108	Deep red	Picrylmethylaniline, 2,4,6-Trinitro-phenyl-methylaniline, $(NO_2)_3C_6H_3.NMePh$. — Tbl. fr. alc.
3102	108 u.c.	YT3	Acridine, $C_{13}H_9N$. — \textcircled{P} (1) Vapors fr. melted substance or suspension in boiling aq. exceedingly irritating to mucous membranes! Taste of c. (filtered!) aq. sol. painfully burning. (Abt. No. 4 in pungency T. 2.29.) Powder on contact w. skin causes persistent painful irritation. — B.p. a. 360°. V.d.s. to i. aq.; e.s. alc., eth. Vol. w. st. — (2) Dis. abt. 2 mg. in 1 cc. HNO_3 or H_2SO_4 & dil. w. 20 cc. aq. Pale yel. sol. shows fine GT1-BGT1 fluor. when held before black back-

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
			<i>ground. The fluor. of HCl sol. is much weaker. — B.Pk melts w. decn. at v. high temp. — [Cf. Ber., 17, 438, for other methods for identification & determination.]</i>
3103	108	Light yellow	Diphenyldiacetylene Picrate, [Ph.C : C.C : C.Ph].Pk. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I.
3104	108-8.5	Gold-yel.	β -o-Hydroxyazoxybenzene, $\text{HO.C}_6\text{H}_4.\text{N}_2\text{O.Ph}$. — \textcircled{P} S. in NaOH sol. w. gold-yel. color. — Ndl. fr. pet.-eth. E.s. eth., chlf., h. lgr.; d.s. c. alc. — Reduction gives o-aminophenol (No. 2.2890) & aniline (No. 2.1235).
3105	108-9	Golden	3-Methyl- β -hydroxyazobenzene, $\text{Me}(\text{HO}).\text{C}_6\text{H}_4.\text{N}_2\text{Ph}$. — \textcircled{P} S. in conc. H_2SO_4 w. brown color. — Lft. Sbl. E.s. alc., eth., chlf., warm dil. NaOH. — Acetyl deriv.** yel. ndl. fr. dil. alc., m.p. 87-8°.
3106	109.5	Or.-brown	1-(α)-Nitrosonaphthalol(?) [†] , β -Naphthoquinoneoxime, $\text{O}:\text{C}_{10}\text{H}_8:-\text{NOH}$. — \textcircled{P} Sol. in ac. ac. gives black ppt. w. FeCl_3 , i. dil. ac. ac. or aq., but s. w. deep blue color in aniline. — Lft. or pr. fr. alc., eth., bz., lgr. Alm. i. c. aq.; v.d.s. h. aq.; s. in 42 pt. alc. at 13°; v.s. h. alc.; v.s. eth., bz. E. vol. w. st., but resiniified by h. aq. when impure. — Alk. salts green. NaA pptd. fr. alc. sol. by alc. NaOH sol. — Oxidn. by dil. HNO_3 gives nitronaphthalol, yel. cryst. fr. alc., m.p. 103°.
3107	109	Gold-yellow	4-Nitro-m-toluidine, $(\text{NO}_2)(\text{NH}_2).\text{C}_6\text{H}_4.\text{Me}$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Lft. fr. h. aq. E.s. h. aq. E.s. alc., eth., chlf., bz.
3108	109	Pale yellow	4-Aminoantipyrine, $[\text{NPh.NMe.CMe}:\text{C}(\text{NH}_2).\text{CO}]$. — Cryst. fr. bz. E.s. aq., alc., bz.; d.s. eth. — In alc. sol. benzaldehyde gives yel. lft. of benzylidene deriv., d.s. alc., m.p. 173°. — Acetyl deriv.** e.s. aq.; d.s. eth., bz.; m.p. 197°. — B.Pk, lft., v.d.s. aq.; e.s. alc.; m.p. 144°.
3109	110-1	Red	6-Nitro-2-aminophenol, $(\text{NO}_2)(\text{NH}_2).\text{C}_6\text{H}_4.\text{OH}$. — Ndl. fr. dil. alc. V.d.s. c. aq.; e.s. alc.; v.s. eth., chlf., bz.
3110	110-1	Or.-yel.	2,3,2',3'-Tetramethylazobenzene, $\text{Me}_2.\text{C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_4.\text{Me}_2$. — \textcircled{P} Azo comp. — Ndl. E.s. alc., eth., bz.
3111	109-11	Gold-yel.	Retenequinoneimide, $\text{O}:\text{C}_{10}\text{H}_{14}:\text{NH}$. — \textcircled{P} Sol. in strong acids dark violet! — E.s. most organic solvents. Warming w. alk. or long exposure to moist air gives retenequinone (Cf. Vol. I), or. ndl., d.s. h. alc., ammon.
3112	110	Yellow	α -Aminobenzylalcohol Picrate, $\text{HO.CH}_2.\text{C}_6\text{H}_4.\text{NH}_2.\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.690. — Ndl.
3113	110	Lemon-yellow	β -Isoamylnaphthalene Picrate, $\text{C}_{10}\text{H}_{18}.\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate of h.c., b.p. 288-92°. — Ndl. fr. alc.
3114	110-1	Yellow	2,4,6-Trinitromethylaniline, $(\text{NO}_2)_3.\text{C}_6\text{H}_2.\text{NHMe}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc. V.s. bz.
3115	111-2	Brown	p-Nitromethylbenzoylanilide, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NMe}(\text{CO.Ph})$. — \textcircled{P} Nitro comp.‡ — Fr. fr. dil. alc.
3116	111	Lemon-yellow	2,5,6-Trinitrothymol, $(\text{NO}_2)_3.\text{C}_6\text{H}_2(\text{C}_6\text{H}_5)^4(\text{OH})^2$. — Ndl. fr. aq.
3117	111-2	Yellow	Furfuralazin, $\text{C}_6\text{H}_5.\text{CH}:\text{N.N}: \text{CH.C}_6\text{H}_5$. — [Fr. hydrazine & furfural.] — Cryst. fr. alc.
3118	112	Dark red	Acetone-m-nitrophenylhydrazone, $\text{Me}_2\text{C}:\text{N.NH.C}_6\text{H}_4.\text{NO}_2$. — \textcircled{P} Hydrazone. — Ndl. fr. dil. alc. V.d.s. h. aq.; e.s. h. alc., eth., bz.
3119	110-2; 113-4	Yellow	Benzeneazotriphenylmethane, $\text{Ph.N}_2.\text{CPh}_3$. — \textcircled{P} Azo comp. — Cryst. fr. alc., eth., bz. — Dec. at 120° to N ₂ & tetraphenylmethane.
3120	112	Yellow	Dinitropseudocumenol, $(\text{NO}_2)_2.\text{C}_6\text{H}_4.\text{Me}_2^{(1,2,4)}(\text{OH})^2$. — \textcircled{P} S. in alk. w. deep red color. — Cryst. I. aq.; e.s. alc., eth., chlf. Decd. by boiling w. aq.

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(ORDER II.)

No.	Melting-point (C. ^o). ⁵	Color.	COLORED SOLID COMPOUNDS.
3121	112	Gold-yellow	o-Nitrooxanilic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{NH.CO.CO}_2\text{H}$. — \textcircled{P} Decd. to oxalic ac. & o-nitroaniline (No. 2.2945) by protracted boiling w. aq. — Long ndl. fr. h. aq.; d.s. c. aq., eth.; e.s. alc.
3122	112	Light yellow	5-Nitro-3-amino-1,2-xylene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Ndl. fr. alc. — Acetyl deriv.,** white ndl., m.p. 230–1°.
3123	113	Red	2-Nitro-4-aminophenoylether, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{OEt}$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Lust. pr. fr. alc. V.d.s. c. alc.; e.s. eth., chlf.
3124	113–4	Yellow	2,4-Dinitroethylaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{NHEt}$. — \textcircled{P} Boiled w. conc. KOH gives ethylamine (No. 2.1062), & 2,4-dinitrophenol (No. 2.3126). — D.s. eth., CS.
3125	114 u.c.	OY	m-Nitroaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NH}_2$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Flat ndl. fr. h. dil. alc. — D.s. aq.; s. alc., eth. — \textcircled{D} Mix 0.05 g. substance w. 6 drops ac. anhydride & heat to boiling. Boil product w. 4 cc. aq. Filter sol. h., if not clear. Cool & shake persistently. Wash ppt. on filter w. 1 cc. c. aq. Recryst. fr. 3 cc. h. aq. Dry on tile 15 min. at 100°. The product, m-nitroacetanilide, is obtained in white granular form, m.p. 149.7–150.4° u.c.
3126	114 u.c.	(Y-GY)T2	2,4-Dinitrophenol, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{OH}$. — \textcircled{P} Nitro comp. [‡] — 0.03 g. dis. in 10 cc. N/10 alk. w. YT1 color. — Alc. sol. colored OR by FeCl_3 . — Rectangular plates fr. h. aq. or alc. Odorless cold. Sat. aq. sol. tasteless. V.d.s. c. aq.; e.s. h. aq.; s. alc. — \textcircled{D} Heat moderately in a t.t. for 30 sec. 0.2 g. substance & 5 drops benzoyl chloride. Wash product w. 5 c. c. aq. Heat to boiling w. 5 cc. dil. ammon. Cool well & filter. Wash residue w. 2 cc. c. acetone. Dry 15 min. at 100°. 2,4-dinitrophenylbenzoate, the product, cryst. in Y pr., m.p. 213.7° u.c., v. sl. decn.
3127	114	Yellowish	6-Nitro-3-amino-1,2-xylene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Ndl. fr. alc. — Acetyl deriv.,** pale yellowish ndl., m.p. 149–50°.
3128	114–5	Yellow	p-Nitrophenylnitropropylene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}=\text{C}(\text{NO}_2)\text{Me}$. — \textcircled{P} Nitro comp. [‡] — Ndl. fr. alc. — Oxidn. by KMnO_4 gives p-nitrobenzoic ac. (No. 2.425).
3129	114; 116–7	Red	3-Nitro-p-toluidine, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Pr. fr. alc. V.d.s. h. aq.; e.s. alc. Vol. w. st. — Reduction gives 3,4-toluylenediamine (No. 2.707).
3130	115–6	Yellow or red-yel.	4,4'-Dimethyldiazoaminobenzene, $\text{Me.C}_6\text{H}_4\text{N.NH.C}_6\text{H}_4\text{Me}$. — Ndl.
3132	115	Lemon-yellow	4-Nitroresorcinol, $\text{NO}_2\text{C}_6\text{H}_3\text{(OH)}_2$. — \textcircled{P} Nitro comp. [‡] giving or-yel. sol. in alk. — Hair-like ndl.
3133	115	Deep yellow	β -Methylnaphthalene Picrate, $\text{C}_{11}\text{H}_{10}\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Ndl.
3134	115	Pale yellow	Dinitrotetraethylbenzene, $(\text{NO}_2)_2\text{C}_6\text{Et}_4$. — \textcircled{P} Nitro comp. [‡] — Rhomb. pr. fr. alc.
3135	115; 113	Light yellow	o-Nitrohydrocinnamic Ac., $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Nitro comp. [‡] & acid. Ndl. fr. h. aq. — Reduction by Sn & HCl gives hydrocarbostyrene (No. 2.2282).
3136	115	Yellow	2,5-Dinitro-4-amino-1,3-xylene, $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}_2$. — \textcircled{P} Nitro comp. [‡] & prim. amine. [¶] — Ndl. E.s. alc. — \textcircled{D} Acetyl deriv.,** ndl. d.s. alc., m.p. 226°.
3137	115	Yellow	2-Nitro-1-diacetaminonaphthalene, $(\text{NO}_2)\text{C}_6\text{H}_4\text{N}(\text{C}_6\text{H}_4\text{O})_2$. — \textcircled{P} Nitro comp. [‡] — Triclin. pr. fr. gl. ac. ac. — Boiled w. alc. ammon. gives monoacetyl deriv., m.p. 190°.
3138	115; (118)	Yel.-green or steel-blue	Methyl-p-nitrosoaniline, $\text{NO.C}_6\text{H}_4\text{NHMe}$. — Lft. fr. bz. showing bluish reflections, but green by transmitted light! D.s. aq. w. yel. or green color. E.s. dil. NaOH; repptd. by CO_2 .

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3139	115-6	Green	5-Nitroso-o-toluidine, $(NO)(NH_2)C_6H_4Me$. — \oplus Boiling w. NaOH gives NH ₃ (T. 2.7) & o-cresol (Vol. I). — Ndl. fr. bz. Somewhat s. aq.; e.s. alc., eth., chlf.; d.s. lgr. Deflagrates freely on ignition.
3140	116-7	Or.-yel.	α -Methylnaphthalene Picrate, $C_{11}H_{10}Pk$. — \oplus Picrate* of h.c. of Vol. I. — Long fine ndl. fr. alc.
3141	116-7	Pale yellow	9(?) - Nitrophenanthrene, $NO_2C_{10}H_8$. — \oplus Sol. in conc. H ₂ SO ₄ w. blood-red color, changing to green on warming. — Ndl. fr. alc. D.s. c. alc., eth.; e.s. bz., chlf.
3142	116	Yellow	p-Dihydroxyazoxybenzenedimethylether, $MeO.C_6H_4.N.O.C_6H_4.OMe$. — Melts sharply to turbid liquid becoming suddenly clear at 134°! — Cryst. fr. alc.
3143	115; 117	Yellow	p-Dimethylaminoazobenzene, $Ph.N:N.C_6H_4.NMe_2$. — [An intermediate in color industry.] — Lft. fr. alc. — Hydrochloride forms hair-like purple-red ndl. — Sn & HCl should reduce to aniline (No. 2.1235) & dimethyl-p-phenylenediamine (No. 2.560).
3144	116-0.5	Pale straw-yellow	2,3,2',3'-Tetramethylazoxybenzene, $Me_2C_6H_3.N.O.C_6H_4.Me_2$. — Ndl. s. alc.; v.s. eth., bz.; e.s. pet.-eth.
3145	117-8	Carmine-red	Benzaldehyde-m-nitrophenylhydrazone, $(NO_2)C_6H_4.NH.N:-CH.Ph$. — \oplus Hydrazone.†† — Ndl. fr. h. alc.; i. aq.
3146	117.5	Red	Ethyl Formazylformate, $(Ph.N:N)(Ph.NH.N:)C.CO_2Et$. — Lft. fr. h. alc. I. alk.; d.s. lgr. — Sapn. (T. 2.26) by boiling alc. NaOH to formazylformic ac. (No. 2.3417), & ethyl alc.
3147	117.5	Light yellow	2,4-Diaminoazobenzene, Chrysoidine, $(NH_2)_2C_6H_4.N_2Ph$. — [Commercial Chrysoidine, the dyestuff, is B.HCl.] — \oplus Sol. in x.s. conc. HCl, carmine-red. — Cryst. fr. aq. in "threads." D.s. aq.; more s. alc., eth.; e.s. chlf. — B.HCl, blackish cryst. giving red powder s. in aq. w. deep OR color. — Diacetyl deriv., ndl. fr. gl. ac. ac., m.p. 250.5°. — [Cf. No. 3.509 for description of the commercial dyestuff.]
3148	117.5-8u.c.	YT3	(ms)-9-Methylacridine, $Me.C_12H_8N$. — \oplus Dis. abt. 2 mg. in 1 cc. sulphuric, hydrochloric, or nitric ac., & dil. w. 20 cc. aq. The pale yel. sol. shows strong green to blue-green fluor. when viewed against black background! — Ndl. fr. alc.; plates fr. lgr. I. aq. Nearly odorless. — B.Mel (T. 2.37) (fr. components at 100°), silky red ndl. fr. aq., m.p. 185° d.
3149	117	Light yellow	2-Methylphenazine, Tolazine, $Me.C_12H_8N_2$. — \oplus S. in conc. H ₂ SO ₄ w. blood-red color. — Sbl. in ndl. B.p. abt. 350° d. D.s. lgr., h. aq.; e.s. alc., eth., chlf. S. w. yel. color in conc. HCl; pptd. unchanged by boiling aq. — B.Pk, cryst. spheroids fr. bz., m.p. abt. 168°.
3150	117-8	Yellow	m-Nitrohydrocinnamic Ac., $NO_2C_6H_4.CH_2CH_2CO_2H$. — \oplus Nitro comp.‡ & acid. — Ndl. fr. aq. D.s. c. aq.; e.s. eth.; less s. alc., bz.
3151	117-8	Yellow	2,4-Dinitro-1-methylstyrene, $(NO_2)Me.C_6H_4.CH:CH.NO_2$. — \oplus Nitro comp.‡ — Ndl. E.s. alc.; i. lgr.
3152	118-9	Red	5-Nitro- α -naphthylamine, $NO_2C_{10}H_8.NH_2$. — \oplus Nitro comp.‡ & prim. amine. — Small lust. ndl. fr. h. aq.
3153	118-9	Brick-red	4-Nitro-3-amino-1,2-xylene, $(NO_2)(NH_2)C_6H_4Me_2$. — \oplus Nitro comp.‡ & prim. amine. — Cryst. fr. alc. — Acetyl deriv.,** white ndl. fr. alc., m.p. 160°.
3154	118	Or.-red	4-Nitro-o-anisidine, $NO_2C_6H_4.(MeO)^1(NH_2)^2$. — \oplus Nitro comp.‡ & prim. amine. — Ndl. fr. aq. E.s. h. alc., eth. [D.R.P., 98,637.]

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34.
† = T. 2.35. ** = T. 2.1. ||| = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3155	118.5	Or.-red	6-Amino-3,4'-dimethylazobenzene, $(\text{NH}_2)\text{Me.C}_6\text{H}_3.\text{N}_2.\text{C}_6\text{H}_4.\text{Me.}$ — \textcircled{P} \textcircled{O} Azo comp., w. p-toluidine (No. 2.566) & 3,4-toluyl-enediamine (No. 2.707) as reduction products. — Lust. ndl. E.s. h. alc.; v.s. bz. Salts give greenish sol. — Acetyl deriv.,** felted yel. ndl. fr. alc. & gl. ac. ac., m.p. 157°.
3156	118	Orange	β -Dimethylnaphthalene Picrate, $\text{C}_{12}\text{H}_4.\text{Pk.}$ — \textcircled{P} \textcircled{O} Picrate* of h.c. of b.p. 264–6°. — Fine pr.
3157	118	Yellowish	(β)-2,4'-Dinitrodiphenylmethane, $(\text{NO}_2\text{C}_6\text{H}_4)_2\text{CH}_2.$ — \textcircled{P} Nitro comp.† — Monoclin. pr. fr. bz.
3158	118	Yellow	4-Nitrocresol(2), $\text{NO}_2\text{C}_6\text{H}_4.\text{Me(OH).}$ — \textcircled{P} Nitro comp.‡ s. alk. — Ndl. fr. lgr. D.s. c. aq., lgr.; e.s. alc., eth., bz.
3159	117-9; 119-20	Red	Formazan, $\text{Ph.NH.N:CH.N:N.Ph.}$ — \textcircled{P} Sol. in conc. H_2SO_4 dark blue. — Ndl. w. violet reflections fr. dil. MeOH. E.s. alc.; d.s. lgr. KOH colors alc. sol. dark red. — $\text{B}.\text{AgNO}_3$, cryst. brick-red ppt.
3160	119-20	Red	p-Nitrophenylhydrazine Picrate, $\text{C}_6\text{H}_4\text{O}_2\text{N.Pk.}$ — Picrate* of No. 2.3383. — Ndl. fr. h. aq. S. h. alc.; i. lgr., bz.
3161	119	Dark orange	Trimethylnaphthalene Picrate, $\text{C}_{12}\text{H}_4.\text{Pk.}$ — \textcircled{P} Picrate* of h.c. of b.p. 290° fr. petroleum. — Ndl.
3162	119	Yellow	2,5,2',5'-Tetramethylazobenzene, $\text{Me}_2\text{C}_6\text{H}_3.\text{N}_2.\text{C}_6\text{H}_3.\text{Me}_2.$ — \textcircled{P} Azo comp. — Ndl.
3163	120	Dark red	4-Nitro-2-amino-2'-methylidiphenylamine, $(\text{NO}_2)(\text{NH}_2).\text{C}_6\text{H}_4.\text{NH.C}_6\text{H}_4.\text{Me.}$ — Cryst. fr. dil. alc.
3164	120-1	Red	m-Nitrobenzaldehydophenylhydrazone, $\text{NO}_2\text{C}_6\text{H}_4.\text{CH:N.NH-Ph.}$ — \textcircled{P} Hydrazone.‡ — Ndl.
3165	120.5	Brown	p-Cyanoazobenzene, $\text{Ph.N}_2\text{C}_6\text{H}_4.\text{CN.}$ — Ndl. fr. bz. I. aq.; e.s. h. alc., eth., bz.
3166	120	Straw-yellow	Nitroso-p-nitroethylaniline, $(\text{NO})\text{EtN.C}_6\text{H}_4.\text{NO}_2.$ — \textcircled{P} Nitroso-amine.§ — Ndl. fr. alc.
3167	120-1; 123-3.5c.	Light or.-yel.	Methylformazyl, $\text{Me.C(:N.NH.Ph).N:N.Ph.}$ — \textcircled{P} Reduces alc. AgNO_3 sol. — Lust. ndl. w. bluish reflections fr. alc. E.s. h. alc., c. chlf., bz.
3168	121 u.c.	YT1; GY, T2-3	Picric Ac., 2,4,6-Trinitrophenol, $(\text{NO}_2)_3\text{C}_6\text{H}_3.\text{OH.}$ — Yel. lf. fr. h. aq. Cryst. fr. h. conc. HCl are colorless, but become yel. on exposure to air. Lemon yel. pr. fr. eth. 100 pt. aq. dis.: at 20°, 1.22 pt.; at 100°, 6.33 pt. 100 g. satd. abs. alc. sol. contain 5.92 g. at 14.8°. 100 g. bz. dis.: at 20°, 9.56 g.; at 75°, 96.77 g. 1000 cc. eth. (sp. gr. 0.721) at 13° dis. 10.8 g.; or, when satd. w. aq., at 15°, 51.2 g. D.s. lgr. Odorless. Not vol. w. st. Deflagrates when strongly heated. Alk. salts readily detonated by percussion. \textcircled{P} (1) Aq. sol. tastes intensely bitter & dyes the skin, wool, or silk, bright yel.! — (2) Boil sol. of 1 mg. substance in 5 cc. aq. w. 5 drops 1% KCN sol. for 30 sec. Sol. shows R color. ("Isopurpuric ac. react.") — As dyestuff, is described as No. 3.947. — \textcircled{P} Convert into any of the charac. organic picrates referred to in T. 2.23.
3169	121	Green	1-Nitroso-2-ethylaminonaphthalene, $\text{NO.C}_6\text{H}_4.\text{NHEt.}$ — Tbl. fr. bz.
3170	122-3	Or.-red	1,2,6-Trimethylnaphthalene Picrate, $\text{C}_{12}\text{H}_4.\text{Pk.}$ — \textcircled{P} Picrate* of h.c. of orange flower odor, b.p. 154–6° (15 mm.).
3171	122	Or.-yel.	Pyrocatechol Picrate, $\text{o-C}_6\text{H}_4(\text{OH})_2.\text{Pk.}$ — \textcircled{P} Picrate* of species of Vol. I.
3172	122	Yellow	4,6-(or 3,5-)Dinitroguaiacol, $(\text{NO}_2)_2\text{C}_6\text{H}_3.(\text{OMe})(\text{OH}).$ — \textcircled{P} Nitro comp.‡ — Lft. fr. dil. alc.
3173	122	Yellow	$\text{o-Nitroformanilide, NO}_2\text{C}_6\text{H}_4.\text{NH(CHO).}$ — \textcircled{P} Nitro comp.‡ — Long ndl. fr. alc. E.s. h. aq., alc.; v.s. bz.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3174	122	Light yellow	1,3,5-Trinitronaphthalene, $(NO_2)_3C_{10}H_4$. — \oplus Nitro comp.‡ — Cryst. fr. chlf. Lft. w. serrate edges fr. alc. E.s. alc., chlf., gl. ac. ac.
3175	122	Pale yellowish	3,5-Dinitrocresol(4)methylether, $(NO_2)_2C_6H_3(OMe)Me$. — \oplus Nitro comp.‡ — Ndl.
3176	122-3	Yellowish	2,9-Dimethylacridine, $Me_2C_{12}H_7N$. — Ndl. or pr. fr. alc. E.s. alc., bz.; less s. eth. Vol. w. st. — B.Pk, brown cryst. fr. alc.
3177	122.1 u.c.	GYT3	s-Trinitrobenzene, $(NO_2)_3C_6H_3$. — Plates or pr. fr. bz. Odorless. Taste of c. sat. aq. sol. bitter (No. 4 on scale of T. 2.29). 100 pt. of following solvents dis. at 16°: MeOH, 4.9 pt.; aq., 0.04 pt.; EtOH, 1.9 pt. — \oplus NaOH sol. dis. w. RO color. — \oplus Dis. 0.05 g. in 15 cc. h. lqr. Add 5 drops aniline. Filter hot. Allow salt to cryst. slowly. Filter. Dry on tile at 55° for 5 min. The addition product cryst. in O ndl., m.p. 124.1° u.c.
3178	123	Red	Benzeneazo- α -naphthylamine, $Ph.N_2C_6H_4NH_2$. — \oplus Azo comp. & prim. amine.¶ — Lust. ndl. fr. dil. alc. Mod. s. alc., eth., bz. Spectrum, Ber. 22, 2069. — Acetyl deriv.,** pale yel. lft. fr. alc., m.p. 233°, e.s. alc., d.s. bz.
3179	123	Or.-red	6-Nitro-4-amino-1,3-xylene, $(NO_2)(NH_2)C_6H_3Me_2$. — \oplus Nitro comp.‡ & prim. amine.¶ — Ndl. E.s. h. alc. — Acetyl deriv.,** m.p. 159-60°.
3180	123-4	Or.-yel.	Retene Picrate, $C_10H_8P_k$. — \oplus \ominus Picrate* of h.c. of Vol. I. — Ndl. fr. alc. + picric ac.
3181	123	Gold-yel.	α -Aminoazobenzene, $NH_2C_6H_4N_2Ph$. — \oplus Azo comp. & prim. amine.¶ — Lust. ndl. fr. dil. alc.
3182	123-5	Dark red	Anilinetrinitroaniline, $Ph.NH_2 + (NO_2)_3C_6H_3NH_2$. — Lust. pr. Partly decd. by boiling alc.
3183	124 u.c.	Red, or or.-red	1,3,5-Trinitrobenzeneaniline, $C_6H_3(NO_2)_3 + Ph.NH_2$. — Lft. S. in warm bz., h. alc. Loses aniline on washing w. aq. or alc.
3184	124-4.5 u.c.	OYT1	Picrolonic Ac., 1,p-Nitrophenyl-3-methyl-4-isontropyrizolone-(5), [¶] $N(C_6H_4NO_2)N:CM_2C:(NO_2H)CO^3$. — Melts w. violent efferv.! D.s. aq., chlf., eth.; v.d.s. bz.; s. alc. Taste v. faintly bitter & sour. Odorless. — \oplus Gives many stable cryst. alkaloidal salts; e.g. brucine salt, m.p. 176-6°. (Cf. No. 2.956, & T. 2.23 & T. 2.39.)
3185	124-5	Light straw-yel.	Succinazone, $Ph.NH.N:CH.CH_2CH_2CH:N.NH.Ph$. — \oplus Hydrazone.‡‡ — Silky lft. fr. alc. — I. aq.; e.s. alc., eth.
3186	124d.	Light yellow	α -Nitrophenyllactone, $^FO.CH(C_6H_4NO_2).CH_2CO^3$. — \oplus Turns blue in melting! — Monoclin. cryst. fr. chlf. D.s. abs. alc., eth.; e.s. chlf., bz. — Boiling w. aq. gives CO ₂ , α -nitrostyrene (No. 2.1462), & some indigo.
3186-I	124	Yellow	Isosafroldioximeperoxide, $^FO.O.N:CM_2(H_4C:O_2:C_6H_5)C:N^3$. — [Fr. nitrous ac. & isosafrol.] — Ndl. D.s. alc.; i. alk.; v.s. bz.
3187	125-6	Or.-yel.	p-Aminoazobenzene, $NH_2C_6H_4N_2Ph$. — Rhomb. ndl. or pr. fr. alc. showing bluish reflections. B.p. a. 360° undecd. V.d.s. h. aq.; more s. eth., h. alc. Dyes wool ← Y (T. 2.16). — B.HCl, blue-violet ndl. or scales fr. h. dil. HCl. (Dyestuff described as No. 3.932). — \oplus Reduce 1 g. B.HCl disd. in 20 cc. h. 50% alc. by T. 3.21. The aniline formed (No. 2.1235) is separated by adding 2.5 g. NaOH to redried mixt. & disdg. w. steam; the p-phenylenediamine (No. 2.877) by extraction of alk. residue w. eth.
3188.	abt. 125	Or.-yel.	1-Nitro- β -naphthylamine, $(NO_2)(NH_2)C_{10}H_4$. — \oplus Nitro comp.‡ & prim. amine.¶ — Lust. ndl. S. h. aq.; e.s. alc.

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(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3189	125	Yellowish	3,1 2 -Dinitrostyrene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{CHNO}_2$. — \textcircled{P} Nitro comp.‡ — Lft. fr. alc. R.d.s. h. aq., alc.
3190	125-6	YT1	Quinine Picrate, $\text{C}_{20}\text{H}_{24}\text{O}_4\text{N}_2\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.947. — Fine ndl. fr. gl. ac. ac.
3191	d. 120-30	Y & GYS1	Benzoquinoneoxime, p-Nitrosophenol, $\text{HON:C}_6\text{H}_4:\text{O}$. — Y ndl. fr. bz. + acetone. Small yel.-green lft. fr. chilled bz. sol., becoming GYS1 after drying. Mod. s. aq. (pale green sol.); e.s. alc., eth., acetone, alk.; less s. gl. ac. ac., & still less in hydrocarbons. Color of dil. acetone sol. YT1. Odor, faint, pungent & quinone-like. Taste, sour. \textcircled{P} (1) Gives blue color in Liebermann's react. (T. 2.18), & GYS1 sol. in T. 2.15 w. diphenylamine. — (2) 5 mg. dissd. in 1 cc. 10% NaOH gives OR-RO sol. — (3) Tollen's reagt. in T. 2.30 gives dark-brown sol., but no ppt. — (4) T. 2.21 gives Ag mirror & dark brown ppt. \textcircled{D} Dis. 0.05 g. in 1 cc. HNO_3 (sp. gr. 1.42). A dark color develops & disappears. Add 1 cc. aq. & cool. Filter. Wash ppt. of yellowish lust. lft. w. 2 cc. aq. Recryst. fr. 5 cc. boiling aq., cooling slowly. Filter. Wash w. 1 cc. aq. Dry 15 min. at 100°. The product, 2,4-dinitrophenol, is obtained in YT2 lft., m.p. 113-3.5° u.c.
3191-1	126.5-7	Garnet-red	α -Azoxynaphthalene, $\text{C}_{10}\text{H}_7\text{N}_2\text{O.C}_1\text{H}_7$. — \textcircled{P} S. in conc. H_2SO_4 w. red-violet color, slowly changing to blue. — Rhomb. cryst. fr. slow evapn. of alc. sol.
3192	126-7	Y-YT1	Narceine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.2315. — Fine ndl. fr. h. 10% ac. ac.
3193	126d.	Yellow	Paucine, $\text{C}_{27}\text{H}_{33}\text{O}_4\text{N}_5$. — [Alkaloid fr. nut of Pentaclethra macrophylla fr. Congo region.] — \textcircled{P} FeCl_3 , colors aq. sol. dark green. — Lft. fr. h. aq., soon becoming greenish fr. decompr. S. in aq. NaOH, sol. quickly becoming brown. — Picrate (T. 2.23), garnet-red pr., d.s. c. aq., d. abt. 220°.
3194	126-7	Yellow	β -p-Nitrophenyldiaminodi-p-tolylmethane, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}[\text{C}_6\text{H}_4-\text{Me}(\text{NH}_2)_2]$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Lft. fr. alc. E.s. bz., h. alc.; d.s. lgr. — Diacetyl deriv.** yel. granules fr. dil. alc., m.p. 136°.
3195	126-7	Yellow	3,5-Dinitro-1,2-xylenol(4), $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{Me}_2(\text{OH})$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. alc.
3196	126-7	Lemon-yellow	Trinitro-m-cresol-naphthalene, $[(\text{NO}_2)_3\text{Me.C}_6\text{H}_3\text{OH}]\text{C}_1\text{H}_8$. — Fine ndl. fr. acetone. — \textcircled{P} \textcircled{D} Decd. at once by aq. into its components.
3197	126	Light yellow	cis-o-Dinitrostilbene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{CH.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. gl. ac. ac. I.c. aq. — Dibromide, m.p. 215°.
3198	126-7	Pale yellow(?)	β -Nitrophenanthrene, $\text{NO}_2\text{C}_1\text{H}_8$. — Oxidn. by CrO_3 in gl. ac. ac. gives or.-yel. ndl., m.p. 260-6°.
3199	126	Dark brown-violet; alm. black	Isatine- α -anilide, $[\text{CO.C}_6\text{H}_4\text{N}:\text{C}(\text{NH.Ph})]^+$. — \textcircled{P} NaOH added to yel. brown alc. sol. gives intense blue gradually fading. — Ndl. fr. bz. E.s. h. alc., eth., bz. Sol. in bz. raspberry-red. — Phenylhydrazine yields cinnabar-red phenylhydrazone.
3200	127	Red	Pr-2-phenylindole Picrate, $\text{C}_{16}\text{H}_{11}\text{N.Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.2407. — Pr. fr. alc. Mod. s. alc., eth., bz. Decd. by aq.
3201	127	Light red	Furfurol-p-nitrophenylhydrazone, $\text{C}_6\text{H}_4\text{O.CH:N.H.C}_6\text{H}_4\text{NO}_2$. — \textcircled{P} Hydrazone.††
3202	127	Or.-yel.	2,3-Dinitroaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{NH}_2$. — \textcircled{P} Boiled w. KOH evolves NH, (T. 2.7). — Lust. ndl. fr. alc. E.s. alc.; s. eth.
3203	127	Pale yellow	Tetrinitromethylaniline, "Tetril," $(\text{NO}_2)_4\text{C}_6\text{H}_2\text{NMe}(\text{NO}_2)_2$. — [Powerful explosive.] — \textcircled{P} \textcircled{D} Boiling w. KOH sol. (T.

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
			2.26; Ber., 19, 2126) gives methylamine (No. 2.1059) & picric ac. (No. 2.3168). — Cryst. I. aq.; v.d.s. c. alc. Deflagrates at 186°.
3204	127 sl. d.	Yellow	p-Benzoylphenylhydrazine, Ph.CO.C ₆ H ₄ NH.NH ₂ . — ⊕ Prob. reduces Tollen's reag. in T. 2.30. — Ndl. V.s. alc., eth., bz. — B.HCl, e.s. h. aq., alc.; i. HCl. — Gives hydrazone w. benzaldehyde, tbd., d.s. h. alc., m.p. 188°.
3205	127-8	Lemon-yellow	5-Nitro-o-toluidine, NO ₂ .C ₆ H ₄ .Me(NH ₂). — ⊕ Nitro comp.‡ & prim. amine. ¶ — Ndl. V.d.s. h. aq.
3206	125-6; 129	Red	2,4,2',4'-Tetramethylazobenzene, Me ₂ .C ₆ H ₄ .N ₂ .C ₆ H ₄ .Me ₂ . — ⊕ ⊖ Azo comp. — Ndl. E.s. alc., eth. May be distilled.
3207	128 u.c.	YO-OY	Acetaldehyde-p-nitrophenylhydrazone, Me.CH:N.NH.C ₆ H ₄ -NO ₂ . — ⊕ Hydrazone. §§ — Lust. cryst. fr. h. 30% alc. V.d.s. aq.; e.s. alc., bz., eth.
3208	128	Or.-yel.	4-Amino-3,4'-dimethylazobenzene, (NH ₂)(Me).C ₆ H ₄ .N ₂ .C ₆ H ₄ -Me. — ⊕ ⊖ Azo comp., reduction w. Sn + HCl giving o-toluidine (No. 2.1262), & 2,5-toluenediamine (No. 2.638). — Ndl. fr. lgr. — B.HCl, vermillion ndl.
3209	127-8; 128-9	Yellow-brown	3,4-(or 4,5-)Dinitroveratrol, (NO ₂) ₂ .C ₆ H ₃ .(OMe). ^{1,2} . — ⊕ Nitro comp.‡ — Ndl. 100 pt. 95% alc. dis. 0.3892 pt. at 15°.
3210	128	Yellow	Acetone-2,4-dinitrophenylhydrazone, (NO ₂) ₂ .C ₆ H ₄ .NH.N:CMe ₂ . — ⊕ Hydrazone. §§ — Lust. tbd. fr. alc. D.s. alc., lgr.; e.s. chlf.
3211	128.5	Gold-yel.	Retenequinoneoxime, O:C ₁₂ H ₁₆ :NOH. — ⊕ T. 2.17 gives hydroxylamine & retenequinone (Vol. I). — Ndl. fr. alc.
3212	128	Sulphur-yellow	Benzeneazosalicycaldehyde, Ph.N ₂ .C ₆ H ₄ .(OH)(CHO). — Cryst. fr. alc. E.s. h. alc.; v.s. eth., chlf., bz. — Phenylhydrazone deriv., m.p. 200°.
3213	128	Green-yel.	2-Nitronaphthol(1), NO ₂ .C ₁₀ H ₇ .OH. — ⊕ Nitro comp.‡ — Lft. D.s. dil. alc.; less s. aq. Vol. w. st. Salts are red.
3214	129	Light red	3,5-Dinitromethyl-p-toluidine, (NO ₂) ₂ .Me.C ₆ H ₄ .NHMe. — ⊕ Nitro comp.‡ — Ndl. fr. alc. — Nitroso deriv., m.p. 128°.
3215	129	Brown	4-Nitro-3-aminophenolmethyleneether, (NO ₂)(NH ₂).C ₆ H ₃ .OMe. — ⊕ Nitro comp.‡ & prim. amine. ¶ — Sbl. in yel. lft.
3216	129	Or.-red	p-Nitrohippuric Ac., NO ₂ .C ₆ H ₄ .CO.NH.CH ₂ .CO ₂ H. — ⊕ Nitro comp.‡ & acid. Pr. fr. h. aq., separating at first in oily drops. E.s. h. aq., alc., eth.; r.d.s. c. aq. — BaA ₂ .4H ₂ O, long yellowish ndl.
3217	128-30	Golden	3-Methyl-4-hydroxyazobenzene, Me(OH).C ₆ H ₄ .N ₂ .Ph. — ⊕ ⊖ Azo comp. — Cryst. I. c. aq.; d.s. h. aq.; e.s. alc., eth., chlf., bz.; less s. lgr.; e.s. dil. alk. — Acetate, yel. tbd. fr. dil. alc., m.p. 81-2°.
3218	129	Green	Benzyl-p-nitrosoaniline, Ph.CH ₂ .NH.C ₆ H ₄ .NO. — Lft. w. bluish reflections fr. alc. D.s. c. alc., eth.; e.s. bz. Boiled w. NaOH gives NH ₃ , benzaldehyde, p-aminophenol, & p-nitrophenol.
3219	130	Red-brown	Pentamethylrosaniline, (Me ₂ N.C ₆ H ₄) ₂ .C(OH)(C ₆ H ₄ .NHMe). — ⊕ S. in HCl w. red-violet color. — Melts under aq. I. aq., lgr., eth.; s. alc. w. violet color. Dyes tannin-mordanted cotton intense violet! Dry salts, amorph. brassy colored masses.
3220	130	Orange	2,3,5-Trinitromethyl-p-toluidine, (NO ₂) ₃ .Me.C ₆ H ₄ .NHMe. — Ndl. fr. alc. V.s. bz., chlf.
3221	130	Yellow	p-Nitrophenyldibenzylamine, NO ₂ .C ₆ H ₄ .N(CH ₂ .Ph) ₂ . — ⊕ Nitro comp.‡ — Ndl. fr. alc. D.s. c. alc.; e.s. eth., bz.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34.
¶ = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3222	129; 131	Yellow	m-Nitrodiazoaminobenzene, $\text{NO}_2\text{C}_6\text{H}_4\text{NH}_2\text{Ph}$. — Thick pr. fr. eth.
3223	130	Gold-yel.	2,3,4,6-Tetranitrophenol, $(\text{NO}_2)_4\text{C}_6\text{H}_2\text{OH}$. — \oplus Nitro comp.‡ — Ndl. w. violet reflections fr. ac.-eth. Sometimes explodes in melting. E.s. aq.; v.d.s. bz., lgr. Highly explosive! Colors wool (T. 2.16) deep reddish yel.
3224	131	Red	o-Tolueneazo- β -naphthol, $\text{Me.C}_6\text{H}_4\text{N}_2\text{C}_10\text{H}_7\text{OH}$. — \oplus \ominus Azo comp. — Lust. cryst. fr. gl. ac. ac. I. c. dil. NaOH.
3224-1	131	Red	Nitro-5-amino-1,2,3,4-tetramethylbenzene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{Me}_4$. — \oplus Nitro comp.‡ & prim. amine.¶ — Lust. ndl. S. conc. HCl, but reptd. by aq.
3225	131	Garnet-red	o-Azophenoldiethylether, o-Azophenetole, $\text{EtO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OEt}$. — \oplus Azo comp. — B.p. 240° d. I. aq. E.s. conc. HCl, but reptd. by aq.; s. alc., eth.
3226	131-2	Yellow	Methyl-p-tolylketonehydrazone, $\text{Me.C}(\text{N.NH}_2)\text{C}_6\text{H}_4\text{Me}$. — \oplus Hydrazone.†† — Cryst. fr. alc. D.s. alc., eth.; e.s. h. gl. ac. ac.
3227	131	Yellowish	1,2-Naphthacridine, $\text{C}_{17}\text{H}_{11}\text{N}$. — \oplus Conc. H_2SO_4 gives green fluor. sol.! — Cryst. E.s. alc., w. yel. color & blue fluor.! S. bz., eth.
3228	131.5	V. pale yellow	o-Hydroxybenzylanilinenitrosamine, $\text{HO.C}_6\text{H}_4\text{CH}_2\text{N}(\text{NO})\text{Ph}$. — \oplus Nitrosamine.§ — Pr. fr. dil. MeOH. D.s. c. lgr.; mod. s. alc. — Reducn. w. Zn dust & ac. ac. gives a hydrazine.
3229	131.5; 134c.	Yellowish	2-Methylacridine, $\text{Me.C}_6\text{H}_4\text{N}$. — \oplus Vapors provoke sneezing. Sol. in dil. H_2SO_4 fluor. blue-green. — E.s. alc., eth., bz. — $\text{B}_2\text{H}_6\text{PtCl}_6$, yel., d.s. aq.
3230	132	Yellow	Glycericaldehydephenylosazone, $\text{CH}_2(\text{OH}).\text{C}(:\text{N.NHPh}).\text{CH}(:\text{N.NHPh})$. — Lft. E.s. alc., eth., bz. Decd. at 170°. Acts as reducing agent.
3231	130; 133	Pale greenish yellow	Methylauramine, $\text{MeN.C}(\text{C}_6\text{H}_4\text{NMe})_2$. — \oplus Dyes tannin-mordanted cotton yel. — Cryst. fr. alc. Alm. i. aq.; e.s. alc. — B.HCl , yel. lft., d.s. c. aq., e.s. alc., m.p. 225°. — B.Pk , yel.-red lft., d.s. alc., d. 225°. [Ethylauramine is said to form colorless pr. fr. alc. & to melt at 130-1°.]
3232	133	Red	Naphthalantracene Picrate, $\text{C}_{14}\text{H}_{10}.2\text{Pk}$. — \oplus \ominus Picrate* of h.c. of Vol. I. Ndl. fr. bz. Decd. by alc.
3233	133; 134; 129	Red-gold	Benzeneazonaphthol(2), $\text{Ph.N}_2\text{C}_6\text{H}_4\text{OH}$. — \oplus Azo comp. s. in conc. H_2SO_4 w. fuchsine-red color! — Cryst. w. beetle-wing reflections fr. alc. S. eth., bz., lgr.; i. NaOH. — Ac. anhydride gives acetate, dark orange scales, m.p. 117°.
3234	133	Yellow or orange	p-Nitrodiphenylamine, $\text{NO}_2\text{C}_6\text{H}_4\text{NHPH}$. — \oplus Nitro comp.‡ — Lft. fr. alc.
3235	133-4	Steel-blue	Phenol Blue, Phenolindophenol, $\text{O:C}_6\text{H}_4:\text{N.C}_6\text{H}_4\text{NMe}_2\text{H}_2\text{O}$. — \oplus S. HCl w. blue color. — Pr. fr. aq. — For spectrum, cf. Ann., 289, 129.
3236	134-5	Ruby-red	Formazylimethylketone, $\text{Me.CO.C}(\text{N:NPh})(:\text{N.NHPh})$. — Pr. w. steel-blue reflections fr. alc. D.s. alc.; e.s. bz.
3237	134-5	Garnet-red	p-Tolueneazonaphthol(2), $\text{Me.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_3\text{OH}$. — \oplus Azo comp. — Cryst. w. green metallic luster fr. gl. ac. ac. E.s. alc., bz.; i. dil. NaOH. — Acetate,** deep red pr. fr. bz., m.p. 99°.
3238	134	Or.-yel.	p-Nitrobenzalhydrazine, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{N.NH}_2$. — \oplus Alc. sol. colored red by HgO . — Pr. fr. alc. Alc. sol. w. acid yields p,p'-dinitrobenzalzin, yel. ndl. fr. ac. ac., m.p. 290°.
3239	134.5	Gold-yel.	Xanthonephenylimide, $[\text{O.C}_6\text{H}_4\text{C}(\text{N:NPh})\text{C}_6\text{H}_3]^2$. — \oplus Yel. sol. in conc. HCl fluor. green, & on boiling w. aq. gives aniline & xanthone. — Cryst. fr. alc. I. alk.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3240	134	Yellow	4-Nitrocoumarone, $\text{C}_6\text{H}_3(\text{NO}_2)\cdot\text{CH}:\text{CH}\cdot\text{O}^2$. — Ndl. fr. dil. alc.
3241	134	Yellow	p-Dihydroxyazoxybenzenediethylether, $\text{EtO}\cdot\text{C}_6\text{H}_4\cdot\text{N}_2\text{O}\cdot\text{C}_2\text{H}_5\cdot\text{OEt}$. — \textcircled{P} Melts to turbid liq. which clears at 165°.
3242	134	Yellow	m-Nitroformanilide, $\text{NO}_2\cdot\text{C}_6\text{H}_4\cdot\text{NH}(\text{CHO})$. — \textcircled{P} Nitro comp.‡ — D.s. eth., lgr.
3243	134-5d.	Pale yellowish	5-Nitrosoresol(2), Toluquinoneoxime, $\text{HON}:\text{C}_6\text{H}_4\text{Me}:\text{O}$. — \textcircled{P} Prob. gives Liebermann's react. (T. 2.18). — Long ndl. fr. aq. D.s. c. aq.; e.s. h. aq.; e.s. alc., eth., chlf.; less s. bz.
3244	135-6d.	Dark red	3-Nitro-4-aminophenol, $\text{NO}_2\cdot\text{C}_6\text{H}_3\cdot(\text{NH}_2)(\text{OH})$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. aq. E.s. h. aq., alc.; d.s. lgr. — Reduced by SnCl_2 & HCl (T. 2.40) to No. 2.932.
3245	135	Straw-yellow	4,8-Dinitronaphthol(1), $(\text{NO}_2)_2\cdot\text{C}_{10}\text{H}_4\cdot\text{OH}$. — \textcircled{P} Nitro comp.‡ — Ndl. D.s. aq.; s. org. solvents.
3246	135-6d.	Golden	2,5-Dinitrohydroquinone, $(\text{NO}_2)_2\cdot\text{C}_6\text{H}_3\cdot(\text{OH})_2\cdot\text{H}_2\text{O}$. — \textcircled{P} Sol. in ammon. purple-red after boiling, leaving cryst. w. green metallic luster on evapn. — Cryst. fr. aq. D.s. c., e.s. h. aq.; s. alc., eth. — Oxidn. by HNO_3 gives nitranilic ac. (2.3469).
3247	135	Yellow	Trinitro diphenylamine, $(\text{NO}_2)_3\cdot\text{C}_{12}\text{H}_7\cdot\text{NH}$. — Ndl. fr. gl. ac. ac.; e.s. alc., bz.
3248	136-7	Orange	3,5,3',5'-Tetramethylazobenzene, $\text{Me}_2\cdot\text{C}_6\text{H}_3\cdot\text{N}_2\cdot\text{C}_6\text{H}_3\cdot\text{Me}_2$. — \textcircled{P} Azo comp. — Ndl. E.s. alc.
3249	136	Dark brown	$\alpha\beta$ -Azonaphthalene, $\text{C}_{10}\text{H}_7\cdot\text{N}_2\cdot\text{C}_{10}\text{H}_7$. — \textcircled{P} Azo comp. s. conc. H_2SO_4 w. violet color. — Lft. fr. gl. ac. ac. w. steel-blue reflections.
3250	136	Brownish yellow	2,4,6-Trinitro-m-toluidine, $(\text{NO}_2)_3\cdot(\text{NH}_2)\cdot\text{C}_6\text{H}_4\cdot\text{Me}$. — \textcircled{P} Boiled w. NaOH sol. gives NH_3 (T. 2.7) & trinitroresol (No. 2.3086). — S. c. alk. w. red color! Cubes fr. eth. E.s. alc., eth.
3251	136.5	Gold-yel.	o-Nitrodicinnamylvinylketone, $\text{Ph}\cdot\text{C}_6\text{H}_4\cdot\text{CO}\cdot\text{C}_6\text{H}_4\cdot\text{C}_6\text{H}_4\cdot\text{NO}_2$. — \textcircled{P} Nitro comp.‡ Cryst. fr. acetone. D.s. alc., eth.; e.s. chlf., bz.
3252	136	Lemon-yellow	Auramine, $\text{HN}:\text{C}(\text{C}_6\text{H}_4\cdot\text{NMe}_2)_2$. — [Cf. No. 3.1085 for description of the commercial dyestuff, which is the hydrochloride of this species.] — Lft. fr. alc. I. aq. — $\text{B}\cdot\text{HCl}\cdot\text{H}_2\text{O}$, yel. lft., m.p. 267-8°, d.s. c. aq.; dyes tannin mordanted-cotton pure yel. — $\text{B}\cdot\text{Pk}$ (T. 2.23), yel. lft., i. c. aq., d.s. c. alc., m.p. 230-6°.
3253	137; (134-5)	Nearly black	2-Nitro-p-phenylenediamine, $\text{NO}_2\cdot\text{C}_6\text{H}_4\cdot(\text{NH}_2)_2$. — Ndl. w. green reflections.
3254	137	Or.-yel.	2,5-Dinitroaniline, $(\text{NO}_2)_2\cdot\text{C}_6\text{H}_3\cdot\text{NH}_2$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Silky ndl. fr. alc. E.s. alc.
3255	137	Gold-yel.	3-Nitro-5-amino-1,2,4-trimethylbenzene, $(\text{NO}_2)_2\cdot(\text{NH}_2)\cdot\text{C}_6\text{H}_3\cdot\text{Me}_3$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Ndl. D.s. h. aq.; e.s. alc. Vol. w. st.
3256	137	Yellow	5-Nitromethyl-o-toluidine, $(\text{NO}_2)\text{Me}\cdot\text{C}_6\text{H}_4\cdot\text{NHMe}$. — Nitro comp.‡ & sec. amine. — Tbl. fr. alc. Alm. i. dil. H_2SO_4 .
3257	137.5	Yellow	3,5-Dinitrohydro-p-cumaric Ac., $(\text{NO}_2)_2\cdot(\text{HO})\cdot\text{C}_6\text{H}_3\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{CO}_2\text{H}$. — \textcircled{P} Nitro comp. & acid. — Fern-like lft. fr. aq. V.d.s. c. aq.; e.s. h. alc. — $\text{Ag}\bar{\text{A}}$, ppt., e.s. h. aq., crystg. on cooling in red-yel. ndl. — $\text{Me}\bar{\text{A}}$ (fr. $\text{Ag}\bar{\text{A}}$ & MeI), ndl. fr. dil. alc., m.p. 87°.
3258	137-8	Y	Quinidine Picrate. — \textcircled{P} Picrate* of No. 2.935. — Cryst. fr. h. 95% alc.
3259	138	Deep red	2-Benzeneazonaphthol(1), β -Naphthoquinonephenylhydrazone, $\text{HO}\cdot\text{C}_{10}\text{H}_4\cdot\text{N}_2\cdot\text{Ph}$. — \textcircled{P} Sbl. in or.-red ndl. w. green reflections. S. conc. H_2SO_4 w. violet-red color. — I. aq.; s. h. alc.;

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(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
			alm. i. dil. ac. or alk. — Acetate, silky or. ndl. fr. bz., m.p. 120-1 $^{\circ}$.
3200	138	Red	Tetramethylnaphthalene Picrate, $C_{14}H_{16}Pk$. — \textcircled{P} Picrate* of h.c. fr. petroleum of m.p. -20 $^{\circ}$ & b.p. 320 $^{\circ}$. — Ndl.
3261	138	Saffron-yellow	6-Nitro-m-toluidine, $(NO_2)Me.C_6H_4.NH_2$. — \textcircled{P} Nitro comp.† & prim. amine. — Ndl. E.s. ac. giving unstable salts. Not vol. w. st.
3262	138	Yellow	2,6-Dinitroaniline, $(NO_2)_2C_6H_4.NH_2$. — \textcircled{P} Nitro comp.‡ & prim. amine. — Long ndl. S. in 192 pt. 95% alc. at 21 $^{\circ}$.
3263	138	Yellow	2,4,6-Trinitrodimethylaniline, $(NO_2)_3C_6H_3.NMe_2$. — \textcircled{P} Nitro comp.‡ — Tbl. fr. bz. S. h. alc.; d.s. eth.
3264	133; 144	Light yellow	2,5-Dinitrosotoluene, $(NO)_2C_6H_5.Me$. — \textcircled{P} Odor quinone-like (pungent). E. vol. w. st. I., except in gl. ac. ac. — Fuming HNO ₃ oxid. to dinitrotoluene (No. 2.1549).
3265	138.5	Pale yellow	p-Aminoazoxybenzene, $NH_2.C_6H_4.N.O.Ph$. — \textcircled{P} Distn. gives aniline (No. 2.1235), & azobenzene (No. 2.2935). — Rhomb. tbl. fr. dil. alc. 100 pt. aq. at 21 $^{\circ}$ dis. 4.20 pt. S. alc.; less s. eth. — Reduced by Sn + HCl to aniline & p-phenylenediamine (No. 2.877).
3266	138	Lemon-yellow	2,4-Dinitronaphthol(1), $(NO_2)_2C_{10}H_8.OH$. — [Alk. salts form commercial dyestuff, Martius' or Naphthol Yellow described as No. 3.945-1] — \textcircled{P} Dyes wool Y-, completely discharged by Rongalite in T. 2.16-a, b. — Ndl. fr. alc. Alm. i. h. aq.; d.s. alc., eth., bz. Not vol. w. st.
3267	139	Brown-red	5-Nitro-4-amino-1,2-xylene, $(NO_2)(NH_2).C_6H_4.Me_2$. — \textcircled{P} Nitro comp.† & prim. amine. — Pr. fr. alc. — Acetyl deriv.** pale yel. ndl., m.p. 107 $^{\circ}$.
3266	139-40	Gold-yel.	Nitrotriphenylamine, $NO_2.C_6H_4.N.Ph_3$. — Lust. lft. fr. dil. ac. ac. I. aq.
3269	137; 139-40	Yellow	5-Nitro-6-hydroxyquinoline, $NO_2.C_9H_4.N.OH$. — \textcircled{P} Dil. alc. sol. colored reddish by FeCl ₃ . Dec. carbonates. — Ndl. I. aq., chlf. eth.; e.s. alk., min. acids, h. alc. — Ba salt, silky or. red filaments, d.s. c. aq.
3270	140-1	Red	3,4,3',4'-Tetramethylazobenzene, $Me_2C_6H_4.N_2C_6H_4.Me_2$. — \textcircled{P} Azo comp. — Ndl.
3271	140-1	Lemon-yellow	Acetyl-o-nitrophenylhydrazine, $(C_6H_5O)NH.NH.C_6H_4.NO_2$. — Ndl. fr. alc. E.s. h. aq., alc., bz.
3272	140	Light yellow	Phthalimidine Picrate, C_9H_7ONPk . — \textcircled{P} \textcircled{D} Picrate* of No. 2.2195. — Tbl. or pr. fr. alc. D.s. c. aq., alc., bz.
3273	140	Light yellow	2-Benzylpyridine Picrate, $C_{10}H_{11}N.Pk$. — \textcircled{P} \textcircled{D} Picrate of No. 2.1430. — Cryst. fr. h. alc.
3274	140	Green	5-Nitrosoethyl-o-toluidine, $(NO)Me.C_6H_4.NHET$. — Lft. w. blue reflections fr. bz.
3275	141-2	Red	Benzeneazo- β -phenylnaphthylamine, $Ph.N_2.C_{10}H_4.NHPh$. — \textcircled{P} Azo comp. Ndl. of metallic luster fr. gl. ac. ac. — Boiled w. 5 pt. gl. ac. ac. + 1-1½ pt. conc. HCl gives aniline & $\alpha\beta$ -naphthophenazine (No. 2.3291).
3276	141-2	Ruby-red	Formazylphenylketone, $Ph.CO.C(N:NPh):N.NHPh$. — \textcircled{P} Ndl. w. metallic luster s. in conc. HCl w. red-violet color. D.s. c. alc.; e.s. eth., chlf., bz.
3277	141	Dark red	N-Methylcarbazole Picrate, $C_8H_7N.Pk$. — \textcircled{P} \textcircled{D} Picrate of No. 2.1764. — Ndl. S. alc.
3278	141	Light red	o-Toluidonaphthoquinone(1,4), $C_7H_7NH.C_{10}H_8O_2$. — Ndl. fr. alc. I. alk.
3279	141	Orange	1,4-(α)-Dimethylnaphthalene Picrate, $C_{12}H_{12}Pk$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3280	141	Dark yellow	Di-o-nitrobenzylnitromethane, $(NO_2.C_6H_4.CH_3)_2.CH.NO_2$. — Cryst. D.s. aq.; alc., eth.; e.s. chlf.
3281	141	Yellow-brown	Piperonyleneacetonephenylhydrazone, $CH_2:O.C_6H_4.CH:CH-CH:CH.CMe:N.NHPH$. — $\textcircled{P} \textcircled{D}$ Hydrazone. \ddagger — Ndl. fr. alc.
3282	141	Y	Narcotine Picrate. — $\textcircled{P} \textcircled{D}$ Picrate* of No. 2.952.
3283	141	Yellow	N-Acetylisatine, $C_6H_4O_2.N.C_6H_4.O$. — Ndl. fr. bz. — Boiling w. HCl gives ac. ac. & isatine (No. 2.3633).
3284	141-2	Yellow	p-Nitrobenzil, $NO_2.C_6H_4.O_2$. — Lft. fr. eth. S. in 30 pt. boiling 85% alc.; more s. eth., chlf.
3285	142-3	Red-yellow	α -Dinaphthylmethane Picrate, $C_{12}H_{10}.2Pk$. — $\textcircled{P} \textcircled{D}$ Picrate* of h.c. s. in 15 pt. boiling alc., m.p. 109°. — Pr. fr. chlf.
—	142	Orange	Methyl 3-Amino-4-hydroxybenzoate. — Cf. No. 2.784.
3286	142-3	Orange	4-Nitro-2-aminophenol, $(NO_2)(NH_2).C_6H_4.OH$. — \textcircled{P} Taste v. sweet. — Cryst. w. $1H_2O$ in pr. of m.p. 80-90°. D.s. c. aq.; v.s. alc., eth. — Ag \bar{A} .H \bar{A} , yel.-brown ppt. crystg. in lft.
3287	142-3	Or.-yel.	2,6-Dimethylnaphthalene Picrate, $C_{12}H_{12}.Pk$. — $\textcircled{P} \textcircled{D}$ Picrate* of h.c., lft. fr. alc. w. orange flower odor, m.p. 110-1°. — Ndl. fr. alc.
3288	142	Brownish-yellow	5-Nitro-2-amino-1,4-xylene, $(NO_2)(NH_2).C_6H_3.Me_2$. — \textcircled{P} Nitro comp. \ddagger & prim. amine. \ddagger — E.s. alc., eth.; d.s. lgr.
3288	140-2; 142.7c.	Amber-yellow	m-Nitrobenzamide, $NO_2.C_6H_4.CONH_2$. — $\textcircled{P} \textcircled{D}$ Sapn. (T. 2.26) products: m-Nitrobenzoic ac. (No. 2.139) & NH $_4$. — Ndl. fr. aq. B.p. 310-5°.
—	142.5	Broken light yellow	Colchicine. — Cf. No. 2.2152.
3290	142	Light yellow	2,4-Dinitroresorcinol, $(NO_2)_2.C_6H_3.(OH)_2$. — [“Solid Green” (No. 3.1131).] Lft. fr. dil. alc. V.s. alc. D.s. c. aq., alc.; i. eth., bz. A rather strong acid. Deflagrates when strongly heated. Warming w. dil. HNO $_3$ gives trinitroresorcinol (No. 2.3419). — K \bar{A} .H \bar{A} O, yel. ndl., v.s. aq., i. abs. alc.
3291	142.5	Lemon-yellow	$\alpha\beta$ -Naphthophenazine, $C_{10}H_8N_2$. — Pr. fr. bz. Sbl. in long flat ndl. V.d.s. alc., eth., bz. Salts w. acids decd. by aq.
3292	142d.	Yellow	o-Benzoquinonedioxime, $HON:C_6H_4:NOH$. — \textcircled{P} Sol. in alk. blood-red! — Ndl. S. aq., bz.
3293	142-3	Yellow	6-Nitrocresol(2), $(NO_2)(Me).C_6H_4.OH$. — \textcircled{P} Taste intensely sweet! — Woolly ndl. fr. aq. V.d.s. c. aq.; e.s. alc., eth.
3294	143.5	Red	5-Nitro- β -naphthylamine, $NO_2.C_10H_8.NH_2$. — \textcircled{P} Nitro comp. \ddagger & prim. amine. \ddagger — Ndl. fr. alc. E.s. h. alc., bz. — Benzoyl deriv., m.p. 181.5°.
3295	143.5	Or.-red	3,2',4'-Triaminoazobenzene, $NH_2.C_6H_4.N_2.C_6H_4.(NH_2)_2$. — [In commercial Bismarck Brown.] Azo comp. \parallel & prim. amine, \ddagger dyeing wool yel. or red-yel. changing to red-brown when dipped in HCl. — Cryst. w. $\frac{1}{2}C_6H_4$ fr. bz. E.s. alc., eth.; i. lgr. — Boiled 15 min. w. 1 pt. NaA + 20 pt. ac. anhydride gives triacetyl deriv., fine yel. ndl. fr. abs. alc., m.p. 264°.
3295	143	Chrome-red	2-Nitrobenzidine, $(NO_2)(NH_2).C_6H_4.C_6H_4.NH_2$. — Ndl. — Reduced by Sn + HCl to amino deriv. of m.p. 134°.
3297	143u.c.	OY	Phenanthrene Picrate, $C_{12}H_{10}.Pk$. — $\textcircled{P} \textcircled{D}$ Picrate* of h.c. of Vol. I. — Long hair-like ndl. fr. h. alc. S. in abt. 37 pt. alc. at 15°; less s. in presence x.s. picric ac.; e.s. eth., bz.
3298	143	Yellow	3-Nitro-p-benzoyltoluide, $NO_2.C_6H_5.Me(NH.CO.Ph)$. — \textcircled{P} Nitro comp. \ddagger Long ndl. E.s. gl. ac. ac., h. alc.
3299	143	Light yellow	Nitro-o-acetanisidide, $(NO_2).C_6H_4.(MeO)(NH.C_6H_4.O)$. — \textcircled{P} Nitro comp. \ddagger — Ndl. fr. alc.

Explanation of typographical signs used in this Division: * = T. 2.39. \ddagger = T. 2.21. \ddagger = T. 2.36. \parallel = T. 2.34.
 \ddagger = T. 2.35. $\ddagger\ddagger$ = T. 2.1. $\ddagger\ddagger$ = T. 2.17.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3300	143 u.c.	Green or steel-blue	p-Nitrosophenylaniline, $\text{NO.C}_6\text{H}_4.\text{NHPH}$. — (P) Sol. in conc. H_2SO_4 , O-RO; in bz., green; in alc., OYSI. — Tbl. fr. bz. brilliant dark steel-blue by reflected, & green by transmitted light. O-colored sol. in NaOH gives aniline, etc., on boiling.
3301	144.5	Purple-red	o-Azobiphenyl, $^{\text{C}}\text{C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_4$. — Ndl. fr. alc.
3302	144	Red-yel.	2-Nitro- α -naphthylamine, $\text{NO}_2.\text{C}_{10}\text{H}_8.\text{NH}_2$. — (P) Boiled w. KOH sol. gives NH_3 (T. 2.7) & 2-nitronaphthol(1) (No. 2.3213). — Monoclin. pr. fr. alc.
3303	144	Or.-yel.	4,4'-Azotoluene, $\text{Me.C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_4.\text{Me}$. — (P) (D) Azo comp. — Ndl. fr. lgr. E.s. eth., lgr.; less s. alc.
3304	144-5	Gold-yellow	8-Nitronaphthol(2), $\text{NO}_2.\text{C}_{10}\text{H}_8.\text{OH}$. — (P) Nitro comp.‡ & phenol. — Long ndl. fr. aq.; e.s. alc.
3305	144	Yellow	2,3-Dinitrophenol, $(\text{NO}_2)_2.\text{C}_6\text{H}_4.\text{OH}$. — (P) Nitro comp.‡ & phenol. — Ndl. fr. aq. V.s. eth., h. alc. — Methyl ether, m.p. 118°.
3306	141; 143; 144-6	Yellow	p-Aacetaminoazobenzene, $(\text{C}_2\text{H}_5\text{O})\text{NH.C}_6\text{H}_4.\text{N}_2.\text{Ph}$. — (P) (D) Azo comp. reducing w. SnCl_2 & HCl mainly to aniline (No. 2.1235) & p-phenylenediamine (No. 2.877). — Ndl. fr. dil. alc.
3307	144	Yellow	4-Nitro-1,1-diacetaminonaphthalene, $\text{NO}_2.\text{C}_{10}\text{H}_8.\text{N}(\text{C}_2\text{H}_5\text{O})_2$. — Boiled w. alc. ammon. gives monoacetyl deriv. of m.p. 190°.
3308	144	Light yellow	1,3-(γ)-Dinitronaphthalene, $(\text{NO}_2)_2.\text{C}_{10}\text{H}_8$. — (P) Nitro comp.‡ — Ndl. fr. alc. Sbl. in small ndl.
—	143-5	Yellowish	Glyoxylic acidphenylhydrazone. — Cf. No. 2.149.
3309	144	Yellowish	Acet-3,4-dinitroanilide, $(\text{C}_2\text{H}_5\text{O})\text{NH.C}_6\text{H}_4.(\text{NO}_2)_2$. — (P) Nitro comp.‡ — Rhomb. cryst. E.s. h. alc.
3310	144-6	Red	2-O-Tolueneazonaphthol(1), $\text{Me.C}_6\text{H}_4.\text{N}_2.\text{C}_{10}\text{H}_8.\text{OH}$. — (P) (D) Azo comp. — E.s. w. dark red color in alc., bz.; less s. lgr. Sol. colored blue by strong acids! — Lust. ndl.
3311	145	Ruby-red	4-Methylbenzene-azo- α -naphthylamine, $\text{Me.C}_6\text{H}_4.\text{N}_2.\text{C}_{10}\text{H}_6.\text{NH}_2$. — (P) (D) Azo comp. — Lft. I. aq.; s.h. alc., bz. — B.HCl , black-blue ndl., m.p. 162-4°.
3312	145	Red-brown	$\alpha\alpha$ -Binaphthyl Picrate, $\text{C}_{20}\text{H}_{14}.2\text{Pk}$. — (P) (D) Picrate* of h.c. of Vol. I. — Ndl. fr. bz. Dec. in air.
3313	abt. 145d.	Yellow or orange	Berberine, $\text{C}_{20}\text{H}_{17}\text{O}_4\text{N}$. — [Alkaloid in Berberis vulgaris, Hydrastis canadensis, etc.] Color varies w. method of preparation. Cryst. w. $6\text{H}_2\text{O}$. Melts to brown amorph. mass. S. in 4.5 pt. aq. (?) at 21° (J. Chem. Soc., 38, 169) w. yel. or or. color; s. in 100 pt. c. aq.; e.s. h. aq.; alm. i. eth., bz., lgr. — (P) (1) Salts have intense yellow color & extremely bitter taste! — (2) Sol. of I in KI gives brown ppt. of B.HI.I_2 , crystg. fr. h. alc. in long brown adamanantine ndl., i. aq. or c. alc. — (3) Warmed w. conc. H_2SO_4 gives olive-green color (Delicacy, 1:10,000). — (4) Fragment of NaNO_2 , stirred into sol. in conc. H_2SO_4 gives violet streak (Delicacy 1:10,000). — (5) Cl-aq. poured upon v. dil. aq. sol. mixed w. equal vol. conc. H_2SO_4 gives purple zone lasting for a day. (Distinct w. 1:100,000 sol.). — (6) B.HNO_3 soon separates in yel. ndl. when 3 cc. HNO_3 (sp. gr. 1.185) is added to 10 cc. dil. aq. sol. of B. — [Cf. Arch. Pharm., 1902, 146 for procedure for detection in plants & additional reactions.]
3314	145	Gold-yellow	Acetyl-m-nitrophenylhydrazine, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NH.NH}(\text{C}_2\text{H}_5\text{O})$. — (P) Should reduce Tollen's reagt. after sapon. (T. 2.26 & 2.30). — Lft. fr. h. aq. E.s. h. aq.; c. alc.
3315	145	Lemon-yellow	2,4,7-Trinitronaphthol(1), $(\text{NO}_2)_3.\text{C}_{10}\text{H}_8.\text{OH}$. — (P) Nitro comp.‡ pr. fr. bz. or alc. contain solvent and which effloresce in air. — Na salt, S-yel. ndl. alm. i. c. aq. — Oxid. by HNO_3 (Cf. T. 1.905-3) gives 4-nitrophthalic ac. (No. 2.215).

(ORDER II.)

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
3316	145	Intensely yellow	Desyl-p-toluide, $\text{Ph.CO.CH}(\text{NH.C}_6\text{H}_4.\text{Me})\text{Ph}$. — \textcircled{P} \textcircled{O} Sapn. by alc. KOH (T. 2.26) gives benzoic ac., etc. — Ndl. fr. alc. D.s. c. alc.; eth.; e.s. bz.
3317	145	Yellow	4,4'-Diaminohydrazobenzene, Diphenine, $\text{NH}_2.\text{C}_6\text{H}_4.\text{NH.NH-C}_6\text{H}_4.\text{NH}_2$. — [Possibly colorless when pure.] Cryst. D.s. c. aq.; s. h. aq.; e.s. alc., eth. Oxidn. gives benzoquinone. — B_2HCl , red scales d.s. aq.
3318	145-8	Garnet-red	Benzeneazohydroquinone, $\text{Ph.N}_2.\text{C}_6\text{H}_4(\text{OH})_2$. — Ndl. fr. dil. ac. ac. S. alc.; e.s. eth., bz.; i. lgr.
3319	146.5 u.c.	Yellow	p-Nitroaniline, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NH}_2$. — \textcircled{P} Nitro comp. & prim. amine. Ndl. fr. dil. alc. S. in 1250 pt. aq. at 18.5°, or 45 pt. boiling; s. alc., eth. Not vol. w. st. Odorless. Tasteless. — \textcircled{O} Heat to boiling 4 drops ac. anhydride + 0.05 g. substance. Boil w. 10-15 cc. aq. Filter. Wash w. 2 cc. c. aq. Recryst. product fr. 5 cc. h. aq. Dry on tile 15 min. at 100°. p-Nitroacetanilide, the product, forms colorless ndl., m.p. 207° u.c.
3320	146	Yellow	9-Nitroanthracene, "Nitrosoanthron," $\text{C}_{10}\text{H}_7.\text{NO}_2$. — Long ndl. fr. alc. V.s. bz., CS_2 ; i. alk. — \textcircled{O} Oxidn. by CrO_3 in gl. ac. ac. sol. gives anthraquinone (T. 1.1011).
3321	146d.	Yellowish	p-Acetaminohydrazobenzene, $\text{Ph.NH.NH.C}_6\text{H}_4.\text{NH(C}_2\text{H}_5\text{O})$. — Lft. fr. dil. alc. I. aq.; e.s. alc., eth. — Prob. reduces Tollen's reagt. in T. 2.30, being oxid. to corresponding azo comp. by air.
3322	146-7; 148	Yellow	2-Ethylquinoline Picrate, $\text{C}_{11}\text{H}_{11}.\text{N.Pk}$. — \textcircled{P} \textcircled{O} Picrate* of No. 2.1398. — Ndl. D.s. aq.
3323	147-8	Yellow	(γ)-2,5-Dinitro, 1,4-xylene, $(\text{NO}_2)_2.\text{C}_6\text{H}_3.\text{Me}_2$. — \textcircled{P} Nitro comp. — Long glassy ndl. fr. alc. D.s. c. alc., eth. — \textcircled{O} Reduced to No. 2.3288 by alc. $(\text{NH}_4)_2\text{S}$.
3324	147; 142-3	Gold-yellow	p-Nitrophenylbenzylamine, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NH.CH}_2.\text{Ph}$. — \textcircled{P} Nitro comp. — Plates fr. dil. alc. E.s. alc. — Nitroso deriv., m.p. 107.5°.
3325	147	Light yellow	5-Nitronaphthol(2), $\text{NO}_2.\text{C}_{10}\text{H}_4.\text{OH}$. — \textcircled{P} Nitro comp. — Ndl. fr. aq. E.s. alc.
3326	147	Yellowish	Dinitrobenzil, $(\text{NO}_2)_2.\text{C}_{14}\text{H}_8\text{O}_2$. — \textcircled{P} Nitro comp. — Lft. S. in 52.5 pt. boiling, or 290 pt. c. alc. [An octahedral form, No. 2.2084, melts at 131°.]
3320	148-8.5	Gold-yellow	Acetone-p-nitrophenylhydrazone, $\text{Me}_2\text{C}:\text{N.NH.C}_6\text{H}_4.\text{NO}_2$. — \textcircled{P} Hydrazone. — Ndl. fr. alc. S. h. aq.; e.s. alc.
3329	148	Yellow	Methylglyoxalosazone, $\text{Me.C}(\text{N.NH.Ph}).\text{CH}:\text{N.NHPH}$. — \textcircled{P} Sol. in conc. H_2SO_4 , olive-green, changing to slate-blue. — Fine ndl. fr. dil. alc. D.s. h. alc.
3330	148-9	Leather-yellow	Methylglyoxalphenylhydrazone, $\text{Me.CO.CH}:\text{N.NHPH}$. — \textcircled{P} Hydrazone. — Lust. cryst. fr. alc. D.s. h. aq. or NaOH ; e.s. h. alc.
3331	148	Straw-yellow	α -Acetofuranesemicarbazone, $\text{C}_6\text{H}_5\text{O.C}(\text{N.NH.CO.NH}_2).\text{Me}$. — \textcircled{P} Semicarbazone. — Ndl. D.s. alc.
3332	148d.	Yellow	p-Nitrodiazoaminobenzene, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NH.N}_2.\text{Ph}$. — \textcircled{P} Boiled w. dil. H_2SO_4 evolves nitrogen & gives phenol (Vol. I), & p-nitraniline (No. 2.3319). — Silky ndl. fr. bz.
3333	146-8; 15	Yellow	3,3'-Diaminoazoxybenzene, $\text{NH}_2.\text{C}_6\text{H}_4.\text{N.O.C}_6\text{H}_4.\text{NH}_2$. — Ndl. fr. alc. Scales fr. toluene. — B_2HCl , cryst. powd., i. HCl. — Diacetyl deriv., ochre colored powd., d.s. gl. ac. ac., m.p. 254°.
3334	150	Dark red	N-Methylindole Picrate, $\text{C}_8\text{H}_7.\text{N.Pk}$. — \textcircled{P} \textcircled{O} Picrate* of No. 2.2823. — Pr. fr. eth. V.s. h. bz.; less s. eth.
3335	150d.	Brown-yellow	Tetranitro- $\beta\beta$ -binaphthyl, $(\text{NO}_2)_4.\text{C}_{20}\text{H}_{10}$. — Amorph. powd. V.d.s.

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 1 = T. 2.35. ** = T. 2.1. ¶ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3336	150(r.h.)	Yellow	α -3-Nitrocoumaric Ac., $\text{NO}_2\text{C}_6\text{H}_3(\text{OH})\text{CH}:\text{CH.CO}_2\text{H}$. — \textcircled{P} Nitro acid,† — Pr. fr. alc. Heated w. aq. or alc. gives anhydride. — $\text{BaA.3}\frac{1}{2}\text{H}_2\text{O}$, amorph. ppt. crystg. fr. h. aq. in fine red ndl.
3337	150-1	Yellow	p-Diketohexamethyleneosazone, $[\text{C}(\text{:N.NHPh}).(\text{CH}_2)_4\text{C}(\text{Ph}-\text{NH.N}):]$. — Pr. fr. alc.
3339	147-8; 152d.	Yellow or yellow-green	2-Nitrosophthalol(1), β -Naphthoquinoneoxime, $\text{NO.C}_{10}\text{H}_8\text{OH}$. — \textcircled{P} Sol. undecd. w. intense red color in conc. H_2SO_4 . — Ndl. fr. bz. Alm. i. c. aq.; e.s. alc., less s. bz., chlf., eth., lgr. — Boiled w. conc. HCl gives hydroxylamine (Cf. T. 2.17)! — E. oxid. to 2-nitronaphthalol(1) (No. 2.3213) by alk. ferricyanide (Ber., 25, 973).
3340	151	Red	Bz-3-Methylindole Picrate. — Picrate of No. 2.1589.
3341	151	Or.-red	4'-Hydroxy-4-methylazobenzene, $\text{HO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{Me}$. — \textcircled{P} \textcircled{D} Azo comp. — Monoclin. pr. w. blue reflections. D.s. h. aq.; v.s. alc., eth., bz., alk. — Ammon. sol. w. AgNO_3 gives ppt. of lust. or. ndl. — B.HCl , carmine-red powd., completely decd. by aq., m.p. 169°.
3342	151	Scarlet	Benzeneazo- α -phenylnaphthylamine, $\text{Ph.N.C}_{10}\text{H}_8\text{NHPh}$. — \textcircled{P} \textcircled{D} Azo comp. — Lit. fr. alc. E.s. bz.; d.s. lgr.
3343	151	Brown-yellow	p-Nitromethylaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NHMe}$. — Pr. w. violet reflections fr. alc. D.s. lgr. — HNO_2 gives nitroso deriv. (No. 2.3071).
3344	150-1; 156; 144	Yellow	3,3'-Diaminoazobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{NH}_2$. — \textcircled{P} \textcircled{D} Azo comp. & prim. amine, — Ndl. S. alc.; less s. eth., chlf., pet.-eth. — B_2HCl , golden lft. — Diacetyl deriv.,** or. ndl. fr. aniline, m.p. 247°.
3345	150-2	Green	1-Nitroso-2-naphthylamine, $\text{NO.C}_{10}\text{H}_8\text{NH}_2$. — Ndl. fr. dil. alc. D.s. h. aq.; e.s. alc., eth., dil. ac. — Heated w. aniline & gl. ac. ac. gives anilinonaphthoquinoneanilide, red ndl. fr. h. alc., m.p. 187° c.
3346	150-2	Green	Aminophenylindoline, Induline 3B, $\text{C}_{10}\text{H}_8\text{N}_2$. — Shimmering pr. fr. alc. — $\text{B.HCl.3}\frac{1}{2}\text{H}_2\text{O}$, alm. completely pptd. fr. sol. by HCl . I. c. alc.
3347	151	Moss-green	5-Nitroso-o-methyltoluidine, $(\text{NO})\text{Me.C}_6\text{H}_4\text{NHMe}$. — \textcircled{P} Boiled w. NaOH sol. (Cf. T. 2.26) gives methylamine (No. 2.1059) & nitroso-o-cresol. — Lust. lft. fr. bz.
3348	152w. efferv.	Dark red, alm. black	Glyoxalosotetrazone, $[\text{CH}:\text{N.NPh.NPh.N:N:CH}]^2$. — Lft. fr. alc.
3349	152	Orange	4-Hydroxyazobenzene, $\text{HO.C}_6\text{H}_4\text{N}_2\text{Ph}$. — \textcircled{P} Azo comp. — Pr. fr. alc. V.s. alc., eth.; v.d.s. h. aq.; s. ammon. — Acetyl deriv.** (fr. ac. anhydride), or. cryst. fr. eth.-alc., m.p. 84-5°.
3350	152; 148	Yellow	Phenylglyoxalosazone, $\text{Ph.C}(\text{:N.NHPh}).\text{CH}: \text{N.NH.Ph}$. — Lft. I. aq.; e.s. eth., bz., h. alc.
3351	152	Yellow	Xanthonephenylhydrazone, $\text{O}:(\text{C}_6\text{H}_5)_2:\text{C}:\text{N.NHPh}$. — Ndl. fr. alc. Sol. in conc. H_2SO_4 yel. w. green fluor.!
3352	153	Red	o-Nitrobenzaldehydephenylhydrazone, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}:\text{N.NHPh}$. — \textcircled{P} Hydrazone,†† — Ndl. Alm. i. aq., lgr.; d.s. alc., eth.
3353	153-4d.	Brown	Perezoneoxime, $\text{C}_{10}\text{H}_8\text{O}_2\text{N}$. — \textcircled{P} E.s. alk. w. blue color! — Flat ndl. fr. dil. alc. Sbl. I. aq.; e.s. alc., eth. Sol. in alc., chlf., or bz., are purple-red.
3354	153-4u.c.	Dark red; or.-yel.	Nitrosocotoxin, $\text{Ph.CO.C}_6\text{H}_3(\text{OH})_2(\text{OMe})(\text{NO})$. — Ndl. Dark-red & stable fr. conc. sol. in gl. ac. ac.; or.-yel., efflorescing w. loss of weight when crystd. fr. dil. sol. in same solvent. Alm. i. aq., eth. mod. s. h. alc.; e.s. bz.
3355	153; 141	Orange	o-Azophenoldimethylether, o-Azoanisole, $\text{MeO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OMe}$. — \textcircled{P} Azo comp. — Pr. fr. MeOH. Dist. D. vol. w. st. E.s. alc., eth. bz., chlf.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3356	154d.	Red	2,3-Diphenylindole Picrate. — Picrate of No. 2.2018. — Ndl. fr. bz. E.s. alc., bz.
3357	154	Lemon-yellow	3,4-Dinitroaniline, $(NO_2)_2.C_6H_4.NH_2$. — \textcircled{P} Nitro comp.† & prim. amine, ¶ evolving NH_3 w. boiling KOH sol. (T. 2.7). — Fine ndl. fr. aq.
3358	154	Yellow	Nitro-m-toluenediamine, $(NO_2)(NH_2)_2.C_6H_4.Me$. — Ndl. w. violet reflections fr. aq. — R.d.s. h. aq.; s. h. alc. Salts decd. by aq.
3359	154	Yellow	o-Nitrodiphenyldiacetylene, $NO_2.C_6H_4.C:C.C:CPH$. — \textcircled{P} Sol. in conc. H_2SO_4 , brown-red. — Lst. fr. alc. Sinters at 145°. S. alc., eth.
3360	154	Light yellow	m-Nitrotriphenylurea, $NO_2.C_6H_4.NH.CO.PH_3$. — \textcircled{P} Prob. dis. in warm conc. H_2SO_4 w. deep blue color. — Fine ndl. E.s. alc.; d.s. eth., lgr.; i. aq.
3361	154	Light yellow	(γ), or 1,4,5-Trinitronaphthalene, $(NO_2)_3C_{10}H_4$. — Lust. lft. S. at 18.5°: in 95 pt. bz.; 260 pt. eth.; or 894 pt. 90% alc.
3362	154-5	Sulphur-yellow	Benzophenone-p-nitrophenylhydrazone, $Ph_2C:N.NH.C_6H_4.NO_2$. — Ndl. I. h. aq.; s. acetone, bz., alc. S. alc. NaOH w. strawberry-red color.
3363	155	Dark garnet red	p-Nitrobenzaldehydophenylhydrazone, $NO_2.C_6H_4.CH:CH:N.NHPh$. — Hydrazone.‡ — Cryst. w. bluish reflections.
3364	155	Or.-red	4-Benzeneazo-3-methyl-1-phenylpyrazolone(5), $C_{10}H_{14}ON_4$. — \textcircled{P} Azo comp. — Lust. ndl. fr. alc. S. in 505 pt. alc. I. aq. dil. ac. Sol. in boiling dil. NaOH & repprd. by ac. — Reduced by Zn dust + gl. ac. ac. to aniline & No. 2.106.
3365	155-6	Golden-yellow	αβ-Dinaphthyl Picrate, $C_{20}H_{14}Pk$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I.
3366	152-5; 155; 158; 160	Yellow	1-Xylosephenylosazone, $CH_2(OH).(CH.OH)_2.C:(N.NHPh)-CH:NHPh$. — Silky ndl. D.s. aq.; s. eth., acetone. Sol. levorotatory.
3367	abt. 155	Light yellow	Formaldehyde-2,4-dinitrophenylhydrazone, $CH_2:CH:N.NH.C_6H_4-(NO_2)_2$. — Cryst. fr. alc. — V.d.s. eth.
3368	155-6	Yellowish	6-Nitrosothymol, Thymoquinoneoxime, $(NO)Me(C_6H_5)(HO)-C_6H_5$. — \textcircled{P} S. w. red-yel. color in NaOH sol. — Ndl. V.d.s. h. aq.; e.s. alc., chlf., eth. — Treatment w. conc. HNO_3 gives dinitrothymol (No. 2.2908).
3369	156	Red	β-Diazoaminonaphthalene, $C_{10}H_7.NH.N_2.C_{10}H_7$. — \textcircled{P} Dis. in conc. H_2SO_4 w. violet color.
3370	152-3; 159.5	Red	3-Nitro-1,2-dihydroxynaphthalene, $NO_2.C_10H_8(OH)_2$. — \textcircled{P} Nitro comp.† Fine ndl. fr. aq. Sbl. in long red ndl.! — Tbl. fr. alc. or bz. D.s. h. aq.; e.s. alc.
3371	156	Red	6-Nitro-o-tolidine, $(NO_2)Me^*(NH_2)^4.C_6H_2.C_6H_4.Me^*(NH_2)^4$. — Ndl. fr. alc. R.d.s. c. alc.; more s. bz.; d.s. eth.; i. lgr.
3372	156	Red	2-Diethylaminoanthraquinone, $NEt_2.C_10H_8O_2$. — Ndl. S. in abt. 200 pt. c. alc.
3373	156-7	Yel.-red	2,4-(α)-Dinitrodiphenylamine, $(NO_2)_2.C_6H_4.NHPh$. — Reduction by H_2S in alc. NH ₃ sol. gives aminonitrodiphenylamine, red cryst., m.p. 125° (Ber., 28, 2971).
3374	155.5-6.5u.c.	Yellow	Dimethylamine Picrate, $Me_2NH.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1061. — Lust. scales or ndl. S. in 56 pt. aq.
3375	156-7	Yellow	Benzalacetonephenylhydrazone, $Ph.CH:CH.C:(N.NHPh)Me$. — \textcircled{P} Hydrazone.‡ — Flat ndl. I. aq., alk.; d.s. c., e.s. h. alc.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34. ¶ = T. 2.35. ** = T. 2.1. ‡ = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3376	156.5	Yellow	p-Hydroxyazoxybenzene, $\text{HO.C}_6\text{H}_4.\text{N}_2\text{O.Ph}$. — \textcircled{P} Boiled w. NH_4Cl sol. & Zn dust gives aniline (No. 2.1235), & p-aminophenol (No. 2.963). — Ndl. fr. bz. E.s. alc., eth., chlf.; d.s. c. bz.
3377	156	Yellow	l-Xylose-p-nitrophenylhydrazone, $\text{CH}_2(\text{OH}).(\text{CH.OH})_2.\text{CH}:\text{N}-\text{NH.C}_6\text{H}_4.\text{NO}_2$. — Cryst. S. alc.
3378	156	Yellow	l-Gulosephenylosazone, $\text{CH}_2(\text{OH}).(\text{CH.OH})_2.\text{C}(\text{:N.NHPh}).\text{CH}:\text{N.NHPh}$. — Cryst. flock fr. dil. alc. Mod. s. h. aq. $[\alpha]_D = +46^\circ$ ($C = 0.4$ in MeOH). — The corresponding d,l-deriv. is less s. in aq., w. m.p. 157–9°. — [Identical w. the osazones of l-gulose & l-sorbinose.]
3379	156	Yellow	Apiosephenylosazone, $\text{CH}_2(\text{OH})_2.\text{C}(\text{OH}).\text{C}(\text{:N.NHPh}).\text{CH}:\text{N.NHPh}$. — S. h. aq., eth.; e.s. alc., acetone.
3379-I	156–7	Yellow	2-Nitro-3-aminobenzoic Ac., $(\text{NO}_2)(\text{NH}_2).\text{C}_6\text{H}_3.\text{CO}_2\text{H}$. — Ndl. fr. aq. V.s. h. aq., c. alc., eth.; r.d.s. c. aq. Boiled w. KOH sol. gives NH_3 & 2-nitro-3-hydroxybenzoic ac. — $\text{KA}_2\text{H}_2\text{O}$, deep red tbl. — $\text{Ba.A}_2\text{H}_2\text{O}$, ndl., v.s. c. aq.
3380	156	Light greenish yellow	o-Diphenyleneazone, Phenazone, $[\text{C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_4]$. — Fine ndl. fr. alc. B.p. a. 360° alm. undecd. E.s. alc., bz.; v.s. chlf.; d.s. lgr.; more s. eth. — \textcircled{P} Picrate (T. 2.23), m.p. 194° .
3381	157d.	Or.-red	Diphenylcarbazone, Ph.N:N.CO.NH.NHPh . — Ndl. fr. bz. E.s. alc., chlf., bz.; s. unchanged in conc. H_2SO_4 . — Reduced by Zn dust + NaOH to diphenylcarbazide (No. 2.248).
3382	157.5	Bordeaux-red	o-Nitrocinnamicaldehydephenylhydrazone, $\text{NO}_2.\text{C}_6\text{H}_4.\text{CH}:\text{CH}:\text{N.NHPh}$. — \textcircled{P} Hydrazone. — Ndl. fr. alc. I. aq.
3383	157d.	YO, or O	p-Nitrophenylhydrazine, $\text{NO}_2.\text{C}_6\text{H}_4.\text{NH.NH}_3$. — \textcircled{P} Reduces Tollen's reagt. instantly in T. 2.30. — In T. 2.16: (a) dyes wool YOSI; (b) Rongolite discharge, YT2; (c) H_2SO_4 , unchanged; (d) NaOH , dark red-brown. — Lust. ndl. fr. h. alc. Color fr. 30% alc., O; fr. 95% alc., YO. — D.s. aq.; s. alc., eth., chlf.; i. bz. Odor aromatic. — \textcircled{P} Add 1 drop benzaldehyde to sol. of 0.03 g. in 4 cc. h. dil. alc. (2:1). Dis. ppt. by heating. Allow cryst. to separate. Wash w. 5 cc. dil. alc. (2:1). Recryst. fr. 9 cc. boiling dil. alc. Wash w. 3 cc. dil. alc. Dry at 100° . The resulting phenylhydrazone forms fine YO ndl., m.p. 190° u.c.
3384	157	Brown	Pr-2,3-Dimethylindole Picrate, $\text{C}_{10}\text{H}_{11}\text{N.Pk}$. — \textcircled{P} \textcircled{P} Picrate* of No. 2.1911. — Lust. ndl. fr. alc. Alm. i. aq.
3385	157	Brown-yellow	Dinitro-o-acetanisidine, $(\text{NO}_2)_2(\text{C}_6\text{H}_3\text{O.NH}).\text{C}_6\text{H}_3.\text{OMe}$. — Pr. fr. alc. D.s. c. alc.
3386	157	Or.-yel.	3-Methyl-1-phenyl-4-isonitrosopyrazolone(5), $[\text{NPh.N:CMe.C}(\text{:NOH}).\text{CO}]$. — \textcircled{P} "Strong ac." w. yel.-red salts. — Ndl. fr. ac. ac. V.d.s. aq.; e.s. h. gl. ac. alc.; s. eth. Sbl. abt. 100° .
3387	157d.	Golden	4-Nitroso-1-methylnaphthylamine, $\text{NO.C}_{10}\text{H}_8.\text{NHMe}$. — \textcircled{P} Boiling w. NaOH sol. gives methylamine (No. 2.1059), & 4-nitrosonaphthol(1), (No. 2.3602). — Cryst. fr. bz. I. aq. — B.HCl , fine yel.-green ndl.
3388	157	Yellow	Methyl 2,4,6-trinitrobenzoate, $(\text{NO}_2)_3.\text{C}_6\text{H}_3.\text{CO}_2\text{Me}$. — D.s. pr. plates.
3389	157d.	Light yellow	3-Nitro-2-hydroxynaphthoquinone(1,4), $(\text{NO}_2)(\text{HO}).\text{C}_{10}\text{H}_4\text{O}_2$. — Lft. E.s. alc., eth., h. aq.; d.s. chlf., bz. Long boiling w. aq. dec. to HCN & phthalic ac. — AgA , dark yel. cryst. e.s. h. aq. — \textcircled{P} V.e. oxid. by boiling dil. HNO_3 to phthalic ac. (T. 1.318–1).
3389-I	157–8	Light yellow	2,6-Dinitro-p-toluic Ac., $(\text{NO}_2)_2.\text{C}_6\text{H}_3\text{Me}.\text{(CO}_2\text{H})$. — \textcircled{P} Acid & nitro comp. — Lft. fr. h. aq. D.s. c. aq.; e.s. alc., eth. Sbl. — $\text{CaA}_2\text{H}_2\text{O}$, thick red pr.
3390	157	Greenish dark yel.	Diethylanilinoxide Picrate, $\text{C}_{10}\text{H}_{16}\text{ON.Pk}$. — Pr. w. violet reflections fr. alc.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3391	158	Red	3-Nitronaphthoquinone(1,2), $\text{NO}_2\text{C}_{10}\text{H}_8\text{O}_2$. — Cryst. D.s. aq., eth.; s. h. alc., bz.; v.s. h. gl. ac. ac. — (D) Short boiling in alc. sol. w. 2-3 mol. aniline gives anilide, lust. red ndl. fr. xylene, m.p. abt. 250°.
3392	157-9	Or.-yel.	p-Dihydroxyazobenzenediethylether, p-Azophenetole, $\text{EtO-C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OEt}$. — (D) Azo comp. — Lft. D.s. c. alc.; e.s. chlf., eth. Dist. undecd.
3393	158	Gold-yellow	Phenanthrenequinoneoxime, $\text{HON:C}_{10}\text{H}_8\text{O}$. — (D) S. conc. H_2SO_4 w. blood-red color; s. h. NaOH sol. w. green color. — Ndl. fr. alc. E.s. h. alc. — Gives T 2.17-b.
3394	158	Bronze	Formazyl Cyanide, $\text{CN.C}_{10}\text{H}_8\text{N}_4$. — (D) Sol. in conc. H_2SO_4 , dark blue; in other solvents, deep red! — Lft. fr. alc. D.s. c. alc., eth., lgr.
3395	158-9	Yellow	6-Aminoquinoxaline, $\text{NH}_2\text{C}_6\text{H}_4$. — (D) Eth. sol. fluor. yellow-green! Sol. in conc. mineral ac., intense violet, becoming brown-red on diln. w. aq. — Ndl. fr. eth. Sbl. alm. undecd. E.s. aq., alc., chlf.; less s. eth., bz. — B.HCl , brown-red lft. w. green reflections, d. abt. 215°.
3396	158-9w. efferv.	Yellow	o-Aminocinnamic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{CH:CHCO}_2\text{H}$. — (D) Solutions fluor. intense blue-green! — Ndl. D.s. c. aq.; s. h. aq., alc., eth. — BaA_2 , r.d.s. h. aq.
3397	158-9	Yellow	6-Nitro-3-aminodurene, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{Me}_3$. — (D) Prim. amine. — Ndl. fr. dil. alc.
3398	158d.	Yellowish(?)	Choline Picrolonate, $\text{C}_6\text{H}_5\text{ON.Pk}$. — Picrate* of deliq. syrupy base. — Loses aq. at 130°. Dec. at 241-5°.
3399	159	Red	5,5'-Diamino-2,2'-azotoluene, $(\text{NH}_2)_2\text{C}_6\text{H}_3\text{N}_2\text{C}_6\text{H}_3(\text{NH}_2)_2$. — (D) Azo comp. — Cryst. fr. alc. — [Said to also cryst. in modifications w. m.p. 132-3° & 142-5°.] — S. h. aq.; e.s. h. alc. — B.2HCl , brown lft. fr. h. aq. 100 pt. boiling aq. dis. 0.168 pt.
3400	159	Orange	(γ)-2,3-Dinitromethyl-p-toluidine, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{Me}(\text{NHMe})$. — Lft. w. bluish reflections fr. chlf. D.s. c. alc.; lgr.; e.s. h. bz., chlf.; s. eth. — Nitrosamine, yel. ndl. fr. alc., m.p. 128° (Ber., 30, 840).
3401	159-60	Ocher-yellow	p-Aminobenzeneazo-α-naphthylamine, $\text{NH}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{NH}_2$. — (D) Azo comp. & prim. amine. — Ndl. E.s. alc., chlf., bz. — Salts s. aq. w. red color. Boiled w. FeCl_3 sol. gives pungent odor of benzoquinone & α-naphthoquinone (Vol. I). — [An intermediate in dyestuff industry.]
3402	159	Yellow	3,5-Dinitroaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{NH}_2$. — (D) Nitro comp.† & prim. amine. — Ndl. fr. h. aq.; e.s. alc., eth.; less s. bz.
3403	159-60	Yellow	Pyrazole Picrate, $\text{C}_6\text{H}_4\text{N.Pk}$. — (D) Picrate* of No. 2.1650-1. — Ndl.
3404	abt. 160u.c.	Deep viol.-red	2-Aminofluorenone, $[\text{C}_9\text{H}_7(\text{NH}_2)\text{CO.C}_6\text{H}_4]$. — (D) Sol. in HCl colors pine splinter brilliant red! — Melts to deep red liq. Pr. fr. alc. Alm. i. aq.; v.d.s. pet.-eth.; mod. s. h. alc., eth. — B.HCl , yel. pr. decd. by aq.
3405	158.5- 161.5d.	Garnet-red & gold-yel.	p-Nitrobenzenediazoamino-p-toluene, $\text{NO}_2\text{C}_6\text{H}_4\text{N}_2\text{H.C}_6\text{H}_4\text{Me}$. — Dimorphous. Thick lust. garnet-like cryst. fr. eth. or h. abs. alc. The yel. form is produced in fine ndl. by adding aq. to alc. sol. It is unstable, changing in contact w. solvents to red form. — E.s. eth., chlf.; d.s. c. bz.
3406	abt. 160 (r.h.)	Or.-yel.	1-Arabinosephenylosazone, $\text{CH}_2(\text{OH}).(\text{CH}_2\text{OH})_2\text{C}(\text{N.NHPh})-\text{CH: N.NHPh}$. — Cryst. fr. acetone or h. aq. I. c. aq., eth., bz., lgr.; s. alc., acetone, pyridine, h. aq.; i. alk. — $[\alpha]_D = +1^\circ, 10'$ (fr. 0.2 g. in 4 cc. pyridine + 6 cc. abs. alc.

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(ORDER II.)

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
3407	160-1	Or.-yel.	Chrysoquinonemonoxime, $\text{HON:C}_{18}\text{H}_{16}\text{O}$. — \textcircled{P} S. in conc. H_2SO_4 w. intense violet-red, or in dil. NaOH w. yel.-brown color! — Cryst. fr. gl. ac. ac. Mod. s. alc., bz., gl. ac. ac.
3408	d. abt. 160w. eferv.	Dark yellow	Nitranilicacidquinone, $\text{C}_{12}\text{H}_8\text{O}_{10}\text{N}_2$. — [Fr. nitrous ac. & benzoquinone.] — \textcircled{P} FeCl_3 , colors alc. sol. intense blood-red. — Oblique pr. fr. aq. V.s. alc.; v.d.s. chlf., bz.
3408-1	d. 160	Yellow	Triethylamine Picrolonate, $\text{Et}_3\text{N:C}_{10}\text{H}_8\text{O}_4\text{N}_4$. — \textcircled{P} \textcircled{D} Salt of No. 2.1095 w. No. 2.3184. — Cryst. S. in 63 pt. boiling, or 536 pt. c. aq.
3409	160-1	Green	6-Nitroso-3-methylamino-1,2-xylene, $(\text{NO})(\text{NHMe})\text{C}_6\text{H}_3\text{Me}_2$. — Ndl. w. metallic luster fr. alc. E.s. alc., bz.
3410	161d. (r.h.)	Red	Nitroformazan, $\text{Ph.NH.N:C}(\text{NO}_2)\text{N:N.Ph}$. — Ndl. fr. alc.
3411	161	Yel.-red	4-Nitro-m-phenylenediamine, $\text{NO}_2\text{C}_6\text{H}_4\text{(NH}_2)_2$. — \textcircled{P} Protracted boiling w. NaOH sol. gives NH_3 (T. 2.7) & nitro-o-aminophenol. — Somewhat s. aq.; more s. alc., eth.
3412	161-2u.c.	O-YO	Acenaphthene Picrate, $\text{C}_{12}\text{H}_{10}\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Ndl. fr. h. alc. D.s. c. alc.
3413	161-5	Pale yel.	1,6-Dinitronaphthalene, $(\text{NO}_2)_2\text{C}_{10}\text{H}_6$. — \textcircled{P} Nitro comp.‡ — Ndl. E.s. alc.
3414	162	Golden	trans-oo'-Diaminostilbene $\text{NH}_2\text{C}_6\text{H}_4\text{CH:CH.C}_6\text{H}_4\text{NH}_2$. — \textcircled{P} Alc. sol. fluor. strongly blue-violet! — Lust. pr. fr. alc. — Distn. of B w. B.HCl yields indole (No. 2.1546) & aniline (No. 2.1235).
3415	162u.c.	Y-*	p-Nitrodimethylaniline, $\text{NO}_2\text{C}_6\text{H}_4\text{NMe}_2$. — \textcircled{P} Nitro comp.‡ giving Ag mirror & brownish ppt. in T. 2.21. Lust. ndl. fr. h. alc. S. alc.; e.s. conc. HCl ; s. h. ac. ac.; i. aq. Odorless. Tasteless. — \textcircled{D} Treat 0.2 g. disd. in 5 cc. conc. HCl w. 1 g. gran. Sn. After Sn is disd., ppt. boiling hot sol. w. H_2S . Filter hot. Evap. to dryness on aq.-bath. The residue, consisting of dimethyl-p-phenylenediamine hydrochloride, gives the striking color react. described as prelim. test for No. 2.560.
3415-1	161-3	Yellow	Hyoscynamine Picrate. — Picrate of No. 2.777.
3416	163-4	Red	Diethylpicramide, 2,4,6-Trinitrodiethylaniline, $(\text{NO}_2)_3\text{C}_6\text{H}_3\text{-NET}_2$. — \textcircled{P} \textcircled{D} Boiling w. KOH sol. (T. 2.26) gives diethylamine (No. 2.1068-1) & picric ac. (No. 2.3168). — Cryst. V.s. bz.
3417	163d. (r.h.)	Cherry-red	Formazylformic Ac., $(\text{Ph.N:N})(\text{Ph.NH.N}): \text{C.CO}_2\text{H}$. — \textcircled{P} Sol. in KOH is blood-red. — Ndl. w. bluish reflections fr. alc. D.s. alc., eth.; v.s. chlf., bz. Dec. to CO_2 & No. 2.3159 in melting. — AgA, dark green-brown ppt. w. green reflections.
3418	163.5	Yellow	Naphthalene Styphnate, $[\text{C}_{10}\text{H}_8].[(\text{NO}_2)_2\text{C}_6\text{H}_4\text{(OH)}_2]$. — \textcircled{P} \textcircled{D} Styphnate* of h.c. of Vol. I. — Ndl. fr. acetone.
3419	162; 163.5	Yellow	Trinitroocinol, $(\text{NO}_2)_3\text{C}_6\text{Me(OH)}_2$. — \textcircled{P} Explodes feebly just above m.p. Sol. colored brown by FeCl_3 . — Long ndl. D.s. c. aq.; e.s. h. aq. (pptd. fr. sol. by min. ac.); e.s. h. bz.; less s. eth. — PbA, cryst. yel. ppt. — Naphthalene deriv., $\text{C}_{10}\text{H}_8 + \text{C}_7\text{H}_6\text{O}_3\text{N}_2$, yel. ndl., m.p. 120°.
3421	163	Pale yellow	Nitropapaverine, $\text{NO}_2\text{C}_{10}\text{H}_{20}\text{O}_4\text{N.H}_2\text{O}$. — Delicate pr. fr. dil. alc. In light soon becomes yel. I. aq., alc. Mod. s. h. alc.; e.s. chlf.; s. in 3100 pt. eth.
3422	163.4	Yellowish	Picryl Benzoate, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{CO}_2\text{Ph}$. — Clear glassy pr. fr. bz.
3423	163	Green, or blue	p-Nitrosophenyl-p-toluidine, $\text{NO.C}_6\text{H}_4\text{NH.C}_6\text{H}_4\text{Me}$. — Cryst. fr. bz. in green lf. Also observed as blue pr. S. alc., eth., chlf.; less s. bz.; v.d.s. lgr. — Boiled w. NaOH sol. gives p-toluidine (No. 2.566), & nitrosophenol.
3424	164	Gold-yellow	4-Nitronaphthol(1), $\text{NO}_2\text{C}_{10}\text{H}_8\text{OH}$. — \textcircled{P} Nitro comp.‡ — Ndl. fr. h. aq. E.s. alc. Not vol. w. st. Salts red.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3425	164.5	Deep yellow	2,4-Dinitrocinol, $(NO_2)_2.C_6H.Me(OH)_2$. — \textcircled{P} Nitro comp. exploding when rapidly heated. — Lft. alm. i. c. aq.; s. h. aq.; e.s. eth., h. bz.; s. 18 pt. alc. at 15° ; alm. i. lgr.
3426	164	Yellow	3,5-Dinitropyrocatechol, $(NO_2)_2.C_6H_4(OH)_2$. — \textcircled{P} Nitro comp. — Ndl. fr. alc. — W. cooled nitrosulphuric ac. gives nitranilic ac. (No. 2.3469).
3428	164	Yellow	d-Gulosephenylosazone, d-Idosephenylosazone, d-Sorbinosephenylosazone, $CH_2(OH).(CH.OH)_2.C(:N.NHPh).CH:N-NHPh$. — Ndl. D.s. aq.; s. warm alc., acetone; i. bz., chlf., eth.
3430	165	Dark red	p-Picrylaminophenolmethyleneether, $(NO_2)_2.C_6H_3.NH.C_6H_4.OMe$. — Silky ndl. fr. alc. + ac. ac. D.s. eth.
3431	165	Red	Phenanthrenequinonephenyldihydrazone, $C_{14}H_8O.N.H.Ph$. — \textcircled{P} S. in conc. H_2SO_4 w. violet color. — Lust. cryst. fr. alc. Mod. s. h. alc.
3432	165d.	Deep garnet-red, or golden	Benzeneazopyrocatechol, $Ph.N_2.C_6H_4(OH)_2$. — Cryst. w. blue reflections fr. alc. Separates in golden lft. fr. quickly cooled alc. sol., gradually changing to red ndl. — E.s. alc.; s. NaOH or Na_2CO_3 sol.!
3433	165-6d.	Ruby-red, or pale yel.	o-Nitrobenzoylformic acid diphenylhydrazone, $NO_2.C_6H_4.C(:N-NHPh).CO_2H$. — \textcircled{P} Acid. — Mic. red ndl. fr. alc.; or pale yel. tbd. fr. ac. ac. — I. aq., bz.; e.s. alc., eth. 100 pt. sat. gl. ac. ac. sol. at 20° contains 1.24 pt. E.s. KOH!
3434	165	Gold-red	Nitroso- α -diethylnaphthylamine, $NO.C_{10}H_8.NEt_2$. — \textcircled{P} Is colored dark blue by conc. H_2SO_4 . — Scales fr. alc.
3435	165	Brown-red	s-Tetramethylanthracenehydride Picrate, $C_{14}H_{10}.Pk$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Lust. ndl.
3436	d. 165	Yellow	Aniline Picrate, $Ph.NH_2.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1235. — Cryst. fr. alc. S. in 222 pt. aq. at 17.5° , or in 11.9 pt. alc. at 15° .
3436-I	165	Yellow	Ethylamine Picrate, $Et.NH_2.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1062. — Cryst. fr. alc. S. in 66.7 pt. aq. or 30.7 pt. alc. at 16° .
3437	165	Yellow	5-Nitroso-1,4-xylenol(2), p-Xyloquinoneoxime, $NO.C_6H_4.Me.(OH)$. — Ndl. Alm. i. aq.; s. alc., eth., bz.
3438	166	Ponceau-red	Formazylglyoxylic Ac., $Ph.NH_2.N:C(N:NPh).CO.CO_2H$. — \textcircled{P} Acid, disg. in alk. w. or.-red, or in conc. mineral ac. w. deep violet color. — Adamantine ndl. fr. alc. E.s. w. red color in alc., eth., chlf. bz.; d.s. aq., lgr. — Boiled w. HCl gives aniline & phenazine.
3439	166d.	Yellow	Nitrohemipinic Ac., $(NO_2)(MeO)_2.C_6H_4.(CO_2H)_2$. — \textcircled{P} Acid & nitro comp. — Glassy pr. w. $1H_2O$ of crystn. (lost at 120°). Heated at m.p. for 1-2 hr. gives anhydride, yel. pr. fr. bz., m.p. 145° . — $Ag\ddot{A}$, yel. ppt. — $Ba\ddot{A}.2H_2O$, yel. ndl. d.s. aq.
3440	167-8	Dark red	Pr-2-Methyl- α -naphthindole Picrate, $C_{13}H_{11}N.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.2096. — Ndl. fr. bz.
3441	167	Brown or yellow	2,4-Dinitromethylidiphenylamine, $(NO_2)_2.C_6H_3.NMePh$.
3441-I	166-7	Yellow	Acetylpropionylphenylosazone, $Me.C(:N.NHPh).C(:N.NHPh)-Et$. — Ndl. fr. bz.
3442	164d.; 166-8	Gold-yellow	d-Erythrosephenylosazone, $CH_2(OH).CH(OH).C(:N.NHPh)-CH:N.NHPh$. — V.d.s. h. aq.; more s. eth., h. bz.; v.s. acetone, alc. Reduces h. Fehling's sol. — [The l- & d-l- osazones are also described as having the m.p. here given.]
3443	166-8d.	Yellow	d,l-Arabinosephenylosazone, $CH_2(OH).(CH.OH)_2.C(:N.NHPh).CH:N.NHPh$. — Ndl. fr. aq.

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(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3444	abt. 166 u.c., d.	Yellow	α -Naphthylamine Picrate, $C_{10}H_8N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.589. — Ndl. fr. dil. alc. Darkens abt. 150°, & is black at m.p.
3445	165; 169-71	Yellow	α -Picoline Picrate, $C_8H_7N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1153. — Ndl. Mod. s. aq.
3445	167	Yellow	2,4,6-Trinitrophloroglucinol, $(NO_2)_3C_6(OH)_3$. — Hexag. cryst. w. $1H_2O$ (lost at 100°) fr. aq. Exploded by heat! E.s. h. aq., alc., eth. Dyes deep yel. — $K_3\bar{A}$, deep yel. cryst.
3447	166; 168	Yellow	3,5-Dinitro-p-toluidine, $(NO_2)_2C_6H_4Me(NH_2)$. — \textcircled{P} Nitro comp.† & prim. amine. — Ndl. fr. CS. D.s. h. alc.; s. in 300 pt. CS; e.s. bz. — \textcircled{D} Oxidn. by CrO_3 mixt. (Ber., 11, 1976) gives chrysanthemic acid (No. 2.3870).
3445	168-9	Red	Picramic Ac., 4,6-Dinitro-2-aminophenol, $(NO_2)_2(NH_2)C_6H_4OH$. — Monoclin. pr. fr. chlf. Mod. s. alc.; s. eth., chlf.; e.s. bz., gl. ac. ac., conc. HCl. 100 pt. aq. at 22° dis. 0.14 pt.
3445	168-9	Red	p-Benzenedisazobenzene, $Ph.N_2C_6H_4N_2Ph$. — \textcircled{P} Azo comp. — Lft. fr. gl. ac. ac. V.d.s. alc., eth., bz. — \textcircled{D} Boiled w. Fe powd. & dil. ac. ac. gives p-phenylenediamine (No. 2.877).
3450	168	Yellow-red	2,2'-Diaminoazotoluene, $(NH_2)MeC_6H_4N_2C_6H_4(NH_2)Me$ [$N_2 = 4,4'$]. — \textcircled{P} \textcircled{D} Azo comp. — Silky ndl. fr. dil. alc. D.s. aq.; e.s. alc., eth. M.p. of diacetyl deriv., 290°.
3451	168	Bronzy	2',4'-Dinitro-4-dimethylaminodiphenylamine, $(NO_2)_2C_6H_4NH-C_6H_4NMe_2$. — Scales fr. alc. — B.HCl, yel. cryst.
3452	168; 170	Yellow	Dinitro-4-aminobenzaldehyde, $(NO_2)_2(NH_2)C_6H_4CHO$. — [D.R.P.] — Cryst. fr. alc.
3453	168; 171	Yellow	2,6-Dinitro-p-toluidine, $(NO_2)_2C_6H_4Me(NH_2)$. — \textcircled{P} Nitro comp.† & prim. amine. — Hair-like ndl. fr. 50% ac. ac.; v.s. alc.; v.d.s. h. aq., CS.
3454	168	Yellowish	4-Nitropyrocatechol, $NO_2C_6H_4(OH)_2$. — \textcircled{P} Sol. in KOH sol. w. purple color. ("Delicate")! — Ndl. E.s. aq., alc., eth.; d.s. bz. — $Ba\bar{A}.3H_2O$, dark red lft., v.d.s. h. aq.
3455	168	Pale yellow	Cinnamicaldehydenaphthaldehyde, $Ph.CH:CH.CH:N.NHPH$. — \textcircled{P} Hydrazone. — Fine plates or ndl. I. aq.
3456	169	Dark red	α -Nitrosodi- α -naphthylamine, $NO.C_{10}H_8.NH.C_{10}H_7$. — Lust. ndl. fr. dil. alc. E.s. alc., bz. Boiled w. dil. H_2SO_4 gives α -naphthylamine (No. 2.589) & 4-nitrosonaphthol(1).
3457	168-70u.c.	OS2	Skatole Picrate, $C_8H_7N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1830. — Lust. ndl. fr. h. aq. Sbl. fr. 140°.
3458	169	Or.-yel.	β -Lapachonoxime, $HON:C_{10}H_8O_2$. — \textcircled{P} Oxime (T. 2.17). — Silky pr. fr. alc. I. NaOH sol. — Benzoyl deriv., golden ndl. fr. alc., m.p. 180-1°.
3458	169	Honey-yel.	6-Nitro-3-hydroxybenzoic Ac., $NO_2C_6H_3(OH)(CO_2H)$. — \textcircled{P} Nitro comp.† & acid. — Ndl. fr. aq. E.s. aq., alc., eth. Ad. sol. colored faintly red-brown by $FeCl_3$. — $Ba\bar{A}.6H_2O$, yel.-red pr., e.s. aq.
3460	168-70	reddish-yellow	5-Aminocoumarin, $[O.CO.CH:CH.C_6H_3(NH_2)]^2$. — \textcircled{P} Prim. amine. — Ndl. V.d.s. c. aq.; e.s. h. aq. alc. — Benzoyl deriv., m.p. 173°.
3461	169	Yellow	p-Aminobenzophenonephenylhydrazone, $NH_2C_6H_4C(=O)N.NH-Ph$. — \textcircled{P} Prim. amine. — Ndl. S. h. alc., chlf.
3462	169.5	Yellow	Dipicrylhydroxylamine, $[(NO_2)_2C_6H_4]_2NOH$. — [Fr. hydroxylamine & picryl chloride.] — Cryst. Sbl.
3463	170-1	Red	Dimethylaminomethylphenazine, $C_8H_7N_2$. — \textcircled{P} S. dil. HCl w. violet color. E.s. in alc. to red-brown sol. w. yel.-red fluor., or in chlf. w. green fluor. — Pr. fr. abs. alc. Garnet-red by transmitted light w. bronzy green reflections.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3464	167; 172-3	Dark brown	Dianilinotoluquinone(2,5)-anilide, $(\text{Ph.NH})_2\text{Me.C}_6\text{HO.(NPh)}$. — Lft. w. bluish reflections. Mod. s. h. alc. or h. ac. ac.
3465	169-71	Coppery lustre	Phenylisorosinduline, $\text{C}_{18}\text{H}_{14}\text{N}_2$. — \textcircled{P} E.s. alc. or bz. w. blue color! — Warts fr. bz. + lgr. D.s. lgr.
3466	170	Or.-yel.	Hexamethylbenzene Picrate, $\text{Me}_6\text{C}_6\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Loses its h.c. at 100-10°. Decd. by alc.
3467	170-1	Pale or.-yel.	γ -Nitrophenanthrene, $\text{NO}_2\text{C}_1\text{H}_8$. — Lft. fr. ac. ac. D.s. alc., eth. — Oxidn. by CrO_3 in ac. ac. gives or.-yel. ndl. m.p. 263° d.
3468	170	Yellow	2,6-Dinitro-4-aminophenol, Isopicramic Ac., $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_3\text{-OH}$. — Ndl. fr. aq. 100 pt. aq. at 22° dis. 0.082 pt., or 0.812 pt. at 100°; v.s. alc. — KA blue-black ndl. fr. alc.; v.s. aq.
3469	deflagrates w.m. at 170	Yellow	3,6-Dinitro-2,5-dihydroxyquinone, Nitranilic Ac., $(\text{NO}_2)_2\text{C}_6\text{O}_2\text{-}(\text{OH})_2$. — Cryst. (in vacuo) in tbl. When containing cryst.-aq. melts a little above 100°. V.s. aq., alc.; i. eth. — Aq. sol. on standing yields HCN & oxalic ac.
3470	170-1	Light yellow	Phenazine, Azophenylene, $[\text{C}_6\text{H}_4\text{N}^+][\text{N.C}_6\text{H}_4^-]$. — \textcircled{P} S. conc. H_2SO_4 w. blood-red color. Colored green by SnCl_4 . — Ndl. V.d.s. aq.; s. in 50 pt. c. alc.; less s. eth., bz. B.p. a. 360°. Sbl. undecd. — B.Pk, yel. ndl. fr. bz.; d.s. c. bz.
3471	168; 172	Light yellow	2-Nitrobenzoyl-p-toluide, $(\text{NO})\text{Me.C}_6\text{H}_3\text{NH(CO.Ph)}$. — Pr. E.s. h. alc.
3472	170-1; 172	Yellowish	Phenylauramine, $\text{PhN:C}(\text{C}_6\text{H}_4\text{NMe}_2)_2$. — Ndl. or cryst. powd. I. aq.; eth.; r.d.s. alc. — B.HCl, red, mod. s. sq., e. decd. by dil. HCl to aniline (No. 2.1235) & tetramethyldiaminobenzophenone (No. 2.2317).
3473	171	Golden	o-Azophenol, 2,2'-Dihydroxyazobenzene, $\text{HO.C}_6\text{H}_4\text{N.C}_6\text{H}_4\text{OH}$. — \textcircled{P} \textcircled{D} Azo comp. Sol. in KOH sol. w. red-yel. color & pptd. by acid. — Lft. fr. eth. I. aq.; s. in 300 pt. c. alc.; e.s. eth. Sbl. — Heated w. conc. HNO_3 gives picric ac. (No. 2.3168).
3474	171-2d.	Yellow	N-Nitrosoindole, $\text{C}_8\text{H}_7\text{N.NO}$. — \textcircled{P} Gives Liebermann's react. (T. 2.18). — Lust. cryst. Alm. i. aq., eth., lgr., bz.; e.s. warm acetone. — Reduction gives indole.
3475	170; 172	Yellow	1,8-(β)-Dinitronaphthalene, $(\text{NO}_2)_2\text{C}_1\text{H}_8$. — [An intermediate in dyestuff manufacture.] — \textcircled{P} Nitro comp.† — Rhomb. striated pr. fr. chlf. At 19° 100 pt. of each of following solvents dis.: 88% alc., 0.1886 pt.; chlf., 1.096 pt.; bz., 0.72 pt. — \textcircled{P} Convert into 1,3,8-trinitronaphthalene (No. 2.2527), by boiling 5 min. w. 5 pt. fuming HNO_3 + 5 pt. conc. H_2SO_4 . Cool. Pour on crushed ice. Wash ppt. w. eth. Cryst. fr. h. HNO_3 or gl. ac. ac. (Ann., 169, 96; Ber., 5, 905).
3476	171d.	Yellow	Methylpicrazide, $\text{Me.N}_2\text{H}_2\text{C}_6\text{H}_4(\text{NO}_2)_2$. — Lust. lft. fr. chlf. Mod. s. alc., eth.; less s. chlf.
3477	172	Yellow	Nitropiperonylic Ac., $\text{CH}_2\text{:O}_2\text{:C}_6\text{H}_4\text{(NO}_2\text{)(CO}_2\text{H)}$. — \textcircled{P} Nitro comp.† & acid. — Ndl. fr. aq. Deflagrates strongly when ignited. D.s. h. aq.; e.s. alc., eth. — $\text{PbA}_2\text{H}_2\text{O}$, lust. yel. ndl., alm. i. aq. — AgA, yel. ndl. or lft.
3478	172	Yellow	Benzooquinonesemicarbazone, $\text{NH}_2\text{CO.NH.N:C}_6\text{H}_4\text{:O}$. — Ndl. E.s. aq., alc., alk. — Boiled w. alk. gives phenol.
3479	172; 166-7	Pale yellow	2,7-Dimethyl-9-phenylacridine, $(\text{Me}_2\text{Ph.C}_1\text{H}_8)$. — \textcircled{P} Sol. in alc. shows bluish, & in acids, strong green fluor.! Addition of ammon. to sol. of B.HCl in dil. alc. changes green to violet fluor.! — Ndl. E.s. bz. w. pale yel. color.
3480	172	Light yellow	Tetranitrodiphenylmethane, $(\text{NO}_2)_4\text{C}_1\text{H}_8$. — Pr. fr. gl. ac. ac. I. alc., eth.

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 ¶ = T. 2.35. ** = T. 2.1. ¶¶ = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3461	173-4	Red	Tetramethyldiaminobenzoquinone, $(Me_2N)_2C_6H_2O_2$. — Tbl. fr. alc.
3452	abt. 173.5	Red	Formazylbenzene, $(PhNH:N)C_6H_4NHPH$. — Lft. w. green metallic reflections fr. alc. D.s. alc.; e.s. eth., chlf.
3483	173-4; 171-2	Brick-red	2,4,5,2',4',5'-Hexamethylazobenzene, Azopseudocumene, $Me_2C_6H_4N_2C_6H_4Me_2$. — \textcircled{P} \textcircled{O} Azo comp. S. undecd. w. blood-red color in conc. H_2SO_4 . — Cryst. fr. gl. ac. ac. 100 pt. alc. at 14.5° dis. 0.382 pt. D.s. eth.; s. bz.
3484	173	Sulphur-yellow	d,L-Phenylalanine Picrate, $(C_6H_{11}O_2N_2)_2Pk$. — \textcircled{P} Picrate* of No. 2.478-1. — Browns at 170°. 100 pt. c. sq. dis. 2.55 pt.; 100 pt. c. alc. dis. 1.3 pt.; less s. eth.
3485	171-4	Yellow(?)	ω -Phenylethylamine Picrate, $PhCH_2CH_2NH_2Pk$. — \textcircled{P} \textcircled{O} Picrate* of No. 2.1257. — Tetrag. pr. fr. alc.
3486	173-4	Steel-blue	p-Nitrosoaniline, $NO.C_6H_4NH_2$. — Curved ndl. fr. bz. — \textcircled{D} Readily reduced by Sn + HCl to p-phenylenediamine (No. 2.877).
3487	174	Purple-brown	p-Picrylaminophenol, $(NO_2)_2C_6H_2NH.C_6H_4OH$. — Ndl. E.s. alc.
3488	173-5; 183c.	Red-brown	α -Aminoazonaphthalene, α -Naphthaleneazo-4- α -naphthylamine, $C_{10}H_7.N_2C_{10}H_4.NH_2$. — \textcircled{P} S. w. dark green color in conc. H_2SO_4 , changing to blue & then to violet by addition of aq. Azo comp. reduced by Sn + HCl to α -naphthylamine (No. 2.589), & 1,4-diaminonaphthalene (No. 2.820). — Ndl. w. green metallic reflections. — Not e.s. alc., eth., bz. Dist. undecd. — Gives purple & golden-brown hydrochlorides.
3489	174 u.c.	Yellow	2,4-Dinitromethylaniline, $(NO_2)_2C_6H_4NHMe$. — \textcircled{P} \textcircled{O} Sec. amine giving nitroso deriv., m.p. 85°. — Ndl. fr. dil. alc. D.s. h. aq.
3490	174-5	Yellowish(?)	2,4,8-Trinitronaphthol(1), $(NO_2)_2C_6H_4OH$. — \textcircled{P} Nitro comp.‡ dyeing wool yellow (T. 2.16-a).
3491	171; 176		2,7-Dimethylacridine, $Me_2C_{12}H_7N$. — Colorless or nearly colorless ndl. fr. dil. alc. — \textcircled{P} E.s. w. green fluor. in bz., conc. H_2SO_4 , or gl. ac. ac.! Sol. in dil. ac. fluor. bluish-green. — $B_2H_6PtCl_6$, cryst. powd., i. aq., alc. — $B.HNO_3$, ndl., s. alc. w. yel. color & green fluor.
3492	d. 175	Red	Picryl- β -naphthylhydrazine, $(NO_2)_2C_6H_2N_2H_2C_{10}H_7$. — Pr. D.s. alc., eth.
3493	175-6	Or.-yel.	Benzaldehydeindogenide, $[NH.C_6H_4.CO.C(CH_2Ph)]^2$. — \textcircled{P} S. conc. H_2SO_4 w. deep red color; sol. alc.-KOH w. green-blue color. — Flat ndl. E.s. alc., chlf.; less s. eth. w. yel. green fluor.!
3484	175	Yellow	6-Amino-2,3-diphenylquinoxaline, $C_8H_5N_2$. — \textcircled{P} S. conc. H_2SO_4 w. deep red color, changing to green & yel.-red on diln.; mod. s. alc., eth., w. yel.-green fluor. — Cryst. fr. alc. — Acetyl deriv.,** silky scales fr. chlf., m.p. 252°.
3495	173; 178	Yellow	5-Nitro-8-hydroxyquinoline, $(NO_2)(HO)C_6H_4N$. — Dyes green w. Fe mordant. — Ndl. Vol. w. st. D.s. alc., eth.; e.s. h. HCl.
3496	175	Yellow	p-Aminobenzaldehydophenylhydrazone, $NH_2C_6H_4CH:N.NH-Ph$. — Lust. lft. fr. alc. E.s. alc., bz.; i. aq., lgr.
3497	175.5	Sulphur-yellow	2,4,6-Trinitroresorcinol, Styphnic Ac., $(NO_2)_3C_6H(OH)_2$. — \textcircled{P} Strong dibasic ac. w. yel. or or. "salts which explode more violently than picrates on ignition." — Hexag. cryst. fr. dil. alc. S. in 156 pt. aq. at 14°, or in 88 pt. at 62°; e.s. alc., eth. Aq. sol. gives no immediate ppt. (unlike picric ac.) w. ammon. $CuSO_4$ sol.
3498	175	Pale yellow	p-Nitrotriphenylurea, $NO_2C_6H_4NH.CO.NPh_3$. — \textcircled{P} Trace of HNO_3 prob. gives blue color w. sol. in warm conc. H_2SO_4 . — Tbl. s. alc.; i. aq.; alm. i. eth., lgr.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3499	174.5-5	Y-GY	Atropine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.797. — Lust. ndl. or plates fr. 1 pt. acetone + 5 pt. aq.
3500	176-7 u.c.	R	Indole Picrate, $\text{C}_8\text{H}_7\text{N.Pk.}$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1546. — Long lust. ndl. fr. bz. Melts to black liq. w. efferv.
3501	176	Red-brown	Pr-2-Methyl- β -naphthindole Picrate, $\text{C}_{11}\text{H}_{11}\text{N.Pk.}$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.2872. — Ndl. fr. bz.
3502	d. 176	Red-brown	Picryl- α -naphthylhydrazine, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{N}_2\text{H}_2\text{C}_6\text{H}_7.$ — Pr. fr. bz.
3503	175-6 w. efferv.	Light yellow	p-Aminocinnamic Ac., $\text{NH}_2\text{C}_6\text{H}_4\text{CH}=\text{CH.CO}_2\text{H.}$ — \textcircled{P} V. unstable, changing to red resin during evapn. of eth. sol. — V.s. h. aq.; e.s. alc., eth.
3504	177-8	Red-brown	4-Nitro-2-aminomethylaniline, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{NHMe.}$ — Lust. ndl. w. blue reflections.
3505	177	Yellow	2,4-Dinitro-2-amino-1,3-xylene, $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_4\text{Me.}$ — Ndl.
3506	177	Light yellow	Formyl-o-nitrophenylhydrazine, $(\text{CHO})\text{NH.NH.C}_6\text{H}_4(\text{NO}_2).$ — \textcircled{P} Sol. colored red to blue-violet by alk. — Ndl. fr. alc. E.s. h. aq., alc.; d.s. eth., bz.
3507	abt. 178	Scarlet	Picrylaniline, Phenyl-2,4,6-trinitrophenylamine, $(\text{NO}_2)_3\text{C}_6\text{H}_2\text{-NH.Ph.}$
3508	178	Or.-yel.	m-Nitrophenylhydroxylamine, $\text{NO}_2\text{C}_6\text{H}_4\text{NH(OH).}$ — Gran. powd. fr. bz. D.s.
3500	176-80	Brown-yel.	Diphenylazophenylene, $[\text{C}_6\text{H}_4\text{NPh.NPh}]$. — \textcircled{P} Dis. in mod. conc. HNO_3 w. fuchsin-red color, soon fading. Heated w. dil. H_2SO_4 + MnO_2 gives pungent odor of quinone. Lust. cryst. fr. alc. D.s. c. alc.; e.s. eth., bz. — Gives in CS_2 sol. w. 6 mol. Br a d.s. Br deriv., crystg. in short ndl., m.p. 243°.
3510	178	Steel-blue	6-Nitroso-m-toluidine, $(\text{NO})\text{Me.C}_6\text{H}_4\text{NH}_2.$ — \textcircled{P} Prim. amine. \textcircled{I} — Ndl. fr. bz. I. aq., lgr.; e.s. alc., eth.
3511	179	Scarlet	2,4-Dinitrophenyl- β -naphthylamine, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.C}_10\text{H}_7.$ — \textcircled{P} Prim. amine. \textcircled{I} — Pr. V.d.s. alc.; s. eth., bz.
3512	179-80	Yellow	β -Naphthylauramine, $\text{C}_{10}\text{H}_7\text{N:C(C}_6\text{H}_4\text{NMe}_2)_2.$ — [D.R.P. 44,077.] Cryst. powd. D.s. alc.
3513	179c., d.	Yellow	Glyoxalosazone, $\text{Ph.NH.N:CH.CH:N.NHPh.}$ — Monoclin. tbl. fr. eth. Alm. i. aq., alk., lgr., dil. min. ac.; s. h. alc., chlf., bz.
3514	178-80d.	Yellow	4-Ethylquinoline Picrate, $\text{Et.C}_6\text{H}_4\text{N.Pk.}$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1425. — Cryst.
3515	179	Light yellow	2,4-Lutidine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1198-1.
3516	179	Light yellow	5,7-Dinitroquinoline, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{N.}$ — Sbl. in glassy ndl. V.s. alc., eth., chlf. — B.HCl , m.p. abt. 86°.
3517	179	Yellowish	p-Dinitrobenzyl, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{C}_6\text{H}_4\text{NO}_2.$ — \textcircled{P} Nitro comp. \ddagger — Ndl. D.s. h. alc.; alm. i. c. alc.; d.s. eth., chlf. — \textcircled{P} Oxidn. by CrO_3 in gl. ac. ac. sol. gives p-nitrobenzoic ac. (No. 2.425).
3518	180	Red-brown	Bis-2,4-Dinitrodiphenylamine, $[(\text{NO}_2)_2\text{C}_6\text{H}_4]_2\text{NH.}$ — \textcircled{P} Dark red sol. in NaOH sol. evolves NH_3 when heated. — Ndl. D.s. alc.
3519	180-1	Or.-red	p-Nitrocinnamicaldehydephenylhydrazone, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}=\text{CH:N.NHPh.}$ — \textcircled{P} Hydrazone. \ddagger — Cryst. fr. abs. alc. I. aq.
3520	deflagrates at 180	Or.-yel.	3,3'-Diazoaminobenzoic Ac., $\text{CO}_2\text{H.C}_6\text{H}_4\text{NH.N}_2\text{C}_6\text{H}_4\text{CO}_2\text{H.}$ — \textcircled{P} Dibasic ac. — Cryst. gran. Alm. i. aq., alc., eth.; s. alk., pptd. unchanged by ac. — Boiled w. HCl gives m-aminobenzoic & m-chlorbenzoic ac. w. evolution of N.

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 ¶ = T. 2.35. ** = T. 2.1. ¶¶ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3521	180	Or.-yel.	Methenyl-bis-3-methyl-1-phenylpyrazolone(5) , $C_{12}H_{18}O_2N_4$. — \textcircled{P} S. in 4% NaOH sol. & pptd. fr. sol. by CO_2 . — Ndl. fr. alc. I. aq.; e.s. alc.; d.s. eth.
3522	180-1 (r.h.).	Yellow	Dibenzaldiphenylhydrotetrazone , $C_{18}H_{12}N_4$. — \textcircled{P} S. in conc. H_2SO_4 w. intense blue color! — Ndl. fr. bz. I. aq., alc.; d.s. h. bz. At m.p. changes gradually to dehydrobenzal-phenylhydrazone, m.p. 207-8 $^{\circ}$ c.
3523	180-1	Yellow	(β)-1,2-Naphthoquinonedioxime , $C_{10}H_8(:NOH)_2$. — \textcircled{P} S. alk. w. red-yel. color. Alc. sol. gives dark red ppt. w. ammon. $AgNO_3$ sol. — Ndl. — Heating w. dil. H_2SO_4 gives anhydride, ndl. fr. lgr., i. alk., m.p. 77 $^{\circ}$.
3524	180	Yellow	2,8-Dimethylquinoline Picrate , $C_{11}H_{11}N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1389. — Pr. fr. chlf. + alc.
3525	180.5-1.5 u.c.	\leftarrow YT1	Homatropine Picrate , $C_{13}H_{12}O_3N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.740. — Lft. fr. h. aq. E.s. h. aq.
3525	180-1u.c.	Y-YT1	Betaine Picrate . — \textcircled{P} \textcircled{D} Picrate* of No. 2.2619. — Fine ndl. fr. h. alc.
3527	180d. w. efferv.	Yellow	Rhamnosephenylosazone , $Me.(CH.OH)_2.C(:N.NHPh).CH:-NHPh$. — Ndl. fr. bz. I. c. or h. aq.; d.s. eth., bz.; more s. h. alc., ac. ac.; e.s. acetone. — Reduces boiling Fehling's sol. — $[\alpha]_D = +1^{\circ}, 24'$ (for 0.2 g. in 4 cc. pyridine + 6 cc. alc.).
3528	180-1	Light yellow	m-Aminocinnamic Ac. , $NH_2C_6H_4.CH:CH.CO_2H$. — \textcircled{P} Acid & prim. amine. — Long ndl. D.s. c. aq.; e.s. h. aq.; e.s. alc., eth. — $B.HCl$ (at 100 $^{\circ}$), lust. ndl. — $Ba\bar{A}_2\cdot 2H_2O$, lft. fr. dil. alc., v.s. aq.
3529	180	Yellowish	2,4,5,7-Tetranitronaphthol(1) , $(NO_2)_4C_{10}H_4.OH$. — \textcircled{P} Nitro comp.† s. w. yel. color in warm alk. — Lust. cryst. fr. gl. ac. ac. D.s. c., e.s. h. gl. ac. ac.; s. in 220 pt. bz. at 18 $^{\circ}$. — $K\ddot{A}\cdot 1\frac{1}{2}H_2O$, dark red pr. of metallic lust. s. in 340 pt. aq. at 19 $^{\circ}$.
3530	180-1	Greenish	α-Naphthostyryl , $[C_10H_8.NH.CO]^+$. — \textcircled{P} Sbl. in yel. ndl. — Ndl. fr. alc. D.s. h. aq.; s. alc.; r.d.s. eth.; i. c. Na_2CO_3 sol.; s. warm HCl . — Boiled w. ac. anhydride gives acetyl deriv., ndl. fr. alc., m.p. 125 $^{\circ}$.
3531	180-1	Beetle-wing lustre	Methylrosinduline , $C_{22}H_{17}N_3$. — \textcircled{P} S. conc. H_2SO_4 w. green color, turning red on diln. w. aq. — Thick pr. fr. bz. + lgr. More s. aq. than dil. alk.; e.s. eth., bz.
3532	181	Red	2-Dimethylaminoanthraquinone , $Me_2N.C_6H_4:(CO)_2:C_6H_4$. — [D.R.P., 108,837.] — \textcircled{P} Dyes wool orange in acid bath. — E.s. alc., gl. ac. ac.; v.s. bz.; alm. i. eth. Salts w. acids colorless & decd. by aq.
3533	181-2u.c.	RO →	Carbazole Picrate , $C_{12}H_9N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.2584. — Ndl. fr. toluene. V.d.s. c. bz., alc., but s. h. Decd. by alk., aq., or much alc. to components.
3534	181-2	Yellowish	3,5-Dinitroanisic Ac. , $(NO_2)_2C_6H_4.(OMe)(CO_2H)$. — \textcircled{P} Acid & nitro comp.† — Alm. i. c. aq.; e.s. h. aq.; e.s. alc. — 6 g. acid + 30 cc. abs. alc. + 0.5 g. Na disd. in 25 cc. alc., after 30 min. gives purple ppt. when treated w. 125 cc. lgr. — W. aq. at 170 $^{\circ}$ gives 2,6-dinitrophenol (No. 2.2926).
3535	181	Yellow	2,4-Dinitrophenyl-p-nitroaniline , $(NO_2)_2C_6H_4.NH.C_6H_4.NO_2$. — E.s. c. gl. ac. ac. (Fusion w. K_2S & S gives black wool dyestuff.)
3536	181-2	Yellow	Formaldehyde-p-nitrophenylhydrazone , $CH_2:N.NH.C_6H_4.NO_2$. — \textcircled{P} Hydrazone.† — Lust. ndl. fr. cooling h. bz. sol.; or shimmering ndl. fr. h. aq., in which it is d.s.
3537	181-2	Yellow	Tetramethyldiaminoacridine , $(Me_2N)_4C_{12}H_7N$. — Ndl. E.s. aq., alc.; d.s. lgr. — $B.HCl$, ndl. — $B_2H_2PtCl_6$, red-brown ndl.
3538	182-3	Red	Azodicarbonanilide , $Ph.NH.CO.N_2.CO.NHPh$. — Ndl. clusters fr. acetone. — V.d.s. aq.; e.s. alc., eth.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3539	182-3	Red-yel.	Fluoranthene Picrate, $C_{12}H_{10} \cdot Pk.$ — \textcircled{P} \textcircled{D} Picrate* of h. c. of Vol. I. — Ndl. s. h. alc.
3539-I	182; 188; 176	Y-OY	2,4-Dinitroaniline, $(NO_2)_2 \cdot C_6H_4 \cdot NH_2$. — \textcircled{P} Boiled w. dil. KOH sol. gives red coloration. Boiled w. conc. KOH sol. is split to NH_2 & 2,4-dinitrophenol, No. 2.3126. — Monoclin. cryst. w. bluish reflections. I. c. aq.; d.s. h. aq.; s. in 132.6 pt. alc. at 21 $^{\circ}$.
3540	181.5-2.5 u.c.	YT1	9-Phenylacridine, $Ph \cdot C_13H_8N$. — \textcircled{P} Dis. 2 mg. in 1 cc. conc. HCl & dil. w. 20 cc. aq. The light yel. sol. shows brilliant G-YG fluor. when viewed in strong light w. black background. — B.p. 403-4 $^{\circ}$. E.s. bz.; s. eth.; d.s. alc.; i. aq.
3541	183 u.c.	Red	β -Anilino- α -naphthoquinoneanilide, $C_{22}H_{14}ON_2$. — \textcircled{P} E.s. conc. HCl w. violet color. Salts gold-green to violet-black, sol. undecd. in alc., but hydrolyzed by aq. — Ndl. fr. alc. I. aq.; v.d.s. alc.; more s. eth., chlf., bz., lgr.; i. dil. acids. Unchanged by boiling alk. — Zn dust w. ac. ac. yields naphthalene & aniline.
3542	183	Or.-yel.	Dinitropseudocumidine, $(NO_2)_2 \cdot C_6Me_3(NH_2)$ [Me ₂ :NH ₂ = 1,2,4:5]. — Ndl. fr. alc.
3543	183-4d.	Ocher-yellow	d,l-Ornithine Picrate, $C_6H_{12}O_2N_2 \cdot Pk. \cdot 2\frac{1}{2}H_2O$. — Lust. plates.
3544	182-4	Yellow	4,4'-Diaminoazoxybenzene, $NH_2 \cdot C_6H_4 \cdot N_2O \cdot C_6H_4 \cdot NH_2$. — Ndl. Forms salts w. acids. — Reduced by Sn + HCl to p-phenylenediamine (No. 2.877).
3545	183	Yellow	Acetyl amidrazone, $Me \cdot CO \cdot C(NH_2) : N \cdot NH \cdot Ph$. — Ndl. E.s. acids! D.s. aq., bz., lgr.
3546	183	Light yellow	5-Nitrofuran-3-carbonic Ac., $NO_2 \cdot C_4H_2O \cdot CO_2H$. — \textcircled{P} Acid. — Rectangular tbl. fr. aq. Sbl. in lft. E.s. alc., eth. — Reduction w. Sn & HCl gives succinic ac. (Vol. I), NH_2 , & CO_2 . — Ba $\bar{A}_2 \cdot xH_2O$, yel. lft., r.d.s. c. aq.
3547	183-5d.	Dark red	2,4,6-Trinitrohydrazobenzene, Picrylphenylhydrazine, $(NO_2)_3 \cdot C_6H_3 \cdot NH \cdot NHPh$. — Deflagrates when strongly heated. — Short ndl. fr. acetone. D.s. h. alc.
3548	181; 184c.	Red	Rubazonic Ac., $C_{20}H_{17}O_2N_2$. — \textcircled{P} S. alk. w. deep violet color. Reducing agents give leuco deriv., the color being restored by air. — Ndl. fr. gl. ac. ac. I. aq., dil. acids; d.s. alc., gl. ac. ac. s. bz., eth., chlf.
3549	184-5	Or.-red	Acetophenone-p-nitrophenylhydrazone, $MePh \cdot C : N \cdot NH \cdot C_6H_4 \cdot NO_2$. — \textcircled{P} Hydrazone. \ddagger — Silky ndl. I. aq.; s. alc., bz., acetone. D.s. w. pale rose color in warm NaOH sol., color deepening on addition of alc.
3550	184-5	Orange	$\beta\beta$ -Dinaphthyl Picrate, $C_{20}H_{14} \cdot Pk.$ — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Ndl.
3551	184	Yellow	Dinitrobiphenol, $(NO_2)_2(OH)C_6H_4 \cdot C_6H_4(OH)(NO_2)_2$. — \textcircled{P} S. alk. w. intense blood-red color! — Ndl. fr. bz. Sbl. I. aq.; v.d.s. c. alc., eth. Alk. salts, black, amorph., w. green metallic reflections.
3552	184; 187	Red	p-Methylisatin, $C_9H_7O_2N$. — \textcircled{P} . S. alk. w. deep violet color! — Lit. fr. aq. D.s. c. aq.
3553	185	Fire-red	2,5-(β)-Dinitromethyl-p-toluidine, $(NO_2)_2 \cdot C_6H_3 \cdot Me(NHMe)$. — \textcircled{P} Sec. amine \ddagger gives nitrosamine of m.p. 123-4 $^{\circ}$. — Pr. w. green reflection fr. acetone. V.s. bz., chlf.; d.s. eth., lgr.
3554	185	Or.-yel.	Carbopetrocene Picrate, $C_{14}H_{10} \cdot Pk.$ — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Ndl. Decd. by aq. or alc.
3555	185	Yellow	Formopyrine Picrate, $C_{12}H_{14}O_2M_4 \cdot Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.2357. — Ndl. fr. alc.

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(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3556	185	Yellow	Nitrosocorydaline, $C_{11}H_{12}O_3N.NO$. — \textcircled{P} Prob. gives blue color in T. 2.15. — Ndl. fr. alc. D.s. aq., eth.; e.s. chlf.
3557	185.5	Yellow	5-Nitro-2-acetaminonaphthalene, $NO_2.C_{10}H_8.NH(C_2H_5O)$. — Tbl. fr. alc.; ndl. fr. bz.
3558	184-6u.c.; 188-90c.	Yellowish	6-Nitroveratric Ac., $NO_2.C_6H_5.(OMe)_2^{24}(CO_2H)^1$ — \textcircled{P} Acid & nitro comp.† — Ndl. s. in 25 pt. h. aq.; e.s. alc., eth., bz.; i. lgr. N. Eq. = 227. — Ag \ddot{A} . pale yel. ppt. cryst. fr. h. aq.
3555	186-7	Dark red	Benzaldehyde-o-nitrophenylhydrazone, $Ph.CH:N.NH.C_6H_4-NO_2$. — \textcircled{P} Hydrazone.‡ — Lft. fr. h. bz. I. aq., eth.; d.s. h. alc.
3560	186	Deep red-brown	2,4,6-Trinitrophenylhydrazine, $(NO_2)_3C_6H_3.NH.NH_2$. — \textcircled{P} S. aq. NaOH sol. w. deep blue color! — Cryst. fr. alc. Alm. i. aq., eth., bz.; d.s. c. alc.
3561	186	Yellow	Rhamnose-p-nitrophenylhydrazone, $Me.(CH.OH)_4.CH:N.NH-C_6H_4.NO_2$. — Ndl.
3562	186; 188c.; (196-7)	Yellow	d-Galactosephenylosazone, $C_{12}H_2O_4N_4$. — \textcircled{P} Hydrazone react.‡ — Ndl. fr. alc. Alm. i. c. aq., eth., chlf., bz. $[\alpha]_D = +0.48^\circ$ (in pyridine-alc.).
3563	186	Yellow	4,6-Dinitro-2-amino-1,3-dimethyl-5-tert.-butylbenzene, $(NO_2)_2-(NH_2)Me_2.C_6.(CMe_3)$. — Cryst. powd. Mod. s. alc., bz., lgr.
3564	186u.c. (r.h.)	Pale yellow	p-Nitrobenzoyl-d-(or l)-serine, $NO_2.C_6H_4.CO.NH.CH(CO_2H).-CH_2.OH$. — Sinters abt. 168° u.c. Lust. plates fr. aq.
3565	187-7.5d.	Red	Juglonoxime, $(HON:)C_6H_3O.OH$. — \textcircled{P} S. conc. H_2SO_4 or dil. NaOH w. intense blood-red color! — Lust. ndl. V.s. h. alc., v.d.s. aq.; s. eth.
3567	187	Gold- yellow	m-Nitrobi-o-cresol, $NO_2.C_6H_3Me(OH).C_6H_3Me(OH)$. — \textcircled{P} Phenolic nitro comp.† — Ndl. fr. aq. E.s. c. alc., eth.; less s. bz.
3568	187	Yellow	3-Methylquinoline Picrate, $C_{10}H_8N.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1388. — Ndl.
3569	187-8	Yellow	2-Phenylquinoline Picrate, $C_{12}H_{11}N.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.698. — Lft. fr. alc.
3570	187	Yellow	Tolazon, $[C_6H_4(Me).N:N.C_6H_4(Me)]^2$. — \textcircled{P} Sol. w. yel. color in ac. — Lust. pr. fr. alc. B.p. a. 360° (alm. undec.). E.s. alc.; v.s. chlf., bz.; v.d.s. lgr.
3571	190; 186	Alizarine- red	$\alpha\alpha$ -Azonaphthalene, $C_{10}H_7.N.C_{10}H_7$. — \textcircled{P} \textcircled{D} Azo comp. s. undec. w. blue color in conc. H_2SO_4 , sol. fluor. brick-red at 180°. — Ndl. fr. gl. ac. ac. w. greenish reflections. Sbl. in yel. lft., becoming cinnabar-red on rubbing.
3572	188-9	Or.-yel.	Acetylformazan, $Ph.N(C_6H_5O)N:CH.N:NPh$. — \textcircled{P} S. conc. H_2SO_4 w. cornflower-blue color. — Bronzy ndl. fr. alc. V.s. chlf.
3573	188	Bronze- yellow	Nitrobinaphthyl, $NO_2.C_{20}H_{13}$. — Lust. lft. fr. bz. E.s. h. bz.; less s. alc., eth.
3574	188	Dark yellow	2,4,6-Trinitroaniline, Picramide, $(NO_2)_3C_6H_3.NH_2$. — \textcircled{P} Decd. to NH ₂ & picric ac. (No. 2.3168), by boiling KOH sol. (T. 2.26).
3575	188	Yellow	β -Naphthylpiperidine Picrate, $C_{12}H_7N.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.615. — Cryst. ppt. S. h. alc.; e.s. eth., bz., chlf.; i. lgr.
3576	188	Yellow	Antipyrine Picrate, $C_{11}H_{12}ON_2.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1946. — Ndl. fr. aq.
3577	189-90d.	Red-brown	2-Nitro-p-tolylglycine, $NO_2.C_6H_4Me.NH.CH_2.CO_2H$. — Pr. V.d.s. c. aq., eth.; e.s. h. alc. — Pb $\ddot{A}.H_2O$, purple-red ndl., i. c. aq.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3578	188.5-9.5 u.c.	O	α -Naphthol Picrate, $[C_{10}H_7OH]_2PK$. — \textcircled{P} \textcircled{D} Picrate* of species of Vol. I. — Ndl. fr. dil. alc.
3579	189d.	Lemon-yellow	Nitron, 1,4-Diphenyl-3,5-endoanilohydrotriazole, $C_{20}H_{14}N_4$. — [Used in quant. determination of nitrates. Original portion of description is based on a sample fr. Kahlbaum of OYS1 color & m.p. 179.5-80.5° u.c., which formed garnet-red ndl. when recryst. fr. alc. Remaining portion is fr. Busch (Ber., 38, 858).] — Lft. or tbl. fr. alc. E.s. chlf.; v.d.s. alc., bz.; d.s. eth. Odorless. Taste bitter. Alc. sol. soon becomes RO-ROSI. — \textcircled{D} Add sol. of 0.1 g. substance diss. in 1 cc. 5% ac. ac. to 5 cc. 1% KNO_3 sol. Wash the voluminous nearly white ppt. w. 10 cc. c. aq. Recryst. fr. 25 cc. boiling aq. Wash w. 5 cc. c. aq. Dry at 100°. The product, $B.HNO_3$, melts w. blackening at 259.4° u.c.
3580	abt. 189	Yellow	β -Naphthoquinoneoxime(1)-semicarbazone(2), $HON:C_{10}H_4:N-NH.CO.NH_2$. — \textcircled{P} Semicarbazone.†† — Ndl. S. alc.; i. aq.
3581	190-1	Red	2-Anilinonaphthoquinone, $Ph.NH.C_9H_4O_2$. — \textcircled{P} Dis. w. purple color in alc. KOH. — Lust. ndl. Sbl. undecd. E.s. h. alc.; e.s. eth., bz.; alm. i. lgr.; i. c. NaOH.
3582	190	Copper-red	4,6-Dinitro-2-aminoresorcinol, $(NO_2)_2(NH_2)C_6H_4(OH)_2$. — Lft. fr. abs. alc. Alm. i. aq.; r.d.s. alc.; d.s. dil. ac.; e.s. alk. or conc. H_2SO_4 , being pptd. unchanged fr. latter by aq.
3583	190.5	Or.-red	2,4-Dinitrophenyl- α -naphthylamine, $(NO_2)_2C_6H_4.NH.C_10H_7$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Lust. ndl. I. aq.; d.s. c. alc.; e.s. bz., chlf.
3584	190u.c.	YO-YOS1	Benzaldehyde-p-nitrophenylhydrazone, $Ph.CH:N.NH.C_6H_4-NO_2$. — \textcircled{P} Hydrazone.†† A few mg. dis. in 2 cc. 10% aq. NaOH to OR sol. becoming deep red when diluted w. 4 cc. alc. — Lust. ndl. fr. h. alc. Odorless. I. c. aq.; e.s. alc., bz., lgr., eth.
3585	d. 190u.c.	OY	Guanine Picrate, $C_6H_4ON_2PK$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.2659. — Lust. ndl. fr. h. aq. acidulated w. HCl. Alm. i. c. aq.
3586	189-91u.c., d.	Yellow	Thebaine Picrate. — Picrate* of No. 2.982. — Cryst. powd. fr. h. 10% ac. ac.
3587	190	Yellow	Anhydrodi-o-aminobenzophenone, $C_{10}H_{12}N_2$. — Lust. lft. fr. alc. Sbl. Conc. HCl at 160° gives o-aminobenzophenone (No. 2.3087).
3588	190u.c.	Light yellow	p-Azoxybenzaldehyde, $CHO.C_6H_4.N.O.C_6H_4.CHO$. — \textcircled{P} S. conc. H_2SO_4 w. or. color. — Ndl. fr. bz. D.s. h. aq., c. alc., or lgr.
3589	190.5	Pale yellow	Acetdinitro-p-toluide, $(C_6H_5O.NH)(NO_2)_2.C_6H_5Me$. — Ndl.
3590	d. abt. 190	Yellowish green	Benzidine Picrate, $NH_2.C_6H_4.N_2.C_6H_4NH_2.2PK$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.840. — Cryst. fr. abs. alc. Slowly decd. by dil. alc.
3591	191	Light yellow	Quinaldine Picrate, $C_{10}H_9N.PK$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1376 — Cryst. d.s. aq., c. alc.
3592	191-2	Light yellow	trans-o-Dinitrostilbene, $NO_2.C_6H_4.CH:CH.C_6H_4.NO_2$. — Ndl. fr. chlf. V.d.s. alc., eth., lgr.; more s. h. bz.
3593	192-2.5	Dark red	Bilirubin, $(C_{10}H_{16}O_2N_2)_2$. — [Combined w. Ca in gall stones.] Rhomb. tbl. or pr. fr. chlf. Alm. i. aq.; v.d.s. eth., bz., CS, amyl alc.; sl. more s. alc.; e.s. alk.‡; s. in 567 pt. c. chlf.; s. in 30.9 pt. boiling, or 112.6 pt. c. dimethylaniline fr. which it cryst. in broad columns truncated obliquely at both ends. Solutions are brownish red. Is extracted by dil. aq. NaOH fr. chlf. sol. $CaCl_2$, $BaCl_2$, $PbAc_2$ & $AgNO_3$ give brownish ppt. w. ammon. sol. Gmelin's T. (for bilirubin or biliverdin). — By aid of a pipette bring conc. some HNO_3 beneath aq. sol. of

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§ = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

(ORDER II.)

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
			compound in narrow glass cylinder so that liquids shall only mix v. slightly at contact zone. Sol. above HNO_3 becomes colored in yel.-red, red, violet, blue, & green layers. The react. is still shown at dilution 1 : 80,000, & is not prevented by presence of albumin. — Ehrlich's T. (Z. klin. Medizin [4] (1883), 721). — Add to chlf sol. of compound an equal vol. p-diazobenzensulphonic ac. sol. (freshly prepared fr. 1 g. sulphanilic ac. in 1000 cc. aq. + 15 cc. HCl + 0.1 g. NaNO_2) & enough alc. to give homogeneous sol. Sol. becomes red, & on addition of conc. HCl , violet, & then blue. On adding alk. color becomes red, & w. x.s., green-blue.
3584	192-3d.	Dark red	$\text{o-Nitrophenylglycine}$, $(\text{NO}_2\text{C}_6\text{H}_4)\text{NH}.\text{CH}_2\text{CO}_2\text{H}$. — \textcircled{P} Nitro comp.† & acid. — Pr. fr. alc. D.s. aq., eth.; e.s. h. alc.
3595	192	Orange; egg-yel.	$\text{Methylguanidine Picrate}$, $\text{C}_6\text{H}_5\text{N}.\text{Pk}$. — \textcircled{P} Picrate of strongly alk. deliq. NH : $\text{C}(\text{NH}_2)(\text{NHMe})$. — 4 or 6-sided pleochroic tbl. fr. aq. V.s. aq.
3596	abt. 192d.	Yellow	$\text{o-Nitrobenzaldehyde-2,4-dinitrophenylhydrazone}$, $\text{NO}_2\text{C}_6\text{H}_4\text{-CH: N.NH.C}_6\text{H}_3(\text{NO}_2)_2$.
3597	192-5	Red-brown	$\text{Dinitrophenyl-}\beta\text{-naphthylamine}$, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{N}$. — Cryst. fr. gl. ac. ac.
3598	191-5	Orange	$\text{4-Nitro-}\alpha\text{-naphthylamine}$, $\text{NO}_2\text{C}_6\text{H}_4\text{NH}_2$. — \textcircled{P} Nitro comp.‡ & prim. amine yielding NH_3 & 4-nitro- α -naphthol (No. 2,3424) when boiled (T. 2.26) w. KOH sol. — Thin ndl. fr. alc. More s. h. ac. than h. aq., separating unchanged on cooling.
3599	193-4	Orange	$\text{m-Nitroazobenzene-}\beta\text{-naphthol}$, $\text{NO}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OH}$. — \textcircled{P} Nitro† & azo comp. — Scales w. metallic lustre fr. toluene. — Acetate,** hair-like red ndl. fr. alc., m.p. 161-2°.
3600	193	Gold-yellow	$\text{4,6-Dinitro-m-toluidine}$, $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}$. — \textcircled{P} Nitro comp.† & prim. amine — Small hard cryst. fr. gl. ac. ac. I. aq.; v.d.s. eth., bz.; h. alc.; i. conc. HCl .
3601	193-4	Y	Cinchonine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2,1039-1. — Flat ndl. slowly separating fr. h. dil. ac. ac. sol. after cooling.
3602	abt. 193d.	Pale greenish yellow	$\text{4-Nitrosonaphthol(1)}$, " α -Naphthoquinoneoxime," $\text{NO.C}_6\text{H}_4\text{-OH}$. — Ndl. E.s. alc., eth.; d.s. chlf., h. bz.; e.s. Na_2CO_3 sol. & pptd. by CO_2 ! Salts unstable.
3603	191-2; 193	Green-black	α -Naphtholdisazo-2,4-benzene, $\text{HO.C}_6\text{H}_4(\text{N}_2\text{Ph})_2$. — \textcircled{P} Azo comp. — Bronzy ndl. fr. fusil oil. D.s. alc., lgr.; e.s. bz.; v.s. chlf. — Acetyl deriv., bronzy brown lft. fr. alc., m.p. 159-60°.
3604	194	Or.-yel.	$\text{p-Nitrophenylpyruvic Ac.}$, $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{CO.CO}_2\text{H}$. — [D.R.P. 92,794.] — \textcircled{P} FeCl_3 colors aq. sol. bluish green. — Cryst. w. 1 mol. ac. ac. fr. gl. ac. ac. D.s. aq.; e.s. alc., eth.; d.s. bz., chlf. — Alkali salts deep brown-red.
3605	abt. 194	Yellow	$\text{2,4-Dinitrophenyl-m-nitroaniline}$, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.C}_6\text{H}_4\text{NO}_2$. — Ndl. fr. gl. ac. ac. D.s. h. alc.
3606	194	Yellow	$\text{4-Isonitroso-3-methylpyrazolone}$, " $\text{NH.N : CMe.C(:NOH).CO}$ ". — \textcircled{P} Titrates as monobasic acid. — Silky ndl. S. aq., alc. AgA, dark red ppt., ndl. fr. gl. ac. ac., d. a. 130°.
3607	194	Yellow	$\text{1,3,6,8-Tetranitro-4-naphthylamine}$, $(\text{NO}_2)_4\text{C}_6\text{H}_4\text{NH}_2$. — Ndl. D.s. h. alc., bz.
3608	193-5	Yellow	Dinitro-2-amino-1,3,5-trimethylbenzene, $(\text{NO}_2)_2(\text{NH}_2)\text{Me}_2\text{C}_6$. — Ndl. S. h. alc.; alm. i. h. aq.; s. conc. HCl ! — Acetyl deriv.,** m.p. 275°, s. in 20 pt. h. alc.
3609	194-5	Light yellow	(γ)-1,3,5,8-Tetranitronaphthalene, $(\text{NO}_2)_4\text{C}_10\text{H}_4$. — \textcircled{P} Warmed w. alk. is colored red. Lust. tetrahedra fr. acetone. D.s. alc., chlf.; e.s. acetone; s. conc. HNO_3 or H_2SO_4 . — Oxidn. (Ber., 28, 375) gives p-dinitrophthalic ac. (No. 2,334).

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3610	195-6sl. d.	Carmine-red	1-Tryptophane Picrate, $C_{11}H_{12}O_2N_2.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.2605. — Ndl. & tbl. fr. sq. 0.91 pt. s. in 100 pt. c. sq.; 1 pt. s. in 100 pt. eth.; e.s. alc.
3611	195.5	Red-yel. or red	3,3'-Dinitrodiazoaminobenzene, $NO_2.C_6H_4.N_2.H.C_6H_4.NO_2.$ — \textcircled{P} Warmed w. HCl evolves N. — Pr. V.d.s. alc., eth.; mod. s. alc. — C. conc. HCl gives m-nitroaniline & m-nitrodisazobenzene chloride.
3612	195	Or.-red	Cinnamicaldehyde-p-nitrophenylhydrazone, $Ph.CH:CH.CH:NH.C_6H_4.NO_2.$ — \textcircled{P} Hydrazone†† s. in alc. NaOH w. deep violet-red color! — Cryst. I. aq.; s. alc., acetone, bz.
3613	195	Brown	Benzenedisazobenzene- β -naphthol, $Ph.N_2.C_6H_4.N_2.C_{10}H_8.OH.$ — \textcircled{P} Azo comp. S. conc. H_2SO_4 , w. yel. color. — Bronzy lft. fr. h. gl. ac. ac. I. aq., alk.; d.s. alc. — Reduced by Sn + HCl to aniline, p-phenylenediamine, & amino- β -naphthol.
3614	195	Gold-yellow	Methylene-2,2'-dinitrodiphenyldiamine, $CH_2.(NH.C_6H_4.NO_2)_2.$ — \textcircled{P} Nitro comp. Ndl. fr. alc.; i. aq., c. alc.
3615	195	Yellow	m-Nitrophenylurea, $NO_2.C_6H_4.NH.CO.NH_2.$ — \textcircled{P} Nitro comp.‡ — Cryst. fr. h. aq. — \textcircled{D} Sapn. (T. 2.26) products: NH ₃ , CO ₂ , & m-nitroaniline (No. 2.3125).
3616	195.5	Yellow	8-Nitroacet- β -naphthalide, $NO_2.C_{10}H_8.NH(C_2H_3O).$ — \textcircled{P} Nitro comp.‡ — Ndl. D.s. alc. bz. — \textcircled{D} Sapn. (T. 2.26) products: ac. ac. & No. 2.3073.
3617	195d.	Light yellow	1,6-Dinitronaphthol(2), $(NO_2)_2.C_{10}H_6.OH.$ — \textcircled{P} Nitro comp.‡ — Ndl. V.d.s. boiling aq.; more s. alc.; v.s. eth., chlf. Salts generally v.d.s. aq. — KÄ.2H ₂ O, yel. ndl. — AgA, intense yellowish-red.
3618	195	Light yellow	Trinitro-p-diphenylbenzene, $(NO_2)_3.C_{12}H_11.$ — \textcircled{P} Nitro comp.‡ — Rhomb. ndl. fr. gl. ac. ac. — Reduced by Sn + HCl to comp. of m.p. 169.5°.
3619	d. abt. 195 u.c.	G-BG	Nitrosoantipyrine, $[NPh.CO.C(NO):CMe.NMe].$ — \textcircled{P} Gives heavy Ag ppt. in T. 2.21 (tests w. Tollen's & diphenylamine reags. giving negative results). — Ndl. fr. h. acetone. Turns yel. & then dark brown on heating, finally decomposing w. efferv. D.s. aq., alc., chlf. Sol. in conc. H_2SO_4 , yellow.
3620	195	Dark reddish violet	Hexamethyl-p-rosaniline, $(Me_3N.C_6H_4)_3.C.OH.$ — \textcircled{P} Crystd. salts show beetle-wing colors & dye tannin-mordanted cotton blue-violet. (Cf. No. 3.229 under "Crystal Violets.") — Cryst. fr. bz. contain bz. of cryst. & effloresce in air. I. aq.; s. eth., iqr.; r.d.s. alc.
3621	196	Dark red	p-Nitrotoluicaldehyde-p-nitrophenylhydrazone, $Me.C_6H_4.CH:NH.C_6H_4.NO_2.$ — \textcircled{P} Hydrazone†† — "Ndl. w. green fluor."
3622	196-7	Lemon-yellow	5-Nitro-o-acettoluide, $NO_2.C_6H_4.Me(NH.C_6H_4.O).$ — \textcircled{P} \textcircled{D} Sapn. by boiling mixt. of 1 vol. conc. H_2SO_4 + 3 vol. aq. (Ann., 158, 345, & T. 2.26-b) gives ac. ac. & 5-nitro-o-toluidine (No. 2.3018).
3623	195; 196	Yellow	Dinitrophenolphthalein, $C_{10}H_{12}O_3N.$ — [D.R.P., 52,211.] — Ndl. D.s. alc.; s. alk. w. or.-yel. color.
3623-I	196-6.5	Yellow	2,2'-Dinitrodiazoaminobenzene, $NO_2.C_6H_4.N_2.H.C_6H_4.NO_2.$ — Cryst. D.s. h. alc.
3624	195.5-6 u.c., d.	Y	Codeine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.903. — Short ndl. fr. h. dil. ac. ac.
3625	197	Red	3,3'-Diamino-4,4'-dimethylazobenzene, $(NH_2)Me.C_6H_4.N_2-C_6H_4.Me(NH_2).$ — \textcircled{P} Azo comp. & prim. amine.¶ — Ndl. fr. alc. D.s. aq.; e.s. alc., eth. — B.2HCl, light red tbl. — Diacetyl deriv., ** m.p. 300°.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. ‡ = T. 2.36. ¶ = T. 2.34. ¶ = T. 2.35. ** = T. 2.1. †† = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3626	195; 198	Dark red	4-Nitro-o-phenylenediamine, $\text{NO}_2\text{C}_6\text{H}_3\text{(NH}_2)_2$. — Ndl. — $\text{B.HCl}\text{H}_2\text{O}$, yel.-brown ndl. — $\text{B.H}_2\text{PtCl}_6$, brown-red pr.
3627	198 u.c.	OR-R	Picryl- α -naphthylamine, $(\text{NO}_2)_3\text{C}_8\text{H}_4\text{NH.C}_1\text{0H}_7$. — Ndl. fr. alc., I. aq.
3628	197-8	Or.-yel.	3,3'-Dinitrobiphenyl, $\text{NO}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NO}_2$. — P Nitro comp. — Ndl. fr. alc. I. aq.
3629	197-8; 193-4d.	Yellow	Acetyl-2,4-dinitrophenylhydrazine, $(\text{NO}_2)_2\text{C}_6\text{H}_3\text{NH.NH(C}_2\text{H}_5\text{O)}$. — Silky ndl. fr. dil. alc.; v.d.s. eth., bz.
3630	197	Bronzy	α -Tolylrosinduline, $\text{C}_{20}\text{H}_{11}\text{N}_3$. — [Pat.] Cryst.
3631	189-90(r.h.) 203-4(sl.h.)	Bluish reflections	2-Anilinoaposafranine, "Induline $\text{C}_8\text{H}_5\text{N}_4$," $\text{C}_8\text{H}_5\text{N}_4$. — P Sol. in conc. H_2SO_4 , red-violet. Aq. sol. of salts rose colored & non-fluor. — Shimmering pr. fr. alc. E.s. alc., bz.; d.s. eth., lgr. — B.HCl , ndl. fr. alc. w. green beetle-wing reflections.
3632	198	Red	4-Nitro-4'-aminobiphenyl, $\text{NO}_2\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NH}_2$. — P Nitro comp. † & prim. amine. — Ndl. fr. alc. Alm. i. h. aq.; e.s. h. alc. — P Oxidn. by CrO_3 in gl. ac. ac. gives p-nitrobenzoic ac. (No. 2.425). Reduen. by Sn & HCl should give benzidine (No. 2.840).
3633	198-9	RO-O	Isatine, Anhydride of α -Aminobenzoylformic Ac., $[\text{CO.C}_6\text{H}_4\text{N}:\text{C(OH)}]^2$. — Lust. pointed ndl. fr. h. aq. V.d.s. c. aq.; s. h. aq.; d.s. eth.; e.s. h. alc. P (1) ("Indophenine Reaction"). Dis. 1 mg. in 5 cc. bz. to which a small drop of thiophene has been added, & shake. An opaque sol. is formed in the H_2SO_4 layer, which appears green or blue when diluted w. much more H_2SO_4 . — (2) A few mg. fused w. a little solid KOH emits distinct odor of aniline. — (3) 1 mg. shows ORS1 color on tile when treated w. 1 drop 10% NaOH sol. P Dis. 0.05 g. in 10 cc. aq. Add 2 drops phenylhydrazine & allow to stand some min. Filter off the fine yel. ppt. Wash w. 2 cc. aq. Cryst. fr. 7 cc. boiling 95% alc. Wash the beautiful OY ndl. that separate on cooling w. 1 cc. 95% alc. Dry on tile 15 min. at 100°. The product, isatinephenylhydrazone, melts to clear yel. liq. at 210° u.c. (215° c.)!
3634	198-9	Red-brown	Rosinduline, $\text{C}_{20}\text{H}_{11}\text{N}_3$. — P Sol. in conc. H_2SO_4 , green, becoming red on diln. w. aq. Alc. sol. of salts fluor. fiery red! — Bronzy lft. fr. eth. I. aq.; e.s. alc., eth., bz. — B.HCl . $\frac{1}{2}\text{H}_2\text{O}$, red ndl. w. green reflections.
3635	198d.	Yellow: red	2,4-Dinitrophenylhydrazine, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.NH}_2$. — P Reduces Tollen's reag. in T. 2.30. — Yel. lft. & red pr. I. aq.; v.d.s. c. alc.; s. aq.-NaOH w. deep red color.
3636	198	Yellow	3-Nitro-p-coumaric Ac., $\text{NO}_2\text{C}_6\text{H}_3\text{(OH).CH:CH.CO}_2\text{H}$. — P Acidic nitro comp. † — Ndl. fr. alc. — Salts are red. — EtA, ndl. fr. alc., m.p. 108.5°.
3637	198d.	Yellow	1,8-Dinitronaphthol(2), $(\text{NO}_2)_2\text{C}_{10}\text{H}_8\text{OH}$. — P Acidic nitro comp. † — Lust. ndl. fr. dil. alc. E.s. alc., chlf. — $\text{Ba}_2\text{H}_2\text{O}$, or. ppt. — Oxidn. by alk. KMnO_4 gives 3-nitrophthalic ac. (No. 2.389).
3638	199	Yellow	4,1'-Dinitrostyrene, $\text{NO}_2\text{C}_6\text{H}_4\text{CH:CH.NO}_2$. — P Reacts vigorously w. conc. H_2SO_4 at 100° evolving CO. — Cryst. fr. gl. ac. ac. I. aq.; s. caustic alk.; d.s. organic solvents.
3639	199	Yellow	2-Nitro- α -acetnaphthalide, $(\text{NO}_2)_2\text{C}_{10}\text{H}_8\text{NH(C}_2\text{H}_5\text{O)}$. — Ndl. R.d.s. alc.; more s. gl. ac. ac.
3640	199-200	Yellow	Glycocyamine Picrate, $[\text{NH}_2\text{C}(\text{NH})\text{NH.CH}_2\text{CO}_2\text{H}] \text{Pk}$. — Picrate* of compound s. in 126 pt. c. aq., which at 300° is black without having melted. — Fine ndl. V.d.s. aq.
3641	199-200	Yel.-green	Resacetophenoneoxime, $\text{Me.C}(\text{NOH})\text{O.C}_6\text{H}_4\text{(OH)}_2$.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3642	abt. 200	Red	5-Nitro-1-aminoanthraquinone, $\text{NO}_2\text{C}_4\text{H}_3:(\text{CO})_2:\text{C}_4\text{H}_4\text{NH}_2$. — \textcircled{P} S. in 70% H_2SO_4 w. red color, changing to blue on standing! — Cryst. S. alc.
3643	198-9; 203-4	Yellow	d-Ornithine (α - δ -Diaminovaleric acid) Picrate, $\text{C}_8\text{H}_{12}\text{O}_2\text{N}_2\text{Pk}$. — Picrate* of syrupy comp. e.s. aq., i. eth.
3644	200-1	Yellow	d,l-Arginine (α -Amino- δ -guanidinovalerianate) Picrate, $[\text{C}_8\text{H}_7\text{O}_2\text{N}_4\text{Pk}]_2$. — \textcircled{P} Picrate.* — Lust. pr. 100 cc. aq. dis. 0.22 pt. at 16°.
3645	200	Sulphur-yellow	8-Methylquinoline Picrate, $\text{C}_9\text{H}_8\text{N}\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1382. — Lft. D.s. alc.; less s. eth., bz.
3646	200d.	Yellow	Ethylpicrazide, $\text{Et.NH}_2\text{C}_6\text{H}_5(\text{NO}_2)_2$. — \textcircled{P} Heated w. KOH sol. evolves ethylamine (No. 2.1062). — 6-sided lft. fr. chlf. V.d.s. alc.
3647	200	Yellow(?)	p-Hydroxyphenylethylamine Picrate, $\text{HO.C}_6\text{H}_4\text{CH}_2\text{CH}_3\text{NH}_2\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.912. — Short pr.
3648	abt. 200d.	Yellow	Lactosephenylosazone, $\text{C}_{12}\text{H}_{20}\text{O}_4(\text{:NH}_2\text{Ph})_2$. — Ndl. s. in 80-90 pt. boiling aq.; i. eth., chlf., bz.; e.s. gl. ac. ac. — Levorotatory in gl. ac. ac.
3649	200; 202-3	Red	p-Toluidonaphthoquinone(1,4), $\text{Me.C}_6\text{H}_4\text{NH.C}_1\text{O}_2\text{H}_2$. — \textcircled{P} S. conc. H_2SO_4 w. purple color. — Ndl. I. c. dil. NaOH.
3650	201-2	Yel.-brown	5-Nitro-2-aminophenol, $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{OH}$. — \textcircled{P} S. alk. w. or-red color. — Ndl. fr. aq.
3651	201-2	Yellow	Acenaphthylene Picrate, $\text{C}_{12}\text{H}_8\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of h.c. of Vol. I. — Ndl. V.d.s. c. alc.
3652	200-1u.c., d. (r.h.)	Pale yellow	p-Nitrobenzoyl-d,l-serine, $\text{NO}_2\text{C}_6\text{H}_4\text{CO.NH.CH}(\text{CO}_2\text{H}).\text{CH}_2\text{-OH}$. — Thin ndl. fr. aq. S. in 300-400 pt. c., or 20 pt. h. aq.; e.s. h. alc.; alm. i. eth., pet.-eth. Browns abt. 180° u.c. & melts w. efferv.
3653	202	Red	Amino-2-methylantraquinone, $(\text{NH}_2)\text{Me.C}_4\text{H}_4\text{O}_2$. — \textcircled{P} Lust. red ndl. fr. dil. HCl (the nearly colorless hydrochloride being dect. by aq.). — Sbl. Alm. i. aq.; e.s. w. yel. color in alc., eth., bz. — Acetyl deriv., light red ndl. fr. gl. ac. ac., m.p. 176-7°.
3654	202-3	Garnet-red	2-Aminonaphthoquinone(1,4), $\text{NH}_2\text{C}_{10}\text{H}_8\text{O}_2$. — Ndl. fr. alc., eth., gl. ac. ac. — $\text{B}_2\text{H}_6\text{SO}_4$, reddish ndl.; dect. by aq.
3655	202d.; 198-200	Gold-yellow	Isatoxime, Nitrosoxindole, $[\text{C}_6\text{H}_4\text{C}(\text{:NOH}).\text{C}(\text{OH})\text{:N}]$. — \textcircled{P} S. KOH sol. w. dark brown color. (Not dect. by boiling KOH sol.). — V.d.s. aq.; more s. alc.
3656	abt. 202d. (r.h.)	Yellow	Phenanthrenequinonedioxime, $\text{C}_{10}\text{H}_8(\text{:NOH})_2$. — \textcircled{P} S. alk. or conc. H_2SO_4 . — Pr. fr. h. alc. I. aq.; d.s. h. alc.; v.d.s. chlf., bz.
3657	202-3	Yellow	3,5-Dinitro-2-amino-1,4-xylene, $(\text{NO}_2)_2(\text{NH}_2)\text{C}_6\text{H}_4\text{Me}_2$. — Ndl. fr. gl. ac. ac.
3658	202	Yellow	1,3,5,8-Tetranitro-4-naphthylamine, $(\text{NO}_2)_4\text{C}_{10}\text{H}_4\text{NH}_2$. — Ndl.
3659	203; 204	Dark red	$\beta\beta$ -Azonaphthalene, $\text{C}_{10}\text{H}_7\text{N}_2\text{C}_{10}\text{H}_7$. — \textcircled{P} \textcircled{D} Azo comp. — Pr. fr. chlf. Red-yel. lft. fr. alc. Sbl. abt. 210° in red-yel. scales or yel. ndl. D.s. alc.
3660	203-4d.	Or.-red	1-Tryptophane Picrolonate, $\text{C}_{11}\text{H}_{12}\text{O}_2\text{N}_2\text{C}_6\text{H}_4\text{O}_2\text{N}_4$. — \textcircled{P} \textcircled{D} Salts of No. 2.3184 w. No. 2.2605. — Ndl. clusters. 100 pt. aq. dis. 0.384 pt.; e.s. alc.; less s. eth.
3661	203	Brown-red	Dianilobenzoquinoneanil, $\text{C}_8\text{H}_6\text{ON}_2$. — \textcircled{P} S. w. blood-red color in conc. H_2SO_4 . — Ndl. D.s. alc., eth., bz., lgr.
3662	202; 205	Light orange	2,2'-Dinitro-4,4'-diaminodiphenylmethane, $(\text{NO}_2)_2(\text{NH}_2)_2\text{C}_6\text{H}_4\text{-C}_6\text{H}_4(\text{NO}_2)_2(\text{NH}_2)_2$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — Lust. lft. fr. dil. alc. E.s. h. alc.; d.s. eth.

Explanation of typographical signs used in this Division: * = T. 2.39. ‡ = T. 2.21. ¶ = T. 2.36. || = T. 2.34. ¶ = T. 2.35. ** = T. 2.1. ¶¶ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3663	203	Dark brown	Hexamethylanthracene Picrate. — \textcircled{P} \textcircled{D} Picrate* of h.c. of m.p. 220.
3663-I	203-4	Yellow	8-Hydroxyquinoline Picrate, $\text{HO.C}_6\text{H}_4\text{N.Pk.}$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.36. — Pr. v.d.s. c. alc.; alm. i. bz.
3664	200-5d.	YT1	Heroine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.941. — Mic. hexag. plates fr. h. 50% alc. Darkens on exposure to air & light.
3665	203	Pale yellow	Dinitropodocarpic Ac., $(\text{NO}_2)_2(\text{HO})\text{Me.C}_6\text{H}_{14}\text{CO}_2\text{H.}$ — \textcircled{P} Nitro comp.† & acid. — Octahedral cryst. fr. dil. alc. I. aq.; d.s. chlf., bz.; mod. s. alc. — $\text{K}_2\text{A.5H}_2\text{O}$, dark carmine red salt w. metallic green reflections, v.s. aq.! — $\text{AgA.4H}_2\text{O}$, or. flocks.
—	203	Yellowish(?)	1,3,6,8-Tetranitronaphthalene, $\text{C}_{16}\text{H}_4(\text{NO}_2)_4.$
3667	204d.	Light brown	4,4'-Dihydroxyazobenzene, p-Azophenol, $\text{HO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OH.}$ — \textcircled{P} \textcircled{D} Azo comp. — Cryst. w. $\text{H}_2\text{O}.$ D.s. aq.; e.s. alc.; eth., bz. — Conc. HNO_3 yields 2,4-dinitrophenol (No. 2.3126) only. — Diacetyl deriv., yel. lft. fr. alc., m.p. 119°.
3668	204	Yellow	3-Nitro-2-aminobenzoic Ac., $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{CO}_2\text{H.}$ — \textcircled{P} Nitro† acid. — Ndl. fr. aq. V.s. alc., eth.; less s. bz., chlf. — KA , brick-red cryst., i. alc. — $\text{BaA.2H}_2\text{O}$, purple-red ndl., d.s. c. aq.
3669	205	Red or green	Dimethylnaphtheurhodine, $\text{C}_{18}\text{H}_{18}\text{N}_2.$ — \textcircled{P} Sol. in alc., eth. or toluene fluor. yel. Sol. in gl. ac. ac. is violet-red. — Rhomb. tbl. fr. toluene. Red, by transmitted, red & green by reflected light. Sbl. in woolly flocks.
3670	205	"Brown-violet"	Indigooxime, $\text{C}_4\text{H}_{10}\text{O.(:NOH).}$ — Coppery ndl. S. in dil. alk. w. wine-red color; d.s. alc., eth.; i. bz.
3671	205	Or.-yel.	Picryl-m-nitroaniline, $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.C}_6\text{H}_4\text{NO}_2.$ — Cryst. fr. gl. ac. ac. V.d.s. h. alc.; i. eth.
3672	205	Light brown	3,3'-Dihydroxyazophenol, m-Azophenol, $\text{HO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OH.}$ — \textcircled{P} \textcircled{D} Azo comp. — Lft. fr. dil. alc. V.d.s. aq.; e.s. h. alc.; s. Na_2CO_3 sol. w. brown-yel. color. — Boiled w. Zn dust & aq. gives e. oxidizable hydrazo deriv. (Cf. T. 2.21). — Acetyl deriv.** (fr. h. ac. anhydride + little fused NaA), yel. ndl. fr. dil. alc.; e.s. h. alc., m.p. 137°.
3673	205d.	Brown-yellow	Nitropyrogallol, $\text{NO}_2\text{C}_6\text{H}_3(\text{OH}).$ — \textcircled{P} Gives green color w. FeCl_3 & deep red color w. lime-water. — Ndl. fr. h. aq. Loses 1 mol. aq. of crystn. at 100°. V.d.s. c. aq.; e.s. h. aq.
3674	205-6	Gold-yellow	d-Arginine Picrate, $\text{C}_6\text{H}_5\text{O}_2\text{N}_4\text{Pk.2H}_2\text{O.}$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1007. — Long thin silky ndl. Mod. s. h. aq. S. in 200 pt. aq. at 16°.
3675	205d. (r.h.)	Yellow	d-Glucosephenyllosazone, $\text{CH}_2(\text{OH}).(\text{CH.OH}).\text{C}(\text{:N.NHPh}).-\text{CH}=\text{N.NHPh.}$ — [Identical w. osazone fr. d-mannose or d-fructose.] — Ndl. fr. dil. alc. Alm. i. aq.; s. in abt. 200 pt. c. abs.; more s. h. alc. or acetone. Reduces h. Fehling's sol. $[\alpha]_D = -1^\circ, 32'$ (for 0.2 g. in 4 cc. pyridine + 6 cc. abs. alc. in 1 dm. tube). — [M.p. varies widely w. method of heating & purification, & many discordant values are recorded.]
3676	205.5-6	Yellow	Acetyl-p-nitrophenylhydrazine, $(\text{C}_2\text{H}_5\text{O})\text{NH.NH.C}_6\text{H}_4\text{NO}_2.$ — \textcircled{P} Sol. in dil. aq. — NaOH w. deep red color forming Na salt. — Ndl. fr. dil. alc.
3677	206d.	Dark violet-brown	4-Benzeneazonaphthol(1), $\text{Ph.N}_2\text{C}_6\text{H}_4\text{OH.}$ — \textcircled{P} \textcircled{D} Azo comp. — Lft. fr. bz. w. beetle-wing reflections. S. h. alc., bz.; e.s. dil. Na_2CO_3 ; s. conc. H_2SO_4 w. violet-blue color! — Acetate,** brown-red ndl., e.s. h. alc., m.p. 128°.
3678	205-7	Brown	Benzene-p-disazophenol, $\text{HO.C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{OH.}$ — \textcircled{P} \textcircled{D} Azo comp. — Amorph. powd. I. alc., chlf., toluene. — Diacetyl deriv.,** yel. lft. fr. nitrobenzene, m.p. 246-8°.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3679	205-7	Dark or.-yel.	α -Naphthoquinolinequinone, $C_{14}H_8O_2N$. — Ndl. fr. alc. I. aq.; s. alc., eth., bz., dil. mineral acid.
3680	d. abt. 206	Gold-yellow	Nitrobenzoquinone, $NO_2C_6H_3O_2$. — \textcircled{P} S. alk. w. brown-violet color. Colors skin black. — Cryst. fr. h. alc. V.s. h. aq.; d.s. eth., lgr., bz.; e.s. alc., chlf.
3681	abt. 206d. (r.h.)	Yellow	Maltosephenylosazone, $C_{14}H_{10}O_4N_4$. — Ndl. Warts fr. aq. w. 5-8% aq. of crystn. S. in 75 pt. h. aq., or in 150 pt. h. abs. alc., i. eth.
3682	abt. 206	Yellow	N-Benzoylpseudoisatin, $C_8H_4O_2N.CO.Ph$. — \textcircled{P} Shaken w. bz., conc. H_2SO_4 , & little thiophene gives blue indophenine coloration! — Ndl. fr. gl. ac. ac. V.d.s. alc., eth.; d.s. c. gl. ac. ac.
3683	206	Yellow	2-Nitro-4-aminophenol, $(NO_2)(NH_2).C_6H_4.OH$. — Cryst. in colorless ndl. w. aq. crystn. (lost at 170 $^{\circ}$). D.s. h. aq.; e.s. alc., eth., bz. — [Also described as red ndl., m.p. 126-8 $^{\circ}$. Ber., 27, 196.]
3684	206d.	Yellow	Dinitroacenaphthene, $(NO_2)_2.C_10H_6$. — Fine ndl. I. aq.; alm. i. alc.
3685	207; 215	Light yellow	Methylamine Picrate, $MeNH.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1059. — Pr. or tbd. 1.33 pt. dis. in 100 pt. aq.
3686	207	Light yellow	Glyoxalbisdiphenylhydrazone, $Ph_2N.N:CH.CH:N.NPh_2$. — \textcircled{P} S. in conc. H_2SO_4 w. dark violet, or in warm gl. ac. ac. w. emerald-green color! — Ndl. fr. pyridine or much alc.
3687	207-8	Straw- yellow	Leucoauramine G, $Me.NH.C_6H_5Me.CH(NH_2).C_6H_5Me.NHMe$. — Ndl. fr. bz. [Fr. reduction of auramine G by Zn dust & HCl.]
3688	207	Greenish yellow	Nitrohexamethyleucoaniline, $(NO_2)(NMe_2)_2C_6H_4.CH.(C_6H_4-NMe_2)_2$. — Ndl. fr. toluene. [D.R.P., 82,570.]
3689	208w. efferv.	Dark red	4-Tolueneazonaphthol(1), $Me.C_6H_4.N_2.C_6H_4.OH$. — \textcircled{P} \textcircled{D} Azo comp. — D.s. alc., bz.; e.s. acetone; s. c. dil. NaOH! — Acetate,** ndl., m.p. 101-2 $^{\circ}$.
3690	208d.	Cinnabar- red	Rhamnose-p-nitrophenylosazone, $C_9H_{10}O_3:(N_2H.C_6H_4.NO_2)_2$. — \textcircled{P} S. in NaOH sol. w. deep blue color, changing to dark violet on gentle warming! — Ndl. fr. alc.
3691	208	Gold- yellow	5-Nitro-3-aminobenzoic Ac., $(NO_2)(NH_2).C_6H_3.CO_2H$. — \textcircled{P} Acid. — Small pr. fr. aq. E.s. h. alc.; less s. eth., bz. — $Pb\bar{A}_3.3\frac{1}{2}H_2O$, or.-red ndl., d.s. c. aq. — Me \bar{A} , yel. ndl. fr. aq., m.p. 160 $^{\circ}$.
3692	208	Yellow	3,5-Dinitro-o-toluidine, $(NO_2)_2.C_6H_4.Me(NH_2)$. — Cryst. I. aq.; alm. i. h. alc.; s. in abt. 100 pt. h. toluene.
3693	208.5	Yellow	o-Dinitrodicinnamylvinylketone, $CO(C_6H_4.C_6H_4.NO_2)_2$. — Ndl. fr. gl. ac. ac. Alm. i. alc., eth.; e.s. chlf.
3694	abt. 208	Yellow	Purine Picrate, $C_6H_4N_4.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.2499. — Lft. s. in abt. 20 pt. boiling aq.
3695	208-9d.	Y+	Cinchonidine Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.993. — Lust. ndl. fr. h. 10% ac. ac. Darkens fr. 200 $^{\circ}$.
3696	209	Chrome-red	Nitrochrysene, $NO_2.C_{10}H_{11}$. — Thick pr. E.s. h. nitrobenzene; d.s. c. alc., eth. Sbl. undecd.
3697	209; 212	Pale yellow	p-Nitrocarbanilide, $NO_2.C_6H_4.NH.CO.NHPh$. — \textcircled{P} T. 2.12 prob. gives powerful odor of carbylamine. — Ndl. I. aq. — Sapn. (T. 2.26) by KOH gives aniline, NH ₃ , p-nitrophenol, & CO ₂ .
—	209	Yellowish(?)	α -Aminoacridine. — Cf. No. 2.1008.
3698	207; 212-3	Red-brown or gold- yellow	4-Nitrobenzene-azo-4-hydroxybenzene, $NO_2.C_6H_4.N_2.C_6H_4.OH$. — \textcircled{P} \textcircled{D} Azo comp. — Cryst. fr. toluene. Alm. i. aq.; e.s. alc., bz. — Acetyl deriv.,** or. ndl., m.p. 147 $^{\circ}$.

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 1 = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3699	210-1	Yellow-red	Isatinephenylhydrazone, $C_14H_{10}ON.N.NHPh$. — Ndl. fr. alc.
3700	207-8; 212-3	Yellow	Lepidine Picrate, $C_8H_3N.Pk$. — \textcircled{P} \textcircled{D} Picrate of No. 2.1413. — Cryst. fr. alc.
3701	209-10; 211	Yellow	4,6-Dinitro-o-toluidine, $(NO_2)_2.C_6H_4.Me(NH_2)$.
3702	210	Yellow	Methyl-2,4,2',4'-tetranitrodiphenylamine, $MeN.[C_6H_4.(NO_2)_2]_2$. — Lft. fr. alc. or gl. ac. ac.
3703	210	Yellowish	p-Nitrooxanilic Ac., $NO_2.C_6H_4.NH.CO.CO_2H$. — \textcircled{P} Acid. — Ndl. fr. h. aq.; d.s. c. aq.; s. alc. Cryst. w. $1H_2O$.
3704	211	Yellow	Nitrobisphenanthrene, $NO_2.C_{20}H_{16}$. — \textcircled{P} S. in h. conc. H_2SO_4 w. intense green color. — Pr. fr. bz. V.d.s. alc., eth.
3705	211-2d.	Yellow	p-Nitrophenyisemicarbazide, $NO_2.C_6H_4.NH.NH.CO.NH_2$. — \textcircled{P} S. dil. NaOH w. or.-red color. — Small ndl. fr. aq. E.s. h. aq., alc.; d. s. bz., lgr.
3706	212-3	Light red	2,4,4'-Trinitrohydrazobenzene, $(NO_2)_2.C_6H_4.NH.NH.C_6H_4.NO_2$. — \textcircled{P} S. aq. KOH sol. w. blue color! — Ndl. fr. nitrobenzene. S. h. alc.
3707	212-3	Orange	3-Nitrobenzene-4'-azoaniline, $NO_2.C_6H_4.N_2.C_6H_4.NH_2$. — \textcircled{P} \textcircled{D} Azo comp. — Lft. fr. dil. alc. D.s. alc.; e.s. eth., bz. — Acetyl deriv.** or. scales fr. alc., m.p. 166-7°.
3708	d. 212	Brown	Harmalol, $C_{12}H_{12}ON_2.3H_2O$. — [Fr. Russian Peganum harmala.] — \textcircled{P} D.s. aq. w. yel. color & green fluor. which disappears w. ac. or alk. — Pr. fr. alc. w. greenish reflections. Darkens at 180°.
3709	212-3	Bronzy	p-Tolylrosinduline, $C_{12}H_{11}N_2$. — [Pat.] Fine ndl.
3710	212	Golden	4'-Amino-4-acetaminoazobenzene, $NH_2.C_6H_4.N_2.C_6H_4.NH-(C_6H_5O)$. — \textcircled{P} \textcircled{D} Azo comp. colored red by acids. — Lft. fr. dil. alc.
3711	212d.	Golden	o-Dinitrophenyldiacetylene, $NO_2.C_6H_4.C\equiv C:C.C_6H_4.NO_2$. — Ndl. fr. chlf. V.d.s. h. alc. — \textcircled{D} Convert to diisatogen & indigo as described by Baeyer, Ber., 15, 51.
3712	212d.	Yellow	5- α -Naphthaleneazosalicylic Ac., $C_{12}H_9N_2.C_6H_5(OH)(CO_2H)$. — \textcircled{P} Acid & azo comp. — Ndl. fr. alc. E.s. alc. — NaA, cryst. s. in 1420 pt. c., or in 644 pt. boiling aq.
3713	208-15 according to heating rate	Yellow	2,3,4,6-Tetrannitroaniline, $(NO_2)_4.C_6H.NH_2$. — [High explosive, detonating under falling 5 kg. hammer at 35 cm., but burning without explosion on ignited paper. (Vol. IV, p. 31, Cong. Appl. Chem. of 1912).] — Cryst. fr. xylene. I. aq.; s. in 6 pt. acetone; s. xylene, aromat. nitro compounds.
3714	212d.	Yellow	Phenolphthaleineoxime, $C_{10}H_14O_3(:NOH)$. — Cryst. powd. V.d.s. h. alc.; e.s. NaOH, dil. HCl; i. eth., bz., lgr.
3715	210-5d. (r.h.)	Yellow	d,l-Xylosephenylosazone, $CH_3(OH)(CH.OH)_2.C(:N.NHPh)-CH:N.NHPh$. — Ndl. fr. dil alc. Alm. i. eth., h. aq.; s. in abt. 100 pt. boiling alc.
3716	212-3u.c.	Y-YT1	Creatinine Picrate, $C_6H_5ON_2.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1049. — Ndl. V.d.s. c. aq.
3717	212.5; 214.5	Yellowish	(s)-4,6-Dinitroresorcinol, $(NO_2)_2.C_6H_4(OH)_2$. — Glassy pr. fr. ac.-eth. I. aq.; e.s. eth., chlf.; less s. bz., alc. Sbl. — BaA ₂ , yel. ndl., exploded by heat.
3718	212-3	Light yellow	Toluphanthrazine, $C_{12}H_9N_2$. — [Fr. 3,4-toluylendiamine & phenanthrenequinone.] Hair-like cryst. I. aq.; d.s. alc.; e.s. eth., chlf., bz.
3719	213	Orange	Methylenedinitrodiphenyldiamine, $CH_2.(NH.C_6H_4.NO_2)_2$. [NH: NO ₂ = 1:3]. — Ndl. fr. alc. I. aq., eth., chlf., bz. d.s. alc. Unchanged by h. KOH sol.
3720	214	Golden	α -Nitroacridine, $NO_2.C_12H_9N$. — Lft. D.s. c. alc., eth.; s. chlf. Sbl. — Salts yellow.

No.	Melting-point (C°).	Color.	COLORED SOLID COMPOUNDS.
3721	214	Lemon-yellow	6-Nitro-2-amino-p-toluidic Ac. , $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_3\text{Me}(\text{CO}_2\text{H})$. — \textcircled{P} Acid. — Ndl. fr. aq. E.s. alc., eth. Sbl. — $\text{Ba}\ddot{\text{A}}_3 \cdot 4\text{H}_2\text{O}$, gold-yel. lft. v.d.s. c. aq.
3722	214	Yellow	3-Aminonaphthoic(2) Ac. , $\text{NH}_2\text{C}_{10}\text{H}_8\text{CO}_2\text{H}$. — \textcircled{P} Acid & prim. amine. — Brassy lft. fr. dil. alc. V.s. alc., eth., w. vel. color & green fluor. — Acetyl deriv.,** m.p. 238°. — $\text{Na}\ddot{\text{A}}$, lft. d.s. aq.
3723	211d.; 218d.	Yellow	5-Benzeneazo-4-hydroxybenzenecarbonic(1) Ac. , Benzene-5-azo-salicylic Ac. , $\text{Ph.N}_2\text{C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})$. — \textcircled{P} Acid azo comp. — Ndl. fr. bz. 100 pt. aq. sol. at 16° contain 0.03 pt.; e.s. alc., eth.; d.s. c. bz., chlf. — $\text{Ba}\ddot{\text{A}}_3$ (at 150°), gold-yel. ndl., v.d.s. c. aq.
3724	215-6	Brown-red	2-o-Nitrobenzeneazonaphthol(1) , $\text{NO}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_{10}\text{H}_8(\text{OH})$. — \textcircled{P} \textcircled{D} Azo comp. — Ndl. w. greenish reflections fr. fusel oil. D.s. alc., eth., bz. S. conc. H_2SO_4 w. green color, changing to greenish brown.
3725	215d.	Or.-yel.	Aristolochine , $\text{C}_{12}\text{H}_{22}\text{O}_2\text{N}_2$. — [Poison fr. seeds of Aristolochia clematis, or roots of A. rotunda.] — \textcircled{P} S. in conc. H_2SO_4 w. dark green color! — S. h. aq., alc., eth.; i. bz., pet.-eth.; s. alk. & pptd. by CO_2 . Fusion w. KOH purple-red.
3726	abt. 215u.c.	Yellow	α -(syn)- Benzilosazone , $\text{Ph.C:(N.NHPH).C:(N.NHPH)Ph}$. — Cryst. D.s. alc.; e.s. bz. 100 pt. acetone at 18-19° dis. 1.65 pt.
3727	215-6u.c.	Yellow	Trimethylamine Picrate , $\text{Me}_3\text{N.Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1060. — Ndl. s. in abt. 77 pt. c. aq.
3728	215	Yellow	(s)-4,4'-Dinitrodiphenylamine , $(\text{NO}_2\text{C}_6\text{H}_4)_2\text{NH}$. — \textcircled{P} NaOH sol. has violet color. — Ndl. w. blue reflections fr. alc. Alm. i. aq.; d.s. alc., bz.
3729	215	Yellow	Picryl-p-nitroaniline , $(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NH.C}_6\text{H}_4\text{NO}_2$. — Cryst. fr. h. alc.
3730	216	Or.-red	6,6'-Dinitro-o-tolidine , $(\text{NO}_2)(\text{NH}_2)\text{Me.C}_6\text{H}_4\text{C}_6\text{H}_3\text{Me}(\text{NH}_2)_2\text{NO}_2$. — Cryst. fr. cumene. Alm. i. aq., pet.-eth.; d.s. bz.; v.s. alc. — Sulphate d.s. aq. — Diacetyl deriv. (by boiling w. ac. anhydride), white, e.s. alc., m.p. 275°.
3731	215-7	Brown-red	o-Nitrobenzeneazosalicylic Ac. , $\text{NO}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_3(\text{OH})(\text{CO}_2\text{H})$. — \textcircled{P} \textcircled{D} Acid & azo comp. — Cryst. powd. V.d.s. aq.; e.s. h. alc. — Alk. salts s. aq. w. dark red color.
3732	216	Straw or amber-yellow	$\text{C}\alpha\alpha'\text{-N}\beta\beta'\text{-Naphthacridine}$, $\text{C}_{14}\text{H}_{12}\text{N}$. — [Fr. β -naphthylamine & formic aldehyde.] — \textcircled{P} S. conc. H_2SO_4 w. green fluor. ! — Ndl. s. alc.; d.s. h. bz. Becomes darker in light.
3733	216	Yellow-brown	p-Nitroethylenediphenyldiamine , $(\text{NO}_2\text{C}_6\text{H}_4\text{NH})_2\text{C}_2\text{H}_4$. — Ndl. fr. nitrobenzene. I. alc., eth., bz.
3734	217	Orange	8-Aminonaphthophenazine , $\text{C}_{16}\text{H}_{11}\text{N}_3$. — \textcircled{P} S. conc. H_2SO_4 w. blood-red color, changing to pale yel. on diln. w. aq. — Lft. fr. alc. I. aq.; s. alc., bz., eth. w. yel. color.
3735	216-8	Light yellow	p-Nitrodicinnamylvinylketone , $\text{Ph.C}_6\text{H}_4\text{CO.C}_6\text{H}_4\text{C}_6\text{H}_4\text{NO}_2$. — Ndl. fr. ac. anhydride. I. alc., eth., chlf.; e.s. gl. ac. ac.
3736	217	Yellowish	Acetyl-α-dinaphthylamine , $(\text{C}_6\text{H}_5\text{O})\text{N}(\text{C}_{10}\text{H}_7)_2$. — Ndl. in stellate groups fr. alc.
3737	218	Gold-yellow	N-Methylpyrrolidine Picrate , $\text{C}_6\text{H}_{11}\text{N.Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1085. — Lft. fr. alc.
3738	218	Deep yellow	(γ)-5,8-Dinitro-α-naphthoic Ac. , $(\text{NO}_2)_2\text{C}_{10}\text{H}_8\text{CO}_2\text{H}$. — \textcircled{P} Acid. — Trimetric cryst. fr. alc. Swells up in melting. S. h. aq.; v.s. alc. — $\text{Ca}\ddot{\text{A}}_3 \cdot 7\text{H}_2\text{O}$, broad lust. ndl., v.s. aq. — $\text{Et}\ddot{\text{A}}$, long yel. ndl. fr. alc., m.p. 129°. (Fr. $\text{Ag}\ddot{\text{A}}$ & EtI .)

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¶ = T. 2.35. ** = T. 2.1. ¶ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3739	218-21	Red	Dinitrobenzidine, $(NO_2)(NH_2)C_6H_3C_6H_3(NH_2)(NO_2)$. — \textcircled{P} Pptd. fr. sol. in conc. ac. by aq. — Cryst. aggregates fr. alc.
3740	219-20	Sulphur-yellow	Pyruvic acid-p-nitrophenylhydrazone, $(NO_2)C_6H_4NH.N:CM(CO_2H)$. — \textcircled{P} Hydrazone.††
—	219-20	Pale yellow	3-Nitrophthalic Ac. — Cf. No. 2.389.
3741	220	Reddish	4-Hydroxyazobenzenecarbonic(3') Ac., $HO.C_6H_4N_2C_6H_4CO_2H$. — \textcircled{P} Acid & azo comp. — Cryst. fr. dil. alc. V.d.s. h. aq.; e.s. alc., eth. Taste bitter.
3742	216-20; 221-2	Or.-red	4,4'-Dinitroazobenzene, $NO_2C_6H_3N_2C_6H_3NO_2$. — \textcircled{P} \textcircled{O} Azo comp. reduced by Sn + HCl to p-phenylenediamine (No. 2.877). — Lust. ndl. fr. acetone. S. in 5.2 pt. boiling bz.; d.s. alc., eth.
3743	220d.	Yel.-red	5-Nitro-2-amino-p-toluic Ac., $(NO_2)(NH_2)C_6H_3Me(CO_2H)$. — \textcircled{P} Acid & nitro comp.‡ — Glassy ndl. fr. aq. D.s. c. aq.; e.s. alc., eth.
3744	220; 218.5c.; 222-3c.	Yellow	2-Nitrodiphenyleneketone, $C_6H_4CO.C_6H_3(NO_2)^2$. — \textcircled{P} Nitro comp.‡ — Cryst. fr. alc. Sbl. D.s. c. alc. — Reduction by $(NH_4)_2S$ gives amino deriv., violet-red cryst. fr. alc., m.p. 163° c.
3745	220	Yellow	β -Dinitrofluorenone, $(NO_2)_2C_6H_3CO$. — \textcircled{P} Nitro comp. — Ndl.
3746	abt. 220	Dirty yellow	Picryl-o-nitroaniline, $(NO_2)_2C_6H_3NH.C_6H_4NO_2$. — Ndl. Alm. i. alc.
3747	221	Lemon-yellow	p-Nitrobenzalsemicarbazone, $NO_2C_6H_4CH:N.NH.CO.NH_2$. — \textcircled{P} Semicarbazone.†† — Ndl. fr. aq. Cryst. w. 1H ₂ O.
3748	221-2d.	Straw-yellow	Santoninphenylhydrazone, $C_{10}H_{15}O_3N_2H.Ph$. — \textcircled{P} Hydrzone.†† — Lust. ndl. fr. alc.
3749	220-1; 222- 3.5	Yellow	Isoquinoline Picrate. — \textcircled{P} \textcircled{O} Picrate* of No. 2.1365. — Ndl. D.s. aq., alc.
3750	221d.	Yellow	Pentamethylenediamine Picrate, $NH_2.(CH_2)_5NH_2.2Pk$. — \textcircled{P} \textcircled{O} Picrate* of No. 2.1232. — Thin ndl. or tbl. Alm. i. aq. Melts w. blackening & efferv.
3751	abt. 221d.	V. pale greenish yellow	α -Aminobenzaldehydophenylhydrazone, $NH_2C_6H_4CH:N.NHPH$. — Lft. fr. alc. turning brown in air. D.s. alc., eth.; v.s. acetone.
3752	222	Red	Pyrene Picrate, $C_{14}H_{10}Pk$. — \textcircled{P} \textcircled{O} Picrate* of h.c. of Vol. I. — Long ndl. fr. h. alc. V.s. bz. Stable in h. alc.
3753	222	Vermilion	2,4-Dinitrodiphenylamine, $(NO_2)_2C_6H_4.NH$. — \textcircled{P} Sol. in gl. ac. ac. if decolorized by Zn dust becomes green & then yel. w. NaNO ₂ . — D.s. warts fr. xylene.
3754	d. 222	Light yellow	Dimethylamine Picrolonate, $Me.NH.C_6H_3O_2N_4$. — \textcircled{P} \textcircled{O} Salt of No. 2.1061 w. No. 2.3184. — Fine ndl. s. in 764 pt. c., or 33 pt. boiling aq.; or in 853 pt. c., or 38 pt. h. alc.
3755	223	Red-yel.	1,8-Dinitro-2-naphthylamine, $(NO_2)_2C_6H_3NH_2$. — \textcircled{P} Nitro comp.‡ & prim. amine.¶ — 6-sided tbl. V.d.s. alc., eth., c. chlf.; e.s. acetone.
3756	223-4	Yellow	3,4'-Dinitrodiazoaminobenzene, $NO_2C_6H_4N_2.NH.C_6H_4NO_2$. — Ndl. fr. alc. — Decd. by HCl in tube at 100° to m- & p-nitraniline, m- & p-chlorbenzene, & N.
3757	220; 225	Yellow	3,5-Tetranitrobiphenol(4), $(NO_2)_2(OH)C_6H_2C_6H_2(OH)(NO_2)_2$. — \textcircled{P} Nitro comp.‡ & phenol. — Ndl. — Diacetyl deriv.** yel. ndl., m.p. 236°.
3758	223.5u.c.; (230c.)	Yellow	1-Nitroanthraquinone, $NO_2C_6H_3:(CO)_2:C_6H_4$. — Sbl. in lft. Cryst. fr. gl. ac. ac. D.s. alc., eth., gl. ac. ac.; more s. chlf., bz. — Fusion w. KOH gives alizarine (Vol. I).

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3759	223	Brassy	a-Dimethyldiaminoquinoxazone, $C_4H_{12}N_2O_2$. — (P) D.s. h. aq. w. blue color! Sol. in alc., red-violet. — Pr. fr. alc. Sbl. I. alk.
3760	223	Blue	Indoxin, $C_{18}H_{12}O_4N_2(?)$. — (P) S. alk. w. emerald-green color! — Mic. ndl. w. coppery reflections fr. chlf. + lgr. V.s. alc., eth., bz.
3761	224.5	Red	3-Nitro-4-aminocinnamic Ac., $(NO_2)(NH_2)C_6H_4CH:CH.CO_2H$. — (P) Acid & nitro comp. — Ndl. fr. h. aq. E.s. h. alc.; less s. aq.; alm. i. bz., lgr.
3762	224d.	Dark ruby-red	α -Nitroanthragalloi, α -Nitro-1,2,3-trihydroxyanthraquinone, $NO_2.C_18H_8O_4$. — (P) Sol. in alk. green; in conc. H_2SO_4 red to brown. — Pr. w. $1C_6H_6$ fr. bz. + alc. E.s. alc.; alm. i. aq., pet.-eth.; s. bz., eth., chlf. — Oxidn. by HNO_3 (T. 1.905-3) gives phthalic ac. (Vol. I).
3763	224-5	Yellow-red	Dinitro- β -naphthylamine, $(NO_2)_2C_{10}H_8NH$. — Ndl. fr. h. cumene. V.d.s. alc., eth.
3764	216(sl.h.), 232(r.h.)	Red-brown	2,4-Dinitrobenzaldehydephenylhydrazone, $(NO_2)_2C_6H_4CH:N-NHPh$. — (P) Alc. sol. w. a drop or two sodium alcoholate sol. becomes blue. — Ndl. fr. acetone. I. aq.; d.s. alc., eth., bz.
3765	224	Yellow	uns.-Dimethylguanidine Picrate, $C_6H_8N_2Pk$. — Small ndl. fr. aq.
3766	224(r.h.)	Light yellow	Anthraquinoneoxime, $C_{18}H_8O(:NOH)$. — (P) S. alk. w. red-brown color. — Ndl. fr. dil. alc. E.s. alc. Sbl.
3767	225	Brown-red	4,6-Dinitro-3-aminophenol, $(NO_2)_2(NH_2)C_6H_3OH$. — (P) Boiling w. aq. — KOH sol. gives NH_3 (T. 2.7) & dinitroresorcinol. — Cryst. w. metallic lustre fr. chlf. V.d.s. aq., chlf.; more s. alc., eth.
3768	223; 225	Red-brown	Salicycaldehyde-p-nitrophenylhydrazone, $HO.C_6H_4CH:N-NH.C_6H_4.NO_2$. — (P) S. alk. w. deep red color. Hydrazone. — Pr. w. bluish reflections fr. alc. V.d.s. aq.; e.s. alc.
3769	225d.	Yellow	p-Nitrophenylglycine, $NO_2.C_6H_4.NH.CH_2CO_2H$. — [D.R.P., 88,433.] — (P) Acid. — Cryst. fr. aq. Sinters at 210°. Melts w. brisk efferv.
3770	225	Yellow	α -Naphthylauramine, $C_{10}H_8N:C(C_6H_4.NMe_2)_2$. — [D.R.P., 44,077.] — Cryst. powd. D.s. alc.
3771	225u.c., d.	Sulphur-yellow	d-Argentine Picrolonate, $C_{10}H_8O_4N_2.C_6H_4O_4N_2$. — (P) Salt of No. 2.1007 & No. 2.3184. — Small ndl. fr. h. aq. s. in 1124 pt. c. aq., or 2885 pt. c. alc.
3772	226	Greenish brown	Diformazyl, $C_8H_2N_8$. — (P) S. conc. H_2SO_4 w. indigo-blue color! — Lft. w. metallic reflections. D.s. c. alc.; v.s. chlf., bz.; i. lgr.
3773	227; (235)	Dark red	Aminotoluquinonedi-p-tolylimide, $(NH_2)Me.C_6H_4(:N.C_6H_4.Me)_2$. — [Fr. oxidation of p-toluidine.] — (P) S. conc. H_2SO_4 w. greenish-blue color, changing to Bordeaux red on heating. — Plates fr. xylene. E.s. gl. ac. ac.; d.s. alc.
3774	227	Or.-yel.	Acetaminodisazobenzene, $(C_6H_5O)NH.C_6H_4.N_2.C_6H_4.N_2Ph$. — (P) Colored deep green by conc. H_2SO_4 ! — Lft. fr. alc.
3775	227-8	Yellow	4,4'-Diaminostilbene, $NH_2.C_6H_4CH:CH.C_6H_4NH_2$. — (P) Prim. amine! giving transient blue-green color w. $FeCl_3$ in alc. sol. — Lust. cryst. fr. alc. Sbl. in lft. D.s. bz., h. aq.; mod. s. wood spirits. — B_2HCl , lust. yel. or brown fluorescing tbd. fr. dil. HCl; d. abt. 243°.
3776	228-9	Violet-red	2,4'-Dihydroxy-1,1'-azonaphthalene, $HO.C_{10}H_8N_2.C_{10}H_8OH$. — (P) Azo comp. — I. aq., dil. ac. or alk.; v.d.s. alc.; s. conc. H_2SO_4 w. violet color!

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34.
 ¶ = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3778	228	Yellow	Benzylamidine Picrate, Ph.C(:NH)(NH ₂). — ⊕ ⊖ Picrate* of No. 2.679. — Ndl.
3779	228	Pale yellow	C ₂₈ -N ₂ -Dinaphthacridine, C ₂₁ H ₁₂ N. — ⊕ V.d.s. h. alc. w. yel. color & intense blue fluor.! — Cryst. — B.HNO ₃ , or.-yel. ndl. s. h. alc. w. yel. color & green fluor.
3780	228-30; 224	Red	3,3'-Dinitro-4,4'-diaminodiphenylmethane, [(NO ₂)(NH ₂).C ₆ H ₄ .]—CH ₃ . — Ndl. fr. gl. ac. ac. I. alc., bz., chlf.
3781	229-30	Red-brown	4-Nitro-4'-azodimethylaniline, NO ₂ .C ₆ H ₄ .N ₂ .C ₆ H ₄ .NMe ₂ . — ⊕ ⊖ Azo comp. — Mic. ndl. fr. alc. D.s. h. alc., bz., eth. — B.HCl, lust. steel-blue ndl. — B ₂ H ₆ PtCl ₆ , gran. red ppt.
3782	230-1	Red	Phenylinduline, Phenylanilinoaposafranine, C ₂₀ H ₂₂ N ₄ . — ⊕ Sol. in conc. H ₂ SO ₄ blue! — Translucent tbl. fr. bz. + MeOH. — B.HCl, gran. fr. alc. w. green metallic lustre. I. aq.; s. w. fuchsine-red color in alc.!
3783	230	Red	cis-2,2'-Diaminostilbene, NH ₂ .C ₆ H ₄ .CH:CH.C ₆ H ₄ .NH ₂ . — ⊕ Prim. amine. — Ndl. fr. aq.
3784	230	Red	Dinitrochrysouquinone, (NO ₂) ₂ .C ₁₀ H ₈ O ₂ . — Ndl. S. h. alc.; less s. eth., bz.
3785	230-1	Brick-red	2-Azooquinoline, NC ₆ H ₄ .N ₂ .C ₆ H ₄ N. — Lft. Sbl. S. h. alc. w. deep red color. Salts are ptd. by x.s. acid fr. sol. in mineral ac. & decd. by much aq.
3786	darkens at 230	YO-O	Brucine Picrate. — ⊕ ⊖ Picrate* of No. 2.956. — Mic. octahedra fr. h. dil. ac. ac. Is entirely decd. abt. 275°.
3787	229.5-30.5	Or.-yel.	Diacetyl-p-nitrophenylhydrazone, Me.CO.C(:N.NH.C ₆ H ₄ .NO ₂).-Me. — ⊕ S. NaOH sol. w. deep violet-red, or in h. alc. w. or.-red color. — Ndl. w. bluish reflections. R.d.s. alc., bz.
3788	230	Brass-yellow	Dipicrylethylenediamine, [(NO ₂) ₂ .C ₆ H ₄ .NH] ₂ .C ₆ H ₄ . — Lft. fr. nitrobenzene. I. alc., eth., bz.
3788-I	230	Yellow	4-Nitro-3-hydroxybenzoic Ac., (NO ₂)(HO).C ₆ H ₄ .CO ₂ H. — ⊕ Acid & nitro comp.† — Long lft. V.d.s. h. aq. — Ba ₂ H ₂ O, yel.-red lft., or red ndl., alm. i. aq.
3789	230	Straw-yellow	4,5-Dinitronaphthol(1), (NO ₂) ₂ .C ₁₀ H ₈ .OH. — ⊕ Nitro comp.‡ & phenol. — Ndl. D.s. aq.; s. organic solvents.
3790	230	Light yellow	α-Diaminomethylphenylacridine, (NH ₂) ₂ .C ₂₀ H ₁₂ N. — ⊕ S. alc., eth., bz. w. yel. color & green fluor. — Cryst. fr. eth. I. aq. — [Pat.]
3791	d. 230	Light yellow	d,L-Lysine (α-Diaminocaproic Ac.) Picrate, [NH ₂ .CH ₂ .(CH ₂) ₃ .CH(NH ₂).CO ₂ H].Pk. — ⊕ Picrate* of syrupy ac. of alk. react. — Mic. ndl. fr. aq. S. h. aq.; v.d.s. abs. alc.; i. eth., bz.
3792	228; 232	Yellowish	5-Aminonaphthoic(2) Ac., NH ₂ .C ₁₀ H ₈ .CO ₂ H. — [D.R.P., 92,995.] — ⊕ Prim. & prim. amine. — Lft. alm. i. aq.; i. eth., bz.; mod. s. h. alc.
3793	230	Green-yellow	Methylphenyl-m-nitrophenylurea, PhMeN.CO.NH(C ₆ H ₄ .NO ₂). — ⊕ Nitro comp.† — Ndl. E.s. h. alc. — Sappn. (T. 2.26) should give: methylaniline (No. 2.1249); m-nitroaniline (No. 2.3125); CO ₂ .
3794	232 u.c.	YO-O	Picryl-β-naphthylamine, (NO ₂) ₂ .C ₆ H ₄ .NH.C ₁₀ H ₇ . — D.s. pr. fr. gl. ac. ac.
3795	232-3	Brown-yellow	Dianilinotoluquinone(2,5), (Ph.NH).Me.C ₆ HO ₂ . — ⊕ S. w. blood-red color in conc. H ₂ SO ₄ . — Ndl. fr. alc. Alm. i. c. alc.; more s. h. gl. ac. ac.
3796	232	Lemon-yellow	Methylenedinitrodiphenyldiamine, CH ₂ (NH.C ₆ H ₄ .NO ₂) ₂ ; [NH:-NO ₂ = 1:4]. — Ndl. I. aq., eth.; v.d.s. h. alc.
3797	233d.(r.h.)	Yellow	4,4'-Dinitrodiazoaminobenzene, NO ₂ .C ₆ H ₄ .N.NH.C ₆ H ₄ .NO ₂ . — ⊕ S. h. dil. KOH w. violet color! — V.d.s. h. alc., eth.; d.s. chlf., bz. — Charac. Na salt forms blue ndl. alm. i. aq.,

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3798	233d.	Yellow(?)	or c. aq. containing NaOH; dec. in air. — C. conc. HCl splits to p-nitroaniline (No. 2.2945) & p-nitrodiazobenzene chloride. m.s.-Methylphenanthridine Picrate, $C_{11}H_{11}N.Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1052.
3799	233; (170)	Yellowish	o-Nitrocarbanilide, $NO_2.C_6H_4.NH.CO.NHPh.$ — \textcircled{P} Prob. gives strong carbarylamine odor in T. 2.12. — Ndl. I. c. aq.; alm. i. c. alc.; e.s. h. alc.
3800	234; 235; 229-30	Garnet-red	Phenylrosinduline, $C_{20}H_{16}N_2.$ — \textcircled{P} Sol. w. pure green color in conc. H_2SO_4 of less than 95.2%, but w. brown color in stronger acid. — Cryst. Alm. i. aq.; r.d.s. alc.; more s. bz. — Neutral salts, red; acid salts, green. — $B.HCl.1\frac{1}{2}H_2O$, dyes wool like fuchsine.
3801	234-5	Brick-red	p-Nitrobenzeneazo-(2)-naphthol(1), $NO_2.C_6H_4.N_2.C_10H_8.OH.$ — \textcircled{P} \textcircled{D} Azo comp. — Ndl. w. greenish reflections. D.s. h. alc., eth., chlf., bz.; e.s. h. xylene. — Acetyl deriv.,** brick-red ndl. fr. gl. ac. ac., m.p. 179.5°.
3802	234-5	Brown-yellow	4'-Nitro-4-acetaminoazobenzene, $NO_2.C_6H_4.N_2.C_6H_4.NH-(C_6H_5O).$ — \textcircled{P} \textcircled{D} Azo comp. — Ndl. fr. alc.; d.s. aq., alc. — [Pat.]
3803	234-5	Gold-yellow	Quinophthalon, Quinoline Yellow, $[O.CO.C_6H_4.C(CH_2C_6H_5N)]^2.$ — \textcircled{P} Dyes wool bright yel. without mordant in dil. sol.! — Ndl. fr. alc. Sbl. undecd. I. aq.; v.d.s. eth.; d.s. h. alc.; s. h. ac. ac.; s. without deen. in conc. $H_2SO_4.$ — Decd. by fuming HCl at 240° to quinaldine (No. 2.1376) & phthalic ac. (Vol. I).
—	234-5u.c.	GYT3	p-Nitrobenzoic Ac. — Cf. No. 2.425.
3804	233-5d.	Yellow(?)	Ethylenediamine Picrate, $C_6H_4N_2.2Pk.$ — \textcircled{P} \textcircled{D} Picrate* of No. 2.1130. — Lft. D.s. aq.
3805	236	Red	α -Diaminoanthraquinone, Anthracene Orange, $(NH_2)_2.C_14H_8O_2.$ — Cryst. fr. eth. Sbl. in garnet ndl. w. greenish reflections. V.d.s. aq.; mod. s. alc., eth., chlf., bz. — Treatment w. alc. & nitrous ac. gives anthraquinone (Vol. I).
3806	237d.	Red	Azobenzene-2,3'-dicarboxic Ac., $CO_2H.C_6H_4.N_2.C_6H_4.CO_2H.$ — \textcircled{P} \textcircled{D} Azo comp. & ac. — Ndl. fr. gl. ac. ac. R.d.s. alc., eth.; alm. i. lgr., bz.
3807	237-8	Brown-red	p-Azobenzaldehyde, $CHO.C_6H_4.N_2.C_6H_4.CHO.$ — \textcircled{P} Azo comp. — Lft. Sbl. fr. 220°. Alm. i. except in nitrobenzene or h. amyl alc. — \textcircled{D} Yields a charac. bisphenylhydrazone, dark red ndl. fr. nitrobenzene, m.p. 278.5° d.; s. w. intense blue color in conc. $H_2SO_4.$
3808	237.5u.c.	Yellow; brown-yel.	2'-Methyl-3'-amino-1,2-naphthacridine, $Me(NH_2).C_{17}H_{14}N.$ — \textcircled{P} E.s. alc. w. or.-yel. color & yel.-green fluor.! — Salts w. mineral ac. are red, dyeing tannin-mordanted cotton or.-yel. — Cryst. fr. xylene. V.d.s. eth., toluene.
3809	237d.	Dark yellow	o-Azobenzoic Ac., $CO_2H.C_6H_4.N_2.C_6H_4.CO_2H.$ — \textcircled{P} \textcircled{D} Acid & azo comp. — Alm. i. h. aq.; i. bz.; s. alc. — $Ag_2\bar{A}$, amorph. red-yel. ppt.
3810	237	Lemon-yellow	2,4-Dinitro- α -naphthylamine, $(NO_2)_2.C_{10}H_8.NH_2.$ — \textcircled{P} \textcircled{D} Treatment w. aq.-KOH sol. dec. (even in cold) giving NH ₂ & dinitro- α -naphthol (No. 2.3266). — Ndl. fr. alc. D. s.h. aq.
3811	237	Yellow	7-Methylquinoline Picrate. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1402. — Pr. V.d.s. h. alc., bz.
3812	237sl.d.	Yellow	β -8-Hydroxyquinolinecarboxic Ac., $HO.C_6H_5N.CO_2H.$ — \textcircled{P} FeCl ₃ colors deep yel. aq. sol. violet-red to deep brown. — Long pr. w. 1 mol. aq. crystn. (lost at 100°). D.s. c. aq., alc., bz. — \textcircled{D} Dec. smoothly to CO ₂ & 8-(o)-hydroxyquinoline (No. 2.36) at 237-50°.

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 ¶ = T. 2.35. ** = T. 2.1. # = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3813	d. abt. 238	Brown-yellow	Benzoquinoneoximesemicarbazone, $\text{HO.N:C}_6\text{H}_4:\text{N.NH.CO.-NH}_2$. — Ndl. fr. gl. ac. ac. I. aq., alc., eth.
3814	238d.	Light yellow	p-Dipicrylamine, Hexanitrodiphenylamine, $(\text{NO}_2)_3\text{C}_6\text{H}_2\text{NH-C}_6\text{H}_3(\text{NO}_2)_3$. — [An explosive; an intermediate in mfg. of dyestuff "Aurantia" & sulphur colors.] — Strongly acid! Pr. fr. gl. ac. ac. Alm. i. aq., eth. — NH ₄ salt $(\text{NO}_2)_3\text{C}_6\text{H}_4\text{NH-NH}_3$ ("Aurantia") is dyestuff described as No. 3.897.
3815	238	Yellow	Anthramine, $\text{C}_{14}\text{H}_8\text{NH}_2$. — [Reduc. prod. of aminoanthraquinone.] — ② Alc. sol. fluor. green! Nitrous fumes passed into alc. sol. give charac. cryst. red ppt.! Fusion w. arsenic ac. gives deep blue mass! — Alm. i. aq., alk.; d.s. in most solvents. — B.HCl, lft., d.s. dil. HCl. — Acetyl deriv.,** silvery lft., s. alc. w. blue fluor., m.p. 240°.
3816	238	Yellow	d,L-Phenylalanine Picrolonate, $\text{C}_9\text{H}_{11}\text{O}_2\text{N.C}_{10}\text{H}_8\text{O}_4\text{N}_4$. — ② ① Salt of No. 2.478-1 & No. 2.3184. — Lft. or pr. browning abt. 220°. 100 pt. c. aq. dis. 0.19 pt.; 100 pt. c. alc. dis. 0.309 pt., alm. i. eth.
3817	238	Yellowish(?)	1,6-Dinitro-2-naphthylamine, $(\text{NO}_2)_2\text{C}_10\text{H}_4\text{NH}_2$. — ② ① Nitro comp.† & prim. amine. — V.d.s. aq., alc., bz. — May be changed to 1,6-dinitronaphthalene (No. 2.3413), through diazo deriv. (Ber., 17, 1172).
—	238d.	Yellow in thick layers	Harmaline. — Cf. No. 2.1032.
3818	239d.	Yellow	Diacetylphenylosazone, $\text{Me.C:(N.NHPh).C:(N.NHPh)Me}$. — ② Gives Bülow's reac. (T. 2.11). — Mic. plates fr. bz. Alm. i. aq., alc.; v.d.s. eth., chlf. Sol. in conc. H ₂ SO ₄ , brown, soon becoming dirty wine-red, or green in thin layers.
3819	238-40	Yellow	Phenanthroline Picrate, $\text{C}_{12}\text{H}_8\text{N}_3\text{Pk}$. — ② ① Picrate* of No. 2.684. — Mic. pr. V.d.s. h. alc.
3820	240	Light brownish red	α-Nitro-o-aminocinnamic Ac., $(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{CH:CH.CO}_2\text{H}$. — ② Nitro comp.† & acid. — Ndl. D.s. aq.; i. bz., eth., lgr.; e.s. alc.
3821	d. abt. 240	Yellowish	p-Benzoquinonedioxime, $\text{HO.N:C}_6\text{H}_4:\text{N.OH}$. — Long fine ndl. becoming white in drying; also short colorless ndl. E.s. conc. ammon. — C. fuming HNO ₃ , gives p-dinitrobenzene (No. 2.2319). — Boiled 10 min. w. acetic anhydride + NaÅ gives antidiacetyl deriv., lft. fr. toluene, m.p. 190° d.
3822	240d.	Yellow	Trinitro-α-naphthylamine, $(\text{NO}_2)_3\text{C}_6\text{H}_4\text{NH}_2$. — Lft. fr. toluene. Alm. i. alc., eth., bz.; v.d.s. toluene.
3823	d.w.m. abt. 240	Yellow	p-Azoxybenzoic Ac., $\text{CO}_2\text{H.C}_6\text{H}_4\text{N}_2\text{O.C}_6\text{H}_4\text{CO}_2\text{H}$. — ② ① Azoxy acid. — Amorph. powd. I. alc.; s. pyridine. — BaÅ (at 120°), dark yel. tbl., i. aq.
3824	241; 242-3	Ruby-red	1-Aminoanthraquinone, $\text{NH}_2\text{C}_6\text{H}_4:(\text{CO})_2\text{C}_6\text{H}_4$. — Iridescent ndl. fr. dil. HCl or dil. alc. Sbl. in ndl. I. aq.; e.s. alc., eth., chlf. bz. Separates in nearly colorless ndl. fr. sol. in h. conc. HCl as hydrochloride, which is decd. by aq. — Acetyl deriv.** (by heating w. ac. anhydride & NaAc.), or-red, e.s. alc., m.p. 202°; 215° c.
3825	236-7; 246	Garnet-red	Azophenine, $(\text{Ph.NH})_2\text{C}_6\text{H}_4:(\text{NPh})_2$. — ② S. conc. H ₂ SO ₄ w. violet color, suddenly changing to blue at 300°! — Monoclin. ndl. fr. aniline. I. alc., eth., alk.; s. chlf.
3826	241	Gold-yellow	4,4'-Diaminoazobenzene, $\text{NH}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4\text{NH}_2$. — [Intermediate in dyestuff mfg.] — ② ① Azo comp. — Ndl. fr. dil. alc. D.s. aq., bz., lgr.; e.s. alc. — Monacid salts are green, diacid salts, red.
3827	239; 241-2; 243	Yellow; brown; steel-blue	p-Tetramethyldiaminoazoxybenzene, $\text{Me}_2\text{N.C}_6\text{H}_4\text{N}_2\text{O.C}_6\text{H}_4\text{NMe}_2$. — ② Azoxy comp. — Cryst. fr. alc. D.s. aq.; more s. h. alc., bz. Salts decd. by aq. — Reduction by Sn & HCl gives dimethyl-p-phenylenediamine (No. 2.560).

No.	Melting-point (C. ^o).	Color.	COLORED SOLID COMPOUNDS.
3828	242-3	Yellow	s-($\alpha\beta$)-Dinaphthazine, $C_{20}H_{12}N_2$. — Sol. in alc. or bz. fluor. blue; that in gl. ac. ac., green. — Long ndl. fr. h. alc.
3829	240; 245	Yellow	7-Aminonaphthoic(2) Ac., $NH_2C_{10}H_8CO_2H$. — \textcircled{P} Acid & prim. amine. \textcircled{I} — Lft. E.s. h. ammon.
3830	244-5d.	Dark red	o-Nitrobenzene-4-azonaphthol(1), $NO_2C_6H_4N_2C_10H_8OH$. — \textcircled{P} \textcircled{I} Azo comp. \textcircled{II} s. conc. H_2SO_4 w. violet-blue color! — Bronzy ndl. fr. h. xylene. D.s. c. alc., eth., chlf.
3831	244d.	Or.-yel.	3-(β)-Nitroalizarine, "Alizarine Orange," $(NO_2)(HO)_2C_6H_3(CO)_2C_6H_4$. — [Cf. No. 3.1122 for description as dyestuff.] — Long ndl. fr. bz. Sbl. w. sl. decn. in yel. lft. D.s. aq.; s. chlf.; s. NaOH w. purple-red color! — Diacetate, ** yel. ndl. fr. bz., m.p. 218°.
3832	244	Yellow-brown	3-Amino-2,7-dimethylacridine, $(NH_2)Me_2C_{12}H_8N$. — [D.R.P., 107,626.] — \textcircled{P} Dyes tannin-mordanted cotton clear or.-yel. Dis. in alc. or eth. w. yel. color & green fluor.! — Cryst. fr. boiling toluene. Acetyl deriv., ** yel. cryst. powd., m.p. 258°.
3833	d. 244	Pale yellow	Methylamine Picrolonate, $MeNH_2C_{10}H_8O_2N_4$. — \textcircled{P} \textcircled{I} Salt of No. 2.1059 w. No. 2.3184. — Ndl. s. in 1073 pt. c., or 369 pt. boiling aq.; s. in 4717 pt. c., or 133 pt. boiling alc.
3833-I	244d.	Pale yellow	Ethylamine Picrolonate, $EtNH_2C_{10}H_8O_2N_4$. — \textcircled{P} \textcircled{I} Salt of No. 2.1062 w. No. 2.3184. Tbl. S. in 93 pt. boiling, or 3846 pt. c. aq. Browns abt. 220°.
3834	245	Red	Phenanthrenequinone-p-nitrophenylhydrazone, $[C:(N.NH-C_6H_4NO_2)C_6H_4C_6H_3CO]^+$. — Ndl. fr. xylene. V.d.s. alc., eth., lgr. S. alc. KOH w. violet color!
3835	246	Light yellow	m-Nitrobenzaldehydesemicarbazone, $NO_2C_6H_4CH:N.NH.CO-NH_2$. — \textcircled{P} Semicarbazone. \textcircled{II} — Ndl. fr. alc.
3835	247	Nearly black	α -Naphthylrosinduline, $C_{12}H_9N_3$. — [Pat.] — \textcircled{P} Ndl. w. metallic lustre fr. toluene which dis. w. grass-green color in conc. H_2SO_4 ! — Conc. HCl at 210° gives α -naphthylamine (No. 2.589), & rosindone.
3837	247	Or.-red	m-Nitrobenzaldehyde-p-nitrophenylhydrazone, $NO_2C_6H_4CH:N.NH.C_6H_4NO_2$. — \textcircled{P} Hydrazone, s. in NaOH sol. w. pale violet color, changing to deep violet-red on dilution w. alc. — Lft. D.s. h. alc., bz., lgr.; e.s. acetone.
3838	247	Yellowish	2,4-Dinitro-1-acetaminonaphthalene, $(NO_2)_2C_{10}H_8NH(C_2H_5O)$. — \textcircled{P} \textcircled{I} Boiling w. NaOH sol. (T. 2.26) gives NH ₃ ac. ac., & No. 2.3266.
3839	abt. 247d.	Greenish yellow	α -Naphthoquinonesemicarbazone, $O:C_{10}H_8:N.NH.CO.NH_2$. — Cryst. fr. gl. ac. ac. I. aq., alc. — Action of alk. gives α -naphthol (Vol. I).
3840	248-9	Dark red-brown	Benzoidole, Aposafranone, $C_{12}H_9ON_2$. — \textcircled{P} Gran. w. green metallic lustre, d.s. h. aq. w. fuhsine-red color, & s. conc. H_2SO_4 w. green color changing to rose on diln. — E.s. alc., i. alk.
3841	248	Pale yellow	α -Azoxybenzoic Ac., $CO_2H.C_6H_4.N_2O.C_6H_4CO_2H$. — \textcircled{P} \textcircled{I} Azoxy comp. \textcircled{II} & ac., reducing boiling Fehling's sol. — Lft. V.d.s. h. aq., bz., chlf.; d.s. eth., c. alc.
3842	249	Red	p-Nitrobenzaldehyde-p-nitrophenylhydrazone, $NO_2C_6H_4CH:N.NH.C_6H_4NO_2$. — \textcircled{P} Hydrazone. \textcircled{II} s. w. pale violet color in aq.-NaOH, changing to deep violet-blue on diln. w. alc. — Flat ndl. w. violet reflections. I. aq.; v.d.s. alc., bz.; v.s. acetone.
3843	249-50	Light red	Dinitropurpuroxanthine, $(NO_2)_2C_{12}H_8O_4$. — Ndl. fr. gl. ac. ac. S. aq.; more s. alc., eth., gl. ac. ac. — BaA, dark red ndl.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34. ¶ = T. 2.35. ** = T. 2.1. \textcircled{I} = T. 2.17.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3844	249-50	Or.-red	p-Azobiphenyl, $\text{Ph.C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_4.\text{Ph}$. — \textcircled{P} \textcircled{D} Azo comp. — Plates fr. bz. I. alc.; e.s. eth.
3845	249	Or.-brown	p-Nitrobenzeneazo- β -naphthol, $\text{NO}_2.\text{C}_6\text{H}_4.\text{N}_2.\text{C}_10\text{H}_8.\text{OH}$. — \textcircled{P} \textcircled{D} Azo comp. — Tbl. w. metallic lustre fr. toluene. I. alc. or aq.-KOH; s. alc. = KOH. — Acetate,** flat or. ndl. fr. alc., v.d.s. alc., m.p. 192-3°.
3845	d. 246-52	Yellowish(?)	d-Lysine(α -Diaminocaproic Ac.) Picrolonate, $\text{NH}_2.\text{CH}_2.(\text{CH}_2)_3.\text{CH}(\text{NH}_2)(\text{CO}_2\text{H})$. — \textcircled{P} \textcircled{D} Salt of syrupy alkaline lysine w. No. 2.3184. — Cryst. fr. aq. E.s. aq.; less s. alc.
3847	d. 250	Yellowish	Pentamethylenediamine Picrolonate, $\text{NH}_2.\text{CH}_2.(\text{CH}_2)_5.\text{CH}_2.\text{NH}_2.2\text{C}_10\text{H}_8\text{O}_4\text{N}_4$. — \textcircled{P} \textcircled{D} Salt of No. 2.1232 w. No. 2.3184. D.s. aq., alc.
3848	d. 250	Yellowish	Tetramethylenediamine Picrate, $\text{NH}_2.\text{CH}_2.(\text{CH}_2)_4.\text{CH}_2.\text{NH}_2.2\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1201. — Silky triclin. ndl. Alm. i. c. aq.
3849	251	Light yellow	Biacridonyl, $\text{C}_{18}\text{H}_{14}\text{O}_2\text{N}_2$. — Tbl. fr. chlf. + gl. ac. ac. I. aq., alc.; d.s. gl. ac. ac.; e.s. chlf. — Ignition w. Zn dust gives acridine (No. 2.3102)!
3850	251-2	Light yellow	β -Naphthoquinoline Picrate, $\text{C}_{12}\text{H}_8\text{N.Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.716. — Cryst. ppt.
3851	d. 255-2	Light yellow	Trimethylamine Picrolonate, $\text{Me}_3\text{N.C}_10\text{H}_8\text{O}_4\text{N}_4$. — \textcircled{P} \textcircled{D} Salt of No. 2.1060 w. No. 2.3184. — Rhomb. tbl. s. in 1121 pt. c., or 166 pt. boiling aq.; s. in 794 pt. c., or 233 pt. boiling alc.
3852	252	Red	Nitrochrysoquinone, $\text{NO}_2.\text{C}_{16}\text{H}_8\text{O}_2$. — Ndl. fr. alc.
3853	deflagrates at 252	Yellow	Lysine Picrate, $\text{NH}_2.\text{CH}_2.(\text{CH}_2)_5.\text{CH}(\text{NH}_2)(\text{CO}_2\text{H})$. — \textcircled{P} \textcircled{D} Picrate* of alkaline syrupy lysine. — S. in 185 pt. aq. at 21°.
3854	253-4	Or.-brown	p-Nitrobenzeneazosalicylic Ac., $\text{NO}_2.\text{C}_6\text{H}_4.\text{N}_2.\text{C}_6\text{H}_3(\text{OH})(\text{CO}_2\text{H})$. — \textcircled{P} \textcircled{D} Acid & azo comp. — Ndl. fr. dil. ac. ac. E.s. alc.
3855	248-50; 253-5	Black; dark violet	$\alpha\beta$ -Naphthinduline, $\text{C}_{20}\text{H}_{17}\text{N}_2$. — \textcircled{P} Alc. sol. of salts fluor. fiery red! — Cryst. w. brassy lustre fr. bz. D.s. alc., eth., bz.
3856	254	Dark brown-red	3-Aminophenanthrenequinone, $\text{NH}_2.\text{C}_6\text{H}_3\text{O}_2$. — Ndl. fr. alc.
3857	254	Yellow	1,4-Dimethylxyloquinone-2,5-dioxime, $\text{Me}_2.\text{C}_6\text{H}_4.(\text{NOH})_2$. — \textcircled{P} S. alk. w. yel. color. — Cryst. fr. gl. ac. ac. I. aq.; v.d.s. h. alc., bz.; more s. h. alc. — $\text{BaA}_2\text{H}_2\text{O}$, ppt. of silky ndl. — \textcircled{D} Completely dect. to CO_2 & 8-hydroxyquinoline (No. 2.36) by distn.
3858	254-6 (in s.t.)	Light yellow	8-Hydroxyquinolinecarbonic(4) Ac., $\text{HO.C}_6\text{H}_4.\text{C}_3\text{H}_4\text{N}(\text{CO}_2\text{H})$. — \textcircled{P} FeCl_3 gives green color to aq. sol. — Cryst. powd. V.d.s. h. aq., bz.; more s. h. alc. — $\text{BaA}_2\text{H}_2\text{O}$, ppt. of silky ndl. — \textcircled{D} Completely dect. to CO_2 & 8-hydroxyquinoline (No. 2.36) by distn.
3859	255	Dark blue	Aminohydroxyanthraquinolinequinone, Alizarineblueamide, $(\text{NH}_2)(\text{HO}).\text{C}_{17}\text{H}_8\text{O}_2\text{N}$. — Ndl. fr. bz. V.d.s. alc.; d.s. c. bz., eth.; i. h. alk. Decd. by boiling dil. H_2SO_4 to alizarine blue & NH_4 .
3860	256	Golden	3,5-Dinitro-2-aminobenzoic Ac., $(\text{NO}_2)_2(\text{NH}_2).\text{C}_6\text{H}_3\text{CO}_2\text{H}$. — \textcircled{P} Acid giving NH_4 & 3,5-dinitrosalicylic acid (No. 2.257), when boiled w. NaOH sol. — Scales fr. alc. D.s. alc. — MeA , yel. lft., d.s. h. alc., m.p. 165°.
3861	256d.	Lemon-yellow	α -Nitrobenzaldehydesemicarbazone, $\text{NO}_2.\text{C}_6\text{H}_4.\text{CH}:\text{N.NH.CO}-\text{NH}_2$. — \textcircled{P} Semicarbazone.†† S. in h. NaOH sol. w. red color. — Ndl. fr. aq. D.s. h. alc.; i. eth.
3862	256-7	Brassy	Phenylmauvein, $\text{C}_{10}\text{H}_8\text{N}_4$. — \textcircled{P} S. conc. H_2SO_4 w. grass-green color, becoming blue on diln. w. aq. — Cryst. fr. bz.
3863	256-7d. u.c.	Y	Brucine Picrolonate, $\text{C}_{12}\text{H}_8\text{O}_4\text{N}_2.\text{C}_{10}\text{H}_8\text{O}_4\text{N}_4$. — \textcircled{P} \textcircled{D} Salt of No. 2.956 w. No. 2.3184. — Mic. cryst. (fr. components in alc. sol.). I. aq., alc. — Darkens fr. 214°.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3864	257d.	Red	d-Glucose-p-nitrophenylosazone, $C_8H_{10}O_4(N_2H_4C_6H_4NO_2)_2$. — \textcircled{P} S. in aq.-alc. NaOH sol. w. deep indigo-blue color! — Powd. fr. pyridine + eth. V.d.s.
3865	257	Red-yel.; gold-yel.	2-Nitroanthrenequinone, $NO_2C_{14}H_8O_2$. — Lft. fr. gl. ac. ac. Alm. i. alc. — \textcircled{D} Oxidn. w. CrO_3 mixt. gives 4-nitro-diphenic ac., ndl. fr. h. aq., m.p. 217 $^{\circ}$.
3866	d. 257	Yellowish(?)	Epiguanine Picrate, $C_8H_4ON_4Pk$. — \textcircled{P} \textcircled{D} Picrate of No. 2.2658. — Lft. or ndl. clusters. S. in 2740 pt. aq. at 18 $^{\circ}$. Melts w. efferv.
3867	258d.	Red	Lactose-p-nitrophenylosazone, $C_{12}H_{20}O_4(N_2H_4C_6H_4NO_2)_2$. — \textcircled{P} S. in NaOH sol. w. deep blue color! — Powd. fr. pyridine + eth. V.d.s.
3868	257-9	Cinnabar-red	Methylrosindone, $C_{17}H_{12}ON_2$. — Ndl. w. metallic reflections fr. alc. + bz. R.d.s. alc., eth., bz. The yel.-red sol. fluor. brick-red.
3869	259; 261-2	Red	Rosindone, $C_{12}H_8ON_2$. — 6-sided tbl. fr. toluene + alc. R.d.s. h. alc.; e.s. ac. ac.; i. NaOH sol. Conc. H_3SO_4 gives dark green sol. becoming orange-red on diln.
3870	259	Gold-yellow	3,5-Dinitro-4-aminobenzoic Ac., Chrysanic Acid., $(NO_2)_2(NH_2)C_6H_3CO_2H$. — \textcircled{P} Boiled w. KOH sol. gives NH_3 & 3,5-dinitro-p-hydroxybenzoic ac. (No. 2.442). — Cryst. fr. aq. or alc. Alm. i. c. aq.; d.s. h. aq.; mod. s. h. alc. — Ag \bar{A} , bulky yel. ppt. — Me \bar{A} , golden lft., m.p. 144 $^{\circ}$.
3871	260	Yellow	p-Succinodinitroanilide, $NO_2C_6H_4NH.CO.CH_2CH_2CO.NH-C_6H_4NO_2$. — Long ndl. fr. aniline. I. aq.; alm. i. eth., chlf., bz., gl. ac. ac.; v.d.s. alc. — Treatment w. Sn & HCl gives succinic ac. (Vol. I) & p-phenylenediamine (No. 2.877).
3872	abt. 260d.	Yellow	Isatinsemicarbazone, $NH_2CO.NH.N:C_6H_4ON$. — Ndl. fr. alc.; d.s. eth., bz., chlf.
3873	abt. 260	Pale yellow	α -Dinitroanthraquinone, $NO_2C_{14}H_8O_2$. — Mic. monoclin. cryst. fr. chlf. Alm. i. aq., eth.; v.d.s. alc., bz.; sl. more s. chlf.
3873-I	d. 260	Pale yellow	Diethylamine Picrolonate, $Et_2NH.C_6H_4O_2N_4$. — \textcircled{P} \textcircled{D} Salt of No. 2.1068-1 w. No. 2.3184. — Ppt. S. in 402 pt. boiling, or 3788 pt. c. aq.
3874	261d.	Red	Maltose-p-nitrophenylosazone, $C_{12}H_{20}O_4(N_2H_4C_6H_4NO_2)_2$. — Powd. fr. pyridine + eth. V.d.s.
3875	261	Red-brown	N-o,p-Dinitrophenylaminoindazole, $(NO_2)_2C_6H_4N_3$. — [Pat. Dyestuff intermediate.] — Cryst.
3876	261d.	Yellow	Nitrosodi- α -naphthylamine, $NO_2N(C_{10}H_7)_2$. — \textcircled{P} Prob. gives Tests 2.15 & 2.18! — Cryst. powd. D.s. alc.; e.s. chlf., bz. — Isomerized by Fischer & Hepp react. to deriv. of m.p. 169 $^{\circ}$, dark red ndl. fr. alc. (Ann., 243, 301).
3877	261	Yellow	Hexanitrodiphenylamine, m-Dipicrylamine, $(NO_2)_2C_6H_4NH-C_6H_4(NO_2)_2$. — Cryst. fr. gl. ac. ac. I. aq., alc., eth.
3878	263	Or.-yel.	p-Dinitrostilbene, $NO_2C_6H_4CH:CH.C_6H_4NO_2$. — \textcircled{P} Nitro comp., \ddagger s. conc. H_3SO_4 w. cherry-red color.
3879	263-4d.	Gold-yellow	Neurine Picrate, $CH:CH.NMe.Pk$. — \textcircled{P} \textcircled{D} Picrate* of v. toxic hygroscopic ptomaine. — Groups of feathery ndl. 100 pt. aq. at 23 $^{\circ}$ dis. 1.09 pt.; d.s. c. alc.; i. eth., bz., pet.-eth.; e.s. h. alc.
3880	d. 263	Yellow	Tetramethylenediamine Picrolonate, $NH_2(CH_2)_2CH_2.NH_2-ZPk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1201. — Ndl. V.d.s. aq., alc.
3881	263 (browning)	Light yellow	5-Nitro-2-aminobenzoic Ac., $(NO_2)(NH_2)C_6H_4CO_2H$. — \textcircled{P} Acid & nitro comp., \ddagger Boiled w. conc. KOH gives NH_3 &

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 \textcircled{P} = T. 2.35. \textcircled{D} = T. 2.1. $\textcircled{\textcircled{P}}$ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
			5-nitrosalicylic ac. (No. 2.416). — Ndl. E.s. alc., eth., h. aq.; d.s. c. aq.; alm. i. bz., chlf. — BaA ₃ .3H ₂ O, yel. ndl. or pr.; e.s. c. aq.; d.s. h. aq.
3882	264d.	Light red	4-Nitro-2-aminobenzoic Ac., (NO ₂)(NH ₂).C ₆ H ₄ .CO ₂ H. — (P) Acid & nitro comp.† — Ndl. — MeA (by boiling w. MeOH & H ₂ SO ₄), dark or. ndl., m.p. 157°, s. alc., lgr.
3883	265	Yellow	Dicyandiamidine Picrate, NH:C(NH ₂).NH.CO.NH ₂ .Pk. — (P) (D) Picrate* of No. 2.769.
3884	265	Yellow(?)	Adenine Picrolonate, C ₆ H ₄ N ₄ .C ₁₀ H ₈ O ₄ N ₄ . — (P) (D) Salt of No. 2.1057 w. No. 2.3184. — Cryst.
3885	266	Orange	4,4'-Tetramethyldiaminoazobenzene, Me ₂ N.C ₆ H ₄ .N ₂ .C ₆ H ₄ .NMe ₂ . — (P) (D) Azo comp. — Ndl. w. steel-blue reflections fr. bz. D.s. alc., eth.; e.s. h. bz. — B.Pk.C ₂ H ₅ OH, leaf-green ndl. fr. alc.
3886	267d.	Dark red	Diaminodiphenyleneazon, [C ₆ H ₄ (NH ₂).N:N.C ₆ H ₄ (NH ₂)].2H ₂ O. — Pr. fr. v. dil. alc. E.s. alc.; d.s. eth., bz.; i. lgr.
3887	267-70	Gold-yellow	Chrysaniline, Diaminophenylacridine, C ₁₁ H ₁₄ N ₂ . — [Salts used as dyestuff under name "Phosphine" (No. 3.1052-3).] — (P) Dyes wool golden-yel. — Cryst. fr. dil. alc. w. 2H ₂ O, or fr. bz. w. 1C ₆ H ₄ . — Alm. i. aq.; mod. s. alc. — B.HNO ₃ , ruby-red ndl., v.d.s. aq. — B.2Pk.H ₂ O (at 100°), ruby-red ndl. fr. alc.
3888	268	Yellow	2-Amino-7-hydroxyphenazine, C ₁₂ H ₉ ON ₂ . — (P) S. alc. or eth. w. green fluor. — Sbl. in ndl. — Diacetyl deriv.,** greenish-yel. ndl., m.p. 258°.
3889	abt. 268d.	Yellow	m-Nitrobenzaldehyde-2,4-dinitrophenylhydrazone, NO ₂ .C ₆ H ₄ -CH:N.NH.C ₆ H ₃ (NO ₂) ₂ . — V.d.s.
3890	268-9 (in s.t.)	Yellowish	Pyrococil, α -Pyrrolecarboxylic acid anhydride, [N(C ₆ H ₅).CO] ² . — [Fr. dry distn. of gelatine.] — Pearly lt. I. aq.; v.d.s. alc., eth.; s. undec. c. conc. H ₂ SO ₄ . Boiled w. KOH sol. gives salt of corresponding ac. (No. 2.356).
3891	268-70	Light yellow	1,3-Dinitro-2-hydroxyanthraquinone, (NO ₂) ₂ (HO).C ₁₀ H ₆ O ₂ . — Ndl. D.s. aq., alc., eth.; e.s. aniline w. brown-red color. — Boiling w. NaOH gives β -nitroalizarine (No. 2.3831). — KA, red lt. d.s. aq.
3892	269-70	Light yellow	Nitro-2-methylantraquinone, (NO ₂)Me.C ₁₀ H ₆ O ₂ . — (P) S. conc. H ₂ SO ₄ w. yel. color, changing to brown on warming, & giving purple ppt. on diln. w. aq. which dis. w. violet-blue color in alk. — Sbl. in alm. colorless ndl. V.d.s. alc., eth.; d.s. chlf., bz.
3893	270-1u.c.	Y-YO	o-Picrylaminobenzoic Ac., (NO ₂) ₂ C ₆ H ₄ .NH.C ₆ H ₄ .CO ₂ H. — (P) S. alk. w. deep red color. — Ndl. fr. cooling gl. ac. ac. sol.
3894	270	Dark brownish violet	Alizarine Blue, 3,4-Dihydroxyanthraquinolinequinone, C ₁₇ H ₁₄ O ₄ N. — (P) S. in ammon. w. blue color. — Ndl. fr. bz., or by sublimation. Vapors or.-red. I. aq.; alm. i. alc., eth.; s. h. bz. — Oxidn. by HNO ₃ gives phthalic ac. — B.Pk (T. 2.23), dark or.-red ndl. fr. bz., decd. by aq., m.p. 245°.
3895	270-3	Yellowish(?)	Cytosine Picrolonate, C ₆ H ₅ ON ₂ .C ₁₀ H ₈ O ₄ N ₄ . — (P) (D) Salt of No. 2.1056 w. No. 2.3184. — Ndl. V.d.s. alc.
3896	272-3	Gold-yellow	Dinitrobi-o-cresol, (NO ₂) ₂ .C ₆ H ₄ O ₂ . — Ndl. fr. gl. ac. ac. Sbl. w. decn. I. aq., alc., eth.; d.s. bz.
3897	d.w. frothing 272-4	Yellowish(?)	Guanidine Picrolonate, CH ₂ N ₄ .C ₁₀ H ₈ O ₄ N ₄ . — (P) (D) Salt of deliq. alk. guanidine w. No. 2.3184. — Fine ndl. fr. h. aq. D.s. c. aq.
3898	274	Red	Aminophenazine, NH ₂ .C ₁₂ H ₉ N ₂ . — (P) Solutions fluor. or.-red. — Ndl. w. bronzy lustre fr. alc. Sbl. in long red ndl.
3899	275	Orange	3-Nitrophenanthrenequinone, NO ₂ .C ₁₀ H ₈ O ₂ . — Ndl. fr. gl. ac.

(ORDER II.)

No.	Melting-point (C.).	Color.	COLORED SOLID COMPOUNDS.
3900	276-7d.	Salmon	Macrurindisazobenzene , $(\text{HO})_2\text{C}_6\text{H}_4\text{CO.C}_6(\text{OH})_2(\text{N}_2\text{Ph})_2$. — \textcircled{P} Azo comp. — Hair-like ndl. fr. nitrobenzene. I. alc., bz.
3901	272; 280	Brown	3-Dinitrobiphenol(4) , $(\text{NO}_2)_2(\text{HO})\text{C}_6\text{H}_4\text{C}_6\text{H}_4(\text{OH})(\text{NO}_2)_2$. — Cryst. fr. gl. ac. ac. I. alc.; d.s. gl. ac. ac.
3902	276d.	Gold-yellow	5,7-Dinitro-8-hydroxyquinoline , $(\text{NO}_2)_2(\text{HO})\text{C}_8\text{H}_4\text{N}$. — D.s. lft. Forms yel. salts.
3903	275-8	Gold-yellow	Diacetaminoazoxybenzene , $(\text{C}_2\text{H}_5\text{O})\text{NH.C}_6\text{H}_4\text{N.O.C}_6\text{H}_4\text{NH-C}_2\text{H}_5\text{O}$. — Hair-like ndl. V.d.s. c. alc.
3904	276	Lemon-yellow	2'-Methyl-3'-amino-9-phenyl-1,2-naphthacridine , $\text{C}_{14}\text{H}_{14}\text{N}_2$. — [Pat.] — \textcircled{P} D.s. alc. w. yellow & green fluor., changing to or. on addition of HCl. — Ndl. fr. aniline.
3905	abt. 277; 277-8(r.h.)	Gold-yellow	Bis-p-dimethylaminobenzylidene-p-phenylenediamine , $\text{NMe}_2\text{-C}_6\text{H}_4\text{CH:N.C}_6\text{H}_4\text{N:CH.C}_6\text{H}_4\text{NMe}_2$. — \textcircled{P} Dyes tannin-mordanted cotton red-brown. — Ndl. fr. dimethylaniline. I. aq., alc. Decd. by boiling aq. to p-dimethylaminobenzaldehyde & p-phenylenediamine. — $\text{B.} 2\text{HCl.} 5\text{H}_2\text{O}$, brown-red or steel-blue ndl., v.d.s. c. aq., or w. $1\text{H}_2\text{O}$ in light red mic. ndl.
3906	277-9d.	Dark brown-red	p-Nitrobenzeneazo(4)-naphthol(1) , $\text{NO}_2\text{C}_6\text{H}_4\text{N}_2\text{C}_{10}\text{H}_8\text{O.H}$. — \textcircled{P} \textcircled{D} Azo comp. — Ndl. w. steely lustre fr. nitrobenzene. V.d.s. h. alc., etc. — Reduction w. Sn & HCl gives No. 2.877 & 1,4-aminonaphthol(1). — Acetyl deriv.,** ruby ndl., m.p. 165-6°.
3907	279	Brown	2(3)-Aminophenanthrophenazine , $\text{C}_{18}\text{H}_{13}\text{N}_3$. — \textcircled{P} Sol. in conc. H_2SO_4 , carmine-red; in eth., yel. w. green fluor. — Cryst. powd. fr. phenol & alc. D.s. usual solvents. Sbl.
3908	280	Dark yellow	3,6-Diaminophenazine , $(\text{NH}_2)_2\text{C}_6\text{H}_4\text{N}_2$. — \textcircled{P} Sol. in conc. H_2SO_4 , green, becoming blue, violet, & red on diln. Salts dye silk red. — Ndl. fr. h. aq. D.s. c. aq.; v.s. alc., eth. — B.Pk, brown d.s. ndl., m.p. abt. 330°.
3909	279-80w. efferv.	Dark yellow	Adenine Picrate , $\text{C}_6\text{H}_5\text{N}_3\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate of No. 2.1057. — Mic. pr. fr. h. aq. Also cryst. w. $1\text{H}_2\text{O}$ in silky aggregates. S. in 3500 pt. c. aq.; e.s. alc.
3911	280	Egg-yellow	α-8-(o)Hydroxyquinolinecarbonic Ac. , $\text{HO.C}_6\text{H}_4\text{C}_8\text{H}_5\text{N}(\text{CO}_2\text{H})$. — \textcircled{P} FeCl_3 colors aq. sol. green. — Cryst. d.s. h. aq., alc., eth.; i. gl. ac. ac., lgr., bz. — Br-aq. in acid sol. gives CO_2 & dibromoxyquinoline, m.p. 193°. — Dry distn. gives 8-hydroxyquinoline (No. 2.36).
3912	280	Yellow	β-Dinitroanthraquinone , (Fritzsche's reagent), $(\text{NO}_2)_2\text{C}_{14}\text{H}_6\text{O}_3$. — Ndl. fr. h. gl. ac. ac. D.s. alc., eth., chlf., bz.; mod. s. h. gl. ac. ac. Sbl. in alm. colorless lft. — \textcircled{D} Sol. of 9 pt. anthracene + 10 pt. dinitroanthraquinone in 100 pt. boiling toluene gives violet rhomb. lft. of anthracene deriv., $\text{C}_{14}\text{H}_{10}\text{C}_{14}\text{H}_6(\text{N}_2\text{O}_2)_2\text{O}_2$, on cooling.
3913	275-86d. u.c.	Y	Strychnine Picrate , $\text{C}_{27}\text{H}_{25}\text{O}_2\text{N}_3\text{Pk}$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1047. — Powdery ppt. Alm. i. aq., alc., chlf.
3914	280	Pale yellow	Dinitrobinaphthyl , $(\text{NO}_2)_2\text{C}_{20}\text{H}_{12}$. — Ndl. V.d.s. h. bz. or gl. ac. ac.; i. most solvents.
3915	281	Yellowish	3,6-(s)-Diaminoacridine , $(\text{NH}_2)_2\text{C}_{15}\text{H}_7\text{N}$. — Ndl. fr. alc. E.s. alc.; d.s. bz.
3916	283	Brown	α-Trinitronaphthoic Ac. , $(\text{NO}_2)_3\text{C}_{10}\text{H}_4\text{CO}_2\text{H}$. — \textcircled{P} Acid of intensely bitter taste. — Long ndl. fr. alc. E.s. h. eth. — $\text{CaA.} 5\text{H}_2\text{O}$, lft. or ndl., e.s. h. aq. — EtA, brown ndl. fr. alc., m.p. 131°.
3917	283-4	Yellow	α-($\alpha\beta$)-Dinaphthazine , $\text{C}_{20}\text{H}_{12}\text{N}_2$. — \textcircled{P} S. undec. in conc. H_2SO_4 w. violet-blue color! S. bz. w. green fluor.! — Ndl. fr. gl. ac. ac. Alm. i. alc., bz., gl. ac. ac., eth. Volatile undec.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.34. ¶ = T. 2.35. ** = T. 2.1. §§ = T. 2.17.

(ORDER II.)

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3918	283-5d.	Carmine-red	2,4-Dinitrobenzaldehyde-p-nitrophenylhydrazone , $(NO_2)_2.C_6H_4.-CH:N.NH.C_6H_3.NO_2$. — \textcircled{P} S. w. red color in conc. H_2SO_4 . — Ndl. fr. nitrobenzene. I. aq., dil. NaOH; d.s. alc., acetone. (Not hydrolyzed by h. HCl & ac. ac.)
3919	284	Reddish yellow	3-Nitro-4-aminobenzoic Ac. , $(NO_2)(NH_2).C_6H_4.CO_2H$. — \textcircled{P} Boiling w. v. conc. KOH sol. (T. 2.26) gives NH_3 & nitro-p-hydroxybenzoic ac. (No. 2.294). — Ndl. fr. alc. I. aq.; d.s. h. alc. — $Ba\bar{A}_2.5H_2O$, or. cryst., d.s. h. aq.
3920	abt. 285w. efferv.	Light yellow	Tropidine Picrate , $C_2H_{12}N.Pk$. — \textcircled{P} \textcircled{D} Picrate* of No. 2.1205. — Pr. fr. h. aq.
3921	285-6	Yellow	Tetramido-β-naphthylamine , $(NO_2)_4.C_{10}H_{11}N$. — Cryst. grains fr. h. nitrobenzene. V.d.s. alc., eth., bz.; e.s. h. nitrobenzene.
3922	286-7	Yellow-white	5-Nitronaphthoic(2) Ac. , $NO_2.C_10H_8.CO_2H$. — \textcircled{P} Acid & nitro comp. — Ndl. V.d.s. h. aq.; d.s. alc.
3923	285-7	Yellow	Indophenazine , $C_{16}H_{12}N_2$. — \textcircled{P} S. mineral ac. w. red-brown color. — Ndl. fr. alc. Sbl. D.s. alc.; e.s. eth., h. chlf., bz.; alm. i. alk. — $Ag\bar{A}$ (100°), red-brown gelat. ppt. — Acetyl deriv., ** ndl. fr. dil. alc., m.p. 202°.
3924	286-8	Green	Induline 6B , Anilinophenylaminophenylinduline , $C_{16}H_{12}N_2(?)$. — \textcircled{P} Sol. in aniline purple-blue! Ndl. w. metallic lustre fr. aniline. — $B.HCl$, green cryst. s. conc. H_2SO_4 w. intense blue-green color; i. usual solvents.
3925	289d.	Gold-yellow	4-(α)-Nitroalizarine , $(NO_2)(HO)_2.C_14H_8O_2$. — [Pat.] — \textcircled{P} S. KOH sol. w. blue-violet color. — Ndl. fr. alc. or gl. ac. ac. Sbl. w. carbonization. — Oxidn. w. HNO_3 (Ann., 201, 353) gives phthalic ac. (Vol. I).
3926	290	Intense yellow	2,7-(α)-Dinitrofluorenone , $C_18H_8(NO_2)_2.CO.C_6H_4(NO_2)^2$. — Ndl. fr. gl. ac. ac. V.d.s. h. alc.; e.s. xylene.
3927	290	Yellow	Benzil-p-nitrosazone , $Ph.C(:N.NH.C_6H_4.NO_2).C(:N.NH.C_6H_4-NO_2).Ph$. — Cryst. powd. fr. pyridine + eth. D.s. except in pyridine & nitrobenzene.
3928	290-5	Yellow-red	4-Nitro-1-aminoanthraquinone , $(NO_2)(NH_2).C_{14}H_8O_2$. — Ndl. fr. epichlorhydrine. I. aq.; d.s. alc. Sol. in conc. H_2SO_4 , yel.
3929	294; (264)	Yellow	6-Aminonaphthophenazine , Phenonaphtheturhodine , $NH_2-C_{12}H_8N_2$. — \textcircled{P} Sol. fluor. green! — Cryst. fr. alc. D.s. c. alc., eth., bz. — $B_2H_2PtCl_6$ (100°), d.s. red ndl.
3930	295	Red	Naphthindone , $C_{18}H_{16}ON_2$. — Tbl. w. brassy lustre fr. cumene. — Ignition w. Zn dust gives No. 2.3828.
3931	295	Brownish	8-Nitronaphthoic(2) Ac. , $NO_2.C_10H_8.CO_2H$. — \textcircled{P} Acid & nitro comp. — Ndl. fr. alc. Alm. i. aq.; s. in 600 pt. c. alc.; d.s. eth., chlf., bz. — $Me\bar{A}$, pale yel. ndl. fr. alc., m.p. 112°.
3932	Blackens w.m. abt. 300	Y Bkn.	Citrazinic Ac. , 2,6-Dihydroxypyridinecarboxylic(4) Ac. , $C(CO_2H)-:CH(C(OH):N.C(OH):CH^2)$. — \textcircled{P} Heat to boiling on crucible cover 5 or 10 drops of mixt. of 1 drop c. sat. aq. $NaNO_2$, 1-5 cc. aq. Place abt. 10 mg. acid on dry part of cover and v. gradually stir h. sol. into it w. rod. Spread mixt. over entire surface of still warm cover & allow to stand until cool. Dis. residue in abt. 25 cc. c. aq. Sol. will be of intense pure blue (B) color! — Powder. Alm. i. h. aq. or h. alc.; e.s. alk. or alk. carbonates. Sol. in ammon. shows pale violet fluor. Odorless.
3933	296u.c. explodes w.m. (307.6c.)	Yellow	Tetranitroanthraflavlic Ac. , $(NO_2)_4(HO)_2.C_14H_8O_2$. — Ndl. Mod. s. h. aq. w. red color; s. alc., eth.
3934	298d.	Yellow-red	4-Nitro-3-aminobenzoic Ac. , $(NO_2)(NH_2).C_6H_4.CO_2H$. — \textcircled{P} \textcircled{D} Acid. — Cryst. fr. alc. D.s. aq.; mod. s. alc. Boiling w. KOH sol. (Cf. T. 2.26) gives NH_3 & No. 2.3788-1. — Reduced by Sn & HCl to No. 2.362. — $Ba\bar{A}_2.2H_2O$, yel.-red pr., d.s. h. aq.; v.d.s. c. aq.

No.	Melting-point (C. $^{\circ}$).	Color.	COLORED SOLID COMPOUNDS.
3935	abt. 300	Yellow	δ -Dinitroanthraquinone, $(NO_2)_2C_{14}H_8O_2$. — Scales fr. gl. ac. ac. S. alc., gl. ac. ac.
3936	302	Red	2-Aminoanthraquinone, $NH_2C_{14}H_8(CO)_2$; C_8H_4 . — Sbl. in small ndl. I. aq., alk., eth.; mod. s. alc., bz. — N_2O_2 passed into alc. sol. gives ppt. of brown-yel. flocks, m.p. 238–40°. — Fusion w. KOH followed by oxidn. of product by air gives blue vat-dyestuff, indanthrene. (D.R.P., 129,845, 129,847, 129,848.)
3937	301–3; 294; 290	Gold-yellow	2,7-Dinitrophenanthrenequinone, $(NO_2)_2C_{14}H_8O_2$. — Lst. fr. gl. ac. ac. V.d.s. gl. ac. ac. or alc.
3938	305	Garnet-red	α -Quinophthalin (α -Imide of Quinophthalone), $C_{15}H_{12}ON_2$. — Cryst. fr. alc. V.d.s. alc., bz., chlf., eth. — $B_2H_6PtCl_6$ (T. 2.14), ndl., m.p. 278°.
3939	307	Yellow-red	Phenylisophtalylcarbazolequinone, $[C_6H_5NH.C_6H_4O_2]^+$. — \oplus S. conc. H_2SO_4 w. red-violet color. — Pointed ndl. D.s. alc.; s. h. gl. ac. ac., h. bz.
3940	311d.	Red	Glyoxal-bis-p-nitrophenylhydrazone, $NO_2C_6H_4NH.N:CH.CH:NH.C_6H_4NO_2$. — \oplus S. alc.-KOH w. deep blue color! — V.d.s. except in warm nitrobenzene.
3941	310–5d.	Black-brown	Isosafranimone, $C_{15}H_{12}ON_2$. — Ndl.
3942	d.w. efferv. 311–5	Yellow	Guanidine Picrate, $HN:C(NH_2)_2Pk$. — \oplus Picrate* of s., alk., deliq. base. — Rosettes of mic. ndl. S. in 2630 pt. aq. at 9°. D.s. alc., eth.
3943	320d.	Pale yellow	m-Azoxybenzoic Ac., $(CO_2H)C_6H_4N_2O.C_6H_4(CO_2H)$. — \oplus \oplus Azoxy acid. Mic. cryst. I. aq.; d.s. alc., eth., — $Ag_2\bar{A}$, floc. ppt.
3944	abt. 325	Light yellow	Amino-s- $\alpha\beta$ -naphthazine, $NH_2C_{20}H_{11}N_2$. — \oplus S. conc. H_2SO_4 w. blue-violet color. — Ndl. w. greenish reflections fr. nitrobenzene. Sbl. Alm. i. bz., alc., these sol. showing bright yel.-green fluor.
3945	337d.	Sulphur-yellow	Dinitrobianthryl, $(NO_2)_2C_{20}H_{16}$. — Small ndl. E.s. chlf., bz. — \oplus Oxidn. by CrO_3 in gl. ac. ac. sol. gives anthraquinone (Vol. I).
3946	354(c.)	Yellow	9-Acridone, $[C_8H_4NH.C_6H_4CO]^+$. — \oplus Alc. sol. shows intense blue fluor! Thick ndl. fr. alc. S. h. alc., gl. ac. ac.; alm. i. eth., chlf., bz. — \oplus Ignition w. Zn dust gives acridine (No. 2.3102).
3947	350–5u.c.	Bronzy; red-brown	Phenanthroneazine, $C_{20}H_{11}ON$. — [Fr. phenanthrenequinone & phenylhydrazine.] Ndl. w. green metallic reflections fr. h. xylene. Sbl. w. sl. decn. above m.p. in cryst. red-brown scales. D.s. nitrobenzene, xylene; alm. i. most solvents. S. conc. H_2SO_4 w. violet color, changing to blue & dull green-brown on contact w. $K_2Cr_2O_7$ cryst. Conc. HNO_3 dis. w. fuchsine-like violet-red color, changing to green-blue on cautious diln. w. aq., more aq. giving ppt. of green-blue flocks w. copper-red reflections.
3948	390–2d. (r.h. in s.t.)	BVS3	Indigotine, Indigo Blue, $C_{16}H_{10}O_2N_2$. — [Dyestuff. Cf. No. 3.130.] — Sbl. w. violet vapor giving rhomb. cryst. Streak coppery-red. I. aq., alc., eth., dil. ac. or alk.; s. h. chlf. or aniline.
3949	440–1	Yellow	Phenanthroneazine, Diphenanthryleneazotide, $C_{20}H_{11}N_2$. — \oplus S. conc. H_2SO_4 w. intense blue color! — Lust. greenish ndl. fr. nitrobenzene. Sbl. in broad yel. ndl. I. gl. ac. ac.; v.d.s. eth.; s. nitrobenzene, aniline.

Explanation of typographical signs used in this Division: * = T. 2.39. † = T. 2.21. § = T. 2.36. || = T. 2.24.
 ¶ = T. 2.55. ** = T. 2.1. §§ = T. 2.17.

ALPHABETICAL INDEX.

This index includes the names of all compounds described in the tables of Volume II, and also serves as a general index to the non-tabular sections of the volume.

A reference numeral directly following the name of an indexed compound is its "Serial number"; but wherever such a numeral is preceded by "p" the reference is to a page number.

In seeking for any specific description through the index, it is best to look first for the colloquial or "trivial" name of the compound, if it has one which is widely used. Structural names, when there is no special reason for departing from the rule, have usually been constructed in accordance with the principles of terminology followed in the last edition of Beilstein's Handbuch. Synonyms are used sparingly. Such literal or syllabic prefixes as *o*, *m*, *p*, *d*, *l*, *dl*, *i*, *n*, *s*, *as*, *cis*, *trans*, *fum.*, *mal.*, *O*, *C*, and the *Greek letters*, have *not* been treated as integral parts of the names in which they occur in fixing the sequence of words in the alphabetical arrangement. In numerical prefixed and suffixes numerals set off with hyphens refer to *following*, and those inclosed by parentheses to *preceding*, word elements. In indexing names of compounds long dashes are employed to represent words, or parts of words, repeated from corresponding positions from lines above. When such dashes stand for parts of words only, the points of division are shown in the first word of the series by hyphens.

A

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