

### III. *On some of the Products of the Decomposition of Nitrotoluylic Acid.*

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IN a paper read before the Chemical Society, and published in the third volume, p. 421, of the Transactions of that body, I described the mode of preparation and properties of two new organic acids, which, from their general analogy to benzoic and nitro-benzoic acids, were considered to be the representatives of those acids in the succeeding or *Toluylic* series, and were named accordingly *Toluylic* ( $C_{16}H_8O_4$ ) and Nitrotoluylic ( $C_{16}\left\{\begin{smallmatrix} H_7 \\ NO_4 \end{smallmatrix}\right\}O_4$ ) acids.

These two acids, interesting as supplying two members of a well-defined series of acids, and as establishing a parallelism between the benzoyl and toluylic groups, were formed by the oxidation of *Cymol*, a hydrocarbon ( $C_{20}H_{14}$ ) occurring with *cuminol*, an oxyhydrocarbon ( $C_{12}H_{12}O_2$ ) in the oil of cumen (*Oleum cumini*), from which source alone they have hitherto been obtained.

The original object in submitting cymol to the action of fuming nitric acid was the formation of the substitution compound  $C_{20}\left\{\begin{smallmatrix} H_{13} \\ NO_4 \end{smallmatrix}\right\}$ , from which, by the action of reducing agents, it was anticipated that an organic base, analogous to those formed from nitrobenzole (aniline)\*, from nitrotoluole (toluidine)†, and from nitrocumole (cumidine)‡, would have been obtained: the unexpected formation however of a crystalline acid gave a different turn to the investigation, and the search for cumidine ( $C_{20}H_{15}N$ ) was for the time laid aside.

I have at various times since the publication of this first paper returned to the study of the action of nitric acid on cymol, and have made many attempts, by employing acids of various degrees of strength, by keeping down the temperature by surrounding the vessels with freezing mixtures, &c., to check the action of the acid before it had produced its maximum degree of oxidation on the oil; yet though I have in this way obtained several beautifully crystallized intermediate products, and amongst them a compound homologous to *dinitrobenzole*, in which two equivalents of the hydrogen of the oil have given place to two of hyponitric acid, viz.  $C_{20}\left\{\begin{smallmatrix} H_{12} \\ 2NO_4 \end{smallmatrix}\right\}$ , still the principal

\* ZININ, *Annal. der Chem. und Pharm.* Bd. xlv. p. 283.

† MUSPRATT and HOFMANN, *Mem. Chem. Soc.* vol. ii. p. 373.

‡ NICHOLSON, *Chem. Soc. Quart. Journ.* vol. i. p. 2.

product of the action has been nitrotoluylic acid; and I have not hitherto succeeded in obtaining the first substitution compound  $C_{20} \left\{ \begin{smallmatrix} H_{13} \\ NO_4 \end{smallmatrix} \right\}$ , and of the intermediate substances above alluded to, quantities insufficient for a thorough investigation of their nature.

The study of the products of the decomposition of nitrotoluylic acid has been attended with more successful results: some new and interesting compounds have been obtained, and as these, viewed in connection with those described in my former paper, and with those of CHANCEL obtained in his researches on nitrobenzoic acid\*, complete the proofs of the parallelism of the benzoyl and toluyll groups, I have thought that a brief account of them might be deemed not unworthy of the notice of the Royal Society.

#### *Preparation of Nitrotoluylamide.*

Among the salts analysed in order to determine the atomic weight of nitrotoluylic acid was the ethyl compound, prepared by submitting an alcoholic solution of the acid to a current of dry hydrochloric acid gas till copious fumes were evolved, and then distilling. Some attempts were made to convert this ether into nitrotoluylamide by saturating its alcoholic solution with dry ammoniacal gas, and setting it aside in a well-stopped bottle; after some months however the anticipated transformation had not taken place. Equally unsuccessful was the attempt to prepare this amide by the action of heat on nitrotoluylate of ammonia, as FIELD† procured *cuminamide* from cuminate of ammonia; an explosion invariably took place, however carefully the heat was applied. This investigation was resumed by repeating the experiment with the ether in sealed tubes exposed to heat in a water-bath. About 5 grms. of the crystallized salt were placed in a stout glass tube, 2 feet long and three quarters of an inch in diameter; the tube was then filled to within 2 inches of the top with strong aqueous ammonia, sealed, and submitted to a boiling temperature in a deep water-bath. After forty-eight hours' continuous action the upper part of the tube had become filled with brilliant yellow crystalline plates, which were easily removed on breaking the tube from the solid and unaltered ether below. Two or three crystallizations sufficed for their purification. They were dried *in vacuo* over sulphuric acid, and then slowly brought to their fusing-point in the air-bath. The crystals began to fuse at 245° FAHR.; at 255° they were fairly liquid, and at 260° they emitted copious vapours.

*Analysis.*—The combustion was made in the usual manner with oxide of copper:

I. .524 grm. yielded 1.029 grm. carbonic acid and .220 grm. water.

II. .420 grm. gave .886 grm. carbonic acid and .187 grm. water.

\* Compt. Rend. vol. xxviii. p. 293; LAURENT and GERHARDT, C. R. 1849, p. 177; Journ. Pr. Chem. xlvii. p. 140; Ann. Ch. Pharm. lxxii. 274.

† Mem. Chem. Soc. vol. iii. p. 404.

These numbers furnish the following per-centages of carbon and hydrogen :—

	I.	II.
Carbon . . . .	53·55	53·69
Hydrogen. . . .	4·66	4·61

The formula of nitrotoluylamide,  $C_{16}\left\{\begin{smallmatrix} H_6 \\ NO_4 \end{smallmatrix}\right\}O_2, NH_2$ , or if we adopt the theory of KOLBE, that toluylic acid like benzoic acid is the oxide of a conjugate radical  $(C_{14}H_7)C^2$



requires

Carbon . . . .	53·33
Hydrogen . . . .	4·44

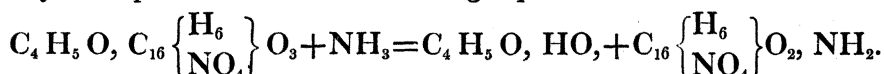
*Properties.*—Nitrotoluylamide is in its properties exactly analogous to nitrobenzamide, from which it differs in composition by  $(C_2H_2)$ , the constant ascending or descending term of these organic series. It is very sparingly soluble in cold water, more so in hot; but very soluble in alcohol, ether, and pyroxylic spirit. Its alcoholic solution crystallizes spontaneously in long, brilliant, yellow needles. Its fusing-point is about  $245^\circ \text{FAHR.}$ ; that of *nitrobenzamide* was determined by CHANCEL to be under  $212^\circ$ ; when heated with a concentrated solution of potash, it is converted into nitrotoluylate of that base.

By the process above described for procuring this substance, it can only be obtained in very limited quantities, and the process is by no means unattended with danger, from the liability of the tube to burst under the great pressure to which it is exposed; and whilst searching for some better method of procuring it in quantities sufficient to study the products of its decomposition, I became acquainted with the investigations of CHANCEL upon the nitro-compounds derived from benzoic acid\*, and upon trying the method there proposed by him for the preparation of nitrobenzamide, I succeeded, after many fruitless trials, in obtaining the new amide in considerable quantities. The purified ethyl compound was dissolved in ordinary spirits of wine, and aqueous ammonia added until a precipitate began to form: the solution was transferred to a stout green glass bottle, and the stopper being well-secured, it was plunged into a water-bath and kept for several days at a temperature of about  $140^\circ \text{FAHR.}$  The action proceeded very slowly; with about 15 grammes of ether it was usually completed in eight or ten days. The liquid, from being nearly colourless, was now of a deep yellow colour, and no precipitation took place on the addition of a large quantity of water. The ammoniacal alcohol was distilled off, and the residual crystalline mass redissolved in weak alcohol and boiled with animal charcoal; from the filtered solution pure nitrotoluylamide separated in crystals of remarkable size and brilliancy.

The formation of nitrotoluylamide by the action of ammonia on nitrotoluylate of

\* LAURENT and GERHARDT, C. R., 1849, 177, Annual Rep. Prog. Chem. vol. iii. part 1. p. 223.

oxide of ethyl is represented in the following equation :—



*Action of Reducing Agents on Nitrotoluylamide.*

When an alcoholic solution of nitrotoluylamide is submitted to the action of hydrosulphate of ammonia, the reaction is of a complicated nature, the alcohol itself undergoes decomposition, and its elements take a part in the formation of the new products. But an aqueous solution of the amide undergoes under the influence of this reducing agent a very interesting decomposition, precisely analogous to that of nitrobenzamide under similar circumstances. Ten grammes of the substance were dissolved in three pints of boiling distilled water, and agitated while still hot with strong solution of hydrosulphate of ammonia; the liquid was then thoroughly saturated with sulphuretted hydrogen, the bottle well stopped, and kept for several days plunged in a water-bath at the temperature of about 140° FAHR. The solution, which had now acquired a dark brown colour, was slowly evaporated on the water-bath; large quantities of sulphur separated during the evaporation, which were from time to time removed. When the liquid was reduced to about six ounces it was filtered and set aside. In a few hours a mass of fawn-coloured crystals was deposited; these were redissolved in water and boiled twice with animal charcoal. On cooling, a crop of beautiful pale-yellow cubical crystals made their appearance; these were again digested with animal charcoal, and from the filtered solution, fine, large, nearly colourless and transparent rhombic prisms of the new substance were obtained. They were dried first between folds of bibulous paper, and then *in vacuo* over sulphuric acid, after which no further diminution in weight took place on reducing them to powder and heating them for two hours in an air-bath at their melting-point, viz. 240° FAHR. Unlike therefore the homologous compound obtained by CHANCELLOR from nitrobenzamide, this substance contains no water of crystallization. When the fused mass was removed from the air-bath, at the moment of solidification, it burst with a slight explosion into a heap of glistening crystalline plates.

*Analysis.*—The combustion was made in the usual manner with oxide of copper.

I. 442 grm. produced 1.035 grm. carbonic acid and .270 grm. water.

II. 4057 grm. yielded .948 grm. carbonic acid and .249 grm. water, numbers which correspond centesimally to

	I.	II.
Carbon . . . .	63.86	63.72
Hydrogen . . . .	6.79	6.82

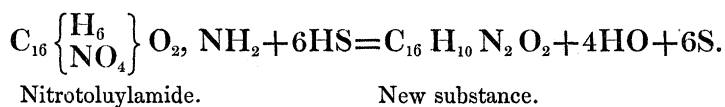
On heating a portion of the substance with soda-lime in a green glass retort, ammonia was first plentifully disengaged; but on raising the heat a yellowish oily liquid distilled over, which in the course of half an hour solidified into a crystalline mass, having the peculiar smell of the *aniline* class of organic bases. Although no doubt was entertained that this substance was *toluidine*, it was thought desirable to deter-

mine the fact experimentally. Three or four grammes of the new substance were therefore distilled with caustic potash, and the solid product, after having been washed two or three times with water, was saturated with an alcoholic solution of oxalic acid. It was evaporated to dryness on the water-bath, then redissolved in boiling alcohol and filtered; the crystals which separated on cooling were washed with cold distilled water, then redissolved in boiling water, and decomposed by caustic potash; some clear oily drops separated, which were dissolved in ether, and the ethereal solution evaporated to dryness. The residue was dissolved in hydrochloric acid, and mixed with an alcoholic solution of bichloride of platinum; a fine orange-coloured crystalline mass was obtained, which was washed with ether and dried on the water-bath. It was decomposed by ignition in a porcelain crucible.

*Analysis*.—595 grm. yielded 1872 grm. of metallic platinum = 31.46 per cent. The formula of the double chloride of platinum and toluidine ( $C_{14}H_9N$ ,  $HCl$ ,  $PtCl_2$ ) requires 31.6 per cent. of platinum.

The crystalline body obtained by the distillation of the new substance with caustic potash was therefore evidently *toluidine*.

The decomposition which nitrotoluylamide undergoes under the influence of hydrosulphate of ammonia is in exact accordance with the reactions constantly observed between this reducing agent and nitrated organic bodies, and is represented in the following equation:—



The proportion per cent. of carbon and hydrogen required by the formula

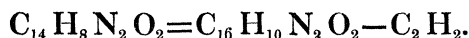


are,

Carbon . . . . .	64.00
Hydrogen . . . . .	6.61

which agree very closely with those obtained by the combustion of the new substance with oxide of copper.

This substance has two homologues in the benzoyl or immediately preceding series, viz. *carbamide*—*carbanilide*, a neutral body discovered by Dr. HOFMANN\*, and the true *urea* of the aniline series discovered by M. CHANCEL†, both represented by the formula

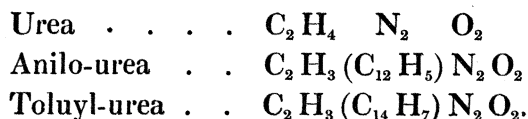


The origin of the crystalline compound under examination, which is perfectly analogous to that of *aniline urea*, renders it at once probable that it is the analogue of the latter, an assumption which was fully corroborated by the study of its properties and chemical deportment.

\* Chem. Soc. Quart. Journ. vol. ii. p. 36.

† Compt. Rend. xxviii. p. 293; LAURENT and GERHARDT, C. R. 1849, p. 177; Journ. Pr. Chem. xlvii. p. 140; Ann. Ch. Pharm. lxxii. p. 274.

It has all the chemical functions of ordinary urea, its taste is cooling and slightly bitter; it is very soluble in water, alcohol, and ether. Its aqueous solution may be kept for some time without change, but after some weeks it acquires a brownish colour, and flocculent matter is deposited. It is a true organic base, and forms crystalline compounds with several acids. On dissolving some of the crystals in weak nitric acid and placing the solution under a bell jar over sulphuric acid, the whole soon became a crystalline mass; and when the small quantity of adhering nitric acid was removed by affusion with cold water, and the crystals dissolved and boiled with animal charcoal, the nitrate was obtained by spontaneous evaporation in large and nearly colourless thin plates, the taste of which was cooling and slightly bitter, hardly to be distinguished from that of nitre. By dissolving some of the crystals in dilute oxalic acid, and placing the filtered solution over sulphuric acid, a crystalline mass was obtained, from which the oxalate of the new base was obtained in long, transparent, and nearly colourless needles, by washing away as before the excess of acid, redissolving in water, and allowing the solution to crystallize spontaneously. The limited quantity of this substance at my disposal, and the very great difficulty of preparing it, prevented my making a quantitative determination of any of its salts. When heated a little above its fusing-point the base takes fire, burning with a red smoky flame and leaving much carbon. I propose for it the name of *toluyl-urea*; its relation with CHANCEL's compound and with the urea type appears from the following synopsis:—



*Product of the Decomposition of Toluyl-urea.*

On heating some of the pulverized crystals strongly with caustic potash, ammonia was first disengaged, and then *toluidine* distilled over; when, however, an aqueous solution was boiled with a solution of caustic potash till ammonia ceased to be evolved and the potash then neutralized with an acid, the acid being left slightly in excess, a clear yellow liquid was obtained, from which after a few hours' standing there was deposited a group of magnificent golden-yellow needles, which only required a second solution and crystallization to render them perfectly pure. They were dried first between folds of filtering paper, then reduced to powder and thoroughly dried in the air-bath, after which they underwent no further loss of weight by being fused.

*Analysis.*—This was made in the ordinary manner by combustion with oxide of copper.

I. .3355 grm. produced .780 grm. carbonic acid and .188 grm. water.

II. .4042 grm. gave .939 grm. carbonic acid and .224 grm. water,

corresponding centesimally to

	I.	II.
Carbon . . . .	63.46	63.22
Hydrogen . . . .	6.22	6.15

By heating with soda-lime toluidine distilled over and solidified in the receiver. The new body thus formed is a well-defined acid, producing a series of crystalline salts; those of potash, soda, and ammonia, crystallized with difficulty; the lime and baryta salts, though exceedingly soluble, were obtained after long standing over sulphuric acid in fine acicular tufts. The lead salt was obtained by dropping the ammonia compound into solution of acetate of lead; it fell as a white powder, which when boiled with a large quantity of distilled water, and filtered, separated, as the solution cooled, in the form of long, brilliant, transparent, and nearly colourless needles with tufted ends. By decomposing this lead salt by dilute sulphuric acid, the new acid was again obtained in long needles having a fine crimson colour. On adding a solution of the ammonia salt to a solution of nitrate of silver, a copious gray crystalline precipitate fell, which, when washed with cold distilled water, and then redissolved in a large quantity of boiling water, furnished the silver salt of the new acid in the form of thin plates, of a glistening and brilliant appearance. After a second crystallization the salt was dried in the dark *in vacuo* over sulphuric acid, and then submitted to analysis.

*Analysis*.—I. .495 grm. gave .672 grm. carbonic acid and .148 grm. water.

II. .318 grm. left by ignition in a porcelain capsule .132 grm. of metallic silver.

III. .337 grm. left .1409 grm. of silver,

results which correspond in 100 parts to

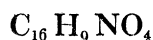
	I.	II.	III.
Carbon . . . .	37.02		
Hydrogen. . . .	3.32		
Silver . . . . .		41.51	41.81.

If we assume that this new acid is formed from toluyl-urea by the elimination of one equivalent of ammonia and by the assumption of the elements of two equivalents of water, the substance thus hypothetically obtained will be represented by the formula



and on comparing the numbers obtained by the analysis of the acid and its silver salt with those calculated from the above formula, we find such an agreement as to place beyond doubt the accuracy of the assumption.

Thus, the formula



requires

Carbon . . . . .	63·57
Hydrogen . . . . .	5·95

and that of  $\text{AgO}$ ,  $\text{C}_{16}\text{H}_8\text{NO}_3$  requires

Carbon . . . . .	37·21
Hydrogen . . . . .	3·10
Silver . . . . .	41·86

The numbers actually obtained from the analysis of the acid and its silver salt are, for the acid,—

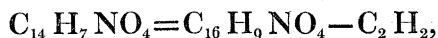
*Mean.*

Carbon . . . . .	63·34
Hydrogen . . . . .	6·18

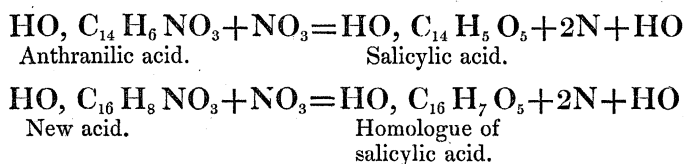
for the silver salt,—

Carbon . . . . .	37·02
Hydrogen . . . . .	3·32
Silver . . . . .	41·66

This acid, for which I propose the provisional name of *Carbo-toluylic acid*, has several representatives in the *benzoyl* series, viz. *anthranilic acid*, obtained by FRITZSCHE by the action of caustic potash on indigo; *benzamic acid*, the product of the action of hydrosulphuric acid on nitrobenzoic acid discovered by ZININ; and, lastly, *carbanilic acid*, prepared by CHANCEL, by treating nitrobenzamide with hydrosulphuric acid. Of these three acids, which are all represented by the formula



*benzamic* and *carbanilic* acids have lately been found by M. GERLAND\* to be identical, while *anthranilic acid* is a body of a different type. It remains to be examined, to which of these two acids, whether to *anthranilic* or to *carbanilic* acids, the new acid above described corresponds. The mode of its generation would render it probable that it is homologous to *carbanilic acid*, whence the provisional name I have assigned to it; should it however prove on further examination to correspond rather to *anthranilic acid*, then the convertibility of this acid into *salicylic acid* by means of nitrous acid, first suggested by Dr. HOFMANN†, and lately experimentally confirmed by M. GERLAND‡, would appear to indicate that the analogous transformation of the new acid would furnish a series of new substances which at present are entirely wanting, namely, the proper homologues of *salicylic acid* and its derivatives. Thus:—



The extreme difficulty however of preparing the different substances described in

\* Quart. Journ. Chem. Soc. vol. v. p. 133.

† Ibid. vol. iii. p. 235.

‡ Ibid. vol. v. p. 133.



this paper, and more especially the costly nature of the hydrocarbon (cymol), which constitutes the (at present) only known source of the parent acid, viz. nitrotoluylic acid, form unfortunate barriers to the continuation of the investigation. *Cymol*, however, doubtless exists among the liquid products of the distillation of coal; indeed, Mr. MANSFIELD, in his valuable memoir on Coal Tar\*, describes a fluid which distils over between 170° and 172° (338° to 342° FAHR.), "which bears so strong a resemblance to cymol in odour, and in other properties, as to induce a belief that this body is identical with the hydrocarbon existing in cumin oil." It is my intention to search for it in this fluid, and should it prove possible to obtain it cheaply and in abundance, the experiment above suggested, and many others of equal scientific interest, will be made, the results of which I shall hope to be allowed the honour of submitting to the Royal Society.

In conclusion, I will give a tabular view of the corresponding members of the benzoyl and toluyll groups, the parallelism of which in the present and previous memoir I have had the pleasure of establishing.

Benzoyl series. HO, C <sub>14</sub> H <sub>5</sub> O <sub>3</sub> . . . . . Benzoic acid.	Toluyll series. HO, C <sub>16</sub> H <sub>7</sub> O <sub>3</sub> Toluylic acid.
HO, C <sub>14</sub> { <sup>H<sub>4</sub></sup> <sub>NO<sub>4</sub></sub> }O <sub>3</sub> . . . . . Nitrobenzoic acid.	HO, C <sub>16</sub> { <sup>H<sub>6</sub></sup> <sub>NO<sub>4</sub></sub> }O <sub>3</sub> Nitrotoluylic acid.
C <sub>14</sub> { <sup>H<sub>4</sub></sup> <sub>NO<sub>4</sub></sub> }O <sub>2</sub> , NH <sub>2</sub> . . . . . Nitrobenzamide.	C <sub>16</sub> { <sup>H<sub>6</sub></sup> <sub>NO<sub>4</sub></sub> }O <sub>2</sub> , NH <sub>2</sub> Nitrotoluyllamide.
C <sub>2</sub> H <sub>3</sub> (C <sub>12</sub> H <sub>5</sub> )N <sub>2</sub> O <sub>2</sub> . . . . . Anilo-urea.	C <sub>2</sub> H <sub>3</sub> (C <sub>14</sub> H <sub>7</sub> )N <sub>2</sub> O <sub>2</sub> Toluyll-urea.
HO, C <sub>14</sub> H <sub>6</sub> NO <sub>3</sub> . . . . . Carbanilic acid.	HO, C <sub>16</sub> H <sub>8</sub> NO <sub>3</sub> Carbo-toluylic acid.

\* Chem. Soc. Quart. Journ. vol. i. p. 244.