UNITED STATES PATENT OFFICE.

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PROCESS OF MAKING BARBITURIC ACIDS.

No. 814,496.

Specification of Letters Patent. .

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To all whom it may concern:

Be it known that I, Otto Wolfes, doctor of philosophy, a subject of the German Emperor, residing at the city of Darmstadt, in the Empire of Germany, have invented certain new and useful Improvements in the Process of Manufacturing Barbituric Acids, of which the following is a description.

My invention relates to the manufacture of barbituric acids; and it consists of certain novel processes particularly pointed out in the concluding claims.

From experiments made by Traube, Conrad, and myself it is known that cyanacetic ester and its substitution products may be condensed with urea, guanidin, and similar bodies to derivates of pyrimidin, one amid of the urea, &c., combining with the cyanogen group of the ester, while the carboxethyl in separating alcohol participates with the second amid of the ring formation. The process is represented by the following equation:

$$\begin{array}{c|c}
x & CN & NH_2 \\
COOR & NH_2 \\
\hline
 & NH \\
\hline
 & NH \\
\hline
 & CO + ROH \\
\hline
 & NH \\
\hline
 & CO + ROH
\end{array}$$

It has been discovered by me that compounds with two cyanogen groups—as, for instance, the malonitrile—may be condensed with urea and with compounds constituted in an analogous way—as, for example, guanidin and the derivatives and homologues of urea and guanidin—if condensing means (viz., alkali metals, their amids, or alcoholates) are employed; but even without the employment of such condensing means the combination is successful if the bodies are permitted to act on each other for a prolonged time at a high temperature or under pressure.

My process as outlined above and em-45 bodied in various forms in the following ex- of crystallization and melts at 240° centigrade 90

amples may be expressed by the following formula:

In the above formula R and R, indicate hydrogen atoms or simple alkyl radicals, and 60 X indicates a bivalent body, such as oxygen and imino, (NH radical.) The bodies obtained in this way may be easily converted into the corresponding barbituric acids by an agent separating ammonia. Such agent may 65 be sulfuric acid, hydrochloric acid, or nitric acid or even alkalis. (the latter, however, being less suitable.)

I will now describe the manner in which I at present prefer to practice my invention, 70 giving, by way of illustration, a number of examples thereof; but it will be understood that various modifications and changes, both as to materials and treatment, may be made without departing from the spirit of my invention and without exceeding the scope of

my claims.

First example: A solution of 9.6 kilograms guanidinhydrochlorate (one molecule) in alcohol is mixed with a solution of 2.3 kilo-80 grams sodium (one molecule) in fifty liters of alcohol, the separated chlorid of sodium drawn off, and the solution of the free guanidin then heated with 12.2 kilograms (one molecule) diethylmalonitrile for five hours 85 to 100° centigrade. After being cooled off the reactionary mass, fine colorless needles, is drawn off and carefully washed with alcohol. The new product contains alcohol of arrestallizations.

while foaming up. It is dissolved easily in water and in diluted acids even in the cold. In cold alcohol it can be dissolved only with difficulty. This triiminodiethyl-barbituric acid (diiminodiethylmalonyl-guanidin) is converted smoothly into diethylbarbituric acid by being heated for three hours to 120° centigrade with diluted hydrochloric acid in a closed receptacle. When cooled off, the product congeals to a crystalline mass. Separate from the mother-liquor, wash with water, and again crystallize from hot water. The thus-purified substance melts at 191° centigrade and has all the properties of the

15 known diethylbarbituric acid. Second example: 12.2 kilograms diethylmalonicacidnitrile are heated with six kilograms urea and 2.3 kilograms sodium in fifty liters of absolute alcohol for four hours in an 20 autoclave to 100° centigrade. After adding six liters of glacial acetic acid the alcohol is evaporated. The residue is heated for an hour with about one hundred liters of water and is then left standing over night. After 25 being dissolved in two molecules of warm diluted hydrochloric acid and being precipitated with ammonia the base crystallizes in tufted aggregates of crystal. It is a 5-diethyl-2 oxy-4, 6-diiminopyrimidin. It 30 melts, while developing basic vapors, at about 272°, (277° corr.) The chlorid is dissolved with more difficulty in water than the sulfate or nitrate. In alkalis the base is soluble, but not in ammonia. From hot water it 35 may be crystallized over again. If boiled with five times the amount of thirty per cent. sulfuric acid, it is converted also into diethylbarbituric acid. The purifying and isolating

o Third example: 9.4 kilograms monoethyl-malonitrile are heated for five hours with an equivalent amount of guanidin in alcoholic solution in an autoclave to 100° centigrade. Lustrous small leaves are separated contain-

ing alcohol of crystallization. After being crystallized over again from alcohol and being dried at 100° centigrade they melt at 189°, (190° corr.) They can be easily dissolved in cold water and hot alcohol. If this compound is heated for several hours in a closed with diluted hydrochloric acid (1:3) to

vessel with diluted hydrochloric acid (1:3) to 120°, it is converted into the monoethylbar-bituric acid. This acid melts at 194° corr., (Conrad and Guthzeit giving 190° corr.) It

55 reacts and tastes rather strongly acid just as the barbituric acid itself. It differs thereby distinctly from the diethylbarbituric acid, which tastes bitter and possesses a reaction which is acid only to a very small degree.

Fourth example: 6.6 kilograms malonitile are heated to 100° centigrade for one hour in an autoclave with an equivalent

amount of guanidin prepared from 9.6 kilograms hydrochlorate and 2.3 kilograms sodium in eighty liters of alcohol. When cooled 65 off, colorless small needles are separated, the amount of which needles is about 6.5 kilograms when dried. The melting-point of this triiminobarbituric acid is 248°, (corr. 252°.) The base forms with diluted cold 7° mineral acid salts, which are dissolved only with difficulty. With nitrate of sodium and glacial acetic acid a light violet nitroso body is formed at once. It has a remarkable resistance against warm strong sulfuric acid or 75 boiling sodium hydroxid. By prolonged and highly heating it with acids barbituric acid is obtained therefrom.

Having thus fully described my invention, what I claim, and desire to secure by Letters 80

Patent, is—
1. The process of manufacturing a barbituric acid consisting in the condensation of a malonitrile of the general formula

with a substance having the general formula

$$X = C \begin{cases} NH_2 \\ NH_2 \end{cases}$$

whereby a 4.6 diimino-pyrimidin of the general formula

$$\begin{array}{c|c}
R & CNH-NH \\
\hline
R_1 & CNH-NH
\end{array}$$

is produced, the latter being subsequently converted by saponification into a 4-6-dioxy-pyrimidin.

2. The process of manufacturing a barbituric acid consisting in condensing a malonitiele with a substance of the general formula

$$X = C \begin{cases} NH_2 \\ NH_2 \end{cases}$$

and converting the iminobarbituric acid thus formed into barbituric acid by saponification.

3. The process consisting in condensing diethylmalonitrile with guanidin with the aid 115 of sodium alcoholate, and subsequently saponifying to produce barbituric acid.

In testimony whereof I have signed my name to this specification in the presence of two subscribing witnesses.

OTTO WOLFES.

Witnesses:

MAX CONRAD, WALTER HOUSING.