The present invention relates to improvements in the production of ammonium nitrate compositions, and is especially applicable where it is required that the composition should be in a form in which the crystals are tightly compacted together, for instance when it is to be formed into compressed masses or into granules produced by comminution to a limited extent of larger compressed forms.

At temperatures ranging from about -18° C. to the transition point occurring at 32.3° C. the stable crystal form of ammonium nitrate is a rhombohedral pyramidal form of specific gravity 1.716, designated by the Roman numeral IV; between the transition point occurring at 32.3° C. and the other transition point occurring at 84° C. the stable crystal form is rhombohedral of specific gravity 1.654, designated by the Roman numeral III, and between the last mentioned transition point and the one occurring at 125° C. the stable crystal form is tetragonal designated by the Roman numeral II.

In the production and storage of compact forms of ammonium nitrate compositions, for example explosive or power gas generating charges, whether produced by casting or by compression, it has always been a matter of difficulty to ensure the maintenance of the external form and internal structure of the compact charge owing to the discontinuous density changes that occur as a result of subjecting the compact charge to temperature changes passing in either direction through the transition point occurring at 32.3° C. Owing to the large difference in specific gravity between the forms III and IV, as a result of such temperature changes the compact ammonium nitrate compositions may crack or crumble internally or may shrink or expand so as to part from or damage some other object which it is required to maintain fixedly in contact with the charge. Such occurrences may lead to serious consequences in connection with the use of the charge: for instance crumbling or cracking increases the burning surface of a power gas generating charge and hence the rate at which pressure is developed by it. The transition of ammonium nitrate occurring at 32.3° C. is facilitated by the presence of moisture.

It is known that in solid solutions of potassium nitrate in ammonium nitrate obtained either by co-crystallisation of solutions made by dissolving the two salts in water, or by solidification of fused mixtures of the two salts, the transition point occurring at 32.3° C. is lowered and the transition point occurring at 84° C. is raised. It has been stated that solid solutions containing more than 8% potassium nitrate prepared by either of the aforesaid methods never become completely converted from form III to form IV however far below 32.3° C. the temperature is reduced.

Campbell and Campbell (Canadian Journal of Research, volume 24B July 1946, pages 93 to 108), who have recently studied this phenomenon, state that solid solutions obtained by either of these two methods can be kept indefinitely at room temperature with ammonium nitrate in form III without undergoing transition into form IV and that the solid solutions show no discontinuous volume change between approximately 1° C. and 100° C. In order to induce the transformation of form III into IV they state that it is necessary to cool the solid solution to a temperature below 1° C., but indicate that once it has taken place the transformation of form IV into form III in a solid solution containing 5% potassium nitrate is brought about by heating it to approximately 20° C. These observers suggest that even when the proportion of potassium nitrate in the solid solution is as high as 10% the transformation of form III into form IV may occur eventually if the solid solution is kept long enough at a sufficiently low temperature, for instance --10° C., particularly if it is moist. These observers further state that in order to lower the transition point below 32.3° C. it is quite insufficient merely to mix the potassium nitrate with the ammonium nitrate, and that the mass must be melted up or crystallised from a mixed aqueous solution of the two salts.

It has also been proposed in British Patent No. 573,147 inter alia to produce ammonium nitrate explosive or propellant compositions showing relatively small density changes when exposed to temperature cycles, by a process comprising fusing together, in the presence of added water, a mixture containing the ammonium nitrate and a minor proportion of potassium nitrate, or evaporating to dryness an aqueous solution of the said mixture, and subjecting the resulting product in either case to such thorough drying conditions that the water content is less than 0.1%.

The production of ammonium nitrate compositions suitable for use, for instance as explosive, propellant and power gas pressure producing charges, smoke compositions, insecticidal and fungicidal vaporizing charges and the like by casting fusion mixtures is of somewhat limited...
applicability, for instance on account of the possibility it affords for ingredients of the wholly or partly molten composition to separate from one another, e.g., by segregation of solid ingredients or of immiscible liquid phases and has the disadvantage of a certain decomposition hazard encountered at the high temperatures. The preparation of solid solutions of potassium nitrate in ammonium nitrate by co-crystallisation from a solution of the two salts is a troublesome operation since it is difficult to obtain stable products of uniform composition and since the solid solution is more difficult to dry off than either of its components.

These disadvantages are avoided by the method employed in the present invention, which is applicable to the manufacture of compositions suitable for use e.g. as explosive propellant, gas pressure, smoke, insecticidal or insect repelling or fungicidal charges, or as fertilizers, which may include other appropriate inorganic and/or organic constituents e.g. trichloroethylene and other volatile bodies, alkali metal chlorates, ammonium bichromate, phosphates, chlorides, sulphates, oxalates, vapoourisable insecticides, fungicides and insect repellents, waxes and so forth.

As a result of our X-ray investigations we have made the remarkable and unexpected discovery that after a simple mixture of powdered potassium nitrate with solid particles of ammonium nitrate has been maintained at a temperature higher than 32.3° C. which may be up to about 100° C., in the absence of any appreciable percentage of moisture the potassium nitrate is effective in stabilising the ammonium nitrate crystal as modification III at temperatures from 32° C. down to at least 0° C. This by maintaining the mixture for a sufficient length of time at a temperature above 32.3° C. the discontinuous density changes ordinarily experienced on passing ammonium nitrate compositions through a temperature cycle passing through 32.3° C. are avoided, so that for instance compressed charges of the compositions will neither crumble nor crack.

According to the present invention the process for the production of compositions comprising ammonium nitrate and potassium nitrate comprises mixing together ammonium nitrate and powdered potassium nitrate and maintaining the mixture thus formed within a temperature range of about 32° C. and about 100° C. for a time long enough to ensure that the ammonium nitrate is stabilised substantially as modification III.

The ammonium nitrate used for mixing with the potassium nitrate may be either in the form of powder or small granules. At least 5% potassium nitrate reckoned on the sum of the ammonium nitrate and potassium nitrate is found to be required to attain stabilisation of the ammonium nitrate in form III, where stability is required at temperatures of the order of -10° C., but quantities say from 1% upwards may be sufficient for practical purposes where such low temperatures are not reached e.g. to prevent the breaking up of granular fertilisers or compressed pellets, under appropriate circumstances. In some cases it is possible to obtain the stabilisation of the ammonium nitrate in form III even if other additional solid components are present during the heat treatment with potassium nitrate e.g. inorganic salts, such as chromates, phosphates; ammonium oxalate, guanidine nitrate or nitroguanidine. In other cases it may be necessary to carry out the potassium nitrate treatment before the other solid components are introduced, e.g. waxes. The progress of the stabilisation action of the potassium nitrate may be followed by continuous or intermittent cycling in compressed pellets made from samples taken, from time to time, from the material being exposed to temperatures in the range between 32° C. and about 100° C. Since this temperature cycling of the compressed mass itself involves exposure of the mixture to a temperature above 32° C. thus tending to complete the action if the time of exposure of the mixture has been insufficient, it is safer to apply a test that does not involve repeated cycling, for instance to determine the temperature-time heating or cooling curve of a sample so as to determine the presence or absence of a transition point, or to plot a volume-temperature curve by observing the density changes by means of a dialtometer so as again to see whether or not a transition point is present. Best of all however is to subject samples taken from time to time to X-ray examination. The X-ray diffraction pattern of ammonium nitrate powder in the form IV differs markedly from that of ammonium nitrate powder in form III, and the presence of for example about 10 per cent of potassium nitrate has been found by us to have no noticeable effect on the respective patterns. The diffraction pattern ultimately attained by the method of our invention is almost indistinguishable from that obtained from a cooled and powdered fusion mixture containing the same proportion of ammonium nitrate and potassium nitrate. By comparison of photographs it is therefore easy to find the minimum time of exposure of the mixture to the temperature, between about 32° C. and about 100° C. beyond which further time of exposure produces no appreciable further change in the X-ray pattern.

At temperatures of 40-50° C. complete stabilisation may be expected to take place in about 10-30 hours depending on the degree of subdivision of the ammonium nitrate and the potassium nitrate and the intimacy of the primary admixture, even if the mixture is agitated only before its exposure to the temperature above about 32° C. The moisture content may also have some effect on the time required, but the reaction proceeds readily even when recently dried ammonium nitrate and potassium nitrate are used, and even in the presence of drying agents such as dehydrated ammonium oxalate. According to a modification of the invention, in place of potassium nitrate there may be used other water soluble potassium salts that react metathetically with an excess of ammonium nitrate to form potassium nitrate; for example, potassium sulphate, potassium chloride, potassium dihydrogen ortho phosphate, potassium oxalate, and potassium chromate. In this case the presence of at least a small amount of moisture during the mixing is advantageous.

When potassium nitrate is used it is desirably in finely powdered condition. The ammonium nitrate may be in ordinary crystal powder condition, but the invention is also applicable for the treatment of ammonium nitrate granules produced by atomisation processes. The compositions produced according to the process of the present invention have been found to be particularly suitable in the production of compressed charges comprising ammonium nitrate and potassium nitrate.
The invention is illustrated by the following examples in which the parts are parts by weight except where otherwise indicated.

**Example I**

The ammonium nitrate was in the form of crystal clusters passing a 60 mesh British Standard Specification screen and contained 0.1-0.2% moisture. 450 parts of this ammonium nitrate were ground together with 50 parts pulverised potassium nitrate having a grit fine enough to pass a 40 mesh B. S. S. screen, until the whole of the product passed through a 100 mesh B. S. S. screen, the mixing being conducted at room temperature. An X-ray powder photograph of the mixture at this stage was the same as that of a sample of ammonium nitrate in form IV, and even after storage at 18°C. of a sample of the mixture for two weeks the X-ray photograph was still substantially the same. The rest of the freshly ground material was heated to 45°C. and stored at that temperature for 18 hours, and then allowed to cool and was kept for a week at room temperature. On examination the X-ray powder photograph was substantially identical with that obtained from a solid solution of potassium nitrate and ammonium nitrate of the same percentage composition prepared by fusion and cooling, and was also substantially identical with that of a sample of ammonium nitrate III. Further heating at 45°C. for 6 hours produced no detectable difference in the X-ray photograph. Pellets of the composition 1:4:4 in diameter compressed under a pressure of 2 tons per sq. in. showed a diametric expansion of only 1.56% after 16 temperature cycles between 0 and 50°C. and did not crumble or crack. Compressed pellets made from a pulverised commercial fusion mixture of the same percentage composition showed about the same diametric expansion when similarly cycled. The pulverulent mixture may be mixed with e.g. potassium chromate and ammonium bichromate and gamma hexachloro-cyclohexane and compressed into pellet form, to give an insecticidal vapour generating pellet which will not crumble with temperature variations on storage, or it may be mixed with trinitrotoluene and pelletted to give an Amatol explosive that will resist volume changes tending to shifter it on alternating hot and cold storage.

**Example II**

The ammonium nitrate used was spray crystallised material passing a 16 mesh and retained on a 25 mesh B. S. S. screen, and had an initial natural bulk density of 0.917 grams per c.c. Four temperature cycles between 0°C. and 50°C. sufficed to reduce the natural bulk density of a sample of this material to 0.778 grams per c.c. and to cause serious cracking and disintegration of the granules. The remainder of the granular material was mixed with 10% of its weight of potassium nitrate, 10% of this product was passed through a 200 mesh B. S. S. screen, after the granules had been moistened with 2% of their weight of a saturated solution of ammonium nitrate in water, these operations being conducted at room temperature. The moistening of the ammonium nitrate granules was intended to cause them to be practically identical with that of a congealed fused mixture of ammonium nitrate and potassium nitrate in the proportions 90:10.

**Example III**

450 parts crystalline ammonium nitrate as used in Example I were ground together with 50 parts potassium oxalate with a grit fine enough to pass a 170 mesh B. S. S. screen until the whole passed through a 100 mesh screen, these operations being conducted at room temperature. The ground mixture was then heated to 80°C. and maintained at that temperature for 24 hours, and a sample was taken. The remainder of the mixture was maintained for another 6 hours at the same temperature and was then cooled, and a further sample was taken. Each of the samples was compressed at 3½ tons per sq. in. into a cylindrical mould of 1/2” diameter. Each pellet was then subjected to 16 temperature cycles between 0-50°C. and it was found that their initial diameters had increased by only 1.3% in each case. X-ray powder photographs of the hot stored mixtures after standing at ordinary temperature for some days showed that the ammonium nitrate had been stabilised in the III form.

The pulverulent composition may be used as a constituent of compressed gas pressure producing charges containing say potassium chromate or ammonium bichromate as sensitisier of thermal decomposition as an added ingredient.

**Example IV**

865 parts crystalline ammonium nitrate as used in Example I were ground together with 60 parts of potassium nitrate of a grit fine enough to pass a 170 mesh B. S. S. screen and 75 parts anhydrous ammonium oxalate of the same grit as the potassium nitrate, until the mixture all passed a 100 mesh screen. These operations were conducted at room temperature. The ground mixture was then heated to 45°C. and maintained at that temperature for 18 hours. After cooling and standing for 4 to 5 days at 15-17°C. the X-ray diffraction pattern was found to be practically identical with that of a congealed fused mixture of ammonium nitrate and potassium nitrate in the proportions 90:10. The product may be used for all purposes similar to that indicated for the product of Example III, which it resembles in enabling a compressed pellet of the gas pressure producing
charge to be produced which will resist the usual disruptive effect of temperature cycling.

We claim:

1. A process for the production of a composition comprising ammonium nitrate and potassium nitrate which comprises mixing together ammonium nitrate and powered potassium nitrate and maintaining the mixture thus formed within a temperature range of about 32° C. and about 100° C. for a time long enough to ensure that the ammonium nitrate is stabilized substantially as modification III which is that modification having a rhombic crystal form, a specific gravity of 1.654 and stable between the transition points normally occurring at 32.5° and 84° C.

2. A process as claimed in claim 1 wherein the ammonium nitrate used for mixing with the potassium nitrate is in the form of powder.

3. A process as claimed in claim 1 wherein the ammonium nitrate used for mixing with the potassium nitrate is in the form of small granules.

4. A process as claimed in claim 1 wherein at least 8 per cent potassium nitrate is present reckoned on the sum of the ammonium nitrate and potassium nitrate where stability of the ammonium nitrate in modification III is required at temperatures of the order of —10° C.

5. A process as claimed in claim 1 wherein the mixture of ammonium nitrate and potassium nitrate maintained within a temperature range of about 32° C. and about 100° C. for the time aforementioned includes at least one additional solid component.

6. A process as claimed in claim 5 wherein the additional solid component is an inorganic salt.

7. A process as claimed in claim 6 wherein the inorganic salt is selected from the group consisting of chromates and phosphates.

8. A modification of the process as claimed in claim 1 wherein a water-soluble potassium salt other than potassium nitrate is used which reacts metaethetically with an excess of ammonium nitrate to form potassium nitrate.

9. A process as claimed in claim 8 wherein the water-soluble potassium salt is selected from the group consisting of potassium sulphate, potassium chloride, potassium dihydrogen orthophosphate, potassium oxalate, and potassium chromate.

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JOHN WHITSTONE.

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