Optimal Parameters for Obtaining Diammonium Phosphate from off-balance Phosphate Raw Materials of the Karatau Basin

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Abstract: The article contains information about substandard material resources of phosphorite raw materials of the Republic of Kazakhstan. Samples of phosphate-siliceous raw materials of the Zhanatas Deposit were studied for the content of P$_2$O$_5$. To obtain diammonium phosphate (DAP), off-balance phosphorite from the Zhanatas Deposit was used, containing P$_2$O$_5$-16.18%. Technological solutions for the enrichment of substandard phosphate raw materials "Zhanatas" and the production of DAP from evaporated and non-evaporated WPA are proposed. Information on mineral petrographic analysis and the chemistry of the DAP production process based on the results of X-Ray Diffraction, X-Ray Phase Analysis, and SEM is provided.

Keywords: off-balance phosphorites, Karatau basin, diammonium phosphate, WPA

1. Introduction

In market conditions, part of the reserves of mineral raw materials is uncompetitive, and leads to a reduced quality of raw materials, and due to increased economic costs. Mineral resources of any state are the material basis for economic development. The mineral resource base of the Republic of Kazakhstan, by the richness of its mineral resources and their diversity, is included in the group of world leaders [1]. Along with the decrease in the quality of mineral raw materials, there is a sufficient replenishment of extracted minerals [1, 2]. This leads to an excess of their consumption over the increase, problematic issues of providing existing enterprises with proven reserves and the feasibility of organizing new production facilities [2, 3].

Since the middle of the last century, it has become necessary to enrich low-grade phosphorite ores by removing impurities to increase the content of macro and micro components that affect the quality of the fertilizer obtained from them and improve their suitability for processing [4-6].

The mineralogical composition of phosphate raw materials largely depends on the origin of the rock. Unlike apatite rocks of igneous origin, phosphorites are sedimentary rocks that were formed close to the earth's surface at low pressure and temperature, which had a great influence on the chemical composition and variety of impurities in the ore rock [7-9].

Currently, the production of phosphorites by the open method is widely developed all over the world. However, underground mining methods are also used [5, 10]. One of the largest deposits of the Karatau basin, which provides it with phosphate raw materials in an underground way, is the Chulaktau Deposit, where phosphor ore is extracted from the ground and mining operations are carried out at a depth of 360 m below ground level [5, 11].

The concentrated ore horizon consists of phosphorites and phosphorite conglomerates, phosphorite and siliceous shales, and phosphorite-bearing dolomites containing layers of phosphoritic juices. There are two phosphorite horizons with an average thickness of the lower layer of 7.5 m, and the upper one of 18 m [7, 11]. Both of these layers are separated by an 18 m thick phosphorite-shale layer. The structure of phosphorite is a crystalline, latent crystalline, granular, oolitic composition [10-12].

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The growing demand for industrial mineral raw materials, against the background of declining quality and depletion of material and energy resources, requires a comprehensive development of the mineral resource base, the search for new large and rich in the target component of deposits, the involvement in the exploitation of non-traditional types of minerals, the attraction of new technologies for processing low-quality ores with the solution of environmental problems [4].

Current global trends in the mineral resource sector of the economy are characterized by widespread depletion of profitable mineral reserves due to significant production volumes characterized by more complex mining and geological conditions for conducting exploration and development of deposits and their corresponding rise in price [3, 4].

Kazakhstan has a strong mineral resource base of the phosphorous industry, which is able to meet the country’s domestic needs and exports for the long term. It is based on the reserves of micrograin phosphorites of the Karatau phosphorite basin, which are characterized by a relatively high content of phosphorus pentoxide (24%) [2, 5]. The disadvantages of these ores include their very difficult enrichability and the inaccessibility of most of their reserves for open-pit mining [5]. The low quality of the obtained concentrates (25-27% P₂O₅ at 35-40% for the main world producers) makes the products of Kazakhstani producers unclaimed on the world market [5, 6]. As a result, the main problem that needs to be solved is the creation of modern, more efficient technologies for ore enrichment not only of micrograin phosphorites, but also of nodular phosphorites from the Chilisai Deposit in Western Kazakhstan [1, 5]. This will allow us to organize the production of high-quality phosphate raw materials for the production of phosphorus-containing fertilizers that meet international standards and ensure export to the countries of the Commonwealth of Independent States, as well as to the markets of Eastern Europe and Asia [5, 7].

The most common phosphorus fertilizer in the world - diammonium phosphate (DAP, diammophos), contains both nitrogen and phosphorus, the popularity of which is associated with ease of use, a relatively high content of active substance (N+P₂O₅) with good physical properties and the absence of nitrates, as well as chlorine [2, 7]. DAP is produced in a single technological process by neutralizing wet-process phosphoric acid (WPA) with an excess of ammonium. DAP is especially useful for sugar-containing fruits and root crops, since phosphorus in its composition saves the accumulation of a fairly high concentration of sugar, which improves the quality of fruit and vegetable products [5-7].

DAP has a wide range of applications. In addition to acidic soils, it is also used on other types of arable land. Depending on the composition of the fertile layer, its dosage is selected individually for each type of soil [2, 6]. Additional application of nitrogen is carried out to equalize the ratio of components to the equilibrium state. There are no special restrictions on the timing of DAP application. It is applied both during spring plowing operations as a pre-sowing fertilizer, and throughout the growing season of various plant groups [6, 8]. One of the most important positive qualities of DAP is its versatility and low price. The use of DAP is economically feasible, since one ton of this fertilizer can replace at least 2.5-3 tons of simple superphosphate [2, 8].

The purpose of this work was to study the composition and properties of WPA obtained from low-quality phosphate-siliceous raw materials, studies were conducted to determine the mineral, petrographic and chemical compositions of a representative sample of phosphate-siliceous off-balance raw materials from the Zhanatas deposit.

2. Materials and methods

To obtain DAP from low-quality phosphate-siliceous raw materials, studies were conducted to determine the mineral, petrographic and chemical compositions of a representative sample of phosphate-siliceous off-balance raw materials from the Zhanatas deposit.

The data of the chemical composition analysis and the applicable regulatory and technical documentation for research are shown in Table 1.
Table 1. Chemical composition of the Zhanatas Deposit phosphate-siliceous off-balance raw materials

| Name of the defined characteristics, units of measurement | The norm according to the Normative document | Actual values based on test results | Normative documents for test methods |
|----------------------------------------------------------|---------------------------------------------|-----------------------------------|------------------------------------|
| Mass fraction of boron, %                                 | -                                           | 0.0366                            | -                                  |
| Mass fraction of potassium, %                            | -                                           | 0.51                              | GOST 5382-91                       |
| Mass fraction of calcium, %                              | -                                           | 3.65                              | GOST 4013-82                       |
| Mass fraction of magnesium, %                            | -                                           | 1.305                             | GOST 5382-91                       |
| Mass fraction of manganese, %                            | -                                           | 0.0942                            | GOST 5382-91                       |
| Mass fraction of arsenic, %                              | -                                           | 0.018                             | GOST 23581-9-79                    |
| Mass fraction of phosphorus, %                           | -                                           | 7.5                               | GOST 5382-91                       |
| Mass fraction of fluorine, %                             | -                                           | 0.004                             | GOST 5382-91                       |
| Mass fraction of phosphorus oxide, %                     | -                                           | 17.4                              | GOST 5382-91                       |
| Mass fraction of potassium oxide, %                      | -                                           | 0.156                             | GOST 5382-91                       |
| Mass fraction of magnesium oxide, %                      | -                                           | 4.76                              | GOST 5382-91                       |
| Insoluble residue, %                                     | -                                           | 30.3                              | GOST 5382-91                       |

Mineral petrographic analysis of off-balance phosphate-siliceous shales of the Zhanatas deposit was carried out in transmitted light using MIN – 8 and MIN – 9 polarization microscopes.

Samples of off-balance Zhanatas macroscopic phosphorites are more uniform and are represented by black carbonaceous shales. Under the microscope, the rock has an oolitic structure. In transmitted light, oolites are black due to the presence of a significant amount of organic matter. Quartz grains are slightly metamorphosed and rounded.

The oolites and the cementing phase are co-ordinated with francolite. In the form of borders, fibrous chalcedony is marked. There are rolled fragments of feldspars and leaves of mica (sericite). Organic substances mainly compose the central part of oolites, and they are less common in the cementing mass. The sample contains cubic pyrite crystals, around which iron hydroxides develop, their content is less than 1%. The sample contains the following minerals: phosphorite-47.9%, dolomite-12.25%, calcite-5.04%, quartz-22.8%.

Figure 1. Phosphorite Zhanatas:
1-quartz; 2-chalcedony; 3-calcite;
4-francolite; 5-pyrite

Figure 2. Phosphorite Zhanatas:
1-quartz; 2-chalcedony; 3-calcite;
4-francolite; 5-pyrite

The studies on IR-Fourier spectrometer spectra obtained in the spectral range 4000-300 cm\(^{-1}\) when using drugs in the form of tablets prepared by pressing 2 mg of sample and 200 mg KBr. When studying the phosphate-siliceous raw materials of the Zhanatas Deposit, it was found that it contains in its composition (Figure 3):
- Fluorapatite Ca\(_5\)(PO\(_4\))\(_3\)F– 1094, 1045, 965, 603, 577, 570, 471 cm\(^{-1}\) [13, 16];
- Quartz SiO$_2$ – 798, 779, 694, 521, 471, 396 cm$^{-1}$ [14, 15];
- Dolomite CaMg[CO$_3$]$_2$ – 1454, 880, 729 cm$^{-1}$ [14];
- Dolomite CaMg[CO$_3$]$_2$ – 1432, 880, 729 cm$^{-1}$ [16];

In addition, as shown in figure 3, IR spectral analysis has established the possibility of the presence of:
- Calcite CaCO$_3$– 1432, 714 cm$^{-1}$ [14, 16];
- Siderite FeCO$_3$– 1432, 865 cm$^{-1}$ [14].

![Figure 3. IR spectral analysis of the Zhanatas Deposit phosphate-siliceous off-balance raw materials](image)

The research was carried out on a D8 Advance (Bruker) diffractometer, $\alpha$-Cu. Processing of the obtained diffractogram data and calculation of interplane distances were performed using the EVA software. Sample decoding and phase search were performed using the Search/match program using the PDF-2 powder diffractometric database. Analyses have established that the studied phosphate-siliceous raw materials contain (Figure 4).

![Figure 4. X-ray phase analysis of the Zhanatas Deposit phosphate-siliceous off-balance raw materials](image)

During the experiments, a production acid of the following composition was used: P$_2$O$_5$-21%, F-1.64%, SO$_3$-2.28%, CaO-0.2%, Fe$_2$O$_3$-0.53%, Al$_2$O$_3$ -0.19%, suspensions -2.28% Na$_2$O-0.19%. Return dry sodium salts obtained as follows: the precipitated silica precipitate from H$_3$PO$_4$ is decomposed with sulfuric acid, then after decomposition is treated with water and the resulting solutions are evaporated to dry sodium salts. The content of Na$_2$O in the dry salts is equal to 40.4%.
Na$_2$SiF$_6$ is precipitated from acid heated to 343 K. After the salt is introduced, the contents are stirred for a specified time and cooled to 298 K.

The acid was evaporated in a Wurtz flask, which was placed in a water bath. The discharge in the system was recorded by a pressure gauge. At the end of the experiment, the acid and sediment are weighed. The precipitate is washed with alcohol and analyzed by X-ray analysis and qualitative analysis for P$_2$O$_5$ and SO$_3$. The evaporated acid is analyzed for the content of P$_2$O$_5$, SO$_3$, F, Na, with the determination of its density and viscosity.

The effect of the amount of sodium salts on the degree of acid defluorination and on the size of Na$_2$SiF$_6$ crystals, the method of WPA cooling, and the mixing time of the reagents were studied.

a) Influence of the rate of consumption of sodium salts

When determining the effect of the salt norm in the range of 100 to 200% of the stoichiometry, the salts were introduced for 4 min with continuous stirring, then the resulting pulp was stirred for another 5 min, and the acid was slowly cooled to 298 K [9, 12]. According to the results shown in Table 2, it can be seen that at a rate of 100%, the degree of desulfurization is 77%.

The introduction of a small excess of precipitator (the norm of 120-140%) sharply increases the degree of acid defluorination by 6.5%. Further treatment of the norm does not lead to a significant effect [17].

| Precipitation rate, % | Residual content, % | Degree of defluorination, % | The average size of the crystals, micron |
|-----------------------|---------------------|-----------------------------|----------------------------------------|
| 100                   | 0.382               | 77.0                        | 6.5                                    |
| 120                   | 0.252               | 84.5                        | 18.0                                   |
| 140                   | 0.241               | 85.3                        | 15.0                                   |
| 160                   | 0.232               | 86.0                        | 9.0                                    |
| 200                   | 0.212               | 87.3                        | 7.6                                    |

An excess of a small amount of sedimentation agent also has a positive effect on the average size of crystals. The largest crystals correspond to the norm of salts of 120-140% (0.3-0.35 sodium per 100 kg of acid) [12, 18]. It should be noted that at these standards, individual crystals have a size of up to 22-23 microns.

b) Influence of the cooling method.

The experiments were carried out at a rate of consumption of sodium salts of 120-140%, changing the conditions for the introduction of the precipitator and the method of cooling acid solutions after the introduction of the precipitator. As expected, a sharp cooling of the acid, after the introduction of the precipitator, contributes to the formation of small crystals (Table 3). The largest crystals [8, 9, 12] Na$_2$SiF$_6$ fall out after precipitation for 4 min, followed by mixing for 5 minutes and slow cooling to 90-100 min. At the same time, the crystal sizes in the range of the norm of sodium salts of 120-140% are equal to 18 and 15 microns, respectively [18].

The effect of the flow rate, injection conditions, and cooling method of sodium salts on the degree of desfluorination and the size of Na$_2$SiF$_6$ crystals is shown in Table 3.

| Process parameter | Norm of sodium salts | Content of F, % | Degree of defluorination, % | The average size of crystals, micron | Content of F, % | Degree of defluorination, % | The average size of crystals, micron |
|-------------------|----------------------|----------------|-----------------------------|-----------------------------------|----------------|-----------------------------|-----------------------------------|
| Salt injection - 4 min | Slow cooling (90-100 min) | 0.232 | 84.5 | 18.0 | 0.24 | 85.3 | 15.0 |
| Mixing - 5 min | Cooling abrupt (10 min) | 0.31 | 81.0 | 9.0 | 0.28 | 83.0 | 7.8 |
From the analysis of Table 3, it can be seen that the degree of defluorination of the precipitator introduction conditions and the method of acid cooling does not have a significant effect [17, 22]. However, the growth of Na$_2$SiF$_6$ crystals is affected by both the time of introduction of the precipitator and the method of cooling the pulp. The best conditions for removing fluoride from an acid are:

- rate of consumption of sodium salts 120%;
- time of introduction of the precipitator 4 min;
- slow mixing time of the reaction pulp is 90-10 min, at which the degree of defluorination is 83-85.3%, and the crystal size is about 18 micron.

To obtain complex fertilizers and double superphosphate by the chamber method from Karatau phosphorites, WPA of 40-50% P$_2$O$_5$ concentration is required [9, 21]. This concentration can be achieved by evaporation of WPA [2, 9]. Since WPA is contaminated with various admixtures of metals, the process of evaporation is complicated by precipitation on heating surfaces of precipitation consisting mainly of calcium sulfate, silicofluorides, iron phosphates, aluminum, etc. Evaporation of WPA from Karatau phosphorites due to the presence of Mg and F in it, as mentioned above, is possible only up to a concentration of 38-39% P$_2$O$_5$.

The authors [4, 12] conducted studies on the evaporation of pre-defluorination WPA from Karatau phosphorites, from a concentration of 26-52% P$_2$O$_5$. The following parameters are set as optimal conditions: the system discharge is 400 mmHg and the temperature is 353-363 K.

### Table 4. Influence of temperature and time of the process of WPA defluorination with aqueous solutions of sodium salts

| T, K | T, min | The composition of the defluorinated WPA with lye F, Na$_2$O, K$_2$O % | Degree of defluorination, % | The average size of crystals, micron |
|------|--------|------------------------------------------------|-----------------------------|-----------------------------------|
| 298  | 10     | 0.24, 0.0835, 0.539 | 49.3 | 32.4 |
| 298  | 30     | 0.26, 0.0895, 0.604 | 53.6 | 38.2 |
| 298  | 45     | 0.27, 0.0933, 0.607 | 51.3 | 31.5 |
| 298  | 60     | 0.25, 0.0860, 0.512 | 46.7 | 34.8 |
| 313  | 10     | 0.28, 0.096, 0.607 | 54.6 | 38.3 |
| 313  | 30     | 0.285, 0.097, 0.604 | 52.3 | 36.5 |
| 313  | 45     | 0.29, 0.099, 0.608 | 50.3 | 34.8 |
| 313  | 60     | 0.30, 0.103, 0.610 | 48.3 | 33.3 |
| 323  | 10     | 0.32, 0.108, 0.600 | 52.3 | 36.5 |
| 323  | 30     | 0.34, 0.115, 0.604 | 50.3 | 34.8 |
| 323  | 45     | 0.342, 0.116, 0.610 | 49.0 | 33.3 |
| 323  | 60     | 0.33, 0.114, 0.613 | 47.0 | 32.4 |
| 333  | 10     | 0.348, 0.118, 0.604 | 50.0 | 34.8 |
| 333  | 30     | 0.36, 0.120, 0.609 | 47.6 | 33.3 |
| 333  | 45     | 0.366, 0.122, 0.619 | 46.5 | 32.4 |
| 333  | 60     | 0.37, 0.125, 0.620 | 45.6 | 31.8 |
| 343  | 10     | 0.388, 0.134, 0.604 | 46.3 | 31.3 |
| 343  | 30     | 0.39, 0.134, 0.614 | 45.6 | 30.8 |
| 343  | 45     | 0.393, 0.135, 0.615 | 45.3 | 30.3 |
| 343  | 60     | 0.394, 0.136, 0.618 | 44.3 | 30.0 |
Table 5. Changes in the content of P$_2$O$_5$, F, and Na$_2$O as a function of temperature at a time of 30 min and an excess of 140% sodium salts from stoichiometry

| Content of components in acid, % | 323 | 328 | 333 | 338 | 343 | 348 | 353 | 358 | 363 | 368 |
|---------------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| P$_2$O$_5$                      |     |     |     |     |     |     |     |     |     |     |
| 18.0                            | 18.2| 18.5| 18.9| 19.0| 19.2| 19.3| 19.4| 19.6| 19.8|     |
| 18.6                            | 18.7| 18.8| 19.1| 19.2| 19.3| 19.4| 19.5| 19.8| 19.9|     |
| 19.1                            | 19.2| 19.3| 19.5| 19.7| 19.8| 19.9| 20.0| 20.1| 20.2|     |
| 19.5                            | 19.5| 19.7| 19.9| 20.1| 20.1| 20.4| 20.4| 20.6| 20.7|     |
| 19.9                            | 20.1| 20.2| 20.4| 20.5| 20.6| 20.7| 20.7| 20.9| 21.1|     |
| F                               |     |     |     |     |     |     |     |     |     |     |
| 0.19                            | 0.20| 0.21| 0.21| 0.22| 0.23| 0.23| 0.24| 0.25| 0.25| 0.26 |
| 0.195                           | 0.21| 0.22| 0.23| 0.23| 0.24| 0.24| 0.25| 0.26|     |     |
| 0.20                            | 0.21| 0.215| 0.229| 0.236| 0.236| 0.242| 0.25| 0.261| 0.269|     |
| 0.21                            | 0.215| 0.217| 0.238| 0.239| 0.240| 0.244| 0.253| 0.263| 0.272|     |
| 0.22                            | 0.218| 0.219| 0.230| 0.232| 0.243| 0.247| 0.259| 0.267| 0.275|     |
| Na$_2$O                         |     |     |     |     |     |     |     |     |     |     |
| 0.71                            | 0.73| 0.74| 0.75| 0.76| 0.77| 0.79| 0.79| 0.81| 0.82|     |
| 0.72                            | 0.74| 0.74| 0.75| 0.76| 0.78| 0.79| 0.79| 0.80| 0.81| 0.83 |
| 0.72                            | 0.75| 0.76| 0.76| 0.78| 0.79| 0.80| 0.81| 0.83| 0.83|     |
| 0.73                            | 0.75| 0.77| 0.77| 0.79| 0.80| 0.81| 0.82| 0.83| 0.84|     |

For experiments, an acid of the following composition was used (Table 6), obtained from phosphate-siliceous raw materials of the Zhanatas Deposit.

Table 6. Physico-chemical characteristics of WPA

| Type of WPA     | P$_2$O$_5$ | SO$_3$ | F | Na | CaO | MgO | viscosity | density |
|-----------------|------------|--------|---|----|-----|-----|-----------|---------|
| Non-defluorination WPA | 23.4 | 2.56 | 1.91 | 0.36 | 0.20 | 1.25 | 1.00 | 1.379 |
| Defluorination WPA    | 23.4 | 4.25 | 0.47 | 0.77 | 0.25 | 1.40 | 1.09 | 1.395 |

Precipitation occurs when both non-defluorination and defluorination WPA is evaporated. When evaporating defluorination WPA, the amount of sediment is much greater than when evaporating defluorination WPA, as can be seen from Figure 5.

Figure 5. The dependence of the precipitation, the concentration of WPA: 1- defluorination WPA; 2- non-defluorination WPA

Figure 6. Changes in the viscosity of the acid depending on the concentration of P$_2$O$_5$: 1- defluorination WPA; 2- non-defluorination WPA
Analyzing the obtained results with the studies given in [22], in which WPA is obtained from Khibinsk apatite concentrate by the semi-hydrate method, it is possible to state the comparability of the obtained values. The difference is that in our case, WPA with a lower content of 23.4% $P_2O_5$, obtained from off-balance phosphorites of the Zhanatas deposit, is used by the dihydrate method.

The determination of the viscosity of cold acids is very difficult, which may be due to the precipitation of newly formed precipitation when the acids are cooled. Especially large crystals and a large number of them fall out when using evaporated non-defluorinated WPA [18, 21]. Despite the high viscosity of the evaporated defluorinated WPA, which is probably due to an excess of sodium salts for defluorinated, the acid is mobile in the hot state [21]. The density of the acid solutions as non-defluorinated and defluorinated about the same. It is known that their concentration increases monotonically as shown in Figure 7.

Figure 8 shows data on changes in the content of sulfates, sodium, and fluorine in WPA during its evaporation.

![Figure 7. Change in the density of WPA when it is evaporated: 1- defluorination WPA; 2- non-defluorination WPA](image1)

![Figure 8. Change of $SO_3$ in acid during its evaporation: 1- defluorination WPA; 2- non-defluorination WPA](image2)

![Figure 9. Change of F and Na in acid during its evaporation: 1- defluorination WPA; 2- non-defluorination WPA](image3)
Decomposition of phosphorus-containing off-balance raw materials with a mixture of aqueous solutions of sulfuric and phosphoric acids, using the dihydrate method of WPA production, is carried out by the reaction:

$$\text{Ca}_5(\text{PO}_4)_3 \text{F} + 5\text{H}_2\text{SO}_4 + n\text{H}_3\text{PO}_4 + \text{aq} = (n + 3)\text{H}_3\text{PO}_4 \times 2\text{H}_2\text{O} + \text{HF} + \text{aq}$$

Carbonates of off-balance phosphate raw materials decompose similarly, with the release of CO$_2$ into the gas phase. The main amount of CO$_2$ is released due to the dissolution of dolomite and calcite by sulfuric and phosphoric acids [4, 19, 24]:

$$\text{CaMg}((\text{CO}_3)_2 \times 2\text{H}_2\text{SO}_4 + \text{aq} = \text{CaSO}_4 \times 2\text{H}_2\text{O} + \text{Mg SO}_4 + 3\text{CO}_2 + \text{aq}$$

The process of dissolution of carbonates is one of the fastest in the decomposition of off-balance raw materials and leads to the formation of foam. The magnesium contained in the raw material is completely dissolved into the form of magnesium sulfate and phosphate in WPA [19]. In addition, the magnesium goes to WPA so upon dissolution of the silicate MgSiO$_4$:

$$\text{Mg}_2\text{SiO}_4 + 2\text{H}_2\text{SO}_4 = 2\text{MgSO}_4 + \text{SiO}_2 + 2\text{H}_2\text{O}$$

Similarly, calcium silicate decomposes. When silicates are dissolved, silicon dioxide passes into the WPA [19, 23]. When aluminum silicates are dissolved in WPA, SiO$_2$ and ions of sodium, potassium, aluminum, iron, and others are released. It should be noted that the impurities of aluminum and iron contained in off-balance phosphate raw materials are almost completely transferred to WPA. Iron and aluminum compounds in WPA from off-balance phosphorites react with phosphate ions by the reaction:

$$\text{R}_2\text{O}_3 + 2\text{H}_3\text{PO}_4 = 2\text{RPO}_4 + 3\text{H}_2\text{O}$$

The solubility of iron phosphates in WPA is relatively low, which can lead to the formation of supersaturated solutions of iron phosphate. In the presence of 2-3% SO$_4^{2-}$ supersaturated WPA containing iron phosphates are stable for a long time. Precipitation of iron phosphates from the WPA is possible during long-term storage of phosphoric acid at low temperatures [20].

The solubility of aluminum phosphates in WPA is much higher than that of iron phosphates. Because of this and their relatively low content in raw materials, aluminum phosphates are usually not deposited in the production of WPA.

Part of the hydrogen fluoride released during the decomposition of raw materials reacts with silicic acid to form a gaseous SiF$_4$ by the reaction [18, 19]:

$$\text{SiO}_2 + 4\text{HF} = \text{SiF}_4 + 2\text{H}_2\text{O}$$

The other part reacts with aluminum compounds to form an AlF$_{3-x}$ type of WPA, where $x = 1 \div 6$.

In the dihydrate process, 5-12% of the fluorine coming from the raw material is removed in gaseous form from the reaction system for absorption purification [20, 21]. The amount of fluoride gases released depends on the temperature of the process, the concentration of sulfuric acid and the concentration of WPA, and increases with their increase. When the gases are cooled, SiF$_4$ undergoes partial hydrolysis, with the formation of a SiO$_2$ precipitate:

$$3\text{SiF}_4 + n\text{H}_2\text{O} = \text{SiO}_2 + n\text{H}_2\text{O} + 2\text{H}_2\text{SiF}_6$$

The release of silica gel can become a problem for the operation of absorption equipment, leading to clogging, and therefore the absorption treatment scheme is organized in such a way that at the initial stage
of absorption, with a high degree of irrigation, the main amount of fluoride gases is absorbed in the flues and the floor absorber. HF and SiF₄ present in WPA form fluoroaluminaic acid by reaction:

\[
2\text{HF} + \text{SiF}_4 = \text{H}_2\text{SiF}_6
\]

Fluoroaluminaic acid interacting with Na⁺ and K⁺ ions forms poorly soluble sodium and potassium silicofluorides by reaction:

\[
(\text{Na, K})_2\text{SO}_4 + \text{H}_2\text{SiF}_6 = (\text{Na, K})_2\text{SiF}_6 + \text{H}_2\text{SO}_4
\]

Which are capable of forming stable supersaturated solutions that lead to deposits on the walls of filtration equipment pipelines, especially at a sharp temperature drop.

The physical and chemical processes that occur during the preparation of DAP are determined primarily by the reactions of phosphoric acid neutralization. Changes in the composition of the ammonium phosphate pulp during ammonization affect the following technological parameters [19, 20]:
- solubility and digestibility of the reaction products;
- rheological properties of the pulp;
- partial pressures NH₃ and F above the pulp.

During the ammonization process, when the pH increases to 2.5, water-soluble NH₄H₂PO₄, NH₄HSO₄, (NH₄)₂SiF₆ [20, 25] are formed by the reactions:

\[
\begin{align*}
\text{H}_2\text{PO}_4 + \text{NH}_3 &= \text{NH}_4\text{H}_2\text{PO}_4 \\
\text{H}_2\text{SO}_4 + \text{NH}_3 + \text{NH}_4\text{H}_2\text{PO}_4 &= \text{NH}_4\text{HSO}_4 + \text{NH}_4\text{H}_2\text{PO}_4 \\
\text{H}_2\text{SiF}_6 + 2\text{NH}_3 &= (\text{NH}_4)_2\text{SiF}_6
\end{align*}
\]

In addition, citrate-soluble iron and ammonium compounds are formed:

\[
(\text{Fe, Al})_3(\text{H}_3\text{O})\text{H}_8(\text{PO}_4)_6 6\text{H}_2\text{O} + 9\text{NH}_3 = (\text{Fe, Al})_3\text{NH}_4\text{H}_2\text{PO}_4 6\text{H}_2\text{O} + 9\text{NH}_3
\]

When neutralizing a magnesium-containing WPA, it also forms:

\[
(\text{Fe, Al})_3(\text{H}_3\text{O})\text{H}_8(\text{PO}_4)_6 6\text{H}_2\text{O} + 9\text{NH}_3 + \text{H}_2\text{SiF}_6 = 3\text{Mg(Fe, Al)NH}_4(\text{HPO}_4)_2\text{F}_2 + 6\text{NH}_4\text{H}_2\text{PO}_4 + \text{SiO}_2 + 5\text{H}_2\text{O}
\]

It is noted that the compound (Fe, Al)₃NH₄H₂PO₄₆·x·H₂O crystallizes well, is easily filtered and separated [19, 24]. The compound (Fe, Al)NH₄HPO₄F₂ is amorphous and forms colloidal non-settling and poorly filtered sediments [19, 25]. The Mg(Fe, Al)NH₄(HPO₄)₂F₂ salt is least soluble and well crystallized [19, 20, 25].

\[
(\text{Fe, Al})_3\text{NH}_4\text{H}_8(\text{PO}_4)_6 6\text{H}_2\text{O} + (\text{NH}_4)_2\text{SiF}_6 + 3\text{NH}_3 = 3\text{NH}_4\text{H}_2\text{PO}_4 + 3(\text{Fe, Al})\text{NH}_4\text{HPO}_4\text{F}_2 + \text{SiO}_2 + 4\text{H}_2\text{O}
\]

\[
6\text{Mg(Fe, Al)NH}_4(\text{HPO}_4)_2\text{F}_2 + (\text{NH}_4)_2\text{SiF}_6 + 4\text{NH}_3 + 2\text{H}_2\text{O} = 6\text{Mg(Fe, Al)(NH}_4)_2(\text{HPO}_4)_2\text{F}_2 + \text{SiO}_2
\]

On the basis of the above reactions occurring during ammonization and neutralization, larger crystals are formed that allow for a more softened filtration regime of the resulting product.

### 4. Conclusions

Comparing the results obtained for phosphoric acid defluorination with solutions of sodium salts and dry return sodium salts, it is clear that the degree of H₃PO₄ defluorination at the same rate of consumption of sodium salts and other equal conditions is almost the same and is about 84%. When using aqueous solutions of sodium salts to precipitate fluoride from acid, the concentration of P₂O₅ in the defluorination acid decreases by 2-2.5%, while the use of dry sodium salts almost does not affect the concentration of P₂O₅ in the final acid. However, the sizes of the Na₂SiF₆ crystals that fall out are not the same. The
largest crystals of 50-53 microns are observed when phosphoric acid is defluorinated with aqueous solutions of sodium salts. When fluorine is deposited with dry sodium salts, the crystals do not exceed 18-20 microns. During the evaporation of defluorinated WPA the amount of precipitating sludge in 3.5-4 times less than the evaporation acid. Values viscosity and density non-defluorination and defluorination WPA at the same concentration of P₂O₅ is about the same.

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