Topochemical Transformations in Sodium-Bismuth-Silicate System at 100–900 °C

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Abstract. The authors have developed a method for producing highly dispersed sillenite bismuth silicate in the system Na₂O-Bi₂O₃-SiO₂ (NBS) from water solutions of organosilicon monomers (sodium methylsiliconate) and bismuth nitrate. The paper studies the phase composition and microstructure of the synthesized NBS material at different temperatures. The morphology of crystals in the NBS material and the peculiarities of its thermal-oxidative breakdown are investigated. X-ray diffraction spectra obtained using a CuKα-source are used to evaluate the crystal lattice spacing and to analyze the broadening of the maximum-intensity diffraction line for this crystal with due consideration of crystal indices h, k, l by the approximation method to determine the dimensions of the coherent scattering region and microdistortions of the crystal lattice Δa/a. The authors established that the silicate shell on Bi₁₂SiO₂₀ particles is close to the silicates with continuous chain radicals [SiO₃]∞²⁻, and a part of them are bridges between the bismuth silicate particles.

Keywords: Sillenite · Phase composition · Microstructure · Crystal morphology · Thermal treatment · Crystal lattice microdeformation

1 Introduction

The development of highly dispersed metal-organosiloxane fillers with modified surface allows solving a multitude of important problems in the field of radiation materials science (Pasechnik 2006). The promising approach is to use water-soluble chemically active organosiloxanes as the basis for production of metal oligomers. A new technological approach to the solution of the stated complex problem is required.

As of today, the chemistry of organosiloxane compounds of bismuth attract particular attention. This is conditioned by multiple valuable properties of organosilicon compounds (high thermal stability, hydrophobicity, dielectric characteristics and resistance to a range of aggressive media). Besides, bismuth atoms have large capture cross-section of gamma-radiation, which is almost the same as for lead atoms in a wide energy spectrum. The presence of vacant 3d-orbitals in silicon atoms conditions high reactivity of bond ≡Si-OH in silicate minerals.
2 Materials and Methods

The authors have developed a method for producing highly dispersed sillinite bismuth silicate in system Na₂O-Bi₂O₃-SiO₂ from water solutions of organosilicon monomers (sodium methylsiliconate) and bismuth nitrate (Yastrebinskii et al. 2018).

The amounts of the components were calculated with the aim of producing stable bismuth silicate Bi₁₂SiO₂₀ (6Bi₂O₃ · SiO₂).

At 100 °C we have obtained highly dispersed (0.2–0.3 μm) hydrophobic NBS material that is insoluble in water (NBS wetting angle is 122°). The density is 3780 kg/m³.

According to mass-spectroscopy, NBS material had the following composition (expressed as oxides), wt%: Na₂O - 23.83; Bi₂O₃ - 59.70; SiO₂ - 16.47.

Differential thermal analysis (DTA) and thermogravimetric analysis (TGA) of specimens were performed on a STA-449 F1 Jupiter derivatograph (Germany). X-ray diffraction (XRD) analysis of phases and structure was performed on a ARL™ X’TRA Powder Diffractometer (Switzerland) with Cu kx source (λ_kx = 1.542 Å) using a nickel filter. The infrared spectra were obtained on a Specord-75IR spectrometer (Germany). The material microstructure was studied by raster electron microscopy (REM) in the modes of reflected (back-scattered) and secondary electrons.

3 Results and Discussion

Phase Composition and Microstructure of Mineral Phases in NBS Material Synthesized at 100 °C. XRD phase analysis using the Powder Diffraction File (PDF) and literature data (Gorshkov and Timashev 1981; Gorelik et al. 2002) allowed detecting the formation of three amorphous-crystalline mineral phases:

1. Metastable bismuth silicate Bi₂SiO₅ (d = 3.0379 Å/l = 100%; 3.7169 Å; 2.7223 Å) with tetragonal crystal system (a = 3.802; c = 15.134 Å), with the amorphous ring of about 3 Å.

2. Bismuth oxide α-Bi₂O₃ (d = 3.2596 Å/l = 100%; 3.2596 Å; 1.9625 Å) with monoclinic crystal system (a = 5.8499; b = 8.1698; c = 7.5123 Å) with the amorphous ring of about 3 Å.

3. Bismuth organosilicate H₃C(SixBi₅Oz)Na with the amorphous ring of 10–12 10–12 Å and clear X-ray reflection at d = 11.4513 and 5.7090 Å. However, the precise determination of this composition using PDF failed.

Fig. 1. IR spectrum of synthesized NBS material
The results of IR-spectroscopy, the silicate phases in NBS powder synthesized at 100 °C have linear structure. The splitting of the absorption bands in the range of 1000–1100 cm\(^{-1}\) that is typical for the siloxane bond indicates the presence of several types of siloxane phases (Fig. 1).

**Morphology of Crystals in Synthesized NBS Material.** According to REM, NBS material synthesized at 100 °C contained particle agglomerations of irregular shape with the size of 0.8–2.5 μm (Fig. 2).

**Defectiveness of Crystals in NBS Material Subjected to Thermal Treatment.** X-ray diffraction spectra obtained using a CuK\(_{α}\)-source were used to evaluate the crystal lattice spacing and to analyze the broadening of the maximum-intensity diffraction line for this crystal with due consideration of crystal indices h, k, l by the approximation method to determine the dimensions of the coherent scattering region and microdistortions of the crystal lattice Δa/a.

At 100–300 °C XRD analysis detected amorphous-crystalline bismuth organosilicate \(\text{H}_3\text{C}(\text{Si}_x\text{Bi}_y\text{O}_z)\text{Na}\) with the amorphous ring of 10–12 Å and clear X-ray reflection at \(d = 11.4810\) Å and 5.7020 Å.

At the temperature of 200 °C, bismuth silicate \(\text{Bi}_2\text{SiO}_5\) in the mixture of minerals in terms of X-ray parameters approaches to the benchmark silicate of this composition (card no. 36-288 PDF: \(d = 3.0400\) Å (I = 100%, hkl = 103)).

In the temperature interval of 300–400 °C, metastable bismuth silicate \(\text{Bi}_{12}\text{Si}_{0.87}\text{O}_{20}\) with cubic crystal lattice in the synthesized dry mix transforms into stable bismuth silicate \(\text{Bi}_{12}\text{SiO}_{20}\) also with cubic lattice.

In the temperature interval of 300–500 °C, the density of dislocations in the structure of bismuth silicate \(\text{Bi}_{12}\text{SiO}_{20}\) crystals was lowering, while at the temperature higher than 650 °C, it was conversely rising up. The increase of the temperature from 300 to 500 °C improves the parameters of elementary crystal lattice of bismuth silicate \(\text{Bi}_{12}\text{SiO}_{20}\) by 0.0357 Å and 0.0268 Å, as compared to the benchmark crystal. The volume of elementary crystal cell in this temperature interval grows by 2% and amounts to 1040.5870 Å\(^3\).
4 Conclusion

In the study, a method for producing highly dispersed sillenite bismuth silicate in the system Na$_2$O-Bi$_2$O$_3$-SiO$_2$ (NBS) from water solutions of organosilicon monomers (sodium methylsiliconate) and bismuth nitrate was developed. The paper studied the phase composition and microstructure of the synthesized NBS material at different temperatures. The crystalline structure of the substance and the presence of silicate amorphous phase in it were discovered. The paper revealed the morphology of crystals in the synthesized NBS material and the peculiarities of its thermal-oxidative breakdown; the silicate shell on the particles of Bi$_{12}$SiO$_{20}$ was close to continuous chain radicals [SiO$_3$]$_{3/2}^-$ and a part of them played the role of bridges between bismuth silicate particles. The determination of physicochemical properties of modified Bi$_{12}$SiO$_{20}$ sillenite crystals was of appreciable significance.

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