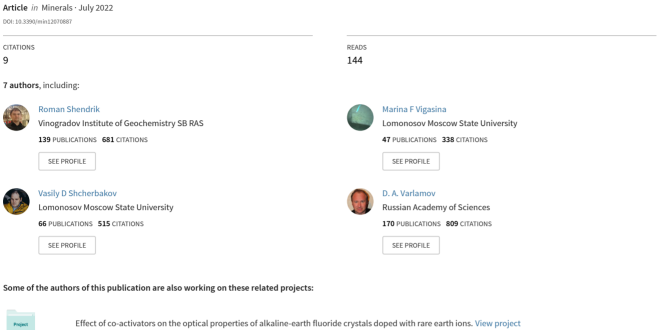
Crystal Chemistry, Isomorphism, and Thermal Conversions of Extra-Framework Components in Sodalite-Group Minerals







Изучение петрологии зон термального воздействия гранитоидных интрузий на породы гранулитовых комплексов на примере Центральной Зоны Лапландского гранулитового комплекса. View project





Article

Crystal Chemistry, Isomorphism, and Thermal Conversions of Extra-Framework Components in Sodalite-Group Minerals

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Abstract: Isomorphic substitutions of extra-framework components in sodalite-group aluminosilicate minerals and their thermal conversions have been investigated using infrared, Raman, electron spin resonance (ESR), as well as ultraviolet, visible and near infrared (UV–Vis–near IR) absorption spectroscopy methods and involving chemical and X-ray diffraction data. Sodalite-related minerals from gem lazurite deposits (haüyne, lazurite, and slyudyankaite) are characterized by wide variations in S-bearing extra-framework components including SO₄²⁻ and various polysulfide groups (S₂•-, S₃•-, S₄•- radical anions, and S₄ and S₆ neutral molecules) as well as the presence of CO₂ molecules. Heating at 700 °C under reducing conditions results in the transformation of initial S-bearing groups SO₄²⁻ and S₃•- to a mixture of S²⁻, HS⁻, S₂•-, and S₄•- and transformation of CO₂ to a mixture of CO₃²⁻ and C₂O₄²⁻ or HC₂O₄- anionic groups. Further heating at 800 °C in air results in the decomposition of carbonate and oxalate groups, restoration of the SO₄²⁻ and S₃•- groups, and a sharp transformation of the framework. The HS⁻ anion is stable only under reducing conditions, whereas the S₃•- radical anion is the most stable polysulfide group. The HS⁻-dominant sodalite-group mineral sapozhnikovite forms a wide solid-solution series with sodalite. The conditions required for the formation of HS⁻- and CO₂0-bearing sodalite-group minerals are discussed.

Keywords: sodalite group; isomorphism; solid solutions; infrared spectroscopy; electron (UV-Vis) spectroscopy; Raman spectroscopy; electron spin resonance; photoluminescence

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1. Introduction

Minerals belonging to the sodalite group are microporous cubic or pseudo-cubic alumino- and beryllosilicates which occur in different kinds of alkaline magmatic and metasomatic rocks. Some of them (sodalite, haüyne, and nosean) belong to important rock-forming minerals.

The unit cell parameters of the "idealized sodalite (**SOD**-type) framework" deposited in the Database of Zeolite Structures are a = 8.9561 Å; V = 718.4 Å³; and space group Im-3m (aristotype) [1,2]. However, due to different schemes of the Si/Al ordering as well as specific features of the occupation of the intra-framework cavities (sodalite cages) by extra-framework components, the observed symmetry is usually lower. A distinctive feature of sodalite-type compounds is a three-dimensional system of channels consisting of sodalite cages and running along different directions. The extra-framework constituents in sodalite-group minerals are more diverse than in the members of the related

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cancrinite group and include different cations (Na+, [N(CH₃)₄]+, Ca²⁺, Mn²⁺, Fe²⁺, and Zn²⁺), anions (Cl⁻, F⁻, OH⁻, HS⁻, S²⁻, and SO₄²⁻), radical anions (S₃•-), and neutral molecules (H₂O, CO₂, S₆) as species-defining components. Some subordinate, admixed components (K⁺, H⁺, S₂•-, S₄•-, SO₄•2-, S₃, S₄, MoO₄²⁻, WO₄²⁻, AsO₄3-, and COS) were identified in sodalite-group minerals using a multianalytical approach involving different spectroscopic methods [3–6].

To date, many dozens of chemically different microporous compounds with the SOD-type framework have been synthesized. Interest in such compounds is caused by their specific properties, which make it possible to consider them as advanced materials with technologically important properties. The hydrothermal method was applied to obtain sodalite-type compounds with aluminosilicate frameworks as well as frameworks containing W-, Mo-, Ga-, Be-, Ge-, P-, or As-centered tetrahedra, with a wide variety of extra-framework cations and anions [7]. Microporous compounds belonging to the topological type of sodalite are advanced materials that can be used as pigments, sorbents for water purification and gas sorption, matrices for immobilization of radioactive isotopes and heavy metals, hydrogen and methane storage, superconductors, catalysts, membranes for separation of gases and desalination of seawater, etc. [8–37].

This paper provides new data on the isomorphism of extra-framework components in some sodalite-group minerals as well as mechanisms for their thermal conversions.

2. Materials and Methods

The studied samples are listed below. Their empirical formulae are partly taken from literature sources. Chemical data for newly analyzed samples are given in Table 1. Samples 1–6 and 9 were collected at the Malo-Bystrinskoe gem lazurite deposit, Baikal Lake area, Siberia, Russia.

Sample 1 is bluish lilac S₄-bearing haüyne. The mineral forms granular aggregate (4 cm across) embedded in calciphyre. The associated minerals are calcite, diopside, and pyrite. The charge-balanced empirical formula is (Table 1; the contents of CO₂ and HS-were estimated from the IR spectrum; for the identification of H₂S and S₄, see below): Na_{6.24}Ca_{1.49}K_{0.05}(Si_{5.99}Al_{6.01}O₂₄)(SO₄)_{1.50}(H₂S)_{0.075}(S₄)_{0.09}Cl_{0.10}F_{0.24}(CO₂)_{0.16}·nH₂O (Z = 1).

Sample 2 is deep blue haüyne. It forms single-crystal grains up to 1 cm across in association with calcite, phlogopite, and accessory sphalerite, Zn-bearing spinel, and grossular. The empirical formula of Sample 2 is (Table 1): Na_{6.28}Ca_{1.47}K_{0.17}(Si_{6.05}Al_{5.92}Fe_{0.03}O₂₄)(SO₄²⁻)_{1.68}S⁰_{0.40}Cl_{0.06}(CO₂)_{0.06}·nH₂O, where S⁰ is all sulfide sulfur provided that it occurs in neutral molecules.

Sample 3 is lilac haüyne forming a rim around light gray fine-grained aggregates of earlier silicates up to 2 cm across. The associated minerals are lazurite, sodalite, diopside, and calcite. The empirical formula is Na_{6.39}K_{0.06}Ca_{1.57}(Si_{6.08}Al_{5.92}O₂₄)(SO₄)_{1.78}(S₄)_{0.03}(S²⁻)_{0.02}Cl_{0.07}(CO₂)_{0.15}·nH₂O (Z = 1) [4]. The mineral is cubic, with the unit cell parameter a = 9.076 Å.

Sample 4 is light blue SO₃•-bearing haüyne with the empirical formula Na_{6.45}K_{0.03}Ca_{1.35}(Si_{6.07}Al_{5.93}O₂₄)(SO₄)_{1.35}(SO₃•-)_{1.35}S₂•-0.02Cl_{0.16}(CO₂)_{0.02}·nH₂O (Z = 1) [4]. The unit cell parameter is a = 9.067 Å. The mineral forms grains up to 5 mm in calciphyre, in association with pyrite.

Sample 5 is bright blue haüyne with the empirical formula Na_{6.45}K_{0.01}Ca_{1.36}(Si_{6.06}Al_{5.94}O₂₄)(SO₄)_{1.58}S⁰_{0.43}Cl_{0.09}(CO₂)_{0.02}·nH₂O, where S⁰ is all sulfide sulfur provided that it occurs in neutral molecules. The unit cell parameter is a = 9.071 Å. The mineral forms grains in coarse-grained calciphyre consisting mainly of calcite, with subordinate diopside.

Sample is the holotype specimen slyudyankaite of Na₂₈Ca₄(Si₂₄Al₂₄O₉₆)(SO₄)₆(S₆)_{1/3}(CO₂)·2H₂O [38]. The mineral is triclinic, space group: *P*1, *a* = 9.0523 (4) Å, b = 12.8806 (6) Å, c = 25.681 (1) Å, α = 89.988(2)°, β = 90.052(1)°, γ = 90.221(1)°, 2994.4 ų. The empirical and V(2) $Na27.57Ca4.05K0.11(Si24.52Al23.48O96)(SO4)6.06S^02.42Cl0.12(CO2)1.43 \cdot 2.21H2O$ (Z = 1), where $S^02.42$ is toMinerals 2022, 12, 887 3 of 31

tal sulfide sulfur, mainly occurring as S₆ and subordinate S₄ neutral molecules, according to the structural data. The color of slyudyankaite is nonuniform: from blue-green in the major part of grains to pink and yellow in thin peripheral zones. The mineral forms isolated anhedral equant grains up to 0.5 cm across. The associated minerals are diopside, calcite, fluorapatite, phlogopite, lazurite, and pyrite.

Sample 7 is gray nosean forming crystals up to 0.5 cm across and elongate twins on (111) up to 1 cm long in cavities of nosean sanidinite. The sample originates from the In den Dellen (Zieglowski) pumice quarry, 1.5 km NE of Mendig, Laach Lake (Laacher See) paleovolcano, Eifel region, Rhineland-Palatinate, Germany. The associated minerals are sanidine, augite, annite, and zircon. The empirical formula of Sample 7 is [3]: $(H_3O)_xNa_{5.72}K_{0.64}Ca_{0.33}(Si_{6.43}Al_{5.51}Fe_{0.06}O_{24})(SO_4)_{1.21}Cl_{0.11}F_{0.09}(CO_2)_{0.11}\cdot nH_2O$ (Z=1).

Sample 8 is the holotype specimen of sapozhnikovite Nas(Al₆Si₆O₂₄)(HS)₂ from a hydrothermally altered urtite-like rock at Karnasurt Mountain, Lovozero alkaline massif, Kola Peninsula, Russia [39]. The mineral forms isolated colourless to pale grayish anhedral equant grains up to 0.5 cm across. The empirical formula is Na_{7.73}Al_{6.08}Si_{5.97}O₂₄(HS)_{1.25}Cl_{0.60}·0.16H₂O (Z = 1). Sapozhnikovite is cubic, $P\bar{4}3n$, with a = 8.91462(7) Å.

Sample 9 is the neotype specimen of lazurite with the empirical formula $(Na_{6.97}Ca_{0.88}K_{0.10})_{7.96}(Si_{6.04}Al_{5.96}O_{24})(SO_4)_{1.09}(S_3^{\bullet}-)_{0.55}S^{2}-_{0.05}Cl_{0.04}\cdot 0.72H_2O$ [5]. The associated minerals are calcite and diopside as well as accessory dolomite, forsterite, and phlogopite. The crystal structure of Sample 9 is characterized by commensurate and incommensurate modulations; the *a* parameter of the cubic sub-cell is equal to 9.087(3) Å.

Sample 10 is the holotype specimen of the recently IMA-approved new F-dominant sodalite-group mineral bolotinaite (IMA 2021-088). The mineral occurs as isolated yellow elongate interpenetration twins up to 1 mm long in cavities of a volcanic ejectum of trachitoid sanidinite collected in the In den Dellen pumice quarry, Eifel region, Rhineland-Palatinate, Germany. The associated minerals are sanidine, nepheline, annite, and zircon. The empirical formula of Sample 10 is (Na5.92K0.82Ca0.10H0.08)(Si6.33Al5.67O24)(SO4)0.17F0.84Cl0.16(CO2)0.38·3.96H2O.

Sample 11 is blue sodalite forming grains up to 3 cm across in a cancrinite-sodalite pegmatite uncovered at the western slope of Mount Kobelikha, Vishnevye Gory, Vishnevogorskiy syenite-miaskite complex, South Urals, Russia. The associated minerals are potassic feldspar, cancrinite, annite, muscovite, different zeolites, as well as accessory zircon, thorite, ilmenite, columbite-(Fe), Nb-bearing rutile, a pyrochlore-group mineral, various sulfides, etc. The composition of Sample 10 is close to the end-member formula Nas(Si₆Al₆O₂₄)Cl₂ [3].

Sample 12 was collected from the same area as sapozhnikovite (Sample 8), north slope of Karnasurt Mountain located in the northern part of the Lovozero alkaline massif. This sample includes several similar but not identical to each other pieces of hydrothermally altered highly agpaitic urtite-like rocks containing rock-forming minerals of the sodalite–sapozhnikovite solid-solution series. Their share in a rock varies from 5 to 20 vol.%. Other rock-forming minerals are nepheline (the major constituent: > 60 vol.%), aegirine, albite, potassic feldspar, natrolite, and kyanoxalite in different proportions; accessory minerals are murmanite (sometimes with lomonosovite relics), fluorapatite, and loparite-(Ce). Sodalite–sapozhnikovite series minerals form white or colourless, with strong orange fluorescence in longwave UV light (λ = 330 nm), grains up to 1 cm across. The representative chemical compositions of these minerals are given in Table 1.

Chemical analyses of haüyne Samples 1 and 2 (Table 1) were carried out using a Tescan VEGA-II XMU INCA Energy 450 (TESCAN, Brno, Czech Republic) (EDS mode, 20 kV, 120–150 pA, beam size 120 nm, excitation zone < 5 μ m). The following standards were used: CaF₂ for F, albite for Na, synthetic Al₂O₃ for Al, wollastonite for Ca, potassium feldspar for K, SiO₂ for Si, Fe metal for Fe, and FeS₂ for S. Chemical analyses of sodalite–sapozhnikovite series minerals (Sample 12, Table 1) were carried out using a Jeol JSM-6480LV scanning electron microscope equipped with an INCA-Wave 500 wave-

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length-dispersive spectrometer (Laboratory of Analytical Techniques of High Spatial Resolution, Department of Petrology, Geological Faculty, Moscow State University), with an acceleration voltage of 20 kV, a beam current of 10 nA, and a 10 μ m beam diameter. The standards used are as follows: jadeite for Na, Al and Si, FeS₂ for Fe, ZnS for S, and NaCl for Cl. Contents of other elements with atomic numbers > 6 are below detection limits.

Additionally, chemical compositions of 55 samples of minerals belonging to the sodalite–sapozhnikovite solution series were investigated.

Table 1. Chemical composition (wt.%) of haüyne from the Malo-Bystrinskoe gem lazurite deposit, Baikal Lake area, Siberia, Russia obtained in this work. Representative chemical analyses of sodalite–sapozhnikovite series minerals (Sample 12) from Karnasurt Mt., Lovozero massif, Kola Peninsula, Russia obtained in this work.

Sar	1			2					
		Bluish lilac			Deep blue				
		18.04			17.92				
	K ₂ O			0.21			0.74		
	CaO			7.83			7.60		
	Al ₂ O ₃			28.63			27.79		
	Fe ₂ O ₃			0			0.19		
	SiO ₂			33.65			33.46		
	CO ₂ a			0.67			0.25		
	SO ₃ b			14.58			15.33		
	F			0.42			0		
	Cl			0.32			0.31		
_	O=Cl,F			-0.25			-0.07		
	Total			104.10		103.52			
Analysis No.	1	2	3	4	5	6	7	8	
			Cont	ents, wt. %					
Na ₂ O	23.60	24.20	24.60	24.28	24.49	24.34	24.01	24.21	
Al ₂ O ₃	30.84	31.50	31.37	31.38	31.69	31.29	31.16	31.33	
Fe ₂ O ₃	0.36	-	0.27	-	-	0.13	-	-	
SiO ₂	36.03	36.88	36.95	36.61	36.82	36.50	36.48	36.26	
HS*	2.78	3.10	3.52	3.74	4.17	4.48	4.86	5.27	
Cl	3.97	3.41	3.32	3.07	2.68	2.31	1.85	1.35	
-O=(Cl,HS)	-1.57	-1.52	-1.60	-1.59	-1.61	-1.60	-1.60	-1.58	
Total	96.01	97.57	98.43	97.49	98.24	97.45	96.76	96.84	
	Form	ula Calculated	on the Basis of	Al + Fe + Si = 1	2 Atoms Per F	ormula Unit			
Na	7.56	7.61	7.72	7.68	7.68	7.71	7.56	7.70	
Al	6.00	6.02	5.99	6.03	6.04	6.02	6.02	6.05	
Fe	0.04	-	0.03	-	-	0.02	-	-	
Si	5.96	5.98	5.98	5.97	5.96	5.96	5.98	5.95	
S	0.84	0.91	1.03	1.11	1.22	1.33	1.45	1.57	
Cl	1.11	1.06	0.91	0.85	0.73	0.64	0.51	0.38	
S+Cl	1.95	1.97	1.94	1.96	1.95	1.97	1.96	1.95	

^a CO₂ contents corresponding to CO₂ molecules were determined from the IR spectra using a procedure described in [3]; ^b All sulfur is given as SO₃, which results in high total sums for the samples bearing sulfide groups. * Recalculated from values for S measured by electron microprobe. Dash means Fe content below detection limit. Analyses are ordered by increase of the S: Cl ratio.

The diffuse absorption spectra in the near infrared, visible, and ultraviolet (NIR/Vis/UV) ranges were measured at room temperature using a Lambda 950 spectro-photometer (Perkin-Elmer, Shelton, CT, USA). In particular, the absorption spectra of platelets of Sample 1 (about 1 mm thick) were measured in the transmission mode through a circular diaphragm 0.8 mm in diameter. The spectra were measured in an integrating sphere at room temperature. To do this, small crystals of the studied mineral were placed in a quartz test tube, transparent in the region of 250–2000 nm, and the test tube was placed in an integrating sphere.

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The ESR spectra were measured with a RE-1306 X-band spectrometer (KBST, Smolensk, Russia) with a frequency of 9.3841 GHz at room temperature and 9.1841 GHz at 77 K. For low-temperature measurements, a quartz ampoule with the sample was placed in a flooded cryostat.

Heating of Samples 3 and 4 was carried out according to the following scheme. The samples were placed in a muffle furnace in air and sequentially heated every 50 °C for 10 min. After holding at each of the temperatures, ESR and NIR/Vis/UV diffuse absorption spectra were measured.

Thermal conversions of other studied samples were carried out in two stages. In the first stage, the samples were heated for three days at 700 $^{\circ}$ C in reducing atmosphere (over the Fe-FeS buffer). Thereafter, parts of the preheated samples were annealed at 800 $^{\circ}$ C in air for 24–36 h.

The powder X-ray diffraction study was carried out at room temperature with an automatic powder diffractometer (D8 ADVANCE, Bruker, Germany) equipped with a Göbel mirror. The powder X-ray diffraction patterns were obtained in step scan mode (in the 2θ range of 10 to 80°) using CuK α radiation at an accelerating voltage of 40 kV, with a current of 40 mA, time per step of 1 s, and a 2θ step of 0.02° . Calculations of interplanar distances were performed using the software that was delivered with the diffractometer (DIFFRAC Plus Evaluation package EVA, Bruker AXS, Bruker, Germany). Unit cell parameters were refined using the (440), (611), (622), and (721) reflections.

3. Results

3.1. Isomorphism of Extra-Framework Components in Cubic Sodalite-Group Minerals

Raman spectra are very sensitive to anionic groups and neutral molecules containing sulfide sulfur. Representative Raman spectra of cubic sodalite-group minerals are given in Figures 1–3.

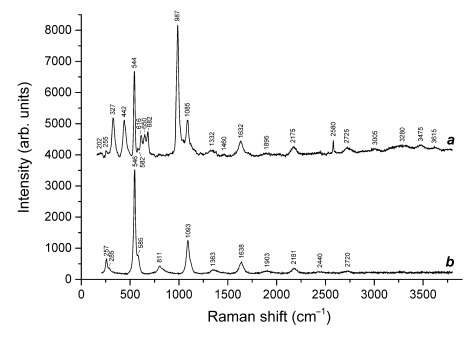


Figure 1. Raman spectra of (a) S4-bearing haüyne (Sample 1) and (b) lazurite (Sample 9).

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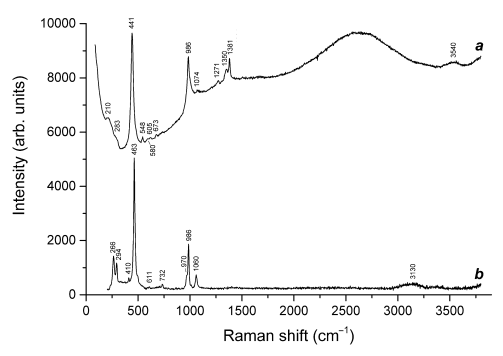


Figure 2. Raman spectra of (a) bolotinaite (Sample 10) and (b) sodalite (Sample 11).

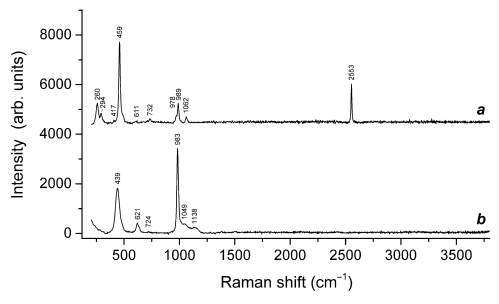


Figure 3. Raman spectra of (a) sapozhnikovite (Sample 8) and (b) nosean (Sample 7).

The assignment of bands in the Raman spectra of cubic sodalite-group minerals shown in Figures 1–3 (Table 2) was made in accordance with [3,4,38–50].

Table 2. Assignment of Raman bands of cubic sodalite-group minerals.

		Raman S	hift (cm ⁻¹)			<u></u>		
		Samp	ole No.			Assignment		
1	7	8	9	10	11			
202w	_	-	-	210	_	Combination of low-frequency lattice modes and/or trans-S4 bending		
								mode
255	-		257	-	-	S ₃ •- bending mode (v ₂)		
-	-	260	-	-	266	Bending vibrations of the [ClNa ₄] ³⁺ and [(HS)Na ₄] ³⁺ clusters		
	294	204 205	285w	202	_	Combination of low-frequency lattice modes involving Na+ cations		
		263W	5w 283w		and/or S₀ bending mode			

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327	-	-	-	-	-	cis-S ₄ mixed (bending + stretching) v ₄ mode
	-	417w	-	-	410w	Bending vibrations of the aluminosilicate framework
442	439		-	441s	-	SO_4^{2-} [the $E(v_2)$ mode] and/or $\delta[O-Si(Al)-O]$ bending vibrations
	-	459s	-	-	463s	Stretching vibrations of the [ClNa ₄] and [(HS)Na ₄] clusters
544s	-	-	546s	548w	-	S ₃ •- symmetric stretching (v ₁) mode
582w			585			S ₃ •- antisymmetric stretching (v ₃), possibly, overlapping with the
J02W			363			stretching band of S ₂ *-
_	_	_	_	580w,	_	S ₂ •- stretching mode
				605w		
	-	611w	-	-	611w	Overtone of vibrations involving Na ⁺ cations?
616	621	-	-	-	-	SO_4^{2-} bending vibrations [$F_2(v_4)$ mode]
650	-	-	-	-	-	gauche-S4 symmetric stretching A1(v1) mode
682	-	-	-	673w	-	trans-S₄ symmetric stretching v₃ mode
	724w	732w	-	-	732w	Mixed vibrations of the aluminosilicate framework
987s	983s	978, 989	-	986s	970, 986	SO_4^{2-} symmetric stretching vibrations [$A_1(v_1)$ mode]
1053w	1049	1062			1060	Stretching vibrations of the framework and/or CO ₃ ²⁻ symmetric
1055W	03W 1049 1062	-	_	1060	stretching vibrations	
	-	-	-	1074	-	HF libration
1085	-	-	1093s	-	-	$S_3^{\bullet-}$ overtone $(2 \times v_1)$
	1138		_	_	_	SO_4^{2-} asymmetric stretching vibrations [$F_2(v_3)$ mode], possibly, over-
	1136	-	-	-	-	lapping with $S_2^{\bullet-}$ overtone (2 × v_1)
	-	-	-	1271	-	CO ₂ Fermi resonance
1332	-	-	-	-	-	Overtone of the cis-S ₄ antisymmetric stretching mode ($2 \times v_3$)
	-	-	-	1350	-	H ⁺ translation
	-	-	1363	-	-	$S_3^{\bullet-}$ combination mode $(2v_1 + v_2)$
	-	-	-	1381	-	CO ₂ Fermi resonance
1460w	-	-	-	-	-	CO ₃ asymmetric stretching mode
1632	-	-	1638	-	-	S₃•- overtone (3 × v₁)
1895	-	-	-	-	-	S ₃ •- combination mode $(3 \times v_2 + v_1)$
2175	-	-	2181	-	-	$S_3^{\bullet-}$ overtone $(4 \times v_1)$
-	-	-	2440w	-	-	S ₃ •- combination mode $(4 \times v_2 + v_1)$
	-	2553	-	-	-	HS⁻ stretching mode
2580	-	_	-	-	-	H ₂ S symmetric stretching mode
2725	-	-	2720	-	-	S₃•- overtone (5 × v₁)
3005, 3280	-	-	-	-	3130	H₃O+ stretching mode
3475, 3615	-	-	-	3540	-	H ₂ O stretching vibrations

Except for a very weak band at 285 cm⁻¹ (combination of low–frequency lattice modes involving Na⁺ cations), all bands in the Raman spectrum of lazurite (Sample 9: curve b in Figure 1) correspond to the S3^{•-} radical anion [5]. Weak bands of sulfate groups are not observed in the Raman spectrum of Sample 9 because of their overlapping with the bands of S3^{•-}.

Analogous bands of the trisulfide radical anion are present in the spectrum of bluish lilac haüyne (Sample 1), which contains trace amounts of S3 $^{\bullet-}$ (curve a in Figure 1). Additionally, in the Raman spectrum of Sample 1, bands of SO $^{42-}$ groups are observed at 987 cm $^{-1}$ [symmetric stretching vibrations, $A_1(v_1)$ mode], 616 cm $^{-1}$ [bending vibrations, $F_2(v_4)$ mode], and 442 cm $^{-1}$ [bending vibrations, the $E(v_2)$ mode]. A series of distinct Raman bands (at 327, 650, and 682 cm $^{-1}$) corresponding to S4 molecules having cis-, gahche- and trans-conformations, respectively, as well as bands of carbonate groups (at 1953 and 1460 cm $^{-1}$), H_2S molecules (at 2580 cm $^{-1}$), and hydronium cations H_3O^+ (at 3005 and 3280 cm $^{-1}$) (Table 2) are the specific feature of Sample 1 distinguishing it from all other haüyne samples studied in this work and elsewhere [1,2,48]. The presence of hydronium in Sample 1 is in agreement with the deficit of extra-framework cations determined by the electron microprobe analysis (Table 1).

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Ideally, sodalite, bolotinaite and sapozhnikovite are the end-members of the solid-solution system with the general formula $[Na_7(\square,Na)](Si_6Al_6O_{24})\{Cl,HS,[F_{0.5}(H_2O)_2]\}_2$. However, natural samples of these minerals contain various extra-framework impurities $(K, H_3O^+, SO_4^{2-}, S_2^{\bullet-}, CO_2, S_4, HF, etc.)$ The Raman spectrum of sapozhnikovite in the range of 200–2500 cm⁻¹ is very close to the spectrum of typical sodalite (Figures 2 and 3). The only significant distinctive feature of sapozhnikovite is a strong Raman band at 2553 cm⁻¹ corresponding to stretching vibrations of HS⁻ anions. Bolotinaite is the member of this solid-solution system most enriched in impurities (Table 1). In particular, it contains $S_2^{\bullet-}$, which is the cause of the yellow color of this mineral and its strong luminescence under laser beams (Figure 2).

In the NIR/Vis/UV absorption spectrum of Sample 1, strong bands with the maxima at 0.65, 0.88, 2.05, and 2.38 eV (respectively, 1900, 1400, 605, and 520 nm) as well as absorption growth above 2.7 eV (460 nm) and a weak band at 1.33 eV (920 nm) are observed (Figure 4). The ESR spectrum of Sample 1 contains a sextet of Mn²⁺ ions in the low-spin state (S = $\frac{1}{2}$ [51]) with g = 2.001, a strong single band with g = 2.002, and a weak band with g = 2.036 (Figure 5). Most probably, Mn²⁺ ions occur in calcite forming inclusions in haüyne individuals.

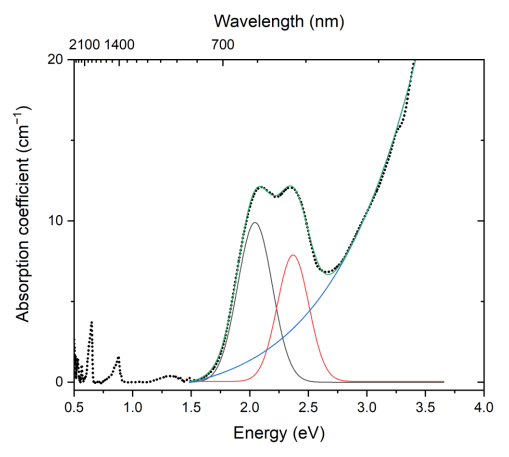


Figure 4. Absorption spectrum of haüyne (Sample 1, dotted curve). The black and red solid lines show the deconvolution of the doublet in the region of 2.0–2.5 eV into two peaks, and the total decomposition curve is shown with green color.

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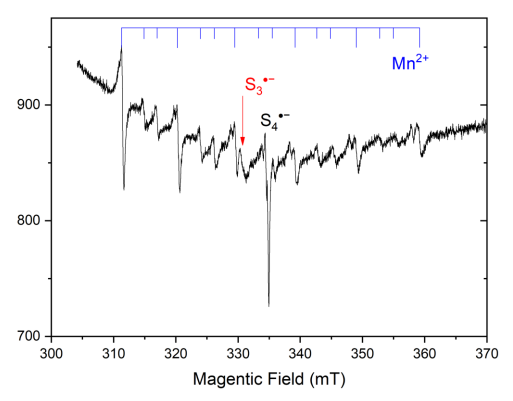


Figure 5. ESR spectrum of haüyne (Sample 1) measured at room temperature.

The absorption bands with maxima at 0.65 and 0.88 eV are associated with the vibrational modes of water in the studied mineral. The absorption bands with maxima at 2.05 and 2.38 eV, which impart a blue color to the studied mineral, can be associated with the presence of $S_3^{\bullet-}$ radical anions, which are easily detected by ESR [4,5], while the lilac hue can be associated with the presence of S_4 molecules (red chromophore). In the samples under study, the ESR signal characteristic of $S_3^{\bullet-}$ (with g=2.036) is practically not observed, which indicates a very low concentration of $S_3^{\bullet-}$ radical anions in Sample 1. The absorption spectrum of $S_3^{\bullet-}$ radical anions is characterized by a plateau at about 1.77 eV. However, the shape of the absorption spectrum of Sample 1 differs from the characteristic absorption of $S_3^{\bullet-}$ of lazurite [4,5]. Thus, the absorption band at 2.05 eV is not related to $S_3^{\bullet-}$ radical anions. On the other hand, in a number of works [40,41,52,53] it is noted that absorption bands at 2.05 and 2.38 eV, as well as a much less intense band about 1.33 eV, can be attributed to the absorption of *trans*- and *cis*-conformers of the neutral S_4 molecule. The authors of [40] calculated the oscillator strengths f for these conformals, which are equal to 0.07 and 0.047, respectively.

The observed intense ESR signal with a g-factor of 2.002 can be attributed to the presence of some amount of $O^{\bullet-}$ radical anions (Figure 5). Previously, blue sodalite, in which a similar ESR signal and absorption spectrum were observed, was studied in [54,55], but the nature of the color of the mineral was not fully elucidated in this work.

When this sample is heated above a temperature of 500 °C, the studied sodalite loses its color. Along with this, an absorption band appears at 3.1 eV (400 nm) upon excitation, in which intense luminescence is observed with a maximum at 1.9 eV (650 nm). Such luminescence is characteristic of S₂•- radical anions [4].

3.2. Crystal Chemistry, Isomorphism, and Thermal Conversions of Haüyne

The IR spectra of initial and heated haüyne (Samples 2 and 5) are given in Figures 6–8. The spectra of initial samples contain bands of O–H stretching and bending vibrations of H₂O molecules (in the ranges of 3300–3700 and 1640–1670 cm⁻¹, respectively), antisymmetric vibrations of the CO₂ molecules (at 2341 cm⁻¹), asymmetric stretching

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 $[F_2(v_3) \text{ mode } 1135\text{--}1137 \text{ cm}^{-1}]$ and bending $[F_2(v_4) \text{ mode, } 616 \text{ cm}^{-1}]$ vibrations of the SO₄ tetrahedra, as well as stretching (1002–1004 cm⁻¹), O–(Si,Al)–O (650–730 cm⁻¹), and Si–O–Al (in the range of 390–450 cm⁻¹) modes of the aluminosilicate framework.

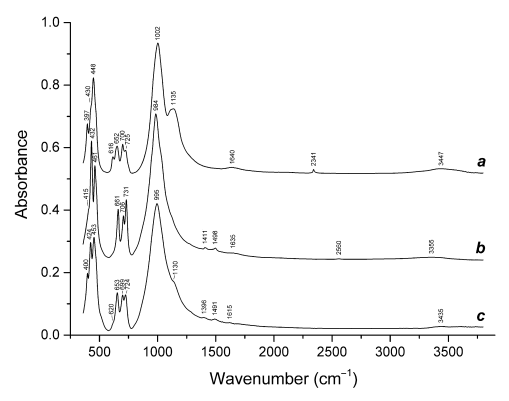


Figure 6. Infrared spectra of (a) initial haüyne (Sample 2), (b) Sample 2 preheated for three days at 700 °C, over the Fe-FeS buffer, and (c) preheated Sample 2 additionally annealed at 800 °C in air for one day.

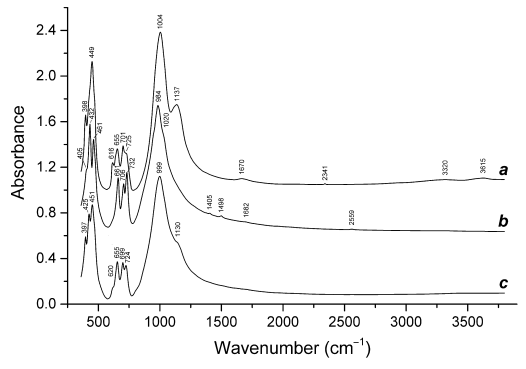


Figure 7. Infrared spectra of **(a)** initial haüyne (Sample 5), **(b)** Sample 5 preheated for three days at 700 °C over the Fe-FeS buffer, and **(c)** preheated Sample 5 additionally annealed at 800 °C in air for one day.

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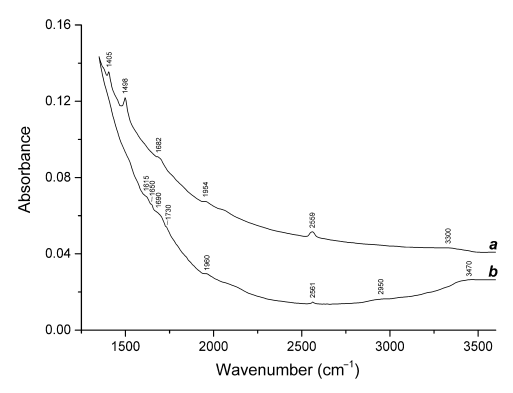


Figure 8. Infrared spectra in the 1350–3600 region of (**a**) haüyne (Sample 5) preheated for three days at 700 °C over the Fe-FeS buffer, and (**b**) preheated Sample 5 additionally annealed at 800 °C in air for one day.

Heating results in the lowering of intensities of the bands of H₂O and disappearance of the bands of SO₄²⁻ and CO₂. The samples preheated under reducing conditions change color from blue to yellow. Their IR spectra contain a weak band at 2559–2560 cm⁻¹, which is due to stretching vibrations of the HS⁻ anion [39]. The weak bands at 1405–1411 and 1498 cm⁻¹ in the IR spectra of the preheated samples correspond to asymmetric stretching vibrations of carbonate groups required for the charge compensation. The weak band at 1954 cm⁻¹ (curve *a* in Figure 8) is an overtone of stretching vibrations of the aluminosilicate framework.

It is worth noting that bands of the isolated S^{2-} anion are absent in the middle IR range, but the transformation of SO_4^{2-} into S^{2-} as a result of annealing of lazurite at 800 °C was reported [56,57]. Thus, the main scheme of transformations in the first step includes reduction of sulfate anions, and formation of S^{2-} and HS^{-} .

Splittings and shifts of IR bands in the preheated samples are mainly due to transformations of the framework. However, the band at 461 cm⁻¹ may be partly due to vibrations of the *trans*-S₄ molecule (C_{2h} conformation) [40].

Annealing of the preheated haüyne samples at 800 °C in air results in the change of color from yellow to blue, disappearance or weakening of the band of HS⁻ and in the case of Sample 5, disappearance of the bands of CO_3^{2-} (Figures 1–3).

As noted above, unlike IR spectra, Raman spectra are sensitive to anionic groups and neutral molecules containing sulfide sulfur. The Raman spectra of Samples 2 and 5 are given in Figures 9 and 10.

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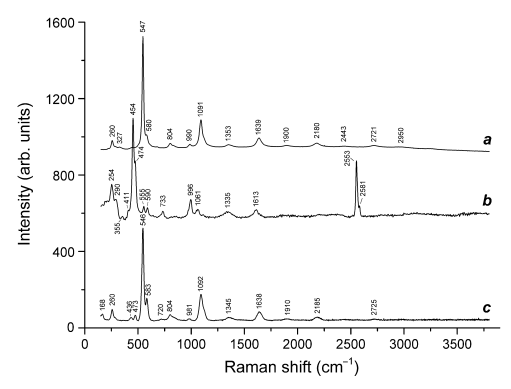


Figure 9. Raman spectra of **(a)** initial Sample 2, **(b)** Sample 2 preheated for three days at 700 °C, over the Fe-FeS buffer, and **(c)** preheated Sample 2 additionally annealed at 800 °C in air for one day.

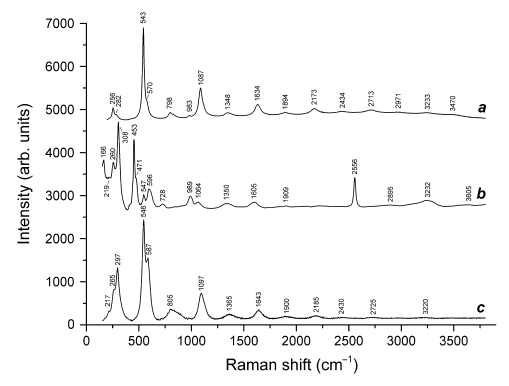


Figure 10. Raman spectra of (**a**) initial haüyne (Sample 5), (**b**) Sample 5 preheated for three days at 700 °C, over the Fe-FeS buffer, and (**c**) preheated Sample 5 additionally annealed at 800 °C in air for one day.

The assignment of bands in the Raman spectra of Samples 2 and 5 and products of their thermal conversions made is given in Tables 3 and 4. The Raman spectra confirm the presence of HS^- and CO_3^{2-} anions in the products of the heating of both samples at 700

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°C, over the Fe-FeS buffer. Additionally, the Raman spectra indicate the formation of oxalate $C_2O_4{}^{2-}$ and monosulfide S_-^{2-} anions, $S_2{}^{\bullet-}$ and $S_4{}^{\bullet-}$ radical anions, and S_4 molecules in the preheated samples. The Raman spectra of both initial haüyne samples and products of their annealing at 800 °C in air contain sets of characteristic bands of $S_3{}^{\bullet-}$ radical anions.

Table 3. Assignment of Raman bands of Sample 2 and products of its thermal conversions.

	Raman Shift (d	cm ⁻¹)		
Initial Sample	Preheated Sample	Sample Heated at 800 °C in Air	Assignment	
-	194w	168w	Combination of low-frequency lattice modes	
-	213w	219	trans-S4 bending mode	
-	254	-	Bending vibrations of the [(HS)-Na ₄] ³⁺ cluster	
260	-	260	S₃•- bending mode (v₂)	
287w	290	-	Low–frequency lattice modes involving Na ⁺ cations and/or S4 ^{•-} bending vibrations	
	-	-	cis-S ₄ mixed v ₄ mode (combined symmetric bending + stretching vibrations)	
327w	355w	-	cis-S₄• mixed v₃ mode	
_	-	380	cis-S₄ mixed v₃ mode	
448w	-	-	SO_4^{2-} [$E(v_2)$ mode] and/or δ [O–Si(Al)–O] bending vibrations	
_	454s	-	Stretching vibrations of the [(HS)-Na ₄] ³⁺ cluster	
_	474	473	S ₆ stretching mode and/or mixed v ₄ mode of <i>trans</i> –S ₄	
547s	-	546s	S ₃ •- symmetric stretching (v ₁) mode	
	555	-	S ₃ neutral molecule and/or <i>gauche-S</i> ₄	
580	-	583	S ₃ • antisymmetric stretching (v ₃), possibly, overlapping with the stretching band of S ₂ •-	
	590	-	Stretching vibrations of the [(S ² -)Na ₄] ²⁺ cluster and/or S ₂ •- stretching mode	
	733	-	O–C–O bending vibrations of oxalate anions	
	_	720	Mixed vibrations of the aluminosilicate framework	
804	-	804	S ₃ •- combination mode (v ₁ + v ₂)	
_	845w	-	C–C stretching vibrations of oxalate anions	
990	996	981w	SO_4^{2-} symmetric stretching vibrations [$A_1(v_1)$ mode] and/or framework stretching vibrations	
-	1061	-	CO ₃ ²⁻ symmetric stretching vibrations	
1091s	-	1092s	S₃•- overtone (2 × v₁)	
-	1335	-	Symmetric C–O stretching vibrations of oxalate anions	
1353	-	1345	$S_3^{\bullet-}$ combination mode $(2v_1 + v_2)$	
-	1613	-	Antisymmetric C–O stretching vibrations of oxalate anions	
1639	-	1638	S ₃ •- overtone (3 × v ₁)	
1900w	-	1910w	S ₃ •- combination mode $(3 \times v_2 + v_1)$	
2180	_	2185	$S_3^{\bullet-}$ overtone $(4 \times v_1)$	
2443w	_	-	S ₃ *- combination mode $(4 \times v_2 + v_1)$	
_	2553s	-	HS ⁻ stretching mode	
_	2581w	-	H ₂ S symmetric stretching mode	
2721	-	2725	$S_3^{\bullet-}$ overtone $(5 \times v_1)$	
2950w	_	-	S ₃ *- combination mode ($5 \times v_1 + v_2$)	
			` ′	

Note: w—weak band, s—strong band.

Table 4. Assignment of Raman bands of Sample 5 and products of its thermal conversions.

	Raman Shif	t (cm ⁻¹)	
Initial	Preheated	Sample Heated at	Assignment
Sample	Sample	800 °C in Air	
-	166	-	Combination of low-frequency lattice modes
-	219w	217w	trans-S4 bending mode
-	260	-	Bending vibrations of the [(HS)-Na ₄] ³⁺ cluster

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256	-	265	S ₃ •- bending mode (v ₂)
282w	-	-	Low–frequency lattice modes involving Na ⁺ cations and/or S4 ^{•-} bending vibrations
_	308s	297s	S ₄ •- bending vibrations and/or <i>cis</i> -S ₄ mixed v ₄ mode (combined symmetric bending + stretching vibrations)
-	453s	-	Stretching vibrations of the [(HS)-Na ₄] ³⁺ cluster
-	471	-	S ₆ stretching mode and/or mixed v ₄ mode of trans–S ₄
543s	547	546s	S ₃ •- symmetric stretching (v ₁) mode
570	-	587	S ₃ *- antisymmetric stretching (v_3), possibly, overlapping with the stretching band of S ₂ *-
_	596	-	Stretching vibrations of the [(S²-)Na ₄]²+ cluster and/or S₂•- stretching mode
-	728	-	O-C-O bending vibrations of oxalate anions
798	-	805	$S_3^{\bullet-}$ combination mode ($v_1 + v_2$)
-	850w	-	C-C stretching vibrations of oxalate anions
983w	989	-	SO_4^{2-} symmetric stretching vibrations [$A_1(v_1)$ mode] and/or framework stretching vibrations
-	1064	-	CO ₃ ²⁻ symmetric stretching vibrations
1087s	-	1097s	$S_3^{\bullet-}$ overtone $(2 \times v_1)$
-	1350	-	Symmetric C–O stretching vibrations of oxalate anions
1348	-	1365	$S_3^{\bullet-}$ combination mode $(2v_1 + v_2)$
-	1605	-	Antisymmetric C–O stretching vibrations of oxalate anions
1634	-	1643	S₃•- overtone (3 × v₁)
1894w	-	1900w	S ₃ •- combination mode $(3 \times v_2 + v_1)$
-	1909w	-	Overtone of SO ₄ ² - symmetric stretching vibrations?
2173	-	2185	$S_3^{\bullet-}$ overtone $(4 \times v_1)$
2434w	-	2430w	$S_3^{\bullet-}$ combination mode $(4 \times v_2 + v_1)$
	2556s	-	HS- stretching mode
2713	-	2725	S₃•- overtone (5 × v₁)
2971w	-	-	S ₃ •- combination mode ($5 \times v_1 + v_2$)
3233w, 3470w	3232, 3605w	3220w	O-H stretching modes

In the original Sample 3, no $S_3^{\bullet-}$ centers were observed before heating. Two absorption bands with maxima at 525 and 585 nm are associated with $S_4^{\bullet-}$ centers in different configurations, and a weak absorption band at 400 nm corresponds to $S_2^{\bullet-}$ centers (Figure 11) [4,58]. Being excited in the 400 nm region, the samples show intense luminescence with a maximum in the 650 nm region associated with these $S_2^{\bullet-}$ radical anions.

In the ESR spectrum, in addition to six lines associated with Mn^{2+} centers in calcite inclusions, there is a signal with *g*-tensor components of 2.034 and 2.021, associated with $S_4^{\bullet-}$ centers (Figure 12) [58]. As a result of heating the sample, the intensity of this signal decreases, together with a weakening of the absorption bands. At temperatures above $400\,^{\circ}\text{C}$, the $S_4^{\bullet-}$ —centers disappear.

The intensities of the absorption band and luminescence associated with $S_2^{\bullet-}$ centers increase in this case. At temperatures above 700 °C, a signal with g_1 = 2.056, g_2 = 2.041, and g_3 = 2.008, associated with $S_2^{\bullet-}$ centers, appears in the ESR spectrum. At the same time, the shape of the absorption band in the region of 600 nm changes, it acquires a table-like shape, and its intensity also begins to grow. The sample acquires an intense blue color, also associated with the $S_3^{\bullet-}$ radical anion [4]. In this case, the absorption band in the region of 400 nm and the luminescence associated with the $S_2^{\bullet-}$ centers begin to decrease. The temperature dependences of the observed effects are shown in Figure 13.

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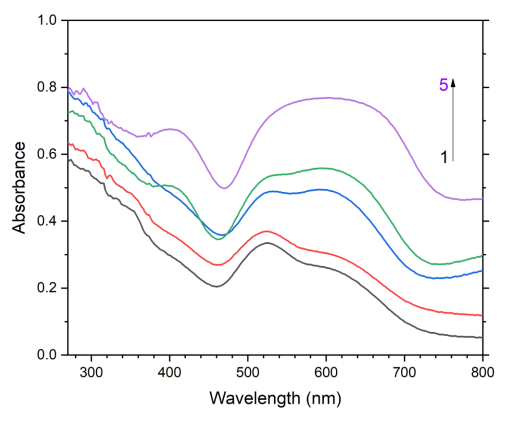


Figure 11. Spectra of diffuse absorption of haüyne (Sample 3): unheated sample (1) and samples heated at 200 °C (2), 400 °C (3), 600 °C (4), and 800 °C (5).

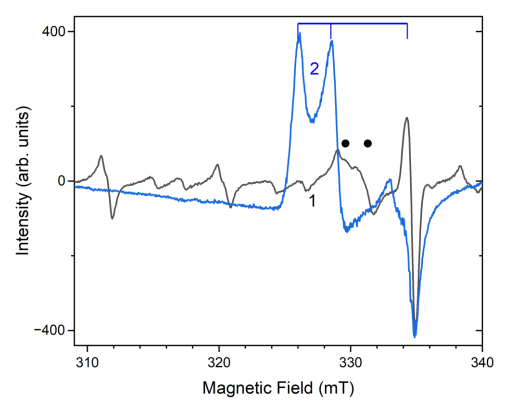


Figure 12. The ESR spectra of unheated haüyne (Sample 3) (1) and Sample 3 heated at a temperature of 800 °C (2). The dots indicate the bands of $S_4^{\bullet-}$ and the vertical lines show the bands of $S_2^{\bullet-}$.

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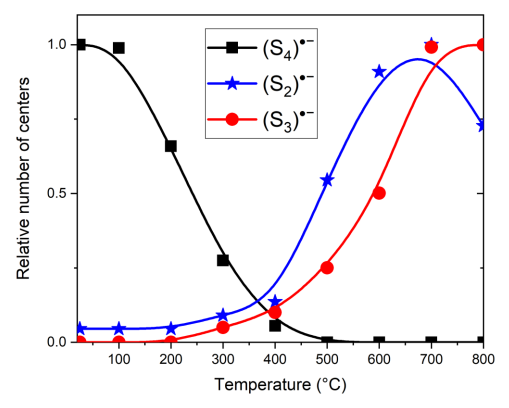


Figure 13. Relative amounts of various sulfur radical anions in haüyne (Sample 3) heated at different temperatures. The values of the relative amounts are normalized to the maximum number of each of the radical anions.

The absorption spectrum of initial sample 4 shows a broad band at 600 nm, which is also associated with $S_3^{\bullet-}$ radical anions, as well as a sharp rise in the region of 280 nm, which can be caused by the absorption of $S_6^{\bullet-}$ radical anions (Figure 14) [39,58,59]. In the original sample 4, a weak signal with the *g*-tensor components $g_1 = 2.045$, $g_2 = 2.034$, and $g_3 = 2.008$, associated with $S_3^{\bullet-}$ radical anions [4], as well as a signal with the *g*-tensor components $g_1 = 2.011$ and $g_2 = 2.006$, associated with the radical anion $S_6^{\bullet-}$ [45,58,60], are observed (Figure 15).

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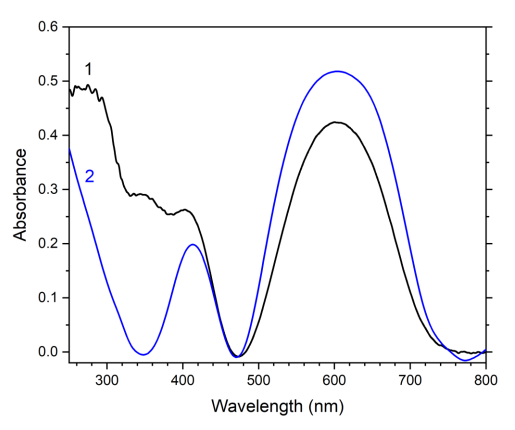


Figure 14. Diffuse absorption spectra of original haüyne (Sample 4) (1) and Sample 4 heated at 800 $^{\circ}$ C (2).

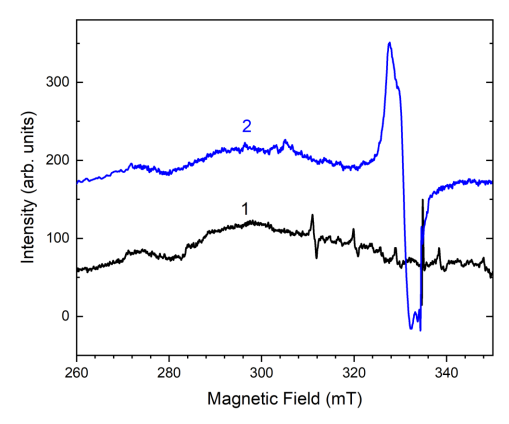


Figure 15. The ESR spectra of original haüyne (Sample 4) (1) and Sample 4 heated at 800 °C (2). The black triangles show the bands of the $S_6^{\bullet-}$ radical anions bands, and the vertical blue lines indicate the bands of $S_3^{\bullet-}$.

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When the sample was heated above 500 °C, the ESR signal with g_1 = 2.011 and g_2 = 2.017 decreases together with a decrease in the rise intensity in the region of 280 nm. Along with this, the sample acquires a saturated blue color, which is accompanied by an increase in the ESR signal with g_1 = 2.045, g_2 = 2.034, and g_3 = 2.008, as well as absorption bands in the region of 600 nm associated with $S_3^{\bullet-}$ radical anions. The temperature dependences built on the basis of the analysis of the absorption and EPR spectra are shown in Figure 16.

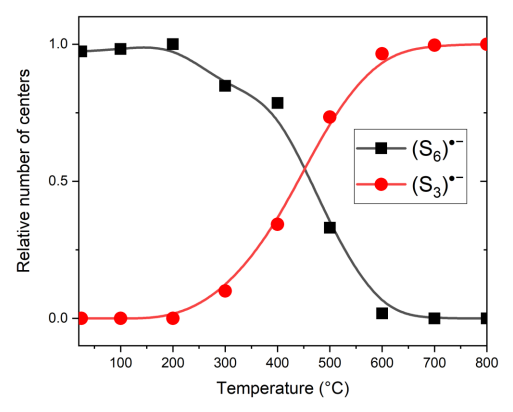


Figure 16. Relative amounts of polysulfide radical anions in haüyne (Sample 3) heated at different temperatures. The values of the relative amounts are normalized to the maximum number of each of the radical anions.

The blue color of Sample 2 is due to the presence of the S₃• radical anions, which are easily identified by an ESR signal with $g_1 = 2.049$, $g_2 = 2.035$, and $g_3 = 2.010$ (Figure 17, curve 1). After heating in reducing atmosphere using the Fe/FeS buffer at 700 °C, the color of the sample changes to yellow, and the ESR signal associated with the S₃• radical anion disappears (Figure 17, curve 2). During subsequent heating at 800 °C, the blue color is partially restored and an ESR signal with $g_1 = 2.049$, $g_2 = 2.035$, and $g_3 = 2.010$ appears (Figure 17, curve 3). However, its intensity is less than that of the original sample.

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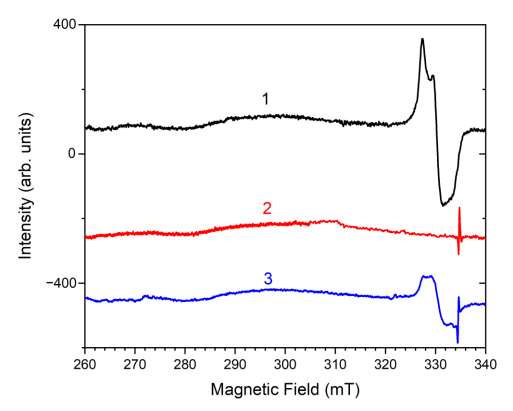


Figure 17. ESR spectra of initial haüyne (Sample 2) (1), and Sample 2 heated at 700 °C under buffer (2) and at 800 °C in air (3).

The observed results show that various polysulfide radical anions are converted into $S_2^{\bullet-}$ and $S_4^{\bullet-}$ radical anions and S_4 neutral molecules during heating at 700 °C in a reducing atmosphere, whereas the $S_3^{\bullet-}$ radical anion is stable during heating at 800 °C in air. These data do not contradict the experimental results obtained in [61,62], where it was shown that various sulfur radicals are converted into $S_2^{\bullet-}$ and $S_3^{\bullet-}$ radical anions when heated.

It should be noted that the values of the g-tensor components of the $S_3^{\bullet-}$ radical anions formed during heating of most S-bearing sodalite-group minerals studied in this work and elsewhere [58] are close to those of the g tensor of $S_3^{\bullet-}$ in haüyne and some synthetic sodalites [4,63]. The only exception is $S_3^{\bullet-}$ radical anion in Sample 3 heated in air: its values of the g-tensor components are close to those of $S_3^{\bullet-}$ or $S_3^{\bullet-}$ occurring in minerals of the cancrinite group [58,63].

Annealing of preheated powdered Sample 5 at 800 °C in air results in gradual enhancement of its unit cell parameter from 8.04 to 8.97 Å, accompanied by its transformation into a new cubic phase with the a parameter growing from 9.05 to 9.08 Å during 32 h (Figure 18). This transformation is accompanied by broadening and shifts of reflections.

The initial phase with a smaller a parameter disappears five hours after the start of annealing. The final product of annealing is pale blue. Annealing of preheated single-crystal grains of Sample 5 at 800 °C during 32 h results in the formation of a greenish blue cubic phase. Based on the above spectroscopic data, one can conclude that the enhancement of the unit cell parameter during annealing is caused by the transformation of small extra-framework anions (S²- and HS⁻) into larger ones (SO₄²- and, to a lesser extent, S₃•-). Thus, the process observed during annealing has the character of a phase transition, and both in the initial and in the final phase, the degree of sulfur oxidation (and, consequently, the unit cell parameter) can vary smoothly within relatively narrow limits.

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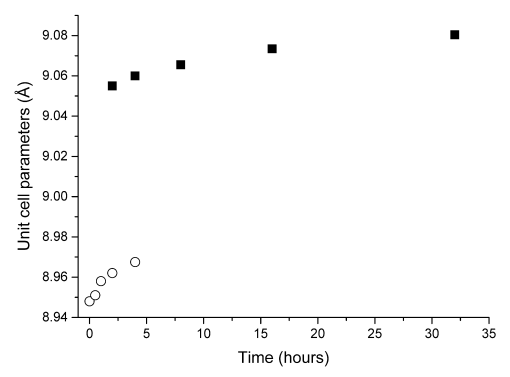


Figure 18. Kinetics of changes of the cubic unit cell parameter of the initial phase of the preheated haüyne (Sample 5, circles) and product of its annealing at 800 °C (squares).

The observed abrupt increase in the unit cell parameter may indicate that the formation of the new phase proceeds by the nucleation mechanism. Partial oxidation of sulfur in the preheated sample at the initial stage of calcination leads to a broadening of the lines in the diffraction patterns, which may indicate the inhomogeneity of the composition and associated mechanical stresses in the crystal.

3.3. Thermal Conversions of Slyudyankaite

Slyudyankaite, ideally Na₂₈Ca₄(Si₂₄Al₂₄O₉₆)(SO₄) $_6$ (So₄) $_6$ (So₂) $_1$ (CO₂)·2H₂O, is a triclinic sodalite-group mineral containing sodalite cages of two kinds. Cages of the first type contain Na⁺ and Ca²⁺ cations and SO₄²⁻ anions as well as trace amounts of S₃•- radical anions. Cages of the second type contain only neutral molecules (cyclic S₆ having chair-like conformation, CO₂, H₂O, and subordinate S₄).

The IR spectrum of slyudyankaite (curve a in Figure 19) contains bands of O-H stretching vibrations of hydrogen bonds of medium strength (3380 to 3610 cm⁻¹), strong hydrogen bond or strong dipole-dipole interactions with CO2 molecules (shoulder at 3240 cm⁻¹), antisymmetric stretching vibrations of free ¹²CO₂ molecules and ¹²CO₂ molecules which are acceptors of strong hydrogen bonds (2341 and 2385 cm⁻¹, respectively), antisymmetric stretching vibrations of the ¹³CO₂ molecules (2275 cm⁻¹), C-O stretching vibrations of trace admixture of O=C=S molecules (a very weak band at 2040 cm⁻¹), bending vibrations of H₂O molecules (at 1632 cm⁻¹ with an indistinct shoulder at ~1680 cm⁻¹ which may correspond to H₂O molecules forming strong hydrogen bonds), asymmetric stretching vibrations of the SO₄²⁻ anionic groups [the degenerate F₂(v₃) mode] (1138 and 1107 cm⁻¹), stretching vibrations of the aluminosilicate framework (1002 cm⁻¹), mixed vibrations of the aluminosilicate framework (in the range of 650-720 cm⁻¹), stretching vibrations of the neutral S₄ molecule having cis conformation (641 cm⁻¹), bending vibrations of the SO_4^{2-} anionic groups [the degenerate $F_2(v_4)$ mode] (614 cm⁻¹), and lattice modes involving bending vibrations of the aluminosilicate framework and librations of SO₄²⁻ and extra-framework molecules (below 550 cm⁻¹). An indistinct Minerals 2022, 12, 887 21 of 31

shoulder at ~465 cm⁻¹ may be partly due to stretching vibrations of the cyclic S₆ molecule having D_{3d} symmetry [41].

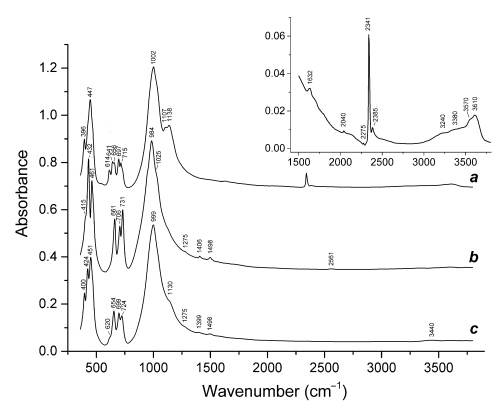


Figure 19. Infrared spectra of (a) initial slyudyankaite (Sample 6), (b) Sample 6 preheated for three days at 700 °C, over the Fe-FeS buffer, and (c) preheated Sample 6 additionally annealed at 800 °C in air for one day. The inset shows the IR spectrum of initial Sample 6 in the range of 1500–3800 cm⁻¹.

Changes in the IR spectrum of slyudyankaite that occur as a result of its calcination are basically similar to those observed during calcination of the haüyne samples described above: the bands of O–H stretching vibrations, CO₂, S₄, and S₆ molecules, and SO₄²⁻ anions disappear or become weaker. In the IR spectrum of Sample 6 preheated at 700 °C under reducing conditions, a weak band of HS⁻ is observed at 2561 cm⁻¹. Both heated samples (preheated and annealed at 800 °C in air) contain bands of CO₃²⁻ anions in the range of 1390–1500 cm⁻¹. Unlike IR spectra of heated haüyne (Samples 2 and 5), spectral curves of both preheated and annealed at 800 °C, slyudyankaite samples have a weak shoulder at 1275 cm⁻¹, which may correspond to the BO₃³⁻ anion. If this assumption is correct, then it follows that initial slyudyankaite contained boron in the four-fold coordination, which could not be detected by means of IR spectroscopy because IR bands of the BO₄ tetrahedron overlap with bands of the aluminosilicate framework.

Raman spectra of slyudyankaite and products of its heating are presented in Figure 20. The assignment of Raman bands is given in Table 5.

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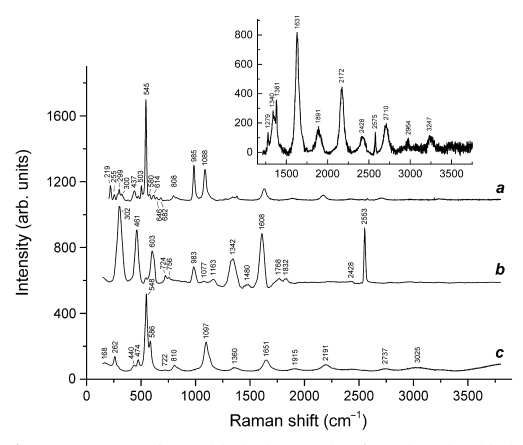


Figure 20. Raman spectra of **(a)** initial slyudyankaite (Sample 6), **(b)** Sample 6 preheated for three days at 700 °C, over the Fe-FeS buffer, and **(c)** preheated Sample 6 additionally annealed at 800 °C in air for one day. The inset shows the Raman spectrum of initial Sample 6 in the range of 1200–3750 cm⁻¹.

Table 5. Assignment of Raman bands of slyudyankaite and products of its thermal conversions.

	Raman Shift	(cm ⁻¹)	
Initial Sample	Preheated Sample	Sample Heated at 800 °C in Air	Assignment
219	-	-	trans-S4 bending
260	-	262	S ₃ •- bending A ₂ (v ₂) and S ₆ (with D _{3d} symmetry) bending
283	-	-	Framework bending vibrations (resonance with a S ₆ bending mode?)
298	302s	-	S ₄ •- bending vibrations
330	-	-	cis-S ₄ mixed v ₄ mode (symmetric bending + stretching)
380w	-	-	cis-S₄ mixed v₃ mode
437	-	440w	SO ₄ [bending E (v ₂) mode] and/or S ₆ (mixed mode)
-	461s	-	[(HS)Na ₄] ³⁺ stretching vibrations
477	-	474	S ₆ stretching mode and/or mixed v ₄ mode of trans-S ₄
503	-	-	Bending vibrations of the framework
545 s	-	548s	S ₃ • symmetric stretching (v ₁) (possibly, overlapping with the stretching band of <i>gauche-</i> S ₄)
580	-	586	S ₃ •- antisymmetric stretching mode (v ₃)
-	603	-	S₂• stretching mode
614	-	-	SO ₄ ²⁻ [bending F ₂ (v ₄) mode]
645	-	-	cis-S ₄ stretching
682	-	-	trans-S₄ symmetric stretching v₃ mode
-	724, 756w	722w	O-C-O bending vibrations of oxalate anions
807	-	810	S ₃ •- combination mode ($v_1 + v_2$)
985s	983	-	$SO_{^{4^{2-}}}[symmetric\ stretching\ A_{^{1}}\ (v_{^{1}})\ mode]$ (possibly, overlapping with the weak

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			band of framework stretching vibrations)
-	1077w	-	CO ₃ ²⁻ symmetric stretching mode
1088 s	-	1097s	S ₃ •- overtone (2 × v_1) [possibly, overlapping with the SO ₄ •- stretching band (v_3 – F_2)]
-	1163	-	$S_2^{\bullet-}$ overtone (2 × v_1)
1279, 1381	-	-	Symmetric stretching vibrations of CO ₂ molecules (Fermi doublet, resonance with the overtone of bending vibrations).
1340	1342	1360	Symmetric C–O stretching vibrations of CO ₂ molecules—involved in strong dipole-dipole interactions and/or symmetric C–O stretching vibrations of acid oxalate anions
-	1480w	-	CO ₃ ²⁻ asymmetric stretching mode
-	1609s	-	Antisymmetric C–O stretching vibrations of acid oxalate anions
1631	-	-	S₃•- overtone (3 × v₁)
-	-	1651	Symmetric C–O stretching vibrations of oxalate anions
-	1768, 1832	-	C=O stretching vibrations of acid oxalate groups
1891	-	1915	$S_3^{\bullet-}$ combination mode $(3 \times v_1 + v_2)$
2172	-	2191	S3 $^{\bullet-}$ overtone (4 × v_1)
2428w	-	-	$S_3^{\bullet-}$ combination mode $(4 \times v_2 + v_1)$
-	2553s	-	HS- stretching mode
2575w	-	-	H ₂ S symmetric stretching mode
2710	-	2737	$S_3^{\bullet-}$ overtone (5 × v_1)
2964 w	-	-	$S_3^{\bullet-}$ combination mode $(5 \times v_1 + v_2)$
3025	-	_	O–H stretching vibrations
3247w	-	-	S₃•- overtone (6 × v₁)

The Raman spectrum of the initial slyudyankaite sample contains bands of SO₄²- anions, S₅•- radical anions, and different neutral molecules (*cis*- and *trans*-S₄, S₆, H₂S, and CO₂). Preheating of slyudyankaite at 700 °C under reducing conditions results in the transformation of S-bearing species into HS⁻, S₂•-, and S₄•- and, possibly, monosulfide anion S²-, which could not be detected by means of vibrational spectroscopy methods but is required for the charge balance. Note that S²- was identified as the main extra-framework anion in the crystal structure of the product of thermal conversions of lazurite under the same conditions [57]. Reduction of CO₂ occurring in initial slyudyankaite results in the formation of carbonate and acid oxalate anions.

Further annealing of preheated slyudyankaite at 800 °C in air results in the disappearance of HS-, S2•-, and S4•-, restoration of the SO4²- and S3•- anionic groups, and transformation of acid oxalate groups into neutral ones. Bands of S3•- in the Raman spectrum of the annealed slyudyankaite are shifted towards higher frequencies as compared to analogous bands of the initial sample, which may indicate their occurrence in a straitened state.

3.4. Sodalite — Sapozhnikovite Solid-Solution Series

Sodalite, ideally Nas(Al₆Si₆O₂₄)Cl₂, and sapozhnikovite, its hydrosulfide-dominant structural analogue with the formula of the hypothetical end-member Nas(Al₆Si₆O₂₄)(HS)₂ form a solid-solution (isomorphous) series in which the Cl⁻:HS⁻ (simply, Cl:S) ratio is the only significant variable. In the studied specimens from Karnasurt Mt. (Lovozero alkaline massif), this ratio varies from [Cl_{1.11}(HS)_{0.84}]_{E1.95} to [(HS)_{1.57}Cl_{0.38}]_{E1.95} (Table 1; Figure 21). Samples with Cl > S (in atom proportions) belong to the mineral species sodalite, whereas samples with S > Cl belong to sapozhnikovite. Slight deficiency of the Cl + S sum in comparison with the ideal value 2.00 atoms per formula unit (Table 1; Figure 21) is caused by the presence of minor amounts of H₂O in these minerals [39], probably together with Cl⁻ and HS⁻ anions in sodalite cages. The presence of water also explains slight deficiency of analytical totals in all electron-microprobe analyses of sodalite–sapozhnikovite series minerals (Table 1).

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In the interval [Cl_{1.11-0.38}(HS)_{0.84-1.57}], this homovalent isomorphous series is continuous (Figure 21). We suggest this interval is caused by geochemical features rather crystal chemical constraints.

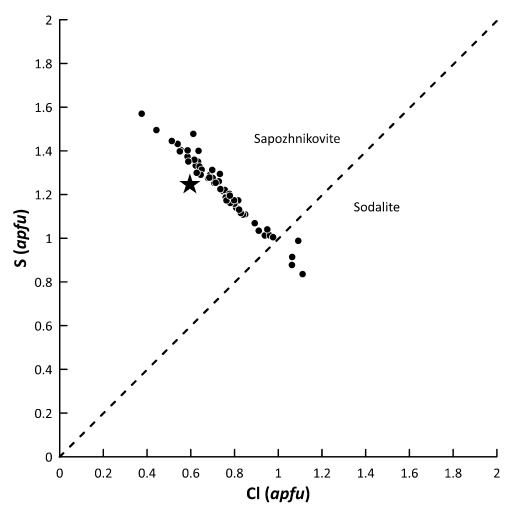


Figure 21. Variations of the contents of Cl and S atoms per formula unit (*apfu*) in minerals belonging to the sodalite–sapozhnikovite solid-solution series from Karnasurt Mt., Lovozero massif, Kola Peninsula, Russia.

4. Discussion

4.1. General Remarks on the Isomorphism of Extra-Framework Components in Sodalite-Group Minerals

The general common formula of cubic alkaline aluminosilicates belonging to the sodalite group is $ABNa_6(Si_6Al_6O_{24})XY$, where the pairs of cationic and anionic extra-framework components AB + XY are $Na_2 + Cl_2$ for sodalite, $Na_2+(HS)_2$ for sapozhnikovite, $K \Box + F(H_2O)_4$ for bolotinaite, $Na_2+(SO_4)(H_2O)$ for nosean, $Ca_2+(SO_4)_2$ for haüyne, and $NaCa+(S_3^{\bullet-})(SO_4)$ for lazurite. However, the application of a complex of spectroscopic methods in this work and elsewhere [3,4,64] shows that the crystal chemistry of subordinated components in cubic sodalite-group aluminosilicates is much more complex. In particular, H^+ , H_3O^+ , CO_3^{2-} , OH^- , $S_2^{\bullet-}$, S_3 , cis- and trans- $S_4^{\bullet-}$, $SO_3^{\bullet 2-}$, $SO_4^{\bullet 2-}$, cis-, trans- and gauche- S_4 , S_6 , CO_2 , CO_3 , and HF were identified as extra-framework species in these minerals. Experiments with annealing show that these components take part in complex thermal conversions involving some additional components, e.g., oxalate anions. Similar transformations can occur in natural mineral assemblages. Thus, the composition of sodalite-group minerals may serve as a marker of mineral-forming media including fugacities of volatile components (H_2O , O_2 , CO_2 , HF, SO_2 , and polysulfide compounds).

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In general, similar processes were observed in sulfur-enriched fluids [61,62]. Apparently, when heated, the channel widths increase, which allows complex anion radicals to migrate and disproportionate with the formation of the most stable radical anion radicals (S₃•- or S₂•-, depending on the redox conditions). Along with this, dehydration of the studied samples also occurs, which also facilitates the migration of polysulfide species.

4.2. Sapozhnikovite as a Marker of Reducing Conditions

Experiments with heating of haüyne under reducing conditions (in the presence of Fe-FeS buffer) show that the main scheme of transformations of extra-framework components is $SO_4^{2-} \rightarrow S^{2-} + 2O_2(gas)$, and a subordinate process is $CO_2 + 2SO_4^{2-} + H_2O \rightarrow 2HS^+ + CO_3^{2-} + 4O_2(gas)$. The conversions $3SO_4^{2-} \rightarrow S_3^{\bullet-} + 5e + 6O_2(gas)$, $2S_3^{\bullet-} \rightarrow S_2^{\bullet-} + S_4^{\bullet-}$, and $2CO_2 + 2e \rightarrow C_2O_4^{2-}$ (e = electron) are the additional channels of haüyne transformations during their heating at 700 °C under reducing conditions.

Thus, HS⁻ and C₂O₄²- anions are stable only under reducing conditions. Data on the occurrence of sapozhnikovite Na₈(Al₆Si₆O₂₄)(HS)₂ confirm this conclusion. Crystallization of aegirine NaFe³⁺Si₂O₆, which precedes formation of sapozhnikovite, could be accompanied by the following redox reactions: Na₂O + 2FeO + 4SiO₂ + CO₂ \rightarrow 2NaFeSi₂O₆ + CO; Na₂O + 2FeO + 4SiO₂ + 0.5CO₂ \rightarrow 2NaFeSi₂O₆ + 0.5C, where C is carbon with the oxidation degree of 0 [39]. During the crystallization of aegirine, iron occurring as Fe²⁺ in the fluid is fixed as Fe³⁺ in the solid phase. Such processes are possible only in peralkaline media with a high Na:Ca ratio (otherwise, hedenbergite CaFe²⁺Si₂O₆ is formed instead of aegirine). Similar reactions, e.g., 2Na₂O + 4FeO + 4SiO₂ + 2SO₃ + 2OH⁻ + O₂ \rightarrow 4NaFeSi₂O₆ + 2HS⁻, could lead to the formation of reduced forms of sulfur, including HS⁻ anions.

The redox processes in sodalite-group minerals discussed above involve species hosted by sodalite cages as well as gaseous O₂. However, in the case of sapozhnikovite, an alternative mechanism including incorporation of HS⁻ as a result of the exchange reaction of sodalite with a late reduced fluid is not excluded.

Unlike CO₂ molecules, carbonate anions rarely occur in sodalite-group minerals in significant amounts. An unusual CO₃²-bearing mineral with the sodalite-type framework, empirical formula (Na_{5.28}K_{0.40}Ca_{1.23}Fe_{0.01})(Si_{5.93}Al_{6.07}O_{24.01})(SO₄)_{0.77}Cl_{0.94}(CO₃)_{0.30} and unit cell parameter a = 9.0352 Å has been discovered in a metasomatic ejectum from Mount Vesuvio, Italy and its crystal structure has been solved [65]. Taking into account above-described thermal transformations of CO₂ in haüyne and the fact that the CO₃-bearing sodalite-group mineral from Vesuvio underwent heating during metasomatism, it is reasonable to suppose that CO₃²- anions in this mineral were formed as a result of the transformation CO₂ + 2SO₄²- + H₂O \rightarrow 2HS⁻ + CO₃²- + 4O₂(gas) in an initial sodalite-group mineral. In the experiments with heating of haüyne under reducing conditions, CO₂ molecules occurring in sodalite cages partly transformed into neutral or acid oxalate anions. In this regard, the association of sapozhnikovite with kyanoxalite (a cancrinite-group mineral containing neutral or acid oxalate anions) is indicative.

Experiments with annealing of sulfate sodalite-group minerals show that under high-temperature oxidizing conditions (in air, at 800 °C) S₃•- is the most stable sulfide species. Annealing of the preheated samples in air at 800 °C results in partial reverse transformations: S₂•- + S₂- + 2O₂(gas) \rightarrow SO₄²-, S₄•- + S₂•- \rightarrow 2S₃•-, as well as subordinate processes S₃•- + 5*e* +6O₂(gas) \rightarrow 3SO₄²- and C₂O₄²- \rightarrow 2CO₂(gas) + 2*e*. The oxidation of S₃•- into SO₄²- can proceed only partly due to the charge-balance requirement.

As noted above, S₃*- is a very strong blue chromophore and even trace amounts of this radical anion are detectable by Raman spectroscopy. However, S₃*- was not detected in sapozhnikovite. The possible cause of this fact is a high concentration of lithium in peralkaline rocks of the Lovozero massif (55 ppm, in average [66], whereas, e.g., in peralkaline rocks of the Khibiny massif, the mean Li content is only 20 ppm). Unlike other metal cations, Li⁺ reacts with S₃*- to form a cyclic radical anion LiS₃* with strong covalent Li–S bonds [67], which results in the immobilization of S₃*-. Kyanoxalite containing trace

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amounts of $S_3^{\bullet-}$ crystallized after sapozhnikovite at the hydrothermal zeolite stage with a lowered activity of lithium and containing trace amounts of $S_3^{\bullet-}$ [58].

4.3. Color Centers in Sodalite-group Minerals

Sodalite-group minerals show a wide range of colors. In particular, yellow, pink, violet, lilac, green, and blue samples are known. In most cases, coloration of sodalite-group minerals is caused by S-bearing extra-framework components. The $S_3^{\bullet-}$ radical anion is a very strong chromophore [45] which causes the deep blue color of lazurite [5]. Blue color of haüyne is usually due to trace amounts of $S_3^{\bullet-}$ [3,4,44,48]. The main yellow chromophore in sodalite-group minerals is the $S_2^{\bullet-}$ radical anion, which occurs in significant amounts in yellow varieties of bolotinaite (IMA 2021-088) earlier described as an "F-rich sodalite-group mineral" [4] as well as in above-described yellow products of heating of haüyne under reducing conditions.

Green chromophores are unknown among extra-framework components in sodalite-group minerals. Green color of some varieties of haüyne and slyudyankaite appears as a result of the presence of both S₃*- and a yellow chromophore (S₂*- and/or S₆). However, S₆ is a rather weak chromophore [41].

Pink color of sodalite-group minerals (in particular, sodalite and tugtupite) may have different causes, including the presence of S_4 or $S_4^{\bullet-}$ [41,45], Cr^{3+} or F-centers [68]. Lilac and violet tints of some haüyne samples from gem lazutite deposits (Sample 1 in this work; see [3,4]) is due to simultaneous presence of S_4 or $S_4^{\bullet-}$ and trace amounts of $S_3^{\bullet-}$.

Color coordinates in the CIE1931 color space chromaticity diagram for the samples studied in this work using NIR/Vis/UV spectroscopy are shown in Figure 22. Coordinates of coloration were calculated for daylight illuminant D with 4500 K temperature.

Lilac-blue color of Sample 1 may be partly due to the presence of two conformers of S4 that absorb mainly in the middle part of the visible range with band maximums shifted in opposite directions (towards blue and red ranges). However, the Raman spectrum of Sample 1 shows a series of characteristic bands of the S3• radical anion, which may be an additional cause of the blue tint.

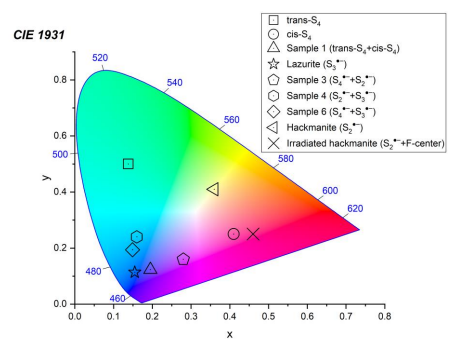


Figure 22. Color space chromaticity diagram for sodalite-group minerals.

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The color change of hackmanite and tugtupite upon irradiation with ultraviolet light from colorless or pale pink to purple is explained by the transfer of an electron from the impurity S_2^{2-} anions to the Cl vacancy, with the formation of color centers (F-centers). The absorption band of the F-centers is located at about 550 nm. The resulting S2*- radical anions practically do not affect the color due to their low concentration and relatively low oscillator strength. When irradiated with visible light, the F-centers disappear due to the phototransfer of an electron back to S₂•-, and the crystals become colorless. This reversible color change is called tenebrescence or photochromism. It is noteworthy that the presence of significant amounts of potassium in the mineral leads to the disappearance of photochromism: in hackmanite and tugtupite, where sodium sharply predominates among the extraframework cations, photochromism is observed, unlike haüyne, despite hauine usually containing a fairly large number of S2^{•-} centers. The causes of this are still being discussed. Perhaps this phenomenon is due to the fact that when sodium is replaced by potassium, either the value of the energy barrier for electron phototransfer from sulfur to a vacancy changes, or the excited state of the F-center enters the conduction band. Radiation coloring of haüyne does not change the visible color caused by F-centers. The presence of Fe³⁺ and Ti³⁺ impurities also affects the intensity of photochromic color and luminescence of S₂•- radical anions [69]

Other possible radical anions in minerals of the sodalite group are $SO_4^{\bullet-}$ μ $SO_3^{\bullet-}$. The most intense absorption band of $SO_4^{\bullet-}$ is located at about 450 nm and the second band is at about 300 nm in Na₂S₂O₈ [70]. It was noted in [71] that there is an electron transfer between Fe²⁺ μ SO₄^{\text{\cdot}-} with the formation of Fe³⁺ μ SO₄²⁻. Thus, the presence of SO₄^{\text{\cdot}-} in the mineral can lead to the loss of the blue color. The radical anion SO₃^{\text{\cdot}-} absorbs in the region of about 280 nm and does not affect the color [70].

5. Conclusions

Application of a complex of spectroscopic methods to sodalite-group minerals and products of their thermal conversions made it possible to identify various S- and C-bearing extra-framework components, including SO_4^{2-} , HS^- , S^{2-} , CO_3^{2-} , $C_2O_4^{2-}$, $HC_2O_4^-$, Cl^- , and F^- anions $S_2^{\bullet-}$, $S_3^{\bullet-}$, $S_4^{\bullet-}$, and $S_6^{\bullet-}$ radical anions as well as CO_2 , H_2O , and S_4 neutral molecules. The composition of extra-framework species in sodalite-group minerals may serve as a marker of mineral-forming media including fugacities of volatile components (H_2O , O_2 , CO_2 , HF, SO_2 , and polysulfide compounds). Complex mutual transformations of these components are observed during heating at high temperatures. The scheme of transformations of extra-framework components in SO_4^{2-} -bearing members of the sodalite group during their heating at 700 °C under reducing conditions includes the elementary processes $SO_4^{2-} \rightarrow S^{2-} + 2O_2(gas)$, $CO_2 + 2SO_4^{2-} + H_2O \rightarrow 2HS^- + CO_3^{2-} + 4O_2(gas)$. $3SO_4^{2-} \rightarrow S_3^{\bullet-} + 5e + 6O_2(gas)$, $2S_3^{\bullet-} \rightarrow S_2^{\bullet-} + S_4^{\bullet-}$, and $2CO_2 + 2e \rightarrow C_2O_4^{2-}$ (e = electron). Subsequent annealing in air at 800 °C results in the partial reverse transformations $S_2^{\bullet-} + S^{2-} + 2O_2(gas) \rightarrow SO_4^{2-}$, $S_4^{\bullet-} + S_2^{\bullet-} \rightarrow 2S_3^{\bullet-}$, $S_3^{\bullet-} + 5e + 6O_2(gas) \rightarrow 3SO_4^{2-}$, and $C_2O_4^{2-} \rightarrow 2CO_2(gas) + 2e$.

Based on these data, it can be concluded that natural HS⁻- and CO₂⁰-bearing sodalite-group minerals can be formed under reducing conditions and as a result of thermal transformation of an initial mineral, respectively. In this respect, the occurrence of the HS⁻-dominant mineral sapozhnikovite in association with the oxalate-rich cancrinite-type mineral kyanoxalite in agpaitic rocks [58] and the occurrence of a CO₃²--bearing sodalite-group mineral in a metasomatic volcanic ejectum [65] are indicative.

Normally, the oxalate group is unstable at temperatures higher than 500 °C. However, it was shown that the occurrence of oxalate anions in wide channels of cancrinite-group minerals is a stabilizing factor. In particular, oxalate anions in kyanoxalite decompose at 700 °C [72]. New data obtained in this work show that oxalate groups in sodalite cages are relatively stable up to 700 °C and decompose at 800 °C. Earlier we have shown that similar stabilization of nitrate anion takes place in cancrinite channels [73].

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Color variations of sodalite-group minerals are mainly caused by the presence of S-bearing chromophores, $S_3^{\bullet-}$ (blue), $S_2^{\bullet-}$ and/or S_6 (yellow), $S_4^{\bullet-}$ and/or S_4 (red). Green color is due to the simultaneous presence of blue and yellow chromophores.

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