Embreyite: structure determination, chemical formula and comparative crystal chemistry

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ABSTRACT

Embreyite from the Berezovskoe, Urals, Russia, was studied by the means of powder X-ray diffraction (XRD), single-crystal XRD, infrared spectroscopy and microprobe analysis. The empirical formula of embreyite obtained on the basis of microprobe analysis is $Pb_{1,29}Cu_{0,07}Cr_{0,52}P_{0,43}O_4$ (without taking into account the presence of H₂O). An examination of single-crystal XRD frames of the tested crystals cut from embreyite intergrowths revealed split reflection spots of weak intensities, even after a long exposure time. The crystal structure of embreyite (monoclinic, C2/m, a = 9.802(16), b = 5.603(9), c = 7.649(12) Å, $\beta = 114.85(3)^{\circ}$ and V = 381.2(11)Å³) has been solved by direct methods and refined to $R_1 = 0.050$ for 318 unique observed reflections. The powder XRD patterns of the holotype embreyite and the fresh material studied are close in both d values and the intensities match the pattern calculated from the structural single-crystal XRD data. The unit-cell parameters were recalculated for the holotype sample using a new cell setting and corresponding hkl indices. The crystal structure of embreyite is based on layers formed by corner-sharing mixed chromate-phosphate tetrahedra and PbO6 distorted octahedra. The interlayer space is filled by disordered Pb^{2+} and Cu^{2+} cations. Generally, the crystal structure of embreyite can be referred to the structural type of palmierite. $\{Pb[(Cr,P)O_4]_2\}$ layers in embreyite are similar in topology to those in yavapaiite-type compounds. The general formula of embreyite can be represented as $(Pb_{\lambda}M_{\nu}^{2+} \square_{1-x-\nu})_{2} \{Pb[(Cr,P)O_{4}]_{2}\}(H_{2}O)_{n}$, where $M^{2+} = Cu$ and Zn and $0.5 \le x+y \le 1$, or, in the simplified form: $(Pb,Cu,\square)_2 \{Pb[(Cr,P)O_4]_2\}(H_2O)_n$. The simplified formula of embrevite is similar in stoichiometry to vauquelinite and may explain the existence of the solid-solution series. The determination of the crystal structure of embreyite may also help to resolve the crystal chemical nature of cassedanneite. The XRD pattern of cassedanneite contains a distinct reflection with d = 13.9 Å, forbidden for the embrevite unit cell. This feature may indicate the doubling of the c unit-cell parameter of cassedanneite in comparison with embreyite. We assume that cassedanneite has structural similarity to embreyite with, presumably, a disordered distribution of Cr and V.

KEYWORDS: embreyite, layered structure, lead, chromate, phosphate, vauquelinite, cassedanneite, oxidation zone, Berezovskoe deposit, Urals.

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Introduction

EMBREVITE was described as a new mineral species in old museum collection samples from the Berezovskoe deposit, Central Urals, Russia by Williams (1972). [Note: Williams reports the locality as "Berezov, Siberia". According to old museum labels, geographical maps and 18-19th century literature, the Urals was typically included in Siberia]. This deposit, discovered in 1745, is located near the city of Ekaterinburg and is currently still being mined for gold. Berezovskoe is worldfamous for historical mineralogical objects because of outstanding, very rich and diverse chromate mineralization in the oxidation zone. Berezovskoe is the type locality of five Pb chromate minerals: crocoite (discovered in 1766), vauquelinite (1818), phoenicochroite (1833), embrevite (1972) and cassedanneite (1988). Crocoite was the first new mineral species described from Russia and chromium was discovered in samples of this mineral from Berezovskoe in 1797 by L.N. Vauguelin (see Pekov, 1998 and references therein).

The chemical composition, powder XRD and physical properties were well determined for the type material of embrevite. However, all of the crystals were too low quality for single-crystal X-ray studies. Crystals of the original embrevite were described as thin plates consisting of subindividuals oriented differently relative to each other and, additionally, split. The idealized formula of the mineral based on chemical data obtained by different methods was reported as Pb₅(CrO₄)₂ $(PO_4)_2$ ·H₂O. All of the analyses of the type material detected Pb²⁺, Cr⁶⁺, P, O and H as the main constituents. Cu²⁺ was also identified in the range of 1.2-2.5 (average 1.7) wt.% CuO (Williams, 1972). Additional recent studies of embrevite from the Berezovskoe ore field (several mines) and two other Ural localities (Mt. Bertevaya near the city of Nizhniy Tagil and Mt. Sukhovyaz within the city of Verkhniy Ufaley) revealed significant chemical variations (in wt.%): PbO 68.1-77.9, CuO 0.6-6.9, ZnO 0.0-1.9, P₂O₅ 6.4-9.7, As₂O₅ 0.0-2.6 and CrO₃ 9.1-15.0 (Kleymenov et al., 2003; Khanin et al., 2015). It was found that embrevite forms a continuous solid-solution series with vauquelinite, despite the difference in powder XRD patterns. Many of the recently obtained electron-microprobe analyses (EMPA) of embreyite are not in good agreement with the original idealized formula of $Pb_5(CrO_4)_2(PO_4)_2 \cdot H_2O$. The following generalized formula was suggested, taking into account all of the additional studies:

Pb₂[Pb_x M_y^{2+} , \Box_{1-x-y}]_{∑1}(CrO₄)(PO₄)(OH_{2(x+y)-1}, H₂O, \Box)_{∑1}, where M^{2+} = Cu or Zn, \Box = vacancy and 0.5 ≤ $x + y \le 1$; simplified as Pb₂(M, \Box)(CrO₄) (PO₄)X, where M = Pb, Cu or Zn and X = OH, H₂O or \Box (Khanin *et al.*, 2015). However, the latter formula could not be proven due to the lack of a single-crystal XRD study.

Embrevite is a member of a rather small group of lead-containing minerals with $(CrO_4)^{2-}$ anions (Table 1). Note that Pb chromates are the richest group for chromate minerals in general. Chromium mineral diversity is on the lower limit of the common trend as a function of the crustal abundance of Cr (Christy, 2015). A relatively small number of described chromate mineral species to date can be explained generally by the instability of the $(CrO_4)^{2-1}$ anion and the tendency of Cr⁶⁺ to reduce under conventional natural conditions. There have been several attempts to study the crystal structure of embrevite during the more than 40 years since its discovery. They were all unsuccessful due to the low quality of crystals. Crystals suitable for a structure determination were found in samples (Fig. 1) collected in 2011 at Berezovskoe by I.V.P. Herein, we report the determination of the structural features of embrevite and discuss relationships with the other Pb chromate minerals and synthetic phases.

Experimental

Sample description

The sample used in this study was found in the 'Krokoitovyi Shurf' (Crocoite Pit) at Mt. Uspenskava in the area of the former Tsvetnov Mine, the type locality of crocoite. This mine was operated from 1752 to 1802, and is situated in the central part of the Berezovskoe ore field. The richest secondary mineralization is located at a depth of ~ 12 m below surface. It is distributed in an area consisting mainly of partially weathered beresites and listwanites (Fettes and Desmons, 2007) with numerous quartz veins containing abundant cavities. Some of the latter are formed after dissolved pyrite, galena and 'fahlerz', as well as cavernous open-work polycomponent pseudomorphs consisting of the supergene minerals. The assemblage of secondary minerals associated intimately with embreyite examined in the course of this study includes crocoite, vauquelinite (two morphological and colour varieties: greenish-black spherical crystal clusters and olive-green finegrained crusts and kidney-shaped aggregates), pyromorphite, goethite (limonite) and

Mineral	Formula	Ref.
Cassedanneite	$Pb_5(VO_4)_2(CrO_4)_2 \cdot H_2O$ unknown crystal structure	[1]
Chromschieffelinite	$Pb_{10}Te_{\epsilon}O_{20}(OH)_{14}(CrO_4)\cdot 5H_2O$	[2]
Crocoite	$PbCrO_4$	[3, 4]
Embreyite	$Pb_2(Pb,Cu,\Box)[(Cr,P)O_4]_2(H_2O,OH,\Box)]$	[5, 6, *]
Fornacite	$Pb_2Cu(CrO_4)(AsO_4)(OH)$	[6-9]
Georgerobinsonite	$Pb_4(CrO_4)_2(OH)_2FCl$	[10]
Hemihedrite	$Pb_{10}Zn(CrO_4)_6(SiO_4)_2F_2$	[8, 11, 12]
Iranite	$Pb_{10}Cu(CrO_4)_6(SiO_4)_2(OH)_2$	[8, 13–15]
Macquartite	$Pb_7Cu_2(CrO_4)_4(SiO_4)_2(OH)_2$ no single crystal X-ray data	[16, 17]
Phoenicochroite	$Pb_2O(CrO_4)$	[8, 18]
Reynoldsite	$Pb_2Mn_2O_5(CrO_4)$	[19]
Santanaite	$Pb_9^{2+}Pb_2^{4+}O_{12}(CrO_4)$ unknown crystal structure	[20]
Vauquelinite	$Pb_2Cu(CrO_4)(PO_4)(OH)$	[6, 8, 21]

TABLE 1. List of Pb chromate minerals.

References: [1] – Cesbron *et al.* (1988); [2] – Kampf *et al.* (2012*a*); [3] – Quareni and de Pieri (1965); [4] – Effenberger and Pertlik (1986); [5] – Williams (1972); [6] – Khanin *et al.* (2015); [7] – Cocco *et al.* (1967); [8] – Cesbron and Williams (1980); [9] – Ksenofontov *et al.* (2014); [10] – Cooper *et al.* (2011); [11] – Williams and Anthony (1970); [12] – McLean and Anthony (1970); [13] – Bariand and Herpin (1963); [14] – Adib and Ottemann (1970); [15] – Yang *et al.* (2007); [16] –Williams and Duggan (1980); [17] – Cooper and Hawthorne (1994); [18] – Williams *et al.* (1970); [19] – Kampf *et al.* (2012*b*); [20] – Mücke (1972); [21] – Fanfani and Zanazzi (1968); * – this work.

undetermined clay mineral(s). Embreyite occurs as flattened, coarse hexagonal, typically roundish, disc-like crystals up to 0.3 mm and, rarely, up to 1 mm across (Fig. 1). They are usually divergent,



FIG. 1. Aggregates of brown-orange embreyite crystals (in the centre) with crocoite (bright orange prismatic crystals) and pyromorphite (yellow-green) around. Field of view: width = 5.8 mm. Photo: A.V. Kasatkin.

resembling an open book, and combined in groups or crusts up to 3 mm × 3 mm and up to 1 mm thick. The mineral is orange to yellow orange, sometimes brownish-yellow, with a strong greasy lustre. Embreyite is one of the latest minerals of this assemblage overgrowing crocoite and pyromorphite.

Chemical composition

The chemical composition was determined for the embreyite crystal used for the structure determination on a Jeol 733 electron microprobe instrument operating in energy-dispersive mode with an accelerating voltage of 20 kV, a beam current of 2 nA and a beam diameter of 5 μ m. The X-ray acquisition live-time was 30 s. The following standards were used: PbTiO₃ (Pb), Cu metal (Cu), chromite USNM 117075 (Cr) and LaPO₄ (P). The average (four spot analyses) chemical composition (wt.%, ranges in parentheses) is: PbO 74.46 (73.79–75.04), CuO 1.38 (1.18–1.51), CrO₃ 13.40 (13.06–13.71) and P₂O₅ 7.96 (7.78–8.10),

total 97.20. Contents of other elements with atomic numbers higher than carbon are below detection limits. The H_2O content was not determined because of paucity of the material. The empirical formula calculated on the basis of 4 O atoms per formula unit (apfu) (without taking into account the possible presence of some H_2O : see below) is: $Pb_{1,29}Cu_{0,07}Cr_{0,52}P_{0,43}O_4$.

Infrared spectroscopy

In order to obtain the infrared (IR) absorption spectrum (Fig. 2), a powdered sample of embreyite was mixed with dried KBr, pelletized and analysed using a Bruker ALPHA FTIR spectrometer with a resolution of 4 cm⁻¹ and 16 scans. The IR spectrum of a pellet of pure KBr was used as a reference.

Note, most of the Pb–O vibrations are expected to be in the region below 300 cm⁻¹ and were not registered in the region of the IR spectra studied. Absorption bands in the IR spectrum of embreyite and their assignments are (cm⁻¹; s – strong band, sh – shoulder): 3329 and diffuse absorption between 3100 and 3300 [O–H stretching vibrations of H₂O molecules], 1680w [bending vibrations of H₂O molecules], 1680w [bending vibrations of H⁴ cations], 1030sh, 959s [v₃(F₂) – asymmetric P–O stretching vibrations of PO₄⁻⁻ anions], 900sh [v₁(A₁) – symmetric P–O stretching vibrations of PO₄³⁻⁻ anions], 855s, 830sh [v₃(F₂) – asymmetric Cr–O stretching vibrations of CrO₄²⁻⁻ anions], 560sh, 538 [triply degenerate v₄(F₂) O–P–O bending mode of PO_4^{3-} anions], 418w, 383 [lattice modes involving $v_2(E)$ O–P–O bending vibrations, possibly combined with vibration modes of H₂O molecules]. The band assignments were done according to Nakamoto (2009) and Chukanov and Chervonnyi (2016).

Powder XRD

Powder XRD data of embreyite were collected with a Rigaku R-AXIS Rapid II single-crystal diffractometer equipped with cylindrical image plate detector using Debye-Scherrer geometry (d=127.4 mm, CoK α radiation and $\lambda=1.7890$ Å). The powder XRD pattern and unit-cell parameters refined from the powder data are given in Table 2 and compared with the data for the holotype sample reported by Williams (1972).

Single-crystal XRD

Single crystals of embreyite were examined under an optical microscope, mounted on glass fibers with epoxy and tested on a Bruker APEX DUO diffractometer equipped with a micro-focus X-ray tube operated with MoK α radiation ($\lambda = 0.71073$ Å) at 50 kV and 40 mA. An examination of XRD frames of the tested crystals cut from embreyite intergrowths revealed split reflection spots of weak intensities, even after a long exposure time. The quality of the crystals of embreyite made the solution of the crystal structure challenging. An orange translucent platy crystal, 0.14 mm ×



FIG. 2. Infrared (FTIR) spectrum for the embreyite sample studied in this work.

This w	vork	Ho spec	lotype cimen*	This work**		1. 1. 1
I _{obs}	$d_{\rm obs},$ Å	I _{obs}	$d_{\rm obs}, {\rm \AA}$	d_{calc} , Å	Icalc	пкі
6	6.92	3	6.941	6.941	2	001
48	4.738	60	4.751	4.763, 4.741	25, 40	2 01, 110
8	4.377	3	4.378	4.401	1	Ī11
15	3.550	32	3.563	3.561, 3.555	11, 4	$111, \bar{2}02$
15	3.471	28	3.475	3.470	10	002
100	3.157	100	3.167	3.185, 3.158	54, 100	$201, \bar{1}12$
44	2.813	60	2.818	2.822, 2.801	55, 24	311,020
8	2.599	17	2.608	2.620, 2.609, 2.598	4, 3, 4	310, 312, 021
4	2.547			2.542	1	112
1	2.409	4	2.407	2.415	1	<u>2</u> 21
4	2.320	23	2.314	2.314	3	003
8	2.254	32	2.213	2.224	8	400
14	2.194	31	2.187	2.200, 2.197, 2.183, 2.180	12, 3, 5, 3	<u>-</u> 222, 311, <u>3</u> 13, 022
19	2.099	31	2.105	2.105, 2.104	7, 12	4 03, 221
24	1.910	45	1.917	1.917, 1.911	14, 7	113, 204
9	1.848			1.842	2	512
4	1.833			1.838, 1.828	1, 1	4 21, 130
3	1.795			1.795	2	312
7	1.780	26	1.789	1.784, 1.784, 1.780, 1.777	1, 1, 2, 1	023, 314, 222, 404
2	1.760			1.762	1	ī14
2	1.735			1.735	1	004
6	1.692			1.695	4	510
5	1.681			1.683, 1.678	3, 2	4 23, 1 32
2	1.641			1.634	1	<u></u> 602
3	1.617			1.621	2	331
7	1.577			1.580, 1.579, 1.578	1, 3, 2	330, 224, 332
1	1.497			1.501, 1.499	1, 1	<u>4</u> 24, <u>1</u> 33
3	1.410			1.411	2	<u>-</u> <u>6</u> 22
4	1.403			1.401	1	040
4	1.386			1.385	1	422
5	1.378			1.378, 1.377, 1.373	1, 1, 1	515, 133, 041
4	1.335			1.342, 1.339, 1.336	1, 1, 1	531, 601, 240
2	1.325			1.325	2	334
4	1.306			1.307, 1.303	1, 1	<u>5</u> 33, <u>2</u> 42
Unit-c	ell parameters					
Calcu	lated from	Calcul	ated from	Determined by single crysta	ıl XRD data	
pow	der XRD	powe	ler XRD			
	data	(data			
9.	814(6)	9.	75(1)	9.802(16)		<i>a</i> , Å
5.	597(1)	5.6	504(4)	5.603(9)		<i>b</i> , Å
7.	643(5)	7.6	535(9)	7.649(12)		<i>c</i> , Å
11-	4.79(4)	114	1.48(9)	114.85(3)		β, °
38	31.1(4)	37	9.8(9)	381.2(11)		<i>V</i> , Å ³

TABLE 2. P	Powder XRD	data and	unit-cell	parameters	of embreyite	Э.
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*Holotype specimen: I_{obs} and d_{obs} values are after Williams (1972) and unit-cell parameters provided in this table were refined using the *hkl* indices of our calculated XRD pattern (the unit-cell data reported by Williams (1972) are: a = 9.755(3), b = 5.636(3), c = 7.135(3) Å, $\beta = 103.8(3)^{\circ}$ and V = 382 Å³) **XRD pattern calculated from our crystal-structure data. 0.12 mm × 0.07 mm demonstrating the best diffraction pattern (Fig. 3) was chosen for full XRD collection. More than a hemisphere of XRD data were collected with a frame width of 0.5° in ω and 60 s counting time for each frame. The data were integrated and corrected for absorption using a multi-scan-type model using the Bruker programs *APEX* and *SADABS*.

The unit-cell parameters of embrevite were determined and refined by the least-squares technique on the basis of 1334 reflections with 2θ in the range of 8.6-57.4°. The obtained unit-cell parameters (Table 2, 3) are in a good agreement with those previously reported in the $P2_1/m$ space group by Williams (1972). The $|E^2-1|$ parameter was equal to 0.899, which indicated high probability of a centrosymmetric space group confirmed by subsequent structure solution and refinement. The crystal structure of embrevite was solved in C2/musing direct methods and refined to $R_1 = 0.050$ by means of the SHELX program package (Sheldrick, 2015). Crystallographic information is summarized in Table 3. Attempts to solve and refine the crystal structure in the higher symmetry space group, $R\bar{3}m$ $(a = 5.618(9), c = 20.80(4) \text{ Å and } V = 569(2) \text{ Å}^3),$ reported for the structure of rhombohedral polymorph of $Pb_4(PO_4)_2(CrO_4)$ (Barbier and Maxin, 1995) led us to a worse agreement between the crystallographic model and the experimental XRD data $(R_1 = 0.072)$ and in addition significant disorder in the heteropolyhedral layer (see below) not observed in C2/m. Various twin laws in both space groups were tried but none of them appeared



FIG. 3. Typical XRD frame of embreyite collected with width = 0.5° and 60 s of exposure.

TABLE 3. Crystallographic data for embreyite.

Crystal data	
Crystal system	monoclinic
Space group	C2/m
Unit-cell dimensions	
a (Å)	9.802(16)
b (Å)	5.603(9)
c (Å)	7.649(12)
α, β, γ (°)	90, 114.85(3), 90
Unit-cell volume (Å ³)	381.2(11)
Ζ	1
Calc. density $(g \cdot cm^{-3})$	6.564
Absorption coef. (mm^{-1})	58.99
Crystal size (mm)	0.14 imes 0.12 imes 0.07
Data collection	
Temperature (K)	293
Radiation, wavelength (Å)	ΜοΚα, 0.71073
<i>F</i> (000)	636
θ range (°)	4.3-28.7
h, k, l ranges	$-11 \le h \le 12$
-	$-7 \le k \le 5$
	$-10 \le l \le 5$
Total refl. collected	1334
Unique refl. (R_{int})	500 (0.0471)
Unique refl. $F > 4\sigma(F)$	318
Structure refinement	
Refinement method	Full-matrix least-squares on F^2
Weighting coef. a, b	0.0794, 0
Data/restraints/parameters	500/0/55
$R_1 [F > 4\sigma(F)],$	0.0505
$wR_2 [F > 4\sigma(F)]$	0.1278
R_1 all, wR_2 all	0.0845, 0.1435
Goof on \tilde{F}^2	1.006
Largest diff. peak and hole $(e \text{ Å}^{-3})$	1.358, -0.943

to be successful and improved the refinement. Therefore, we provide here a description of the structure of embreyite in the monoclinic space group C2/m as it resulted in a lower *R*-factor, very good fit with powder XRD patterns, better displacement parameters of atoms, and physically realistic interatomic bond-distance values.

A procedure of the structure refinement of embreyite included the following steps: (1) an initial group of the atom sites was determined by direct methods; (2) missing atoms were found from the difference-Fourier maps; (3) the structural model was refined until the atomic arrangements became reasonable; and (4) site occupancies were adjusted in accordance with the chemical data. All positions except the low occupied Cu one (siteoccupation factor = 0.132) were refined

Site	Atom	Wyck.	SOF	x	У	Ζ	$U_{\rm iso}^{}*\!/U_{\rm eq}^{}$
Pb1*	Pb	2d	1	1/2	0	1/2	0.0891(7)
Pb2A**	Pb	4i	0.308	0.7919(5)	1/2	0.8693(4)	0.0729(8)
Pb2B	Pb	8j	0.189	0.7681(3)	0.6375(4)	0.9348(4)	0.0449(7)
Pb2C	Pb	4i	0.108	0.9074(7)	1/2	0.9379(8)	0.0372(12)
Cu1	Cu	2b	0.132	0	1/2	0	0.061(7)*
T^{**}	Cr	4i	0.520	0.9014(3)	0	0.7033(4)	0.0355(8)
	Р		0.432				
01	Ο	8 <i>j</i>	1	0.0009(10)	0.2324(14)	0.7839(12)	0.055(2)
O2	Ο	4i	1	0.7725(14)	0	0.7787(19)	0.057(3)
O3	0	4i	1	0.8271(15)	0	0.4762(16)	0.071(4)

TABLE 4. Fractional atomic coordinates, site occupancy factors (SOF), bond-valence sums (vu) and equivalent or isotropic displacement parameters (Å²) of atoms in the structure of embreyite.

*Bond valence sum is 2.14 vu.

**Overall weighted bond-valence sums for disordered sites: Pb2 = 1.84 vu, T = 5.52 vu. Wyck – Wyckoff positions.

anisotropically. Final atom coordinates and siteoccupation factors are represented in Table 4. Selected bond distance values are in Table 5.

Results

Cation coordination

There are one fully occupied Pb1 site, one tetrahedrally coordinated T site and four oxygen sites in the heteropolyhedral layer in the crystal structure of embreyite. In the disordered interlayer, there is one split Pb2 site and one low occupancy Cu1 position.

Heteropolyhedral layer

The Pb1 site adopts a 6-fold coordination in the first coordination sphere with six short and strong Pb1–O bonds with the average < Pb1–O₆> bond distance of 2.63 Å (Table 5). The resultant Pb1O₆ polyhedron can be described as a distorted octahedron (Fig. 4*a*). There are six more additional oxygen atoms located at larger distances from 3.24 to 3.29 Å thus forming distorted Pb1O₁₂ cuboctahedron. The average < Pb1–O₁₂> bond length of the 12-coordinated Pb1-centred polyhedron is 2.94 Å.

One symmetrically independent tetrahedral *T* site is surrounded by four O atoms (Fig. 4*b*). Calculations of the mineral formula on the basis of O = 4 apfu provided the sum of tetrahedral cations $(Cr_{0.52}^{6+} \text{ and } P_{0.43}^{5+})$ is < 1. The latter prompted us to consider this site as partially occupied. Unconstrained refinement of the occupancies of

the T site led to $(Cr_{0.520}P_{0.432}\square_{0.048})$. Structural containing partially models occupied tetrahedral sites were reported previously for minerals and synthetic compounds, e.g. in byzantievite, Ba₅(Ca,REE,Y)₂₂(Ti,Nb)₁₈(SiO₄)₄[(PO₄), $(SiO_4)]_4(BO_3)_9O_{21}[(OH),F]_{43}(H_2O)_{1.5}$ (REE =rare-earth minerals) (Sokolova et al., 2010) and synthetic $AMo_3P_{5,8}Si_2O_{25}$ ($A = Rb \text{ or } Tl^+$) (Leclaire et al., 1984), where a deficiency of P^{5+} is observed in PO₄ groups to make the formula electroneutral. To the best of our knowledge the crystal structure of embrevite is the first observation in minerals of a

TABLE 5. Selected interatomic distances in Å in the structure of embreyite.

Pb1-O2	2.625(13) × 2	Cu1-O1	$2.234(9) \times 4$
Pb1-O1	2.635(9) × 4		
<pb1-o<sub>6></pb1-o<sub>	2.632	Pb2A-Pb2B	0.998(3)
Pb1-O3	$3.240(8) \times 4$	Pb2A-Pb2C	1.029(6)
Pb1–O3 ⁱ	3.286(16) × 2	Pb2A-Cu1	1.856(5)
<pb1-o<sub>12></pb1-o<sub>	2.943		
12		T-O3	1.576(12)
Pb2A-O3	2.399(12)	T-01	$1.588(9) \times 2$
Pb2A-O1	$2.828(10) \times 2$	T-O2	1.594(14)
Pb2A-O2	$2.873(5) \times 2$	< <i>T</i> -O ₄ >	1.587
Pb2A–O1 ⁱⁱ	2.948(11) × 2		
Pb2A–O1 ⁱⁱⁱ	$2.980(10) \times 2$		
Pb2A–O2 ^{iv}	3.013(15)		
$< Pb2A - O_{10} >$	2.867		

Symmetry codes: (i) -x+1, -y, -z+1; (ii) $x + \frac{1}{2}$, $-y + \frac{1}{2}$, z; (iii) -x+1, -y+1, -z+2; (iv) -x+3/2, $-y + \frac{1}{2}$, -z+2.



FIG. 4. Geometry of the cationic sites in the crystal structure of embreyite. Displacement ellipsoids are drawn at the 50% probability level.

mixed tetrahedral site occupied by Cr^{6+} and P^{5+} . The bond-valence sum (BVS) calculations for the *T* site confirm a mixed 5+/6+-valence occupancy providing the weighted bond-valence sum of 5.52 valence units (vu). The average < T-O> bond length value is 1.59 Å.

Disordered interlayer

The Pb2 site is split over Pb2A. Pb2B and Pb2C positions in the crystal structure of embreyite. Disorder of Pb atoms in the interlayer is rather typical for lead oxysalt layered structures and has been reported e.g. for hydrocerussite (Martinetto et al., 2002), rickturnerite (Rumsey et al., 2012) and $Pb_{21}[Si_7O_{22}]_2[Si_4O_{13}]$ (Siidra *et al.*, 2014*b*). Refinement of the occupancies of the three Pb2 sites with the total occupancy of 0.605 revealed the presence of a vacancy here. Most probably, it can be filled by H₂O molecules evidenced by the presence of typical absorption bands at ~3300 and $\sim 1600 \text{ cm}^{-1}$ in the IR spectrum (Fig. 2). We suggest that the disordered Pb2 sites are the only possible sites for H₂O in the structure of embrevite as we were unable to find other prominent peaks on difference-Fourier maps that could be interpreted as

positions of water molecules. A similar example of complicated water site localization has been reported recently for gianellaite, $[(NHg_2)_2](SO_4)$ $(H_2O)_x$ (Cooper *et al.*, 2016) where H_2O could not be localized in the latter from structural data. However, the presence of water was undoubtedly indicated by absorption bands in the IR spectra.

The most electron-dense Pb2A site is symmetrically coordinated by ten O atoms located at distances varying from 2.40 to 3.01 Å with the average < Pb2A– O_{10} > bond distance of 2.87 Å (Fig. 4*c*).

The chemical analysis reveals $Cu_{0.07}^{2+}$ pfu. A small amount of copper was assigned to the Cu1 site, which can be also considered as one of the Pb2 split positions. Geometrical analysis demonstrates square coordination environments for the Cu1 site (Fig. 4*d*) with oxygen atoms located at a distance of 2.23 Å. However, these long bond-distance values should not be considered literally due to the strong disorder. We suggest that the Cu1 site is the most favoured position for a small amount of Cu in the crystal structure of embreyite because of its typical square coordination environment and structural relation with vauquelinite, discussed below.

The final refinement of the structure of embreyite led to the structural formula of $[Pb_{0.794}Cu_{0.066}]$

 $\begin{array}{l} \label{eq:constraint} & $ \Box_{0.140-x}] \{ [Pb_{0.500}] [(Cr_{0.520}P_{0.432}\Box_{0.048})O_4)] \} (H_2O)_n$ (the presence of H_2O is assumed by IR data), which is in a good agreement with the electron microprobe data on the basis of 4 O atoms per formula unit, i.e. <math>(Pb_{0.79(1)}Cu_{0.07(2)})_{\Sigma 0.86}(Pb_{0.50(2)})_{\Sigma 0.5}$ $(Cr_{0.52(2)}P_{0.43(2)})_{\Sigma 0.95}O_4.$

Crystal structure: description and comparison with palmierite-type compounds

The crystal structure of embrevite is based on {Pb $[(Cr,P)O_4]_2]$ layers formed by corner-sharing mixed chromate-phosphate tetrahedra and Pbcentred polyhedra (Fig. 5a). The interlayer space is filled by disordered Pb^{2+} and Cu^{2+} cations. Generally, the crystal structure of embrevite can be referred to the structural type of palmierite, K₂Pb (SO₄)₂ (Moore, 1973; Tissot *et al.*, 2001) built up from stacking of layers parallel to (001) and formed by corner-sharing distorted Pb2+ polyhedra and SO_4 tetrahedra with K⁺ cations in the interlayer. A variety of cations in different oxidation states are involved in homovalent or heterovalent substitutions observed in the large number of palmieritetype structures with different structural distortions (Lazoryak, 1996).

It is worth considering the low-temperature rhombohedral modification of Pb₄(PO₄)₂(CrO₄) chemically close to embrevite that has been studied by neutron powder diffraction (Barbier and Maxin, 1995). The rhombohedral structure of $Pb_4(PO_4)_2(CrO_4)$ ($R\bar{3}m$, a = 5.5403(1), c =20.4999(4) Å and V = 544.94(1) Å³) is based upon stacking along the c axis of layers composed of (P, Cr)O₄ and PbO₁₂ polyhedra with partially occupied Pb²⁺ sites in the interlayer space, similar to the palmierite-type monoclinic structure of embrevite (Fig. 5b). The structural formula of the rhombohedral modification of Pb₄(PO₄)₂(CrO₄) according to the published data can be written as $2 \times [Pb_{0.83} \square_{0.17}]$ $[Pb_{0.5}(PO_4)_{0.67}(CrO_4)_{0.33}]$, similar to the one determined for embreyite, $[Pb_{0.794}Cu_{0.066} \square_{0.140-x}]$ $\{[Pb_{0.500}][(Cr_{0.520}P_{0.432}\square_{0.048})O_4)]\}(H_2O)_n$. Another palmierite-type compound with disordered cation sites is Sr_{2.67} $\Box_{0.33}(PO_4)_{1.33}(CrO_4)_{0.67}$ (Hartl and Braungart, 1978*a*,*b*). It has a large supercell with a doubling of both the *a* and *c* edges ($R\bar{3}m$, a = 10.91and c = 39.78 Å) which most probably results from ordering of vacancies in the crystal structure. Note that, palmierite-type (Pb,Ba)₃(PO₄)₂ phases structurally related to embrevite were studied previously as ferroelastic materials exhibiting the $R\bar{3}m$ to C2/cferroelastic transition around $T=180^{\circ}$ C, and were considered as model compounds for ferroelastic distortions (Bismayer and Salje, 1981; Aktas *et al.*, 2013; Salje, 2015).

Discussion

Lead chromate minerals

There are 13 lead chromate minerals approved by the International Mineralogical Association to date (Table 1). The crystal structures of cassedanneite $Pb_5(VO_4)_2(CrO_4)_2 \cdot H_2O$ (Cesbron *et al.*, 1988) (see below) and santanaite $Pb_9^{2+}Pb_2^{4+}O_{12}(CrO_4)$ (Mücke, 1972) remain unknown to date. Some of those studied by single-crystal XRD gave poor quality structure refinements for various different reasons: fornacite had $R_1 = 0.10$ (Cocco *et al.*, 1967); reynoldsite, $R_1 = 0.10$ (Kampf *et al.*, 2012b); and vauquelinite, $R_1 = 0.09$ (Fanfani and Zanazzi, 1968). Macquartite, $Pb_7Cu_2(CrO_4)_4$ $(SiO_4)_2(OH)_2$ (Williams and Duggan, 1980) was not studied by single-crystal XRD but assumed to be isostructural with wherryite, Pb7Cu2(SO4)4 $(SiO_4)_2(OH)_2$ (Cooper and Hawthorne, 1994). Lead-containing minerals or synthetic compounds with bichromate $(Cr_2O_7)^{2-}$ groups are unknown. The CrO₄ tetrahedra are isolated in all Pb chromate mineral crystal structures. Most of these minerals (except crocoite, georgerobinsonite and phoenicochroite) contain AO_6 polyhedra with different types of connectivity modes with CrO₄ tetrahedra via common oxygen atoms. CrO₄ groups are monodentate with CuO₆ octahedra in the structure of fornacite and vauquelinite. The structures of hemihedrite and iranite contain CrO₄ groups shared with ZnO₆ (hemihedrite) or CuO₆ (iranite) octahedra also in monodentate mode. However, chromate groups are only shared with Pb-centred polyhedra and do not have common oxygen vertices with TeO₆ and MnO₆ octahedra in the structures of chromschieffelinite and reynoldsite, respectively.

Formula of embreyite

The structural formula obtained for the embreyite sample studied, $[Pb_{0.794}Cu_{0.066} \square_{0.140-x}] \{ [Pb_{0.500}] [(Cr_{0.520}P_{0.432} \square_{0.048})O_4)] \} (H_2O)_n$, is in general agreement with that assumed earlier for this mineral on the basis of different data, $Pb_2[Pb_x M_y^{2+} \square_{1-x-y}]_1(CrO_4)(PO_4)(OH_{2(x+y)-1}, H_2O, \square)_{\Sigma_1}$, where $M^{2+} = Cu$, Zn and $0.5 \le x + y \le 1$, or $Pb_2(M, \square)(CrO_4)(PO_4)X$, where M = Pb, Cu and Zn and X = OH, H_2O and \square (Khanin *et al.*, 2015). The



FIG. 5. General views of the crystal structures of (a) embreyite, (b) synthetic rhomb-Pb₄(PO₄)₂CrO₄ and (c) vauquelinite. Projection of the interlayer block of the structure of vauquelinite on the structure of embreyite is shown in (d).

major discrepancy between them is in the presentation of the tetrahedrally coordinated constituents: the structure determination revealed Cr-P disorder. The results of the structure determination suggest attributing the M^{2+} cations to the second disordered Pb position. Thus, the generalized formula of embreyite can be represented as $(Pb_x M_y^{2+} \square_{1-x-y})_2$ $\{Pb[(Cr,P)O_4]_2\}(H_2O)_n$, where $M^{2+} = Cu$, Zn and $0.5 \le x + y \le 1$, or, in simplified form: (Pb, $Cu, \square)_2 \{Pb[(Cr, P)O_4]_2\}(H_2O)_n$. Minor amounts of vacancies in the tetrahedral site are ignored in the formula for simplification. The powder XRD patterns of the holotype embreyite (Williams, 1972) and our material are close in both d values and intensities of reflections. This observation became a reason to re-calculate unit-cell parameters for the holotype sample (due to minor differences in the unit-cell of our sample in the *c* dimension and β angle values) using our cell setting and corresponding *hkl* indices. Unit-cell parameters and volume obtained for the holotype are very close to those of the embreyite sample described in this present investigation. We recommend this unit-cell setting (with $\beta \approx 114-115^{\circ}$ instead of the original $\beta \approx 104^{\circ}$) and *hkl* indices for the reflections from powder XRD as given in Table 2 for embreyite in general.

Comparison with yavapaiite-type minerals and synthetic compounds

The layers of $\{Pb[(Cr,P)O_4]_2\}$ in embreyite are similar in topology to those in yavapaiite-type

brianite $Na_2Ca[Mg(PO_4)_2]$ compounds, e.g. (Alkemper and Fuess, 1998) and steklite KAl $(SO_4)_2$ (Murashko *et al.*, 2013). The tetrahedron acts as a tridentate-bridging ligand thus leaving its fourth vertex (O_t) as terminal or non-shared and in orientation towards the interlayer space in yavapaiite (Fig. 6a). O, are directed into the layer (Fig. 6b) in embrevite. O, atoms provide the linkage of heteropolyhedral layers with interlayer cations in vavapaiite. However all three O atoms of basal triangular planes of tetrahedra are involved in strong bonding with interlayer cations in embrevite. In this sense the structural architecture of heteropolyhedral layers in embrevite is similar to those of bütschliite-type compounds (Fig. 6c), e.g. K₂Mg $(CO_3)_2$ (Hesse and Simons, 1982) and KBaY $(BO_3)_2$ (Gao *et al.*, 2011), where the tetrahedra are replaced by triangles of carbonate or borate groups with a similar method of bonding with interlayer species. Another type of layer related to heteropolyhedral blocks in embrevite was described previously in the structures of markhininite, TlBi $(SO_4)_2$ (Siidra et al., 2014a) and synthetic, RbEu $(SO_4)_2$ (Sarukhanyan *et al.*, 1983). A^{3+} -centred (A = Bi or REE^{3+}) polyhedra share edges with adjacent SO₄ tetrahedra in contrast to the other yavapaiitetype compounds with corner-sharing octahedra and tetrahedra in heteropolyhedral blocks. Such coordination of sulfate groups leads to more densely packed layered units and, consequently, orientation of the fourth oxygen vertex towards the layer (Fig. 6d). Note, there is also a group of Se-



FIG. 6. Heteropolyhedral yavapaiite-related layers in minerals and synthetic compounds: (*a*) yavapaiite KFe(SO₄)₂, (*b*) embreyite, (*c*) bütschliite $K_2Ca(CO_3)_2$, (*d*) markhininite TlBi(SO₄)₂, (*e*) $K_2Mn(SeO_3)_2$ and (f) RbSc(SeO₃)₂. Black arrows designate the orientation of TO_4 tetrahedra or SeO₃ groups in a direction towards or outwards from the layer. See the text for details.

containing phases related to both yavapaiite- and bütschliite-type compounds. Apical vertices of triangular selenite Se⁴⁺O₃ pyramids are oriented into the centre of the layers identically to (Cr,P)O₄ groups in embreyite (Fig. 6*e*) in the crystal structures of K₂*M*(SeO₃)₂ compounds [M^{2+} = Mg (Hesse and Simons, 1982), Co (Wildner, 1992*a*) and Mn (Wildner, 1992*b*)], whereas the apices of the pyramids are aligned one by one in the same direction emphasizing a non-centrosymmetric character in β-PbNi(SeO₃)₂ (Kovrugin *et al.*, 2015) and RbSc(SeO₃)₂ (Song and Ok, 2015) (Fig. 6*f*).

Comparison with vauquelinite

It is of interest to examine and compare the chemical features and structural architectures of embrevite with vauquelinite, ideally $Pb_2Cu(CrO_4)$ (PO₄)(OH) (Fanfani and Zanazzi, 1968; Cesbron and Williams, 1980), as they form a continuous solid-solution series with the Pb:Cu ratio as the main varying value (Khanin et al., 2015) regardless of significant structural differences. The data from EMPA and powder XRD demonstrate the existence of the margin between these phases as approximately between 0.5 and 0.6 apfu of Cu in the idealized formula of vauquelinite. The recommended simplified formula of embreyite, (Pb,Cu, \square)₂{Pb[(Cr,P)O₄]₂}(H₂O)_n, demonstrates definite similarity in stoichiometry with the vauquelinite formula and explains the existence of the solidsolution series. The Pb:Cu ratio in the point of nucleation of a crystal in a phosphate-chromate mineral-forming system may determine a mineral form (embrevite/vauquelinite) to be crystallized. This may explain the cause and mechanism of the formation of botryoidal aggregates and encrustations up to several mm thick composed by alternating layers of vaquelinite and embrevite (Fig. 7). Such interstratified bi-mineral aggregates are very typical for the chromate assemblages at the Berezovskoe deposit (Williams, 1972; Kleymenov et al., 2003; Khanin et al., 2015). We believe that they may form as a result of oscillatory crystallization processes in the systems chemically close to the margin between the stability fields of embrevite and vauquelinite defined by the Pb:Cu ratio.

Vauquelinite and embreyite demonstrate different, but related, structure types. The former is very close structurally to brackebuschite-group minerals with a crystal structure (Cesbron and Williams, 1980) based on columns of edge-sharing $CuO_4(OH)_2$ octahedra oriented along the *b* axis. CrO₄ and PO₄ tetrahedra are shared with octahedra via monodentate bridging of one of four oxygen vertices. Pb²⁺ cations are located in between the columns and provide three-dimensional integrity of the structure. A closer look at the coordination environments of one of two Pb sites in vauquelinite reveals similarity with the Pb1-centred coordination polyhedron in embrevite. It can be described as a distorted cuboctahedron with six short Pb-O distances (2.40-2.94 Å) and six longer ones (3.01-3.63 Å). PbO_{12} , PO_4 and CrO_4 polyhedra form heteropolyhedral layers in vauquelinite similar to those in embrevite. Additional Cu2+ and Pb2+ cations provide the linkage of these blocks (Fig. 5c) in vauquelinite. The common structural motif of vauquelinite and embrevite suggests the possible location of a small amount of Cu in the structure of the latter in the interlayer space. Embrevite can be considered as an intermediate phase between the structures of vauguelinite and rhombohedral modification of $Pb_4(PO_4)_2(CrO_4)$ (unknown as a mineral), where the disordered interlayer species are represented by a superposition of ordered cations in two different structures related to each other by translation (Fig. 5d).

There is also a link between the lattice dimensions of these minerals (Table 6). The unit-cell parameters of embreyite are close to those of the brackebuschite-related minerals, e.g. tsumebite $Pb_2Cu(PO_4)(SO_4)(OH)$ and arsentsumebite Pb_2Cu (AsO₄)(SO₄)(OH) (Fanfani and Zanazzi, 1968; Zubkova *et al.*, 2002). Remarkably, vauquelinite



FIG. 7. Back-scattered electron image showing rhythmically zoned crust consisting of embreyite (Em) and vauquelinite (Va) layers. Berezovskoe deposit, Urals, Russia. Sample #10393 in the systematic collection of Fersman Mineralogical Museum of the Russian Academy of Sciences, Moscow. Field of view: width = 1.9 mm.

Mineral	Formula	Sp. gr.	Ζ	a (Å)	b (Å)	β (°)	c (Å)	$V(Å^3)$	Ref.
Vauquelinite*	$Pb_2Cu(CrO_4)(PO_4)(OH)$	$P2_1/n$	4	16.110	5.806	110.47	17.366	1521.7	[1]
Fornacite	$Pb_2Cu(CrO_4)(AsO_4)(OH)$	$P2_{1}/c$	4	8.090	5.909	110.0	17.484	785.5	[2]
Tsumebite	$Pb_2Cu(PO_4)(SO_4)(OH)$	$P2_1/m$	2	7.850	5.800	111.0	8.700	369.8	[3]
Embreyite	$(Pb.Cu, \square)_2 \{Pb[(Cr,P)O_4]_2\}(H_2O)_n$	$C2^{j}m$	1	9.802	5.603	114.9	7.649	381.2	[4]
References: [1] - *Unit cell was tr	Fanfani and Zanazzi (1968); [2] – Ksenoft ansformed with a and c axes chosen as the [ntov <i>et al.</i> (20 101] and [10]	14); [3] –] direction	Nichols (1966) is, respectively,); [4] – this w , in the origina	ork. 1 unit-cell repo	rted by Fanfan	i and Zanazzi (1968)

TABLE 6. Crystallographic parameters of vauquelinite, fornacite, tsumebite and embrevite.

and fornacite also belong to the brackebuschite structural family and exhibit doubling of both a and c unit-cell parameters in the former, and the c parameter in the latter. Structural relationships are confirmed by the above-discussed genetic linkage between vauquelinite and embreyite.

Final remarks

The determination of the embrevite structure may help to suggest the crystal chemical nature of cassedanneite. This mineral has also been discovered in an old museum specimen from Berezovskoe. However, to date, no crystals suitable for a structural study are known. Cassedanneite is similar in stoichiometry and powder XRD pattern to embreyite and was considered previously to be an analogue with the VO_4^{3-} anion instead of PO_4^{3-} . The idealized formula of cassedanneite was suggested as $Pb_5(CrO_4)_2(VO_4)_2 \cdot H_2O$; the holotype of this mineral contains 0.61 wt.% CuO and 0.65 wt.% ZnO (Cesbron et al., 1988) not included in the simplified formula. Our recent data for another specimen of cassedanneite from the same locality show up to 3.8 wt.% CuO, concentrations that are similar to those in embrevite. We propose that the structural similarity of cassedanneite and embrevite is related to the disordered distribution of Cr and V. This character of disorder seems to be very likely taking into account the following observations: (1) the existence of the continuous solid-solution series of the brackebuschite-type minerals between vauquelinite $Pb_2Cu(CrO_4)(PO_4)$ (OH), bushmakinite Pb₂Al(VO₄)(PO₄)(OH), ferribushmakinite Pb₂Fe³⁺(VO₄)(PO₄)(OH) and a phase with the hypothetical end-member composition of Pb₂Cu(VO₄)(PO₄)(H₂O). The major substitution scheme is $Cr^{6+} + Cu^{2+} \leftrightarrow V^{5+} + (Al,Fe)^{3+}$ (Khanin and Pekov, 2016a,b). (2) The crystal chemical similarity between Cr6+ and V5+ metals is closer than between Cr^{6+} and P^{5+} .

Our recently obtained data demonstrate that the powder XRD pattern of cassedanneite contains a distinct reflection with d = 13.9 Å, forbidden for the embreyite unit cell (see Table 2). This feature may indicate the doubling of the *c* unit-cell parameter of cassedanneite in comparison with embreyite, which may explain the structural differences between these two minerals.

Hexagonal relationships observed for embreyite and rhombohedral $Pb_4(PO_4)_2(CrO_4)$ indicates the possibility of phase transitions. The behaviour of embreyite at higher temperatures may be somewhat similar to the reported leadhillite \rightarrow susannite

[a = 13.754(5), b = 5.806(6), c = 9.563(3) Å, $\beta = 94.57(17)^{\circ}$ and V = 761.2 Å³]. Sp. gr. - space group.

 $(P2_1/a \rightarrow P3)$ transition at 85°C (Bindi and Menchetti, 2005; Steele *et al.*, 1998) explained by the changes in orientation of sulfate tetrahedra and PbO_n coordination polyhedra. However, the absence to date of pure embreyite material in sufficient quantities prevents additional studies.

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References

- Adib, D. and Ottemann, J. (1970) Some new lead oxide minerals and murdochite from T. Khuni Mine, Anarak, Iran. *Mineralium Deposita*, 5, 86–93.
- Aktas, O., Salje, E.K.H. and Carpenter, M.A. (2013) Resonant ultrasonic spectroscopy and resonant piezoelectric spectroscopy in ferroelastic lead phosphate, Pb₃(PO₄)₂. Journal of Physics: Condensed Matter, 25, 465401.
- Alkemper, J. and Fuess, H. (1998) The crystal structures of NaMgPO₄, Na₂CaMg(PO₄)₂ and Na₁₈Ca₁₃Mg₅(PO₄)₁₈: new examples for glaserite related structures. Zeitschrift für Kristallographie – Crystalline Materials, 213, 282–287.
- Barbier, J. and Maxin, D. (1995) Phase transformation in Pb₄(PO₄)₂CrO₄. Journal of Solid State Chemistry, **116**, 179–184.
- Bariand, P. and Herpin, P. (1963) Une nouvelle espèce minérale: l'iranite, chromate hydraté de plomb. Bulletin de la Societe Francaise Mineralogie et de Cristallographie, 86, 113–135.
- Bindi, L. and Menchetti, S. (2005) Structural changes accompanying the phase transformation between leadhillite and susannite: A structural study by means of *in situ* high-temperature single-crystal X-ray diffraction. *American Mineralogist*, **90**, 1641–1647.
- Bismayer, U. and Salje, E. (1981) Ferroelastic phases in Pb₃(PO₄)₂–Pb₃(AsO₄)₂; X-ray and optical experiments. *Acta Crystallographica*, A37, 145–153.
- Cesbron, F. and Williams, S.A. (1980) Iranitehémihédrite, bellite, phoenicochroite, vauquelinite et fornacite: synthèse et nouvelles données. *Bulletin de Minéralogie*, **103**, 469–477.
- Cesbron, F., Giraud, R., Pillard, F. and Poullen, J.-F. (1988) La cassedannéite, nouveau chromo-vanadate

de plomb de Beresovsk (Oural). *Comptes Rendus de l'Académie des Sciences – Series II*, **306**, 125–127.

- Christy, A. (2015) Causes of anomalous mineralogical diversity in the Periodic Table. *Mineralogical Magazine*, 79, 33–49.
- Chukanov, N.V. and Chervonnyi, A.D. (2016) Infrared Spectroscopy of Minerals and Related Compounds. Springer Verlag, Cham, Switzerland.
- Cocco, G., Fanfani, L. and Zanazzi, P.F. (1967) The crystal structure of fornacite. *Zeitschrift f
 ür Kristallographie*, 124, 385–397.
- Cooper, M.A. and Hawthorne, F.C. (1994) The crystal structure of wherryite, $Pb_7Cu_2(SO_4)_4(SiO_4)_2(OH)_2$, a mixed sulfate-silicate with [⁶¹ $M(TO_4)_2\phi$] chains. *The Canadian Mineralogist*, **32**, 373–380.
- Cooper, M.A., Ball, N.A., Hawthorne, F.C., Paar, W.H., Roberts, A.C. and Moffatt, E. (2011) Georgerobinsonite, Pb₄(CrO₄)₂(OH)₂FCl, a new chromate mineral from the Mammoth – St. Anthony mine, Tiger, Pinal County, Arizona: description and crystal structure. *The Canadian Mineralogist*, **49**, 865–876.
- Cooper, M.A., Abdu, Y.A., Hawthorne, F.C. and Kampf, A.R. (2016) The crystal structure of gianellaite, [(NHg₂)₂](SO₄)(H₂O)_x, a framework of (NHg₄) tetrahedra with ordered (SO₄) groups in the interstices. *Mineralogical Magazine*, **80**, 869–875.
- Effenberger, H. and Pertlik, F. (1986) Four monazite type structures: comparison of SrCrO₄, SrSeO₄, PbCrO₄ (crocoite), and PbSeO₄. Zeitschrift für Kristallographie, **176**, 75–83.
- Fanfani, L. and Zanazzi, P.F. (1968) The crystal structure of vauquelinite and the relationships to fornacite. *Zeitschrift für Kristallographie – Crystalline Materials*, **126**, 433–443.
- Fettes, D. and Desmons, J. (2007) Metamorphic rocks A Classification and Glossary of Terms. Cambridge University Press, UK.
- Gao, J., Song, L., Hu, X. and Zhang, D. (2011) A buetschliite-type rare-earth borate, KBaY(BO₃)₂. *Solid State Sciences*, **13**, 115–119.
- Hartl, K. and Braungart, R. (1978*a*) Strontiumchromat(V, VI), Sr_{2.67} _{0.33}(CrO₄)_{1.33}(CrO₄)_{0.67}, eine Hochtemperaturphase mit Defekt-Bariumphosphat-Struktur. *Zeitschrift für Naturforschung*, B33, 952–953 [in German].
- Hartl, K. and Braungart, R. (1978b) Strontiumphosphatchromat(VI), Sr₃(PO₄)₂·SrCrO₄, eine dimorphe Hochtemperaturverbindung. Zeitschrift für Naturforschung, B33, 954–955 [in German].
- Hesse, K.-F. and Simons, B. (1982) Crystal structure of synthetic K₂Mg(CO₃)₂. Zeitschrift für Kristallographie, 161, 289–292.
- Kampf, A.R., Mills, S.J., Housley, R.M., Rumsey, M.S. and Spratt, J. (2012*a*) Lead-tellurium oxysalts from Otto Mountain near Baker, California: VII. Chromschieffelinite, Pb₁₀Te₆O₂₀(OH)₁₄(CrO₄)

 $(H_2O)_5$, the chromate analog of schieffelinite. *American Mineralogist*, **97**, 212–219.

- Kampf, A.R., Mills, S.J., Housley, R.M., Bottrill, R.S. and Kolitsch, U. (2012*b*) Reynoldsite, $Pb_2Mn_2^{4+}O_5(CrO_4)$, a new phyllomanganate-chromate from the Blue Bell claims, California and the Red Lead mine, Tasmania. *American Mineralogist*, **97**, 1187–1192.
- Khanin, D.A. and Pekov, I.V. (2016a) Minerals with brackebuschite-like structures: a novel solid-solution system involving Cr⁶⁺ and V⁵⁺. *Zapiski Rossiiskogo Mineralogicheskogo Obshchestva*, **145**, 96–112 [in Russian].
- Khanin, D.A. and Pekov, I.V. (2016b) New data on cassedanneite. *International Conference Dedicated to* 300-year Anniversary of the Fersman Mineralogical Museum. Moscow, pp. 188–189 [in Russian].
- Khanin, D.A., Pekov, I.V., Pakunova, A.V., Ekimenkova, I.A. and Yapaskurt, V.O. (2015) Natural system of fornacite–vauquelinite–embreyite solid solutions and variations in the chemical composition of these minerals from occurrences of the Urals. *Zapiski Rossiiskogo Mineralogicheskogo Obshchestva*, **144**, 36–60 [in Russian].
- Kleymenov, D.A., Pekov, I.V., Erokhin, Y.V. and Chukanov, N.V. (2003) New data on embreyite from the oxidation zone of the Berezovskoye gold deposit. Pp. 171–177 in: *Mineralogiya Urala-2003*. Miass, Russia [in Russian].
- Kovrugin, V.M., Colmont, M., Terryn, C., Colis, S., Siidra, O.I., Krivovichev, S.V. and Mentré, O. (2015) pH-Controlled pathway and systematic hydrothermal phase diagram for elaboration of synthetic lead nickel selenites. *Inorganic Chemistry*, 54, 2425–2434.
- Ksenofontov, D.A., Kabalov, Y.K., Pekov, I.V., Zubkova, N.V., Ekimenkova, I.A. and Pushcharovskii, D.Y. (2014) Refinement of the crystal structure of fornacite using the Rietveld method. *Doklady Earth Sciences*, **456**, 520–523.
- Lazoryak, B.I. (1996) Design of inorganic compounds with tetrahedral anions. *Russian Chemical Reviews*, 65, 287–305.
- Leclaire, A., Monier, J.C. and Raveau, B. (1984) A molybdosilicophosphate with an intersecting-tunnel structure which exhibits ion-exchange properties, $AMo_3P_{5.8}Si_2O_{25}$ (A = Rb, Tl). Acta Crystallographica, **B40**, 180–185.
- Martinetto, P., Anne, M., Dooryhée, E., Walter, P. and Tsoucaris, G. (2002) Synthetic hydrocerussite, 2PbCO₃·Pb(OH)₂, by X-ray powder diffraction, *Acta Crystallographica*, C58, i82–i84.
- McLean, W.J. and Anthony, J.W. (1970) The crystal structure of hemihedrite. *American Mineralogist*, 55, 1103–1114.
- Moore, P.B. (1973) Bracelets and pinwheels: A topological-geometrical approach to the calcium

orthosilicate and alkali sulfate structures. *American Mineralogist*, **58**, 32–42.

- Mücke, A. (1972) Santanait, ein neues Bleichromat-Mineral. Neues Jahrbuch für Mineralogie, Monatshefte, 10, 455–458.
- Murashko, M.N., Pekov, I.V., Krivovichev, S.V., Chernyatyeva, A.P., Yapaskurt, V.O., Zadov, A.E. and Zelensky, M.E. (2013) Steklite, KAl(SO₄)₂: A finding at the Tolbachik Volcano, Kamchatka, Russia, validating its status as a mineral species and crystal structure. *Geology of Ore Deposits*, **55** [Special Issue: *Rossiiskogo Mineralogicheskogo Obshchestva*], 594–600.
- Nakamoto, K. (2009) Infrared and Raman Spectra of Inorganic and Coordination Compounds, Part B, Applications in Coordination, Organometallic, and Bioinorganic Chemistry. John Wiley & Sons, Hoboken, USA.
- Nichols, M.C. (1966) The structure of tsumebite. *American Mineralogist*, **51**, 267–267.
- Pekov, I.V. (1998) Minerals First Discovered on the Territory of the Former Soviet Union. OP, Moscow.
- Quareni, S. and de Pieri, R. (1965) A three-dimensional refinement of the structure of crocoite, PbCrO₄. Acta Crystallographica, 19, 287–289.
- Rumsey, M.S., Krivovichev, S.V., Siidra, O.I., Kirk, C.A., Stanley, C.J. and Spratt, J. (2012) Rickturnerite, Pb₇O₄[Mg(OH)₄](OH)Cl₃, a complex new lead oxychloride mineral. *Mineralogical Magazine*, **76**, 59–73.
- Salje, E.K.H. (2015) Modulated minerals as potential ferroic materials. *Journal of Physics: Condensed Matter*, 27, 305901.
- Sarukhanyan, N.L., Iskhakova, L.D. and Trunov, V.K. (1983) Crystal structure of RbEu(SO₄)₂. *Kristallografiya*, **28**, 452–456 [in Russian].
- Sheldrick, G.M. (2015) Crystal structure refinement with SHELXL. Acta Crystallographica, C71, 3–8.
- Siidra, O.I., Vergasova, L.P., Krivovichev, S.V., Kretser, Y. L., Zaitsev, A.N. and Filatov, S.K. (2014a) Unique thallium mineralization in the fumaroles of Tolbachik volcano, Kamchatka Peninsula, Russia. I. Markhininite, TIBi(SO₄)₂. *Mineralogical Magazine*, **78**, 1687–1698.
- Siidra, O.I., Zenko, D.S., Krivovichev, S.V. (2014b) Structural complexity of lead silicates: Crystal structure of Pb₂₁[Si₇O₂₂]₂[Si₄O₁₃] and its comparison to hyttsjoite. *American Mineralogist*, **99**, 817–823.
- Sokolova, E., Hawthorne, F.C., Pautov, L.A. and Agakhanov, A.A. (2010) Byzantievite, Ba₅(Ca,*REE*, Y)₂₂(Ti,Nb)₁₈(SiO₄)₄[(PO₄),(SiO₄)]₄(BO₃)₉O₂₁[(OH), F]₄₃(H₂O)_{1.5}: the crystal chemistry of the only known mineral with the oxyanions (BO₃), (SiO₄) and (PO₄). *Mineralogical Magazine*, **74**, 285–308.
- Song, S.Y. and Ok, K.M. (2015) Modulation of framework and centricity: cation size effect in new quaternary selenites, ASc(SeO₃)₂ (A = Na, K, Rb, and Cs). *Inorganic Chemistry*, **54**, 5032–5038.

- Steele, A.M., Pluth, J.J. and Livingstone, A. (1998) Crystal structure of macphersonite Pb₄SO₄(CO₃)₂(OH)₂: comparison with leadhillite. *Mineralogical Magazine*, 62, 451–459.
- Tissot, R.G., Rodriguez, M.A., Sipola, D.L. and Voigt, J. A. (2001) X-ray powder diffraction study of synthetic palmierite, K₂Pb(SO₄)₂. *Powder Diffraction*, 16, 92–97.
- Wildner, M. (1992a) Isotypism of a selenite with a carbonate: structure of the buetschliite-type compound K₂Co(SeO₃)₂. Acta Crystallographica, C48, 410–412.
- Wildner, M. (1992b) Structure of K₂Mn(SeO₃)₂, a further buetschliite-type selenite. Acta Crystallographica, C48, 595.
- Williams, S.A. (1972) Embreyite, a New Mineral from Berezov, Siberia. *Mineralogical Magazine*, 38, 790–793.

- Williams, S.A. and Anthony, J.W. (1970) Hemihedrite, a new mineral from Arizona. *American Mineralogist*, 55, 1088–1102.
- Williams, S.A. and Duggan, M. (1980) La macquartite: un nouveau silico-chromate de Tiger, Arizona. *Bulletin de Minéralogie*, **103**, 530–532.
- Williams, S.A., McLean, W.J. and Anthony, J.W. (1970) A study of phoenicochroite – its structure and properties. *American Mineralogist*, 55, 784–792.
- Yang, H., Sano, J.L., Eichler, C., Downs, R.T. and Costin, G. (2007) Iranite, CuPb₁₀(CrO₄)₆(SiO₄)₂(OH)₂, isomorphous with hemihedrite. *Acta Crystallographica*, C63, i122–i124.
- Zubkova, N.V., Pushcharovsky, D.Y., Giester, G., Tillmanns, E., Pekov, I.V. and Kleimenov, D.A. (2002) The crystal structure of arsentsumebite, Pb₂Cu[(As,S) O₄]₂(OH). *Mineralogy and Petrology*, **75**, 79–88.