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Lobanovite, K2Na(Fe2+4Mg2Na)Ti2(Si4O12)2O2(OH)4, a new mineral of the astrophyllite supergroup and its relation to magnesioastrophyllite

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5	Lobanovite, K ₂ Na(Fe ²⁺ ₄ Mg ₂ Na)Ti ₂ (Si ₄ O ₁₂) ₂ O ₂ (OH) ₄ , a new mineral
6	of the astrophyllite supergroup and its relation to magnesioastrophyllite
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24 Abstract

Lobanovite, $K_2Na(Fe^{2+}_4Mg_2Na)Ti_2(Si_4O_{12})_2O_2(OH)_4$, is a new mineral of the astrophyllite 25 26 supergroup from Mt. Yukspor, the Khibiny alkaline massif, Kola Peninsula Russia. It has been 27 previously known under the following names: monoclinic astrophyllite, magnesium astrophyllite, 28 magnesiumastrophyllite and magnesioastrophyllite but has never been formally proposed and 29 approved as a valid mineral species by the CNMNC-IMA. It has now been revalidated and 30 named lobanovite after Dr. Konstantin V. Lobanov, a prominent Russian ore geologist who 31 worked in Kola Peninsula for more than forty years (Nomenclature voting proposal 15-B). 32 Lobanovite has been described from pegmatitic cavities on Mt. Yukspor where it occurs as 33 elongated bladed crystals, up to 0.04 mm wide and 0.2 mm long, with straw yellow to orange 34 colour. Associated minerals are shcherbakovite, lamprophyllite, delindeite, wadeite, umbite and 35 kostylevite. Lobanovite is biaxial (–) with refractive indices (λ = 589 nm) α = 1.658, $\beta_{calc.}$ = 1.687, y = 1.710; $2V_{meas}$ = 81.5–83°. Lobanovite is monoclinic, space group C2/m, a 5.3327(2), b 36 23.1535(9), c 10.3775(4) Å, β 99.615(1)°, V 1263.3(1) Å³, Z = 2. The six strongest reflections in 37 38 the X-ray powder diffraction data [d(Å), I, (hkl)] are: 3.38, 100, (003); 2.548, 90, (063); 10.1, 80, 39 (001); 3.80, 60, (042,131); 3.079, 50, (132,062); 2.763, 90, (171). The chemical composition of 40 lobanovite was determined by electron-microprobe analysis and the empirical formula 41 (K_{1.97}Ba_{0.01})_{Σ1.98} (Na_{0.65}Ca_{0.14})_{Σ0.79} (Fe²⁺_{3.18}Mg_{2.02}Na_{1.00}Mn_{0.72})_{Σ6.92}(Ti_{1.99} Nb_{0.06})_{Σ2.05}[(Si_{8.01} 42 $AI_{0.06})_{\Sigma 8.07}O_{24}O_{2}(OH)_{4.03}F_{0.19}$ was calculated on the basis of 30.2 (O + OH + F) anions, with H₂O calculated from structure refinement, $D_{calc} = 3.161 \text{ g cm}^{-3}$. In the structure of lobanovite, the 43 44 main structural unit is the HOH block, which consists of one close-packed O (Octahedral) and 45 two H (Heteropolyhedral) sheets. The M(1–4) octahedra form the O sheet and the T_4O_{12} 46 astrophyllite ribbons and [5]-coordinated Ti-dominant D polyhedra link through common vertices 47 to form the H sheet. The HOH blocks repeat along [001], and K and Na atoms occur at the 48 interstitial A and B sites. The simplified and endmember formulae of lobanovite are

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- 49 $K_2Na[(Fe^{2+},Mn)_4Mg_2Na]Ti_2(Si_4O_{12})_2O_2(OH)_4$ and $K_2Na(Fe^{2+}_4Mg_2Na)Ti_2(Si_4O_{12})_2O_2(OH)_4$,
- 50 respectively.
- 51
- 52 *Keywords*: lobanovite, magnesiumastrophyllite, astrophyllite supergroup, chemical composition,
- 53 crystal-structure refinement

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54 Introduction

55 Lobanovite, $K_2Na(Fe^{2+}_4Mg_2Na)Ti_2(Si_4O_{12})_2O_2(OH)_4$, is a new mineral of the astrophyllite 56 supergroup from Mt. Yukspor, the Khibiny alkaline massif, Kola Peninsula Russia. It has been 57 previously known under the names monoclinic astrophyllite, magnesium astrophyllite, 58 magnesiumastrophyllite and magnesioastrophyllite (Peng and Ma, 1963; X-ray Laboratory, 59 Hubei Geologic College, 1974; Shi et al., 1998; Piilonen et al., 2003a; Sokolova and Cámara, 60 2008; Cámara et al., 2010) but has never been formally proposed and approved as a valid 61 mineral species by the CNMNC-IMA, although it was discovered in the CNMNC era. It has now 62 been revalidated and named lobanovite after Dr. Konstantin Valentinovich Lobanov (Cyrillic: 63 Константин Валентинович Лобанов; b. 1952, Roslavl', Russia), a prominent Russian ore 64 geologist, DSc. (Doctor of Science, Geology and Mineralogy) and an expert in Precambrian 65 metallogeny. It is particularly appropriate to name this mineral from the Kola Peninsula after Dr. 66 Lobanov, as he worked in the Kola Peninsula for the last 40 years, focusing on the rocks of the 67 Kola superdeep borehole and their relation to the deep structure of ore districts (CNMNC-IMA: 68 Nomenclature voting proposal 15-B). The holotype specimen of lobanovite is the microprobe 69 mount of the single crystal that was used for structure refinement by Sokolova and Cámara 70 (2008) and it was deposited in the collections of the Fersman Mineralogical Museum, Russian 71 Academy of Sciences, Leninskii Prospekt, 18/2, 119071 Moscow, Russia, catalogue number 72 4708/1.

In this paper, we give an overview of previous studies of lobanovite and report a
compilation of data from the literature and necessary for a description of lobanovite as a valid
mineral species.

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77 **Previous work**

Semenov (1959) reported on the *astrophyllite with unusually high content of MgO (6.39 wt.%)*from the Khibiny alkaline massif, Kola Peninsula, Russia; the complete chemical analysis was

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80 not given. Peng and Ma (1963) reported a chemical analysis, with the following simplified formula $K_2Na_2(Fe^{2+},Mn)_2(Fe^{2+},Fe^{3+})_3Mg_2Ti_2(Si_4O_{12})_2(O,OH)_3(OH,F)_4$, Z = 2, unit-cell parameters 81 82 a 10.43(2), b 23.00(5), c 5.35(1) Å, β 102°, and a preliminary crystal structure in space group 83 A2/m (atom coordinates were not reported) for the astrophyllite from Khibiny. X-ray Laboratory, 84 Hubei Geologic College (1974) reported the X-ray powder-diffraction data, DTA data and optical 85 properties for the material mentioned above. The crystal structure of monoclinic astrophyllite 86 was refined by Shi et al. (1998) on sample no. 3086 provided by A. Khomyakov from Mt. 87 Yukspor, the Khibiny alkaline massif, Kola Peninsula, Russia. They gave a full description of the crystal structure, an ideal formula K₂NaNa(Fe,Mn)₄Mg₂Ti₂(Si₄O₁₂)₂(OH)₄(OH,F)₂, and unit-cell 88 parameters, a 10.370(3), b 23.129(5), c 5.322(1) Å, β 99.55(2)°, V 1258.8(5) Å³, space group 89 A2, Z = 2, D_{calc} = 3.173 g/cm⁻³ (this formula has an excess charge of 2⁺). The structure data of 90 91 Shi et al. (1998) is in the International Crystal Structure Database (ICSD) under #52032. The 92 ICSD revised #52032 as #56848, with higher symmetry, space group A2/m, and a guite different 93 formula: $K_{1.96}Na_{2.06}Fe_{3.72}Mg_{2.28}Ti_2(Si_8O_{28})(H_2O)_2$, based on the same work of Shi *et al.* (1998). 94 The latter formula is neutral but it is chemically and structurally incorrect as H₂O groups do not 95 occur in magnesium astrophyllite. Piilonen et al. (2003a) considered the structure of magnesium astrophyllite (space group A2) and re-wrote its formula as K₂Na[Na(Fe,Mn)₄Mg₂]Ti₂Si₈O₂₆ 96 97 $(OH)_4 \square$. Although the ICSD #56848 points out that the possible space group might be A2/m, 98 the crystal structure of magnesium astrophyllite has not been re-investigated since the work of 99 Shi et al. (1998). Following the inconsistency between the chemical composition and structure 100 for magnesium astrophyllite in the ICSD (see above), Sokolova and Cámara (2008) refined the 101 crystal structure of magnesium astrophyllite (see below) and reported the chemical composition 102 of the crystal of magnesium astrophyllite used for structure refinement. They confirmed the 103 general topology of the crystal structure described by Shi et al. (1998) and gave the ideal 104 formula of magnesium astrophyllite as follows: K₂Na(Fe₄Mg₂Na)^[5]Ti₂Si₈O₂₆(OH)₄. Cámara et al. 105 (2010) suggested writing the astrophyllite-ribbon component of the formula as $(Si_4O_{12})_2O_2$

106 versus Si_8O_{26} [Piilonen *et al.* (2003a)]. They emphasized that (1) the translational repeat 107 fragment of the ribbon has the formula Si₄O₁₂; (2) there are two ribbons (corresponding to two H 108 sheets) attached to the central O sheet; (3) the O_2 anions do not link to Si, and this information 109 should be preserved in the chemical formula by writing them separately from the silicate radical. 110 Cámara et al. (2010) wrote the formula of magnesiumastrophyllite as K₂Na(Fe₄ 111 $Mq_2Na)Ti_2(Si_4O_{12})_2O_2(OH)_4$. Burke (2008) renamed magnesium astrophyllite and/or 112 magnesiumastrophyllite as magnesioastrophyllite. Sokolova (2012) showed that the topology of 113 the HOH block in *magnesioastrophyllite* is different from all other structures of the astrophyllitesupergroup minerals due to the different chemical composition of the O sheet (Fe²⁺₄Mg₂Na), 114 115 particularly the dominance of Na⁺ at the M(1) site which makes the M(1) polyhedron too large to 116 share edges with Si₂O₇ groups of astrophyllite ribbons and results in a different linkage of H and 117 O sheets, i.e. a different topology of the HOH block. She wrote the ideal formula of magnesioastrophyllite as follows: $K_2Na(Fe^{2+}_4Mg_2Na)Ti_2(Si_4O_{12})_2O_2(OH)_4$ and emphasized that 118 119 the name magnesioastrophyllite is misleading as it is the dominance of Na⁺ at the M(1) site that results in the different topology of the HOH block [not the dominance of Mg^{2+} at the M(4) site]. 120 121 Further on in the paper, we will use only the name lobanovite and compare its properties 122 and structure to those of astrophyllite (Table 1). 123

124 Occurrence and mineral association

125 Sokolova and Cámara (2008) refined the crystal structure and determined the chemical

126 composition of lobanovite on the same single crystal taken from the sample of delindeite in the

- 127 collection of minerals of Adriana and Renato Pagano, Milan, Italy (Collezione Mineralogica,
- 128 sample #6270). This sample came from Alexander Khomyakov and was described in
- 129 Khomyakov (1995). Hence we use his description for the occurrence and associated minerals.

130

Lobanovite was found in pegmatite cavities at Mt. Yukspor, the Khibiny alkaline massif,

131 Kola Peninsula, Russia. Associated minerals are shcherbakovite, lamprophyllite, delindeite,

132 wadeite, umbite, kostylevite and other species [taken from Khomyakov (1995), p. 151].

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136 Physical properties

137 Lobanovite occurs as elongated bladed crystals, with straw yellow (Anthony et al., 1995) to 138 orange colour (Sokolova and Cámara, 2008) up to 0.04 mm wide and 0.2 mm long, with a 139 vitreous luster and a white to vellowish-white streak. The mineral has Mohs hardness of 3, 140 perfect cleavage parallel to (001) and moderate cleavage parallel to (010). Twinning, parting 141 and fracture have not been observed (Anthony et al., 1995). Lobanovite does not fluoresce 142 under SW or LW ultraviolet light, $D_{calc.} = 3.161 \text{ g/cm}^3$ (from the empirical formula of Sokolova 143 and Cámara, 2008). Lobanovite is optically biaxial (–), $\alpha = 1.658$, $\beta = 1.687_{calc.}$, $\gamma = 1.710$ (589 144 nm), with $2V_{\text{meas.}} = 81.5 - 83^\circ$. The pleochroism: X = bright yellow, Y = pale yellowish grey, Z = 145 grey, with the absorption scheme: Z < Y < X and orientation Y = b, $Z \land a = -5$ to -6° (X-ray 146 Laboratory, Hubei Geological College, 1974). DTA curve shows endothermic peaks at 835°C 147 (strong, escape of OH) and 900°C (weak) (X-ray Laboratory, Hubei Geologic College, 1974). 148 The compatibility index was calculated using the chemical analysis and unit-cell 149 parameters of Sokolova and Cámara (2008) and optical properties from X-ray Laboratory, Hubei 150 Geological College (1974): $1 - (K_{c}/K_{c})$ is 0.057 and is rated as good (Mandarino, 1981). 151

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152 Chemical composition

- 153 Sokolova and Cámara (2008) determined the chemical composition of lobanovite using a
- 154 Cameca SX100 electron microprobe (WDS mode, 15 kV, 20 nA, 10 μm beam diameter). Ta, Zr,
- 155 Sn, Zn, Cs, Rb, Sr and Pb were sought but not detected [for details, see Sokolova and Cámara

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157 crystal-structure refinement. The chemical composition of lobanovite is given in Table 2 (mean 158 of 10 analyses) in comparison with that of the X-ray Laboratory, Hubei Geologic College (1974). 159 Sokolova and Cámara (2008) calculated the empirical formula on the basis of 30.2 (O + OH + F) 160 anions which include 30 (O + OH) p.f.u. and 0.2 F a.p.f.u. (atoms per formula unit). The 161 empirical, simplified and endmember formulae are $(K_{1.97}Ba_{0.01})_{\Sigma_{1.98}}(Na_{0.65}Ca_{0.14})_{\Sigma_{0.79}}$ $(Fe^{2+}{}_{3.18}Mg_{2.02}Na_{1.00}Mn_{0.72})_{\Sigma 6.92}(Ti_{1.99}Nb_{0.06})_{\Sigma 2.05}[(Si_{8.01}Al_{0.06})_{\Sigma 8.07}O_{24}]O_2(OH)_{4.03}F_{0.19}, Z = 2,$ 162 K₂Na[(Fe²⁺,Mn)₄Mg₂Na]Ti₂(Si₄O₁₂)₂O₂(OH)₄ and K₂Na(Fe²⁺₄Mg₂Na)Ti₂(Si₄O₁₂)₂O₂(OH)₄, 163 164 respectively. 165 166 X-ray powder-diffraction data 167 X-ray powder-diffraction data are presented in Table 3.

(2008)]. The content of H₂O (wt. %) was calculated by stoichiometry from the results of the

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169 Crystal structure

170 Sokolova and Cámara (2008) refined the crystal structure of lobanovite to $R_1 = 3.2\%$ on the

basis of 1861 unique reflections ($F_{o} > 4\sigma F$) collected on a Bruker AXS SMART APEX

172 diffractometer with a CCD detector and MoKα-radiation: space group C2/m, a 5.3327(2), b

173 23.1535(9), *c* 10.3775(4)Å, β 99.615(1)°, *V* 1263.3(1) Å³, *Z* = 2, $D_{calc.}$ 3.161 g.cm⁻³ (Table 4)

174 from Mt. Yukspor, Khibina alkaline massif, Kola Peninsula, Russia. The main structural unit in

175 the structure of lobanovite is the HOH block which consists of a central close-packed O

176 (Octahedral) and two adjacent H (Heteropolyhedral) sheets. In the O sheet, there are four [6]-

177 coordinated M(1-4) sites occupied by Na, $Fe^{2+}_{1.25}Mn_{0.75}$, $Fe^{2+}_{1.20}Mg_{0.80}$ and $Mg_{1.25}Fe^{2+}_{0.75}$, with

178 <M(1,2,3,4)–O> = 2.372, 2.166, 2.113 and 2.094 Å, respectively (Fig. 1*a*, Table 5). The ideal

179 composition of the O sheet is (Fe²⁺₄Mg₂Na)O₂(OH)₄ p.f.u. In the H sheet, there is one [5]-

180 coordinated *D* site occupied by $Ti_{1.97}Nb_{0.03}$, with $\langle D-O \rangle = 1.907$ Å, and there are two

181 tetrahedrally coordinated T sites occupied primarily by Si with minor AI, with $\langle T-O \rangle = 1.625$ Å. TO_4 tetrahedra form T_4O_{12} astrophyllite ribbons extending along **a**. Astrophyllite ribbons and [5]-182 183 coordinated D polyhedra connect through common vertices to form the H sheet. The ideal 184 composition of the two H sheets is $Ti_2(Si_4O_{12})_2$ p.f.u. An O sheet and two adjacent H sheets link 185 through common vertices of TO₄ tetrahedra, D octahedra and M octahedra to form an HOH 186 block parallel to (001) (Fig. 1a). The HOH blocks alternate with I (Intermediate) blocks along c 187 (Fig. 1b). In the I block, there are two interstitial sites, a [10]-coordinated A site and an [8]-188 coordinated B site occupied by $K_{1.96}Ba_{0.01}\Box_{0.03}$ and $Na_{0.65}Ca_{0.14}\Box_{0.21}$, respectively, with $\langle A-O \rangle =$ 189 3.120 and $\langle B-O \rangle = 2.536$ Å (Table 5). The ideal composition of the I block is K₂Na a.p.f.u. 190 Cations at the A and B sites form a layer parallel to (001) (Fig. 1b). The presence of the partly 191 occupied F site makes 17% of the A and B sites [11]- and [10]-coordinated, respectively (Table 192 5).

193 The general topology of the lobanovite structure is in accord with Shi et al. (1998). The 194 topology of the HOH block in lobanovite differs from all other structures of the astrophyllitesupergroup minerals (for the astrophyllite structure, see Fig. 2a in Sokolova et al., 2015) due to 195 the chemical composition of the O sheet, $Fe_{4}^{2+}Mq_{2}Na$, where Na⁺ is dominant at the M(1) site 196 197 (Sokolova, 2012). Hence the Na octahedron is too large to share edges with Si_2O_7 groups of 198 astrophyllite ribbons resulting in a different linkage of H and O sheets, i.e. a different topology of the HOH block [cf. Si-Si distances associated with the Na octahedron (3.96 Å) and the Fe²⁺ 199 200 octahedron (3.06 Å), Fig. 1a].

201

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245 Figure captions

247	Fig. 1. The crystal structure of lobanovite: (<i>a</i>) the HOH block viewed perpendicular to the plane
248	of the block [after Fig. 6a of Sokolova (2012)]; (b) general view of the crystal structure of
249	lobanovite [after Fig. 2 <i>b</i> of Sokolova (2012)]. The [5]-coordinated Ti polyhedra and Fe ²⁺ - and
250	Mg-dominant octahedra are pale yellow, green and pink; Na octahedra are navy blue; Si
251	tetrahedra are orange; K and Na atoms at the A and B sites are shown as green and blue
252	spheres; OH groups are shown as small red spheres. The position of the cation layer in the ${f I}$
253	(Intermediate) block (m = 1, where m is a number of cation layers) is shown by a turquoise line.
254	

	Lobanovite*	Astrophyllite**
Formula	K ₂ Na(Fe ²⁺ ₄ Mg ₂ Na)Ti ₂ (Si ₄ O ₁₂) ₂ O ₂ (OH) ₄	$K_2NaFe^{2+}_7Ti_2$ (Si ₄ O ₁₂) ₂ O ₂ (OH) ₄ F
System	monoclinic	triclinic
Space group	C2/m	PĪ
a (Å)	5.3327(2)	5.36–5.42
b	23.1535(9)	11.85–11.95
с	10.3775(4)	11.66–11.75
α (°)	90	112.03–113.13
β	99.615(1)	94.52–94.64
Y	90	103.08–103.14
V (Å ³)	1263.3(1)	652.9–668.5
Ζ	2	1
D _{meas.} /D _{calc.} (g cm ⁻³)	n.d./3.161	3.09-3.23/3.35
Strongest refl. in powder diffraction data $d_{obs.}$ (Å) (I)	3.38 (100), 2.548 (90), 10.1 (80), 3.80 (60), 3.079 (50), 2.763 (50)	10.6 (100), 3.51 (80), 2.77 (60), 2.64 (60), 2.57 (60), 1.76 (30)
Optical character	biaxial (–)	biaxial (+)
α	1.658	1.678–1.695
β	1.687	1.703–1.726
Y	1.710	1.733–1.758
2V _{meas} (°)	81.5–83	66–84
Colour	Yellow to orange	Bronze yellow to golden yellow, brown to reddish brown
Pleochroism	X = bright yellow Y = pale yellowish grey Z = grey	X = deep orange red Y = orange yellow Z = lemon yellow

Table 1. Comparison of lobanovite and astrophyllite

*All data are from Sokolova and Cámara (2008) except for powder diffraction data and optical properties (X-ray laboratory, Hubei Geologic college 1974);

**Data from Anthony et al. (1995) except for cell parameters (Piilonen et al., 2003b)

Constituent	wt.%		Range (wt.%)	SD (wt.%)	Probe Standard
	X-ray Lab., Hubei Geologic College (1974)		— Sokolova and	Cámara (20	008)
Nb_2O_5	n.a	0.64	0.29 – 1.22	0.31	$Ba_2NaNb_5O_{15}$
TiO ₂	12.18	13.11	12.47 – 13.48	0.31	titanite
SiO ₂	37.98	39.72	39.43 - 39.92	0.18	diopside
AI_2O_3	1.11	0.24	0.08 - 0.42	0.13	andalusite
Fe_2O_3	2.95	n.d.			
ВаО	n.a.	0.13	0.04 – 0.19	0.04	baryte
FeO	17.91	18.86	17.98 – 21.05	0.84	fayalite
MnO	4.00	4.21	3.10 – 4.59	0.45	spessartine
CaO	1.15	0.65	0.59 – 0.82	0.06	diopside
MgO	6.39	6.72	6.05 - 6.93	0.24	forsterite
K ₂ O	7.28	7.66	7.54 – 7.78	0.07	orthoclase
Na ₂ O	5.38	4.22	4.10 - 4.29	0.07	jadeite
H ₂ O	3.44	3.00*			
F	0.45	0.29	0.17 – 0.43	0.07	F- riebeckite
0 = F	-0.19	-0.12			
Total	100.03	99.33			

Table 2. Chemical composition of lobanovite

n.d. – not detected, n.a. – not analyzed; *calculated from structure refinement.

I _{obs.}	d _{obs.} (Å)	d _{calc.} (Å)	hkl
80	10.1	10.22	001
60	3.80	3.83 3.82	0 4 2 1 3 1
100	3.38	3.41	003
50	3.079	3.11 3.08	132 062
30	2.889	2.882	–1 3 3
50	2.763	2.764	–171
90	2.548	2.554	063
20	2.350	2.369	172
30	2.260	2.265	–173
30	2.040	2.047	005
50	1.818	1.805	1 11 1
50	1.727	1.727	–175
40	1.657	1.654	0 14 0
10	1.591	1.592 1.586	2 10 2 1 7 5
20	1.525	1.517	-176
70	1.463	1.461	007
40	1.404	1.402	176
10	1.373	1.379	-207 2141

Table 3. Powder diffraction data for lobanovite*

*I_{obs.} and $d_{obs.}$ (Å) taken from X-ray Laboratory, Hubei Geologic College (1974); $d_{calc.}$ (Å) and hkl – this work.

•		•		(/
Atom	x	У	Z	$U_{ m eq}$ (Å ²)
M(1)	0	0	0	0.0129(3)
M(2)	1/2	0.07440(2)	0	0.0093(2)
M(3)	0	0.14505(2)	0	0.0091(2)
M(4)	1/2	0.21399(3)	0	0.0098(2)
D	0.43702(8)	0	0.28713(4)	0.0064(2)
T(1)	0.07901(9)	0.21773(2)	0.73587(5)	0.0073(1)
T(2)	0.07573(9)	0.08555(2)	0.72932(5)	0.0076(1)
А	1/2	0.13847(4)	1/2	0.0267(3)
В	0	0	1/2	0.0103(3)
O(1)	0.4035(4)	0	0.1178(2)	0.0107(4)
O(2)	0.1393(3)	0.08202(7)	0.8868(1)	0.0109(3)
O(3)	0.2982(3)	0.05764(7)	0.6609(1)	0.0152(3)
O(4)	0.1966(3)	0.05755(7)	0.3287(2)	0.0170(3)
O(5)	0.0537(3)	0.15325(6)	0.6769(2)	0.0161(3)
O(6)	0.1298(3)	0.21770(6)	0.8930(1)	0.0096(3)
O(7)	0.8141(3)	0.74859(7)	0.6781(1)	0.0139(3)
O(8)	0.3656(3)	0.14762(6)	0.1030(1)	0.0115(3)
Н	0.386(6)	0.142(1)	0.191(1)	0.0138(0)*
F	1/2	0	1/2	0.009(4)*

Table 4. Final atom coordinates and equivalent displacement parameters for lobanovite [from Sokolova and Cámara (2008)]

*U_{iso.}

Site	Refined site- scattering	Assigned site- population	Calculated site- scattering	<Χ–φ> _{calc.} * (Å)	<Χ–φ> _{obs.} (Å)
	(e.p.f.u.)	(a.p.f.u.)	(e.p.f.u.)	()	()
^[6] <i>M</i> (1)	11.0(4)	1.00 Na	11.0	2.400	2.372
^[6] <i>M</i> (2)	50.0(1)	1.25 Fe ²⁺ + 0.75 Mn	51.3	2.169	2.166
^[6] <i>M</i> (3)	39.7(1)	1.20 Fe ²⁺ + 0.80 Mg	40.8	2.126	2.113
^[6] <i>M</i> (4)	34.1(1)	1.25 Mg + 0.75 Fe ²⁺	34.5	2.090	2.094
^[5] D ^[6] D**	44.4(1)	1.97 Ti + 0.03 Nb	44.6	1.891 1.975	1.907 1.952
^[10] A ^[11] A**	37.8(2)	1.96 K + 0.01 Ba + 0.03 □	37.8		3.120 3.127
^[8] B ^[10] B**	12.4(2)	0.65 Na + 0.14 Ca + 0.21 □	10.0		2.536 2.562
F	1.5(1)	0.83 □ + 0.17 F	1.5		

Table 5. Refined site-scattering and assigned site-populations for lobanovite [from Sokolova and Cámara (2008)]

*Calculated by summing constituent ionic radii; values from Shannon (1976), φ = O, OH, F; **F atom is considered a ligand.





Fig. 1

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