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# Vergasovaite Cu<sub>3</sub>O[(Mo,S)O<sub>4</sub>][SO<sub>4</sub>], a new copper-oxy-molybdate-sulfate from Kamchatka

by Elena Y. Bykova<sup>1</sup>, Peter Berlepsch<sup>2</sup>, Pavel M. Kartashov<sup>1</sup>, Joël Brugger<sup>3</sup>, Thomas Armbruster<sup>4</sup> and Alan J. Criddle<sup>5</sup>

## Abstract

Vergasovaite occurs on encrustations of sulfates in fumaroles in the Northern part of the central fumarole field of cone II of the Northern Break of the Large Tolbachik Fissure Eruption (LTFE) on the Kamchatka peninsula. It is associated with chalcocyanite, dolerophanite, euchlorine, fedotovite, tenorite, Cu-bearing anglesite and native gold. Vergasovaite has an ideal formula Cu<sub>3</sub>O[(Mo,S)O<sub>4</sub>][SO<sub>4</sub>]. Its space group is Pnma with a = 7.420(3), b = 6.741(2), c = 13.548(5) Å, V = 677.6(2) Å<sup>3</sup>, and Z = 4. The five strongest lines d<sub>calc</sub> [Å] ( $I_{calc}$ , hkl) in the X-ray powder pattern are: 3.096 (100, 104); 3.377 (69, 020); 2.998 (56, 121); 2.498 (54, 220); and 3.580 (37, 201). The olive-green mineral is transparent and distinctly pleochroic from olive-green to a yellowish to brownish-green. Microprobe analyses revealed a partial substitution of Mo by S and minor V, leading to the empirical formula (Cu<sub>2.82(7)</sub>,Zn<sub>0.10(7)</sub>,  $Pb_{0.01(1)})_{\Sigma = 2.92(9)}O[(Mo_{0.79(7)},S_{0.20(5)},V_{0.04(3)})_{\Sigma = 1.04(3)}O_4][SO_4]. IR spectroscopy showed vergasovaite to be free of H<sub>2</sub>O and OH thus the mineral is a fully oxidized mixed copper-oxy-molybdate-sulfate. Vergasovaite is isostructural with synthetic Cu<sub>3</sub>O[MoO<sub>4</sub>]<sub>2</sub>.$ 

Keywords: vergasovaite, new mineral, molybdate, sulfate, exhalate, Kamchatka.

# Introduction

The occurrence of Mo-minerals on volcanoes is very rare. The most common species are molybdenite MoS<sub>2</sub>, e.g., from the Merapi volcano on Central Java, Indonesia (KAVALIERIS, 1994), at Mount St. Helens, Washington, USA (BERNARD and LE GUERN, 1986), or from Mount Vesuvius (ZAMBONINI, 1935), and ilsemannite Mo<sub>3</sub>O<sub>8</sub>·5H<sub>2</sub>O (BERNARD and LE GUERN, 1986; BERNARD et al., 1990). A uniquely rich association of Mo-minerals was found in the high temperature exhalates of Kudrjavyi volcano (Iturup, Kuril Islands) where Re-rich and Re-free molybdenite, powellite Ca[MoO<sub>4</sub>], molybdoscheelite Ca[(W,Mo)O<sub>4</sub>], tugarinovite MoO<sub>2</sub>, molybdite MoO<sub>3</sub>, and ilsemannite were identified (BYKOVA et al., 1995). The new mixed copper-oxy-molybdate-sulfate described here occurs in fumarolic products of the Large Tolbachik Fissure Eruption (LTFE) on the Kamchatka peninsula (Fig. 1a).

The LTFE is located in the Southern part of the Kliuchevskaya group of volcanoes in central Kamchatka, where the most active volcanoes of the Kurile-Kamchatka region are located. The Kliuchevskaya group of volcanoes consists of three active (Kliuchevskoi, Bezymianny, and Plosky Tolbachik) and some inactive volcanoes (Fig. 1b). The eruptive zone extends linearly over 30 km. As a consequence of enormous lava extrusions, the top of Plosky Tolbachik collapsed, thus forming a caldera with a diameter of 1.7 km and a depth of more than 400 m.

The LTFE has been a source of exhalative

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minerals of unique diversity and abundance, the formation of which was favored by magma saturated with gases and originating at large depth and escaping at high velocity. Up to now, more than 120 minerals have been found from the LTFE (VERGASOVA and FILATOV, 1993) of which 16 are new species. The central fumarole field, located near the top of the crater of the second cone of the Northern Break (Fig. 1c), is of most interest because of its high exhalative activity. Besides



Fig. 1 Occurrence of vergasovaite: (a) Kamchatka; (b) some active volcanoes, calderas, and craters including the Kliuchevskaya group of volcanoes (modified after BRAITSEVA et al., 1995); (c) schematic map of cone II of the LTFE south of Plosky Tolbachik with the central fumarole field; (d) schematic cross section through "Treschina" fumarole.

Morphology Forms	short prismatic   pinacoid: {010}   prisms: {110} {120}   hipyramids: {111} {263} {315}
Colour (megascopic)	olive-green
Streak	light-yellow
Lustre	vitreous
Fluorescence	not observed
Hardness (VHN <sub>25 g</sub> )	mean of 6: 357 g/cm <sup>3</sup> (range 302–413 g/cm <sup>3</sup> )
Hardness (Mohs)	4–5, 5–5.5 on face subparallel (010)
Cleavage	not observed
Parting	not observed
Tenacity	brittle
Fracture	uneven
Density	4.32 g/cm <sup>3</sup> (calculated)
Ore microscopy	
Colour	grey
Internal reflections	ubiquitous, light green to colourless
Anisotropy	masked by internal reflections
Bireflectance	measurable but discernible
Pleochroism (transmitted light)	olive-green to yellowish/brownish-green
Pleochroism (reflected light)	absent
Standard used	SiC
n <sub>calc</sub> (589 nm)	1.87–1.98 (average 1.925)
n <sub>cale</sub> (Gladstone-Dale)	1.819
Mandarino Compatibility Index	poor

Tab. 1 Optical and other physical properties of vergasovaite.

high temperature Zn–Pb–Cu-compounds, the mineral forming elements are Fe, As, P, Bi, Te, Rb, Cr, Se, V, Au, and Mo.

The name for the new mineral was chosen to honour Lidia Pavlovna Vergasova, born 1941, in recognition of her contribution to the mineralogy of volcanic exhalates of Kamchatka in general, and the mineralogy of the Tolbachik region in particular. She described from there many new minerals including alarsite, alumoklyuchevskite, fedotovite, georgbokiite, ilinskite, kamchatkite, klyuchevskite, leningradite, lesukite, piypite, ponomarevite, sophiite, tolbachite, vlodavetsite.

The mineral and the name were approved (proposal 98-009) by the Commission on New Mineral and Mineral Names (CNMMN) of the International Mineralogical Association (IMA). Type Material is deposited at the Fersman Mineralogical Museum in Moscow (No. 2328/1) and at the Naturhistorisches Museum in Basel.

## Occurrence and genetic environment

Vergasovaite was found in 1993 in a fumarole called "Treschina" (Russian for fissure) in the Northern part of the central fumarole field of cone II of the Northern Break of the LTFE on



*Fig. 2* SEM-image of a small and form-rich vergasovaite crystal that grew on a larger crystal displaying simpler forms.

Face	Form	Size	$\phi_{obs}$	$\rho_{obs}$	$\phi_{calc}$	$\rho_{calc}$
(010)	Pinacoid {010}	medium	359.2	90.7	360	90
(110)	Prism {110}	small	238.0	90	241.4	90
(120)	Prism {120}	small	40.1	90	42.5	90
(130)	Prism {130}	large	31.2	90	31.4	90
$(1\overline{3}0)$	"	large	148.8	89.8	148.6	90
$(\bar{1}30)$		medium	329.2	90	328.6	90
(111)	Bipyramid {111}	large	59.7	46.4	61.4	46.0
$(1\overline{1}1)$	" "	small	301.8	49.1	298.6	46.0
$(\overline{1}11)$		large	116.0	46.6	118.6	46.0
(263)	Bipyramid {263}	medium	33.2	50.0	31.4	49.3
$(\bar{3}15)$	Bipyramid [315]	small	279.8	28.2	280.3	29.0
$(\bar{3}\bar{1}5)$	"	medium	259.0	28.0	259.7	29.0

Tab. 2 Vergasovaite measurements on optical goniometer (angles in degrees).

 $a_0: 1: c_0 = 0.546: 1: 0.497$ (X-ray)

 $a_0: 1: c_0 = 0.535: 1: 0.497$ (goniometer)

Kamchatka peninsula (FEDOTOV and MARK-HININ, 1984). Treschina is a wedge-shaped fumarole with a temperature gradient that decreases from the wide to the narrow part (Fig. 1d). This gradient is a function of the width of the fumarole which controls the speed with which the gases escape to the atmosphere and has led to a mineralogical zonation within the fumarole. From the widest and hottest part within the chloride zone, through the sulfate zone to the narrowest and coolest part within the oxide zone, the sequence of representative minerals is: tolbachite CuCl<sub>2</sub>, chalcocyanite Cu[SO<sub>4</sub>], and tenorite CuO.



Fig. 3 Idealized SHAPE-drawing of vergasovaite.

The new mineral occurred on a sulfate encrustation extracted from the sulfate zone, approximately 20 cm below surface, where it formed directly from fumarolic gases by sublimation. The temperature of the gases in the sulfate zone is 150-170 °C. The composition of the gases is as following (approximate values): 80% H<sub>2</sub>O; 12% N<sub>2</sub>; 4% CO<sub>2</sub>; 2% H<sub>2</sub>; 0.5–0.1% HCl > CH<sub>4</sub> > CO; < 0.1% H<sub>2</sub>S > NH<sub>3</sub> > O<sub>2</sub> > F<sub>2</sub> (personal comment S.F. Glavatskih).

Associated minerals are chalcocyanite  $Cu[SO_4]$ , dolerophanite  $Cu_2O[SO_4]$ , euchlorine NaKCu<sub>3</sub>O[SO<sub>4</sub>]<sub>3</sub>, fedotovite  $K_2Cu_3O[SO_4]_3$ , tenorite CuO, Cu-bearing anglesite Pb[SO<sub>4</sub>] and native gold. Vergasovaite grows directly on chalcocyanite and dolerophanite crystal surfaces where it was formed after chalcocyanite and before tenorite.



Fig. 4 BSE-image of vergasovaite: the white inclusions are Cu-bearing anglesite. Sector and oscillatory zonations are visible.

λ (nm)	<b>R</b> 1	<b>R</b> 2
400	10.64	13.03
420	10.39	12.62
440	10.16	12.17
460	9.95	11.86
470	9.85	11.69
480	9.8	11.61
500	9.66	11.4
520	9.55	11.22
540	9.42	11.06
546	9.4	11.01
560	9.35	10.93
580	9.26	10.78
589	9.22	10.76
600	9.16	10.68
620	9.11	10.63
640	9.07	10.52
650	9	10.48
660	9.03	10.47
680	9	10.42
700	9.03	10.45

*Tab. 3* Reflectance data for vergasovaite measured in air (SiC standard).

Optical and other physical properties

The optical and other physical properties of vergasovaite are summarized in table 1. Rare single crystals are flattened, shortened prisms, slightly elongated parallel the *c*-axis and do not exceed 0.3 mm in size. Single crystals are frequently intergrown (Fig. 2), sometimes forming radiating aggregates up to 0.6 mm overgrowing the matrix. The crystal used for the structure determination was also studied using an optical goniometer. The measured forms (Tab. 2) are: a pinacoid {100}; three prisms: {110} {120} {130}; and three bipyramids: {111} {263} {315}. It seems as if the larger crystals display a reduced variety of forms, with the pinacoid and prisms dominating, compared to the smaller crystals. Presumably the latter grew slower and later than the large crystals. Twinning was not observed. An drawing of ideal vergasovaite is shown in figure 3.

Vergasovaite is transparent and has a megascopic olive-green colour. The streak is light-yellow, the lustre is vitreous and no fluorescence was observed. Micro-indentations were done with a load of 25 g on a LEITZ Durimet 2 microhardness tester. Six perfect, slightly fractured indentations on randomly oriented polished surfaces of three grains give a mean VHN of 357 kg/mm<sup>2</sup> with a range of 302–413 kg/mm<sup>2</sup> which corresponds to a Mohs' hardness of 4–5. Measurements perpendicular to a face subparallel to (010) yielded a Mohs' hardness of 5–5.5 (personal comment D.K. Cherbachev).

I <sub>est</sub> <sup>1</sup>	d <sub>meas</sub>		d <sub>calc</sub> <sup>2</sup>	I <sub>calc</sub> <sup>2</sup>	hkl <sup>2,3</sup>
30	3.71		3.71	33	200
10	3.580		3.580	37	201
60	3.391		3.377	69	020
60	3.342		3.360	24	113
20	2 226	ſ	3.258	12	202
30	3.220	í	3.252	32	210
100	3.077		3.096	100	104
30	2.990		2.998	56	121
60	2 5 4 2	ſ	2.558	27	105
00	2.342	Ĩ	2.546	12	123
60	2.500		2.498	54	220
20	2 400	ſ	2.434	9	301
50	2.400	Ĩ	2.398	15	024
10	2.381		2.392	2	115
60	2.275		2.282	31	124
30	2.167		2.171	33	106
10	2.126		2.128	11	131
10	2.057		2.067	4	116
10	1.903		1.906	3	231
10	1.849		1.852	5	232
10	1.764		1.768	7	315
10	1.715		1.7175	5	403
10	1 678	ſ	1.6885	28	040
10	1.078	ĩ	1.6801	15	226
30	1.606		1.6100	9	325
5	1.582		1.5816	18	422
10	1.544		1.5478	16	208
10	1.534		1.5368	2	240
10	1.518		1.5206	10	028
10	1.389		1.3934	11	327
10	1 387	ſ	1.3810	4	513
10	1.507	<u> </u>	1.3734	5	318
<sup>1</sup> Inte	nsities assi	gned	by visual es	timate.	
<sup>2</sup> Calo	culated on	the ba	asis of the st	ructural c	lata
pub	lished by E	BERLE	PSCH et al. (	in press) t	by the
program LAZY PULVERIX (Yvon et al., 1977)					

*Tab. 4* Powder data for vergasovaite obtained with a Gandolfi camera (114.5 mm diameter,  $FeK_{\alpha}$  X-radiation). Intensities were visually estimated.

Using the empirical formula derived from electron microprobe analysis (Tab. 7) and the single-crystal (CAD4 data) cell parameters (Tab. 5), the calculated density of vergasovaite was determined to be 4.32 g/cm<sup>3</sup>. The density could not be measured because grains of suitable size contain inclusions of Cu-bearing anglesite (Fig. 4) and sink in the Clerici liquid.

Indices were chosen using the calculated inten-

for neutral Cu, Mo, S and O.

sities.

Vergasovaite has high refractive indices (> 1.9), thus it was treated as an ore mineral in obtaining its optical data (the mineral reacts with phosphorous imersion liquids and dissolves in selenium-sulfur melts). Under plane polars the colour is grey with ubiquitous internal reflections

	CAD4 data 1	Powder data 1,2		
Crystal system	orthorhombic	orthorhombic		
Space group	Pnma	Pnma		
a [Å]	7.406(1)	7.420(3)		
b [Å]	6.739(2)	6.741(2)		
c [Å]	13.573(2)	13.548(5)		
$V[Å^3]$	677.4(2)	677.6(2)		
Z	4	4		
Orientation c al. (in press) w	> a $>$ b according which corresponds	to BERLEPSCH et to the standard		
space group setting Pnma.				
<sup>2</sup> Cell parameters were obtained by a least squares				
refinement performed with the program from				
APPLEMAN an	d Evans (1973).			

Tab. 5 Cell parameters of vergasovaite.

which are light green to colourless. Internal reflections mask the anisotropy and the bireflectance is measurable but not discernible. Pleochroism in transmitted light changes from olive-green (parallel to the morphological elongation) to a yellowish to brownish green (perpendicular to the morphological elongation). Pleochroism in reflected light is absent.

The refractive indices calculated from the reflectance values at 589 nm (and assuming k = 0) are 1.87–1.98 (average 1.925). The measurements were made on the best of three crystals, but we cannot be sure that the constants (R and n) represent extreme values for the species, nor that the average corresponds to the mean. The reflectance values are listed in table 3.

The mean  $n_{calc}$  obtained from the Gladstone-Dale relationship (MANDARINO, 1979) and using the chemical data given in table 7, the revised Gladstone-Dale constants in MANDARINO (1981), and the calculated density (4.32 g/cm<sup>3</sup>) is 1.819. The Mandarino Compatibility Index is poor which can be explained by the reasons given above as well as the unreliability of the Gladstone-Dale constants, especially for MoO<sub>3</sub>.

The X-ray powder diffraction pattern was obtained with a Gandolfi camera (114.5 mm diameter) using Mn-filtered Fe $K_{\alpha}$  X-radiation ( $\lambda =$ 1.93728 Å). The X-ray powder diffraction data for vergasovaite are listed in table 4. Relative intensities of the reflections were assigned by visual estimate. In table 5 the unit-cell parameters obtained from CAD4 and powder data are listed. Considering the range of Mo <-> S substitution, the agreement between the two cell parameter sets is excellent.

*Tab. 6* Interpretation of the absorption bands in the IR-spectrum (Fig. 5) of vergasovaite.

Peaks [cm <sup>-1</sup> ]	Bond type	Vibration type
462,480	Mo-O	bending
533,602	Cu–O	stretching
624, 637, 670, 690	S-O	bending
817, 882, 928, 963	Mo-O	stretching
1012, 1086, 1130, 1150, 1200	S-O	stretching

## **Chemical composition**

About 1 mg of vergasovaite was mixed with 150 mg KBr and pressed to a 10 mm diameter pellet. The IR-spectrum was measured from 400–4000 cm<sup>-1</sup> against pure KBr on a SPECORD 75 IR-spectrometer. The results showed vergasovaite to be free of  $H_2O$  or OH (Fig. 5). The interpretation of the absorption bands is given in table 6.

The results of the chemical analyses are summarized in table 7.18 analyses were performed on one crystal with a CAMEBAX SX-50 EMP operated at 20 kV, 30 nA, and a spot-size beam. The main elements measured were Cu, Mo, and S, and the minor constituents are Zn, V and Pb. BER-LEPSCH et al. (in press) also found independently FeO  $\leq 0.09$ , TiO<sub>2</sub>  $\leq 0.06$ , TeO<sub>2</sub>  $\leq 0.16$  wt% and by using WDS scans, no additional element with atomic number  $\geq 10$  was detected.

The empirical formula (on the basis of nine oxygens p.f.u., and the mean of 18 analyses on one grain) is  $(Cu_{2.82(7)}, Zn_{0.10(7)}, Pb_{0.01(1)})_{\Sigma = 2.92(9)} O[(Mo_{0.79(7)}, S_{0.20(5)}, V_{0.04(3)})_{\Sigma = 1.04(3)}O_4][SO_4]$  and matches very well with the formula  $Cu_3O$  [( $Mo_{0.742(7)}, S_{0.258(7)}$ )O<sub>4</sub>][SO<sub>4</sub>] obtained from the



*Fig.* 5 IR-spectrum of vergasovaite. The new mineral is hydroxyl- and water-free.

Constituent	Wt%	Ra	nge	Std. dev.	Probe Standard
		Min	Max	$\sigma_{n+1}$	
CuO	49.81	47.88	52.35	1.17	$Cu_2O$ (synthetic)
ZnO	1.76	0.15	4.19	1.26	ZnSe (synthetic)
SO	21.44	19.89	23.12	1.03	Ba[SO <sub>4</sub> ] (synthetic)
MoO	25.29	21.01	28.73	2.29	$Ca[MoO_4]$ (synthetic)
V <sub>2</sub> O <sub>5</sub>	0.88	0.10	1.81	0.59	vanadinite
PbO	0.63	0.14	2.06	0.74	crokoite
Sum	99.81	98.48	99.92	0.44	

Tab. 7 Chemical data for vergasovaite and Cu-bearing anglesite.

Empirical formulae calculated on the basis of nine oxygens p.f.u. (EMP data). N = 18 analyses for all constituents except for PbO (N = 6); acceleration voltage: 20 kV; beam current: 30 kV; beam size: spot analyses.

A:  $(Cu_{2.82(7)}, Zn_{0.10(7)}, Pb_{0.01(1)})_{\Sigma = 2.91(9)}O[(Mo_{0.79(7)}, S_{0.20(5)}, V_{0.04(3)})_{\Sigma = 1.04(3)}O_4][SO_4]$ 

B:  $(Cu_{2.87(6)}, Zn_{0.06(2)})_{\Sigma = 2.92(6)}O[(Mo_{0.79(7)}, S_{0.21(7)}, V_{0.04(3)})_{\Sigma = 1.03(2)}O_4][SO_4]$ 

Formula derived from the crystal structure determination: C:  $Cu_3O[(Mo_{0.742(7)}, S_{0.258(7)})O_4][SO_4]$ 

Simplified formula:

D:  $Cu_3O[(Mo,S)O_4][SO_4]$ 

A, D: this study; B, C, D: BERLEPSCH et al. (in press)

Two analyses of Cu-bearing anglesite (all wt%; same analytical conditions as for vergasovaite): CuO: 2.10, 1.72; ZnO: n.d., 0.06; PbO: 70.83, 70.67; SO<sub>3</sub>: 26.74, 27.37; V<sub>2</sub>O<sub>5</sub>: n.d., 0.01; Sum: 99.68, 99.83. Empirical formula calculated on the basis of four oxygens p.f.u.:  $(Pb_{0.94}, Cu_{0.07})_{1.01}S_{1.00}O_4$ 

crystal structure determination and the previously obtained empirical formula  $(Cu_{2.87(6)}, Zn_{0.06(2)})_{\Sigma} =$  $_{2.92(6)}O$  [ $(Mo_{0.79(7)}, S_{0.21(7)}, V_{0.04(3)})_{\Sigma} = 1.03(2)O_4$ ][SO4] (BERLEPSCH et al., in press). In all cases the simplified formula is  $Cu_3O$ [(Mo,S) O4][SO4].

Both set of analyses show good negative cor-

Camebax: N = 18; R = -0.73 (--) Jeol: N = 57; R = -0.84 (-----) 30 28 MoO<sub>3</sub> (wt%) 26 24 22 20 26 19 20 21 22 23 24 25 SO3 (wt%)

*Fig.* 6 Regression plot for  $MoO_3$  wt% against  $SO_3$  wt%. Vergasovaite displays an unusual substitution of  $S^{6+}$  for  $Mo^{6+}$  in the tetrahedral sites.

relations (Fig. 6; JEOL, N = 57, R = -0.84; CAME-BAX, N = 18, R = -0.73) between MoO<sub>3</sub> and SO<sub>3</sub> values (wt%) which indicates a considerable and unusual substitution of Mo<sup>6+</sup> by S<sup>6+</sup> (BERLEPSCH et al., in press). The Mo–S- and V-inhomogeneities can also be seen in the element distribution maps



*Fig.* 7 Element distribution maps for vergasovaite to visualize sector and oscillatory zonations. Each map has an individual intensity scale which cannot be compared directly to the other ones.

a) hydration	
Tolbachite: $CuCl_2 \rightarrow$ Chloromagnesite: $\rightarrow$ MgCl	Eriochalcite: $CuCl_2 \cdot 2 H_2O$ Bischofite: $MgCl_2 \cdot 6 H_2O$
b) dissolution and re-c	leposition
Piypite: $K_4Cu_4O_2[SO_4]_4 \cdot \rightarrow \cdot (Na,Cu)Cl$ Tenorite:	Chalcantite + Mitscherlichite: $Cu[SO_4] \cdot 5 H_2O + K_2CuCl_4 \cdot 2 H_2O$ Atacamite/paratacamite/
CuO →	botallackite: $Cu_2Cl(OH)_3$

*Tab. 8* Transformation products of some anhydrous minerals after hydration and/or hydrolysis.

in figure 7. Note that the scales are relative ones and that the intensities cannot be compared directly from one element distribution map to another. In vergasovaite we observe both sector and oscillatory zonation, a phenomenon which is well known in crystals that grow rapidly, e.g., nabokoite  $Cu_7[TeO_4][SO_4]_5 \cdot KCl$  and atlasovite  $Cu_6Fe[BiO_4][SO_4]_5 \cdot KCl$  (both POPOVA et al., 1987). From figure 7 the important role of V in the sector zonation is clearly visible. In contrast to this, Mo and S (and Zn) cause the oscillatory zonation (cf. figure 4 too). Due to the distribution of anglesite inclusions in vergasovaite, we suppose Pb is involved in the oscillatory zonation too (Figs 4 and 7). To underline the unusual Pb <-> Cu substitution, two analyses of Cu-bearing anglesite are listed in table 7 as well. The formula of Cu-bearing anglesite derived from the EMP data is  $(Pb_{0.94}, Cu_{0.07})S_{1.00}O_4.$ 

## **Chemically related compounds**

Vergasovaite is isotypic with the fully oxidized synthetic Cu-molybdate Cu<sub>3</sub>O[MoO<sub>4</sub>]<sub>2</sub> described by KIHLBORG and NORRESTAM (1972) which has not previously been found in nature. In vergasovaite complete substitution of S<sup>6+</sup> for Mo<sup>6+</sup> reduces the average T2–O distance from 1.75 to 1.47 Å. The average distance T1–O in the mineral is 1.71 Å which agrees well with 26% S<sup>6+</sup> substitution for Mo<sup>6+</sup>. It is also striking that the Mo–O distances in the synthetic compound are 1.778 Å for T1 and 1.749 Å for T2. Thus the smaller site in the synthetic compound accepts a higher S substitution in the mineral (BERLEPSCH et al., in press).

Apart from this, vergasovaite  $Cu_3O[(Mo,S)O_4]$ [SO<sub>4</sub>] is the first mineral which contains Cu, Mo, S, and O as its main elements, although several sul-

fates and molybdates of copper have been known for a long time. Among the 67 minerals containing Cu and  $SO_4$  as their main elements/element groups, only chalcocyanite Cu[SO<sub>4</sub>] (WILDNER and GIESTER, 1988) and dolerophanite  $Cu_2O$  $[SO_4]$  (EFFENBERGER, 1985) are H<sub>2</sub>O- and hydroxyl-free sulfates of copper. Thus, an end-member compound  $Cu_3O[SO_4]_2$  is obviously not known. Several hydrated and/or hydroxylated sulfates of copper exist including antlerite  $Cu_3SO_4$  $(OH)_4$  (HAWTHORNE et al., 1989) which has the same space group and related lattice parameters as vergasovaite but a very different structure. Only five minerals are known which contain Cu and MoO<sub>4</sub> as their main elements/element groups two of which are hydroxylated molybdates lindgrenite of copper:  $Cu_3[MoO_4]_2(OH)_2$ (HAWTHORNE and EBY, 1985), and szenicsite  $Cu_3[MoO_4](OH)_4$  (FRANCIS et al., 1994). As it is visible from the list presented by VERGASOVA and FILATOV (1993), hydrated and hydroxylated compounds of the LTFE result from hydration or by dissolution and redistribution due to interaction of meteoric waters with primarily anhydrous exhalates (Tab. 8).

### Conclusions

Many of the characteristic properties of vergasovaite are typical of the exhalative minerals found in the LTFE. As a direct consequence of the hightemperature formation of the exhalates,  $H_2O$ - and OH-free (anhydrous) minerals are abundant, i.e. chlorides, sulfates, arsenates, selenates and so on. The formation of hydrated and hydroxylated compounds was discussed in the previous section. Tolbachite CuCl<sub>2</sub>, ponomarevite K<sub>4</sub>Cu<sub>4</sub>OCl<sub>10</sub> and chalcocyanite Cu[SO<sub>4</sub>] are typical indicators of anhydrous associations.

Due to the high temperatures and the presence of (atmospheric) oxygen, the LTFE exhalates are highly oxidized. Thus oxysalts, i.e. oxychlorides, oxysulfates, oxyselenates, are abundant and to these we now add the mixed oxymolybdate-sulfate vergasovaite. Furthermore, high oxidation states of the elements in the various minerals predominate:  $Cu^{2+}$ ,  $Fe^{3+}$ ,  $V^{5+}$ ,  $As^{5+}$ ,  $Mo^{6+}$ . Tenorite CuO, dolerophanite  $Cu_2O[SO_4]$  and melanothallite  $Cu_2OCl_2$  are also indicative of the oxidizing conditions.

Another characteristic feature of the LTFE minerals is related to the quickly changing conditions under which the minerals were formed. In fumaroles, there are distinct and frequent changes in temperature, pressure, redox-potential, and composition of the gas. Newly formed minerals are highly dispersed and sector- and oscillatory zonation are as abundant as are many inclusions. Complex minerals from both the cationic and anionic point of view are characteristic for LTFE, i.e. arsenate-phosphates, chloride-vanadates, -sulfates, and sulfates-fluorides, sulfate-molybdates, oxy-selenate-chlorides, chloro-tellurate-sulfates and -bismuthate-sulfates in combination with Cu and with K, Na, Cs, Rb, Zn, Pb, Mg, Fe, Al, V.

The extreme concentration of certain elements, compared to the average concentrations in other rocks, is supposed to be related to a gradual accumulation of primarily rare elements. This is caused by both the exhalative activity and the interaction with exogenic factors. In particular, it is supposed that repeated cycles of precipitation, dissolution, infiltration, and re-precipitation led to the accumulation and finally to the observed high concentrations of particular elements. For instance the Mo-concentrations of 200-300 ppm in the exhalative Cu-ores are two orders of magnitude higher than the Clarke values for basic rocks: 1.4 ppm (VINOGRADOV, 1962); oceanic and upper crust: 0.8-1 ppm (TAYLOR and MCLENNAN, 1985); or basalts of oceanic islands: 2.4 ppm (SUN and MCDONOUGH, 1989). In addition, an increase of the Mo-concentrations in exhalative Cu-ores of the LTFE was observed within a very short period: from 240 ppm before 1980 (NABOKO and GLAVATSKIKH, 1983) up to 340 ppm in 1988 (NABOKO and GLAVATSKIKH, 1988). Such significant changes in concentrations cannot be explained solely by the change of composition of the gas, which is approximately constant for the short period indicated, but by a gradual accumulation as explained above.

Vergasovaite incorporates all these features: it is a rare, anhydrous, fully oxidized Cu-oxy-molybdate-sulfate with sector and oscillatory zonation that contains inclusions of Cu-bearing anglesite.

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