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# The crystal structure of averievite, $Cu_5O_2(VO_4)_2 \cdot nMX$ : comparison with related compounds

G. L. STAROVA, S. V. KRIVOVICHEV, V. S. FUNDAMENSKY AND S. K. FILATOV

Departament of Crystallography, Saint-Petersburg State University, University Emb. 7/9, St.Petersburg, 199034, Russia

## Abstract

The crystal structure of averievite,  $Cu_5O_2(VO_4)_2$  nMX has been determined. Trigonal system, space group P3, a = 6.375(1), c = 8.399(1) Å, V = 295.6(1) Å<sup>3</sup>, Z = 1,  $D_x = 4.01(1)$  g/cm<sup>3</sup>. The atomic arrangement is characterized by infinite nets parallel to (001) composed of  $[OCu_4]^{6+}$  tetrahedra linked via corners in hexagonal rings. The bases of neigbouring tetrahedra are in one plane and their non-shared corners are turned to the opposite sides. The  $[VO_4]^{3-}$  tetrahedra are attached to the bases of  $[OCu_4]$  tetrahedra. There are large (R >3.2 Å) channels in the structure where large molecular particles can enter. The comparison of the averievite structure with related compounds (in particular, copper oxovanadates) is given from the point of view of  $[OT_4]$  polyion crystal chemistry.

KEYWORDS: averievite, crystal structure.

#### Introduction

AVERIEVITE was found as a product of post eruption volcanic activity of the Main Tolbachik Fracture Eruption (Kamchatka, 1975-76). It was discovered among new exhalation products of Second North Fracture cone. The mineral occurs as rare isolated grains associated with brush-like agregates of piypite, alumokljuchevskite, langbeinite and pyroclastic materials (Vergasova et al., 1996). The averievite crystals are the non-transparent black pseudohexagonal plates up to  $0.3 \times 0.3 \times 0.1 \text{ mm}^3$ . The chemical composition has been studied by electron microprobe analysis (microsond 'Camebax'). The chemical studies were carried out using the following standards: Fe<sub>2</sub>O<sub>3</sub> (Fe), CuO (Cu), Ca-molybdate (Mo), synthetic Cl-apatite (Cl), orthoclase (K), synthetic pollucite (Cs), synthetic V<sub>2</sub>O<sub>5</sub> (V), Rb-lepidolite (Rb). The means (and ranges) from 17 analyses are: CuO 62.01 (63.41-60.06), Fe<sub>2</sub>O<sub>3</sub> 0.62 (1.06-0.35), K<sub>2</sub>O 0.98 (1.27-0.65), Cs<sub>2</sub>O 3.35 (4.11-2.70), Rb<sub>2</sub>O 0.82 (1.13-0.55), V<sub>2</sub>O<sub>5</sub> 23.84 (25.16-22.67), MoO<sub>3</sub> 1.51 (3.13-0.42), Cl 7.56 (7.93-7.26), O=Cl<sub>2</sub> -1.70 [-(1.79-1.64)], sum 98.99 (100.54-97.82). Single crystal structure analysis enabled us to understand and explain the alignment of the elements obtained by the chemical study.

Mineralogical Magazine, June 1997, Vol. 61, pp. 441–446 © Copyright the Mineralogical Society

#### Experimental

The structure analysis of a crystal of size  $0.3 \times 0.2$  $\times$  0.1 mm<sup>3</sup> was carried out using a Weissenberg camera and 4-circle diffractometer 'Syntex P21' and gave the following data: 318 unique reflections (Mo- $K\alpha$  radiation,  $\sin\theta/\lambda < 0.80$ ,  $I > 3\sigma I$ ), trigonal system, space group P3, a = 6.375(1), c = 8.399(1) Å, V =  $295.6(1) \text{ Å}^3$ , Z = 1, D<sub>x</sub> = 4.01(1) g/cm<sup>3</sup>. The structure was solved by direct methods and refined by fullmatrix least-squares method with anisotropic thermal parameters to an R-factor of 0.052. The program complex 'CSD' was used (Akselrud et al., 1989). The positions of oxygen and channel atoms were localized on difference Fourier maps. The final atomic positions are given in Table 1 (tables of the calculated and observed crystal structure factors and anisotropic atomic parameters are available from the author or editorial office).

#### Discussion

From the chemical point of view averievite is a copper(II) oxovanadate(V). Its crystal structure contains the oxygen atoms which do not enter the  $[VO_4]$  groups and form tetrahedral  $[OCu_4]^{6+}$  complexes as central atoms. Therefore, it may be

Atom	x/a	y/b	z/c	U(is/eq)*100
Cul	0	0	0	1.8(3)
Cu2	0.338(2)	0.161(2)	0.280(2)	0.96(11)
Cu3	2/3	1/3	0.5492(9)	2.3(4)
Cu4	1/3	2/3	0.01774	6.333
Cu5	1/3	2/3	0.53143	6.333
VI	2/3	1/3	-0.032(2)	0.8(4)
V2	1	0	0.584(2)	0.4(3)
01	1	0	0.785(9)	3.2(24)
O2	2/3	1/3	0.339(8)	0.887
O3	0	0	0.244(6)	1.013
O4	2/3	1/3	-0.225(8)	0.9(17)
05	0.379(8)	0.186(9)	0.031(4)	2.9(19)
CH	1/3	2/3	0.27140	6.333
C12	1/3	2/3	0.76793	6.333

TABLE1. Atomic parameters for averievite, Cu<sub>5</sub>O<sub>2</sub>(VO<sub>4</sub>)<sub>2</sub>·CuCl<sub>2</sub>

 $U(eq) = \frac{1}{3}[U_{11}a^{*}2a^{2} + \dots 2U_{23}b^{*}c^{*}bc \cos a]$ 

Occupations	
CH	1.02(4) Cl
Cl2	0.97(5) Cl
Cu4	0.69(7) Cu
Cu5	0.32(8) Cu

described using a new crystal chemical approach based on the subdivision in the crystal structures of the polyions formed from oxocentred  $[OT_4]$  tetrahedra (Filatov et al., 1992; Krivovichev et al., 1995). The atomic arrangement in the averievite structure is characterized by infinite two-dimensional nets  $[O_2Cu_5]^{6+}$  parallel to (001). These nets are composed of oxocentred [OCu4]<sup>6+</sup> tetrahedra linked via corners in hexagonal rings (Figs. 1,2). The bases of neighouring tetrahedra are in the (001) plane and their non-shared corners are turned to the opposite sides. The structural formula of nets, according to rules elaborated by Liebau (1985) (see also IUCr recommendations; Lima-de-Faria et al., 1990), is  $\{uB, l_{\infty}^{2}\}[O_{2}Cu_{5}]^{6+}$ . The  $[VO_{4}]$  tetrahedra are attached to the bases of oxocentred tetrahedra. The oxocentred [O<sub>2</sub>Cu<sub>5</sub>]<sup>6+</sup> nets and [VO<sub>4</sub>] tetrahedra may be represented as main structural units in the averievite structure. They form its framework so that there are large channels (R >3.2 Å) with a threefold axis in the centre where large molecular particles can enter. After determination of atomic positions of nets and [VO<sub>4</sub>] groups, the difference Fourier maps show four strong peaks in these channels which have no bonds with the main framework. Distances between the peaks were 1.99, 2.10, 2.13 and 2.18 A. The comparison of the framework chemical formula with results obtained by electron microprobe

analysis showed that there was abundance of Cu atoms and some quantity of Cl, K, Rb and Cs, which do not substitute the framework elements according to crystal chemical reasons. It was logical to place



FIG. 1. Projection of the structure of averievite on (001) plane (representation in  $|OT_4|$  aspect).



Fig. 2. Projection of the structure of averievite on (100) plane (representation in  $[OT_4]$  aspect).

these atoms in the structure channels as CuCl, CuCl<sub>2</sub> or (K,Rb,Cs)Cl. The anion-cation distance for CuCl is 2.05 Å, for CuCl<sub>2</sub> — 2.3 Å and for KCl, RbCl, CsCl — 2.6, 2.7 and 2.9 Å, respectively (Wells, 1986). Refinement of the channel position occupation shows that one Cu(II) atom (in sum) occupies two positions and two chlorine atoms occupy two other ones (Table 1). Therefore, the 'molecules' of CuCl<sub>2</sub>

are located in structure channels of the studied crystal so that its chemical formula is  $Cu_5O_2(VO_4)_2$ ·CuCl<sub>2</sub>. The wide and exotic range of alkali elements in the chemical analyses of averievite can be explained by the location of these atoms in the structure channels. In general, the chemical formula of averievite is  $Cu_5O_2(VO_4)_2 \cdot nMX$ , where *MX* are particles depending on the environment of mineral formation.

The similar nets of oxocentred tetrahedra may be subdivided in the structure of the synthetic compound  $Pb_2Cu_3O_2(NO_3)_2(SeO_3)_2$  (Effenberger, 1986). According to the  $[OT_4]$  crystal chemical approach we describe this structure as composed of oxocentred  $[O_2Cu_3Pb_2]^{6+}$  nets,  $(SeO_3)^{2-}$  and  $(NO_3)^-$  groups. The nets consist of  $[OCu_3Pb]^{6+}$  tetrahedra so that Cu atoms are shared corners and Pb atoms are non-shared ones. The oxygen bases of  $(SeO_3)^{2-}$  pyramids are attached to oxocentred tetrahedra in the same way as  $[VO_4]$ groups in the averievite structure.  $(NO_3)^-$  groups occur in channels of the structure as MX particles in averievite. Topologically equivalent oxocentred  $[O_2Cu_5]^{6+}$  nets are found in the structure of the natural compound NaCu5O2(SeO3)2Cl3 which was also discovered as an exhalation product of the Tolbachik volcano (Shuvalov et al., 1995). But in this compound the orientation of non-shared corners of oxocentred tetrahedra is different from averievite nets.

The left part of the averievite formula is identical to stoiberite,  $Cu_5O_2(VO_4)_2$ , a mineral from exhalation products of the Izalco volcano (Salvador) (Birnie and Hughes, 1979). But X-ray data and the crystal structure of this mineral are different from averievite (Table 2) (Shannon and Calvo, 1973). In stoiberite the 'free' oxygen atoms are also contained and they are also coordinated by four copper atoms. Using  $[OT_4]$  crystal chemistry, we can consider its structure as composed of oxocentred polyion and  $[VO_4]$ groups. But in stoiberite the oxocentred structural unit is a 1-dimensional one (Fig. 3*d*). It consists of two pyroxene-like chains linked by edge linkage.

TABLE 2. Comparison of some parameters for averievite and stoiberite

Parameter	Averievite	Stoiberite
Chemical formula	$Cu_5O_2(VO_4)_2 \cdot nMX$	$Cu_5O_7(VO_4)_2$
Space group	P3	P2/n
a, Å	6.375(1)	15.654(15)
b, Å		6.054(4)
<i>c</i> , Å	8.399(1)	8.385(11)
β, °	-	102.29(12)
Z	1	4
D, g/cm <sup>3</sup>	4.01	4.96
Type of oxocentred	$\{uB, 1^{2}_{\infty}\}[O_{2}Cu_{5}]^{6+}$ sheets	$\{uB, 2^{1}_{\alpha}\}[O_{2}Cu_{5}]^{6+}$ chains
polyions in structure	parallel to (001)	parallel to [010]



FIG. 3. Types of oxocentred tetrahedral [OT<sub>4</sub>] polyions in the structures of copper oxovanadates (see text).

According to the principles of Liebau (1985), this fragment may be described as unbranched zweier double chain {uB,  $2_{\infty}^{1}$ }[O<sub>2</sub>Cu<sub>5</sub>]<sup>6+</sup>. Other known copper oxovanadates are fingerite Cu<sub>11</sub>O<sub>2</sub>(VO<sub>4</sub>)<sub>6</sub> (Hughes and Hadidiacos, 1985; Finger, 1985), the natural compound Cu<sub>4</sub>O<sub>2</sub>(VO<sub>4</sub>)Cl (Starova et al., 1995) and the synthetic compound  $Cu_3O(V_2O_7)(H_2O)$  (or  $Cu_3V_2O_8(H_2O))$  (Leblanc and Ferey, 1990). Their structures also contained 'free' oxygen atoms coordinated by four copper atoms. The analysis of their structures shows that there are 5 different types of oxocentred polyions in copper oxovanadates (Fig. 3): single tetrahedron  $[OCu_4]^{6+}$  in fingerite (Fig. 3a); single pyroxene-like chains in Cu<sub>3</sub>O(V<sub>2</sub>O<sub>7</sub>)(H<sub>2</sub>O) (Fig. 3b); single edgesharing chains in natural  $Cu_4O_2(VO_4)Cl$  (Fig. 3c); double pyroxene-like chains in stoiberite (Fig. 3d); and  $[O_2Cu_5]^{6+}$  sheets in averievite (Fig. 3e). We do not consider other oxovanadates which have no copper (e.g. kombatite, Pb<sub>14</sub>O<sub>9</sub>(VO<sub>4</sub>)<sub>2</sub>Cl<sub>4</sub>) but in their structures the oxocentred polyions may be also subdivided (Krivovichev et al., 1995).

The coordination polyhedra of Cu atoms in averievite are characterized by strong Jahn-Teller distortions. There are two types of Cu coordination: fivefold (a perfect trigonal bipyramid) for Cu1 and Cu3, and fourfold (a distorted square) for Cu2 (Table 3). The trigonal bipyramids are usual for copper vanadates, but more often distorted octahedra occur (Finger, 1985; Leblanc and Ferey, 1990; Shannon and Calvo, 1973). A square occurs only infrequently in copper vanadates. The Cu polyhedra are connected with each other so that the square has common edges with both trigonal bipyramids forming a copper-oxygen complex (Fig. 4). The vanadium tetrahedra are slightly distorted and have typical distances and angles. The coordination of 'free' oxygen atoms O2 and O3 is tetrahedral with mean angles Cu-O-Cu 109.3° and 108.36° for O2 and O3, respectively. The bond distances O--Cu for 'free' oxygen (mean 1.895 Å) are shorter than O-Cu distances for oxygen from  $(VO_4)^{2-}$  groups (2.015 Å) approximately on 6% (0.12 Å).

### Acknowledgements

This work was supported by the ISF, grant NVV000.

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Geometry of copper coordination polyhedra						
Cul-	O1 O5 O3	1.806 2.108 3x 2.048 2.036	O1 - O5 O5 - Cu1 - O3 O5 - Cu1 - O5' O5 - Cu1 - O3	96.99 3x 180.00 118.54 3x 83.01 3x		
Cu2-	· O2 O5 O6 O3	1.88(2) 2.11(4) 1.93(4) 1.892(15) 1.953	$\begin{array}{c} O2 - Cu2 - O5\\ O2 - Cu2 - O6\\ O2 - Cu2 - O3\\ O5 - Cu2 - O3\\ O5 - Cu2 - O6\\ O5 - Cu2 - O3\\ O6 - Cu2 - O3 \end{array}$	99.1(19) 80.4(17) 173.3(17) 179.1(19) 86.8(17) 93.7(15)		
Cu3-	O2 O4 O6	1.76(7) 1.89(7) 2.03(4) 3x 1.948	O2 - Cu3 - O4 O2 - Cu3 - O6 O4 - Cu3 - O6 O6 - Cu3 - O6'	180.0(18) 80.3(16) 3x 99.7(16) 3x 117.2(15) 3x		
Cu4	C11 C12	2.184 1.986 2.085	Cl1 -Cu4 - Cl2	180.00		
Cu5	C11 C12	2.130 2.098 2.114	C11 - Cu5 - Cl2	180.00		
Georr	netry of [VO <sub>4</sub> ] groups					
VI-	O4 O5	1.63(7) 1.67(4) 3x 1.66	O4 - V1 - O5 O5 - V1 - O5'	108.2(23) 3x 110.7(24) 3x 109.5		
V2-	O1 O6	1.69(8) 1.79(4) 3x 1.765	O1 - V2 - O6 O6 - V2 - O6'	110.7(20) 3x 108.3(17) 3x 109.5		
Geometry of oxocentred [OCu <sub>4</sub> ] <sup>6+</sup> tetrahedra						
02–	Cu2 Cu3	1.88(2) 3x 1.76(7) 1.85	Cu2 - O2 - Cu3 Cu2 - O2 - Cu2'	105.3(20) 3x 113.3(20) 3x 109.3		
O3-	Cul Cu2	2.048 1.892(15) 3x 1.931	Cu1 - O5 - Cu2 Cu2 - O5 - Cu2'	99.32 3x 117.4(17) 3x 108.36		

TABLE 3. Interatomic distances (Å) and bond angles (°) for averievite

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[Manuscript received 8 February 1996: revised 3 September 1996]