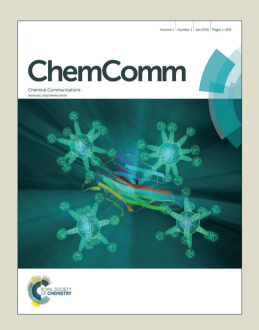
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Oxocentered Cu(II) lead selenite honeycomb lattices hosting Cu(I)Cl₂ groups obtained by chemical vapor transport reactions†

Vadim M. Kovrugin, a,b Marie Colmont, Oleg I. Siidra, Olivier Mentré, Alexander Al-Shuray, Vladislav V. Gurzhiy and Sergey V. Krivovichev

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Chemical vapor transport (CVT) reactions were used to prepare three modular mixed-valent Cu(I)–Cu(II) compounds, $(Pb_2Cu^{2+}{}_9O_4)(SeO_3)_4(Cu^+Cl^2)Cl_5 \qquad (1), \\ (PbCu^{2+}{}_5O_2)(SeO_3)_2(Cu^+Cl_2)Cl_3 \qquad (2), \quad and \quad (Pb_xCu^{2+}{}_{(6-x)}O_2)(SeO_3)_2(Cu^+Cl_2)K_{(1-x)}Cl_{(4-x)} \ (x=0.20) \ (3). \ In their crystal structures chains of anion-centered (OCu^{2+}{}_4) and (OCu^{2+}{}_3Pb) tetrahedra form honeycomb-like double layers with cavities occupied by linear <math display="inline">[Cu^+Cl_2]^-$ groups.

Inorganic copper oxocompounds attract considerable attention due to their interesting structural and physical properties1 as well as mineralogical and geochemical importance.² Of special interest are mixed-valent Cu(I)-Cu(II) systems with separate symmetrically independent monovalent and divalent copper sites due to their contrasted coordinations combined in one crystal structure. Herein we report on the synthesis and characterization of three novel Cu(I)–Cu(II) lead oxoselenite chlorides inspired by mineralogical discoveries in such unusual geological conditions as volcanic fumaroles.^{2g,3} Here copper oxoselenites form from volcanic gases emanating from cooling magmatic chambers deep under the Earth's surface long after the period of eruptive activities. The formation of such Cu compounds in fumaroles provides a useful hint for their synthesis under laboratory conditions, in particular, the chemical vapor transport (CVT) method.⁴ The specific feature of many Cu oxoselenites is the presence in their crystal structures of oxocentered (µ4-O)Cu4 tetrahedral units that polymerize to form extended structural complexes.^{2g,3} . Likely, during these reactions the selenites and metal halides play the role of transport agent.⁵ In order to reproduce natural exhalative chemistry, in this work, we investigate formation of phases in the PbO-Cu²⁺Cl₂-Cu⁺Cl-Cu²⁺O-SeO₂ system containing Pb²⁺ cations that possess stereochemically active 6s2 lone electron pairs favouring formation segregation of structural compartments occupied by these pairs. In addition, the preference of Pb2+ ions for similar OPb4 tetrahedral units is an asset for creation of more complex edifices.

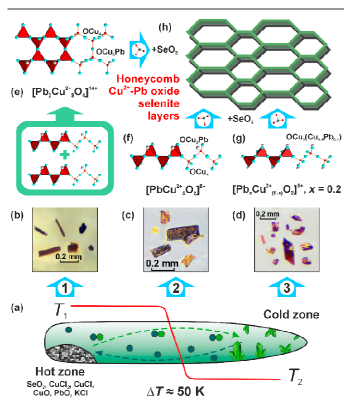


Fig. 1 General scheme of syntheses by the method of CVT reactions (a), the crystals of **1**, **2** and **3** (b–d), and the scheme of formation of honeycomb $Cu^{2\tau}$ –Pb selenite layers (e–h). The types of oxo-centered 1-dimensional units (shown in red) formed by corner-sharing OCu₄ and OCu₃Pb tetrahedra in the structures of **1**, **2** and **3** are shown. (Legend: Cu = cyan balls; Pb = grey balls). See text for details.

Crystallographic information for three novel compounds synthesized by the CVT reactions method (Fig. 1a) is summarized in Table 1. Schematic representations of coordination environments of cations in the crystal structures of **1–3** are shown in Fig. S1, ESI†. In all the compounds, Cu⁺ cations form two relatively short Cu⁺–Cl bonds (2.058–2.118 Å), which result in the formation of tightly bonded

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[CuCl₂] anionic groups that can be considered as separate structural entities. The Cu²⁺ cations have mixed oxochloride coordinations that have previously been observed in Cu oxochloride compounds, 2a with typical trends of Jahn-Teller d⁹ ions. The Cu(1) and Cu(2) sites in 1, the Cu(4) site in 2, and Cu(3) site in 3 form [CuO₄Cl₂] distorted octahedra with four short equatorial Cu-O and two long apical Cu-Cl bonds. The Cu(3) site in 1, the Cu(2) and Cu(3) sites in 2, and the Cu(1) site in 3 are octahedrally coordinated with [CuO₃Cl] squares complemented by two long Cu-Cl bonds. The Cu(4) site in 1 forms a [CuO₄Cl] trigonal bipyramids, whereas the Cu(1) site in 2 and the Cu(4) site in 3 form [CuO₃Cl₂] trigonal bipyramids. The Pb atoms have asymmetrical coordinations consisting of three strong Pb-O bonds (2.356-2.420 Å) located in one coordination hemisphere and four long Pb-Cl in another. This coordination of Pb2+ cations is typical for lead oxohalide compounds⁷ and is consistent with the presence of stereoactive lone electron pairs. In all the compounds under consideration, Se⁴⁺ cations form standard $(SeO_3)^{2-}$ selenite triangular pyramidal oxo-anions (SeO = 1.688-1.726 Å). The structure of 3 has one symmetrically independent K site with the site-occupation factor (s.o.f.) equal to 0.8. Its coordination polyhedron can be described as a distorted hexagonal bipyramid. Bond-valence sums⁸ calculated for all the sites in the crystal structures of 1-3 are in full agreement with their expected oxidations states (in valence units): Cu^+ sites -1.00-1.16, Cu^{2+} sites -1.96-2.14, Pb^{2+} sites -1.95-1.96, Se^{4+} sites -4.03-4.16, K^{+} site -0.80.

Table 1 Crystallographic data for 1, 2, and 3

Emperical formula	1 (Pb ₂ Cu ²⁺ 9O ₄)(Se O ₃) ₄ (Cu ⁺ Cl ₂)Cl ₅	2 (PbCu ²⁺ 5O ₂)(Se O ₃) ₂ (Cu ⁺ Cl ₂)Cl ₃	3 $(Pb_xCu^{2+}_{(6-x)}O_2)$ $(SeO_3)_2(Cu^+Cl_2)$ $K_{(1-x)}Cl_{(4-x)},$ x = 0.20
crystal		C2/m	
system			
space group	10.605(15)	monoclinic	15.116(1)
a (Å)	18.605(17)	18.4956(4)	15.116(1)
b (Å)	6.204(6)	6.1454(1)	6.1850(4)
c (Å)	12.673(11)	15.2985(4)	9.2672(9)
β (deg)	109.87(2)	119.311(1)	95.965(5)
$V(A^3)$	1376(2)	1516.25(6)	861.72(12)
$\rho_{\rm calc}$ (g cm ⁻ ³)	4.514	4.607	3.840
μ (mm ⁻¹)	25.78	25.02	15.52
reflection collected	7722	6910	4817
independent reflections (R_{int})	1301 (0.069)	1924 (0.024)	974 (0.029)
goodness- of-fit	0.791	1.231	1.121
$R_1[I > 2\sigma(I)]^a$	0.0257	0.0260	0.0412
wR_2	0.0349	0.0761	0.1172
R_1 (all data)	0.0475	0.0267	0.0496
wR_2	0.0378	0.0761	0.1176
largest diff.	0.951, -0.988	3.783, -2.942	4.244, -1.592
peak and hole [e Å ⁻³]	·	•	

^a $R_1 = \sum ||F_o| - |F_c|| / \sum |F_o|$; $wR_2 = \{\sum [w(F_o^2 - F_c^2)_2] / \sum [w(F_o^2)^2]\}^{1/2}$.

The high variability of cation coordinations in the structures of 1-3 makes their uniform description in terms of cation coordination polyhedra a difficult task. It is therefore more reasonable to look for more mundane consideration, *e.g.* in terms of cation arrays or coordination of anions. In addition to the O atoms associated with the SeO₃ groups, all three compounds contain additional O_a atoms (oxoanions) not bonded to Se⁴⁺ cations and tetrahedrally coordinated by four metal atoms (Cu and Pb). In the crystal structure of 1 there are two additional O atoms, O(2) and O(4), that form (OCu^{2+}_3Pb) and (OCu^{2+}_4) tetrahedra, respectively. The oxo-centered tetrahedra share common corners to form $[O_4Pb_2Cu^{2+}_9]^{14+}$ double chains depicted in Fig. 1e. These chains are growing parallel to the *b*-axis, and its common value

for the three compounds ($b \sim 6.2$ Å) denote similar arrangement between the oxo-centered building units in the full series. This type of chains of anion-centered tetrahedra is original and has not been observed in inorganic compounds previously. The SeO₃ groups are attached to the triangular bases of oxo-centered tetrahedra that results in formation of complex 1-dimensional $\{[O_4Pb_2Cu^{2+}_9](SeO_3)_4\}^6$ interconnected via Pb-O bonds into 2-dimensional metal-oxide double layers (Fig. 2b). The projection of the layers in the (bc) plane lead to an Honeycomb-like lattice of tetrahedra, even though this idealized vision neglects the disconnections occurring at the oxocentered-oxoanion contacts. In contrast, monometallic [O₂Cu₅] honeycomb-layers are reported in several compounds based upon anion-centered tetrahedra. 9 The layers are surrounded by Cl- anions in the interleaves and accommodate both linear [Cu⁺Cl₂]⁻ anions and Cl⁻ ions in the larger and smaller honeycomb-windows, respectively. The interactions between the Cl ions of the [Cu+Cl2] groups and host cationic networks are restricted to rather weak Cu^{2+} - Cl^- (> 2.975 Å), Pb^{2+} - Cl^- (> 3.396 Å) and K⁺-Cl⁻ (> 3.478 Å) bonds with bond-valences not exceeding 0.08 valence units (Tables S1-S3). Taking into account relative strength of the Cu⁺-Cl bonds and weak interactions between them and the rest of the structure, these units may be considered as guest anions embedded into complex metal oxochloride matrix based upon anion-centered tetrahedra.

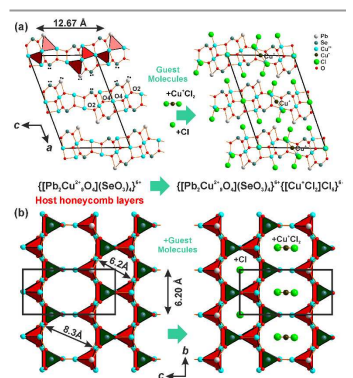


Fig. 2 General projections of the structure of **1** along the b and a axis – (a) and (b). Honeycomb layers of $\{[Pb_2Cu^{2+}{}_9O_4](SeO_3)_4\}^{6+}$ composition are hosts for the $[Cu^{+}Cl_2]^{-}$ guest species localized in the layer cavities. The $\{[Pb_2Cu^{2+}{}_9O_4](SeO_3)_4\}^{6+}$ layers are formed via interconnection of the $[Pb_2Cu^{2+}{}_9O_4]$ oxo-centered chains (red) and isolated SeO $_3$ groups (dark-green). The O(2) and O(4) designated in (a) are central oxygen atoms in OCu $_4$ and OCu $_3$ Pb tetrahedra, respectively.

A very similar 'host-guest' principle is at work in the structures of **2** and **3** as well. Here the (OCu^{2+}_3Pb) and (OCu^{2+}_4) tetrahedra share corners to produce single chains extending along the common *b*-parameter that have the $[O_2PbCu_5]^{8+}$ and $[O_2Pb_xCu^{2+}_{(6-x)}]^{8+}$ compositions, respectively (Fig. 1f,g). Together with SeO₃ groups, these chains form 1-dimensional $\{[O_2M_6](SeO_3)_2\}^{4+}$ metal-oxide (M=Cu, Pb) backbones of the structures that are arranged to form pseudo honeycomb layers. Due to the elementary single-chains, only large honeycomb windows are created that accommodate the $[Cu^+Cl_2]^-$ guest anions (Fig. 3) as observed in **1**. The structure of **3** contains additional K^+ cations located in the interlayer space between the metal

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oxoselenite chloride layers. In all three structures, lone electron pairs on the Pb^{2^+} and Se^{4^+} cations are oriented toward interlayer space, thus conforming the 'chemical scissor' principle of structural organization in compounds with lone-electron-pair cations. 10

It is noteworthy that the compounds 1 and 2 are closely chemically related that can be described by the equation:

$$(Pb_2Cu^{2+}_9O_4)(SeO_3)_4(Cu^+Cl_2)Cl_5(1) + Cu^+Cu^{2+}Cl_3 \rightarrow 2(PbCu^{2+}_5O_2)(SeO_3)_2(Cu^+Cl_2)Cl_3(2)$$

From the structural viewpoint, transition from 1 to 2 is associated with the reconstruction of the metal-oxide backbone, *i.e.* in depolymerization of anion-centered tetrahedra and splitting of the double $\left[O_4Pb_2Cu^{2+}_9\right]^{14+}$ chains into single $\left[O_2PbCu_5\right]^{8+}$ chains. This kind of structural reconstruction accompanied with the inclusion of imaginary ionic component $Cu^+Cu^{2+}Cl_3$ into metal-oxide matrix is in good agreement with the principle of dimensional reduction. It should also be noted that 2 is a synthetic analogue of allochalcoselite, the mineral first described from volcanic fumaroles of the Tolbachik volcano (Kamchatka peninsula, Russia). It should also be noted that 2 is a synthetic analogue of allochalcoselite, the mineral first described from volcanic fumaroles of the Tolbachik volcano (Kamchatka peninsula, Russia).

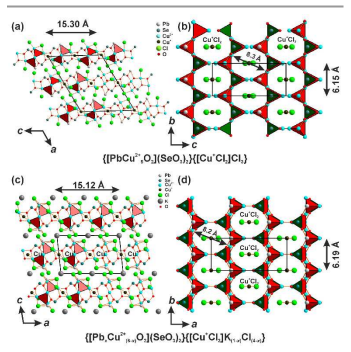


Fig. 3 General projections of the structure of **2** (a, b) and **3** (c, d). In **2**, the OCu_4 and OCu_3Pb tetrahedra (red) form single $[O_2PbCu_5]$ chains, which results in the enlargement of the c-parameter value from 12.67Å (in **1**) to 15.30Å (in **2**). In **3**, K atoms are located in the interlayer under and above the pseudohexagonal voids filled by the $[Cu^*Cl_2]$ groups. The layers (b, d) are characterized by only one type of pores with 8.3 Å and 8.2 Å in diameter in **2** and **3**, respectively.

To summarize, three novel Cu²+-Cu+ Pb oxoselenite chlorides were obtained by the chemical vapor transport reactions, which proves the efficiency of this method for the synthesis of new mixed Cu²+-Cu+ based oxyhalide compounds. These compounds described herein are based upon oxocentered mixed Pb-Cu²+ one-dimensional units of different architectures. These units determine basic topologies of the structures and influence their stability and properties. For instance, in the three compounds only corner-sharing OCu₄ and OCu₃Pb are found into chains and double chains forming hollow voids. It follows that in in all the compounds, the role of the tightly bonded [Cu+Cl₂]- anions is that of guest complexes incorporated inside metal oxide chloride units. The present study also points out that the Cu+Cl₂ groups may serve as transport agents of Cu+ in Cl-rich gaseous environments such as that observed in natural volcanic fumaroles.

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Notes and references

^a Department of Crystallography, St. Petersburg State University, University Emb. 7/9, 199034 St. Petersburg, Russia.

E-mail: s.krivovichev@spbu.ru

- b UCCS, UMR 8181, Université Lille Nord de France, USTL, F-59655 Villeneuve d'Ascq, France.
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Table of contents entry

Three different mixed Cu(I)–Cu(II) Pb selenite chlorides were obtained using chemical vapor transport reactions. Their structures belong to three different but related structure types based on oxo-centered tetrahedral units with Cu^{2+} and Pb^{2+} . Metal-oxide selenite honeycomb layers with pores varying in diameter from 6.2 to 8.3 Å are filled by the $[Cu^+Cl_2]^-$ anions and Cl^- ions.

