Omariniite, Cu$_8$Fe$_2$ZnGe$_2$S$_{12}$, the germanium analogue of stannoidite, a new mineral species from Capillitas, Argentina

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ABSTRACT

Omariniite, ideally Cu$_8$Fe$_2$ZnGe$_2$S$_{12}$, represents the Ge-analogue of stannoidite and was found in bornite-chalcocite-rich ores near the La Rosario vein of the Capillitas epithermal deposit, Catamarca Province, Argentina. The mineral is associated closely with three other Ge-bearing minerals (putzite, catamarcaite, rarely zincobriartite) and bornite, chalcocite, digenite, covellite, sphalerite, tennantite, luzonite, wittichenite, thalcusite and traces of mawsonite. The width of the seams rarely exceeds 60 µm, their length can attain several 100 µm. The mineral is opaque, orange-brown in polished section, has a metallic lustre and a brownish-black streak. It is brittle, and the fracture is irregular to subconchoidal. Neither cleavage nor parting are observable in the sections. In plane-polarized light omariniite is brownish-orange and has a weak pleochroism. Internal reflections are absent. The mineral is distinctly anisotropic with rotation tints varying between brownish-orange and greenish-brown. The average result of 45 electron-microprobe analyses is Cu 42.18(34), Fe 9.37(26), Zn 5.17(43), In 0.20(6), Ge 11.62(22), S 31.80(20), total 100.34(46) wt.%. The empirical formula, based on Σ(Me + S) = 25, is Cu$_{8.04}$(Fe$_{2.03}$In$_{0.02}$)$Σ^{2.05}Zn_{0.96}$Ge$_{1.94}$S$_{12.01}$, ideally Cu$^{+8}$Fe$^{3+2}$Zn$^{2+}$Ge$_{4}^{+2}$S$_{2}$\textsubscript{12}. Omariniite is orthorhombic, space group I$\overline{2}$22, with unit-cell parameters: $a = 10.774(1)$, $b = 5.3921(5)$, $c = 16.085(2)$ Å, $V = 934.5(2)$ Å$^3$, $a:b:c = 1.9981:1.2:9.831$, $Z = 2$. X-ray single-crystal studies ($R_1 = 0.023$) revealed the structure to be a sphalerite derivative identical to that of stannoidite. Omariniite is named after Dr. Ricardo Héctor Omarini (1946–2015), Professor at the University of Salta, for his numerous contributions to the geology of Argentina.

KEYWORDS: omariniite, new mineral species, copper-iron-zinc-germanium-sulfide, electron-microprobe analyses, reflectance data, crystal structure, La Rosario vein, Capillitas district, Catamarca Province, Argentina.

Introduction

OMARINIITE is one of the six germanium-bearing sulfides occurring at Capillitas; among them, putzite, (Cu$_{4.7}$Ag$_{3.3}$)$_{25}$GeS$_6$ (Paar et al., 2004), catamarcaite, Cu$_5$GeW$_8$S$_{25}$ (Putz et al., 2006) and omariniite, previously named “Ge-stannoidite” (Paar et al., 2004; Putz, 2005; Putz et al., 2006), have here their type-locality. Two other species, spryite (Bindi et al., 2016) and zincobriartite (McDonald et al., 2016) were first recognized at Capillitas and described as “As–argyrodite” and Zn-dominant briartite (Putz et al., 2002; Putz, 2005). The full characterization and description as new mineral species were later given by McDonald et al. (2016) and Bindi et al. (2016) who described zincobriartite and spryite, respectively. In addition, the Capillitas epithermal deposit is the type locality for the minerals ishiharaite, ideally (Cu,Ga,Fe,In, Zn)S (Márquez-Zavalía et al., 2015) and lislkirchnerite, ideally Pb$_6$Al(OH)$_8$Cl$_2$(NO$_3$)$_5$·2H$_2$O (IMA No. 2015-064, Effenberger et al., 2015).

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Two further Ge-bearing sulfides – germanite and renierite – are said to occur at Capillitas. They were first reported by Márquez-Zavalía (1988) on the basis of ore microscopic investigations, but have never been analysed by electron microprobe or X-ray techniques. Most probably, they have been misidentified with nekrasovite (“germanite”) and vincieiennite (“renierite”). Both nekrasovite and vincieiennite have been observed frequently in high-sulfidation stage ore from several veins of the Capillitas deposit (Putz, 2005). Thus, the occurrence of germanite and renierite remains uncertain.

According to Höll et al. (2007) Cu-Fe-Zn-Ge-sulfides like germanite and renierite are characteristic for the carbonate-hosted Kipushi-type polymetallic deposits (e.g. Tsumeb, Namibia, and Kipushi, Democratic Republic of Congo) but also occur within high-sulfidation epithermal Cu-Au deposits (e.g. Chelopech, Bulgaria). On the contrary, argyrodite is typically observed in bonanza-grade silver mineralization of the Bolivian “Ag-Sn-belt” (e.g. Porco and Colquechaca Ag-Zn-Pb-Sn deposits, Bolivia; Paar and Putz, 2005) and in vein-type Ag-Pb-Zn (-Cu) deposits (e.g. Freiberg district, Saxony, Germany; Höll et al., 2007). The mineralogy and distribution of germanium at Capillitas is thus unique on a worldwide scale and represents a new mode of the occurrence of this rare metal in epithermal systems. In addition to the six mentioned Ge minerals, Ge contents up to 0.2 wt.% have been found in colusite–nekrasovite (Putz, 2005). This germanium anomaly was discovered during the PhD project of one of the authors (HP), which included a detailed study of the complex mineralogy at Capillitas.

Omariniite is named in honour of Dr. Ricardo Héctor Omarini (1946–2015), Professor at the University of Salta, for his outstanding contributions to the geology of Argentina, especially to the Precambrian basement of the “Formación Puncoviscana”.

The mineral and the mineral name have been approved by the IMA CNMNC (2016-050). Holotype material is housed within the Systematic Reference Series of the National Mineral Collection of Canada, Geological Survey of Canada, Ottawa, Ontario, under catalogue number NMCC 68096. Cotype material is deposited within the collections of the University of Firenze (Italy), the Natural History Museum of London (UK) and in the private collections of two of the authors (HP and WHP).

Location, geology and mineralization

The Capillitas mining district is part of the Farallón Negro Volcanic Complex, which is located in the Province of Catamarca, Argentina. It consists of Miocene extrusive and intrusive subvolcanic rocks within the crystalline basement (metapelites and schists of the Suncho Formation and Capillitas granitic batholith). Porphyry copper-gold deposits (e.g. Bajo de La Alumbrera, Agua Rica) and epithermal vein-type deposits (e.g. Capillitas, Farallón Negro – Alto de la Blenda) are associated with the intrusive rocks, which range from andesitic to dacitic to rhyolitic in composition (Sasso, 1997; Sasso and Clark, 1998).

The Capillitas diatreme represents one of the volcanic centres within the granitic basement block of the Sierra de Capillitas. It is composed of intrusive and volcanioclastic rocks (ignimbrite, rhyolite porphyry, dacite porphyry and tuffs) and is host to the epithermal vein-type Capillitas deposit, which is known mainly for the occurrence of gem-quality rhodochrosite. The epithermal veins are hosted in rhyolite, ignimbrite and granite and show different types of host-rock alteration and both high-sulfidation and intermediate-sulfidation stage mineralization. The polymetallic character of the veins is reflected by its very complex mineralogy with the participation of Cu, Pb, Zn, Fe, As and Sb, associated with W, Bi, Sn, Te, Ag and Au as well as Ge, Cd, In, V, Ni and Tl. Further details on ore deposit geology, alteration and mineralization styles and ore mineralogy are presented elsewhere (e.g. Márquez-Zavalía, 1988, 1999; Paar et al., 2004, Putz, 2005, Putz et al., 2006, Putz et al., 2009).

Cementation processes led to the formation of high-grade copper ores (borrite-chalcocite-digenite) which are restricted to the central part of the deposit in the vicinity of the La Rosario vein. As the old mine workings (“Mina La Rosario” and “Pique Rosario”) dating back to the 19th century are completely collapsed, material of this ore type is restricted to the old dumps nearby. Interestingly, only a few samples of this type of ore contain the unique assemblage of Ge minerals, such as putzite, catamarcaite, zincobriartite and omariniite (putzite-catamarcaite-omariniite-zincobriartite-paragenesis).

Intermediate-sulfidation stage veins hosted within the granitic basement (e.g. the La Argentina vein) are rich in galena, sphalerite and abundant rhodochrosite as the dominating gangue mineral. At the Santa Rita mine, where rhodochrosite-bearing ore of the La Argentina
vein is currently mined for ornamental purposes, small-sized bonanza-grade ore bodies carry a Ag-rich assemblage composed of proustite, pearceite, acanthite, native silver and argyrodite. The latter is locally intergrown with spryite, $\text{Ag}_8(\text{As}_{0.5}^\text{III} \text{As}_{0.5}^\text{V})\text{S}_6$ (Putz et al., 2002; Putz, 2005; Bindi et al., 2016).

The Ge-bearing minerals at Capillitas occur in two genetically different environments. The ore of the intermediate sulfidation stage was dictated by a high activity of silver and arsenic, which led to the formation of argyrodite-spryite. The increased activity of copper within the ores created by supergene enrichment processes, however, led to the formation of copper-dominated phases, amongst them putzite. Further details dealing with the crystallization sequence of the omariniite-bearing ore and the possible conditions of formation are presented in Paar et al. (2004) and Putz et al. (2006).

**Appearance and physical properties**

Omariniite is fairly abundant in those bornite-chalcocite ores which also carry putzite and/or catamarcaite. Within the putzite holotype material, the new mineral frequently, but not exclusively, occurs as: (1) a rim or seam at the contact between putzite and chalcocite, rarely exceeding a width of 60 µm but attaining a length of several hundreds of micrometres (Fig. 1a); and (2) as an envelope

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**Fig. 1. Different associations of omariniite.** (a) Putzite (pu) rimmed by omariniite (om) in association with chalcocite (cc) containing relics of bornite (bn); late-stage sphalerite (sp) in narrow veinlets is crosscutting chalcocite and the Ge sulfides (sample PR/C2). Air, uncrossed polars. (b) Omariniite (om) is seaming catamarcaite (cat) which locally shows subhedral to euhedral crystal forms. These composite grains represent inclusions within putzite (pu) which is strongly replaced by chalcocite (cc) and late-stage sphalerite (sp); zincobriartite (zbt) occurs in traces (sample PR/C3P). Air, uncrossed polars. (c) Rim of omariniite (om; with weak pleochroism) in association with putzite (pu), catamarcaite (cat), chalcocite (cc), bornite (bn) and sphalerite (sp); (sample PR/C2). Air, uncrossed polars. (d) Same field of view as in panel (c) omariniite (om) shows its characteristic anisotropy with orange-brown to greenish grey rotation-tints (sample PR/C2). Air, crossed polars.
around the anhedral to subhedral inclusions of catamarcaite within putzite and chalcocite (Fig. 1b), where it is sometimes also associated with zincobriartite. The rims are usually composed of a mosaic of intensely twinned, anhedral subgrains (Figs. 1c and 1d); individual grains without twinning are rare and measure up to 30 µm × 70 µm. An exceptional single grain measures 200 µm × 100 µm. Wittichenite, tennantite, thalcusite and rare grains of mawsonite are also observed within this ore type but never in contact with omariniite.

The holotype material of catamarcaite also contains omariniite. Within this material, the new mineral occurs as (3) an envelope (generally <10 µm in thickness) around catamarcaite inclusions within chalcocite/digenite/covellite/sphalerite in association with luzonite (see fig. 3a in Putz et al., 2006).

### Table 1. Reflectance data and colour values for omariniite

| λ (nm) | Omariniite | Stannoidite |
|--------|------------|-------------|
|        | R1        | R2          | R1   | R2   |
| 400    | 22.0      | 21.9        | 17.5 | 18.9 |
| 420    | 20.9      | 21.5        | 17.4 | 19.2 |
| 440    | 20.2      | 21.0        | 18.2 | 20.3 |
| 460    | 19.7      | 21.0        | 19.5 | 21.7 |
| 480    | 19.5      | 21.0        | 20.8 | 23.0 |
| 500    | 19.4      | 21.1        | 21.8 | 24.2 |
| 520    | 19.75     | 21.0        | 22.8 | 25.3 |
| 540    | 20.4      | 21.8        | 23.7 | 26.3 |
| 560    | 21.2      | 22.2        | 24.6 | 27.1 |
| 580    | 22.1      | 22.8        | 25.4 | 27.8 |
| 600    | 23.2      | 23.5        | 26.2 | 28.4 |
| 620    | 24.2      | 24.2        | 27.1 | 29.1 |
| 640    | 25.3      | 25.0        | 27.8 | 29.5 |
| 660    | 26.2      | 25.7        | 28.7 | 30.0 |
| 680    | 27.2      | 26.6        | 29.6 | 30.5 |
| 700    | 28.2      | 27.4        | 30.3 | 30.8 |

| C illumin | x   | 0.328 | 0.322 | 0.339 | 0.335 |
|           | y   | 0.320 | 0.321 | 0.343 | 0.342 |
|           | Y%  | 21.4  | 22.4  | 24.4  | 26.8  |
|           | Lambda d | 597 | 590 | 578 | 577 |
|           | Pe%  | 5.9  | 4.7  | 14.8  | 13.8  |

| A          | x   | 0.467 | 0.461 | 0.471 | 0.468 |
|            | y   | 0.403 | 0.406 | 0.414 | 0.415 |
|            | Y%  | 22.0  | 22.8  | 25.1  | 27.4  |
|            | Lambda d | 616 | 616 | 587 | 586 |
|            | Pe%  | 8.0  | 5.3  | 20.8  | 18.9  |

**Optical properties**

In polished sections using plane-polarized light the colour of omariniite is orange-brown which is enhanced in oil. Pleochroism is weak, as also is the bireflectance. Between crossed polars omariniite is distinctly anisotropic, with rotation tints changing from brownish-orange to greenish-brown. No internal reflections were observed.

Reflectance measurements were made using a J&M TIDAS diode array spectrometre between 400 and 700 nm at intervals of 0.8135 nm. The data

![Fig. 2. Reflectance spectra, in air, for omariniite and stannoidite.](image-url)
were reduced to 20 nm intervals and are summarized in Table 1. A comparison of omariniite with stannoidite (Fig. 2) shows characteristic differences between the two species which allows them to be easily distinguished using optical methods. Both species show an increase of the reflectance values with increasing wavelength. The maxima for these minerals are towards the red end of the visible spectrum with dominant wavelengths ($\lambda_\alpha$) relative to the C illuminant of 590–597 nm (omariniite) and 577–578 nm (stannoidite) which explains the orange-brown colours of the species in reflected light. However, the reflectance values for omariniite are generally markedly lower than those observed for stannoidite.

Chemical composition

Omariniite was analysed with a JEOL Superprobe JXA-8600 (controlled by an LINK-eXL system, WDS mode, operated at 25 kV and 35 nA, with 20 s and 10 s as counting times for peak and background, respectively), installed at the University of Salzburg, Austria. The following standards and X-ray lines were used: chalcopyrite (CuK$\alpha$, FeK$\alpha$), synthetic ZnS (ZnK$\alpha$, SK$\alpha$), Ge metal (GeK$\alpha$) and synthetic InAs (InL$\alpha$). Tin was sought but not detected. The raw data were processed with the on-line ZAF-4 program.

Several aggregates of omariniite in various polished sections of the putzite holotype material were analysed and found to be homogeneous. The results show only minor variation of the chemical composition (Table 2). The average result of 45 analytical points of three grain aggregates is (wt.%): Cu 42.18(34), Fe 9.37(26), Zn 5.17(43), In 0.20(6), total 100.34(46). Using the ideal formula of stannoidite, Cu$_8$Fe$_2$Zn$_2$Sn$_2$S$_{12}$ and $\Sigma$ Me + S = 25, this corresponds to Cu$_{8.04}$Fe$_{2.03}$In$_{0.02}$Zn$_{0.96}$Ge$_{1.94}$S$_{12.01}$. The ideal chemical formula of omariniite, with valence states for the elements in agreement with those reported for stannoidite by Yamanaka and Kato (1976), can be expressed as Cu$_{x}$Fe$_{y}$Zn$_z$Ge$_t$S$_{12}$ which requires Cu 41.83, Fe 9.19, Zn 5.38, Ge 11.95, S 31.65, total 100.00 wt.%

| Element | Cu     | Fe     | Zn     | In     | Ge     | S     | Total |
|---------|--------|--------|--------|--------|--------|-------|-------|
|         | 42.18(34) | 9.37(26) | 5.17(43) | 0.20(6) | 11.62(22) | 31.80(20) | 100.34(46) |

X-ray crystallography and crystal-structure determination

A few omariniite crystals were handpicked from a polished section under a reflected light microscope and examined by means of an Oxford Diffraction Xcalibur 3 CCD single-crystal diffractometer using graphite-monochromatized MoK$\alpha$ radiation (see Table 3 for details). Single-crystal X-ray diffraction intensity data were integrated and corrected for standard Lorentz-polarization factors with the CrysAlis RED package (Oxford Diffraction, 2006). The program ABSPACK in CrysAlis RED (Oxford Diffraction, 2006) was used for the absorption correction. A total of 1238 unique reflections was collected. The statistical tests on the distribution of $|E|$ values ($E^2-1 = 0.697$) indicated the absence of an inversion centre. Systematic abscences were consistent with the space group $I\overline{2}2_1$.

Given the similarity in the unit-cell values and in the space groups, the structure was refined starting from the atomic coordinates reported for stannoidite (Kudoh and Takéuchi, 1976) using the program Shelx-97 (Sheldrick, 2008). The site occupation factor (s.o.f.) at the seven cation sites was allowed to vary (Zn vs. structural vacancy) using scattering curves for neutral atoms taken from the International Tables for Crystallography (Wilson, 1992). After several cycles and a careful check of the bond distances at the different sites, the positions were found fully occupied by the atomic species given in Table 4 and their s.o.f. were fixed to 1. The electron density was only found higher than 26 (pure occupation by Fe) at the M5 site, and all the In$^{3+}$ found by electron microprobe was thought to occur at this site. Table 3 reports details of the selected crystal, data collection and refinement. Final atomic coordinates and equivalent isotropic displacement parameters are given in Table 4, whereas selected bond distances are shown in Table 5. A list of the observed and calculated structure factors and crystallographic information file are deposited with the Principal Editor of Mineralogical Magazine at http://www.minersoc.org/pages/e_journals/dep_mat.html.
Description of the structure and relationships with other species

The crystal structure of omariniite (Fig. 3) can be considered as a sphalerite derivative. The omariniite structural topology is also derivable from the stannite structure by substituting Cu atoms for a set of Ge atoms in stannite and adding excess Cu atoms in a set of tetrahedral vacancies (Fig. 4). A comparison of the bond distances for the different tetrahedra occurring in stannoidite and omariniite is presented in Table 5. It appears evident that the Ge-tetrahedron (i.e., M6) is much smaller (mean bond distance 2.248 Å) than the Sn-homologue in stannoidite (mean bond distance 2.40 Å; Kudoh and Takéuchi, 1976) and it is close to that observed in a.

### Table 3. Crystal and experimental data for omariniite.

| Crystal data            |          |
|-------------------------|----------|
| Crystal size (mm)       | 0.05 × 0.06 × 0.07 |
| Cell setting, space group| Orthorhombic, I222 |
| a (Å)                   | 10.774(1) |
| b (Å)                   | 5.3921(5) |
| c (Å)                   | 16.085(2) |
| V (Å³)                  | 934.5(2) |
| z                       | 2        |

| Data collection and refinement |          |
|-------------------------------|----------|
| Radiation, wavelength (Å)     | MoKα, λ = 0.71073 |
| Temperature (K)               | 293      |
| θ max (°)                     | 59.97    |
| Measured reflections          | 10125    |
| Unique reflections            | 1238     |
| Reflections with Fo > 4σ(Fo)  | 432      |
| R_int                         | 0.0423   |
| Rσ                            | 0.0268   |
| Range of h, k, l              | −13 ≤ h ≤ 13, −7 ≤ k ≤ 7, −22 ≤ l ≤ 22 |
| R [Fo > 4σ(Fo)]               | 0.0226   |
| R (all data)                  | 0.0273   |
| wR (on F²)                    | 0.0462   |
| Goof                          | 0.990    |
| Number of least-squares parameters | 62 |
| Max. and min. resid. peak (e Å⁻³) | 0.43 (at 0.29 Å from S2) |
|                               | −1.82 (at 0.96 Å from M6) |

### Table 4. Wyckoff positions, site occupation factors (s.o.f.), fractional atomic coordinates, and equivalent isotropic displacement parameters (Å²) for the selected omariniite crystal.

| Atom | Wyckoff | s.o.f. | x     | y     | z     | Uiso  |
|------|---------|-------|-------|-------|-------|-------|
| M1   | 2a      | Zn1.00| 0     | 0     | 0     | 0.0238(4) |
| M2   | 2b      | Cu1.00| ½     | 0     | 0     | 0.0208(4) |
| M3   | 2c      | Cu1.00| 0     | 0     | ½     | 0.0238(4) |
| M4   | 4f      | Cu1.00| 0.25821(14) | 0     | ½     | 0.0196(3) |
| M5   | 4i      | Fe0.97(1)In0.03| 0     | 0     | 0.32997(11) | 0.0242(5) |
| M6   | 4j      | Ge1.00| 0     | ½     | 0.17651(9) | 0.0234(3) |
| M7   | 8k      | Cu1.00| 0.24616(11) | 0.0105(2) | 0.16974(6) | 0.0245(2) |
| S1   | 8k      | S1.00| 0.1218(3) | 0.2463(7) | 0.09836(18) | 0.0368(7) |
| S2   | 8k      | S1.00| 0.3797(4) | 0.7553(8) | 0.08501(18) | 0.0413(9) |
| S3   | 8k      | S1.00| 0.1293(3) | 0.7482(8) | 0.2446(3) | 0.0415(7) |
TABLE 5. Comparison of the bond distances (Å) in the
crystal structure of stannoidite (1; Kudoh and
Takéuchi 1976) and omariniite (2; this study).

|        | 1        | 2        |
|--------|----------|----------|
| M1–S1 (×4) | 2.346    | 2.447(3) |
| M2–S2 (×4) | 2.259    | 2.258(4) |
| M3–S2 (×4) | 2.297    | 2.292(4) |
| M4–S1 (×2) | 2.303    | 2.459(4) |
| M4–S2 (×2) | 2.356    | 2.404(4) |
| M5–S2 (×2) | 2.391    | 2.377(4) |
| M5–S3 (×2) | 2.293    | 2.381(4) |
| M6–S1 (×2) | 2.419    | 2.275(4) |
| M6–S3 (×2) | 2.376    | 2.220(3) |
| M7–S1     | 2.261    | 2.175(4) |
| M7–S2     | 2.454    | 2.454(4) |
| M7–S3     | 2.341    | 2.244(4) |
| M7–S3     | 2.223    | 2.311(4) |

Due to the iso-electronic nature of its constituent elements (Fe = 26, Cu = 29, Zn = 30, Ge = 32) together with the ambiguity in their valence states, the metal partitioning in omariniite is, however, not straightforward. According to Brese and O’Keeffe (1991), the ideal Me–S distance (in Å) in a regular tetrahedron decreases following the sequence: 2.466/In³⁺, 2.450/Sn⁴⁺, 2.416/Fe²⁺, 2.370/Cu⁺, 2.346/Zn²⁺, 2.266/Fe³⁺, 2.220/Ge⁴⁺, 2.116/Cu²⁺. Consequently, Ge⁴⁺ should be hosted within the smallest tetrahedron (i.e. M6), whereas Fe³⁺ should be hosted at the other small tetrahedral sites. Taking into account this approach for both stannoidite and omariniite structures, a bond valence balance give the following values for the M sites (values for stannoidite and omariniite, respectively, are given in parentheses in valence units – vu): M1 (2.04, 1.52), M2 (1.36, 1.36), M3 (1.22, 1.23), M4 (1.13, 0.86), M5 (2.48, 2.26), M6 (4.59, 3.72), M7 (1.19, 1.28). Recent studies demonstrated the presence of Cu⁺ and Fe³⁺ in chalcopyrite (e.g. Goh et al., 2006) and the corresponding bond-valence sums (BVS) at Cu and Fe sites are 1.18 and 3.15 vu, respectively, with corresponding bond distances of 2.312 and 2.248 Å (Knight et al., 2011). Consequently, the bond-valence sums at Cu sites (M2, M3, M4, M7) in both the stannoidite and omariniite structures probably agree with the presence of Cu⁺. Such an oversaturation at Cu tetrahedral sites have been observed in other sulfides and sulfosalts and it could probably be related to the non-accuracy of the bond parameter for Cu–S bond given by Brese and O’Keeffe (1991) and to the covalent nature of the bonds. On the contrary, the observed strong undersaturation at M1 (in omariniite) and M5 (both stannoidite and omariniite) have to be taken into account. They could be either related to some limits in the crystal structure refinement or to a different
metal partitioning. A possible hypothetical new partition could be Cu⁺ at M1 (BVS 0.82), Fe³⁺ at M2 (BVS 3.07), Fe³⁺ at M3 (BVS 2.76), Cu⁺ at M4 (BVS 0.86), (Cu⁺₅Zn²⁺₅) at M5 (BVS 1.41), Ge⁴⁺ at M6 (BVS 3.72) and Cu⁺ at M7 (BVS 1.28), giving rise to the formula Cu⁺₈Fe³⁺₂ZnGe⁴⁺₂S₁₂ (Z = 2).

Calculated X-ray powder diffraction data (d in Å) for omariniite are given in Table 6. Intensities and dhkl values were calculated using the Powdercell 2.3 software (Kraus and Nolze, 1996) on the basis of the structural model given in Table 5; only reflections with Icalc > 1 are listed. The strongest reflections are given in bold.

| Irel | dcalc (Å) | h k l |
|------|-----------|------|
| 2    | 8.9515    | 1 0 1|
| 4    | 4.8001    | 1 0 3|
| 1    | 4.4757    | 2 0 2|
| 2    | 4.1356    | 1 1 2|
| 100  | 3.1063    | 2 1 3|
| 2    | 2.8018    | 3 1 2|
| 6    | 2.6960    | 0 2 0|
| 6    | 2.6935    | 4 0 0|
| 6    | 2.6808    | 0 0 6|
| 1    | 2.4583    | 2 1 5|
| 24   | 1.9055    | 4 2 0|
| 23   | 1.9010    | 0 2 6|
| 23   | 1.9001    | 4 0 6|
| 12   | 1.6248    | 2 3 3|
| 12   | 1.6237    | 6 1 3|
| 11   | 1.6181    | 2 1 9|
| 4    | 1.5531    | 4 2 6|
| 3    | 1.3480    | 0 4 0|
| 3    | 1.3468    | 8 0 0|
| 3    | 1.3440    | 0 0 12|
| 4    | 1.2361    | 6 3 3|
| 4    | 1.2336    | 2 3 9|
| 4    | 1.2332    | 6 1 9|
| 6    | 1.0994    | 4 4 6|
| 6    | 1.0989    | 8 2 6|
| 5    | 1.0963    | 4 2 12|
| 2    | 1.0375    | 2 5 3|
| 2    | 1.0366    | 1 0 13|
| 2    | 1.0354    | 6 3 9|
| 2    | 1.0322    | 2 1 15|

*Intensities and dhkl values were calculated using Powdercell 2.3 software (Kraus and Nolze, 1996) on the basis of the structural model given in Table 5; only reflections with Icalc > 1 are listed. The strongest reflections are given in bold.

Tsumeb and Khusib Springs carry significant Ge (0.5–0.7 wt.% in Khusib Springs and up to 2.2 wt.% in Tsumeb, respectively; Melcher, 2003).

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