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Lead-antimony sulfosalts from Tuscany (Italy). XXII. Marcobaldiite, ~Pb12(Sb3As2Bi)Σ6S21, a new member of the jordanite homologous series from the Pollone mine, Valdicastello Carducci --Manuscript Draft--

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DIPARTIMENTO DI SCIENZE DELLA TERRA



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Manuscript

Dear Sir,

we would like to publish our paper in European Journal of Mineralogy.

The paper, entitled "Lead-antimony sulfosalts from Tuscany (Italy). XXII. Marcobaldiite, ~Pb₁₂(Sb₃As₂Bi)₂₆S₂₁, a new member of the jordanite homologous series from the Pollone mine, Valdicastello Carducci" has not been previously published and it will not be submitted elsewhere for publication while it is in review for *European Journal of Mineralogy*.

Marcobaldiite is a new N = 3.5 homologue in the jordanite homologous series and shows a new kind of crystal structure. The approval of marcobaldiite as a valid mineral species has been voted by the IMA CNMNC (IMA 2015-109).

Yours faithfully,

Cristian Biagioni

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- 3 Carducci
- 4

5 **Running title**: Marcobaldiite, a new jordanite homologue

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Lead-antimony sulfosalts from Tuscany (Italy). XXII. Marcobaldiite, ~ $Pb_{12}(Sb_3As_2Bi)_{\Sigma 6}S_{21}$, a new member of the jordanite homologous series from the Pollone mine, Valdicastello Carducci

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Abstract: The new mineral species marcobaldiite, ~ $Pb_{12}(Sb_3As_2Bi)_{\Sigma 6}S_{21}$, has been discovered in a 47 single specimen collected in a quartz vein embedded in tourmaline-bearing schists from the Pollone 48 mine, Valdicastello Carducci, Pietrasanta, Apuan Alps, Tuscany, Italy. It occurs as a black blocky 49 prismatic crystal, up to 1 cm in size, with a metallic luster, associated with Sb-rich tennantite. Under 50 the ore microscope, marcobaldiite is white, with a distinct anisotropism, with grey – bluish grey 51 rotation tints. Polysynthetic lamellar twinning is common and characteristic. Reflectance 52 percentages for the four COM wavelengths are $[R_{\min}, R_{\max}(\%), (\lambda)]$: 31.6, 40.1 (470 nm), 30.9, 39.6 53 (546 nm), 30.4, 38.5 (589 nm), and 30.0, 37.6 (650 nm). Electron microprobe analyses gave 54 (average of 10 spot analyses - in wt%): Pb 64.05(34), As 4.51(8), Sb 9.10(14), Bi 4.24(6), S 55 17.24(19), total 99.14(53). On the basis of $\Sigma Me = 20$ apfu, the empirical formula is 56 Pb_{11,98(4)}Sb_{2,90(3)}As_{2,33(3)}Bi_{0,79(1)}S_{20,80(20)}. The main diffraction lines, corresponding to multiple *hkl* 57 indices, are [d in Å (relative visual intensity)]: 3.568 (ms), 3.202 (ms), 3.016 (ms), 2.885 (ms), 58 59 2.233 (vs), 2.125 (s), 1.848 (s), and 1.775 (s). The crystal structure study gives a triclinic unit cell, space group $P\bar{1}$, with a = 8.9248(9), b = 29.414(3), c = 8.5301(8) Å, $\alpha = 98.336(5)$, $\beta = 118.175(5)$, 60 $\gamma = 90.856(5)^\circ$, V = 1944.1(3) Å³, Z = 2. The crystal structure has been solved and refined to $R_1 =$ 61 62 0.067 on the basis of 6193 reflections with $F_0 > 4\sigma(F_0)$ and 363 refined parameters. Marcobaldiite is a new N = 3.5 homologue of the jordanite homologous series, characterized by the 1:1 alternation of 63 64 two kinds of layers of distorted octahedra, three- and four-octahedra thick, i.e. of the kirkiite and 65 jordanite types, respectively. Layers are connected by one atomic layer of three bicapped trigonal prismatic Pb atoms and one triangular pyramidal As atom in split position. The name of this new 66 mineral species honours the mineral amateur Marco Baldi (b. 1944) for his contribution to the 67 knowledge of the mineralogy of the pyrite \pm baryte \pm iron oxide ore deposits from southern Apuan 68 Alps. 69

Key-words: marcobaldiite; new mineral; sulfosalt; crystal structure; lead; antimony; arsenic;
bismuth; Pollone mine; Apuan Alps.

73 **1. Introduction**

The jordanite homologous series, first defined by Makovicky (1989), is a group of lead sulfosalts having general formula $Pb_{4N-2}Me^{3+}_{6}S_{4N+7}$, where $Me^{3+} = As$, Sb, Bi (Makovicky *et al.*, 2006). Three mineral species belonging to this series are reported in Moëlo *et al.* (2008), *i.e.* jordanite, geocronite, and kirkiite. The possibly related mineral tsugaruite, ideally Pb₄As₂S₇ (Shimizu *et al.*, 1998), has a different crystal structure (Matsushita *et al.*, 2014), even if its details have not been published yet.

Jordanite homologues are characterized by two short axes (~8.5 and ~9.0 Å) and a longer one, 80 whose length depends on the homologue order, ranging from ≈ 26 Å (N = 3 - kirkiite) to ≈ 32 Å (N81 = 4 – jordanite and geocronite). The N = 4 isotypic pair jordanite (Pb₁₄As₆S₂₃) – geocronite 82 (Pb14Sb6S23) is well-known from various localities world-wide, and, considering also synthetic 83 products (Jambor, 1969), displays a complete solid-solution. On the contrary, kirkiite, ideally 84 85 Pb₁₀As₃Bi₃S₁₉, is exceedingly rare, having been described only in two localities, the Greek type locality (the Pb-Zn ore deposit of Aghios Philippos, near Kirki, Thrace; Moëlo et al., 1985), and the 86 high-T fumaroles of La Fossa crater, Vulcano, Aeolian Islands, Italy (Borodaev et al., 1998; Pinto et 87 al., 2006). 88

89 In order to fully characterize the occurrence of geocronite from the Pollone mine, near the small hamlet of Valdicastello Carducci, Apuan Alps, Italy, where this mineral has been known 90 since the first description given by Kerndt (1845), Biagioni et al. (2016a) examined both historical 91 92 and new samples from this locality, studying them through single-crystal X-ray diffraction and electron-microprobe analysis. During this investigation, a mineral was found having the two short 93 unit-cell parameters close to those of jordanite homologues and the longer one intermediate 94 between those of the N = 3 and N = 4 members, *i.e.* ≈ 29 Å. The crystallographic and chemical 95 studies allowed its identification as a new homologue belonging to the jordanite series, *i.e.* 96 97 marcobaldiite, with N = 3.5 homologue number. The new mineral and its name have been approved by the CNMNC-IMA (2015-109). The holotype material is deposited in the mineralogical 98 99 collections of the Museo di Storia Naturale, Università di Pisa, Via Roma 79, Calci, Pisa, Italy, under catalogue number 19709. The name is in honour of the mineral collector Marco Baldi (b. 100 101 1944), for his contribution to the knowledge of the mineralogy of the pyrite \pm baryte \pm iron oxide 102 ore deposits from southern Apuan Alps. Marco Baldi was the first to describe the actual occurrence of geocronite crystals from Valdicastello Carducci (Baldi, 1982). His contribution favored the 103 mineralogical rediscovery of the Pollone mine, which since then has been explored by mineral 104 collectors and mineralogists, leading to the discovery of several rare and new mineral species, 105

mainly represented by lead sulfosalts (sterryite, parasterryite, carducciite, meerschautite, and
polloneite – Moëlo *et al.*, 2011; Biagioni *et al.*, 2014; Biagioni *et al.*, 2016b; Topa *et al.*, 2017).

108 The aim of this paper is the full description of the new mineral species marcobaldiite and the 109 discussion of its crystal-chemistry in the framework of the jordanite homologous series.

110

111 **2. Geological setting**

The baryte + pyrite ± (Pb-Zn-Ag) Pollone ore deposit (latitude 43°57'47"'N, longitude 112 10°16'19"E) is located near the small hamlet of Valdicastello Carducci, Pietrasanta, Tuscany, Italy. 113 The ore bodies are hosted within a Paleozoic metavolcanic-metasedimentary sequence, 114 115 metamorphosed up to greenschist facies conditions during the Alpine orogeny. Estimates of P-T 116 conditions indicate metamorphic temperatures up to 350° C with a pressure of 0.35 GPa. Higher T values (450°C) were recorded for mineralizing fluids (Costagliola et al., 1998). Biagioni et al. 117 118 (2016a) described the main geological and structural features of the Pollone mine and distinguished three kinds of occurrence: i) lenses of microcrystalline baryte + pyrite, with the local presence of 119 120 interlayers/lenses of galena + sphalerite, folded and partially re-worked during the late tectonic stage evolution of Apuan Alps metamorphic complex (e.g., Carmignani & Kligfield, 1990); ii) 121 122 quartz lensoid veins, oriented SW-NE, with coarse-grained masses of galena, sphalerite, and sulfosalts; and *iii*) extension quartz \pm baryte \pm sulfides veins, trending N140-170, embedded both in 123 124 the country rocks and in the ore bodies.

125 Twenty-four different sulfosalts have been identified in the Pollone ore deposit (Table 1). The typical occurrence is represented by the microcrystalline baryte + pyrite ore bodies, where sulfosalts 126 are scattered as interstitial grains or as euhedral prismatic to acicular individuals. Rarely, they are 127 related to small quartz + baryte veinlets and small fissures occurring in these ore bodies. This kind 128 of occurrence is typical for the lead-silver sulfosalts typical of the Pollone mine. The quartz \pm 129 baryte ± sulfides extension veins represent another source of sulfosalts. Actually, the well-130 crystallized specimens of jordanite and geocronite were found in the vugs of these veins. Finally, 131 some Ag sulfosalts have been identified in the quartz lensoid veins oriented SW-NE (i.e., diaphorite 132 and pyrargyrite – Frizzo & Simone, 1995). 133

Marcobaldiite was found in a quartz extension vein embedded in tourmaline-rich schist, occurring in the Stanzone tunnel, one of the stopes of the Pollone mine. Its crystallization is related to the circulation of hydrothermal fluids during the Tertiary Alpine tectono-metamorphic events, favoring the remobilization of elements from the country rocks and the ore bodies and their crystallization in the extension veins (Biagioni *et al.*, 2016a).

140 **3. Occurrence and mineral description**

Marcobaldiite was identified in only one sample collected in quartz extension veins; in the same vein Sb-rich tennantite was identified. It occurs as a black prismatic crystal measuring $10 \times 3 \times 3$ mm. The presence of striations suggests the occurrence of a polysynthetic twinning, confirmed by reflected light microscopy (see below). It is brittle, with an irregular fracture. The streak is black, the luster is metallic.

Micro-indentation measurements carried out with a VHN load of 15 g give a mean value of 147 182 kg mm⁻² (range: 170 – 195 kg mm⁻²), corresponding to a Mohs hardness of ~ $3-3\frac{1}{2}$. Density 148 was not measured; calculated density is 6.56 g cm⁻³, on the basis of the empirical formula.

In plane-polarized incident light, marcobaldiite is white in color, distinctly bireflectant. Between crossed polars, it is anisotropic, with grey to bluish-grey rotation tints. Internal reflections were not observed. Polysynthetic twinning is common and characteristic (Fig. 1). The reflectance values of marcobaldiite were determined in air with WTiC as standard (Table 2).

153

154 **3.1. Chemical analysis**

A grain of marcobaldiite was analyzed with a Superprobe JEOL JXA 8200 electron microprobe (Eugen F. Stumpfl laboratory, Leoben University, Austria). The operating conditions (WDS mode) were: accelerating voltage 20 kV, beam current 10 nA, beam size 1 μ m. Standards (element, emission line) were: galena (Pb *M* α), stibnite (Sb *L* α), GaAs (As *L* α), pyrite (S *K* α), and Bi₂Te₃ (Bi *M* α). Counting times were 30 s on the peak and 15 s on the right and left backgrounds. The studied grain proved to be very homogeneous. Chemical data are given in Table 3.

161 On the basis of $\Sigma Me = 18$ atoms per formula unit (*apfu*), the empirical formula of 162 marcobaldiite is Pb_{11.98}Sb_{2.90}As_{2.33}Bi_{0.79}S_{20.80}. The idealized stoichiometric chemical formula of the 163 studied sample is Pb₁₂(Sb₃As₂Bi)_{$\Sigma 6$}S₂₁, corresponding to (in wt%) Pb 64.02, Sb 9.40, As 3.86, Bi 164 5.38, S17.34, sum 100.00.

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166 **3.2. Crystallography**

167 X-ray powder diffraction (XRPD) pattern of marcobaldiite was collected using a 114.6 mm
168 Gandolfi camera with Ni-filtered Cu *K*α radiation. The observed pattern, compared with the
169 calculated one (obtained using the software PowderCell; Kraus & Nolze, 1996) is reported in Table
170 4. Owing to the multiplicity of indices for the majority of the observed diffraction lines, the unit-cell
171 parameters were not refined from X-ray powder diffraction data.

172 Single-crystal X-ray intensity data collection was performed using a Bruker Smart Breeze 173 diffractometer (50 kV, 30 mA) equipped with a CCD 4k low noise area detector. Graphite

monochromatized Mo Ka radiation was used. The detector-to-crystal working distance was 50 mm. 174 6008 frames were collected in ω scan modes, in 0.25° slices. Exposure time was 25 s per frame. The 175 data were integrated and corrected for Lorentz-polarization, background effects, and absorption, 176 using the package of software Apex2 (Bruker AXS Inc., 2004), resulting in a set of 6794 177 independent reflections. The refined unit-cell parameters are a = 8.9248(9), b = 29.414(3), c =178 8.5301(8) Å, $\alpha = 98.336(5)$, $\beta = 118.175(5)$, $\gamma = 90.856(5)^{\circ}$, V = 1944.1(3) Å³. The statistical tests 179 on |E| values ($|E^2 - 1| = 0.993$) suggested the possible centrosymmetricity of marcobaldiite. 180 Consequently, the crystal structure was solved in the space group $P\overline{1}$ using Shelxs-97 (Sheldrick, 181 2008). Scattering curves for neutral atoms were taken from the International Tables for 182 183 Crystallography (Wilson, 1992).

In a first step, the positions of the heavier atoms (Pb, Bi, Sb) were found; through successive 184 difference-Fourier maps, the other atom positions were located. In the crystal structure of 185 marcobaldiite, there are eighteen independent cation positions (one of them split) and twenty-one S 186 sites. Among cation sites, twelve Pb positions and six (Sb/As) sites were identified on the basis of 187 site scattering and Me-S distances. The site occupancy factors (s.o.f.) of the Pb positions were 188 initially refined using the scattering curve of Pb $vs \square$ and they were found to be fully occupied. The 189 190 six (Sb/As) positions were refined using the scattering curves of Sb vs As. Whereas four of them are mixed (Sb/As) positions, with variable As and Sb contents, Sb4 was found to be completely 191 occupied by Sb only, whereas an atom heavier than Sb partially occupies the Pb9 site. 192 Consequently, the s.o.f. of this site was modeled using the curve of Pb and Sb, giving a s.o.f. 193 Pb_{0.61}Sb_{0.39}. After several cycles of isotropic refinement using *Shelxl*-2014 (Sheldrick, 2015), the R_1 194 converged to 0.087. By refining the anisotropic displacement parameters for cations only, the 195 196 refinement yielded a R_1 value of 0.071.

Chemical analysis showed the presence of 0.79 Bi apfu. Owing to the similar scattering 197 factors of Pb (Z = 82) and Bi (Z = 83), it is not possible to distinguish these two atoms on the basis 198 of the site scattering. On the contrary, their bond valence sums (BVS), calculated using the bond 199 200 parameters of Brese & O'Keeffe (1991), were taken into account. An examination of the BVS at the different Pb and (Pb/Sb) sites suggests that Bi is not preferentially partitioned at any specific 201 position, but it would be be statistically distributed among some Pb positions. Bismuth was located 202 in the four Pb sites having the highest BVS values, *i.e.* Pb1, Pb8, Pb15, and Pb16, with a Bi content 203 not over-passing 0.15 apfu in each position. The BVS at the Pb9 position agrees with a mixed 204 205 (Pb,Sb) occupancy, and not with a mixed (Bi,Sb) or (Pb,Sb,Bi) occupancy.

The final anisotropic model for all the atom positions gave $R_1 = 0.067$ for 6193 reflections with $F_0 > 4\sigma(F_0)$ and 363 refined parameters. Further details of data collection and refinement are given in Table 5. Atomic coordinates and equivalent isotropic displacement parameters ofmarcobaldiite are given in Table 6. The unit cell is represented in Fig. 2.

210

211 **4. Crystal structure description**

212 **4.1. General organization**

Marcobaldiite is the N = 3.5 member of the jordanite homologous series. Following the modular description given by Makovicky *et al.* (2006) for the N = 3 homologue kirkiite, the crystal structure of marcobaldiite (Fig. 3) can be described as formed by a stacking sequence of distorted layers of octahedra and trigonal prismatic layers parallel to {010} and sharing S atoms along their boundaries. N = 3 kirkiite-type layers and N = 4 jordanite-type layers, three- and four-octahedra thick respectively, alternate along [010] in a 1:1 ratio, separated by one atomic slab of three bicapped trigonal prismatic (Pb,Bi) atoms and one triangular pyramidal As atom (split position).

Alternatively, the crystal structure can be described as formed by (111) slabs of distorted PbS archetype (*e.g.*, Moëlo *et al.*, 1985; Makovicky, 1989), or (210) slabs of SnS archetype. The broadness of the two kinds of octahedral layer corresponds to three and four subcells of SnS or PbS. Within each layer, lone-electron-pairs of trivalent Sb and As, at the opposite side of shortest (*i.e.* strongest) (Sb,As)–S bonds, are directed towards a common oblique interspace, to form so-called lone-electron-pair micelles (Fig. 3).

226

4.2. Cation coordinations and site occupancies

Among cation positions, there are twelve Pb or mixed (Pb,Bi) sites, one pure Sb position, a mixed (Pb,Sb) site, one pure split As position, and three mixed (Sb/As) sites with different As/(As+Sb) atomic ratios. Cation characteristics (coordination, mean bond distance, and bond valence sum) are summarized in Table 7, while bond valence sums on S atoms are given in Table 8.

Lead atoms are six- to eight-fold coordinated, with average bond distances ranging from 232 2.996 Å for the six-fold coordinated Pb10 site to 3.075 Å for the eight-fold coordinated Pb16 233 position. Pb3, Pb6, Pb10, and Pb11 have a distorted octahedral coordination; actually, the 234 coordination number of the latter site is increased to seven when S21b site is occupied. These Pb 235 236 sites have a pure Pb occupancy, in agreement with their BVS ranging from 1.95 to 1.99 valence unit 237 (v.u.). Pb18 have a six-fold coordination corresponding to a trigonal prismatic polyhedron; two additional very long Pb–S bonds, ranging between 3.83 and 3.91 Å, give rise to a bicapped trigonal 238 prism. The Pb atom hosted at this position is underbonded, with a BVS of 1.75 v.u. A seven-fold 239 coordination is shown by Pb1, Pb2, Pb7, Pb8, and Pb14 (when S21a is occupied). The occupancies 240 of these sites are represented by Pb only (Pb2, Pb7, and Pb14 sites) or by Pb and Bi atoms (Pb1 and 241

Pb8), with a 85:15 atomic ratio. These agree with the BVS calculation, resulting in values ranging
from 1.95 to 2.02 *v.u.* for the pure Pb sites and BVS of 2.10 and 2.08 *v.u.* for the mixed (Pb,Bi).
Finally, the Pb15 and Pb16 sites have a bicapped trigonal prismatic coordination and a mixed
(Pb_{0.85}Bi_{0.15}) site occupancy. Their BVS values are both 2.10 *v.u.*

Five Me^{3+} positions occur in the crystal structure of marcobaldiite, *i.e.* Sb4, Sb12, As5, As13, 246 and the split site As17. Taking into account the shortest Me-S distances (= the strongest bonds: Me-247 S < 2.70 Å), these sites display a trigonal pyramidal coordination, with Me occupying the apex of 248 the pyramid. The coordination sphere is completed by three additional longer bonds. Sb4 is a pure 249 Sb position, with an average $\langle Me-S \rangle$ distance of 2.504 Å and a BVS of 2.94 v.u.. Antimony is 250 dominant also at the Sb12 position, having a refined site occupancy factor (s.o.f.) of Sb_{0.98}As_{0.02}. 251 The average $\langle Me-S \rangle$ distance is 2.462 Å and the BVS indicates moderate overbonding (3.14 v.u.). 252 As5, As13, and As17 are three As-dominant positions showing increasing As contents. Indeed, the 253 refined s.o.f. point to As_{0.63}Sb_{0.37}, As_{0.91}Sb_{0.09}, and As_{1.00}, respectively. Actually, the As17 site is 254 split into two sub-positions (As17a and As17b) having a partial occupancy (0.75 and 0.25, 255 256 respectively) in order to avoid too-short Me-Me distances. S21 is also split in two sub-positions S21a and S21b, with s.o.f. 0.75 and 0.25, respectively. As a consequence, As17a, shift towards the 257 258 N = 3 layer, is bound to S21a, and As17b to S21b.

The refined s.o.f. at As5 and As13 result in relatively high BVS (3.15 and 3.11 *v.u.*, respectively). The higher As contents are coupled with decreasing mean $\langle Me-S \rangle$ distances (2.325 and 2.262 Å for As5 and As13, respectively). The mean bond distances observed at the As17 subsites are shorter than the ideal one, owing the average nature of the ligand positions, related to the statistical occupancy of such positions. This explains the too high BVS obtained (3.22 and 3.52 *v.u.*).

The crystal structure of marcobaldiite is completed by the mixed (Pb,Sb)9 position, within the N = 4 layer, having a distorted octahedral coordination and an average bond distance of 2.922 Å. The refined s.o.f. points to a site occupancy Pb_{0.61}Sb_{0.39}. The calculated BVS, 2.35 *v.u.*, fits the theoretical one, *i.e.* 2.39 *v.u*.

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270 5. Crystal-chemistry of jordanite homologues

271 **5.1. Structural formula**

As described above, the crystal structure of marcobaldiite (Fig. 3) can be considered as a 1:1 alternation of N = 3 (kirkiite-type) and N = 4 (jordanite-type) octahedral layers, separated by a layer formed by Pb trigonal prismatic polyhedra and one As trigonal pyramid (Fig. 2). The N = 3 layer of marcobaldiite has chemical composition [Pb₃(Pb_{0.85}Bi_{0.15})(Sb_{1.37}As_{0.63})_{$\Sigma 2$}S₈]^{-1.85}. This chemistry can

be compared with that given by Makovicky *et al.* (2006) for the N = 3 octahedral layer in kirkiite, 276 *i.e.* $[Pb_3(Pb_{0.75}Bi_{0.25})(As_{1.42}Bi_{0.58})_{\Sigma 2}S_8]^{-1.75}$. Figure 4 (top) shows the comparison between the N = 3277 layers in marcobaldiite and kirkiite, whereas the site occupancies are compared in Table 9. The N =278 4 layer (Fig. 4, bottom) has chemical composition $[Pb_4(Pb_{0.85}Bi_{0.15})(Pb_{0.61}Sb_{0.39})(Sb_{1.07}As_{0.93})_{\Sigma 2}S_{10}]^{-1.46}$ 279 and can be compared with that of jordanite isotypes from the Pollone mine (Biagioni et al., 2016a), 280 having a chemistry $[Pb_5(Pb_{0.50}Sb_{0.50})(Sb_{0.61-1.23}As_{1.39-0.77})_{\Sigma 2}S_{10}]^{-1.5}$. Finally, the layer composed by Pb 281 trigonal prisms and split As site has chemical composition [Pb(Pb_{0.85}Bi_{0.15})₂AsS₃]^{+3.3}, to be 282 compared with $[Pb_3AsS_3]^{+3}$ in jordanite homologues and $[Pb_2(Pb_{0.50}Bi_{0.50})AsS_3]^{+3.5}$ in kirkiite. 283

The crystal-chemical formula of marcobaldiite can be obtained by the sum of the chemical 284 compositions of the N = 3, N = 4 and the trigonal prismatic atomic layer: 285 $[Pb_{3}(Pb_{0.85}Bi_{0.15})(Sb_{1.37}As_{0.63})_{\Sigma 2}S_{8}]^{-1.85} + [Pb_{4}(Pb_{0.85}Bi_{0.15})(Pb_{0.61}Sb_{0.39})(Sb_{1.07}As_{0.93})_{\Sigma 2}S_{10}]^{-1.46}$ 286 + $[Pb(Pb_{0.85}Bi_{0.15})_2AsS_3]^{+3.30}$ Pb8(Pb0.85Bi0.15)4(Pb0.61Sb0.39)(As2.56Sb2.44) \$\subset 5.00S21, = 287 is that 288 Pb_{12.01}(Sb_{2.83}As_{2.44}Bi_{0.60})_{25.99}S₂₁, to be compared with the formula obtained through chemical analysis, *i.e.* $Pb_{11,98}(Sb_{2,90}As_{2,33}Bi_{0,79})_{\Sigma 6,02}S_{20,80(20)}$. It is close to the ideal stoichiometric formula 289 290 $Pb_{12}(Sb_3As_2Bi)S_{21}$.

All known compositions of jordanite homologues have been represented in the As-Sb-Bi sub-system (Fig. 5). Taking into account the ideal formulae of geocronite, $Pb_{14}Sb_6S_{23}$, and As-rich kirkiite, $Pb_{10}As_4Bi_2S_{19}$, one can see that the ideal formula of marcobaldiite is quite identical to the 1:1 mixture of these homologues, having formula $Pb_{24}(Sb_6As_4Bi_2)S_{42} = 2 \times Pb_{12}(Sb_3As_2Bi)S_{21}$.

Since Bi is not dominant at any site of the crystal structure of marcobaldiite, the structural formula derived through the single-crystal study could be reduced to a stoichiometric one taking into account the coupled substitution $Pb^{2+} + Sb^{3+} = Bi^{3+} + Pb^{2+}$. Indeed, *ca.* 0.60 Bi *apfu* are distributed among four Pb positions (*i.e.*, Pb1, Pb8, Pb15, and Pb16), whereas Sb is replaced by *ca.* 0.60 Pb *apfu* at the Pb9 position. Subtracting Bi, the structural formula become Pb₁₂(Sb_{3.44}As_{2.56})₂₆S₂₁.

301

5.2. Bismuth in the jordanite homologous series

The occurrence of relatively high contents of Bi in this new mineral species is a rare feature shown by jordanite homologues. Indeed, the findings of high Bi contents in these minerals are limited to three localities, *i.e.* Aghios Philippos, Kirki, Thrace, Greece (Moëlo *et al.*, 1985; Moëlo *et al.*, 1990); La Fossa crater, Vulcano, Aeolian Islands, Italy (Borodaev *et al.*, 1998; Pinto *et al.*, 2006); and the Pollone mine, Apuan Alps, Tuscany, Italy.

Kirkiite shows Bi contents up to 15.5 and 18.27 wt% in samples from Kirki and Vulcano,
respectively (Moëlo *et al.*, 1985; Pinto *et al.*, 2006), corresponding to a maximum content of 3.09

Bi apfu, close to the ideal formula Pb₁₀As₃Bi₃S₁₉ (Fig. 5). Moreover, Moëlo et al. (1990) reported 310 the identification of an Sb-bearing variety of kirkiite, with two samples having average chemical 311 compositions of Pb_{10,21}(As_{3.48}Sb_{1.36}Bi_{1.16})S_{19.70} and Pb_{10,27}(As_{3.89}Sb_{1.05}Bi_{1.05})S_{19.90} (Fig. 5), ideally 312 close to $Pb_{10}(As_4SbBi)S_{19}$. This is the lowest Bi content reported for the N = 3 homologue of the 313 jordanite series. The solution of the crystal structure of kirkiite by Makovicky et al. (2006) could 314 suggest the cross-substitution of Pb and Bi at the Pb4 and Pb6 positions, with Bi being 315 preferentially partitioned at the Bi1/As1 split site. The chemical variability reported by Moëlo et al. 316 (1985, 1990) could be interpreted in the light of this crystal-chemical behaviour, with the 317 318 composition of the Sb-rich variety corresponding to the occurrence of a mixed (Bi_{0.50}Sb_{0.50}) occupancy at the Bi1/As1 site of Makovicky et al. (2006). In conclusion, available chemical data 319 320 suggests that Bi is always present in the N = 3 homologue kirkiite.

On the contrary, the N = 4 homologues are usually Bi-poor, with Bi contents less up *ca*. 0.50 wt% (*e.g.*, Pollone mine, Apuan Alps, Italy – Biagioni *et al.*, 2016a; Darasun ore deposit, Eastern Transbaikalia, Russia – Bryzgalov *et al.*, 2011). The highest Bi contents were reported by Moëlo *et al.* (1985) on a sample of jordanite associated with type kirkiite, containing up to 4.6 wt% Bi, corresponding to an ideal formula Pb₁₄As₅BiS₂₃ (Fig. 5).

According to Figure 5, the microprobe analysis of marcobaldiite agrees with a mixture of geocronite and (Sb,As)-rich kirkiite. According to its crystal structure, the kirkiite-type component is enriched in Sb relatively to As.

The description of marcobaldiite, characterized by a significant Bi content, similar to that 329 reported for Bi-rich jordanite from Kirki, poses the question about the role of this element in the 330 stabilization of the N = 3.5 homologue. As a matter of fact, one could note that Bi-rich jordanite and 331 kirkiite coexist without this intermediate new homologue and could hypothesize that the three 332 pnictogens As, Sb, and Bi are necessary for the crystallization of marcobaldiite. However, Moëlo et 333 al. (1990) found Sb-rich kirkiite, showing the simultaneous presence of these three elements; also in 334 this case, no hints about the occurrence of marcobaldiite were found. In addition, the recent finding 335 of the As-rich analogue of marcobaldiite, arsenmarcobaldiite (Biagioni et al., 2016c), substantially 336 337 devoid of Bi (Bi ≈ 0.30 wt%), seems to discard the hypothesis that this element plays a key role in the formation of the N = 3.5 homologue. 338

339

340 5.3. Conditions of formation

The geological history of the baryte + pyrite ore deposit of the Pollone mine is very complex and its origin is still debated (Carmignani *et al.*, 1976; Costagliola *et al.*, 1998). Recently, Biagioni *et al.* (2016a) provided new details for understanding the evolution of the Pollone ore deposit. Indeed, the baryte + pyrite ore bodies have been involved in the Alpine tectono-metamorphic events affecting the rocks belonging to the Apuan Alps metamorphic complex, indicating a pre-Alpine origin of the mineralization. During its late stage evolution, the formation of extension vein systems favoured the local remobilization of the ore bodies, giving rise to the peculiar sulfosalt assemblages occurring at the Pollone mine.

The occurrence of Bi is an unusual feature in the framework of the mineralogy of sulfosalts 349 from Apuan Alps, since up to now very few Bi sulfosalts have been identified from the 350 hydrothermal veins (e.g., Orlandi et al., 2010); at the Pollone mine, small amounts of Bi have been 351 352 reported in sterryite (up to 0.26 wt% - Moëlo et al., 2011) and jordanite/geocronite (up to 0.47 wt% 353 - Biagioni et al., 2016a). According to Moëlo et al. (1985), the elemental association of Pb, Bi, and 354 As is typical of thermo-chemical conditions such as high T and weak $f(S_2)$ favouring the incorporation of As as an anion in the crystallizing sulfides (e.g., As-bearing pyrite or arsenopyrite). 355 356 On the contrary, a high $f(S_2)$ could allow the sulfidation of As and its occurrence in sulfosalts; such a high $f(S_2)$ values could be achieved through the interaction between the hydrothermal solution and 357 358 a pre-existing sulfide ore deposit. At the Pollone mine, the late recrystallization stage of the baryte + pyrite ore could have locally mobilized As under high $f(S_2)$ conditions, favouring its sulfidation to 359 360 As³⁺ (Biagioni *et al.*, 2016a), and thus promoting the crystallization of complex Sb-As-(Bi) sulfosalt species. Since it is likely that the stabilization of the N = 3.5 homologue marcobaldiite is not related 361 to the occurrence of Bi, its crystallization could be related to small local changes in thermo-362 chemical conditions as well as Pb/(Sb,As,Bi) ratio, sufficient to induce the formation of distinct 363 sulfosalts. 364

However, the question of the stabilization of marcobaldiite remains open, as this phase has 365 not been observed in experimental studies nor in association with jordanite and kirkiite at Aghios 366 Philippos, Kirki, Greece. Whereas kirkiite and jordanite have been synthesized (e.g., Walia & 367 Chang, 1973), no hints of the N = 3.5 homologue were found in experimental studies. However, it 368 should be noted that some old chemical analyses of jordanite homologues fit better with the 369 composition $Pb_{12}(Sb/As)_6S_{21}$ than with that of the N = 4 homologues. As an example, Fisher (1940) 370 371 reported that the old formula of jordanite was given Pb₄As₂S₇, corresponding to one third of the ideal $Pb_{12}As_6S_{21}$ formula of a N = 3.5 homologue. Likely, such discrepancies were related to 372 373 inaccuracies in chemical analyses; the same composition reported for phase A of Walia & Change (1973) can be recalculated, on the basis of 6 (As+Bi) apfu, as $Pb_{11.57}(As_{4.5-3.9}Bi_{1.5-2.1})S_{20.57}$, closer to 374 Pb₁₂(As,Bi)₆S₂₁ than Pb₁₀(As,Bi)₆S₁₉. In this case, the X-ray powder diffraction pattern of phase A 375 seems to agree with that of kirkiite and not with that of marcobaldiite, even if the diffraction 376 377 patterns of the jordanite homologues are very similar.

Owing to such limitations in old chemical analyses and the similarity in the optical properties as well as in the X-ray powder diffraction patterns with other jordanite homologues, it is likely that the presence of marcobaldiite in sulfosalt assemblages could have been overlooked. Nevertheless, at Kirki, Bi-rich jordanite appears directly in epitactic overgrowth on kirkiite, without any intermediate phase (Moëlo *et al.*, 1985). The best way to detect it is to perform single-crystal Xray diffraction studies and systematic and accurate electron-microprobe analyses.

384

385 6. Conclusion

The description of marcobaldiite represents a new case of homology in the sulfosalt realm and increases the crystal-chemical complexity of the jordanite homologous series. Makovicky *et al.* (2006) predicted the possible existence of N = 1, N = 2, and N = 5 homologues within the jordanite series. Even if they have not been found yet, the discovery of the N = 3.5 homologue allows us to hypothesize the existence of other kinds of combination of the N = 3 and N = 4 octahedral layers, similarly to what happens in the sartorite homologous series (*e.g.*, Moëlo *et al.*, 2008).

This finding is a further confirmation of the great mineralogical variability of the sulfosalt assemblages from the hydrothermal veins of the Apuan Alps metamorphic complex. The factors controlling the sulfosalt crystallization are not only related to the occurrence of minor components (*e.g.*, Ag, Cu, Hg, Tl), filling specific atom positions, and peculiar $f(O_2)$ and $f(S_2)$ conditions, promoting the crystallization of oxysulfosalts or persulfosalts; indeed, it seems likely that very subtle variations in the Pb/(Sb,As,Bi) atomic ratios in the crystallizing medium can allow the formation of distinct and original mineral species.

399

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489 **Table captions**

490 **Table 1** – Sulfosalts identified in the Pollone ore deposit. In bold, minerals having their type 491 locality at the Pollone mine.

- 492 **Table 2** Reflectance data (%) for marcobaldiite in air.
- 493 **Table 3** Electron microprobe analysis of marcobaldiite: chemical composition as wt% (10 spot
- analyses) and number of atoms per formula unit (apfu) on the basis of $\Sigma Me = 18$ apfu.
- **Table 4** –X-ray powder diffraction data for marcobaldiite. Intensities and d_{hkl} were calculated using
- 496 the software *PowderCell* 2.3 (Kraus and Nolze, 1996) on the basis of the structural model given in
- 497 Table 5. The six strongest reflections are given in bold. Only reflections with $I_{calc} \ge 5$ were
- 498 reported, if not observed. Observed intensities were visually estimated (vs = very strong; s = strong;
- 499 ms = medium-strong; m = medium; mw = medium-weak; w = weak; vw = very weak).
- Table 5 Crystal data and summary of parameters describing data collection and refinement for
 marcobaldiite.
- **Table 6** Atomic coordinates, site occupancy factor (s.o.f.), and equivalent isotropic displacement parameters ($Å^2$) for marcobaldiite.
- 504 **Table 7** Coordination number (C.N.), mean bond distance $\langle Me-S \rangle$ (Å), and bond-valence sums
- 505 (BVS in valence unit, *v.u.*) for cation sites in marcobaldiite.
- **Table 8** Bond valence sums (BVS in valence unit, *v.u.*) for S atoms in marcobaldiite.
- 507 Table 9 Comparison between site labels and site occupancy factors (s.o.f.) in marcobaldiite,
 508 kirkiite, and jordanite/geocronite isotypes.
- 509

510 Figure captions

- 511 Figure 1 Reflected light microscopy image of marcobaldiite, showing polysynthetic twinning.
- 512 Holotype material, Natural History Museum of Pisa University, catalogue number # 19709.
- 513 **Figure 2** Projection along **c** of the crystal structure of marcobaldiite.
- **Figure 3** Modular organization of the crystal structure of marcobaldiite. Alternating oblique N = 3
- and N = 4 layers are separated by single slabs TP of trigonal prismatic Pb and split As17a/As17b.
- 516 The three shortest (Sb,As)–S and (Pb,Sb)–S bonds have been enhanced. Shaded ellipses: lone-
- 517 electron-pair micelles of trivalent Sb and As.
- **Figure 4** Comparison between N = 3 and N = 4 octahedral layers of marcobaldiite with those
- 519 occurring in kirkiite (a) and jordanite/geocronite (b).

Figure 5 – Projection in the As-Sb-Bi sub-system of various compositions within the jordanite homologous series. Homologues: N = 3, pink squares (kirkiite); N = 4, yellow triangles (jordanite and geocronite); N = 3.5, blue lozenges (marcobaldiite and arsenmarcobaldiite). EPMA: microprobe analysis; STR: crystal structure, with L3: composition of the kirkiite sub-part (= L3 layer + intermediate slab), and L4: composition of the geocronite sub-part.

527	Table 1 – Sulfosalts	identified ir	the	Pollone	ore	deposit.	In	bold,	minerals	having	their	type
528	locality at the Pollone	mine.										

Mineral	Chemical formula	Mineral	Chemical formula
Boulangerite	Pb ₅ Sb ₄ S ₁₁	Owyheeite	Ag ₃ Pb ₁₀ Sb ₁₁ S ₂₈
Bournonite	CuPbSbS₃	Parasterryite	Ag4Pb20(Sb,As)24S58
Carducciite	(AgSb)Pb ₆ (As,Sb) ₈ S ₂₀	Polloneite	$AgPb_{46}As_{26}Sb_{23}S_{120}$
Chovanite	Pb ₂₈ (Sb,As) ₃₀ S ₇₂ O	Proustite	Ag ₃ AsS ₃
Diaphorite	Ag ₃ Pb ₂ Sb ₃ S ₈	Pyrargyrite	Ag ₃ SbS ₃
Ferdowsiite	$Ag_8Sb_5As_3S_{16}$	Seligmannite	CuPbAsS₃
Fettelite	[Ag ₆ As ₂ S ₇][Ag ₁₀ HgAs ₂ S ₈]	Sterryite	$Cu(Ag,Cu)_3Pb_{19}(Sb,As)_{22}(As-As)S_{56}$
Geocronite	Pb14Sb6S23	Tennantite	$Cu_6Cu_4(Fe,Zn)_2As_4S_{13}$
Jordanite	$Pb_{14}As_6S_{23}$	Tetrahedrite	Cu ₆ Cu ₄ (Fe,Zn) ₂ Sb ₄ S ₁₃
Marcobaldiite	$Pb_{12}(Sb_3As_2Bi)_{\Sigma 6}S_{23}$	Twinnite	PbSbAsS ₄
Meerschautite	(Ag,Cu)5.5Pb42.4(Sb,As)45.1S112O0.8	Xanthoconite	Ag ₃ AsS ₃
Meneghinite	CuPb13Sb7S24	Zinkenite	Pb ₉ Sb ₂₂ S ₄₂

Table 2 – Reflectance data (%) for marcobaldiite in air.

λ/nm	R _{min}	R _{max}	λ/nm	R _{min}	R _{max}
400	31.2	35.2	560	30.6	39.0
420	32.3	39.7	580	30.4	38.5
440	32.7	41.7	589	30.4	38.5
460	31.8	40.2	600	30.1	38.4
470	31.6	40.1	620	30.1	38.1
480	31.8	40.1	640	30.2	38.0
500	31.4	39.9	650	30.0	37.6
520	31.1	39.5	660	29.9	37.6
540	31.0	39.5	680	29.6	36.9
546	30.9	39.6	700	29.1	36.4

Element	wt%	range	e.s.d.
Pb	64.05	63.52 – 64.46	0.34
Bi	4.24	4.16 – 4.35	0.06
Sb	9.10	8.89 – 9.38	0.14
As	4.51	4.35 – 4.62	0.08
S	17.24	16.89 – 17.42	0.19
Total	99.14	98.29 - 100.02	0.53
apfu (<i>Me</i> = 18)			
Pb	11.98	11.91 – 12.06	0.04
Bi	0.79	0.77 – 0.80	0.01
Sb	2.90	2.84 – 2.96	0.03
As	2.33	2.27 – 2.40	0.03
S	20.80	20.47 - 21.11	0.20
Ev (%)	0.8	-0.5 – 2.6	1.1
F_{V} (%) = [Σ (val-	$(-) - \Sigma(y)$	al-)] x 100 / Σ(va	I_)

Table 3-Electron microprobe analysis of marcobaldiite: chemical composition as wt% (10 spot533 analyses) and number of atoms per formula unit (apfu) on the basis of $\Sigma Me = 18$ apfu. 534

Ev (%) = $[\Sigma(val+) - \Sigma(val-)] \times 100 / \Sigma(val-).$

535

Table 4 –X-ray powder diffraction data for marcobaldiite. Intensities and d_{hkl} were calculated using the software *PowderCell* 2.3 (Kraus and Nolze, 1996) on the basis of the structural model given in Table 5. The eight strongest reflections are given in bold. Only reflections with $I_{calc} \ge 5$ were reported, if not observed. Observed intensities were visually estimated (vs = very strong; s = strong; ms = medium-strong; m = medium; mw = medium-weak; w = weak; vw = very weak).

icului	n suo	$m_{\rm S}, m = m_{\rm S}$	Jului	1, 111 w –	mearun	ii wea	$\mathbf{K}, \mathbf{w} = \mathbf{w}$	can, v	$w = v c_1$	y weak).
	lobs	dobs	Icalc	dcalc	hkl	lobs	dobs	<i>I</i> calc	d_{calc}	hkl
	W	4.45	8	4.406	-2 1 1	VW	2.553	8	2.568	-1 -10 2
	W	3.835	25	3.832	210	W	2.489	5	2.496	-282
	147	2 727	6	3.756	2 -3 0	VW	2.442	3	2.434	-2 10 1
	vv	3.737 l	5	3.735	-2 -1 2	mw	2 388	2	2.391	082
			45	3.692	220	IIIVV	2.300 }	2	2.385	-2 -10 2
			18	3.683	-2 -2 2	VW	2.347	1	2.342	1 -9 2
			20	3.676	0 -3 2	W	2.294	1	2.297	162
		ſ	7	3.596	012		(ŕ 5	2.241	-1 10 2
	me	3 568	21	3.591	2 -4 0			48	2.240	2 -2 2
	1115	3.300	25	3.577	-2 -3 2	VC	2 222	6	2.240	-2 -11 2
		l	40	3.555	0 -4 2	v5	2.233	6	2.229	2 10 0
		(17	3.448	022			5	2.204	0 -12 2
			17	3.431	-2 -4 2		l	46	2.203	-422
	m	3.404	26	3.398	2 -5 0		ĺ	7	2.118	370
			11	3.398	0 -5 2	~	2 1 2 5	54	2.110	-204
		l	14	3.384	-232	5	2.125	9	2.107	-2 -12 2
			28	3.314	240		l	12	2.094	2 11 0
	m	2.256	19	3.274	032	VW	2.072	13	2.074	0 -13 2
	[[]	3.200 -	52	3.259	-2 -5 2	W	2.035	3	2.028	380
		ſ	100	3.219	090	W	2.020	5	2.022	2 -13 0
	ms	3.202 {	27	3.219	0 -6 2	mw	1.980	6	1.986	-2 -13 2
		Ĺ	30	3.208	-242	2014	1.050	- 20	1.957	-4 -7 2
		ſ	29	3.109	250	IIIVV	1.950 2	6	1.956	0 -14 2
	m	3.075 {	9	3.088	042	2014	1 011	8	1.916	420
		l	30	3.075	-2 -6 2	IIIVV	1.911 -	6	1.906	2 -14 0
		ſ	28	3.032	0 -7 2		ſ	- 20	1.895	-2 -9 4
	ms	3.016	12	3.023	-252	m	1.878	6	1.880	0 12 2
		l	35	2.991	2 -7 0		l	. 6	1.868	-4 -2 4
	me	2 885 ∫	13	2.908	260		(- 9	1.847	-2 13 2
	1115	2.003 ح	36	2.902	052			17	1.843	2 -11 2
		ſ	10	2.846	0 -8 2	S	1.848 <	7	1.839	4 -7 0
	m	2.832	5	2.841	-3 -2 2			16	1.835	272
		l	20	2.839	-262		l	、 7	1.811	0 -7 4
	W	2.789	28	2.795	2 -8 0	VW	1.797	6	1.798	024
	VW	2.756	4	2.755	1 -5 2	S	1.775	5	1.778	0 -8 4
	2014	2 707	22	2.721	062	W	1.753	5	1.749	-2 14 2
	IIIW	2.101 1	5	2.720	122	W	1.716	12	1.705	-4 11 2
	VW	2.652	19	2.662	-272	W	1.661	12	1.658	-292
	W	2.606	6	2.612	2 -9 0	w	1.608	12	1.609	0 18 0

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Crystal data	
Crystal size (mm ³)	0.09 x 0.08 x 0.04
Cell setting, space group	Triclinic, P-1
<i>a</i> (Å)	8.9248(9)
b (Å)	29.414(3)
<i>c</i> (Å)	8.5301(8)
α (°)	98.336(5)
β (°)	118.175(5)
γ (°)	90.856(5)
V (Å ³)	1944.1(3)
Z	2
Data collection and refinement	
Radiation, wavelength (Å)	Μο <i>Κ</i> α, λ = 0.71073
Temperature (K)	293
2θ _{max} (°)	50.29
Measured reflections	41210
Unique reflections	6790
Reflections with $F_0 > 4\sigma(F_0)$	6188
R _{int}	0.0409
Ro	0.0269
	$-10 \le h \le 10,$
Range of <i>h</i> , <i>k</i> , <i>l</i>	$-34 \le k \le 34,$
	−10 ≤ / ≤ 10
$R[F_{o} > 4\sigma(F_{o})]$	0.0668
R (all data)	0.0702
wR (on F_0^2)	0.2228
Goof	1.093
Number of least-squares parameters	363
Maximum and	10.88 (at 0.97 Å from Pb5)
minimum residual peak (<i>e</i> Å ⁻³)	-3.76 (at 1.05 Å from S18)

Table 5 – Crystal data and summary of parameters describing data collection and refinement for
 marcobaldiite.

Site	s.o.f.	x/a	y/b	z/c	U _{eq}
Pb1	Pb _{0.85} Bi _{0.15}	0.8503(1)	0.38701(4)	0.4904(2)	0.0199(3)
Pb2	Pb _{1.00}	0.8518(1)	0.38638(4)	0.9939(2)	0.0199(3)
Pb3	Pb _{1.00}	0.2253(1)	0.49413(4)	0.7368(2)	0.0213(3)
Sb4	Sb _{1.00}	0.2412(2)	0.50143(6)	0.2478(2)	0.0116(4)
As5	As _{0.63(3)} Sb _{0.37(3)}	0.3689(3)	0.39399(8)	0.0047(3)	0.0171(8)
Pb6	Pb _{1.00}	0.3821(2)	0.39554(4)	0.5075(2)	0.0258(3)
Pb7	Pb _{1.00}	0.7725(1)	0.16711(4)	0.8456(1)	0.0192(3)
Pb8	Pb _{0.85} Bi _{0.15}	0.7695(1)	0.16641(4)	0.3381(2)	0.0194(3)
Pb9	Pb _{0.61(1)} Sb _{0.39(1)}	0.0830(2)	0.05688(5)	0.9425(2)	0.0230(3)
Pb10	Pb _{1.00}	0.0780(1)	0.05992(4)	0.4471(2)	0.0195(3)
Pb11	Pb _{1.00}	0.6128(1)	0.04819(4)	0.9512(2)	0.0201(3)
Sb12	Sb _{0.98(3)} As _{0.02(3)}	0.5985(2)	0.05375(6)	0.4500(2)	0.0146(6)
As13	$As_{0.91(3)}Sb_{0.09(3)}$	0.2850(3)	0.15729(9)	0.8444(4)	0.0157(9)
Pb14	Pb _{1.00}	0.2937(2)	0.16075(4)	0.3477(2)	0.0243(3)
Pb15	Pb _{0.85} Bi _{0.15}	0.4812(2)	0.27712(4)	0.2586(2)	0.0238(3)
Pb16	Pb _{0.85} Bi _{0.15}	0.4818(2)	0.27741(4)	0.7431(2)	0.0249(3)
As17a	AS0.75(1)	0.9939(4)	0.30213(11)	0.2707(4)	0.0146(7)
As17b	As _{0.25(1)}	0.9756(12)	0.2494(3)	0.2340(14)	0.0146(7)
Pb18	Pb _{1.00}	0.0467(2)	0.27511(4)	0.7796(2)	0.0347(4)
S1	S _{1.00}	0.6054(8)	0.3514(2)	0.0996(8)	0.0161(12)
S2	S _{1.00}	0.6316(8)	0.3389(2)	0.6100(8)	0.0162(12)
S3	S _{1.00}	0.0088(8)	0.4443(2)	0.3618(8)	0.0175(13)
S4	S _{1.00}	0.9991(8)	0.4492(2)	0.8435(8)	0.0150(12)
S5	S _{1.00}	0.6084(8)	0.4537(2)	0.4324(8)	0.0159(12)
S6	S _{1.00}	0.6113(8)	0.4537(2)	0.8754(8)	0.0145(12)
S7	S _{1.00}	0.2267(8)	0.3527(2)	0.1134(9)	0.0216(13)
S8	S _{1.00}	0.2236(9)	0.3538(2)	0.7092(9)	0.0227(14)
S 9	S _{1.00}	0.5456(8)	0.1986(2)	0.9932(8)	0.0150(12)
S10	S _{1.00}	0.5942(8)	0.2136(2)	0.5278(8)	0.0166(12)
S11	S 1.00	0.8941(9)	0.1089(2)	0.1302(9)	0.0189(13)
S12	S 1.00	0.8730(8)	0.1013(2)	0.6092(8)	0.0148(12)
S13	S _{1.00}	0.1968(8)	0.0032(2)	0.2106(9)	0.0169(12)
S14	S _{1.00}	0.1985(8)	0.0050(2)	0.7444(8)	0.0167(12)
S15	S _{1.00}	0.4821(8)	0.0967(2)	0.6306(8)	0.0153(12)
S16	S _{1.00}	0.4785(8)	0.0960(2)	0.1916(8)	0.0159(12)
S17	S _{1.00}	0.1671(9)	0.1958(2)	0.6078(9)	0.0236(14)
S18	S 1.00	0.1711(8)	0.1962(2)	0.0017(9)	0.0205(13)
S19	S _{1.00}	0.8506(8)	0.2722(2)	0.3915(9)	0.0222(14)
S20	S _{1.00}	0.8518(8)	0.2734(2)	0.9742(9)	0.0196(13)
S21a	S _{0.75(1)}	0.2244(10)	0.2630(3)	0.3651(10)	0.0111(15)
S21b	S _{0.25(1)}	0.233(3)	0.2882(10)	0.381(3)	0.0111(15)

547 Table 6 – Atomic coordinates, site occupancy factor (s.o.f.), and equivalent isotropic displacement
 548 parameters (Å²) for marcobaldiite.

Table 7 – Coordination number (C.N.), mean bond distance $\langle Me-S \rangle$ (Å), and bond-valence sums 550 551 (BVS – in valence unit, v.u.) for cation sites in marcobaldiite.

Site	s.o.f.	C.N.	<me-s></me-s>	BVS	Site	s.o.f.	C.N.	<me-s></me-s>	BVS
Pb1	Pb _{0.85} Bi _{0.15}	VII	3.031	2.10	Pb10	Pb _{1.00}	VI	2.996	1.95
Pb2	Pb _{1.00}	VII	3.050	1.95	Pb11	Pb _{1.00}	VI	3.016	1.95
Pb3	Pb _{1.00}	VI	3.012	1.99	Sb12	$Sb_{0.98}As_{0.02}$	+	2.462	3.14
Sb4	Sb _{1.00}	+ *	2.504**	2.94	As13	$As_{0.91}Sb_{0.09}$	III	2.262	3.11
As5	$As_{0.63}Sb_{0.37}$	III	2.325	3.15	Pb14	Pb _{1.00}	VII	3.046***	2.02
Pb6	Pb _{1.00}	VII	3.055***	1.97	Pb15	Pb _{0.85} Bi _{0.15}	VIII	3.073	2.10
Pb7	Pb _{1.00}	VII	3.024	2.00	Pb16	Pb _{0.85} Bi _{0.15}	VIII	3.075	2.10
Pb8	Pb _{0.85} Bi _{0.15}	VII	3.013	2.08	As17a	As _{0.75}	Ш	2.233	3.22
Pb9	$Pb_{0.61}Sb_{0.39}$	VI	2.922	2.35	As17b	As _{0.25}	III	2.191	3.52
Pb10	Pb _{1.00}	VI	2.996	1.95	Pb18	Pb _{1.00}	VI	3.023	1.75

* 3 short and 3 longer bonds. ** Mean of the 3 short bonds. *** Calculated assuming the full occupancy of S21b and S21a for Pb6 and Pb14, respectively

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553 Table 8 – Bond valence sums (BVS – in valence unit, v.u.) for S atoms in marcobaldiite.

Site	Site BVS		BVS
S1	2.06	S12	2.09
S2	2.06	S13	1.93
S3	2.02	S14	2.01
S4	2.03	S15	1.97
S5	1.91	S16	1.92
S6	1.95	S17	1.88
S7	1.96	S18	1.95
S8	1.89	S19	2.21
S9	1.96	S20	2.13
S10	1.98	S21a	2.11
S11	2.00	S21b	2.08

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557 Table 9 – Comparison between site labels and site occupancy factors (s.o.f.) in marcobaldiite,
558 kirkiite, and jordanite/geocronite isotypes.

Marcobaldiite		Kirkiite		Jordanite/geocronite	
Site	s.o.f.	Site	s.o.f.	Site	s.o.f.
Pb1	Pb _{0.85} Bi _{0.15}	Pb4	Pb _{0.75} Bi _{0.25}		
Pb2	Pb _{1.00}	Pb3	Pb _{1.00}		
Pb3	Pb _{1.00}	Pb1	Pb _{1.00}		
Sb4	Sb _{1.00}	Bi1/As1	Bi _{0.58} /As _{0.42}		
As5	As _{0.63} Sb _{0.37}	As2	As _{1.00}		
Pb6	Pb _{1.00}	Pb2	Pb _{1.00}		
Pb7	Pb _{1.00}			Pb7	Pb _{1.00}
Pb8	Pb _{0.85} Bi _{0.15}			Pb8	Pb _{1.00}
Pb9	Pb _{0.61} Sb _{0.39}			Pb2/Sb2	Pb _{0.50} /Sb _{0.50}
Pb10	Pb _{1.00}			Pb1	Pb _{1.00}
Pb11	Pb _{1.00}			Pb3	Pb _{1.00}
Sb12	Sb _{0.98} As _{0.02}			Sb4	Sb _{0.61-0.98} As _{0.39-0.02}
As13	As _{0.91} Sb _{0.09}			As6	As1.00-0.75Sb0.00-0.25
Pb14	Pb _{1.00}			Pb5	Pb _{1.00}
Pb15	Pb _{0.85} Bi _{0.15}	Pb7	Pb _{1.00}	Pb9	Pb _{1.00}
Pb16	Pb _{0.85} Bi _{0.15}	Pb6	Pb _{0.5} Bi _{0.5}	Pb10	Pb _{1.00}
As17	As _{1.00}	As3	As _{1.00}	As11	As _{1.00}
Pb18	Pb _{1.00}	Pb5	Pb _{1.00}	Pb12	Pb _{1.00}

559 Note: s.o.f. for kirkiite and jordanite/geocronite after Makovicky *et al.* (2006) and Biagioni *et al.* (2016a), 560 respectively.

Biagioni *et al*.

Figure 1 – Reflected light microscopy image of marcobaldiite, showing polysynthetic twinning.
Holotype material, Natural History Museum of Pisa University, catalogue number # 19709.



Figure 2 – Projection along **c** of the crystal structure of marcobaldiite.



Figure 3 – Modular organization of the crystal structure of marcobaldiite. Alternating oblique N = 3and N = 4 layers are separated by single slabs TP of trigonal prismatic Pb and split As17a/As17b. The three shortest (Sb,As)–S and (Pb,Sb)–S bonds have been enhanced. Shaded ellipses: loneelectron-pair micelles of trivalent Sb and As.



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Figure 4 – Comparison between N = 3 and N = 4 octahedral layers of marcobaldiite with those 575 occurring in kirkiite (top) and jordanite/geocronite (bottom). 576



Figure 5 – Projection in the As-Sb-Bi sub-system of various compositions within the jordanite homologous series. Homologues: N = 3, pink squares (kirkiite); N = 4, yellow triangles (jordanite and geocronite); N = 3.5, blue lozenges (marcobaldiite and arsenmarcobaldiite). EPMA: microprobe analysis; STR: crystal structure, with L3: composition of the kirkiite sub-part (= L3 layer + intermediate slab), and L4: composition of the geocronite sub-part.



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