# Article



# Bulachite, $[Al_6(AsO_4)_3(OH)_9(H_2O)_4]\cdot 2H_2O$ from Cap Garonne, France: Crystal structure and formation from a higher hydrate

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# Abstract

Bulachite specimens from Cap Garonne, France, comprise two intimately mixed hydrated aluminium arsenate minerals with the same Al:As ratio of 2:1 and with different water contents. The crystal structures of both minerals have been solved using data from low-dose electron diffraction tomography combined with synchrotron powder X-ray diffraction. One of the minerals has the same powder X-ray diffraction pattern (PXRD) as for published bulachite. It has orthorhombic symmetry, space group *Pnma* with unit-cell parameters a = 15.3994(3), b = 17.6598(3), c = 7.8083(1) Å and Z = 4, with the formula  $[Al_6(AsO_4)_3(OH)_9(H_2O)_4] \cdot 2H_2O$ . The second mineral is a higher hydrate with composition  $[Al_6(AsO_4)_3(OH)_9(H_2O)_4] \cdot 8H_2O$ . It has the same *Pnma* space group and unit-cell parameters a = 19.855(4), b = 17.6933(11) and c = 7.7799(5) Å i.e. almost the same b and c parameters but a much larger a parameter. The structures are based on polyhedral layers, parallel to (100), of composition  $[Al_6(AsO_4)_3(OH)_9(H_2O)_4]$  and with H-bonded H<sub>2</sub>O between the layers. The layers contain [001] spiral chains of edge-shared octahedra, decorated with corner connected AsO<sub>4</sub> tetrahedra that are the same as in the mineral liskeardite. The spiral chains are joined together by octahedral edge-sharing to form layers parallel to (100). Synchrotron PXRD patterns collected at different temperatures during heating of the specimen show that the higher-hydrate mineral starts transforming to bulachite when heated to 50°C, and the transformation is complete between 75 and 100°C.

**Keywords:** bulachite, Cap Garonne, new mineral, crystal structure, hydrated aluminium arsenate, electron diffraction, synchrotron powder XRD

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# Introduction

Bulachite, from Neubulach, northern Black Forest, Germany was first described by Walenta (1983). It occurs as aggregates of fine white needles encrusting quartz and associated with arsenocrandallite, malachite, bariopharmacosiderite and goethite. Walenta provided results from chemical and thermogravimetric analyses and powder X-ray diffraction (PXRD), that established the ideal composition as Al<sub>2</sub>(AsO<sub>4</sub>)(OH)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub> and the symmetry as orthorhombic with unit-cell parameters a = 15.53, b = 17.78 and c = 7.03 Å. A second occurrence of bulachite from Sardinia, Italy, was reported by Frau and Da Pelo (2001). The Sardinian specimens were collected at a mine dump from old workings in an arsenopyrite lens in the Sa Bidda Beccia area located ~20 km southwest of Cagliari in southern Sardinia. Here the mineral occurs as whitish, satiny, polycrystalline aggregates, usually on dark-brown layers of mansfieldite and goethite encrusting quartzite clasts. Scanning electron microscope (SEM)

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As part of an ongoing characterisation of hydrated Al arsenate minerals (Grey *et al.*, 2013, 2014, 2015, 2016*a,b*) we studied specimens of bulachite from Cap Garonne where the mineral forms patchy crusts of ultrathin lath-like crystals on yellow crystals of pharmacoalumite (Fig. 1). The ruler-shaped laths are typically up to 50 µm long by 0.4 µm wide and only 0.1 µm thick. A surprising result was that PXRD patterns of the specimens, while having a number of sharp peaks in common with the pattern reported by Walenta (1983), did not have the strongest bulachite peak at 7.78 Å but had a number of broad peaks not present in the Walenta pattern, including a strong peak at  $d \approx 10$  Å. Electron diffraction (ED) patterns on flat laths gave in-plane orthogonal dimensions of 17.7 Å × 7.8 Å with the 7.8 Å axis aligned along

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Fig. 1. Patchy regions of bulachite (white) on yellow pharmacoalumite from Cap Garonne. FOV 15 mm. Museums Victoria specimen M55455.

the long dimension. The ED patterns assisted in the indexing of the PXRD pattern, giving an orthorhombic unit cell with  $a \approx 19.9$ ,  $b \approx 17.7$  and  $c \approx 7.8$  Å and possible space groups *Pnma* or *Pn*<sub>21</sub>*a*.

An exception to the PXRD findings was obtained in a synchrotron PXRD data collection made on small clumps of crystal aggregates packed in a capillary. Rotation of the capillary during the experiment accidentally caused the bulk of the sample to move along the capillary out of the beam, so the diffraction pattern was obtained only for crystals from the outer surface of the small clumps that had adhered to the inner wall of the capillary. The resulting PXRD pattern was the same as that reported by Walenta (1983) for bulachite. This experiment suggested that the crusts of crystals have bulachite only at the surface and a different phase in the bulk, with the possibility that bulachite has formed from the bulk specimen by reaction at the surface. To test this hypothesis we obtained synchrotron PXRD data collections on a specimen that was heated at different temperatures. Above 50°C the diffraction pattern corresponding to the abovereported cell started changing and between 75°C and 100°C it had transformed completely to the pattern reported by Walenta (1983) for bulachite.

Electron diffraction patterns obtained for the sample heated to 100°C gave an orthorhombic cell with  $a \approx 15.4$ ,  $b \approx 17.7$  and  $c \approx 7.8$  Å and possible space groups *Pnma* or *Pn2*<sub>1</sub>*a*, i.e. the unit cell is the same as for the phase before heating, except for a large decrease in *a*. The *a* axis is normal to the ruler-shaped laths. The large decrease in *a* under very mild heating conditions is consistent with a layer structure having interlayer water molecules which are evolved on heating above ~50°C, resulting in a contraction of the interlayer separation.

We report here the application of synchrotron PXRD and precession ED methods to determine the structures of both bulachite and the higher-hydrate precursor mineral that transforms to bulachite on dehydration.

# Experimental

The specimens for study were collected by G.F. in Salle B, South mine, Cap Garonne, Var, France (43°4'53"N, 6°1'55"E; Favreau and Galea-Clolus, 2014). The specimens were checked for



Fig. 2. SEM back-scattered electron image of clumps of fibrous bulachite crystals from Cap Garonne. Museums Victoria specimen M55455.

chemical purity by energy-dispersive X-ray analysis in a scanning electron microscope (Fig. 2). The only metals detected were Al and As, in an atomic ration of 2:1, consistent with the formula obtained by Walenta (1983). The sample is preserved in the mineralogical collections of Geosciences, Museums Victoria, under registration number M55455. A sample of the type specimen of bulachite that was studied by Walenta (1983) was supplied by Thomas Theye. It was in the form of sub-millimetre tufts of densely packed fibrous crystals, see figure 4 in Walenta (1983).

# Powder X-ray diffraction

Laboratory PXRD scans showed that even light grinding of the specimens resulted in loss of long-range ordering (severe peak broadening) so for the synchrotron experiments, small unground clumps of subparallel fibres, such as shown in Fig. 2 were packed into a 0.5 mm diameter quartz capillary. As noted previously, in one of the earlier data collections, it was found that during rotation of the capillary the clumps moved along the capillary and out of the beam. Subsequent samples were prepared by packing a few mm of small clumps in a 0.5 mm quartz capillary then inserting a 0.3 mm capillary to the edge of the specimen, to stop any movement. The data collections were conducted on the powder diffraction beamline at the Australian Synchrotron (Wallwork et al., 2007). High energy 21 keV X-rays were used to reduce fluorescence due to As. The wavelength was 0.59028(6) Å, calibrated to NIST standard LaB<sub>6</sub> 660b. The capillary was positioned in the diffractometer rotation centre and spun at ca. 1 Hz. The X-ray beam was aligned to coincide with the diffractometer centre and slits were used to restrict the beam width to 2 mm. The data were collected using a Mythen position sensitive detector (Schmitt et al., 2003) covering 80°20 with a step size of 0.00375°20. Heating was supplied by an Oxford Cryostream system. The first data collection was at close to ambient (25°C). The temperature was then increased in steps of 25°C to 100°C and at each temperature pairs of data sets were collected at two detector positions 0.5° apart in order to cover the gaps between the detector modules.

Acquisition time at each position was 300 s. The data pairs were merged into single files using in-house data processing software, *PDViPcR*, available at the beamline. After the 100°C collection the sample was cooled to ambient and a dataset obtained. A dataset was also collected on an empty capillary so that correction for the capillary contribution could be applied in the calculation of pair distribution functions (PDF) from the datasets.

Le Bail fitting of the PXRD data to obtain refined unit-cell parameters was made using *JANA2006* (Petříček *et al.*, 2014). Manual backgrounds were used with interpolation between 40 points and a pseudo-Voigt peak shape function. It was necessary to include a parameter for selective peak broadening for reflections with h = 2n+1.

Pair distribution functions (PDFs) were generated from the synchrotron datasets, with  $Q_{max} = 16.9 \text{ Å}^{-1}$ . The raw diffraction data were corrected for capillary contribution, reduced to structure functions and then Fourier transformed to the PDFs, G(r) (Egami and Billinge, 2003). PDF calculations were conducted using *PDFgui* (Farrow *et al.*, 2007).

Gandolfi PXRD patterns were obtained on small clumps of crystals of the Neubulach type specimen studied by Walenta (1983). The Gandolfi patterns were obtained on a Rigaku XtaLAB Synergy diffractometer at the School of Chemistry, University of Melbourne. The instrument employs a HiPix CCD Plate detector and CuK $\alpha$  radiation.

#### Transmission electron microscopy

Crystal aggregates were sonicated in ethanol and the suspension was dispersed onto a holey carbon grid for transmission electron microscopy (TEM). The TEM studies were conducted using a JEOL 4000EX microscope with a  $LaB_6$  source and a spherical aberration coefficient of 1.06 mm. The microscope was operated at 400 kV, using parallel illumination geometry to minimise beam damage. TEM images were recorded on a Gatan slow scan CCD camera.

The new technique of low-dose electron diffraction tomography (Kodjikian and Klein, 2019) was applied to crystals from a specimen that had been heated to 100°C and had a PXRD pattern corresponding to bulachite (Walenta, 1983). Crystal aggregates were lightly crushed in an agate mortar in absolute ethanol and a droplet of the suspension was deposited on carboncoated copper grid. The ED data collection was made using a PhilipsCM300ST TEM, equipped with a TVIPS F416CMOS camera, a Nanomegas Spinningstar device and a Bruker Silicon Drift Detector for EDS. The diffraction intensities were measured on one selected particle (atomic composition from EDS: Al = 64.6% and As = 35.3%, oxygen was detected but not quantified).

The tilt step between two frames of the tomography was chosen as 1°, the precession angle was 1.05°. A data set of 105 frames with an exposure time of 0.5 s per frame was recorded from a single crystal of approximate size of 1  $\mu$ m × 0.5  $\mu$ m. During the automated data acquisition the beam was blanked from the sample between the exposures in order to irradiate the crystal only during the exposure time of the diffraction patterns. The dose rate was 0.056  $e^{-1}$ .s<sup>-1</sup>.Å<sup>-2</sup> and the total dose was 3  $e^{-1}$ .Å<sup>-2</sup>. Data analysis of the recorded patterns was performed with the software *PETS2.0* (Palatinus *et al.*, 2019) and *Jana2006* (Petříček *et al.*, 2014).

# Crystal-structure determinations

The crystal structure of the sample heated to 100°C (= bulachite) was independently solved from the low-dose electron diffraction

tomography data and from the synchrotron PXRD data. The individual ED patterns of the tomography series were treated for data reduction in *PETS 2.0.* Sections of the reciprocal lattice showed reflection conditions leading to the extinction symbol Pn-a consistent with space groups Pnma and  $Pn2_1a$ .

Superflip in JANA2006 (Petříček et al., 2014) was used to obtain a model in space group  $Pn2_1a$  from the PXRD data whereas the ED data were used in Superflip and in SIR2014 (Burla et al., 2015) where both space groups were tested and the best fit was found for a Pnma model. Both the ED and PXRD-derived models had the same topology of edge-shared Al-centred octahedra connected to AsO<sub>4</sub> tetrahedra by corner sharing. Based on the ED model, a centre of symmetry was located for the Pn2<sub>1</sub>a model and the atomic coordinates were transformed to the centrosymmetric space group setting for refinement against the synchrotron PXRD data.

The crystal structure of the higher-hydrate mineral was established by recognising that it had the same polyhedral layers as in bulachite, but with an increased layer separation due to the presence of interlayer water molecules. The x coordinates for the bulachite model were scaled by the ratio of the two *a* axes to establish the polyhedral layers in the a = 19.9 Å cell. The PXRD pattern for this model was calculated and found to have only a fair agreement with the experimental pattern. We examined the possibility that the transformation of the a = 19.9 Å phase to the a = 15.4 Å phase involved not only a contraction of the interlayer spacing but also a sliding of the layers relative to one another, as we found in heating studies on other hydrated aluminium arsenate minerals (Grev et al., 2016b). In space group Pnma, the layers can be displaced relative to one another along [001], with no distortion of the polyhedra, by changing the z coordinate for all atoms by the same amount. We ran a series of tests where the z coordinate for all atoms was changed in increments of 0.01. It was found that changing z in a positive sense improved the fit to the experimental PXRD pattern, and the profile R factors decreased with increasing  $\Delta z$  up to 0.09 then increased with further change in z. The value of  $\Delta z$  was then fixed at 0.09, and difference-Fourier maps were used to locate the interlayer water molecules.

# Structure refinements

The *Pnma* structural model for bulachite was refined both with the synchrotron PXRD data and by dynamical refinement of

 Table 1. Synchrotron PXRD data collection and Rietveld refinement details for bulachite.

Crystal data	
Formula	[Al <sub>6</sub> (AsO <sub>4</sub> ) <sub>3</sub> (OH) <sub>9</sub> (H <sub>2</sub> O) <sub>4</sub> ]·2H <sub>2</sub> O
Unit-cell parameters: a, b, c (Å)	15.3994(3), 17.6598(3), 7.8083(1)
Volume (ų), Z	2123.46(7), 4
Space group	Pnma
Data collection and refinement	
X-ray radiation source, wavelength (Å)	Australian Synchrotron, 0.59028(6)
2θ range, step size (°)	1.50-104.82, 0.00375
Resolution for structure analysis (Å),	1.0, 1158
number of contributing reflections	
Number of profile parameters	8
Number of structural parameters/restraints	63/38
Profile function	Pseudo-Voigt
Background	Interpolation between 40 selected points
Rp, Rwp, GoF, R <sub>F</sub>	3.73, 5.09, 4.29, 10.7

 Table 2. Refined atomic coordinates, isotropic displacement parameters and calculated bond-valence sums (BVS) for bulachite.

Atom x		У	Ζ	B (Å <sup>2</sup> )	BVS 5.22	
As1	0.82245(18)	0.00921(19) 0.3827(4)		0.0368(6)		
As2	0.9057(3)	1/4	0.5824(6)	0.0368(6)	5.24	
Al1	0.6329(6)	0.0696(5)	0.2444(17)	0.0368(6)	3.11	
Al2	0.8397(6)	0.0964(5)	0.7535(16)	0.0368(6)	2.96	
Al3	0.7207(6)	0.1692(6)	0.4971(16)	0.0368(6)	2.70	
01	0.7276(5)	-0.0061(6)	0.2856(14)	0.0186(15)	2.19	
02	0.9002(6)	0.0190(6)	0.2359(12)	0.0186(15)	1.32	
03	0.8160(7)	0.0885(5)	0.5006(13)	0.0186(15)	2.00	
04	0.8459(7)	-0.0642(5)	0.5099(13)	0.0186(15)	1.86	
05	0.8043(5)	1/4	0.5078(18)	0.0186(15)	2.28	
06	0.9752(8)	1/4	0.4186(14)	0.0186(15)	1.32	
07	0.9217(6)	0.17267(14)	0.7019(7)	0.0186(15)	1.82	
OH1	0.7085(9)	0.1522(7)	0.250(2)	0.0186(15)	0.98	
OH2	0.6300(10)	0.0952(9)	0.4819(19)	0.0186(15)	0.98	
OH3	0.9259(8)	0.0208(7)	0.772(3)	0.0186(15)	1.10	
OH4	0.7386(8)	0.1541(7)	0.740(2)	0.0186(15)	1.01	
OH5	0.6390(13)	1/4	0.511(3)	0.0186(15)	1.00	
Ow1	0.5229(9)	0.1135(7)	0.226(3)	0.0186(15)	0.55	
Ow2	0.8415(13)	0.1119(9)	0.9944(19)	0.0186(15)	0.51	
W1	0.93	1/4	0.018	0.074(11)		
W2	0.3652	3/4	0.0187	0.074(11)		

the ED data. The latter will be reported separately. Here we describe the PXRD refinement, using the synchrotron data to a resolution of 1.0 Å (the peaks become very weak and broad at higher resolutions). The Rietveld data collection and refinement conditions are given in Table 1.

The profile parameters were established using a Le Bail structureless fit to the PXRD pattern. Manual backgrounds were automatically located at 40 positions in *JANA2006*. The unit-cell parameters, a zero-shift, the Lorentzian parameters *LX* and *LY*, and a parameter to deal with anisotropic peak broadening for reflections *hkl* with h = 2n+1 were refined, giving Le Bail fit indices of Rp = 2.60, Rwp = 3.73 and Goodness of fit, GoF = 3.13.

The atomic coordinates were refined using soft distance restraints for the As–O and Al–O distances of 1.670(1) Å and 1.900(5) Å, respectively. The restrained distances used are the mean of As–O and Al–O distances obtained from single-crystal refinements of other aluminium arsenate minerals (Grey *et al.*, 2013, 2015, 2016*a*). Angle restraints were also used, for the AsO<sub>4</sub> tetrahedra, O–As–O = 109.47(2)°. Group isotropic



**Fig. 3.** Low-angle region of the synchrotron PXRD pattern for the Cap Garonne specimen. The sharp reflections all index as (0kl) based on cell parameters b = 17.7 and c = 7.8 Å. The first reflection is (200) where a = 19.9 Å.

displacement parameters, *B*, for the metal atoms, polyhedral anions and interlayer water molecule oxygen atoms were refined. Rietveld refinement in *JANA2006* converged at fit indices of Rp = 3.73, Rwp = 5.09 and GoF = 4.29. The refined structural parameters are reported in Table 2. The atom labels are the same as used for other hydrated Al arsenate minerals (Grey *et al.*, 2015, 2016a) where OH = hydroxyl, Ow = coordinated H<sub>2</sub>O and W = H-bonded H<sub>2</sub>O. Polyhedral bond distances, cation–cation distances and likely H-bonded pairs are given in Table 3.

For the higher-hydrate mineral the structural parameters were not refined, apart from establishing the magnitude of layer sliding along [001] as discussed above. The synchrotron PXRD pattern for this mineral (Fig. 3) is of lower quality than that for bulachite. In particular, all reflections with non-zero *h* are broadened and the reflections with *h* odd are broader than those with *h* even. This was dealt with by combining anisotropic Lorentzian particle size broadening (parameter *LXe*) with anisotropic peak broadening for reflections with h = 2n+1. As for bulachite, the PXRD data was truncated at 1 Å resolution, 40 manual backgrounds were located and profile parameters were established with a Le bail fit, giving fit indices of Rp = 2.88, Rwp = 4.06 and GoF = 3.40. The interlayer water molecules were located in difference-Fourier

Table 3. Polyhedral bond distances, cation-cation distances (d in Å) and likely H-bonded pairs in bulachite.

As1-01	1.667(8)	Al2-01	1.918(13)	As1-Al1	3.290(10)	Potential H-bonded pairs			
As1-02	1.667(9)	Al2-03	2.014(16)	As1–Al1	3.223(13)	DonorAcceptor			
As1-03	1.679(9)	Al2-07	1.890(11)	As1–Al2	3.291(12)	Ow1…02	2.55	OH2…O2	2.84
As1-04	1.672(10)	Al2–OH3	1.889(16)	As1–Al2	3.276(10)	Ow1…04	2.77	OH3…O2	2.79
<as1-0></as1-0>	1.671	Al2–OH4	1.864(16)	As1–Al3	3.352(10)	Ow106	2.77	OH4…O4	2.92
		Al2–Ow2	1.900(19)			Ow2…02	2.65	OH4…O7	2.86
As2–05	1.666(10)	<al2-φ></al2-φ>	1.912	As2–Al2	3.190(10)	Ow2…04	2.99	OH5…O2	2.55
As2-06	1.668(12)			As2–Al2	3.190(10)	Ow2…07	2.82	OH5…O4	2.77
As2–07 x2	1.673(5)	Al3–03	2.046(14)	As2–Al3	3.255(10)	W1…07	2.82	OH5…O6	2.77
<as2-0></as2-0>	1.670	Al3-05	1.924(11)	As2–Al3	3.255(10)	W2…O6	2.58		
		Al3–OH1	1.96(2)						
Al1–01	2.005(13)	Al3–OH2	1.917(18)	Al1–Al2	2.962(12)				
Al1-04	1.862(17)	Al3–OH4	1.94(2)	Al1–Al3	2.970(16)				
Al1–OH1	1.868(15)	Al3–OH5	1.905(17)						
Al1–OH2	1.91(2)	<al3-φ></al3-φ>	1.949	Al2–Al3	3.004(15)				
Al1–OH3	1.847(15)								
Al1–Ow1	1.869(16)			Al3–Al3	2.854(14)				
<al1-ф></al1-ф>	1.894								

Notes:  $\Phi$  is 0, OH or Ow.



Fig. 4. (a) Electron diffraction (0kl) reciprocal lattice section for the Cap Garonne specimen. (b) Image of flat lath crystal used to obtain the ED pattern. Museums Victoria specimen M55455.

maps. The agreement factors for the Rietveld fitting of the PXRD data are Rp = 5.57, Rwp = 7.43 and GoF = 6.23. Because the polyhedral layers are built from the refined bulachite structure, the polyhedral bond distances are the same as reported in Table 2 for bulachite. The higher hydrate is a potential new mineral and further crystallochemical details await the preparation of a new mineral proposal to the International Mineralogical Association Commission on New Minerals, Nomenclature and Classification (IMA-CNMNC).

### Results

The low-angle region of the PXRD pattern for the Cap Garonne specimen is shown in Fig. 3. The indexed sharp peaks in the pattern are also present in the bulachite pattern reported by Walenta (1983) but the other peaks, which are much broader, have no counterpart in the bulachite pattern. The sharp reflections all correspond to a single reciprocal lattice section, (0kl). This was the most commonly encountered pattern in ED studies of the sample. The ED (0kl) pattern is shown in Fig. 4a. It is obtained when the



Fig. 5. Low-angle region of synchrotron PXRD patterns for the Cap Garonne specimen heated at temperatures 25, 50, 75 and 100°C. Indexing of the 100°C pattern shown.



**Fig. 6.** Gandolfi PXRD patterns for two separate tufts of the Neubulach type specimen of bulachite (Cu $K\alpha$  radiation).

platy laths lie flat on the TEM grid. Indexing of the ED pattern gives cell parameters b = 17.7 Å and c = 7.8 Å. Systematic absences in the pattern for (0*kl*) reflections with k+l=2n+1 indicate an *n*-glide plane. The associated image of the diffracting crystal is shown in Fig. 4b. It confirms that the direction of the 7.8 Å axis is parallel to the long dimension of the lath. Aided by the ED results the PXRD could be fully indexed and a Le Bail refinement gave the orthorhombic unit-cell parameters a = 19.855(4), b = 17.6933(11) and c = 7.7799(5) Å, with possible space groups *Pnma* or  $Pn2_1a$ .

PXRD patterns corresponding to heating of the Cap Garonne specimen are presented in Fig. 5. At 50°C there is the appearance of a peak at d = 7.8 Å (arrowed), which is the strongest peak in the pattern of bulachite (Walenta, 1983). Note that this peak is present in the room-temperature pattern but it is very weak and broad. At 75°C there is a dramatic change in the pattern with a strong increase in the 7.8 Å peak and a marked diminution in the 10 Å peak and other peaks present in the room-temperature pattern. At 100°C the transformation is complete and all peaks present match those reported by Walenta (1983) for bulachite.

Electron diffraction patterns obtained on the sample heated to 100°C were indexed with unit-cell parameters a = 15.4, b = 17.7 and c = 7.8 Å. Using these as starting parameters in a Le Bail fit to the 100°C PXRD pattern gave the refined unit-cell parameters a = 15.3994(3), b = 17.6598(3) and c = 7.8083(1) Å. The systematic extinctions in the ED patterns correspond to space groups *Pnma* or *Pn*2<sub>1</sub>*a*, the same as for the a = 19.9 Å mineral. As mentioned in the previous section the fit indices were Rp = 2.60, Rwp = 3.73 and GoF = 3.13. For comparison we re-ran the Le Bail fitting to the 100°C dataset starting with the unit-cell parameters reported by Walenta (1983). There was a marked deterioration to the fit, with fit indices Rp = 6.17, Rwp = 10.49 and GoF = 9.76. With the higher resolution of the synchrotron data (Walenta used a Debye-Scherrer camera with FeK $\alpha$ . radiation), serious misfits are evident using the Walenta cell, due to the incorrect *c* parameter.

The possibility existed that the bulachite specimen from Cap Garonne was crystallochemically different from the bulachite from Neubulach. To check this we obtained a portion of the specimen characterised by Walenta (1983). We were able to confirm, by obtaining Gandolfi PXRD patterns on different crystal aggregates, that both the a = 19.9 Å and a = 15.4 Å phases are present in the



Fig. 7. Rietveld observed (red points) and calculated (black line) PXRD patterns for bulachite from Cap Garonne. Short bars show the positions of the Bragg reflections. The difference plot is in blue.



Fig. 8. [001] projection of the layer structure for bulachite. Dotted lines show likely interlayer H-bonds. Dark blue spheres are water molecules, Al-centred octahedra are yellow and AsO<sub>4</sub> tetrahedra are mauve.

Neubulach sample, as shown in Fig. 6. For the 19.9 Å phase, the reflections involving h are very broad, including the strong characteristic (200) reflection at  $d \approx 10$  Å, so small amounts of this phase are difficult to detect. Frau and da Pelo (2001) noted that their PXRD pattern had an anomalously high background suggestive of the presence of amorphous phase and broad peaks. Our PXRD heating experiments (Fig. 5) show that the 19.9 Å phase begins to transform to bulachite within a few minutes at 50°C, so it is likely that at suitable humidity conditions, the transformation could occur at ambient temperatures over long periods. The dehydration of the 19.9 Å phase will start at the surface of the mineral crusts and move inwards under diffusion control, so it is not unexpected to find a mix of the two phases in the crusts. It is possible that the specimen that Walenta used to obtain a PXRD pattern came from the upper region of the surface aggregates where dehydration of the 19.9 Å phase had occurred.

As described in the Experimental section, the crystal structure of bulachite was determined in space group *Pnma* using low-dose electron tomography data and refined using Rietveld treatment of synchrotron PXRD data. The data collection and refinement details are given in Table 1 and the Rietveld fit to the PXRD data is shown in Fig. 7.

Projections of the crystal structures along the 7.8 Å axis are shown in Figs 8 and 9 for bulachite and the higher hydrate, respectively, while Fig. 10 shows a view of the structures normal to the (100) layers. This Figure shows that the polyhedral layers are built from edge-shared Al-centred octahedra that form spiral chains along [001], the same as occurs in liskeardite (Grey *et al.*, 2013). The spirals are joined together along [010] by edge-sharing of octahedra in the mirror plane at  $y = \frac{1}{4}$ . The spirals are decorated with AsO<sub>4</sub> tetrahedra that corner-connect to the AlO<sub>6</sub> octahedra in the same manner as in liskeardite.

The close relationship of the two structures to that of liskeardite is illustrated by comparing the PDFs for the three minerals in



Fig. 9. [001] projection of the layer structure for the higher hydrate mineral. Blue spheres are interlayer water molecules, Al-centred octahedra are yellow and  $AsO_4$  tetrahedra are mauve.



**Fig. 10.** Projection approximately along [100] of the structure of bulachite, showing spirals along [001] of edge-shared Al-centred octahedra (yellow) and corner-connected  $AsO_4$  tetrahedra (mauve).

Fig. 11. The first three strong peaks in the PDFs, at 1.8. 2.7 and 3.2 Å correspond to As–O + Al–O, O–O and As–Al + Al–Al distances, respectively. The O–O at 2.7 Å is strong despite the low scattering of oxygen, because fortuitously the O–O distances in both the AsO<sub>4</sub> tetrahedra and the AlO<sub>6</sub> octahedra are almost the same.

The composition of the layers is readily established from the refinement using bond-valence sums (BVS) to distinguish the different anions. BVS values were calculated using the parameters of Gagné and Hawthorne (2015). The values, in valence units, are given in Table 2. Taking into account the site multiplicities, they show clearly that the layers contain 12 oxygens, 9 hydroxyls and 4 coordinated  $H_2O$ . The O2 and O6 sites, which contain unshared tetrahedral oxygen atoms, have low BVS values of 1.32, while the sites O4 and O7 are also slightly low. However,

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**Fig. 11.** Experimental PDFs for (*a*) the higher hydrate mineral and (*b*) bulachite formed at 100°C, compared with the calculated PDF for liskeardite (*c*).

as shown by the H-bond couples in Table 3, oxygen atoms at these sites are acceptor atoms for several H bonds from OH, Ow and W donor atoms. Taking into account the numbers of O, OH and Ow gives the layer formula  $Al_6(AsO_4)_3(OH)_9(H_2O)_4$ , with Z = 4. The same layer formula applies to the higher hydrate mineral. Adding in the H-bonded  $H_2O$  (=W) gives the overall formula for bulachite of  $[Al_6(AsO_4)_3(OH)_9(H_2O)_4]\cdot 2H_2O$ , while the formula for the higher hydrate is  $[Al_6(AsO_4)_3(OH)_9(H_2O)_4]\cdot 8H_2O$ .

**Supplementary material.** To view supplementary material for this article, please visit https://doi.org/10.1180/mgm.2020.52

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