Article



Twinning and incommensurate modulation in baumoite, $Ba_{0.5}[(UO_2)_3O_8MO_2(OH)_3](H_2O)_{\sim 3}$, the first natural Ba uranyl molybdate

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ABSTRACT

Baumoite, Ba_{0.5}[(UO₂)₃O₈Mo₂(OH)₃](H₂O)_{~3}, is a new mineral found near Radium Hill, South Australia, where it occurs in a granite matrix associated with baryte, metatorbernite, phurcalite and kaolinite. Baumoite forms thin crusts of yellow to orange–yellow tabular to prismatic crystals. The mineral is translucent with a vitreous lustre and pale yellow streak. Crystals are brittle, the fracture is uneven and show one excellent cleavage. The Mohs hardness is ~2½. The calculated density is 4.61 g/cm³. Optically, baumoite crystals are biaxial (–), with $\alpha = 1.716(4)$, $\beta = 1.761(4)$, $\gamma = 1.767(4)$ (white light); and $2V_{calc} = 42.2^{\circ}$. Electron microprobe analyses gave the empirical formula Ba_{0.87}Ca_{0.03}Al_{0.04}U_{2.97}Mo_{2.02}P_{0.03}O₂₂H_{11.99}, based on 22 O atoms per formula unit. The eight strongest lines in the powder X-ray diffraction pattern are [d_{obs} Å (I) (hkl)]: 9.175(39)(121), 7.450(100)(020), 3.554(20)(221), 3.365(31)(004, 202), 3.255(31)(123, 302), 3.209(28)(124), 3.067(33)(303, 222, 322) and 2.977(20)(142). Single-crystal X-ray studies ($R_1 = 5.85\%$ for 1892 main reflections) indicate that baumoite is monoclinic, superspace group X2/m(a0g)0s with $X = (0, \frac{1}{2}, 0, \frac{1}{2})$, with unit-cell parameters: a = 9.8337(3), b = 15.0436(5), c = 14.2055(6) Å, $\beta = 108.978(3)^{\circ}$, V = 1987.25(13) Å³ and Z = 4. The crystal structure is twinned and incommensurately modulated and is based upon sheets of U⁶⁺ and Mo⁶⁺ polyhedra of unique topology. Four independent cationic sites partially occupied by Ba atoms are located between the sheets, together with H₂O molecules.

Keywords: baumoite, new mineral species, barium uranyl molybdate, incommensurately modulated structure, twinning, Radium Hill (Received 6 December 2018; accepted 6 March 2019; Accepted Manuscript online: 12 April 2019; Associate Editor: Ian T. Graham)

Introduction

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The Olary Province of South Australia hosts a diversity of small uranium deposits (Dickinson *et al.*, 1954; Campana and King, 1958). These include Radium Hill, Crocker's Well, Mt Victoria and several smaller prospects. Most of the deposits are low grade and the only significant production has been from Radium Hill which was Australia's first uranium mine, discovered in 1906 and operated until 1961. The deposits are intrusive-type deposits, associated with Mesoproterozoic intrusives, mainly granite, alaskite, pegmatite and migmatite. The primary minerals are davidite-(La) and brannerite and only a small number of secondary uranium minerals have been recorded. Secondary uranium minerals have also been reported from numerous pegmatites and granites in the region and it is one of these that has produced the specimens containing baumoite. Baumoite is one of nine uranyl molybdate minerals and the first natural Ba, uranyl

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molybdate. Its structure is twinned and incommensurately modulated, one of several U^{6+} modulated structures that have been discovered recently along with swamboite-(Nd) and shinkolobweite (Olds *et al.*, 2017; Plášil *et al.*, 2017; Plášil, 2018). Despite the weak intensities of the satellite reflections (only those of the 1st order were detected) on the CCD frames, the refinement in superspace converged to the reasonable values and it provided a reasonable structure model. Baumoite is named for its chemical composition. The mineral and its name have been approved by the Commission on New Minerals, Nomenclature and Classification of the International Mineralogical Association (IMA2017-054, Elliott *et al.*, 2017). The holotype specimen of baumoite is deposited in the mineral collection of the South Australian Museum, Adelaide, South Australia (registration number G34697).

Occurrence

The Olary Domain is contained within the Curnarnona Province that extends across northeastern South Australia and western New South Wales. The geology comprises a late Palaeoproterozoic metasedimentary and metavolcanic succession (Willyama Supergroup)

Fig. 1. Scanning electron microscopy photomicrograph showing crystals of baumoite. The field of view is 100 mm across.

with some meta-intrusives and early Mesoproterozoic volcanics and granitoid intrusives (Stevens, et al., 1990; Flint and Parker, 1993). The lithological and geophysical character of the Willyama Supergroup in the Olary Domain differs from the remainder of the Province with a higher proportion of shallow water sediments, lesser volcanics, widespread metasomatism, particularly albitisation, and an abundance of syntectonic to late-tectonic granitoids (Callen, 1990; Forbes, 1991; Flint and Parker, 1993). Rocks in the Radium Hill area are metasedimentary quartz-feldspathic gneiss, composite gneiss, quartz-feldspar-biotite schist and amphibolite of the Willyama Supergroup. Granitoid intrusives crop-out to the northwest of Radium Hill. Baumoite was found as thin crusts of minute crystals in thin seams on specimens collected in the late 1980s from a weathered granite outcrop, 4 km NW of the Radium Hill mine. Associated minerals are baryte, metatorbernite, phurcalite and kaolinite. The granite comprises pink orthoclase, quartz and muscovite. Baumoite has resulted from the alteration of

baryte and primary U and Mo minerals by oxidising groundwaters. No primary U and Mo minerals have been observed in specimens containing baumoite, however, rocks in the surrounding area contain uraninite, davidite-(La) and molybdenite.

Appearance, physical and optical properties

Baumoite occurs as yellow to orange-yellow tabular to prismatic crystals up to 0.11 mm in size (Fig. 1). Crystals are translucent with a vitreous lustre, pale yellow streak, and show bright yellow green fluorescence under shortwave ultraviolet illumination. The Mohs hardness is $\sim 2\frac{1}{2}$. The density could not be measured as it exceeds that of available heavy liquids; the calculated density is 4.61 g/cm³ from the empirical formula and 4.68 g/cm³ from the ideal formula. Baumoite is brittle, the fracture is uneven and shows one excellent cleavage. Baumoite is optically biaxial (-), $\alpha = 1.716(4)$, $\beta = 1.761(4)$ and $\gamma = 1.767(4)$ measured in white light. Calculated 2V is 42.2°. The Gladstone-Dale compatibility index is 0.009, classed as excellent (Mandarino, 1981).

Infrared spectroscopy

The infrared spectrum (Fig. 2) of powdered baumoite in the range 4000 to 650 cm⁻¹ was obtained using a Nicolet 5700 FTIR spectrometer equipped with a Nicolet Continuum IR microscope and a diamond-anvil cell. The spectrum was interpreted with regard to the theoretical and experimental papers published by Hardcastle and Wachs (1990), Fomichev et al. (1992), Čejka (1999), Fedoseev et al. (2001), Sidorenko et al. (2005), Frost et al. (2008), Maczka et al. (2009), Nakamoto (2009), Zhang and Guo (2012) and Barik et al. (2014). Moreover, we used data taken from Katscher et al. (1990), who reviewed infrared and/or Raman spectra of $(MoO_x)^{n-}$ units in synthetic and mineral phases. Two bands at 3699 (vw) and 3489 (sb) and a shoulder at 3263 cm^{-1} are assigned to the v OH stretching vibrations of hydrogen bonded to water molecules and hydrogen bonded



Fig. 2. Fourier-transform infrared spectrum of powdered baumoite.



Table 1. Compositional data for baumoite.

Constituent	Weight %	Range	Standard deviation
BaO	9.88	8.80-11.38	0.73
CaO	0.12	0.08-0.19	0.03
Al_2O_3	0.14	0.07-0.39	0.07
UO ₃	62.80	60.50-65.61	1.29
MoO ₃	21.56	20.36-22.64	0.67
P_2O_5	0.15	0.06-0.81	0.18
H_2O (calc)*	7.99		
Total	102.64		

*calculated to obtain charge balance.

hydroxyls. According to Libowitzky (1999), approximate O–H···O hydrogen bond lengths vary from >3.2 to 2.73 Å. A medium strong band at 1616 cm⁻¹ is attributed to the v₂ (δ) H₂O bending vibration and weak bands at 1417 (shoulder), 1171 (vvw), 1100 (sh), 1030 (w-m), 1013 (w-m) may be connected with U–OH bending vibrations. A shoulder at 934 cm⁻¹ and a very strong band at 851 cm⁻¹ together with a shoulder at 880 cm⁻¹ are attributed to the v (Mo–O) stretching vibrations (Katscher *et al.*, 1990). However, an overlap (a coincidence) of this vibration with the v₃ (UO₂)²⁺ antisymmetric stretching vibration is assumed. A shoulder at 787 cm⁻¹ is connected with the v₁ (UO₂)²⁺ antisymmetric stretching vibration. According to Bartlett and Cooney (1989), approximate U–O bond lengths in uranyl, inferred from the wavenumbers of the (UO₂)²⁺ stretching vibrations, are 1.82 Å (based on v₃) and 1.82 Å (based on v₁). Observed lengths are somewhat

Table 2. Powder X-ray diffraction data for baumoite.

I _{obs}	/ _{calc} *	$d_{\rm obs}$	d _{calc}	h k l
39	31	9.175	9.269	101
100	100	7.450	7.548	020
10	4	6.707	6.733	101
6	1	5.784	5.853	121
	2		4.966	201
4	1	4.642	4.634	202
4	1	4.084	4.041	122
3	1	3.938	3.949	222
9	11	3.764	3.774	040
17	1	3.705	3.635	041
20	19	3.554	3.559	221
4	1	3.456	3.501	140
31	12, 16	3.365	3.378, 3.367	004,202
31	10, 6	3.255	3.264, 3.279	1 2 3, 3 O ⁻ 2
28	36	3.209	3.232	124
33	11, 24, 11	3.067	3.090, 3.075, 3.008	3 0 3, 2 2 2, 3 2 2
20	17	2.977	2.963	142
1	1	2.826	2.817	203
	1	2.694	2.698	124
2	1	2.587	2.601	225
3	2	2.497	2.512	242
1	1	2.439	2.475	342
1	1	2.283	2.283	125
1	1	2.243	2.249	226
3	1	2.040	2.042	106
2	1	1.993	1.975	444
3	1	1.966	1.967	227
2	1	1.862	1.862	265
1	1	1.806	1.810	206
1	1	1.789	1.792	423
1	1	1.728	1.730	541

*Calculated intensities were obtained using the program *Rietica* (Hunter, 1998). The strongest lines are given in bold higher than the U–O lengths in uranyl ~1.8 Å, proposed by Lussier *et al.* (2016). A strong band at 723 cm⁻¹ is attributable to the v (Mo₂O₂) stretching vibration of the Mo = O₂ = Mo unit bridging two Mo-octahedra and thus forming (Mo₂O₂) units (Katscher *et al.*, 1990).

Chemical composition

The chemical composition of baumoite (20 points on one crystal aggregate) was determined using a Cameca SXFive electron microprobe operating in the wavelength-dispersive mode with an acceleration voltage of 20 kV, beam current of 20 nA, and 5 µm beam diameter. Data were processed using the $\varphi(\text{gz})$ correction procedure of Pouchou and Pichoir (1985). Analytical data (average of 20 points) are given in Table 1. The standards employed were: baryte (Ba), wollastonite (Ca), almandine (Al), U-metal (U), Mo-metal (Mo), and apatite (P). The empirical formula, calculated on the basis of 22 oxygen atoms and with H₂O calculated to obtain charge balance, is Ba_{0.87}Ca_{0.03} Al_{0.04}U_{2.97}Mo_{2.02}P_{0.03}O₂₂H_{11.99}. The simplified formula is BaU₃ Mo₂O₁₆(H₂O)₆.

X-ray crystallography

Powder X-ray diffraction

Powder X-ray studies (Table 2) were carried-out using a Rigaku Hiflux Homelab diffractometer, with monochromatised CuK α radiation ($\lambda = 1.541870$ Å). Unit-cell parameters refined using the Le Bail profile-fitting method (Le Bail *et al.*, 1988; Hunter, 1998) and starting from the unit-cell parameters determined from the single-crystal study (see below), are a = 9.921(1), b = 15.080(1), c = 14.295(2) Å, $\beta = 109.279(5)^{\circ}$ and V = 2018.7(3) Å³.

Single-crystal X-ray diffraction

The single-crystal X-ray study was carried-out using intensity data collected at 293 K on a Rigaku SuperNova diffractometer (MoKa radiation from the microfocus X-ray tube, $\beta = 0.71070$ Å) equipped with an Atlas S2 CCD detector using a crystal $0.065 \text{ mm} \times 0.045 \text{ mm} \times 0.01 \text{ mm}$ in size. Initial attempts to solve and refine the structure based on the preliminary data from an Oxford Diffraction Xcalibur E diffractometer (MoKa radiation from the conventional X-ray tube, $\lambda = 0.71070$ Å) equipped with an EoS CCD were done in the monoclinic space group C2/m, but this refinement returned only poor R values $(R_1 \approx 0.10 \text{ with a significantly large GoF} \approx 4)$, thus representing an average structure. By the careful inspection of the diffraction data it was found that the structure of baumoite is twinned and incommensurately modulated. The satellite reflections can be described by the modulation vector $\mathbf{q} = 0.718\mathbf{a}^* + 0.2803\mathbf{c}^*$. Twin domains are related by the two-fold rotation along [001] axis represented by the twinning matrix $\begin{bmatrix} -1 & 0 & -0.451/0 & -1 & 0/0 \end{bmatrix}$ 0 1], which leads to a partial overlap of the reflections. Corrections for background, Lorentz, and polarisation effects were applied to the data during reduction in the CrysAlis package (Rigaku, 2018). The final data set was integrated as an hklm file (for the commensurate 3 + 1 case) using CrysAlis based on the list of reflections generated in Jana2006 (Petříček et al., 2014). A correction for absorption was done using spherical harmonics by the Jana2006 program. Refinement of the modulated structure in the superspace group X2/m(a0g)0s with $X = (0,\frac{1}{2},0,\frac{1}{2})$ was started using the atom coordinates obtained from the average



Fig. 3. Bond distances around U1, U2 and U3 sites in the incommensurately modulated structure of baumoite as a function of the internal coordinate t.



Fig. 4. Bond distances around the Mo1 and Mo2 sites in the incommensurately modulated structure of baumoite as a function of the internal coordinate t.

structure solved by *Superflip* (Palatinus and Chapuis, 2007). One modulation wave was set for the refinement. The modulations in the structure of baumoite were described using both continuous (harmonics) and discontinuous (crenel-like; Petříček *et al.*, 2016) modulation functions. There are strong positional

modulations of the U, Mo and O atoms that are linked to previously mentioned atoms and a strong occupational modulation of the Ba atoms and also less positional modulation of the Ba atoms and O atoms of the H_2O sites. We have used the recently developed refinement technique in which the refinement is repeated



Fig. 5. Bond distances around the Ba sites in the incommensurately modulated structure of baumoite as a function of the internal coordinate t. The bluish region highlights the range of the most typical Ba–O distances in the crystalline solids.



Fig. 6. Refined occupancies of the Ba sites in the incommensurately modulated structure of baumoite as a function of the internal coordinate t.

Table 3. Crystallographic and reinement parameters for baumor	nement parameters for baumoite	refinement	and	Crystallographic	3.	Table
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Crystal data		
Chemical formula	Ba _{0.56} U ₃ Mo ₂ O _{19.72} H _{8.4}	
Temperature (K)	293	
Cell setting	Monoclinic	
Superspace group	$X2/m(\alpha 0\gamma)0s$	
a, b, c (Å)	9.8337(3), 15.0436(5), 14.2055(6	
β (°)	108.978(3)	
V (Å ³)	1987.25(13)	
Ζ	4	
Modulation wave vector	q = 0.718(4) a *+ 0.280(2) c *	
Calculated density (g cm ⁻³)	4.61	
$\mu \ (mm^{-1})$	26.777	
Data collection		
Diffractometer	Rigaku SuperNova with Atlas S2 CCD	
Radiation type, wavelength (Å)	ΜοΚα, 0.71073	
No. of measured, independent and observed	41859, 6953, 4521	
$[l > 2\sigma(l)]$ reflections		
Range of <i>h, k, l, m</i>	$-13 \rightarrow h \rightarrow 13$	
	$-19 \rightarrow k \rightarrow 19$	
	$-18 \rightarrow l \rightarrow 18$	
	$-1 \rightarrow m \rightarrow 1$	
T _{min} , T _{max}	0.462, 1	
No. of main reflections	2442	
No. of 1 st order satellite reflections	4511	
Criterion for observed reflections	l > 3σ(l)	
R _{int}	0.064	
Refinement		
Refinement on	F	
R _{obs} , wR _{obs} (all reflections)	0.0702, 0.0771	
R _{obs} , wR _{obs} (main reflections)	0.0585, 0.0645	
R_{obs} , wR_{obs} (satellites) 1 st order	0.0855, 0.0936	
S _{obs} /all	2.71/2.33	
No. of parameters	252	
$\Delta \rho_{\rm max}, \Delta \rho_{\rm min} \ (e^- \ A^{-s})$	9.51, -5.57	
Twin fraction	0.950(4)/0.050(4)	
Weights	$1/(\sigma^2(F) + 0.0001F^2)$	

several times starting from small randomly chosen modulation amplitudes. Crystallographic details, data collection and refinement parameters for the average and incommensurate structures, respectively, are given in Table 3. Fractional coordinates and atom displacement parameters, Fourier amplitudes of the occupational modulation are given in the accompanying crystallographic information files, which have been deposited with the Principal Editor of *Mineralogical Magazine* and are available as Supplementary material (see below). Selected interatomic distances as functions of *t* are displayed on Figs 3, 4 and 5; refined occupancies of Ba sites are displayed on Fig. 6.

Results and discussion

Description of the structure

The structure of baumoite contains three symmetrically distinct U sites, two Mo sites, four Ba sites and 22 O sites. All U sites in the structure are [7]-fold coordinated in the form of a pentagonal bipyramid, consisting of two strong U–O bonds, i.e. forming uranyl ion $(UO_2)^{2+}$, and five weaker bonds distributed in the equatorial plane of the pentagonal bipyramid (Fig. 3). All U in baumoite is present as hexavalent. The two Mo sites are [6]-coordinated in the form of irregular, distorted octahedra (Fig. 4). The Mo1 and Mo2 polyhedra share their common edge to form $MO_2\varphi_{10}$ dimers. The baumoite structure is based upon uranyl-molybdate sheets. The fundamental building blocks of the sheets are trimers of edge-sharing U1, U2 and U3 polyhedra,



Fig. 7. Approximant structure of baumoite at t=0. Uranyl-molybdate sheets, composed of uranyl pentagonal bipyramids (yellow) and Mo⁶⁺-octahedra (red), are alternating interlayer, where Ba atoms (green) and H₂O molecules (Ow1-4) are localised.

which further share a common edge with other symmetrically related U3 polyhedra to form the six-membered clusters (Figs 7, 8*a*). These clusters are connected through the O11 equatorial atom of the U1 bipyramid to form irregular chains propagated along [$\overline{101}$]. These chains of polyhedra are connected through Mo₂ φ_{10} dimers in a stair-case-like way (Fig. 8*a*).

Adjacent sheets are separated from each other at an interplanar distance of 7.5 Å (corresponding to half of the **b**-cell parameter). Between the sheets four symmetrically independent Ba atoms and four symmetrically independent H₂O molecules are localised. All the Ba atoms are strongly occupationally and positionally modulated (Figs 5 and 6). The O sites of the corresponding H₂O molecules are both occupationally and positionally modulated; the amplitude of their modulation depends on the presence of the corresponding Ba (to which they are coordinated) or presence of the under-and above-laying U-Mo sheet fragments.

Baumoite topology

Sheets in the structure of baumoite are topologically unique, when considering an approximant, idealised structure (Fig. 7). Another structure with similar high U:Mo ratio is a synthetic phase with composition Ag₁₀[(UO₂)₈O₈(Mo₅O₂₀)] (Krivovichev and Burns, 2003); however, in that phase there are infinite complex chains of edge-sharing pentagonal uranyl bipyramids which are linked through Mo₂O₇ dimers, as a part of Mo₅O₂₀ interconnecting adjacent sheets (with interplanar spacing ≈ 6.5 Å) (Fig. 8*c*). Another structure containing similar sheets with isolated dimers of Mo-octahedra (within the sheets) is that of synthetic K₂Na₈(UO₂)₈Mo₄O₂₄[(S, Mo)O₄] (Krivovichev, 2014). All the mentioned molybdates are monoclinic.

Incommensurate modulation in baumoite most probably results from long-range positional ordering of Ba, U and Mo atoms. The nature of the baumoite structure resembles the behaviour of composite structures, known for sulfosalts (Petříček *et al.*,



Fig. 8. Structural sheets in uranyl molybdates. (*a*) Uranyl molybdate sheet in baumoite at t = 0. UO₇ bipyramids (yellow) share edges and vertices to form irregular infinite ribbons interconnected through dimers of Mo polyhedra (red), running through as stairs-like (represented by a dashed line). (*b*) Corresponding representation of the baumoite uranyl-anion topology. (*c*) Sheet in synthetic Ag₁₀[(UO₂)₈O₈(Mo₅O₂₀)] (Krivovichev and Burns, 2003).

1991; Makovicky *et al.*, 2011; Jaszczak *et al.*, 2016). Such structures are based on two, interpenetrating lattice-periodic structures that are mutually incommensurate. Nevertheless, in the case of baumoite, it was not possible to check for such a possibility by the refinement due to limited quality of the data (only satellites of the first order were observed).

The idealised formula of baumoite inferred from the results of the structure refinement and bond-valence considerations is $Ba_{0.5}[(UO_2)_3O_8MO_2(OH)_3](H_2O)_{\sim 2.72}$ with Z = 4. However, from the refinement, it is clear that the picture is much more complex, as some of the O atoms within the equatorial plane of the U polyhedra are in fact protonated (as OH) and some of them can also be H₂O in some regions of the superspace. This corresponds to the regions where Ba positions are not fully occupied, therefore the structure sheets tend to be electroneutral with an approximate composition $[(UO_2)_3O_6MO_2(OH)_4(H_2O)_{\sim 1.0}]^0$.

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