*The Canadian Mineralogist* Vol. 44, pp. 1197-1205 (2006)

# REFINEMENT OF THE CRYSTAL STRUCTURE OF BILLIETITE, Ba [(UO<sub>2</sub>)<sub>6</sub> O<sub>4</sub> (OH)<sub>6</sub>] (H<sub>2</sub>O)<sub>8</sub>

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# Abstract

The crystal structure of billietite, Ba [(UO<sub>2</sub>)<sub>6</sub> O<sub>4</sub> (OH)<sub>6</sub>] (H<sub>2</sub>O)<sub>8</sub>, orthorhombic, space group *Pbn*<sub>2</sub>, *a* 12.0941(8), *b* 30.211(2), *c* 7.1563(5) Å, *V* 2614.7(4) Å<sup>3</sup>, *Z* = 4, has been refined to an *R*<sub>1</sub> index of 3.4% for 4918 unique observed ( $|F_0| > 4\sigma F$ ) reflections measured with MoK $\alpha$  X-radiation and a diffractometer fitted with a CCD detector. The structure consists of anionic [(UO<sub>2</sub>)<sub>6</sub> O<sub>4</sub> (OH)<sub>6</sub>]<sup>2-</sup> sheets, linked by interstitial Ba atoms and (H<sub>2</sub>O) groups. There are two crystallographically distinct structural sheets and eight interlayer (H<sub>2</sub>O) groups in billietite. Four (H<sub>2</sub>O) groups are bonded to the interlayer Ba atoms, and the remaining four (H<sub>2</sub>O) groups are not bonded to a cation, but are held in the structure by a network of hydrogen bonds.

Keywords: billietite, crystal structure, hydrogen bonding, uranyl oxide hydrate of Ba, uranium.

#### Sommaire

Nous avons affiné la structure cristalline de la billietite, Ba [(UO<sub>2</sub>)<sub>6</sub> O<sub>4</sub> (OH)<sub>6</sub>] (H<sub>2</sub>O)<sub>8</sub>, orthorhombique, groupe spatial *Pbn*2<sub>1</sub>, *a* 12.0941(8), *b* 30.211(2), *c* 7.1563(5) Å, *V* 2614.7(4) Å<sup>3</sup>, *Z* = 4, jusqu'à un résidu *R*<sub>1</sub> de 3.4% en utilisant 4918 réflexions uniques observées ( $|F_o| > 4\sigma F$ ) et mesurées avec rayonnement MoK $\alpha$  et un diffractomètre muni d'un détecteur CCD. La structure est faite de feuillets anioniques [(UO<sub>2</sub>)<sub>6</sub> O<sub>4</sub> (OH)<sub>6</sub>]<sup>2–</sup> interconnectés par des atomes de Ba et des groupes H<sub>2</sub>O interstitiels. Il y a deux sortes de feuillets, cristallographiquement distincts, et huit groupes H<sub>2</sub>O interfoliaires. Quatre des groupes H<sub>2</sub>O sont liés à des atomes de Ba entre les couches, et les quatre autres ne sont pas liés à un cation, mais font partie de la structure grâce à un réseau de liaisons hydrogène.

(Traduit par la Rédaction)

Mots-clés: billietite, structure cristalline, liaisons hydrogène, oxyde hydraté uranylé de Ba, uranium.

#### INTRODUCTION

Uranyl oxide hydrate minerals are common constituents of the oxidized parts of uranium deposits (Frondel 1958, Finch & Ewing 1992), as well as soils contaminated by actinides (Roh *et al.* 2000, Yamakawa & Traina 2001). Uranyl phases form by alteration of both  $UO_2$  and spent nuclear fuel under conditions similar to those expected in the proposed geological repository at Yucca Mountain, Nevada (Finch & Ewing 1991, Forsyth & Werme 1992, Wronkiewicz *et al.* 1992, 1996, Johnson & Werme 1994, Finn *et al.* 1996). Studies of mineral

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occurrences can be used to test the extrapolation of results of short-term experiments to periods relevant to nuclear-waste disposal (Ewing 1993) and to assess models used to predict the long-term behavior of spent nuclear fuel in a geological repository (Bruno *et al.* 1995).

Billietite contains Ba, isotopes of which can occur as fission products in spent nuclear fuel. A phase with a structure similar to that of billietite, but that contains Cs, Ba, and Mo in addition to U, was reported in an investigation of oxidative corrosion of spent nuclear fuel in groundwater (Buck et al. 1998). Burns et al. (1997a) argued that uranyl phases, including uranyl oxide hydrates, that form owing to alteration of nuclear waste may incorporate various radionuclides in addition to uranium, thereby affecting radionuclide mobility. Uranyl oxide hydrates containing Sr and Cs have been documented (Hill & Burns 1999, Cahill & Burns 2000, Burns & Li 2002), and incorporation of trace levels of Np into the sodium analogue of the uranyl oxide hydrate compreignacite has recently been reported (Burns et al. 2004).

The crystal structures of most of the uranyl oxide hydrate minerals are now known: the structures of schoepite, metaschoepite, masuyite, protasite, becquerelite, compreignacite, curite, fourmarierite, wölsendorfite, vandendriesscheite, agrinierite and uranosphaerite have been reported (Hughes et al. 2003, and references therein). The structure of billietite was determined and refined to an agreement index (R) of 13.9% by Pagoaga et al. (1987) and is similar to the structures of becquerelite, protasite, compreignacite, masuyite, richetite, and agrinierite, each of which contain sheets of uranyl pentagonal bipyramids based upon the α-U<sub>3</sub>O<sub>8</sub> anion topology (Burns 1999). Pagoaga et al. (1987) located only four (H<sub>2</sub>O) groups pfu (per formula unit), rather than eight as indicated by previous and subsequent studies (Brasseur 1949, Frondel & Cuttita 1953, Christ & Clark 1960, Čejka et al. 1998). Because of the importance of (H<sub>2</sub>O) to the structural stability and bonding in uranyl minerals (Finch et al. 1996, 1998, Finch 1997), we have redetermined the crystal structure of billietite in order to establish the role of the interlayer (H<sub>2</sub>O) groups and to derive the scheme of interlayer hydrogen-bonding.

#### EXPERIMENTAL METHODS

The specimen containing billietite was obtained from Mr. Forrest Cureton and is from Shaba, Democratic Republic of Congo. Several crystals were removed from the specimen and examined in crosspolarized light. Most crystals examined show signs of twinning; an untwinned crystal with sharp extinction and uniform optical properties was selected for study. The crystal was mounted on a Bruker PLATFORM threecircle goniometer equipped with a 1K SMART CCD (charge-coupled device) detector and a crystal-todetector distance of 5 cm.

The data were collected using monochromatic MoK $\alpha$  X-radiation and frame widths of 0.3° in  $\omega$ , with 20 s used to acquire each frame. A complete sphere of three-dimensional data was collected to  $\sim 57^{\circ} 2\theta$ . The final unit-cell dimensions (Table 1) were refined on the basis of 6558 reflections using least-squares techniques. Comparison of the intensities of equivalent reflections collected at different times during the data collection showed no evidence of significant change, and there was no streaking present in the diffraction pattern (cf. Pagoaga et al. 1987). The data were reduced and corrected for Lorentz, polarization and background effects using the Bruker program SAINT. An empirical absorption-correction was done on the basis of 3724 intense reflections; the crystal was modeled as a {010} plate, and reflections with a plate-glancing angle of less than 3° were discarded from the dataset, which lowered R(azimuthal) from 31.8 to 6.2%. A total of 31,321 reflections was collected, of which 6814 were omitted because of a plate-glancing angle less than 3°. Merging of equivalent reflections gave 6104 unique reflections, with 4918 classed as observed  $(|F_0| \ge 4\sigma_F)$ .

#### STRUCTURE REFINEMENT

Scattering curves for neutral atoms, together with anomalous dispersion corrections, were taken from the *International Tables for X-Ray Crystallography*, *Vol. IV* (Wilson 1992). The Bruker SHELXTL Version 5 system of programs was used for refinement of the crystal structure.

Refinement of the structure was initiated in space group  $Pbn2_1$  using the structure reported by Pagoaga *et al.* (1987) as the starting model. Following refinement of the positional parameters and isotropic-displacement parameters for all atoms, the agreement index  $R_1$  was 6.3% for the observed reflections. Conversion of the cation-displacement parameters to an anisotropic form, together with refinement of the entire model, resulted in an  $R_1$  index of 5.0%. Inspection of difference-Fourier maps at this stage revealed seven symmetrically distinct (H<sub>2</sub>O) groups that were not present in the model of

TABLE 1. INFORMATION PERTAINING TO THE STRUCTURE REFINEMENT OF BILLIETITE

a (Å)	12.0941(8)	Crystal size (mm)	0.36 x 0.18 x 0.015
b (Å)	30.211(2)	Total reflections	24507
c (Å)	7.1563(5)	Unique reflections	6104
V (Å <sup>3</sup> )	2614.7(4)	R <sub>int</sub> (%)	6.2
Space group	Pbn2 <sub>1</sub>	Unique $ F_{o}  \ge 4\sigma_{F}$	4918
F(000)	3480	Final <i>R</i> <sub>1</sub> (%)	3.4
µ (mm⁻¹)	38.62	S	1.03
$D_{calc}$ (g/cm <sup>3</sup> )	5.253	Flack parameter	0.06(1)

Unit-cell contents: Ba [(UO2)6 O4 (OH)6] (H2O)9

Pagoaga et al. (1987); insertion of the corresponding O-atoms into the model, together with full-matrix leastsquares refinement of all variable parameters, resulted in a final  $R_1$  index of 3.4% for the 4918 unique observed  $(|F_0| \ge 4\sigma_F)$  reflections, and a goodness-of-fit (S) of 1.03. A model including anisotropic displacement of the anions was tried, but it did not lower the final  $R_1$ index, and refinement of site occupancies of the (H<sub>2</sub>O) groups did not produce significant shifts from full occupancy. In the final cycle of refinement, the average parameter shift/esd was 0.000, and the maximum peaks in the final difference-Fourier maps were 1.6 and -2.8  $e/Å^3$ . The final positional and anisotropic-displacement parameters are given in Tables 2 and 3, and selected interatomic distances and angles are given in Table 4. A bond-valence table, calculated with the universal U-O curve of Burns et al. (1997b) and the curves of Brown & Altermatt (1985) for Ba, is given in Table 5. Structure factors may be obtained from the Depository of Unpublished Data, CISTI, National Research Council of Canada, Ottawa, Ontario K1A 0S2, Canada.

# DESCRIPTION OF THE STRUCTURE

#### Cation coordination

There are six symmetrically distinct U sites, all occupying general positions in space group  $Pbn2_1$ .

TABLE 2. COORDINATES AND EQUIVALENT ISOTROPIC-DISPLACEMENT
PARAMETERS OF ATOMS IN THE STRUCTURE OF BILLIETITE

	×	У	z	U <sub>eq</sub>
<i>U</i> (1)	0.55842(4)	0.74671(2)	0.00092(7)	0.0110(1)
U(2)	0.74723(3)	0.73833(2)	0.5391(1)	0.0116(1)
U(3)	0.93753(4)	0.74265(2)	0.99884(7)	0.0108(1)
U(4)	0.55865(4)	0.00466(2)	0.01958(6)	0.0113(1)
U(5)	0.94118(4)	0.00557(2)	0.01401(7)	0.0115(1)
U(6)	0.74929(3)	0.01411(2)	0.4481(1)	0.0113(1)
Ba	0.63786(6)	0.87627(2)	0.3155(2)	0.0206(2)
O(1)	0.5213(7)	0.6903(3)	0.952(1)	0.0200(19)
O(2)	0.5990(6)	0.8024(3)	0.058(1)	0.0176(18)
O(3)	0.7431(6)	0.6804(3)	0.599(1)	0.0183(19)
O(4)	0.7464(6)	0.7936(3)	0.445(1)	0.0171(17)
O(5)	0.9753(6)	0.6866(3)	0.952(1)	0.0169(18)
O(6)	0.8993(6)	0.7983(3)	0.052(1)	0.0202(19)
O(7)	0.6062(7)	0.9477(3)	0.032(1)	0.0232(19)
O(8)	0.5115(7)	0.0616(3)	0.011(1)	0.0166(17)
O(9)	0.8965(7)	0.9494(3)	0.980(1)	0.0230(22)
O(10)	0.9835(7)	0.0624(3)	0.039(1)	0.0190(19)
O(11)	0.7381(6)	0.9538(3)	0.473(1)	0.0194(22)
O(12)	0.7547(6)	0.0733(3)	0.432(1)	0.0116(16)
OH(13)	0.6016(6)	0.7238(2)	0.322(1)	0.0139(16)
O(14)	0.5935(7)	0.7558(3)	0.691(1)	0.0157(19)
O(15)	0.9038(7)	0.7496(3)	0.688(1)	0.0107(19)
OH(16)	0.8912(6)	0.7211(2)	0.322(1)	0.0139(16)
OH(17)	0.7464(6)	0.7244(3)	0.968(1)	0.0184(21)
O(18)	0.5779(6)	0.0136(2)	0.326(1)	0.0166(17)
OH(19)	0.6147(6)	0.0210(3)	0.697(1)	0.0137(18)
OH(20)	0.8880(6)	0.0229(3)	0.691(1)	0.0128(18)
O(21)	0.9161(6)	0.0043(3)	0.325(1)	0.0205(18)
OH(22)	0.7492(6)	0.0272(3)	0.076(1)	0.0152(18)
OW(23)	0.7739(7)	0.8733(4)	0.995(2)	0.0302(21)
<i>OW(</i> 24)	0.9220(7)	0.6312(3)	0.314(2)	0.0319(22)
OW(25)	0.4129(8)	0.8883(3)	0.202(1)	0.0327(24)
OW(26)	0.9179(8)	0.1144(4)	0.690(1)	0.0382(26)
OW(27)	0.5645(8)	0.6345(3)	0.300(2)	0.0411(26)
OW(28)	0.7419(8)	0.6249(5)	0.025(2)	0.0539(34)
OW(29)	0.634(1)	0.8602(4)	0.696(1)	0.0523(30)
<i>OW</i> (30)	0.864(1)	0.8755(3)	0.390(1)	0.0452(32)

Five of the six U sites are surrounded by seven anion sites in pentagonal-bipyramidal arrangements. Each of these  $U\phi_7$  polyhedra [ $\phi$ : (OH) and O] consists of two apical anions at distances of 1.78–1.83 Å from the central U atom, forming the (U<sup>6+</sup>O<sub>2</sub>)<sup>2+</sup> uranyl cation, and five equatorial anions [(OH) and O] at distances in the range 2.10–2.69 Å. The U(2) site is surrounded by six anion sites arranged at the vertices of a distorted octahedron. There are two uranyl O-atoms at distances

TABLE 3. ANISOTROPIC-DISPLACEMENT PARAMETERS FOR THE CATIONS IN THE STRUCTURE OF BILLIETITE

	<i>U</i> <sub>11</sub>	$U_{22}$	U <sub>33</sub>	U <sub>12</sub>	U <sub>13</sub>	U <sub>23</sub>
U(1) U(2) U(3) U(4) U(5) U(6) Ba	0.0093(2) 0.0079(2) 0.0083(2) 0.0091(2) 0.0093(2) 0.0067(2) 0.0185(3)	0.0153(2) 0.0154(3) 0.0156(2) 0.0172(2) 0.0173(2) 0.0166(3) 0.0213(4)		-0.0001(2) 0.0006(2) -0.0004(2) 0.0003(2) 0.0002(2)	0.0004(2)	0.0006(2) 0.0005(3) 0.0004(2) 0.0005(2) 0.0006(2)

TABLE 4. SELECTED INTERATOMIC DISTANCES (Å) AND ANGLES (°) IN BILLIETITE

$\begin{array}{l} U(1)-O(1)a\\ U(1)-O(2)\\ U(1)-O(14)a\\ U(1)-O(15)b\\ U(1)-OH(17)a\\ U(1)-OH(13)\\ U(1)-OH(16)b\\ \\ \\ \\ \end{array}$	1.795(9) 1.799(8) 2.276(8) 2.300(8) 2.382(8) 2.46(1) 2.583(8) 1.797 2.400	$\begin{array}{c} U(4){-}O(8)\\ U(4){-}O(7)e\\ U(4){-}O(7)e\\ U(4){-}O(18)f\\ U(4){-}O(18)f\\ U(4){-}OH(22)\\ U(4){-}OH(19)a\\ U(4){-}OH(19)f\\ \leq U(4){-}O_{W}^{2} \\ \leq U(4){-}O_{W}^{2} \end{array}$	1.813(8) 1.816(9) 2.220(9) 2.225(8) 2.437(7) 2.456(7) 2.571(8) 1.814 2.382
U(2)-O(4) U(2)-O(3) U(2)-O(15) U(2)-O(14) U(2)-OH(13) U(2)-OH(16) U(2)-O(17)	1.802(9) 1.803(9) 2.198(8) 2.217(8) 2.387(9) 2.391(8) 1.802 2.458 3.097(9)	$\begin{array}{l} U(5){-}O(9)g\\ U(5){-}O(10)\\ U(5){-}O(21)h\\ U(5){-}O(22)\\ U(5){-}O(22)\\ U(5){-}O(22)a\\ U(5){-}O(20)h\\ U(5){-}O_{10}b\\ U($	1.798(9) 1.799(9) 2.214(9) 2.24(1) 2.451(7) 2.455(7) 2.572(8) 1.798 2.386
(U(3)O(6)c U(3)O(5) U(3)O(14)d U(3)O(14)d U(3)OH(17) U(3)OH(16)c U(3)-OH(13)d <u(3)-oh(13)d <u(3)-q<sub>M&gt;</u(3)-q<sub></u(3)-oh(13)d 	1.783(8) 1.786(8) 2.274(7) 2.335(8) 2.387(8) 2.47(1) 2.561(8) 1.784 2.405	$\begin{array}{l} U(6){-}O(12)\\ U(6){-}O(11)e\\ U(6){-}O(21)\\ U(6){-}O(18)\\ U(6){-}O(19)\\ U(6){-}O(120)\\ U(6){-}O(22)\\ U(6){-}O(22)\\ < U(6){-}O_{U^{>}}\\ < U(6){-}\phi_{eq} \end{array} \right\}$	1.793(8) 1.83(1) 2.222(8) 2.422(7) 2.422(7) 2.431(7) 2.693(8) 1.811 2.404
Ba-OW(29) Ba-OW(23)a Ba-OW(23)a Ba-O(11) Ba-O(5)b Ba-O(2) Ba-O(2) Ba-O(2) Ba-O(4) Ba-O(4) Ba-O(7) <ba-φ></ba-φ>	2.77(1) 2.79(1) 2.82(1) 2.87(1) 2.902(8) 2.931(8) 2.931(8) 2.957(8) 2.968(9) 2.968(8) 2.885	O(1)a-U(1)-O(2) O(8)-U(4)-O(7)e O(4)-U(2)-O(3) O(9)g-U(5)-O(10) O(6)c-U(3)-O(5) O(12)-U(6)-O(11)e	177.5(4) 179.0(4) 171.6(4) 177.7(3) 178.6(4) 177.1(3)

a: x, y, z - 1; b: x - ½, 1½ - y, z - ½; c: x, y, z + 1; d: x + ½; 1½ - y, z + ½; e: x, y = 1, z; f: 1 - x, -y, z - ½; g: x, y - 1, z - 1; h: 2 - x, -y, z - ½; i: 1 - x, 1 - y, z + ½;

of 1.80 Å and four meridional O and (OH) anions at distances in the range 2.20–2.39 Å. Calculation of the incident valence at the U sites using the coordination-

TABLE 5. BOND-VALENCE ANALYSIS\* FOR THE STRUCTURE OF BILLIETITE

	<i>U</i> (1)	U(2)	U(3)	<i>U</i> (4)	U(5)	<i>U</i> (6)	Ba	Σ	
O(1) O(2)	1.63 1.62	1.63					0.18	1.63 1.80 1.63	
O(3) O(4) O(5) O(6)		1.63	1.66 1.67				0.16 0.19	1.78 1.85 1.67	
O(7) O(8) O(9)			1.07	1.57 1.58	1.62		0.15 0.16	1.72 1.74 1.62	
O(10) O(11) O(12) OH(13)	0.44	0.57	0.37		1.62	1.52 1.64	0.21	1.62 1.73 1.64 1.38	
O(14) O(15) OH(16)	0.64 0.61 0.35	0.37 0.77 0.8 0.56	0.57 0.64 0.44					1.98 2.05 1.35	
OH(17) O(18)	0.52		0.51	0.71		0.67		1.03 2.08	
OH(19)				0.45		0.48		1.29	
OH(20)				0.00	0.45 0.36	0.47		1.28	
O(21)					0.72	0.71		2.11	
OH(22) OW(23) OW(24)					0.46	0.68		0.24	1.19 0.24 0
OW(25) OW(26) OW(27)							0.21	0.21 0 0	
OW(28) OW(29) OW(30)							0.28 0.26	0 0.28 0.26	
Σ	5.80	5.96	5.85	5.82	5.89	5.76	2.04		

\* calculated using the parameters of Brown & Altermatt (1985) for Ba<sup>2+</sup> and of Burns et al. (1997b) for U<sup>6+</sup>, and given in vu (valence units).

independent parameters for the U–O bond (Burns *et al.* 1997b) indicates that U is in the hexavalent state at all sites (as indicated also by the presence of uranyl anions around all U sites). Calculation of the incident valence at the U(2) site using the parameters for the  $U_r$ – $\phi_4$  (<sup>[6]</sup>U– $\phi$ ) bond (Burns *et al.* 1997b) gave an incident bond-valence of 5.97 vu (valence units), indicating that a coordination number of [6] is appropriate at the U(2) site. The structure of billietite shows distinct pseudosymmetry in which the cations at the U(2) and U(6) sites are (pseudosymmetrically) related. Examination of Table 4 shows that the closest six U– $\phi$  distances involving each of these sites are similar, but the seventhlongest distances are very different: U(2)–OH(17) = 3.097(9) Å, whereas U(6)–O(22) = 2.693(8) Å.

The Ba atom is coordinated by six uranyl-O atoms at distances from 2.77 to 2.99 Å, and four (H<sub>2</sub>O) groups at distances from 2.77 to 2.86 Å. The Ba coordination (Table 4) is similar to that in protasite (Pagoaga *et al.* 1987) and guilleminite (Cooper & Hawthorne 1995).

# Bond topology

There are two symmetrically and stereochemically distinct sheets of uranyl polyhedra in the crystal structure of billietite (Fig. 1). The first sheet (Fig. 1a) consists of three uranyl polyhedra, U(1), U(2) and U(3), that share edges and corners to form a sheet identical to that in ianthinite (Burns *et al.* 1997c) and wyartite (Burns & Finch 1999). Chains of edge-sharing U(1) and U(3) pentagonal bipyramids extend in the *c* direction and are linked by sharing corners and by sharing edges with U(2) octahedra. In terms of the chain types used by Miller *et al.* (1996) to describe sheet-anion patterns, this arrangement forms a PU'PU'...sequence, where U'

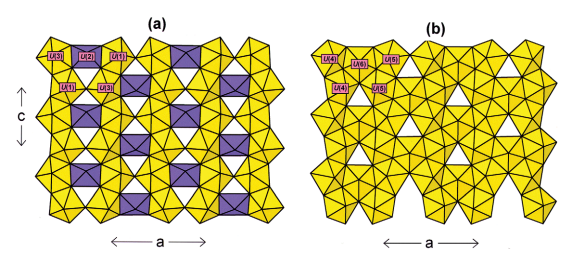


FIG. 1. The structural sheets of uranyl polyhedra in billietite projected along the *b* axis; (a) sheet of U(1), U(2) and U(3) polyhedra; (b) sheet of U(4), U(5), and U(6) polyhedra; U(2): mauve; other *U* polyhedra: yellow.

denotes a modification of the U chain type (Miller et al. 1996). The second sheet (Fig. 1b) also consists of three uranyl polyhedra, U(4), U(5) and U(6), that share edges and corners to form a sheet identical to that in protasite. becquerelite and compreignacite (Burns 1998). Separate chains of edge-sharing U(4) pentagonal bipyramids and U(6) pentagonal bipyramids extend in the c direction and are linked by sharing corners with each other and through the sharing of edges with U(6) pentagonal bipyramids, forming a PUPU... sequence (Miller et al. 1996). The (OH) groups in the second sheet (Fig. 1b) occur at the vertices of the unoccupied triangles, and the (OH) groups in the first sheet (Fig. 1a) occur at the vertices of the unoccupied triangle pointing downward in the figure. Thus the distribution of the (OH) groups in each sheet is the same, although the coordination number of OH(17) is one less than the coordination number of the pseudosymmetrically related OH(22) owing to the coordination difference between U(2) and U(6) (Table 5).

These sheets alternate along the *b* direction (Fig. 2) and are linked by interstitial [10]-coordinated Ba atoms and hydrogen bonding involving interstitial (H<sub>2</sub>O) groups. The linkage between sheets is shown in more detail in Figure 3. There is a pseudo-mirror plane orthogonal to *b* whereby U(1) is related to U(4), U(2) to U(6), and U(3) to U(5). As indicated above, the principal difference between the two sheets involves the difference in coordination of U(2) and U(6), but inspection of Figure 3 shows that, in addition to this difference in coordination, there are significant geometrical differences in the tilting of the polyhedra in each sheet.

### Interlayer $(H_2O)$ groups

There are eight O-atom sites in the interlayer of billietite, and bond-valence calculations indicate that they are all are (H<sub>2</sub>O) groups (Table 5). The OW(23), OW(25), OW(29) and OW(30) groups are bonded to Ba, whereas OW(24), OW(26), OW(27) and OW(28) are not. The arrangement of the (H<sub>2</sub>O) groups in billietite is similar to the interlayer configuration in becquerelite (Burns & Li 2002).

### HYDROGEN BONDING IN BILLIETITE

Hydrogen-bonding plays a key role in the structural connectivities and stabilities of hydrated minerals (Hawthorne 1992), and nearly all minerals containing the uranyl ion also contain essential (H<sub>2</sub>O) groups (Burns *et al.* 1996, Finch 1997). Because H atoms can rarely be located in difference-Fourier maps for highly absorbing materials, the accurate determination of U-O bond lengths (especially those for the uranyl ion) provides the best information for deriving hydrogen-bond interactions between the structural unit and interstitial (interlayer) (H<sub>2</sub>O) groups in U minerals (Finch *et al.* 1996), although the difference in X-ray

scattering for U and O can make accurate determination of the short uranyl (U–O) bond-length difficult (Finch *et al.* 1999).

# Hydroxyl groups

There are six symmetrically distinct (OH) groups in the structural unit of billietite, and inspection of the bond-valence table (Table 5) indicates that each must act as a bond-valence donor to one or more interstitial (H<sub>2</sub>O) groups. In most cases, inspection of the (OH)–OW distances unambiguously indicates the acceptor (H<sub>2</sub>O) groups (Table 6). Note that OW(28) is an acceptor for hydrogen bonds from both OH(17) and OH(22), and that the corresponding OH–OW distances are significantly longer (~3.00 Å) than for those (H<sub>2</sub>O) groups that accept just one hydrogen bond from the (OH) groups (~2.75 Å).

#### $(H_2O)$ groups: general considerations

There are eight symmetrically distinct ( $H_2O$ ) groups in the interlayer region of billietite. As noted above, five of these groups act as bond-valence acceptors for hydrogen bonds from (OH) groups of the uranyl sheet. The remaining ( $H_2O$ ) groups [OW(23), OW(25), OW(29) and OW(30)] bond to the interstitial Ba atom. Note that OW(25) bonds to Ba *and* is a hydrogen-bond acceptor, and is the only ( $H_2O$ ) group in billietite to exhibit this coordination.

# $(H_2O)...O$ hydrogen bonding

Inspection of the bond-valence table for billietite (Table 5) shows that there are twelve O atoms [O(1) to O(12)] that receive significantly less than 2.0 vu from their coordinating U and Ba atoms, and hence must be hydrogen-bond acceptors. Thus at least twelve donor bonds involving interstitial (H<sub>2</sub>O) groups must also involve O atoms of the uranyl sheet; these interactions may be identified from the observed OW-O distances. The O(6), O(9) and O(11) atoms each have only one (H<sub>2</sub>O) group within range of strong to medium hydrogen-bonding: OW(23), OW(23) and OW(30), respectively (Table 6). As OW(23) is involved in two medium-strength hydrogen bonds, with O(6) and O(9), O(2) and O(7) must accept hydrogen bonds from OW(29) and OW(25), as these are the only other (H<sub>2</sub>O) groups within range. The O(1)-OW(30) distance is 2.79 Å, whereas the other O(1)–OW distances are greater than 3.0 Å; as the O(1) anion shows a considerable bond-valence deficiency, if one considers only its coordinating U atoms, a strong hydrogen-bonding interaction is required, and hence we assign this interaction to O(1)-OW(30) (Table 6). As OW(30) is now involved in two hydrogen bonds, O(4) must be an acceptor anion for OW(29) (Table 6). This leaves O(5) and O(8), with distances to OW(24) and OW(25) between 2.83 and

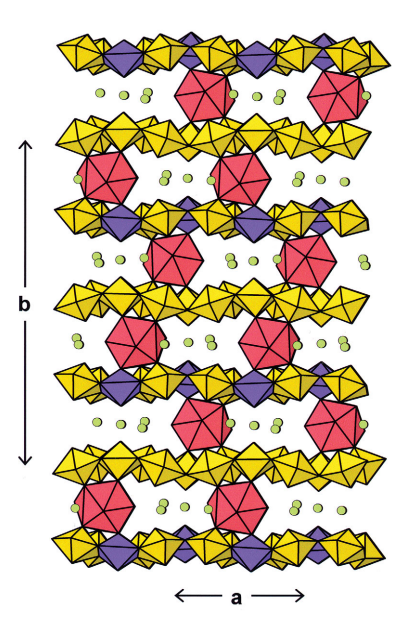


FIG. 2. The crystal structure of billietite projected down the c axis. Legend as in Figure 1; the Ba polyhedra are shown in red, and the interstitial (H<sub>2</sub>O) groups are shown as green circles.

3.15 Å; considering the bond-valences incident at O(5) (1.85 vu) and O(8) (1.74 vu) from their coordinating U and Ba atoms, the O(8)–OW interaction was assigned to the shorter distance [O(8)–OW(25) 2.83 Å]. The latter assignment forces O(3) to be an acceptor for OW(26). The O(10) and O(12) are assigned as acceptors to the

closest remaining (H<sub>2</sub>O) groups OW(27) and OW(24), respectively. All assignments are listed in Table 6.

# $(H_2O)...(H_2O)$ hydrogen bonding

All  $(H_2O)$  groups involved only in hydrogen bonds were assumed to be [4]-coordinated (by H atoms). This

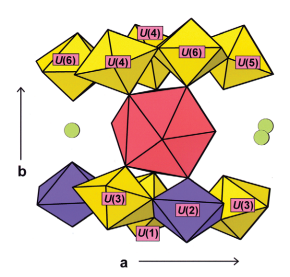
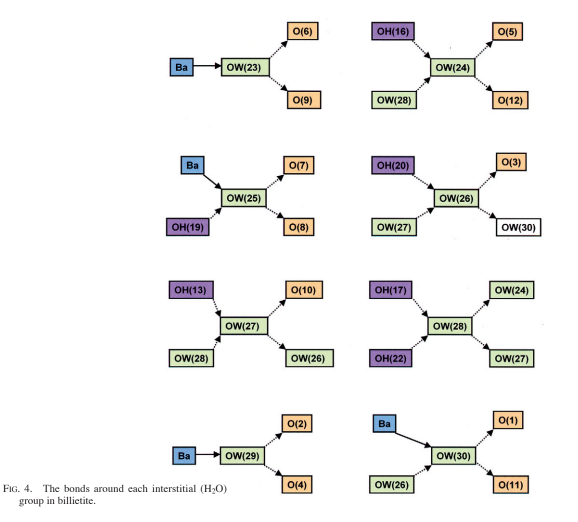


FIG. 3. Details of the linkage of polyhedra around the *Ba* site in billietite; legend as in Figure 2.



being the case, only one arrangement of hydrogen bonds is possible; the relevant interatomic distances are given in Table 6. The arrangement of bonds around each  $(H_2O)$  group in billietite is shown in Figure 4.

#### ACKNOWLEDGEMENTS

We thank Mr. Forrest Cureton for the crystal used in this study, Thomas Albrecht-Schmidt, J. Sejkora and Associate Editors Uwe Kolitsch and Sergey V. Krivovichev for their comments on the manuscript, and the distant pen of Bob Martin for his inimitable clarifications. This research was funded by the Environmental Management Sciences Program of the United States Department of Energy (DE–FG07–97ER14820) grant to PCB, and the Natural Sciences and Engineering Research Council of Canada grants to FCH and a International Postdoctoral Fellowship to RJF.

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- Received July 8, 2005, revised manuscript accepted June 1, 2006.