

Hydrogen bonding in borcarite, an unusual borate-carbonate mineral

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Abstract

The crystal structure, including hydrogen positions, of borcarite, $\text{Ca}_4\text{Mg}[\text{B}_4\text{O}_6(\text{OH})_6](\text{CO}_3)_2$, monoclinic, $a = 17.840(4)$, $b = 8.380(2)$, $c = 4.445(1)$ Å, $\beta = 102.04(3)^\circ$, $V = 649.9(3)$ Å³, space group $C2/m$, has been refined by full-matrix least-squares methods to $R = 2.5\%$ and $wR = 3.0\%$ for 1020 unique observed [$|F| \geq 5\sigma(F)$] reflections collected using Mo- $K\alpha$ X-radiation. The H positions were located on difference-Fourier maps and were refined using the 'soft' constraint that O-H distances are ~ 0.96 Å. The 4:4T FBB (fundamental building block) of the borcarite structure contains four $\text{B}\phi_4$ tetrahedra ($\phi = \text{unspecified ligand}$) which share corners to form a four-membered polyhedral ring. Borcarite is the only mineral known to contain this FBB. The FBBs do not polymerize, but each shares three anions with an $\text{Mg}\phi_6$ octahedron on either side, forming rods of composition $[\text{MgB}_4\text{O}_6(\text{OH})_6]^{4-}$ along the c -axis. The rigidity of these rods is enhanced by hydrogen bonding, and individual rods are connected through $\text{Ca}\phi_8$ polyhedra, $\text{C}\phi_3$ triangles and hydrogen bonds.

KEYWORDS: borcarite, crystal structure, hydrogen bond, borate, Russia.

Introduction

BORCARITE, $\text{Ca}_4\text{Mg}[\text{B}_4\text{O}_6(\text{OH})_6](\text{CO}_3)_2$, is a rare borate-carbonate mineral from Siberia, where it occurs as dense masses and as veins in kotoite marble and in ludwigite-szaibelyite-magnetite rock (Pertzev *et al.*, 1965). The crystal structure of borcarite was solved by Yamnova *et al.* (1976), but they did not report the H-atom positions. We are currently studying the infrared spectra of borate minerals in the principal OH-stretching region (i.e. Burns and Hawthorne, 1994a). As a part of this work, we have refined the structure of borcarite including the H positions, and we report the results of the refinement here.

Experimental

Samples of borcarite from Yakutskaya ASSR, Russia, were obtained from the Canadian Museum of Nature

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(samples 53411 and 53412). A small crystal fragment was mounted on a Nicolet $R3m$ automated four-circle diffractometer. Twenty-six reflections over the range $5^\circ \leq 2\theta \leq 35^\circ$ were centred using graphite-monochromated Mo- $K\alpha$ X-radiation. Unit-cell dimensions (Table 1) were derived from the setting angles of the twenty-six automatically aligned reflections by least-squares refinement. Data were collected using the $\theta/2\theta$ scan method with a 2.2° 2θ scan-range and a variable scan-rate ranging from 4 to 29.3° $2\theta/\text{min}$. A total of 1969 reflections was measured; the index ranges $0 \leq h \leq 25$, $-11 \leq k \leq 11$, $-6 \leq l \leq 6$ were covered and reflections forbidden by the C-centring were omitted. Two standard reflections were measured every fifty reflections; no significant change in their intensities occurred during data collection. An empirical absorption correction based on 36 psi-scans for each of 10 reflections over the range $9^\circ \leq 2\theta \leq 56^\circ$ was applied, reducing R (azimuthal) from 2.11 to 1.23%. The data were corrected for Lorentz, polarization and background effects; of the 1969 reflections measured, there were 1020 unique reflections ($R_{\text{int}} = 1.55\%$) classed as observed [$|F| \geq 5\sigma(F)$].

TABLE 1. Miscellaneous information for borcarite

a (Å)	17.840(4)	Crystal size (mm)	0.04 × 0.15 × 0.12
b (Å)	8.380(2)		
c (Å)	4.445(1)	Total ref.	1969
β (°)	102.04(3)	$ F_o \geq 5\sigma(F)$	1020
V (Å ³)	649.9(3)	R_{int}	1.55%
Space group	$C2/m$		
$F(000)$	548	Final R	2.5%
D_{calc}	2.790 g/cm ³	Final wR	3.0%
μ	1.79 mm ⁻¹		
Unit cell contents $2(\text{Ca}_4\text{Mg}[\text{B}_4\text{O}_6(\text{OH})_6](\text{CO}_3)_2)$			
$R = \Sigma(F_o - F_c) / \Sigma F_o $			
$wR = [\Sigma w(F_o - F_c)^2 / \Sigma F_o^2]^{1/2}$, $w = 1/\sigma^2(F)$			

Structure refinement

Scattering curves for neutral atoms, together with anomalous-dispersion corrections, were taken from Cromer and Mann (1968) and Cromer and Liberman (1970), respectively. The Siemens SHELXTL PLUS (PC version) system of programs was used throughout this study. The R indices are of the form given in Table 1 and are expressed as percentages.

Refinement of the structure was done in the space group $C2/m$ using the atomic parameters of Yamnova *et al.* (1976) as the starting model. Refinement of the positional and isotropic-displacement parameters gave an R index of 4.3%. Conversion to an anisotropic-displacement model, together with refinement of all parameters, gave an R index of 2.6%. At this stage of the refinement, a difference-Fourier map was calculated, and the positions of two H atoms were located. Subsequent cycles of refinement showed that the H positions were not realistic, as indicated by anomalously short O–H bond lengths, a common feature of H positions refined using X-ray data. The 'soft' constraint that O–H distances should be ~ 0.96 Å was imposed by adding extra weighted observational equations to the least-squares matrix. Only the O–H distance is constrained, and each H position is free to seek its optimum position around the oxygen atom. Neutron-diffraction studies (e.g. Hamilton and Ibers, 1968) of structures containing hydrogen bonds have shown that donor-H bond lengths do vary somewhat with the H-acceptor bond length. However, the donor-H bond lengths obtained via unconstrained refinement of X-ray data normally fall well outside the range found using neutron data, and this makes the hydrogen bonding in the structure difficult to interpret. Neutron-diffraction studies of borate minerals are limited due to extreme absorption effects. Therefore, the most appropriate way to

obtain hydrogen bonding information is via constrained refinement of X-ray data. We have found (Burns and Hawthorne, 1993*a,b*, 1994*b,c,d*) that this approach gives realistic hydrogen-bonding schemes for various borate minerals. Refinement of all parameters gave a final R index of 2.5% and a wR index of 3.0%. A refinable weighting scheme of structure factors and a correction for isotropic extinction were tried, but did not improve the refinement results. Final positional and displacement parameters are given in Table 2, selected interatomic distances and angles in Table 3, and a bond-valence analysis is given in Table 4. Observed and calculated structure factors have been deposited with the editor of *Mineralogical Magazine* and are available on request.

Discussion

Borate fundamental building block. There are two symmetrically distinct B positions in the borcarite structure, and both are tetrahedrally coordinated by (O,OH). The borate fundamental building block (FBB) in this structure is a $[\text{B}_4\text{O}_6(\text{OH})_6]^{6-}$ boron-oxygen ring (Fig. 1) made up of four $\text{B}\phi_4$ tetrahedra. Each $\text{B}\phi_4$ tetrahedron shares corners with two adjacent $\text{B}\phi_4$ tetrahedra, forming a four-membered polyhedral ring, and the borate FBBs are isolated (i.e. are not polymerized) in the structure. According to the rules of Christ and Clark (1977), the shorthand notation for this FBB is 4:4T, and borcarite is the only mineral that is known to contain this FBB.

The $\text{B}1\text{O}_2(\text{OH})_2$ and $\text{B}2\text{O}_3(\text{OH})$ tetrahedra have $\langle \text{B}-\text{O} \rangle$ distances of 1.483 and 1.484 Å, respectively, values which are within the range of $\langle {}^{14}\text{B}-\text{O} \rangle$ observed in minerals. The $\text{B}1\phi_4$ tetrahedron shows typical bond-length ranges (Table 3). However, the $\text{B}2\phi_4$ tetrahedron is considerably more distorted than is usual for $\text{B}\phi_4$ tetrahedra in minerals; there is one short B–O distance (1.397 Å, see Table 3) and three

TABLE 2. Final atomic parameters for borcarite

	<i>x</i>	<i>y</i>	<i>z</i>	* <i>U</i> _{eq}	** <i>U</i> ₁₁	<i>U</i> ₂₂	<i>U</i> ₃₃	<i>U</i> ₁₂	<i>U</i> ₁₃	<i>U</i> ₂₃
Ca	0.15543(2)	0.23368(5)	0.1548(1)	90(1)	81(2)	77(2)	109(2)	-8(2)	15(1)	-12(2)
Mg	0	0	0	53(4)	46(7)	72(7)	40(7)	0	10(5)	0
B1	0	0.2400(4)	½	130(8)	124(13)	125(14)	139(13)	0	23(10)	0
B2	0.0948(2)	0	0.6171(8)	80(8)	65(14)	84(14)	82(14)	0	-5(11)	0
C	0.3048(2)	0	0.2515(7)	106(8)	83(13)	125(14)	92(13)	0	-24(10)	0
O1	0.0994(1)	0	0.3071(5)	85(6)	91(10)	97(10)	67(9)	0	15(7)	0
O2	0.05513(8)	0.1427(2)	0.7204(3)	81(4)	88(7)	77(7)	73(7)	17(5)	4(5)	1(5)
OH3	0.04149(9)	0.3385(2)	0.3095(3)	97(4)	105(7)	90(7)	100(7)	-12(6)	30(5)	14(6)
O4	0.2150(1)	0.3654(2)	0.6487(4)	201(5)	143(8)	206(9)	221(9)	57(7)	-36(7)	-118(8)
OH5	0.1743(1)	0	0.8236(5)	102(6)	71(10)	137(11)	93(10)	0	7(8)	0
O6	0.3441(2)	0	0.0398(6)	176(7)	246(13)	155(12)	156(12)	0	105(10)	0
H1	0.005(1)	0.355(4)	0.127(4)	+300						
H2	0.212(2)	0	0.698(9)	+300						

**U*_{eq} = *U*_{eq} Å² × 10⁴
 ***U*_{ij} = *U*_{ij} Å² × 10⁴
 + fixed during refinement

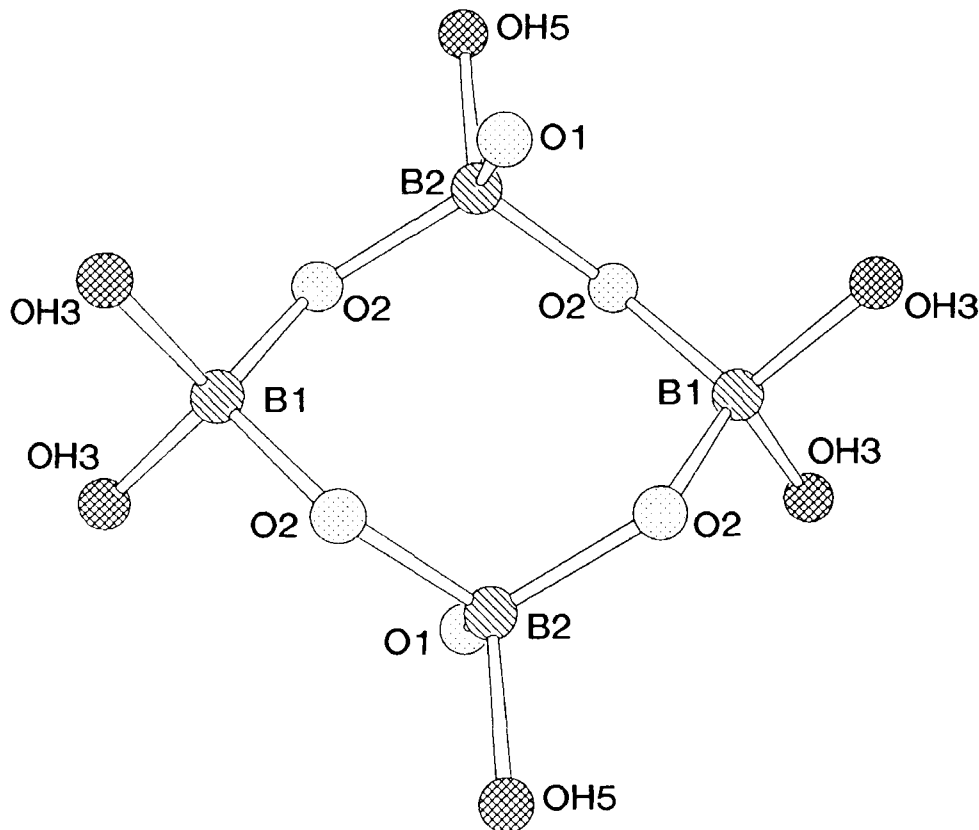


FIG. 1. The borate FBB in borcarite. B atoms are ruled with parallel lines, OH anions are cross-hatched, and O atoms are stippled.

TABLE 3. Selected interatomic distances (Å) and angles (°) for borcarite

Ca—O1	2.361(2)		B1—O2,f	1.480(2)	× 2
Ca—OH3	2.441(2)		B1—OH3,f	1.486(3)	× 2
Ca—O4	2.489(2)		<B1—O>	1.483	
Ca—O2a	2.464(2)				
Ca—O4a	2.899(2)		B2—O1	1.397(4)	
Ca—O4b	2.439(2)		B2—O2,h	1.509(3)	× 2
Ca—OH5a	2.514(2)		B2—OH5	1.520(4)	
Ca—O6c	2.394(1)		<B2—O>	1.484	
<Ca—O>	2.500				
			C—O6	1.285(3)	
Mg—O1,d	1.999(2)	× 2	C—O4b,i	1.289(3)	× 2
Mg—O2a,e,f,g	2.110(2)	× 4	<C—O>	1.288	
<Mg—O>	2.073				
Hydrogen bonding					
H1—OH3	0.94(2)		H1...OH3j	1.95(2)	
OH3—OH3j	2.839(3)		OH3—H1—OH3j	157(3)	
H2—OH5	0.95(5)		H2...O4b,i	2.49(2)	× 2
OH5—O4b,i	3.363(3)	× 2	OH5—H2—O4b,i	152(3)	× 2
Mg ϕ_6 octahedron					
O1,d—O2a,g	2.827(3)	× 4	O1,d—Mg—O2a,g	86.9(1)	× 4
O1,d—O2e,f	2.984(3)	× 4	O1,d—Mg—O2e,f	69.0(1)	× 4
O2a,e—O2g,f	2.391(3)	× 2	O2a,e—Mg—O2g,f	93.1(1)	× 2
O2a,e—O2f,g	3.476(3)	× 2	O2a,e—Mg—O2f,g	111.0(1)	× 2
<O—O>	2.915		<O—Mg—O>	90.0	
B1 ϕ_4 tetrahedron					
O2,f—OH3,f	2.430(2)	× 2	O2,f—B1—OH3,f	110.0(1)	× 2
O2,f—OH3,f	2.364(2)	× 2	O2,f—B1—OH3,f	105.7(1)	× 2
O2—O2f	2.470(3)		O2—B1—O2f	113.1(3)	
OH3—OH3f	2.471(3)		OH3—B1—OH3f	112.5(3)	
<O—O>	2.421		<O—B1—O>	109.5	
B2 ϕ_4 tetrahedron					
O1—O2,h	2.457(3)	× 2	O1—B2—O2,h	115.3(2)	× 2
O2,h—OH5	2.399(2)	× 2	O2,h—B2—OH5	104.7(2)	× 2
O1—OH5	2.404(3)		O1—B2—OH5	110.9(3)	
O2—O2h	2.391(3)		O2—B2—O2h	104.8(3)	
<O—O>	2.418		<O—B2—O>	109.3	
C ϕ_3 triangle					
O6—O4b,i	2.217(3)	× 2	O6—C—O4b,i	118.9(2)	× 2
O4b—O4i	2.257(4)		O4b—C—O4i	122.2(3)	
<O—O>	2.230		<O—C—O>	120.0	

a = x, y, z-1; b = 1/2-x, 1/2-y, 1-z; c = 1/2-x, 1/2-y, -z; d = -x, -y, -z; e = -x, -y, 1-z; f = -x, y, 1-z;
g = x, -y, z-1; h = x, -y, z; i = 1/2-x, y-1/2, 1-z; j = -x, y, -z.

longer distances (1.509 to 1.520 Å). The B2 ϕ_4 tetrahedron shares two corners (O2) with B1 ϕ_4 tetrahedra (Fig. 1), and the B2—O2 distance is 1.509 Å. The other two anions in the B2 ϕ_4 tetrahedron are non-bridging. In almost all hydrous borates, non-bridging oxygen atoms bond to a

TABLE 4. Bond-valence* analysis (v.u.) for borcarite

	Ca	Mg	B1	B2	C	H1	H2	Σ
O1	0.345	0.437 \downarrow ^{x2}		0.932				1.714
O2	0.261	0.324 \downarrow ^{x4}	0.745 \downarrow ^{x2}	0.689 \downarrow ^{x2}				2.019
OH3	0.278		0.733 \downarrow ^{x2}			0.80		2.011
						0.20		
O4	0.244				1.314 \downarrow ^{x2}		0.05 \downarrow ^{x2}	1.968
	0.279							
	0.081							
OH5	0.228			0.669			0.90	1.797
O6	0.315				1.328			1.643
Σ	2.031	2.170	2.956	2.979	3.956	1.00	1.00	

* Bond-valence parameters from Brown and Altermatt (1985).

hydrogen atom, forming an hydroxyl group. This is the case for the OH5 anion of the B2 ϕ_4 tetrahedron, and the B2–OH5 bond-length (1.520 Å) is typical. However, B2 also bonds to O1 which is a non-bridging oxygen atom that does not bond to an H atom. The O1 atom bonds to one Ca and one Mg atom as well, and if the B2–O1 distance was \sim 1.48 Å, the bond-valence sum at B2 would be \sim 1.55 v.u., an unrealistically low total. Thus, the O1 atom moves closer to B2, increasing the bond-valence sum at the O1 position to 1.71 v.u. (Table 4).

C ϕ_3 triangle. The structure contains one C position which is coordinated by three coplanar O atoms with a <C–O> distance of 1.288 Å, a value typical for (CO₃) groups in minerals and inorganic structures. The carbonate triangle has two distinct bond lengths (Table 3) and is distorted, with bond angles of 118.90 and 122.2°, giving a <O–C–O> angle of 120.0°.

Ca ϕ_8 and Mg ϕ_6 polyhedra. The borcarite structure has one unique Ca position which is coordinated by six O atoms and two hydroxyl groups. The Ca– ϕ bond-lengths range from 2.361 to 2.899 Å, and the <Ca– ϕ > distance is 2.500 Å. The structure contains one symmetrically distinct Mg position which is octahedrally coordinated by six oxygen atoms. The octahedron is (2+4) distorted, with four long Mg–O2 bonds (2.110 Å) and two short Mg–O1 bonds (1.999 Å), giving a <Mg–O> distance of 2.073 Å. This distortion is due to the bond-valence requirements of the underbonded O1 position (see above, Table 4), which moves closer to Mg to increase the bond-valence sum at that position.

Structure connectivity. In the borcarite structure, the 4:4T borate FBBs are isolated from each other. The four-membered borate polyhedral ring provides

three oxygen atoms that are in appropriate positions to be an octahedral face, and this is the point of attachment to the Mg ϕ_6 octahedron (Fig. 2a). Each FBB bonds to two Mg ϕ_6 octahedra, and each Mg ϕ_6 octahedron bonds to two FBBs, forming infinite rods of composition [MgB₄O₆(OH)₆]^{4–} along the *c*-axis (Fig. 2b). These rods are *C*-centred and are cross-linked via Ca– ϕ bonds in combination with the C ϕ_3 triangles (Fig. 3) and hydrogen bonds (see below).

Hydrogen bonding. The H-atom positions obtained via constrained least-squares refinement of the X-ray data are reasonable both in terms of donor and acceptor bond-lengths and angles (Table 3) and the bond-valence requirements of the donor and acceptor anions (Table 4).

There are two hydrogen positions in the structure of borcarite. The OH3–H1 \cdots OH3 bond has an acceptor distance of 1.95 Å, and is a fairly strong hydrogen bond. This bond bridges between adjacent borate FBBs within the same [MgB₄O₆(OH)₆]^{4–} rod (Fig. 2a), thereby increasing the rigidity of this structural unit. The OH3 anion donates the hydrogen bond and accepts a hydrogen bond from H1 attached to a symmetrically equivalent OH3 position, forming OH3–H1 \cdots OH3–H1 rings (Fig. 2a). There are four hydrogen bonds between each adjacent borate FBB within the [MgB₄O₆(OH)₆]^{4–} rod.

The OH5–H2 \cdots O4 bond bridges between the borate FBBs and adjacent C ϕ_3 triangles (Fig. 3). The H2 position lies on a mirror plane in the space group *C2/m*, and there are two symmetrically equivalent O4 atoms that may act as acceptor anions. The OH5–H2 \cdots O4 bonds both have an acceptor distance of 2.49 Å, thus the bonds are rather weak. It is possible that this hydrogen bond is bifurcated to the two O4 anions, and a difference-

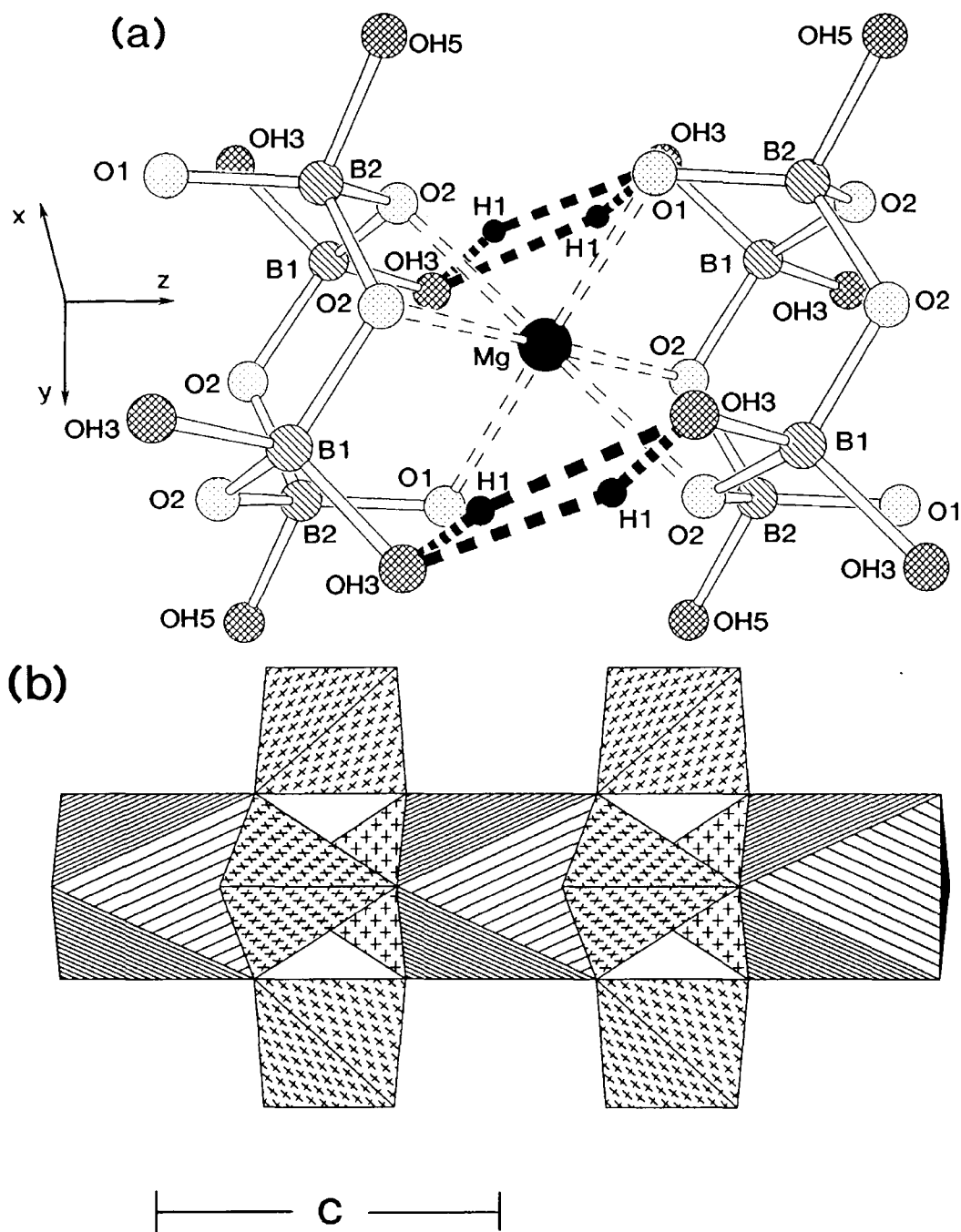


Fig. 2. The structure rod of composition $[\text{MgB}_4\text{O}_6(\text{OH})_6]^{4-}$: (a) the local environment of the borate FBB, B atoms are ruled with parallel lines, the Mg atom is a large solid circle, OH anions are cross-hatched, O atoms are stippled, H atoms are small solid circles; O-H bonds are the short-dashed lines, H...H bonds are the long-dashed lines. (b) the $[\text{MgB}_4\text{O}_6(\text{OH})_6]^{4-}$ rod projected down the *a*-axis. $\text{B}\phi_4$ tetrahedra are shaded with crosses and $\text{Mg}\phi_6$ octahedra are ruled with parallel lines.

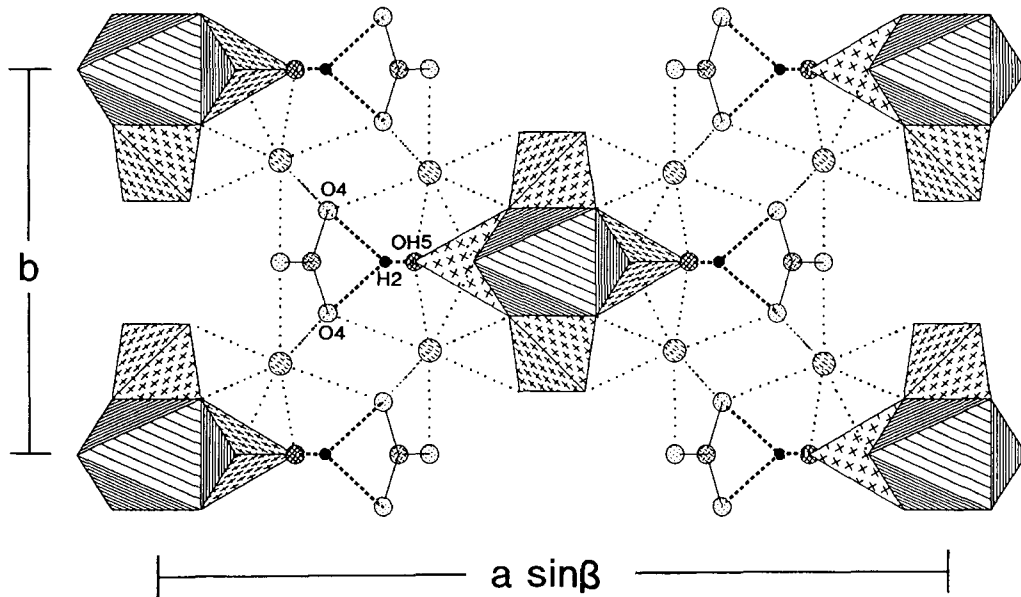


FIG. 3. The borcarite structure projected down the c -axis. $B\phi_4$ tetrahedra are shaded with crosses, $Mg\phi_6$ octahedra are ruled with parallel lines, oxygen atoms are stippled, OH anions are cross-hatched, C atoms are shaded with a herring-bone pattern, Ca atoms are ruled with broken parallel lines and H atoms are shown as small solid circles. Ca- ϕ bonds are dotted, C- ϕ are solid lines, H- ϕ bonds are broken lines.

Fourier map calculated with the H2 atom removed from the structure model supports this hypothesis, as it shows that the electron density associated with the H2 position is somewhat anisotropic, with maximum displacements perpendicular to the mirror plane.

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