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Bavsiite, Ba₂V₂O₂[Si₄O₁₂], mineral data and crystal structure

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Abstract

Bavsiite from the type locality Gun Claim, Yukon Territory, Canada, occurs as millimetre-sized sky-blue platy crystals in a barium-rich low-temperature skarn related to a porphyritic quartz monzonite stock. Associated minerals are alstonite, baryte, celsian, diopside, fresnoite, mica, suzukiite, walstromite, witherite and minerals of the cerchiaraite group. Bavsiite is optical uniaxial (+), with $\omega = 1.725(3)$ and $\epsilon = 1.750(3)$ (589 nm) and pleochroic. Electron microprobe analyses yielded the empirical formula Na_{0.02}Ba_{1.98}Ti_{0.16}Fe²⁺³_{0.05}V⁴⁺_{1.80} Al_{0.05}Si_{4.00}O₁₄ based on 14 oxygen atoms, the simplified chemical formula is Ba₂V₂Si₄O₁₄. Bavsiite is tetragonal, space group *I4/m*, *a* = 7.043(1), *c* = 11.444(2) Å and *Z* = 2 obtained from single crystal data at 100 K, which are in good agreement with cell parameters from powder diffraction data at 293 K: *a* = 7.051(1) Å and *c* = 11.470(1) Å. The eight strongest lines of the powder X-ray diffraction pattern are [*d*, Å (*I*,%)(*hkl*)]: 3.76(30)(112), 3.36(44)(013), 3.004(100)(022), 2.493(43)(220), 2.486(67)(114), 2.286(24)(222), 1.785(39) (116) and 1.763(25)(040). The crystal structure was refined to *R* = 0.0159 based upon 312 unique reflections with *I* > 2 σ (*I*). The crystal structure of bavsiite comprises unbranched single [Si₄O₁₂]⁸⁻ rings connected by [VO₅]⁶⁻ square pyramids and BaO₁₂ polyhedra. It can also be considered as cage–like [Si₄V₂O₁₈]¹²⁻ clusters built by four SiO₄ tetrahedra and two VO₅ square pyramids. These clusters are cross–linked to form a pseudo-two-dimensional network (2D) parallel to (001) containing empty channels along the *a* axis and the 2D networks held together by Ba²⁺ ions located in channels parallel to the *c* axis. The structural formula is Ba₂V₂O₂[Si₄O₁₂]. Bavsiite is polymorphic to suzukiite, BaVSi₂O₇, which is orthorhombic.

Keywords: bavsiite, new mineral, mineral data, crystal structure, Gun Claim, Yukon Territory, Canada

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Introduction

The Gun Claim (sometimes also written Gunn Claim), Yukon Territory (130°0'51"W, 62°50'50"N), Canada is located ~4 km southeast of Wilson lake and 16 km southeast of Itsy Peak. It is the type locality for itsiite (Kampf *et al.*, 2014), meierite (Peterson *et al.*, 2016), pellyite (Montgomery *et al.*, 1972) and the co-type locality of titantaramellite (Alfors and Pabst, 1984). The mineralisation is a barium-rich low-temperature skarn related to a porphyritic quartz monzonite stock. Skarn bodies have developed in limestone adjacent to the igneous contact. The major mineral assemblage includes andradite, baryte, cerchiaraite, gillespite, pellyite, quartz, sanbornite, taramellite and witherite (Montgomery *et al.*, 1972; MacNeil *et al.*, 2013; Kampf *et al.*, 2014).

Occurrence and physical properties

Bavsiite has been approved by the Commission on New Minerals, Nomenclature and Classification of the International Mineralogical Association (IMA2014-019, Bojar and Walter, 2014) and the type specimen is hosted in the mineralogical

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collection of the Universalmuseum Joanneum Graz, Austria (catalogue number 85.282). Bavsiite and suzukiite, which have nearly the same chemical composition, were found in a white, marble-like skarn-type rock. Bavsiite (Fig. 1) occurs as sky-blue, platy crystals up to 0.3 mm across with a thickness of a few hundredth of a millimetre and flattened on {001}. Bavsiite is transparent with a vitreous lustre and is non-fluorescent. The streak is white, the Mohs hardness is ~4 and cleavage parallel to (001) is good. Density was not measured directly due to frequent inclusions in bavsiite. Directly associated minerals are alstonite, baryte, celsian, diopside, fresnoite, mica, suzukiite, walstromite, witherite and minerals of the cerchiaraite group.

The Fourier-transform infrared spectrum was recorded using a Perkin Elmer Spectrum 100 spectrophotometer (ATR diamond cell, resolution 4 cm⁻¹, 20 scans). The infrared spectra for bavsiite and suzukiite are overlaid for comparison and given in Fig. 2. The following bands for bavsiite were observed (in cm⁻¹; s: strong, m: medium, w: weak, sh: shoulder): 1223w, 1068s, 993sh, 916s, 743w, 659s, 584m, 537w, 478s and 427m. The bands in the region from 1200 to 900 cm⁻¹ are related to the stretching of Si–O in the SiO₄ tetrahedron and V–O of the VO₅ pyramid. The region of 740 to 400 cm⁻¹ is related to the bending modes of the SiO₄ and VO₅ structural elements.

Bavsiite is optical uniaxial (+); its optical constants measured with 589 nm on a spindle stage are $\omega = 1.725(3)$ and $\epsilon = 1.750(3)$, the dispersion is weak with r > v. Pleochriosm for ω is medium blue and for ϵ , pale yellow.

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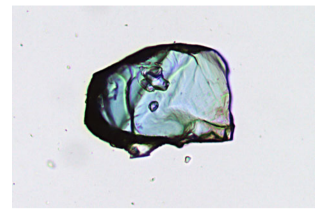


Fig. 1. Transmission-light microscopy image of a single crystal of bavsiite (cat. nr. 85.282). Field of view is 0.2 mm. Photograph: F. Walter.

Composition

Electron microbeam wavelength dispersive analyses were performed on a Jeol 6610 LV electron microscope equipped with an OXFORD 50 mm² energy dispersive and an OXFORD INCA WAVE spectrometer operated at an accelerating voltage of 20 kV and a beam current of 10 nA. Data reduction was done with the *PAP* routine (Pouchou and Pichoir, 1991) implemented in the INCA software of Oxford Instruments. Results of the analyses of baysiite and minerals of its paragenesis are given

Table 1. Representative compositions from electron microprobe wavelength dispersive analyses of bavsiite. Average of 10 spot analyses from five crystal fragments.

Constituent	stituent Wt.% Range		S.D.	Standard	
SiO ₂	33.40	32.67-34.40	0.47	almandine	
TiO ₂	1.75	1.07-2.30	0.06	rutile	
Al_2O_3	0.33	0.25-0.49	0.02	almandine	
FeO	0.30	0.16-0.45	0.02	almandine	
VO ₂	20.69	20.16-21.39	0.35	V metal	
Na ₂ O	0.08	b.d.l0.15	0.02	albite	
BaO	42.28	41.52-43.87	0.67	benitoite	
Total	98.83				

S.D. - standard deviation; b.d.l. - below detection limit.

in Tables 1 and 2. The empirical formula of bavsiite on the basis of 14 oxygen atoms is: $Na_{0.02}Ba_{1.98}Ti_{0.16}Fe_{0.03}^{2+}V_{1.80}^{4+}Al_{0.05}Si_{4.00}O_{14}$. The simplified chemical formula is $Ba_2V_2Si_4O_{14}$ which requires BaO 43.02, VO_2 23.27 and SiO_2 33.72 wt.%. The elements Ca, Mg, Mn and Sr were also measured, but found to be below the detection limits. The chemical composition of suzukiite and its empirical formula is very close to that of bavsiite: $Na_{0.02}Ba_{1.94}Ti_{0.12}Fe_{0.03}^{2+}V^4$ $^+_{1.85}Al_{0.04}Si_{4.01}O_{14}$.

Associated feldspars are very low in Ca and high in Ba with higher Ba content in their cores. Fresnoite analyses $(Na_{0.02}Ca_{0.02}Ba_{0.96}Fe_{0.01}^{2+}V_{0.22}^{4+}Ti_{0.79}Al_{0.01}Si_{1.98}O_8)$ yielded considerable V⁴⁺ (tetravalence from stoichiometric considerations) values.

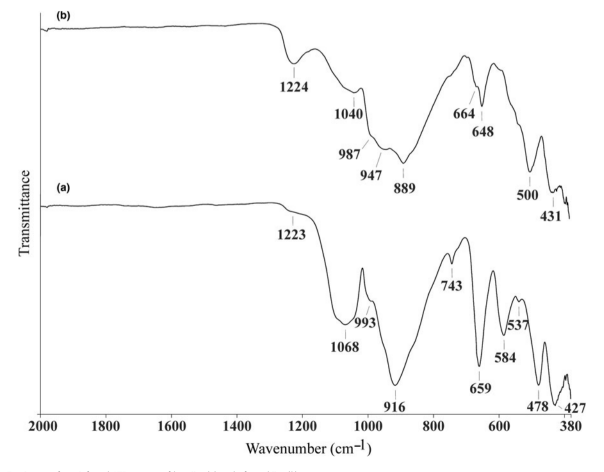


Fig. 2. The Fourier-transform infrared ATR spectra of bavsiite (a) and of suzukiite (b).

Table 2. Representative electron microprobe results from wavelength-dispersive spectroscopy analyses and formula units for bavsiite and minerals of its paragenesis*.

	Celsian [1]	Walstromite [2]	Fresnoite [3]	Diopside [4]	Bavsiite [5]	Suzukiite [6]
Wt.%						
SiO ₂	35.53	40.76	23.99	52.76	33.40	34.14
TiO ₂	b.d.l.	0.16	12.77	b.d.l.	1.75	1.35
Al_2O_3	24.29	b.d.l.	0.09	0.05	0.33	0.25
FeO	0.29	b.d.l.	0.13	4.14	0.30	0.29
VO_2	b.d.l.	b.d.l.	3.71	0.45	20.69	21.81
MgO	b.d.l.	b.d.l.	b.d.l.	15.32	b.d.l.	0.02
CaO	0.17	24.40	0.27	24.38	b.d.l.	0.04
SrO	b.d.l.	0.22	0.09	b.d.l.	b.d.l.	b.d.l.
Na ₂ O	0.10	0.03	0.10	0.13	0.08	0.09
K ₂ O	2.63	b.d.l.	b.d.l.	b.d.l.	b.d.l.	b.d.l.
BaO	35.88	35.10	60.85	b.d.l.	42.28	42.25
Sum	98.89	100.67	102.00	97.23	98.83	100.24
Atoms	per formu	ıla unit				
Si	2.18	3.01	1.98	1.99	4.00	4.01
Ti		0.01	0.79		0.16	0.12
Al	1.76		0.01		0.05	0.04
Fe ²⁺	0.02		0.01	0.13	0.03	0.03
V ⁴⁺			0.22	0.01	1.80	1.85
Mg				0.86		
Ca	0.01	1.93	0.02	0.99		
Sr		0.01				
Na	0.01		0.02	0.01	0.02	0.02
К	0.21					
Ва	0.86	1.02	1.96		1.98	1.94
Sum	5.05	5.98	5.01	3.99	8.04	8.01

*[1] celsian (on the basis of 8 oxygen atoms); [2] walstromite (9 oxygens); [3] fresnoite (8 oxygens); [4] diopside (6 oxygens); [5] bavsiite (14 oxygens, mean value of 10 analyses of 5 crystal fragments); and [6] suzukiite (14 oxygens, mean value of 5 analyses of one crystal fragment),

b.d.l. - below detection limit.

Slightly low analytical sums are caused most probably by a varying probe current.

Powder X-ray diffraction

The powder X-ray diffraction data (Table 3) for baysiite were obtained at 293 K using a Bruker-AXS D8 diffractometer with $CuK\alpha$ radiation. To minimise the background, the sample was placed on a low-background silicon wafer; an internal standard was not used. Unit-cell parameters, derived from whole pattern fitting (Pawley method) with fundamental parameters using TOPAS 4.2 software (Bruker-AXS, 2009) are: a = 7.051(1) Å, c = 11.470(1) Å and V = 570.2(1) Å³, which are in agreement with those obtained from the single-crystal study (see below). Due to the cleavage of baysiite parallel to (001) and due to the sample preparation, the intensities for (002), (004), (006) and (008) of the observed pattern are too high in comparison to the (00l) intensities of the simulated powder pattern, calculated with data from the single-crystal structure refinement (Table 3). In the powder pattern of bavsiite impurities of feldspar, suzukiite and mica are visible (Fig. 3a). For comparison, the simulated powder patterns of bavsiite calculated with parameters of the single-crystal structure refinement at 100 K (Fig. 3b) and of β -BaVSi₂O₇ (Fig. 3c) calculated with parameters of the structure refinement from Liu and Greedan (1994) are overlaid. The powder patterns confirm that baysiite and β-BaVSi₂O₇ have the same crystal structure but are polymorphic to suzukiite (BaVSi₂O₇, orthorhombic) shown by overlaying the

Table 3. Powder X-ray diffraction data (d in Å) for bavsiite.

I _{obs}	I _{calc}	d _{obs}	d_{calc}	h k l
2	3	6.009	6.007	011
107*	9	5.735	5.735	002
10	10	4.986	4.986	110
30	31	3.7630	3.7627	112
17	14	3.5258	3.5257	020
44	16	3.3609	3.3610	013
7	2	3.0398	3.0406	121
	4		3.0406	211
100	100	3.0037	3.0035	022
149*	7	2.8670	2.8674	004
43	35	2.4933	2.4930	220
67	33	2.4860	2.4857	114
24	17	2.2861	2.2863	222
7	3	2.2294	2.2298	130
	2		2.2298	310
11	7	2.2245	2.2246	024
6	2	2.1817	2.1814	015
11	6	2.0783	2.0782	132
	6		2.0782	312
2	<1	1.9278	1.9279	231
	1	1.9278	1.9279	321
3*	<1	1.9118	1.9116	006
6	5	1.8815	1.8814	224
39	13	1.7852	1.7849	116
25	14	1.7627	1.7628	040
	7		1.7602	134
	9		1.7602	314
4	2	1.7412	1.7411	233
	2		1.7411	323
3	3	1.6847	1.6850	042
17	5	1.6807	1.6805	026
5	4	1.5964	1.5963	332
6	3	1.5767	1.5767	240
	3		1.5767	420
2	<1	1.5610	1.5611	143
	1	1.5610	1.5611	413
16	9	1.5203	1.5203	242
	9		1.5203	422
7	6	1.5018	1.5017	044
5	1	1.4540	1.4540	127
	1		1.4540	217
21	7	1.4514	1.4513	136
	6		1.4513	316
8	5	1.4381	1.4379	334
106*	3	1.4338	1.4337	008
2	1	1.3817	1.3816	244
	<1		1.3816	424
8	3	1.3783	1.3779	118
3	1	1.3443	1.3442	152
	2		1.3442	512

 I_{obs} and d_{obs} from measurement at 293 K (Bruker D8 diffractometer, CuK α radiation); d_{calc} calculated with profile fitting using *TOPAS* 4.2 and a = 7.051(1) Å, c = 11.470(1) Å; I_{calc} calculated with atomic parameters from single-crystal structure refinement of bavsiite. * I_{obs} due to preferred orientation of the cleavage plane (001). Scaling is based on I_{obs} (022) = 100

The strongest lines are indicated in bold.

experimental powder pattern of suzukiite crystals from the type locality of bavsiite (Fig. 3d).

Single-crystal X-ray structure refinement

A pale blue platy crystal of bavsiite (Fig. 1) was used for X-ray single-crystal data collection on a Bruker AXS Smart Apex CCD diffractometer equipped with graphite-monochromatised MoKa radiation (0.71073 Å). The data were recorded with the program *SMART*, integrated with the program *SAINT* up to $\theta_{max} = 26.3^\circ$,

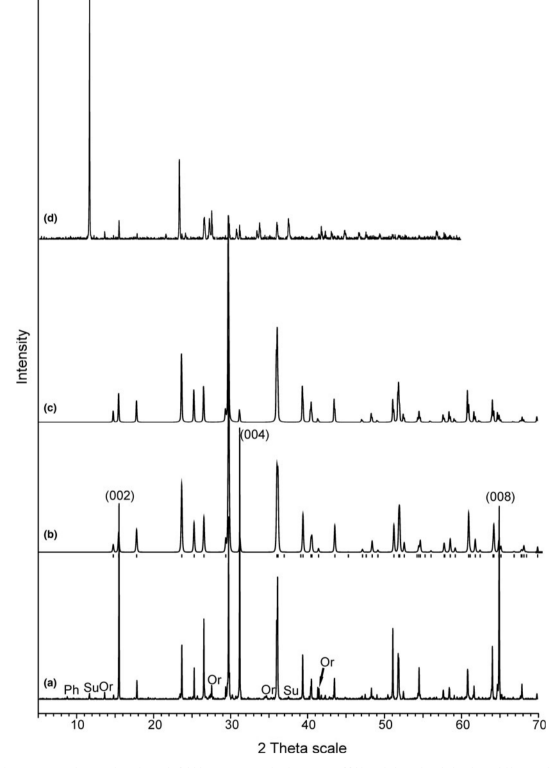


Fig. 3. The powder X-ray patterns (CuK α , 40 kV and 40 mA) of: (*a*) bavsiite, measured with impurities of feldspar (Or), suzukiite (Su) and mica (Ph) – note the preferred orientation effects on the (00*l*) reflections in the experimental pattern of bavsiite; (*b*) calculated powder pattern of bavsiite with data from structure refinement; (*c*) calculated powder pattern of β -BaVSi₂O₇ with parameters of the structure refinement from Liu and Greedan (1994); and (*d*) suzukiite, BaVSi₂O₇, experimental pattern with crystals from the type locality of bavsiite.

and corrected for absorption using the program *SADABS* (Bruker, 2008). The analyses of the intensity data indicate the two possible space groups I4/m and I4/mmm. The merged intensities for I4/m

result in $R_{\text{int}} = 0.033$, $\sigma = 0.015$ and for I4/mmm in $R_{\text{int}} = 0.062$, $\sigma = 0.018$. For baysiite we adopted as a starting point the space group I4/m and the atomic positions as reported for the structural

 Table 4. Crystal parameters, data collection and structure refinement details for bavsiite.

Crystal data	
Ideal formula	$Ba_2V_2O_2[Si_4O_{12}]$
Crystal size (mm)	$0.15 \times 0.11 \times 0.04$
Crystal system, space group	Tetragonal, <i>I</i> 4/ <i>m</i>
Formula weight	712.92
Temperature (K)	100
Unit-cell dimensions (Å)	a = 7.043(1), c = 11.444(2)
<i>V</i> (Å ³)	567.6(2)
Ζ	2
Density (from above formula) (g/cm ³)	4.171
Absorption coefficient (mm ⁻¹)	8.95
Data collection	
Description	Medium blue, platy crystal
Instrument	Bruker AXS Smart Apex CCD
Radiation type, wavelength (Å)	MoKα, 0.71073 Å
F(000)	652
θ range	3.40 to 26.28°
Index ranges	$-8 \le h \le 8, -8 \le k \le 8, -14 \le l \le 14$
Reflections collected/unique	2287/312
$R_{\rm int}, R_{\sigma}$	0.033, 0.015
Ratio of min. to max. transmission	0.636
Refinement	
Refinement method	Full-matrix least-squares on F ²
Data / restraints / parameters	312 / 0 / 32
Twinning by merohedry	010/100/001
BASF parameter	0.290(4)
Extinction	0.0027(4)
Final R indices $[I > 2\sigma(I)]$ (all data)	$R_1 = 0.0159, wR_2 = 0.0399$ (312 data)
GoF on F ²	1.175
Largest diff. peak and hole (e ⁻ Å ⁻³)	0.37 and -0.47

 $R_{1} = \Sigma ||F_{o}| - |F_{c}|| / \Sigma |F_{o}|, \ wR_{2} = [\Sigma (w(F_{o}^{2} - F_{c}^{2})^{2}) / \Sigma (w(F_{o}^{2})^{2})]^{\frac{1}{2}}$

 $w = 1/[\sigma^2(F_0^2) + (0.0179P)^2 + 2.33P], P = (Max(F_0^2, 0) + 2F_c^2)/3$

identic synthetic phase β-BaVSi₂O₇ by Liu and Greedan (1994). The small amounts of titanium and iron in the empirical formula of bavsiite (Table 2) were summarised with full occupancy of vanadium at the V site and all other sites were expected to be fully occupied by silicon, barium and oxygen, respectively. Least-squares structure refinement using anisotropic displacement parameters for all atoms was carried out with the program SHELXL-2013 (Sheldrick, 2015). It converged at satisfactory R values ($R_1 = 0.0279$) but the second parameter in the weighting scheme was unusually large (17.98). Refinement with space group I4/mmm gave nearly the same values for atomic positions and anisotropic displacement parameters but with slightly higher *R* values ($R_1 = 0.0323$) and also an unusually large second weighting parameter (24.32). These results are an indication of a twinning effect in the crystal structure of bavsiite. No splitting of reflections in the data collection was observed, thus a twinning by merohedry is predicted. In Laue group 4/m a mirror *m* parallel to the fourfold axis acts as a twin element and the apparent Laue

Table 6. Selected bond distances (Å) and angles (°) in bavsiite.

Ba–O(1) ×4	3.295(2)	O(2)-V-O(3) ×4	103.29(8)
Ba-O(2) ×4	2.812(4)	O(2)-V-O(2) ´ ×2	153.4(2)
Ba–O(2) ′×4	3.149(4)	O(2)-V-O(2) ″ ×4	86.97(4)
Ba–O(3) ×4	3.562(5)		
		O(1)-Si-O(1) ´	112.1(3)
V-O(2) ×4	1.951(3)	O(1)-Si-O(2) ×2	106.6(2)
V-O(3)	1.608(5)	O(1) ´-Si-O(2) ´ ×2	108.5(2)
		O(2)-Si-O(2)	114.6(2)
		<0-Si-0>	109.5
Si-O(1)	1.609(5)		
Si-O(1) ′	1.616(5)	Si–O(1)–Si´×4	157.9(5)
Si-O(2) ×2	1.594(3)		
<si-0></si-0>	1.603	Shortest OO	2.569

group 4/mmm results. The refinement with space group I4/m using the twinning law $010/100/00\overline{1}$ confirmed the merohedral twinning effect by lowering R_1 to 0.0159 and the presence of twin domains with proportions 0.29 to 0.71. In addition, the weighting scheme shows acceptable values (WGHT 0.0179, 2.33) and a smooth difference-Fourier map shows that absorption effects are well corrected.

Details on data collection, crystallographic data and refinement are given in Table 4. Refined atomic coordinates and displacement parameters are given in Table 5, and selected interatomic distances and angles are listed in Table 6. Bond valences (Table 7) were calculated using the equation of Brown and Altermatt (1985) with the parameters of Gagné and Hawthorne (2015). The crystallographic information files have been deposited with the Principal Editor of *Mineralogical Magazine* and are available as Supplementary material (see below).

Description of the crystal structure

Bavsiite, Ba₂V₂O₂[Si₄O₁₂], crystallises in a tetragonal unit cell of space group *I*4/m with one of the oxygen atoms in a general position whereas all other sites are located at special positions (Table 5). The structure can be considered as consisting of unbranched single $[Si_4O_{12}]^{8-}$ rings connected by $[VO_5]^{6-}$ square pyramids (Fig. 4), or of cage–like $[Si_4V_2O_{18}]^{12-}$ clusters built by four $[SiO_4]^{4-}$ tetrahedra and two $[VO_5]^{6-}$ square pyramids (Fig. 5). These clusters are cross-linked to form a pseudo-two-dimensional network (2D) parallel to (001) containing empty channels along the *a* axis as shown in Fig. 6. There is no connection between the $[VO_5]^{6-}$ square pyramids by themselves and the V–V distance of 3.580 Å sets an upper limit for the channel height (Fig. 6). The 2D networks are held together by Ba²⁺ ions located in channels parallel to the *c* axis (Fig. 7).

The Si–O bond length ranges from 1.594(3) to 1.616(5) Å, average 1.603 Å, and the O–Si–O angles from $106.6(2)^\circ$ to $114.6(2)^\circ$,

 $\label{eq:table 5. Atomic coordinates, equivalent isotropic and anisotropic displacement parameters ({ A}^2) for baysite.$

Atom	X	у	Ζ	$U_{\rm eq}$	U^{11}	U ²²	U ³³	U ²³	U ¹³	U ¹²
Ва	1/2	0	1/4	0.0117(2)	0.0130(2)	0.0130(2)	0.0093(2)	0	0	0
V	0	0	0.3436(1)	0.0078(3)	0.0081(3)	0.0081(3)	0.0072(5)	0	0	0
Si	0.2119(2)	0.2369(2)	0	0.0066(3)	0.0052(7)	0.0055(7)	0.0091(6)	0	0	0.0014(5)
O(1)	0.2682(5)	0.0144(7)	0	0.0283(10)	0.011(2)	0.008(2)	0.065(3)	0	0	0.003(2)
O(2)	0.2098(5)	0.1693(5)	0.3828(2)	0.0280(9)	0.028(2)	0.040(2)	0.016(1)	-0.013(1)	0.007(1)	-0.023(2)
O(3)	0	0	0.2031(4)	0.0165(10)	0.018(2)	0.018(2)	0.014(2)	0	0	0

 $U_{\rm eq}$ is defined as one third of the trace of the orthogonalised $U_{\rm ij}$ tensor

The anisotropic displacement factor exponent takes the form: $-2\pi^2 \left[h^2 a^{*2} U^{11} + ... + 2hka^* b^* U^{12}\right]$

Table 7. Bond-valence sums in valence units (vu) for bavsiite.

	Ва	V	Si	Σ
O(1)	0.07 ×4		1.04 1.02	2.20
O(2)	0.23 ×4 0.10 ×4	0.62×4	1.08 ×2	2.03
O(3)	(0.04 ×4)	1.59		1.59 (1.75)
Σ	1.60 (1.76)	4.07	4.22	

Bonds for O(1): $2 \times Si + 2 \times Ba$; O(2): $1 \times Si + 1 \times V + 2 \times Ba$; O(3): $1 \times V + (4 \times Ba)$

average 109.5°, indicating a nearly regular tetrahedron. The bond angles of the unbranched $[Si_4O_{12}]^{8-}$ ring are 112.1(3)° for O(1)–Si–O(1)' and 157.9(5)° (= 270° – 112.1°) for Si–O(1)–Si'.

Vanadium occurs in a distorted square pyramidal oxygen environment with equatorial bond length of 1.951(3) Å and a very short axial one of 1.608(5) Å. The displacement of the central V atom off the basal square plane towards the axial oxygen O(3) is indicated by the O(2)–V–O(3) basal to axial oxygen angle of $103.29(8)^{\circ}$ and is also visible in Fig. 4.

Ions of Ba^{2+} occupy channels along c and can be considered as 12-coordinated with Ba–O bond lengths ranging from 2.812(4) to 3.295(2) Å and an average of 3.085 Å. A further but very long Ba–O

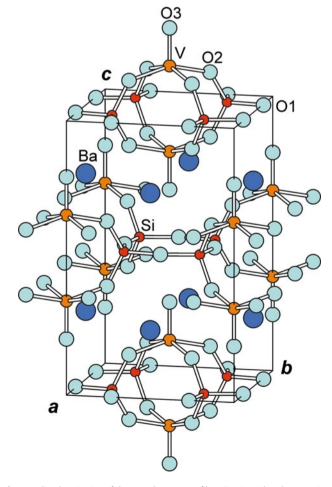


Fig. 4. Inclined projection of the crystal structure of bavsiite. Ba–O bonds are omitted for clarity.

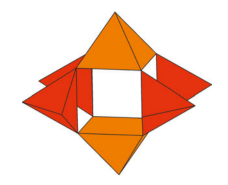


Fig. 5. The cage-like $[Si_4V_2O_{13}]^{12-}$ cluster is a basic structural unit of basisite. The $[SiO_4]$ tetrahedra are coloured red and $[VO_5]$ square pyramids brown.

distance of 3.562(5) Å would sum to a 16-coordination with an even longer average length of 3.205 Å.

Discussion

In the unbranched single $[Si_4O_{12}]^{8-}$ ring the anisotropic parameter U^{33} of the bridging oxygen O(1) is significantly higher than U^{11} and U^{22} generated by the strong bonding to two Si (Si-O(1) = 1.609(5) Å and Si-O(1)' = 1.616(5) Å) in the plane Si-O(1)-Si' parallel to (001) and loosely bonding to two Ba ions (Ba-O(1) = 3.295(2) Å) with c-axis-Ba-O(1) = 29.75° with respect to U^{33} . The silicon tetrahedron in bavsiite is more regular than in β -BaVSi₂O₇ [Si-O(1) = 1.622(5) Å, Si-O(1)' = 1.597(5) Å and $2 \times$ Si-O(2) = 1.603 Å] with Si-O averages of 1.603 Å (bavsiite) and 1.606 Å, respectively. The bond-valence sum in bavsiite for the tetrahedrally coordinated O(1) is high (V = 2.20) whereas the value for O(2) is nearly ideal (V = 2.03). O(2) connects the silicon tetrahedron with the [VO₅]⁶⁻ square pyramid and has an additional two bonds with Ba. Such square pyramids are important structure elements in similar silicates such as β -BaVSi₂O₇, (Liu and Greedan, 1994), suzukiite, BaVSi₂O₇ (Ito *et al.*, 2014), haradaite, SrVSi₂O₇ (Basso *et al.*, 1995) and synthetic

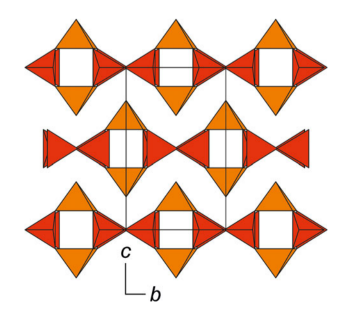


Fig. 6. The cross-linked clusters form a pseudo-two-dimensional network parallel to (001) containing empty channels along the a axis. Ba atoms are omitted for clarity.

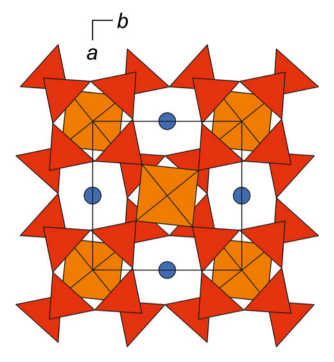


Fig. 7. The crystal structure of bassiite projected along [001] with Ba atoms (blue circles) in the channels along c.

SrVSi₂O₇ (Berger and Range, 1996). In the [VO₅]⁶⁻ square pyramid two types of bonds are visible by their V–O distances: (1) a very short vanadyl bond (shorter than 1.74 Å) and (2) longer equatorial bonds in (V⁴⁺O_n) polyhedra in the range 1.90 to 2.12 Å (Schindler *et al.*, 2000). In bavsiite the vanadyl bond V–(O3) = 1.608(5) Å is longer than in β–BaVSi₂O₇, suzukite and haradaite but nearly equal as in synthetic SrVSi₂O₇: 1.592 (8) Å, 1.593(5) Å, 1.593(9) Å and 1.61(1) Å, respectively. The equatorial bonds in bavsiite and β–BaVSi₂O₇, (1.951(3) Å and 1.936(5) Å, respectively) are shorter than in suzukiite, haradaite and synthetic SrVSi₂O₇ (1.9986(19) Å, 1.990(4) Å and 1.980(5) Å, respectively). In bavsiite the bond-valence sum 1.59 valence units (vu) for O(3) reflects the double bond nature of the vanadyl bond. The nearest further cation bond is to Ba with 3.562(5) Å and would increase the valence value to 1.75 vu for O(3). The tight bonds in the [Si₄V₂O₁₈]^{12–} cluster result in high

The tight bonds in the $[Si_4V_2O_{18}]^{12-}$ cluster result in high valence values for Si and V (4.22 and 4.07 vu, respectively) whereas the Ba atoms in the channels along the c axis are only loosely bonded with valence sums of 1.60 vu (12-coordinated) or 1.76 vu (16-coordinated). The good cleavage in bavsiite parallel to (001), also visible by preferred orientation reflexes (00*l*) in the powder X-ray pattern (Fig. 3), is mainly due to the lengthened Ba–O bonds, which hold together the 2D-network.

According to Strunz and Nickel (2001) bassiite is a cyclosilicate with 4-membered single rings (9.CF.). Bassiite is isotypic to synthetic β -BaVSi₂O₇ (Liu and Greedan, 1994) and polymorphic to suzukiite, BaVSi₂O₇. In the crystal structure of suzukiite (Ito *et al.*, 2014) SiO₄ tetrahedra are connected to corrugated Si₄O₁₂ chains along the *c* axis and VO₅ square pyramids link the chains to form layers parallel to the *ac* plane, which are connected by barium atoms.

Supplementary material. To view supplementary material for this article, please visit https://doi.org/10.1180/mgm.2019.59

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