MINERALOGICAL NOTES

The Unit Cell of Volborthite

ERIK S. LEONARDSEN
Institute of Mineralogy,
OLE V. PETERSEN
Mineralogical Museum,

University of Copenhagen, Denmark

Abstract

Indexed powder diagrams of volborthite, $Cu_3(VO_4)_2 \cdot 3H_2O$, from Monument Valley, Arizona, disclose it to be monoclinic (space group C2, Cm, or C2/m), the unit cell [a=10.604(2), b=5.879(1), c=7.202(2) Å, $\beta=94.81(2)^\circ]^1$ containing two formula units. Calculated density, 3.52 g/cm³, compares with 3.42 g/cm³, the value measured by Guillemin (1956).

Introduction

In the Mineralogical Museum, Copenhagen, Guinier-Hägg powder diffraction exposures have formed part of the routine examination of newly acquired mineral specimens since 1971. Such an examination of volborthite from Monument No. 1 Mine, Monument Valley, Navajo County, Arizona, permitted indexing of the pattern and thus, for the first time, determination of its unit cell. The specimen (M.M. No. 1972.307), according to Witkind (1961), originates from the outermost zone of oxidized minerals of the ore body, the mineral assemblage of which includes carnotite, hewettite, metatyuyamunite, rauvite, tyuyamunite, and volborthite. The specimen consists of a light gray siltstone matrix profusely covered on one side with olive green to citron yellowish-green microcrystalline globules of volborthite, some of which are themselves partly covered with numerous small crystals.

X-Ray Diffraction Data

Guillemin (1956) published an unindexed powder pattern of volborthite, which is reproduced by Joint Committee of Powder Diffraction Standards, card no. 12–523. No other diffraction data were found.

Our pattern is in excellent agreement with this standard, except for some splitting up of the lines due to the Guinier-technique. The pattern was readily indexed by the method of Ito (1950). Least squares refinement gave the following unit cell dimensions: $a_0 = 10.604 \pm 0.002$ Å, $b_0 = 5.879 \pm 0.001$ Å, $c_0 = 7.202 \pm 0.002$ Å, and $\beta = 94.81 \pm 0.02^{\circ}$.

TABLE 1. X-Ray Diffraction Data for Volborthite

JCPDS, card no. 12-523		Present investigation XX				
Int.	d (Å)	Int.	d _{obs} .	10 ⁴ Q _{obs.} (Å ⁻²)	10 ⁴ Q _{calc} .	hkl
100	7.26	10	7.16	195.0	194.2	001 200
20	5.21	3 4	5.291 5.136 4.426	357.2 379.1 510.3	358.2 378.9 508.1	110 20T
10	4.19×		1,81.20	3.003	550.9	111
10	4.03	5	4,103	594.1	595.2 596.6	111
10	3.56	1	3.583	779.0	776.6	002
30	3.08	5	3.090	1047.1	1046.3	202
40	2.98		3.022 2.998	1034.9	1095.4	310 112
		5	2.940	1157.3	1157.3	020
80	2.87	5	2.887	1199.8	1199,8 \$ 1223,1	112
		5	2.859	1223.5	1223.4	202
20	2.71	5	2,722	1349.7	1 1351.4	021
20	2.64	7	2.643	1432.1	1432,9	311
80	2.56	7	2.571	1513.3	1515.5	220
60	2.50	5	2,551	1537.2	1538.6	401 221
					1665.4 1715.6	401
					1739.2	312
80	2.38	7	2,389	1752.2	1747.4 1753.9	003 221
	Z + 30	,	2.309	1734.6	1933.9	022
					1972.9	203
30 10	2.21	1	2.216	2036.3	2004.7	312 402
	A . A 1		2,210	20,000	2059.9	113
	2.12×				2192.7	113
	2,12				2238.4	203
30	2.04	Ł.	2.049	2381.9	2380.7	222
			1		2386.6	402 510
					2590.2	420
					2611.8 2643.6	511 313
					2693.4	1 30
		1	1.9247	2699.3	2695.9 2833.1	421
		1	1.8688	2863.3	2865.5	511 131
		4.6			2872.9	421
					2904.7 2909.7	023
10	1.85	1	1,8521	2915.1	2914.8	40
		1	1.8021	3079.2	3041.9 3083.6	313 512
30	1.78	2	1.7946	3105.0	3106.5	004
30	1.70	3	1.7876	3129.2	3130.2	22
			1.7703	3191.0	3189.8 3224.1	422 600
					3285.5	601
					3287.7 (3395.7	204
		3	1.7170	3392.2	3396.9	221 114
30	1.71		* 7004	al or -	3409.9	330
		2	1.7081	3427.7	3425.8 3445.9	132
			44		3514.3	132
30	1.67		1,6838	3527.1	3526.2	512
					3537-7 3543-9	231

^{*} Lines in position of strong quartz lines.

¹ Numbers in parentheses represent estimated standard deviation in respect to the last decimal cited. (See Am. Mineral. 59, 223.

xx Pattern obtained with Guinier Hägg focusing camera, radius 40.3 mm. $CuK\alpha_I$ ($\lambda = 1.54051$ Å) radiation, Nifilter, and curved crystal monochromator. Internal standard quartz. Intensities visually estimated. All possible Q-values below $Q_{cale} = 3550.0$ have been calculated.

The systematic reflection condition led to the space-groups C2, Cm, or C2/m. Multiple indexed lines were omitted from the least squares refinement. The formula $Cu_3(VO_4)_2 \cdot 3H_2O$, unit cell volume = 447.42 A^3 , and Z=2, gives the calculated density $D_x=3.52 \text{ g/cm}^3$. Guillemin (1956) has measured the density as 3.42 g/cm^3 .

Acknowlegments

The authors wish to thank Mrs. G. Sjørring who typed the table.

References

Guillemin, C. (1956) Contribution a la Minéralogie des Arséniates Phosphates et Vanadates de Cuivre. Bull. Soc. franc. Minéral. Cristallogr. 79, 219-275.

Ito, T. (1950) X-ray Studies on Polymorphism. Tokyo, Maruzen.

WITKIND, I. J. (1961) The uranium-vanadium ore deposit at the Monument No. 1—Mitten No. 2 Mine, Monument Valley, Navajo County, Arizona. U.S. Geol. Surv. Bull. 1107-C.

Manuscript received, July 23, 1973; accepted for publication, October 30, 1973.