WARDSMITHITE, 5CaO·MgO·12B₂O₅·30H₂O, A NEW BORATE MINERAL FROM THE DEATH VALLEY REGION, CALIFORNIA


ABSTRACT

Wardsmithite, 5CaO·MgO·12B₂O₅·30H₂O, is a new borate found at two localities in the Death Valley region, Inyo County, California, on weathered veins of colemanite or priceite in the Furnace Creek Formation (Pliocene). It occurs as nodules and coatings in gowerite, ulexite, and colemanite.

The mineral is hexagonal (or pseudohexagonal) with a platy habit and cleavage {0001}. Typically, it occurs as aggregates of subparallel plates to 75 μm in diameter, uncommonly as single crystals to 15 μm in diameter. The strongest lines in the X-ray powder diffraction pattern (unindexed) are, in Å: 13.5 (100), 12.3 (62), 6.12 (55), 3.358 (51), 4.721 (42), 2.744 (26).

Infrared absorption peaks (in cm⁻¹) are 790, 810, 890, 1000, 1070, 1335, and 1615.

Crystals of wardsmithite are colorless. Optically it is uniaxial (−), with α = 1.490 ± 0.002, ε = 1.476 ± 0.002. Hardness is 2½; specific gravity, 1.88 ± 0.02.

Chemical analysis gave, in weight percent, B₂O₅ 48.58, CaO 16.50, MgO 2.26, H₂O 32.44, sum 99.78. Spectroscopic analysis showed Si, Al, Na, Mn, Ba, Cu, and Pb in small amount.

Wardsmithite is named after Ward C. Smith, geologist of the U.S. Geological Survey.

INTRODUCTION

Further study of the borate minerals produced by weathering of priceite and colemanite in the Furnace Creek Formation shows that the mineral referred to as an "undescribed calcium borate" mineral associated with nobleite (Erd et al., 1961, p. 561) is the first known natural occurrence of 5CaO·MgO·12B₂O₅·30H₂O, wardsmithite. The new mineral is named after Ward C. Smith, geologist of the U.S. Geological Survey, in recognition of his advancement of borate studies in southwestern United States.

Samples containing a small quantity of the mineral were collected in 1956 by McAllister while mapping the geology of the Furnace Creek borate area in the Death Valley region, California, for the U.S. Geological Survey in cooperation with the California Division of Mines and Geology. Preliminary determination of some optical and chemical properties of the mineral by him in 1959 seemed to indicate an undescribed calcium borate; and X-ray diffraction data obtained by Erd were con-
clusive for such a borate. Another occurrence in the area was found by Erd in 1961. Enough wardsmithite for chemical analysis was separated from samples collected in 1965 from the first locality. Type material will be deposited in the U.S. National Museum, Washington, D. C.

Occurrence

Rare and inconspicuous among the efflorescent borate minerals, wardsmithite has been found at two localities about 2 miles apart. The first locality, on the Hard Scramble patented mining claim of the United States Borax & Chemical Corporation, is on the western slope of low foothills of the Black Mountains, 1.9 miles N. 72° W. from Ryan. The second occurrence, which is on the north bank of the wash running northwest from U. S. mineral monument 47 to the junction of California Highway 190 and the road to Dantes View, lies 2.1 miles N. 17° W. from Ryan. It is on the old Neglect claim, which was merged with the Oliver claim when patented as the Oliver Consolidated claim by the United States Borax & Chemical Corporation.

At both places the wardsmithite is on weathered veins of colemanite or priceite in deformed Furnace Creek Formation of Pliocene age. The colemanite veins on the Neglect claim are in lacustrine gypsiferous mudstone and standstone derived from siliceous volcanic rocks and closely associated with some algal limestone. The priceite veins on the Hard Scramble claim are in altered fragmental basalt, which is stratigraphically below the gypsiferous zone and considerably above the zone of commercial borate deposits in the formation.

Wardsmithite at the main occurrence on the Hard Scramble claim is considerably rarer than gowerite, nobleite, microgranular colemanite, and particularly ulexite. These minerals and some ginorite, hungchaoite, rivadavite, macallisterite, kurnakovite, inderite, thenardite, gypsum, and calcite, in various proportions, form aggregates near the surface, in continuation with priceite and colemanite veins below them. A coarse boxwork of calcite or fine-grained colemanite remains on some outcrops. Some of the aggregates enclose remnants of the priceite veins and the transition downward from efflorescent minerals to priceite alone is complete within several tens of centimeters below the surface. The wardsmithite, in exceedingly fine-grained, soft clots, lies on and fills in between gowerite blades radiating from compact cores, and it lies on tufts of ulexite in a characteristic specimen. Some of this ulexite is on the gowerite, and both are attached to solid aggregates of nobleite.

The scanty amount of wardsmithite collected at the Oliver Consolidated locality occurs as very fine-grained nodules, commonly as much as 2 mm in diameter, and irregular coatings on a specimen of coarse cole-
manite and gypsum. The colemanite is anhedral or subhedral; the largest subhedral crystal noted is about 15 mm by 5 mm. Abundant fine-grained gypsum that fills fractures and incrusts surfaces is associated with some fine-grained celestite, meyerhofferite, and inoyoite and microcrystals of colemanite. The wardsmithite formed on surfaces of corroded colemanite and between small spherules of fine-grained ulexite, the most abundant efflorescent borate, on crusts of gypsum.

Crystallography

X-ray data. The X-ray data for wardsmithite are reported in Table 1. No variation in patterns was found for various samples of the mineral from the two localities. Crystals are too small to permit investigation of the crystallography of the unit cell by single-crystal techniques. Considering the optical evidence for hexagonal symmetry, knowing the approximate volume of the unit cell for one formula molecular weight (or multiples of this), and equating either of two sets of spacings with enhanced intensities with the "basal" spacing, we were still unable to index the X-ray powder data on the basis of a hexagonal cell. Attempts to obtain an electron diffraction pattern were also unsuccessful, as grains small enough to be useful in selected-area electron diffraction decomposed in the beam before a pattern could be obtained. We conclude that ward-

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Plus additional lines all with $I<5$

Split of analyzed sample. X-ray diffractometer data, Chart X3205: Cu/Ni radiation, $\lambda_{CuK\alpha}=1.54051$ Å; silicon used as internal standard; scanned at $\frac{1}{4}$° per minute from 2–65° 2θ.

Table 1. X-Ray Diffraction Data for Wardsmithite
smithite is pseudohexagonal and that the true symmetry is monoclinic or lower. Until the true symmetry is established, it is convenient to describe the properties of wardsmithite with reference to a hexagonal cell.

**Morphology.** Wardsmithite crystals are anhedral to subhedral with a hexagonal outline (an individual grain with interfacial angles very near to 60° is shown in the electronmicrograph, Fig. 1) and a platy habit {0001}.

![Fig. 1. Electron micrograph of wardsmithite crystal showing hexagonal outline. Aggregates of smaller crystals of wardsmithite appear as nearly opaque grains on the surface of the large crystal.](image)

Under the microscope, wardsmithite typically appears as aggregates to 75μm in diameter composed of subparallel plates and, less commonly, as single crystals up to 15μm in diameter and 2μm in thickness.

**Physical and Optical Properties**

There appears to be a good cleavage parallel to the "base" {0001}. The hardness (2½) was determined by rubbing wardsmithite on clear surfaces of gypsum and calcite and examining the surfaces under the microscope. The specific gravity, determined by the sink-float method in bromoform-acetone mixtures checked with a Westphal balance, is 1.88±0.02. By a pycnometric method (following Fahey, 1961), the specific gravity of a 400 mg sample was found to be 1.85 at 25°C. There
is excellent agreement between the measured specific gravity (with the value 1.88 giving a better fit), indices of refraction, and chemical composition using the Gladstone-Dale relationship (see Jaffe, 1956). Aggregates of the mineral are white (Munsell color N9) and have a vitreous luster. The mineral is not fluorescent.

Wardsmithite is colorless in transmitted light. It is uniaxial negative (or biaxial negative with a very small 2V) with \( \omega = 1.490 \pm 0.002 \) and \( \epsilon = 1.476 \pm 0.002 \). Hexagonal plates show no birefringence on basal surfaces and have straight extinction when viewed edge-on. Dispersion could not be determined.

**Chemical Properties**

*Analysis.* A 450 mg sample of the mineral from the Hard Scramble locality was obtained by separation in bromoform-acetone mixtures. About 3 percent gowerite, which could not be mechanically or chemically separated, was present in the sample analyzed. This was the only impurity noted optically; no lines attributable to gowerite are present in the X-ray powder pattern of the analyzed sample. The X-ray powder patterns of samples from the two localities are identical; this, together with the optical evidence, strongly suggests that only one phase is present.

A split of the sample that weighed 200 mg was dissolved in 30 ml of 1 N HCl. This solution was adjusted to a pH of 2.5 and then passed through an ion exchange column in order to separate \( \text{B}_2\text{O}_3 \) from the cations (following Martin and Hayes, 1952). Borate was determined by titration in the usual way with standard sodium hydroxide after addition of mannite to the neutralized borate solution. The cations were recovered from the column and determined by the standard methods of quantitative analysis outlined by Hillebrand *et al.* (1953). Water was determined by heating a 150 mg sample to 500°C in a thermostatically-controlled electric furnace and then measuring the loss of weight of the sample. It was found that at higher temperatures (about 700°C) loss of \( \text{B}_2\text{O}_3 \) resulted. The results of the analysis are shown in Table 2. The empirical formula \( 5\text{CaO} \cdot \text{MgO} \cdot 12\text{B}_2\text{O}_3 \cdot 30\text{H}_2\text{O} \) is suggested as the simplest formula containing one atom of Mg, as substitution of Mg for Ca in borates seems to be very limited. The correct structural formula must remain unknown until larger crystals are found that will permit determination of the crystal structure.

*Synthesis.* We did not attempt the synthesis of wardsmithite. A compound close to this composition, \( 4\text{CaO} \cdot 2\text{MgO} \cdot 12\text{B}_2\text{O}_3 \cdot 30\text{H}_2\text{O} \) (her formula doubled for comparison), was prepared by Kurnakova (1953, p. 57–60) by a reaction between inderborite and an aqueous solution of boric acid at 25°C that required 297 days for completion. This compound, plus impurities, was also obtained in 215 days at 25°C by a reaction between aqueous solutions of boric acid, \( \text{Ca(OH)}_2 \), and \( \text{Mg(OH)}_2 \). The only properties given for this compound are the indices of refraction, \( \gamma = 1.538 \) and \( \alpha = 1.486 \), determined by E. E. Burovaya (Kurnakova, 1953). An approximate value for the density, \( 1.98 \pm 4 \text{ g/cm}^3 \), can be calculated from the optical data and the chemical composition using the Gladstone-Dale relationship.

There are no compounds of the type \( \text{MO} \cdot 2\text{B}_2\text{O}_3 \cdot 5\text{H}_2\text{O} \) (wardsmithite
### Table 2. Chemical Analysis of Wardsmithite

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<th>Weight percent²</th>
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² A. C. Vlisidis, analyst. Sample W-16693 (USGS) from the Hard Scramble claim. Spectroscopic analysis by J. L. Harris also showed the following elements, in weight percent: Si, 0.05; Al, <0.001; Na, 0.15; Mn, 0.007; Ba, 0.0005; Cu, 0.02; and Pb, 0.001. For limits of sensitivity see Bastron et al. (1960).

³ Recalculated to 100 percent after subtracting 3 percent gowerite.

⁴ For $5CaO \cdot MgO \cdot 12B₂O₃ \cdot 30H₂O$.

composition in simplest ratio) known in the separate systems CaO-B₂O₃-H₂O or MgO-B₂O₃-H₂O. The synthetic compound $CaO \cdot 2B₂O₃ \cdot 4H₂O$ was reported by Ditte (1883, p. 1668) and the names bechilite and borocalcite have been used for minerals said to have this composition. These “minerals” have been found to be mixtures, and there is no modern confirmation of Ditte’s synthesis (see summaries in Palache et al., 1951, p. 345, 347, 365; Nies and Campbell, 1964, p. 127).

**Solubility.** Wardsmithite is insoluble in methyl alcohol, is very slightly soluble in cold water, slightly soluble in hot water, and readily soluble in cold dilute acids.

**Pyrognostics.** Heated in a closed tube, wardsmithite gives off water, sinters, and then fuses to a gray mass which, with continued heating, becomes a clear glass. There was insufficient sample available for thermal analyses.

**Infrared Absorption Analysis**

Absorption spectra of wardsmithite and of an unidentified dehydration(?) product were prepared by Dr. C. E. Weir and are shown in Figure 2. Two different specimens of wardsmithite were run under three different conditions and all the results agreed within experimental error.
Fig. 2. Infrared spectra of wardsmithite (A) and of an unidentified dehydration(?) product (B) in the region 1800-400 cm⁻¹. C. E. Weir, analyst.

This further demonstrates the homogeneity of the wardsmithite samples used in this study. Infrared absorption peaks (in cm⁻¹) are 790, 810, 890, 1000, 1070, 1335, and 1615. Weir (personal commun., 1966) notes the presence of both boron tetrahedra and triangles in wardsmithite. Tetrahedra are indicated by strong absorption with broad bands below 1000 cm⁻¹, which is characteristic of 4-fold boron, triangles by the broad band at 1335 cm⁻¹. The weak band near 1615 cm⁻¹ is assigned to H-O-H bending. Further interpretation is not possible; the difficulties involved in the interpretation of the infrared spectra of the hydrated borates have been pointed out by Weir (1966, p. 163–164).

The spectra of wardsmithite and the unidentified dehydration(?) product are very similar except that the bands may be shifted slightly and show less band structure for the unidentified material. Also the band near 1615 cm⁻¹ is more shallow for the unidentified material, strengthening the suggestion that this is a dehydration product of wardsmithite. Further description of this material is given below.

**Unidentified Dehydration(?) Product**

An unidentified borate mineral was found in one sample of material from the Hard Scramble locality. It appeared to be a small clump of wardsmithite, about 5 mm in diameter, but having a duller luster. Under
the microscope, however, it was colorless and isotopic with $n = 1.502 \pm 0.002$. There were many anhedral grains less than 1µm in diameter, some bearing a resemblance to rounded hexagonal plates of wardsmithite, but most of the material consisted of aggregates of about the same size range as wardsmithite. The only impurity present was gowerite, less than one percent.

The isotropic borate is amorphous to X-rays. The infrared spectra (Fig. 2), however, are similar to those of wardsmithite and are discussed above. Insoluble or only slightly soluble in cold water, the material is slightly to moderate soluble in hot water and recrystallizes in part to minute rhomb-shaped euhedra having high birefringence and in part to amorphous material at the edge of the drop. The amount of material so dissolved and reprecipitated is too small to permit identification, however. The isotropic borate is easily soluble in cold dilute acids. Addition of a drop of $\text{H}_2\text{SO}_4$ to a solution of the mineral in dilute $\text{HCl}$ gives, on evaporation, crystals of gypsum and sassolite. In a closed tube, the mineral gives off a slight amount of water, then sinters and fuses in the same manner as wardsmithite. The small amount of material precludes further tests, and data are insufficient to characterize this as a distinct mineral species. From the relationship with gowerite, appearance, infrared spectra, and chemical tests, this mineral appears to be a dehydration product of wardsmithite—the first essentially amorphous borate found in nature so far as we are aware. The amorphous material was sought for in other samples of wardsmithite (especially in the sample used for the chemical analysis), but none was found.

Acknowledgments

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References


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