A vibrational spectroscopic study of the borate mineral takedaite Ca$_3$(BO$_3$)$_2$

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**Abstract**

We have studied the mineral takedaite Ca$_3$(BO$_3$)$_2$, a borate mineral of calcium using SEM with EDX and vibrational spectroscopy. Chemical analysis shows a homogeneous phase, composed of Ca. Boron was not detected. A very intense Raman band at 1087 cm$^{-1}$ is assigned to the BO stretching vibration of BO$_3$ units. Additional Raman bands may be due to isotopic splitting. In the infrared spectrum, bands at 1218 cm$^{-1}$ and at 1163, 1262 and 1295 cm$^{-1}$ are assigned to the trigonal borate stretching modes. Raman bands at 712 and 715 cm$^{-1}$ are assigned to the in-plane bending modes of the BO$_3$ units. Vibrational spectroscopy enables aspects of the molecular structure of takedaite to be assessed.

**Keywords:** Takedaite, Borate, Calcium, Raman spectroscopy, Infrared spectroscopy

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**Introduction**

The mineral takedaite Ca$_3$(BO$_3$)$_2$ is the borate mineral of calcium [1]. It is one of several minerals found at Fuka, Okayama Prefecture, Japan [2]. Takedaite is a rare mineral probably formed by boron-rich fluids reacting between crystalline limestone and gehlenite–spurrite skarns [3,4]. Crystal structure has been refined [5]. The mineral is hexagonal with Point Group: 3 2/m and Space Group: R3/c. The unit cell parameters are: $a = 8.638$ Å, $c = 11.85$ Å, $Z = 6$; $V = 765.73$ Å$^3$. The boron isotope ratio (11B/10B) is used to determine the geochemistry of boron minerals including hydroboracite [6–8]. This isotopic ratio may be reflected in the vibrational spectra providing the resolution of the spectrometers is sufficiently high.

Vibrational spectroscopy has been applied to borate glasses [9–12]. However, the number of vibrational spectroscopic studies of borate minerals is few and far between. The number of vibrational spectroscopic studies of borate minerals is quite few and far between [13–16]. The number of Raman studies of borate minerals...
is also very limited [17,18]. There have been a number of infrared studies of some natural borates [19–22]. Most of these references are not new and there have been no recent studies on the vibrational spectroscopy of natural borates. Ross in Farmer’s treatise reported the infrared spectra of several borate minerals [23].

There have been only a few infrared studies of hydroboracite [19,22,24,25] and almost no studies of the Raman spectrum of takedaite [26]. There is some disagreement between researchers as to the position of some selected infrared bands. Moenke [27] and Vlasova and Valyashko [25] show a band at 675 cm\(^{-1}\) which was not observed in the infrared spectrum of Weier [22]. These authors simply reported the infrared data without any assignment of the bands.

Vibrational spectroscopy has proven most useful for the study of borate minerals [28–31]. There are many calcium borate minerals; many contain water and/or hydroxyl units in their structure. There is a vital need to study these calcium borate minerals using vibrational spectroscopy. In the authors view, there is an apparent lack of fundamental knowledge of borate mineral spectroscopy. The authors have made several studies of boron containing minerals [32–35]. The objective of this research is to report the Raman and infrared spectra of takedaite and to relate the spectra to the molecular structure of the mineral. This is the first report of a systematic study of takedaite from Fuka, Japan. It is important to understand the structure of takedaite in order to form nanomaterials based upon its structure. Vibrational spectroscopy enables a better understanding of the structure of takedaite.

**Experimental**

**Samples description and preparation**

The takedaite sample studied in this work forms part of the collection of the Geology Department of the Federal University of Ouro Preto, Minas Gerais, Brazil, with sample code SAB-096. The sample is from the Fuka mine, located in the Okayama Prefecture, Japan. The studied sample occurs as a massive aggregate of micrometric crystals.

The sample was prepared by using of a stereomicroscope Leica MZ4. The takedaite studied in this work occurs as a pure phase. Scanning electron microscopy (SEM) equipment was applied to support the mineral characterization.

**Scanning electron microscopy (SEM)**

Experiments and analyses involving electron microscopy were performed in the Center of Microscopy of the Universidade Federal de Minas Gerais, Belo Horizonte, Minas Gerais, Brazil (http://www.microscopia.ufmg.br). Takedaite crystals were coated with a 5 nm layer of evaporated carbon. Secondary Electron and Backscattering Electrons images were obtained using a JEOL JSM-6360LV equipment. Qualitative and semi-quantitative chemical analyses in the EDS mode were performed with a ThermoNORAN spectrometer model Quest and were applied to support the mineral characterization.

![Fig. 1. Backscattered electron image (BSI) of a takedaite fragment up to 2.0 mm in length.](image1)

![Fig. 2. EDS spectra of takedaite.](image2)

![Fig. 3. (a) Raman spectrum of takedaite over the 100–4000 cm\(^{-1}\) spectral range. (b) Infrared spectrum of takedaite over the 500–4000 cm\(^{-1}\) spectral range.](image3)
**Raman microprobe spectroscopy**

Crystals of takedaite were placed on a polished metal surface on the stage of an Olympus BHSM microscope, which is equipped with 10×, 20×, and 50× objectives. The microscope is part of a Renishaw 1000 Raman microscope system, which also includes a monochromator, a filter system and a CCD detector (1024 pixels). The Raman spectra were excited by a Spectra-Physics model 127 He–Ne laser producing highly polarized light at 633 nm and collected at a nominal resolution of 2 cm⁻¹ and a precision of ±1 cm⁻¹ in the range between 200 and 4000 cm⁻¹. Repeated acquisitions on the crystals using the highest magnification (50×) were accumulated to improve the signal to noise ratio of the spectra. Raman Spectra were calibrated using the 520.5 cm⁻¹ line of a silicon wafer. The Raman spectrum of at least 10 crystals was collected to ensure the consistency of the spectra.

**Infrared spectroscopy**

Infrared spectra were obtained using a Nicolet Nexus 870 FTIR spectrometer with a smart endurance single bounce diamond ATR cell. Spectra over the 4000–525 cm⁻¹ range were obtained by the co-addition of 128 scans with a resolution of 4 cm⁻¹ and a mirror velocity of 0.6329 cm/s. Spectra were co-added to improve the signal to noise ratio.

Spectral manipulation such as baseline correction/adjustment and smoothing were performed using the Spectracalc software package GRAMS (Galactic Industries Corporation, NH, USA). Band component analysis was undertaken using the Jandel 'Peakfit' software package that enabled the type of fitting function to be selected and allows specific parameters to be fixed or varied accordingly. Band fitting was done using a Lorentzian–Gaussian cross–product function with the minimum number of component bands used for the fitting process. The Gaussian–Lorentzian ratio was maintained at values greater than 0.7 and fitting was undertaken until reproducible results were obtained with squared correlations of r² greater than 0.995.

**Results and discussion**

**Chemical characterization**

The SEM image of the mineral sample takedaite studied in this work is shown in Fig. 1. The image shows a takedaite microcrystalline aggregate. Qualitative chemical analysis shows a homogeneous phase, composed of Ca. Carbon was also observed and occurs due to C coating (Fig. 2). Boron was not detected as the atomic number is low.

**Vibrational spectroscopy**

The Raman spectrum of takedaite over the 100–4000 cm⁻¹ spectral range is shown in Fig. 3a. This spectrum shows the position and intensity of the Raman bands. The observed bands are very sharp. No intensity above 1700 cm⁻¹ is observed. The Raman spectrum may be subdivided into sections based upon the type of vibration being analyzed. The infrared spectrum of takedaite over the 500–4000 cm⁻¹ spectral range is illustrated in Fig. 3b. In comparison with the Raman spectrum the bands are broad and overlap. Now in the infrared spectrum, some intensity is observed in the 3000–4000 cm⁻¹ spectral range.

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![Fig. 4.](image1.png)  (a) Raman spectrum of takedaite over the 800–1300 cm⁻¹ spectral range. (b) Infrared spectrum of takedaite over the 500–1600 cm⁻¹ spectral range.

![Fig. 5.](image2.png)  (a) Raman spectrum of takedaite over the 300–800 cm⁻¹ spectral range. (b) Raman spectrum of takedaite over the 100–300 cm⁻¹ spectral range.
The Raman spectrum over the 800–1300 cm\(^{-1}\) spectral range is reported in Fig. 4a. The spectrum is dominated by an intense Raman band at 1087 cm\(^{-1}\). This band is assigned to the BO stretching vibration of BO\(_3\) units. Some low intensity shoulder bands are noted at 1085 and 1091 cm\(^{-1}\). Iliev et al. determined the Raman spectrum of a synthetic cobalt boracite\[36\]. The symmetry species of some vibrational modes were determined. Iliev et al.\[37\] used Raman imaging to show the ferroelectric properties of boracite type compounds. These workers\[37\] showed that boracites exhibit a sequence of transitions from the high temperature paraelectric cubic phase to ferroelectric orthorhombic, monoclinic, trigonal phases, and finally to a monoclinic phase at low temperatures where both ferroelectric and magnetic orders coexist. Kim and Somoano determined the improper ferroelectric transition using Raman spectroscopy\[38\].

In contrast, the infrared spectrum of boracite (Fig. 4b) shows complexity with many overlapping bands, making band attribution difficult. The intense infrared band at 1218 cm\(^{-1}\) with shoulders at 1163, 1262 and 1295 cm\(^{-1}\) is assigned to the trigonal borate stretching modes. The overlapping infrared bands are the infrared BO antisymmetric stretching vibrations. According to Ross [23] (page 220 of this reference), bands between 1300 and 1500 cm\(^{-1}\) are due to the antisymmetric stretching modes of trigonal boron. This is perhaps confirmed by the intensity of the infrared bands in the 1300–1500 cm\(^{-1}\) region. The series of bands from 600 through to 800 cm\(^{-1}\) are related to trigonal borate bending modes. Infrared bands are observed at 677, 694, 707, 712, 728, and 742 cm\(^{-1}\). The infrared bands at 1345, 1403 and 1472 cm\(^{-1}\) are related to the out-of-plane OBO bending modes.

The mineral takedaite has the formula Ca\(_3\)(BO\(_3\))\(_2\) and as such contains no water or hydroxyl groups. The Raman spectrum of takedaite over the 300–800 cm\(^{-1}\) spectral range is shown in Fig. 5a. The Raman spectrum of takedaite over the 100–300 cm\(^{-1}\) spectral range is shown in Fig. 5b. Intense Raman bands are observed at 712 and 715 cm\(^{-1}\) and are assigned to the in-plane bending modes of the BO\(_3\) units. A low intensity Raman band is found at 585 cm\(^{-1}\). A very intense Raman band is observed at 282 cm\(^{-1}\) with shoulders at 276 and 287 cm\(^{-1}\). These bands are attributed to CaO stretching vibrations. The Raman bands at 154 and 159 cm\(^{-1}\) are simply described as lattice vibrations.

The infrared spectrum over the 2600–3800 cm\(^{-1}\) spectral range is given in Fig. 6b. As for the Raman spectrum, the spectrum is very noisy and suffers from a lack of signal. Infrared bands may be resolved at 3280, 3353, 3492 and 3608 cm\(^{-1}\). The infrared spectrum bears a strong resemblance to the Raman spectrum.

The Raman spectrum of takedaite over the 1300–2300 cm\(^{-1}\) spectral range is shown in Fig. 7a. Two intense Raman bands are found at 1436 and 1439 cm\(^{-1}\). These bands are assigned to the antisymmetric stretching vibrations of trigonal boron. These bands
may be due to isotopic splitting. These bands are assigned to the
antisymmetric stretching vibrations of trigonal boron [23] (please
see Table 11.VIII of this reference). Two low intensity Raman bands
are noted at 1556 and 1590 cm⁻¹. Two intense Raman bands are
observed at 1749 and 1753 cm⁻¹ with a shoulder band at
1797 cm⁻¹. These bands are also attributed to the antisymmetric
stretching vibrations of trigonal boron [23]. In the case of the infra-
red spectrum (Fig. 7b), only one infrared band at 1795 cm⁻¹ is
observed. No water bending modes were found. Low intensity
Raman bands at 2148, 2174 and 2199 cm⁻¹ are observed. These
bands may be associated with overtone or combination bands.

Conclusions

There are many borate minerals. Most are either hydrate or con-
tain hydroxyl groups or both. Many of these minerals have never
been analyzed using vibrational spectroscopy. In this instance,
we have studied the anhydrous borate mineral takedaite. Chemical
analysis shows a homogeneous phase, composed by Ca. Vibrational
spectroscopy proves the absence of either water or hydroxyl units
in the structure. The spectra are characterized by a trigonal borate
with intense Raman bands at 712, 715 and 1087 cm⁻¹. Raman
spectroscopy allows the possibility of distinguishing takedaite from
the other boron containing calcium minerals.

Acknowledgements

The financial and infra-structure support of the Discipline of
Nanotechnology and Molecular Science, Science and Engineering
Faculty of the Queensland University of Technology, is gratefully
acknowledged. The Australian Research Council (ARC) is thanked
for funding the instrumentation. The authors would like to
acknowledge the Center of Microscopy at the Universidade Federal
de Minas Gerais (http://www.microscopia.ufmg.br) for providing
the equipment and technical support for experiments involving
electron microscopy.

References

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