Structural characterization of the borate mineral inyoite – CaB₃O₃(OH)₅·4(H₂O)

Ray L. Frost a,⇑, Andrés López a, Ricardo Scholz b, Frederick Theiss a, Geraldo Magela da Costa c

a School of Chemistry, Physics and Mechanical Engineering, Science and Engineering Faculty, Queensland University of Technology, GPO Box 2434, Brisbane, Queensland 4001, Australia
b Geology Department, School of Mines, Federal University of Ouro Preto, Campus Morro do Cruzeiro, Ouro Preto, MG 35400-00, Brazil
c Chemistry Department, Federal University of Ouro Preto, Campus Morro do Cruzeiro, Ouro Preto, MG 35400-00, Brazil

HIGHLIGHTS

• We have studied the mineral Ca(H₄B₃O₇)(OH)₄·4(H₂O) or CaB₃O₃(OH)₅·4(H₂O).
• Using a range of techniques including XRD, SEM, EDX, TG and vibrational spectroscopy.
• Both tetrahedral and trigonal boron units are observed.
• Bands due to the isotopes of boron are observed.
• Aspects of the molecular structure of inyoite are determined.

ARTICLE INFO

Article history:
Received 28 July 2014
Received in revised form 23 September 2014
Accepted 27 September 2014
Available online 5 October 2014

Keywords:
Inyoite
Borate
Calcium
Raman spectroscopy
Infrared spectroscopy

ABSTRACT

We have studied the mineral Ca(H₄B₃O₇)(OH)₄·4(H₂O) or CaB₃O₃(OH)₅·4(H₂O) using electron microscopy and vibrational spectroscopy. The mineral has been characterized by a range of techniques including X-ray diffraction, thermal analysis, electron microscopy with EDX and vibrational spectroscopy. Electron microscopy shows a pure phase and the chemical analysis shows the presence of calcium only. The nominal resolution of the Raman spectrometer is of the order of 2 cm⁻¹ and as such is sufficient enough to identify separate bands for the stretching bands of the two boron isotopes. Raman and infrared bands are assigned to the stretching and bending modes of trigonal and tetrahedral boron and the stretching modes of the hydroxyl and water units. By using a combination of techniques we have characterized the borate mineral inyoite.

Introduction

Borate minerals are an important supplement for different industries [1]. Boron is found in borates of metals, especially of calcium and sodium. Important deposits of boron ores are located in Turkey, mainly in Anatolia [2]; California, USA [3]; Argentina [4] and Bolivia [5]. Borate brine deposits are composed by a complex association of minerals. The mineral inyoite is a hydrated borate mineral of sodium with the formula Na₂B₄O₆(H₂O)₂. It was described as a new mineral by Schaller from the Rich Station, Kramer Borate deposit, Boron, Kramer District, Kern Co., California, USA [6]. Other occurrences were reported from Argentina [7] and Turkey [8].

The mineral occurs as prismatic to tabular crystals and crystalizes with monoclinic symmetry. Space Group is P2₁/a with a = 10.63(2)Å, b = 12.06(2)Å, c = 8.405(2)Å, β = 114°02´ with Z = 4. Inyoite contains the same isolated [B₃O₃(OH)₅]²⁻ polyions that were found in meyerhoffertite and in the synthetic, CaB₃O₅(OH)₅·2H₂O. Such a polyion is formed by two BO₄ tetrahedra sharing a corner and one BO₃ triangle linking the two tetrahedra. Polyions of inyoite are connect to another and to neighboring water molecules by bonding through calcium ions and by hydrogen bonds [9].

There are many borate containing minerals which have yet to have their vibrational spectra measured and the molecular structure assessed in terms of their vibrational spectra. The importance of the mineral inyoite rests with the chemistry of the compound and the potential to synthesize nanomaterials based upon polymerization of borate units. Such compounds have the potential to make especially selected ferroelectric, pyroelectric and piezoelectric properties. The mineral inyoite is a precursor for the synthesis of such nanomaterials. Spectroscopy methods are important tools in the study of complex mineral associations. The objective of this

http://dx.doi.org/10.1016/j.molstruc.2014.09.079
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research is to report the Raman and infrared spectra of inyoite and to relate the spectra to the molecular structure of the mineral. This is the first report of a combined systematic study of inyoite by a wide range of techniques including infrared and Raman spectroscopy. Due to the industrial importance, inyoite was subject of scientific studies in different ways [10].

Experimental

Samples description and preparation

The inyoite 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 SAC-013. The mineral originated from Mount Blanco mine, Mount Blanco, Black Mountains, Death Valley, Inyo County, California [11] in the Puna Austral subprovince [12] and constitutes an evaporate deposit. Detailed study concerning the geology of the deposit and genetic aspects was published [13].

The sample occurs as a single crystal up to 3 cm along the c axis. Cleavage fragments were collected under a stereomicroscope Zeiss Stemi DV4 from the Museu de Ciência e Técnica – UFOP. Scanning electron microscopy (SEM) in the EDS mode was applied to support the mineral characterization.

X-ray diffraction

X-ray diffractogram was obtained in a Shimadzu XRD 6000 diffractometer equipped with an iron tube and a graphite monochromator. The scans were done between 4° and 70° (2θ) with a scanning speed of 0.5 degree/min. Silicon was used as an internal standard. Cell parameters were refined by means of the Jade program using least-square refinement after subtracting the background and the Kα2 contribution and using intensity and angular weighting of the most intense peaks.

Thermogravimetric analysis

Simultaneous thermogravimetric analysis (TGA) and differential thermal analysis (DTA) were performed in a Du Pont SDT2960 module. The temperature ranged from 25 °C to 1000 °C, using a constant flow of synthetic air (100 ml/min) and a heating rate of 20 °C/min.

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

Mineral characterization

X-ray powder diffraction data (Fig. 1) show monoclinic symmetry with space group P21/a. Unit cell parameters are: a = 10.63 Å; b = 12.06 Å; c = 8.405 Å; β = 114.03° and Z = 4, which are in agreement with published data [9].

The SEM image of inyoite sample studied in this work is shown in Fig. 2. The sample corresponds to needle like cleavage fragment up to 2.0 mm. The SEM image shows a homogeneous phase. Qualitative chemical analysis gave Ca as the major element. The presence of C is related to the carbon coating (Fig. 3).

Thermogravimetric analysis

The TGA and DTG curves for the mineral inyoite are shown in Fig. 4. Two mass-loss events occur at 111 °C and 129 °C (33.0%)
with a further mass loss at 403 °C (10.0%). It is proposed that the mass losses at 111 °C and 129 °C are due to the removal of water and the mass loss at 403 °C is attributed to the loss of the hydroxyl units. The theoretical mass losses using a molecular mass of 278 is 25.96% for the water mass loss step and 6.1% for the loss of the hydroxyls.

The following reactions are proposed:

\[ \text{Ca(H}_2\text{B}_3\text{O}_7\text{(OH)}\cdot 4\text{H}_2\text{O} \rightarrow \text{Ca(H}_2\text{B}_3\text{O}_7\text{(OH)} + 4\text{H}_2\text{O}} \]

\[ 2\text{Ca(H}_2\text{B}_3\text{O}_7\text{(OH)} \rightarrow 2\text{Ca(H}_2\text{B}_3\text{O}_6) + \text{H}_2\text{O}} \]

The values of the experimental results are less than the experimental values. One possibility is that there is a greater amount of water present.

Vibrational spectroscopy

The Raman spectrum of inyoite over the 4000–100 cm\(^{-1}\) spectral range is reported in Fig. 5a. The spectrum shows complexity with many bands being observed. This figure shows the position and relative intensities of the Raman bands. It is noteworthy that there are large parts of the spectrum where no intensity is observed. The Raman spectrum is therefore subdivided into sections depending upon the type of vibration being analysed. The infrared spectrum of inyoite over the 4000–500 cm\(^{-1}\) spectral range is displayed in Fig. 5b. The spectrum is not shown below 500 cm\(^{-1}\). The reason for this is that we are using a reflectance technique and the ATR cell absorbs all incident radiation. There are parts of this infrared spectrum where little or no intensity is observed. This spectrum may be thus subdivided into sections depending upon the type of vibration being analysed.

The Raman spectrum of inyoite over the 1450–850 cm\(^{-1}\) spectral range is illustrated in Fig. 6a. The Raman spectrum in this spectral region is dominated by a sharp intense band at 910 cm\(^{-1}\). On the high wavenumber side of this band, bands of significantly lesser intensity are observed at 925, 957, 971, 1013, 1048 and 1062 cm\(^{-1}\) with a broad band at 1204 cm\(^{-1}\). The Raman band at 910 cm\(^{-1}\) is assigned to the BO stretching vibration of the B\(_4\)O\(_{10}\) units. It is probable that there at least 13 BO stretching vibrations based upon a B\(_4\)O\(_{10}\) unit. Whether all these vibrations are coincident are not known but it is likely. The width of the symmetric stretching vibration in the Raman spectrum suggests that these vibrational modes of the BO stretching vibrations are coincident. Further, the existence of two isotopes, also complicates the situation. The nominal resolution of the Raman spectrometer is of the order of 2 cm\(^{-1}\) and as such is sufficient enough to identify separate bands for the stretching bands of the two boron isotopes. The two reduced masses for a pure B–O stretching mode would be \((10 \times 16)/(10 + 16) = 6.154\) for 10-B and \((11 \times 16)/(11 + 16) = 6.518\) for 11-B. The wavenumber is inversely proportional to square root of reduced mass; so the isotopic wavenumber ratio should be \(\sqrt{6.518/6.154} = 1.03\). 10-B is about 20% of natural boron, so a mode that is mostly B–O stretching and that includes significant motion of the B atom (not a breathing mode of a BO\(_3\) trigonal planar unit or a BO\(_4\) tetrahedral unit) should show a large peak for 11-B and a smaller peak at higher wavenumber for 10-B. For example if the sharp Raman peak at 925 in Fig. 3a is from the 11-B component such a mode, then it should have a smaller 10-B satellite near \((1.03) \times (925) = 952\) cm\(^{-1}\), and indeed a small
peak at 955 is observed in the figure. Similar small, higher wavenumber bands are also shown in this figure associated with peaks at 1013 and 1062 cm\(^{-1}\).

The Raman bands at 980, 1013, 1032, 1088, 1140, and 1323 cm\(^{-1}\) are attributed to the BOH in-plane bending modes. It is not known to what the very broad band at 1204 cm\(^{-1}\) is attributed. Iliev et al. determined the Raman spectrum of a synthetic cobalt boracite [14]. The symmetry species of some vibrational modes were determined. These researchers [15] used Raman imaging to show the ferroelectric properties of boracite type compounds. These workers [15] 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 [16]. On the low wavenumber side of the 1039 cm\(^{-1}\) peak, Raman bands with significant intensity are observed at 825 and 925 cm\(^{-1}\). These bands may be attributed to the antisymmetric stretching modes of tetrahedral boron.

The detailed infrared spectrum over the 1050–650 cm\(^{-1}\) spectral range is provided in Fig. 6b. This spectrum displays complexity with many bands being observed. The series of infrared bands at 924, 955, 980 and 1007 cm\(^{-1}\) are attributed to the trigonal borate antisymmetric stretching modes. The infrared band at 955 cm\(^{-1}\) is assigned to the BO stretching mode, the equivalent to the Raman band at 910 cm\(^{-1}\). The series of infrared bands from 677 through to 869 cm\(^{-1}\) are related to trigonal borate bending modes. The infrared bands at 1032, 1076 and 1161 cm\(^{-1}\) are assigned to BOH deformation modes. The infrared bands at around 792 and 804 cm\(^{-1}\) are assigned to water librational modes [17–19].

The Raman spectra in the 900–300 cm\(^{-1}\) and in the 300–100 cm\(^{-1}\) spectral ranges are shown in Fig. 7. Four sharp Raman bands are observed at 390, 599, and 739 cm\(^{-1}\). These bands are simply defined as trigonal and tetrahedral borate bending modes. A series of Raman bands at 160, 172, 182, 192, 206, 258 and 268 cm\(^{-1}\) (Fig. 7b) are due to lattice modes.

The Raman spectrum of inyoite in the 3800–2600 cm\(^{-1}\) spectral range is reported in Fig. 8a. The infrared spectrum of inyoite in the 3800–2500 cm\(^{-1}\) spectral range is reported in Fig. 8b. The formula of inyoite Na\(_2\)B\(_4\)O\(_6\)(OH)\(_2\)·H\(_2\)O is such that both water and hydroxyl stretching bands would be expected in both the Raman and infrared spectra. The difficulty is which band is attributable to which vibration. In the normal course of events, the hydroxyl stretching vibrations occur at higher wavenumbers than the water stretching wavenumbers [20–23]. Further, the widths of the hydroxyl stretching vibrations are narrow compared with the width of the water bands. A sharp Raman band observed at 3444 cm\(^{-1}\) are superimposed upon some broad bands at 2828, 3153 and 3389 cm\(^{-1}\). It is likely that these latter three bands are attributable to water stretching vibrations. The first band is attributed to the stretching vibrations of the hydroxyl units. The Raman bands observed in the Raman spectrum are also observed in the infrared spectrum at 2996, 3109, 3240, 3332, 3403, 3446, 3503 and 3535 cm\(^{-1}\). These infrared bands are ascribed to the stretching vibrations of the water and hydroxyl units. The sharp infrared band at 3535 cm\(^{-1}\) is assigned to the stretching vibrations of hydroxyl units.
The Raman spectrum of inyoite in the 1800–1500 cm\(^{-1}\) spectral range is reported in Fig. 9a. The infrared spectrum of inyoite in the 1800–1550 cm\(^{-1}\) spectral range is reported in Fig. 9b. Raman bands are found at 1656 and 1689 cm\(^{-1}\) and are assigned to water bending vibrations. The observation of two bands is in harmony with the number of water stretching vibrations. The two bands indicate water with different hydrogen bonding. The position of the bands indicates very strong hydrogen bonding in the mineral inyoite. In the infrared spectrum bands are found at 1468 and 1644 cm\(^{-1}\). This latter band is in harmony with the Raman bands and indicates water bending modes associated with strong hydrogen bonding.

**Conclusions**

There are many borate containing minerals which have yet to have their vibrational spectra measured and the molecular structure assessed in terms of their vibrational spectra. In this work we have measured the Raman and infrared spectrum of inyoite, a borate containing mineral. The importance of the mineral inyoite rests with the chemistry of the compound and the potential to synthesize nanomaterials based upon polymerization of borate units. Such compounds have the potential to make especially selected ferroelectric, pyroelectric and piezoelectric properties. The mineral inyoite is a precursor for the synthesis of such nanomaterials.

The inyoite sample studied in this work is from the Salar del Hombre Muerto, La Puna Plateau, Salta Province, Argentina. The borate mineral inyoite has been characterized by a range of complimentary techniques including X-ray diffraction, scanning electron microscopy, energy dispersive X-ray spectroscopy, thermal analysis and Raman spectroscopy at ambient temperatures complimented with infrared spectroscopy. Tentative assignments are made based upon the position and intensity of the infrared and Raman bands. Two boron isotopes are known namely \(^{10}\)B and \(^{11}\)B. The \(^{10}\)B is around 20% in concentration compared with \(^{11}\)B. The Raman spectrum shows a large peak for \(^{11}\)B and a smaller peak at higher frequency for \(^{10}\)B. The sharp Raman peak at 932 cm\(^{-1}\) is from the \(^{11}\)B component such a mode, then it should have a smaller \(^{10}\)B satellite near \((1.03) \times (932) = 980\) cm\(^{-1}\), and indeed a low intensity peak at 980 cm\(^{-1}\) is observed.
Acknowledgments

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.

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