Characterization of the sulphate mineral amarantite – Fe$_3$ (SO$_4$)O $\cdot$ 7H$_2$O using infrared, Raman spectroscopy and thermogravimetry

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We have studied the mineral amarantite Fe$_3$ (SO$_4$)O $\cdot$ 7H$_2$O. The Raman spectrum is dominated by an intense band at 1017 cm$^{-1}$. Very intense Raman band observed at 409 cm$^{-1}$ is assigned to the $\nu_2$ bending modes. Vibrational spectroscopy enables aspects of the molecular structure of amarantite to be ascertained.

**A B S T R A C T**

The mineral amarantite Fe$_3$ (SO$_4$)O $\cdot$ 7H$_2$O has been studied using a combination of techniques including thermogravimetry, electron probe analyses and vibrational spectroscopy. Thermal analysis shows decomposition steps at 77.63, 192.2, 550 and 641.4°C. The Raman spectrum of amarantite is dominated by an intense band at 1017 cm$^{-1}$ assigned to the SO$_4^{2-}$ $\nu_1$ symmetric stretching mode. Raman bands at 1039, 1054, 1098, 1131, 1195 and 1233 cm$^{-1}$ are attributed to the SO$_4^{2-}$ $\nu_3$ antisymmetric stretching modes. Very intense Raman band is observed at 409 cm$^{-1}$ with shoulder bands at 399, 451 and 491 cm$^{-1}$ are assigned to the $\nu_2$ bending modes. A series of low intensity Raman bands are found at 543, 602, 622 and 650 cm$^{-1}$ are assigned to the $\nu_4$ bending modes. A very sharp Raman band at 3529 cm$^{-1}$ is assigned to the stretching vibration of OH units. A series of Raman bands observed at 3025, 3089, 3227, 3340, 3401 and 3480 cm$^{-1}$ are assigned to water bands. Vibrational spectroscopy enables aspects of the molecular structure of the mineral amarantite to be ascertained.

**I N T R O D U C T I O N**

The mineral amarantite Fe$_3$ (SO$_4$)O $\cdot$ 7H$_2$O [1–3] is a sulphate of iron found in arid climates. Amarantite is a very rare mineral and is only found in very dry localities such as Carocoles, Chile [1,4]. The mineral has been identified in iron sulphate deposits in Iran [5]. The mineral is also known from Muturoo, Pine Creek – Muturoo area, Mingary, Olary Province, South Australia, Australia [6]. The mineral is found at Weelhamby Lake, Western Australia and in the Koolyanobbing Greenstone Belt, Western Australia. The reason why the mineral is found in these Australian sites is that these localities are very arid.
Interest in the study of Fe(II) sulphates arises from at least two perspectives. Firstly the observation of sulphates of iron on Mars by recent exploration and secondly by the formation of Fe(ii) and Fe(III) sulphates in evaporite deposits [7]. Studies of these minerals have been undertaken for some considerable time [8–12]. Interest in the chemistry of iron bearing sulphates stems from a number of reasons. Firstly, because of the discovery of jarosites on Mars [13,14]. Such a discovery implies the presence of water on Mars either at present or at some time in the planetary past [15,16]. Interest in such minerals and their thermal stability rests with the possible identification of these minerals and dehydrated para-genetically related minerals on planets and on Mars. There have been many studies on related minerals such as the Fe(ii) and Fe(III) sulphate minerals [17–22]. The formation of these minerals can occur through microbiological reduction of pyrite [23,24]. Fe minerals like Fe⁺, Fe³⁺ hydroxysulfate (copiapite); Fe³⁺ oxohydroxides (goethite); KFe³⁺hydroxysulfate (jarosite) and Fe⁵⁺ hydroxysulfate (schwartzmannite) that may form as weathering products in pyrite oxidation process. Oxidation rates are dependent on temperature, pH, Eh, relative humidity, and the surface area of reactant pyrite.

One of the difficulties associated with the analysis of Fe(III) sulphates and related minerals is that they are often lack crystallinity, making detection using X-ray diffraction (XRD) techniques difficult. Vibrational spectroscopic techniques are most useful in overcoming this difficulty. Raman spectroscopy enables the spectra of such minerals to be readily obtained. Most efflorescent minerals such as jarosite are poorly crystalline as the loss of water from the structure causes the materials to become powdered. Another problem associated with the study of Fe(III) and Fe(ii) sulphates is their thermodynamic stability [25]. Often, the minerals are formed from acid–sulphate rich environments such as acid mine drainage and acid–sulphate soils and as such their solubility is controlled by the climatic conditions in particular temperature. The minerals can precipitate in the day time as the solution evaporates and then redissolves at night when the temperature decreases. Such phenomena results in very complex mineral systems which involve jarosites and other sulphates for example iron(ii) sulphates and potassium sulphate. The ability to be able to easily and readily detect these iron sulphate minerals including jarosite is of importance [26,27].

Sulphate minerals are formed by the oxidation of sulphide minerals by a combination of oxidation, dehydration, and neutralization reactions. In sulphate efflorescent deposits gelatinous hydrated hydroxy sulphates such as the sulphate minerals including jarosites and halotrichites may be formed in solution and precipitate from this solution as a function of the pH and temperature [28]. Such mixed sulphate species will depend upon the temperature and pH of the crystallization [29,30]. A typical deposit is found in the El Jaroso ravine. A recent study characterized jarosites in terms of their UV–Visible and NIR spectroscopy [31] whereas other recent studies have characterized selected evaporate minerals by thermal analysis and Raman spectroscopic techniques [32–35]. Thus it is important to understand the NIR spectroscopy of the efflorescent sulphate minerals jarosites and halotrichites since these minerals are formed in mineral waste deposits, slag wastage and other environmental situations [36–38].

Amarantite is triclinic [2,39,40], which means crystallographically, it has only one symmetry fold. It must be rotated 360° to be exactly the same. Due to it being triclinic it falls into the biaxial optical class, the axis degrees do not equal 90° and the sides of each axis are not the same length. Amarantite is anisotropic, which means, the velocity of light varies with crystallographic direction, and, thus, there is more than one refractive index. The objective of this research is to determine the molecular structure using vibrational spectroscopy and to ascertain the thermal stability of the mineral.

**Experimental**

**Samples description and preparation**

The amarantite sample studied in this work was obtained from the collection of the Geology Department of the Federal University of Ouro Preto, Minas Gerais, Brazil, with sample code SAB-099. The sample is from the type locality for the mineral, the Caracoles deposit, Sierra Gorda district, Chile. The sample was gently crushed and the associated minerals were removed under a stereomicroscope Leica MZ4. Scanning electron microscopy (SEM) was applied to support the chemical characterization and ascertain the morphology of the mineral.

**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). Amarantite crystals were coated with a 5 nm layer of evaporated Au. Secondary Electron and Backscattering Electron images were obtained using a JEOL JSM-6360LV equipment. Qualitative and semi-quantitative chemical analysis in the EDS mode were performed with a ThermoNORAN spectrometer model Quest and was applied to support the mineral characterization.

**Thermogravimetric analysis – TG/DTG**

Thermogravimetric analysis of the amarantite mineral were obtained by using TA Instruments Inc. Q50 high-resolution TGA operating at a 10 °C/min ramp with data sample interval of 0.50 s/pt from room temperature to 1000 °C in a high-purity flowing nitrogen atmosphere (100 cm³/min). A total mass of 36.3 mg of finely ground sample was heated in an open platinum crucible.

**Raman microprobe spectroscopy**

Crystals of amarantite were placed on a polished metal surface on the stage of an Olympus BHS 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 Spectralcalc 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 amarantite sample studied in this work is shown in the supplementary information as Fig. S1. The crystal shows a prismatic form, with perfect cleavage along (010). Qualitative chemical composition shows a homogeneous phase, composed by S and Fe. The EDS spectrum is shown in the supplementary information as Fig. S2.

To date, there have been no thermal analytical studies of amarantite and related minerals [3,41]. There have been some differential thermal analysis studies of a very wide range of sulphates from some years past [42]. The pattern for the thermogravimetric analysis is presented in Fig. 1. The TG curve shows a total mass loss of about 61.3% on heating to 1000 °C and is in agreement with the stoichiometric content of H₂O and SO₄ that correspond to 64.0 wt%. In the temperature range of 25–100 °C, is observed a mass loss of about 5.2%.

The accepted formula by the IMA is Fe₂⁺(SO₄)O·7H₂O. If the formula is written as per Fe atom, then it may be written as Fe³⁺(SO₄)(OH)·3(H₂O).

This mass loss is attributed to the loss of adsorbed water, according to the equation:

Fe³⁺(SO₄)(OH)·3(H₂O) → Fe³⁺(SO₄)(OH)·2(H₂O) + H₂O

The mass loss at 77.63 °C is assigned to the loss of one water molecule. The mass loss of 8.1% would be predicted from the formula Fe³⁺(SO₄)(OH)·2(H₂O). The mass loss at 192.2 results in a mass loss of 16.66%. This mass loss is attributed to the following chemical reaction:

Fe³⁺(SO₄)(OH)·2(H₂O) → Fe³⁺(SO₄)(OH) + 2H₂O

The mass loss of 16.66% fits well with the predicted mass loss of 16.2%. A small DTG peak is found at 550 °C which is assigned to the loss of OH units according to the following equation:

2Fe³⁺(SO₄)(OH) → 2Fe³⁺O(SO₄) + H₂O

In the temperature range of 500–650 °C a mass loss of 30.43%. A strong exothermic reaction is observed at 641.4 °C. This mass loss is attributed to the loss of sulphate.

2Fe³⁺O(SO₄) → Fe₂O₃ + 2SO₃

According to this reaction the mass loss of 36% would be predicted based upon the formula Fe³⁺(SO₄)(OH)/3(H₂O). A mass loss of 30.43% is observed for this step. The final product of the thermal decomposition of amarantite is hematite.

Vibrational spectroscopy

Background spectroscopy of sulphate

The Raman spectroscopy of the aqueous sulphate tetrahedral oxyanion yields the symmetric stretching (ν₁) vibration at 981 cm⁻¹, the in-plane bending (ν₂) mode at 451 cm⁻¹, the antisymmetric stretching (ν₃) mode at 1104 cm⁻¹ and the out-of-plane bending (ν₄) mode at 613 cm⁻¹ [43]. Ross reports the interpretation of the infrared spectra for potassium alum as ν₁, 981 cm⁻¹; ν₂, 465 cm⁻¹; ν₃, 1200, 1105 cm⁻¹; ν₄, 618 and 600 cm⁻¹ [44]. Water stretching modes were reported at 3400 and 3000 cm⁻¹, water bending modes at 1645 cm⁻¹, and water librational modes at 930 and 700 cm⁻¹ [45]. The Raman spectrum of the mineral chalcanthite shows a single symmetric stretching mode at 984.7 cm⁻¹. Two ν₂ modes are observed at 463 and 445 cm⁻¹ and three ν₃ modes at 1173, 1146 and 1100 cm⁻¹. The ν₄ mode is observed as a single band at 610 cm⁻¹. A complex set of overlapping bands is observed in the low wavenumber region at 257, 244, 210, 136 and 126 cm⁻¹. Recently, Raman spectra of four basic copper sulphate minerals, namely antlerite, brochantite, posnjakite and langite, were published [46]. The SO₄ symmetric stretching modes for the four basic copper sulphate minerals are observed at 985, 990, 972 and 974 cm⁻¹. Only the mineral brochantite showed a single band in this region. Multiple bands were observed for these minerals in the antisymmetric stretching region.

Ross also lists the infrared spectra of the pseudo-alums formed from one divalent and one trivalent cation. Halotrichite has infra-
red bands at $v_1$, 1000 cm$^{-1}$; $v_2$, 480 cm$^{-1}$; $v_3$, 1121, 1085, 1068 cm$^{-1}$; $v_4$, 645, 600 cm$^{-1}$. Pickeringite the Mg end member of the halotrichite–pickeringite series has infrared bands at $v_1$, 1000 cm$^{-1}$; $v_2$, 435 cm$^{-1}$; $v_3$, 1085, 1025 cm$^{-1}$; $v_4$, 638, 600 cm$^{-1}$ [44]. These minerals display infrared water bands in the OH stretching, 3400 and 3000 cm$^{-1}$ region; OH deformation, 1650 cm$^{-1}$ region; OH libration, 725 cm$^{-1}$ region. Ross also reports a weak band at $\sim$960 cm$^{-1}$ which is assigned to a second OH librational vibration [44]. As with the infrared spectra, Raman spectra of alums are based on the combination of the spectra of the sulphate and water. Sulphate typically is a tetrahedral oxyanion with Raman bands at 981 ($v_1$), 451 ($v_2$), 1104 ($v_3$) and 613 ($v_4$) cm$^{-1}$ [47].

Some sulphates have their symmetry reduced through acting as monodentate and bidentate ligands [47]. In the case of bidentate behavior both bridging and chelating ligands are known. This reduction in symmetry is observed by the splitting of the $v_3$ and $v_4$ into two components under $C_3v$, symmetry and three components under $C_{2v}$ symmetry. A complex set of overlapping bands is observed in the low wavenumber region with broad bands observed at 257, 244, 210 136 and 126 cm$^{-1}$. Recently, Raman spectra of four basic copper sulphate minerals, namely antlerite, brochantite, posnjakite and langite, were published [46]. The SO symmetric stretching vibrations supports the concept of the non-equivalence of the sulphate units in the structure of amaranthite.

Vibrational spectroscopy of amaranthite

The Raman spectrum of amaranthite over the complete wavenumber range from 100 to 4000 cm$^{-1}$ spectral range is displayed in Fig. 2a. This spectrum shows the position of the peaks and the relative intensities of the bands. It is apparent that there are large parts of the spectrum where no intensity is observed. Therefore the spectrum is subdivided into sections depending upon the type of vibration being studied. The infrared spectrum over the 500–4000 cm$^{-1}$ spectral range is reported in Fig. 2b. This spectrum shows the relative intensity of the infrared bands and their position. There are also large parts of the infrared spectrum where no intensity is observed. Hence, the spectrum is subdivided into sections as a function of the type of vibration being examined.

Raman spectroscopy in the 800–1400 cm$^{-1}$ spectral range

The Raman spectrum of amaranthite in the 800–1400 cm$^{-1}$ spectral range is shown in Fig. 3a. The spectrum shows a very intense band at 1017 cm$^{-1}$ with a low intensity shoulder band at 1006 cm$^{-1}$. This band is assigned to the $SO_4^{2-}$ $v_1$ symmetric stretching mode. A series of bands are observed at 1039, 1054, 1098, 1131, 1195 and 1233 cm$^{-1}$. These bands are assigned to the $SO_4^{2-}$ $v_3$ antisymmetric stretching modes. The observation of more than one band in the symmetric stretching region together with multiple antisymmetric stretching vibrations supports the concept of the non-equivalence of the sulphate units in the structure of amaranthite. The low intensity Raman band at 831 cm$^{-1}$ is probably due to water librational modes. The Raman spectrum of devilline shows a single $SO_4$ symmetric stretching band at 1007 cm$^{-1}$. The band is sharp with a bandwidth of 3.1 cm$^{-1}$. The antisymmetric stretching band for devilline is observed at 1134 cm$^{-1}$. In contrast, the Raman spectrum of cyanotrichite shows a single intense band at 976 cm$^{-1}$ with a broad low intensity band at 966 cm$^{-1}$. Three antisymmetric stretching bands are observed at 1137, 1101 and 1057 cm$^{-1}$. For glaucocerinite bands are observed at 1007 and...
Infrared spectroscopy in the 500–1300 cm\(^{-1}\) spectral range

The infrared spectrum of amaranthite in the 500–1300 cm\(^{-1}\) spectral range is displayed in Fig. 3b. The spectrum consists of a broad spectral profile and is made up of a series of overlapping bands as shown. An intense sharp band is observed at 1000 cm\(^{-1}\) with a shoulder band at 986 cm\(^{-1}\). These bands are assigned to the \(\nu_1\) \(\text{SO}_4^{2-}\) symmetric stretching mode. The infrared bands observed at 1039, 1080, 1107, 1139 and 1196 cm\(^{-1}\) are assigned to the \(\nu_3\) \(\text{SO}_4^{2-}\) antisymmetric stretching vibrations. It is noted the Raman and infrared bands are almost in the same position. Some low intensity infrared bands are observed at 724, 768, 825 and 845 cm\(^{-1}\) and are assigned to water vibrational modes. The two infrared bands at 655 and 661 cm\(^{-1}\) are \(\text{SO}_4^{2-}\) bending modes.

Raman spectroscopy in the 300–800 cm\(^{-1}\) and in 100–300 cm\(^{-1}\) spectral ranges

The Raman spectrum in the 300–800 cm\(^{-1}\) and 100–300 cm\(^{-1}\) spectral range is reported in Fig. 4a and b. The complexity of the antisymmetric stretching region is reflected in the spectra of the \(\nu_2\) bending region. A very intense Raman band is observed at 409 cm\(^{-1}\) with shoulder bands at 399, 451 and 491 cm\(^{-1}\). These bands are assigned to the \(\nu_2\) bending modes. Multiple bands are observed in the 400–500 cm\(^{-1}\) region for many minerals and are attributed to the \(\nu_2\) bending modes. The observation of several bending modes is in agreement with the number of antisymmetric stretching vibrations. Bands are observed at 479, 443 and 408 cm\(^{-1}\) for cyanotrichite, at 450 and 430 cm\(^{-1}\) for devilline, 498 and 471 cm\(^{-1}\) for glaucocerinite, at 475, 445 and 421 cm\(^{-1}\) for serpierite and at 475 and 449 for ktenasite.

The spectra of antlerite and brochantite are similar as are those for posnjakite and langite in this spectral region [46,48]. The antlerite Raman spectrum shows bands at 485, 469, 440 and 415 cm\(^{-1}\) with the latter band having the highest intensity. Bouchard–Abouchacra reported bands for antlerite at 483, 470, 444 and 416 cm\(^{-1}\) in excellent agreement with our data [49]. Raman spectra of brochantite are similar except that additional bands at 517 and 501 cm\(^{-1}\) are observed. The Raman spectra of posnjakite show bands at 511, 482, 447, 422, 386 and 363 cm\(^{-1}\). This complex set of bands in this region is also observed for langite with bands observed at 507, 481, 449, 420, 391 cm\(^{-1}\). The Raman spectrum of wroewulfeite shows a complex set of overlapping bands at 472, 457, 443 and 415 cm\(^{-1}\). The observation of additional bands over and above that which would be predicted for the sulphate anion (a single band at 451 cm\(^{-1}\)), may be attributed to a number of factors including symmetry reduction, local stress in the crystals and crystal orientation effects.

A series of low intensity Raman bands are found at 543, 602, 622 and 650 cm\(^{-1}\) are assigned to the \(\nu_4\) bending modes. The Raman spectrum of ktenasite shows a single peak in the \(\nu_4\) region at 604 cm\(^{-1}\). The Raman spectrum of devilline shows two low intensity bands at 668 and 617 cm\(^{-1}\); cyanotrichite two overlapping bands at 594 and 530 cm\(^{-1}\); the Raman spectrum of glaucocerinite shows two bands at 694, 613 and 555 cm\(^{-1}\). The complexity of the symmetric and antisymmetric bending region shows a reduction in symmetry for these complex sulphate miner-

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**Fig. 4.** (a) Raman spectrum of amaranthite over the 300–800 cm\(^{-1}\) spectral range and (b) Raman spectrum of amaranthite over the 100–300 cm\(^{-1}\) spectral range.

**Fig. 5.** (a) Raman spectrum of amaranthite over the 2600–4000 cm\(^{-1}\) spectral range and (b) infrared spectrum of amaranthite over the 2600–4000 cm\(^{-1}\) spectral range.
Raman spectroscopy in the 2600–3800 cm$^{-1}$ spectral range

The Raman spectrum of amarantite in the 2600–3800 cm$^{-1}$ spectral range is reported in Fig. 5a. Two features are observed: firstly a very sharp band at 3529 cm$^{-1}$ and a broad spectral profile with Raman bands observed at 3025, 3089, 3227, 3340, 3401 and 3480 cm$^{-1}$. The sharp band is assigned to an OH stretching vibration, whereas the broad bands are attributed to water stretching vibrations. The infrared spectrum of amarantite in the 2600–3800 cm$^{-1}$ spectral range is displayed in Fig. 5b. The spectral profile is similar to that in the Raman spectrum. The infrared band at 3519 cm$^{-1}$ is attributed to the stretching vibration of the OH units. The infrared bands at 2944, 3058, 3272 and 3403 cm$^{-1}$ are assigned to the stretching vibrations of water.

Conclusions

The techniques of electron microscopy, thermogravimetry and vibrational spectroscopy have been combined to determine the chemistry of the mineral amarantite. Chemical analysis using an electron probe shows qualitatively that only Fe and S are present. Thermogravimetry proves the thermal decomposition of amarantite takes place in a series of four major steps at 77.63, 192.2, 550 and 641.4 °C with a lesser mass loss step at 550 °C. Raman spectroscopy is a very powerful tool for the study of sulphate minerals. In this work we have used vibrational spectroscopy to study the mineral amarantite, a mineral found in evaporite deposits and land surfaces with extreme aridity. Multiple antisymmetric stretching bands are observed as well as multiple bending modes suggesting a reduction in symmetry of the sulphate in the amarantite structure. The symmetry of the sulphate as observed by the number of bands in the $v_2$, $v_3$ and $v_4$ modes is essentially reduced to $C_{2v}$.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.saa.2013.04.111.

References
