Structure refinement, hydrogen-bond system and vibrational spectroscopy of hohmannite, $Fe_2^{3+}[O(SO_4)_2]\cdot 8H_2O$

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ABSTRACT

The crystal structure of hohmannite, $Fe_2^{3+}[O(SO_4)_2] \cdot 8H_2O$, was studied by means of single-crystal X-ray diffraction (XRD) and vibrational spectroscopy. The previous structural model was confirmed, though new diffraction data allowed the hydrogen-bond system to be described in greater and more accurate detail. *Ab initio* calculations were performed in order to determine accurate H positions and to support the experimental model obtained from XRD data. Infrared and Raman spectra are presented for the first time for this compound and comments are made on the basis of the crystal structure and the known literature for sulfate minerals.

Keywords: hohmannite, crystal structure refinement, *ab initio* calculation, FTIR spectroscopy, Raman spectroscopy.

Introduction

HOHMANNITE, $Fe_2^{3+}[O(SO_4)_2]\cdot 8H_2O$, is a hydrous Fe-sulfate occurring in arid environments. It is found typically as a secondary product, associated with chalcanthite, metahohmannite, fibroferrite, sideronatrite and copiapite (Palache et al., 1951) and originates from the chemical weathering and oxidation of primary Fe sulfides, such as pyrite or pyrrhotite, owing to bacterial action or to airoxidation in hydrothermal and magmatic ore deposits (Bandy, 1938). At present there is an increasing interest in the scientific community for the study of Fe sulfates for various reasons. Firstly, Fe-bearing sulfates have been identified recently on Mars, with wide distributions and, locally, in large amounts, using both remote sensing spectroscopic observations and directsurface explorations by rovers (e.g. Klingelhöfer et al., 2004; Johnson et al., 2007; Vicenzi et al.,

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2007; Lane et al., 2008; Mills et al., 2013; McCollom et al., 2014). Secondly, these sulfates originate as an intermediate step in a complex series of oxidation, dehydration and neutralization reactions culminating in the precipitation of Fe oxyhydroxides (Jerz and Rimstidt, 2003; Burns, 1987); these processes are associated with AMD (acid mine drainage) and result in the outflow of acidic water from metal- or coal-mine wastes. thus representing a significant environmental concern (e.g. Nordstrom et al., 2000; Welch et al., 2007, 2008, 2009). Iron-sulfate concentrations are common in base-metal deposits, coal deposits, tailings and waste-rock piles, and are related spatially to oxidizing Fe-sulfide minerals. Their considerable solubility causes Fe-sulfate minerals to be found in sheltered sites such as rock overhangs (thereby protected from rain dissolution), or during drier times of the year, as a consequence of the migration of Fe-sulfate solutions to the surface and subsequent evaporation (Jerz and Rimstidt, 2003). Therefore these compounds are sensitive indicators of environmental conditions such as pH, relative humidity, oxygen activity and sulfate activity. Hohmannite, together with other metal-bearing sulfates, is, additionally, involved in industrial processes, as a product of the corrosion of carbon steels (Ruhl and Kranzmann, 2012) and as an intermediate product in the treatment of metal residues from hydrometallurgic plants (Ngenda *et al.*, 2009).

Iron-bearing sulfates are difficult to characterize because they occur typically as loose, poorly coherent, powdery masses, making their detection using single-crystal XRD techniques problematic. For this reason, structural investigations on hohmannite are limited. Vibrational spectroscopy techniques are extremely useful in this context (Della Ventura *et al.*, 2013; Ventruti *et al.*, 2013) because they can provide a rapid identification of the sulfate phase from both powders and single crystals; they are also able to provide a detailed characterization of the degree of hydration of the phase, which is essential for a rigorous interpretation of the hydrologic history of the environment.

To date, no spectroscopic studies have been reported for hohmannite. Its crystal structure was solved firstly by Scordari (1978) starting from a three-dimensional Patterson synthesis, whereas the geometry and positions of the hydrogen bonds were inferred only from geometrical considerations. The aim of the present work is to provide a complete and detailed characterization of the sample studied originally by Scordari (1978) by combining modern XRD with periodic *ab initio* calculations to describe the hydrogen-bonding

TABLE 1. Refined cell parameters and data-collection parameters for the hohmannite sample studied by singlecrystal XRD.

Crystal data	
Crystal dimensions (mm)	$0.320 \times 0.310 \times 0.040$
Crystal system	Triclinic
Space group	PĪ
Unit-cell dimension a (Å)	9.1428(2)
b (Å)	10.9346(3)
c (Å)	7.2168(2)
α (°)	90.547(1)
β (°)	90.612(1)
γ (°)	107.375(1)
Volume (Å ³)	688.46(3)
Z, ρ_{calc} (g·cm ⁻³)	2, 2.238
$\mu (\text{mm}^{-1})$	2.50
F(000)	472
Data collection and refinement	
θ range for data collection (°)	2 to 38.0
Index range	$-15 \le h \le 15, -18 \le k \le 18, -12 \le l \le 12$
Completeness	98.0%
Max. and min. transmission	0.629, 0.747
Reflections collected/independent	24,651/7183
R merging $[R_{(int)}]$ (%)	2.24
Reflections used	5787 with $I > 3\sigma(I)$
No. of refined parameters	238
Gof ^a	0.76
$R_1^{\rm b}$ (on F) / $wR_2^{\rm c}$ (on F^2)	0.027/0.029
$(\Delta/\sigma)_{\rm max}$	0.001
$\Delta \rho_{\rm min} / \Delta \rho_{\rm max} (e/Å^3)$	-0.71/0.79

^a =[$\Sigma [w(F_o^2 - F_c^2)^2]/(n-p)$]^{1/2}, where *n* is the number of reflections, *p* is the total number of parameters refined and *w* is the weighting scheme applied;

^b =
$$R_1 = \Sigma[|F_o| - |F_c|]/\Sigma|F_o;$$

^c = $wR_2 = [\Sigma[w(F_o^2 - F_c^2)^2]/\Sigma[w(F_o^2)^2]]^{1/2}.$

system. Vibrational (Fourier transform infrared (FTIR) and Raman spectroscopy) data are also presented and interpreted on the basis of this improved structural model.

Experimental

The crystals investigated in this study were handpicked from the same rock specimen as studied by Scordari (1978) from the Sierra Gorda (Chile) deposit; thus the chemical composition is inferred to be the same as that reported in that study. A single crystal selected on the basis of its purity under a binocular microscope was mounted on a Bruker AXS APEXII diffractometer kappagoniometer equipped with a 1K SMART CCD (charge coupled device) detector with a crystal-todetector distance of 4 cm. The diffraction data were collected using monochromatic MoKa radiation and frame widths of 0.5° in ω , with 30 s acquisition time for each frame. A sphere of threedimensional data was collected by employing an optimization strategy by the Apex program suite (Bruker, 2008) over the interval $4^{\circ} \leq 2\theta \leq 76^{\circ}$. The three-dimensional data were reduced and corrected for Lorentz, polarization and background effects by means of the Bruker program SAINT-IRIX (Bruker, 2008). Empirical absorption corrections were applied to all data on the basis of the intensities of equivalent reflections (Blessing, 1995) by means of the multi-scan method implemented in SADABS (Bruker, 2008). The structure refinement was performed in space group $P\bar{1}$ using the program CRYSTALS (Betteridge et al., 2003) starting from the atomic parameters obtained by Scordari (1978). Reflections with $I > 3\sigma(I)$ were used for the structure refinement. Refined parameters were: atomic positions, atomic displacement parameters and overall scale factor. All atoms apart from hydrogen were refined anisotropically. The hydrogen atoms were located from the difference Fourier maps and refined independently by fixing the O–H distances at 0.85(3) Å and maintaining a dihedral angle H-O-H of ~104°. The isotropic atomic displacement parameters of hydrogen atoms were evaluated as $1.2*U_{eq}$ of the parent oxygen atom. Details on data collection, crystallographic data and refinement are given in Table 1. Atomic positions and displacement parameters for all nonhydrogen atoms are listed in Table 2. The refined positions of hydrogen atoms derived from difference-Fourier syntheses are given in Table 3. Selected interatomic distances are given in Table 4. The electronic structure of the refined structure was calculated by means of a linear combination of atomic orbitals (LCAO) unrestricted Hartree-Fock (UHF) periodic computational scheme, using the *CRYSTAL09* program (Saunders *et al.*, 2003). For each atom a split-valence basis set was used. Optimization of exponents and coefficients was performed for the Gaussian-type valence orbital of S ions. To facilitate the convergence, the number of the unpaired electrons of Fe^{3+} ions was locked in the first cycles of calculation. The cell parameters and the initial hydrogen positions adopted for each quantum-mechanical calculation were those determined by XRD.

For FTIR powder spectroscopy, samples were prepared as KBr pellets by mixing ~0.5 mg of sample, purified under the binocular microscope and verified by XRD, in 150 mg of KBr. Spectra were collected in the 4000–400 cm⁻¹ MIR (midinfrared) range by co-adding 128 scans with a 4 cm⁻¹ nominal resolution on a Nicolet Magna 760 spectrophotometer equipped with a KBr beam splitter, a globar source and a room-*T* DGTS detector. Single-crystal FTIR spectra in the NIR (near infrared) range were obtained from a small cleavage fragment, ~20 µm thick, using a Bruker Hyperion 3000 microscope, equipped with an MCT (mercury-cadmium-telluride) detector in the 7500–4000 cm⁻¹ range by co-adding 128 scans with a 4 cm⁻¹ nominal resolution.

Single-crystal Raman spectra were obtained using a confocal Horiba Jobin Yvon Labram HR Evolution spectrometer equipped with an Olympus BX41 optical microscope and a multichannel air-cooled charge-coupled device (CCD) detector. Unpolarized Raman spectra were excited with the He-Ne 632.8 nm line. The laser power was decreased to <1 mW to avoid potential local heating effects due to heavy light absorption. The sample was inspected optically for any laser damage, and none was observed. Spectra were recorded over the wavenumber range 10 to 4000 cm⁻¹ using an Ultra Low Frequency (ULF) filter. The wavenumber accuracy was ± 0.5 cm⁻¹, and the spectral resolution was <1 cm⁻¹. Baseline subtraction and peak fitting were performed using GRAMS/32 software (Galactic Industries Ltd, Salem, New Hampshire, USA).

The crystal structure of hohmannite

Hohmannite is related to amarantite, $Fe_2^{3+}[O(SO_4)_2]\cdot 7H_2O$ and metahohmannite, $Fe_2^{3+}[O(SO_4)_2]\cdot 4H_2O$ (Bandy, 1938; Strunz and

TABLE 2. Crystallographic coordinates and displacement parameters (Å²) for non-hydrogen atoms in hohmannite.

Atom	x	у	И	$U_{ m iso}$	U^{11}	U^{22}	U^{33}	U^{23}	U^{13}	U^{12}
Fe1	0.40222(2)	0.45707(2)	0.33834(3)	0.0132(1)	0.01412(8)	0.01191(8)	0.01111(8)	-0.00013(6)	-0.00076(6)	0.00003(6)
Fe2	0.66913(3)	0.76067(2)	0.30451(3)	0.0146(1)	0.01654(9)	0.01175(8)	0.01194(8)	0.00038(6)	0.00023(6)	-0.00124(7)
S3	0.14032(4)	0.17441(4)	0.30892(5)	0.0157(1)	0.01514(14)	0.01352(14)	0.01436(14)	-0.00162(11)	-0.00134(11)	-0.00182(11)
$\mathbf{S4}$	0.59142(4)	0.35448(3)	0.01969(5)	0.0134(1)	0.01497(14)	0.01293(13)	0.01162(13)	0.00035(10)	-	0.00330(11)
05	0.16357(16)	0.16955(14)	0.51212(17)	0.0261(6)	0.0250(6)	0.0285(6)	0.0144(5)	0.0003 (4)		-0.0075(5)
06	0.97969(16)	0.15706(14)	0.2686(2)	0.0284(6)	0.0172(5)	0.0269(6)	0.0357(7)	-0.0006(5)		-0.0014(5)
07	0.23354(15)	0.30203(12)	0.24307(18)	0.0221(5)	0.0236(6)	0.0155(5)	0.0195(5)	0.0007(4)		-0.0056(4)
08	0.19269(18)	0.07563(13)	0.2166(2)	0.0279(7)	0.0381(8)	0.0197(6)	0.0254(6)	-0.0027(5)		0.0077(5)
60	0.35349(16)	0.52742(13)	0.08989(18)	0.0243(5)	0.0256(6)	0.0202(5)	0.0201(5)	0.0081(4)		-0.0039(4)
010	0.55299(15)	0.72765(13)	0.06396(17)	0.0235(5)	0.0250(6)	0.0212(5)	0.0157(5)	0.0016(4)		-0.0060(4)
011	0.70509(18)	0.28674(16)	0.0294(2)	0.0306(8)	0.0348(8)	0.0381(8)	0.0283(7)	-0.0017(6)		0.0255(7)
012	0.56049(14)	0.38986(13)	0.21013(16)	0.0197(5)	0.0214(5)	0.0254(6)	0.0131(4)	-0.0042(4)		0.0081(4)
013	0.55748(13)	0.61234(10)	0.42567(15)	0.0144(4)	0.0162(4)	0.0117(4)	0.0117(4)	0.0010(3)		-0.0015(3)
014	0.7418(3)	0.2431(2)	0.4471(3)	0.0499(12)	0.0485(12)	0.0387(10)	0.0640(14)	0.0003(10)		0.0145(9)
015	0.77807(18)	0.92986(13)	0.16374(19)	0.0294(6)	0.0370(8)	0.0202(6)	0.0170(5)	0.0022(4)		-0.0129(5)
016	0.53901(19)	0.86434(14)	0.4068(2)	0.0287(7)	0.0361(8)	0.0235(6)	0.0295(7)	0.0052(5)		0.0129(6)
017	0.4882(3)	0.0398(2)	0.1897(3)	0.0439(11)	0.0538(12)	0.0425(10)	0.0448(10)	0.0145(8)		0.0278(9)
018	0.22425(17)	0.51219(15)	0.4441(2)	0.0281(7)	0.0217(6)	0.0304(7)	0.0330(7)	-0.0086(6)		0.0091(5)
019	0.1124(2)	0.80474(17)	0.2176(3)	0.0386(9)	0.0276(7)	0.0276(7)	0.0592(12)	-0.0045(8)		0.0061(6)
020	0.81760(17)	0.67875(15)	0.1759(2)	0.0280(7)	0.0208(6)	0.0324(7)	0.0283(7)	-0.0102(6)		0.0045(5)
021	0.9830(3)	0.4760(3)	0.2061(4)	0.0669(16)	0.0421(12)	0.0652(16)	0.0878(19)	0.0013(14)		0.0081(11)

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Atom	x	У	Ζ
H22	0.795(4)	0.229(4)	0.538(4)
	0.7707	0.2289	0.6771
H23	0.788(4)	0.222(4)	0.357(4)
	0.8133	0.2222	0.535
H24	0.791(4)	0.923(3)	0.052(3)
	0.7828	0.9296	0.0296
H25	0.842(3)	0.986(3)	0.202(4)
	0.8469	0.0083	0.217
H26	0.519(3)	0.917(3)	0.337(4)
	0.5288	0.9377	0.3341
H27	0.460(3)	0.838(3)	0.459(4)
	0.4480	0.8282	0.4811
H28	0.418(3)	0.067(4)	0.208(5)
	0.3979	0.0670	0.2065
H29	0.516(4)	0.054(4)	0.099(4)
	0.552	0.0862	0.0939
H30	0.152(3)	0.503(3)	0.372(4)
	0.1335	0.4942	0.3612
H31	0.239(3)	0.579(2)	0.491(4)
	0.2442	0.5942	0.5064
H32	0.166(4)	0.782(3)	0.144(4)
	0.1796	0.779	0.1355
H33	0.124(4)	0.876(2)	0.213(5)
	0.1322	0.8966	0.2214
H34	0.904(3)	0.708(3)	0.188(4)
	0.9204	0.7224	0.1931
H35	0.797(3)	0.657(3)	0.076(3)
	0.7906	0.6449	0.0519
H36	0.960(5)	0.429(4)	0.108(4)
	0.9418	0.404	0.1204
H37	0.904(5)	0.494(4)	0.242(6)
	0.9081	0.5122	0.2414

TABLE 3. Experimental and calculated coordinates for hydrogen atoms in hohmannite.

Nickel, 2001; Scordari et al., 2004), and is based structurally on polymerization of centrosymmetrical groups with composition $[Fe_4(H_2O)_8O_2(SO_4)_4]$. These minerals differ chemically in the amount of interstitial H₂O groups, which also result in different packing of the sulfate chains. The structure of hohmannite (Fig. 1), in particular, is characterized by two crystallographically non-equivalent Fe sites, Fe1 and Fe2, surrounded by oxygen and water molecules to form [Fe1O₅(H₂O)] and [Fe2O₃(3H₂O)] octahedra, respectively. Two centrosymmetrically related [Fe1O₅(H₂O)] octahedra share one edge (O13-013), whereas two additional centrosymmetrically related [Fe2O₃(3H₂O)] octahedra link to either end of the shared edges. Two symmetrically unrelated SO₄²⁻ groups join the Fe octahedral by

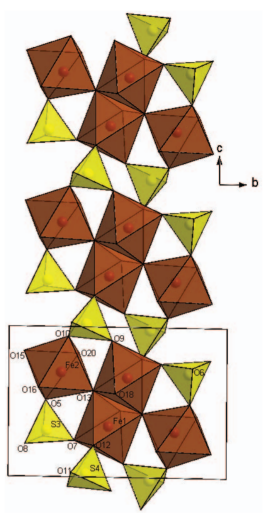


FIG. 1. Crystal structure of hohmannite viewed along the a axis.

corner sharing to form $[Fe_4(H_2O)_8O_2(SO_4)_4]$ clusters that polymerize through the O12 oxygen to form complex chains of Fe–O–S linkages running along the *c* axis. Adjacent chains are linked together by hydrogen bonds involving both the four water molecules belonging to the cluster and the four additional interstitial (H₂O) groups.

The positions of hydrogen atoms found here by difference Fourier (Table 3) confirm that the oxygen atoms O15, O16, O18 and O20 are water molecules linked to Fe^{3+} ions, whereas the oxygen atoms O14, O17, O19 and O21 are interstitial water molecules. Inspection of the data shows that all H…O distances are reasonable

Fe1-O7	2.029(1)	Fe2-O5*	1.978(1)
Fe1-O9	2.054(1)	Fe2-O10	1.998(1)
Fe1-O12	2.035(1)	Fe2-O13	1.869(1)
Fe1-O13	1.954(1)	Fe2-O15	2.099(1)
Fe1-013*	1.947(1)	Fe2-O16	2.013(2)
Fe1-O18	2.049(2)	Fe2-O20	2.060(2)
<fe1-o></fe1-o>	2.011	<fe2-o></fe2-o>	2.003
\$3-05	1.483(1)	S4-09*	1.479(1)
S3-06	1.449(1)	S4-O10*	1.476(1)
S3-O7	1.487(1)	S4-O11	1.447(1)
S3-O8	1.464(2)	S4-O12	1.477(1)
<s3-o></s3-o>	1.471	<s4-o></s4-o>	1.470

TABLE 4. Selected interatomic distances (Å) for the crystal structure of hohmannite.

Notes: * -x, -y, -z

and that all H atoms are involved in hydrogen bonds (Table 5).

Bond-valence sums were calculated using the electrostatic strength function of Brown and Altermatt (1985) and the bond-valence parameters of Breese and O'Keeffe (1991) for all the non-H atoms refined in the present study, and bearing in mind the H-bond contributions on the basis of the H…O computed distances (Ferraris

and Ivaldi, 1988). Table 6 shows that, excluding the contribution of the hydrogen atoms to the bond-valence calculations, the bond-valence sum (in vu, valence units) for O6, O8 and O11 (1.60, 1.54 and 1.61, respectively) is significantly less than the ideal value of 2 vu. When the hydrogen atoms are included in these calculations, the bond-valence sum becomes much more satisfactory. These calculations confirm that: (1) O6, O8

TABLE 5. Hydrogen-bond geometry.

D-H···A	D-H	H···A	D···A
	(Å)	(Å)	(Å)
O14-H22····O19*	0.86(2)	2.03(3)	2.878(3)
O14-H23····O6	0.85(2)	2.17(3)	2.921(3)
O15-H24···O8*	0.82(2)	1.95(3)	2.762(2)
O15-H25····O6 [†]	0.76(2)	1.97(3)	2.710(2)
O16-H26…O17 [†]	0.82(2)	1.81(3)	2.632(2)
O16-H27···O14*	0.79(2)	1.93(3)	2.714(3)
O17-H28····O8	0.79(2)	2.09(3)	2.852(3)
O17-H29····O17*	0.71(2)	2.30(3)	2.896(4)
O18-H30····O21 ^{**}	0.82(2)	1.89(3)	2.712(3)
O18-H31O14*	0.78(2)	1.94(3)	2.708(3)
O19-H32···O11*	0.81(2)	2.02(3)	2.827(2)
O19-H33····O8 [†]	0.76(2)	2.08(3)	2.831(2)
O20-H34…O19 [‡]	0.76(2)	1.89(3)	2.647(3)
O20-H35O7*	0.76(2)	2.38(3)	3.074(3)
O20-H35····O9*	0.76(2)	2.38(3)	2.993(3)
O21-H36····O21*	0.86(2)	2.50(4)	3.029(5)
O21-H37···O20	0.85(2)	2.43(4)	3.042(4)

Notes:* = -x, -y, -z

 $x^{**}_{+} = -1 + x, y, z.$

 $[\]frac{1}{x} = 1 + x, y, z$

 $^{^{\}dagger} = x, 1+y, z$

	05	90	07	08	60	010	011	012	013	014	015	016	017	018	019	020	021	Ω
			0.482		0.451			0.474	0.602					0.457				3.058
Fe2	0.553					0.524			0.743		0.399	0.503				0.443		3.165
	I.404	1.004	1.448	1.541	1.479	1.492 1.613	1.613	1.488										6.072 /
	2.017	1.604	1.930	1.541	1.930	2.016	2.016 1.613	1.962	1.937		0.399	0.503		0.457		0.443		
										0.846					0.154			1.000
		0.129								0.871								1.000
H24 H25		0 167		0.171							0.829 0.833							1.000
												0.793	0.207					1.000
										0.176		0.824						1.000
				0.143									$0.857 \\ 0.109$					$1.000 \\ 1.000$
													0.891					
H30														0.814			0.186	1.000
										0.174				0.826				1.000
				0.148											0.852			1.000
							0.156								0.844			1.000
															0.186	0.814		1.000
			0.100		0.100											0.800		1.000
																	0.086	1.000
H37																0.093	0.907	1.000
	2.017	1.900	2.030	2.003	2.030		2.016 1.769	1.962	1.937	2.067	2.061	2.120	2.064	2.097	2.036	2.150	2.093	

TABLE 6. Bond-valence analysis for the crystal structure of hohmannite.

STRUCTURE REFINEMENT OF HOHMANNITE

and O11 are indeed the acceptor atoms for the hydrogen atoms; (2) the interstitial water molecules O14, O17, O19 and O21 act simultaneously as donors and acceptors; and (3) the refined model, as well as the location for the hydrogen atoms, are correct. A reliable hydrogen-bonding scheme according to the converging results obtained from the different techniques quoted above is illustrated in Fig. 2. The bond-valence results are consistent with the previous study (Scordari, 1978) except for some details. In our scheme the O20-H35 hydrogen bond is probably bifurcated because H35 is almost equidistant from both O7 and O9. Finally, unlike the model proposed by Scordari (1978), the O21 water group acts as donors towards $\overline{O20}$ and O20. These structural data have been tested by making use of the density-functional-based energy minimization calculations. The hydrogen positions found with both techniques are converging (Table 3) even if the calculated O-H bonds (0.95-0.98 Å) are longer than the corresponding ones derived from the difference Fourier map (0.71(2)-0.86(2) Å). These differences are not unexpected and can be explained readily. The theoretical values, in fact, concern distances between nuclear positions, while the experimental values from XRD experiments measure distances between the centroids of the electron density: in

the latter case it is reasonable to expect the electron density of hydrogen to be shifted markedly towards the bonding atom.

Infrared and Raman spectroscopy

The powder FTIR spectrum of hohmannite in the MIR 4000–400 cm⁻¹ range is displayed in Fig. 3; measured band positions (wavenumbers, cm⁻¹) are listed in Table 7, where the band assignments are based on literature data for similar materials (e.g. Omori and Kerr, 1963; Adler and Kerr, 1965; Bishop *et al.*, 2004; Cloutis *et al.*, 2006; Ross, 1974; Lane, 2007; Della Ventura *et al.*, 2013).

In the higher wavenumber, water-stretching region (>2500 cm⁻¹) a very broad absorption extending from 3700 to 2800 cm⁻¹ (Fig. 3) is observed; it peaks around 3438 cm⁻¹ with a shoulder at 3234 cm⁻¹. This broad band is evidently due to the contribution of several overlapping components, including the overtone of the H₂O bending mode ($2v_2$) expected around 3250 cm⁻¹. The unpolarized spectrum collected on a single crystal in the NIR range (Fig. 4), shows a relatively broad and intense absorption due to the ($v_3 + v_2$) H₂O combination at 5115–5230 cm⁻¹. No bands are observed in the 4000–4500 cm⁻¹ range, ruling out any presence

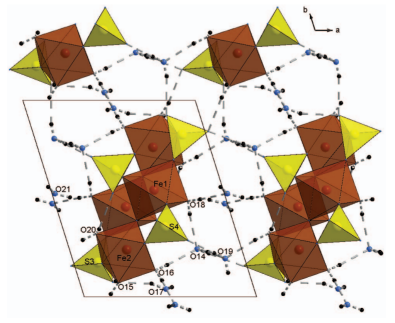


FIG. 2. The hydrogen-bonding system of hohmannite.

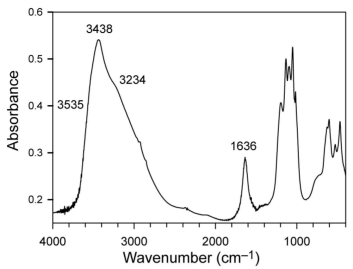


FIG. 3. Room temperature FTIR powder spectrum of hohmannite in the MIR 4000-400 cm⁻¹ region.

of hydroxyl groups in the structure. The first H_2O overtone $(2 \cdot v_3)$ band is observed as a broad and unresolved absorption centred around 6920 cm⁻¹.

In the hohmannite structure there are eight independent H₂O molecules (Fig. 2); these are involved in hydrogen bonds of different strengths with the surrounding oxygens (Table 5), thus reducing their local C_{2v} ideal symmetry (e.g. Nakamoto, 1997). As a consequence, each O–H bond in the H₂O molecule vibrates independently with a frequency related to the local strength of hydrogen bonding. If eight H₂O molecules are considered, 16 asymmetric stretching bands are expected and these overlap to give the broad pattern of Fig. 3. According to the data of Table 5, in the hohmannite structure the different $O-H\cdots O$ hydrogen bond lengths vary in the range 2.632-3.074 Å. If the empirical $O_{donor}\cdots O_{acceptor}$ distance–frequency correlation of Libowitzky (1999) is considered, bands covering a broad range of frequency are expected, between 3500 and 3000 cm⁻¹, and this is what is observed (Fig. 3). The H₂O bending mode occurs as a relatively broad absorption centred at 1636 cm⁻¹ (Fig. 3). The Raman spectrum shows a very weak and broad absorption in the water-

TABLE 7. Positions (wavenumber, cm⁻¹) and proposed assignment for peaks observed in the FTIR and Raman spectra of hohmannite.

Infrared	Raman	Assignment
3438, 3234	3495, 3438, 3292, 3204	$v_3(H_2O), 2 \cdot v_2(H_2O)$
1636		v ₂ (H ₂ O)
1198, 1134, 1096, 1052	1204, 1166, 1125, 1098, 1075, 1058	$v_3(SO_4)$ antisymmetric stretching
1028, 1016	1031, 1018	$v_1(SO_4)$ symmetric stretching
659, 629, 604, 529	659, 628, 605, 580	$v_4(SO_4)$ antisymmetric bending
470	496, 470 400, 334, 258, 245, 227 198, 163, 144, 129, 73, 49	v ₂ (SO ₄) symmetric bending Fe–O,OH modes Lattice modes

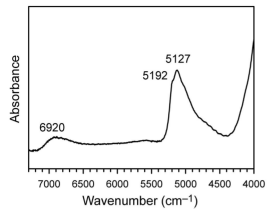


FIG. 4. Single-crystal unpolarized spectrum of hohmannite in the NIR region.

stretching region (Fig. 5) with two components peaked at 3438 and 3204 cm⁻¹, and two evident shoulders at 3495 and 3292 cm⁻¹, respectively.

The powder FTIR spectrum in the low-frequency 1400–400 cm⁻¹ region is given in Fig. 6 in comparison with the Raman spectrum, collected down to 10 cm⁻¹. It is well known that a free SO_4^{2-} ion with ideal T_d symmetry has four modes of vibration (Table 8): the symmetric stretching $v_1(A_1)$ at 983 cm⁻¹, the symmetric bending $v_2(E)$ at 450 cm⁻¹, the antisymmetric

stretching $v_3(F_2)$ at 1105 cm⁻¹ and the antisymmetric bending $v_4(F_2)$ at 611 cm⁻¹ (e.g. Adler and Kerr, 1965; Ross, 1974). In accordance with selection rules, all modes are either Raman alone or Raman and IR active: there are no IR active only or inactive vibrations. In particular, the A_1 non-degenerate symmetric stretching and the doubly degenerate E symmetric bending modes are Raman active only, whereas the triply degenerate asymmetric F_2 stretching and bending modes are both Raman- and infraredactive (Table 8). In a solid sulfate structure, however, the arrangement of the surrounding cations causes distortion of the SO₄ polyhedra away from the T_d site configuration (Adler and Kerr, 1965; Ross, 1974; Della Ventura et al., 2013). As a consequence of this symmetry reduction, we typically observe in the IR pattern a shift of the absorption bands to different energies, the removal of degeneracy (band splitting) and modification in the spectral activity (appearance of ideally non-active IR/Raman modes).

According to Table 8, where a correlation table for the fundamental vibrations of the SO_4^{2-} group in hohmannite, point group C_1 , is presented, 6 IR + 6 Raman bands are expected as both v_3 and v_4 modes, whereas 2 IR + 2 Raman and 1 IR + 1 Raman bands are expected as v_2 and v_1 modes, respectively. Examination of Fig. 6 and

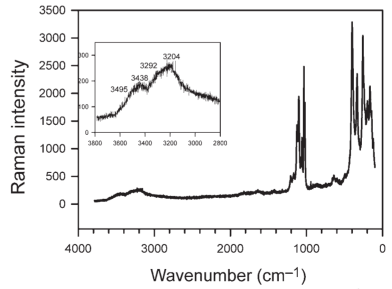


FIG. 5. Unpolarized single-crystal Raman spectrum of hohmannite in the full $4000-20 \text{ cm}^{-1}$ frequency range. The H₂O stretching region is magnified in the inset.

STRUCTURE REFINEMENT OF HOHMANNITE

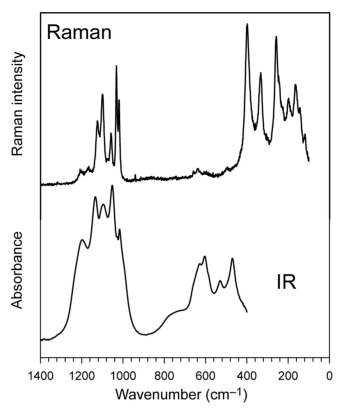


FIG. 6. Comparison between the powder FTIR spectrum and the single-crystal Raman spectrum of hohmannite in the low-frequency region.

Table 7 shows that most of the expected bands are indeed present in the spectra although, due to severe overlapping, some of them appear as shoulders only. In the FTIR pattern, four intense bands are observed in the antisymmetric stretching range at 1198, 1134, 1096 and 1052, while two components at 1028 and 1016 occur as symmetric stretching modes. In the Raman spectrum, three weak bands at 1204, 1166 and 1075, and three intense bands at 1125, 1098 and 1058 cm⁻¹ are

present, while the v_1 modes occur as an intense doublet at 1031 and 1018 cm⁻¹. It is to be noted that, although this mode for the (SO₄²⁻) group is non degenerate (Table 8, see also Adler and Kerr, 1965), the presence of two components in both IR and Raman spectra is in agreement with two independent sulfate sites in the hohmannite structure (Scordari, 1978; this work). The antisymmetric bending modes (v₄) are strong in the IR spectrum whereas they are very weak in the Raman

TABLE 8. SO₄ internal mode analysis in hohmannite by the correlation method.

SO ₄ internal mode	Free ion point group $T_{\rm d}$	Site symmetry C_1	Space group $P\bar{1}$ (C_i)	
ν ₁	A_1	А	$A_{\rm g} + A_{\rm u}$	$A_{\rm g}$ (Raman active)
V ₂	E	2A	$2\dot{A}_{g}+2A_{u}$	$A_{\rm u}$ (IR active)
V3	F_2	3 <i>A</i>	$3A_{g}^{s}+3A_{u}^{s}$	u ()
v ₄	$\bar{F_2}$	3 <i>A</i>	$3A_{g}^{J}+3A_{u}^{J}$	

pattern. The measured Raman range extends at very low wavenumbers ($\leq 10 \text{ cm}^{-1}$) allowing the identification of v_2 (SO₄²⁻), Fe-O, H₂O and lattice modes, following the assignments from previous studies in the literature on sulfates. The v_2 (SO₄²⁻) modes are observed at wavenumbers >400 cm⁻ (e.g. Ross, 1974; Knittle et al., 2001; Murphy et al., 2009; Frost et al., 2013) in the Raman spectrum as very intense absorptions (Fig. 6). Vibrational analysis by the application of the correlation method yields six modes that are Raman active $(6 A_g + 6 A_u)$. The unequivocal assignment of these modes is difficult due to the concomitant occurrence in this spectral range of $v_2(SO_4^{2-})$ and Fe-O,OH modes (Frost et al., 2013; Murphy et al., 2009). Nevertheless, these low wavenumber modes are assigned to the Raman peaks located between 400 and 227 cm⁻¹. Finally, Raman peaks that are found below 200 cm^{-1} can generally be classified as lattice modes (e.g. Frost et al., 2004, 2013).

Conclusion

A new and more accurate structure refinement of the rare Fe sulfate, hohmannite, is reported here, where the hydrogen-bonding scheme has been characterized due to improved XRD data, and confirmed by periodic ab initio calculations. The first vibrational spectroscopic (FTIR + Raman) data for this phase are also reported. The spectra in the principal H₂O stretching region show a multicomponent broad band in the $3700-2800 \text{ cm}^{-1}$ frequency range compatible with the empirical $O_{donor} \cdots O_{acceptor}$ distancefrequency correlation of Libowitzky (1999) based on the structure refinement. Comparison with the spectroscopic data found in the literature (e.g. Omori and Kerr, 1963; Adler and Kerr, 1965; Ross, 1974; Bishop et al., 2004; Cloutis et al., 2006; Lane, 2007; Della Ventura et al., 2013) shows that the vibrational spectra of hohmannite, in particular in the low-frequency $(1400-440 \text{ cm}^{-1})$ region, allow us to distinguish this species from other Fe sulfates. For example, as mentioned above hohmannite may occur associated with amarantite and butlerite. Both FTIR and Raman spectra obtained for hohmannite are significantly different from published spectra of amarantite (Frost et al., 2013) and butlerite/ parabutlerite (Cejka et al., 2011). The distinction between hohmannite and metahohmannite based on spectroscopic data is, however, much more subtle. The behaviour of hohmannite at high

temperature has been examined recently and its decomposition products studied by using a combination of thermogravimetric, XRD and FTIR data (Ventruti et al., 2015). The powder FTIR spectrum observed is not significantly modified in the lower-frequency ($<1300 \text{ cm}^{-1}$) region across the hohmannite-metahohmannite transition which occurs at ~120°C (Scordari et al., 2004, Ventruti et al., 2015). However, the FTIR spectrum in the NIR ($4000-5000 \text{ cm}^{-1}$) region is extremely sensitive to the modification in the H₂O content of the sulfate phase, and can thus be used to distinguish between these two sulfate compounds. Despite these difficulties we remain convinced that it is worth attempting to improve spectroscopic methods to characterize these minerals.

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