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A vibrational spectroscopic study of a hydrated hydroxy-phosphate mineral fluellite, Al₂(PO₄)F₂(OH)·7H₂O



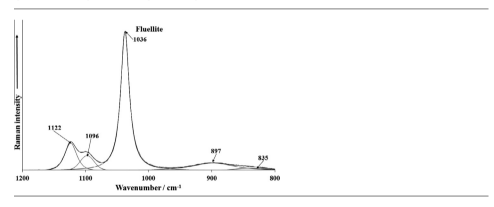
Jiří Čejka ^{a,b}, Jiří Sejkora ^a, Ivo Macek ^a, Ray L. Frost ^{b,*}, Andrés López ^b, Ricardo Scholz ^c, Yunfei Xi ^b

- ^a Department of Mineralogy and Petrology, National Museum, Cirkusová 1740, CZ-193 00 Praha 9, Czech Republic
- b School of Chemistry, Physics and Mechanical Engineering, Science and Engineering Faculty, Queensland University of Technology, GPO Box 2434, Brisbane, Queensland 4001. Australia
- ^c Geology Department, School of Mines, Federal University of Ouro Preto, Campus Morro do Cruzeiro, Ouro Preto, MG 35400-00, Brazil

HIGHLIGHTS

- We have studied Raman and infrared spectra of two well-defined fluellite samples.
- From the Krásno near Horní Slavkov (Czech Republic) and Kapunda, South Australia (Australia).
- Observed bands were assigned to the stretching and bending vibrations of phosphate tetrahedra.
- Approximate O–H···O hydrogen bond lengths were inferred.

GRAPHICAL ABSTRACT



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ABSTRACT

Raman and infrared spectra of two well-defined fluellite samples, $Al_2(PO_4)F_2(OH) \cdot 7H_2O$, from the Krásno near Horní Slavkov (Czech Republic) and Kapunda, South Australia (Australia) were studied and tentatively interpreted. Observed bands were assigned to the stretching and bending vibrations of phosphate tetrahedra, aluminum oxide/hydroxide/fluoride octahedra, water molecules and hydroxyl ions. Approximate O–H···O hydrogen bond lengths were inferred from the Raman and infrared spectra.

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Introduction

Fluellite $Al_2(PO_4)F_2(OH) \cdot 7H_2O$ is very rare, late hydrothermal or supergene mineral formed by alteration of earlier phosphates minerals. It may be found as a colorless to purple-black crystals [1] but usually forms a colorless to white and yellow crystals or powder aggregates in association with fluorapatite, wavellite, cacoxenite,

variscite, strengite, minyulite etc. The size of the crystals is several mm rare up to 1 cm [2]. It occurs in several types of geological environments, for example in complex granitic pegmatites [3,4], phosphatic marbles [5], in lateritic conglomerate and phosphatic sedimentary rocks [6–8] and at ore deposits [1,9–11].

Fluellite was described as a new mineral in 1824 by Lévy [12] without any quantitative chemical tests only with presence of aluminum and fluorine. Further chemical data were presented in 1882 by Groth [13] and proposed the formula AlF₈·H₂O. In 1920 Laubmann and Steinmetz [14] described mineral kreuzbergite from

^{*} Corresponding author. Tel.: +61 7 3138 2407; fax: +61 7 3138 1804. E-mail address: r.frost@qut.edu.au (R.L. Frost).

Oberpfalz, Bavaria as an aluminum phosphate. These two minerals with similar compositions existed till 1940 when Scholz and Strunz [15] carried out qualitative chemical analyses on kreuzbergite and concluded that it has the same composition as fluellite. The name kreuzbergite was discredited and only fluellite remained.

The mineral fluellite has orthorhombic symmetry with space group Fddd. Its crystal structure was solved by Guy et al. [16,17] and consists of octahedrally [Al–O] and tetrahedrally [PO₄] coordinated cations in open framework arrangement within which there are distinct channels containing hydrogen-bonded water molecules. The aluminum atoms are situated at centers of symmetry and are bonded octahedrally to two centro-symmetric pairs of oxygen atoms and one pair of fluorine ions. Raman spectrum of fluellite from Gold Quarry mine, Maggie Creek District, Eureka County, Nevada (USA) was published in the RRUFF's data base (fluellite R070473) without any resolution of band wavenumbers and assignment.

The aim of this paper is to report the Raman spectra of well-defined natural hydrated phosphate minerals, fluellite from two different occurrences, and to relate the spectra to this molecular and the crystal structure. The paper follows the systematic research of the large group of oxyanion containing minerals [18–21], and especially their molecular structure using IR and Raman spectroscopy [22–25].

Experimental

Minerals

The studied samples of the mineral fluellite originated from two different occurrences: greisen Sn-W deposit Krásno near Horní Slavkov [2], western Bohemia, Czech Republic (labelled as CZ) and phosphate deposit Kapunda [26], Mt. Lofty Ranges, South Australia, Australia (labelled as AU). At both occurrences, fluellite forms very brittle, water-clear translucent dipyramidal crystals up to 1 cm (CZ) or only 1 mm in size (AU).

Carefully hand-picked samples were used for X-ray powder diffraction experiments. To minimize the complicated shape of background, the samples studied were placed on a flat low-background silicon wafer. Powder XRD measurements were carried out with Cu K α radiation at a HZG4/Arem diffractometer (50 kV, 40 mA) in the range 5–70° 2θ in the step-scan mode 0.02°/5 s (CZ) and at a Bruker D8 Advance diffractometer (40 kV, 40 mA) in the range 5–70° 2θ in the step-scan mode 0.01°/8 s (AU). The position and intensities of reflections were calculated using the Pearson VII profile shape function in the ZDS program package [27]. The measured patterns were indexed using theoretical pattern calculated from the crystal-structure data of fluellite [16,17]. The unit-cell parameters refined from measured powder XRD using the program of Burnham [28] are compared with published data in Table 1.

The fluellite sample (CZ) was quantitatively analyzed by Cameca SX 100 electron microprobe system in wavelength dispersion mode for chemical composition. Studied sample was mounted into the epoxide resin and polished. The polished surface was coated with carbon layer 250 Å. An acceleration voltage of 15 kV, a specimen current of 10 nA, and a beam diameter of 5 μ m were used. Well-defined natural and synthetic compounds were used as

Table 1Unit-cell parameters of fluellite.

	a [Å]	b [Å]	c [Å]	V [Å ³]
Krásno (CZ)	8.558(1)	11.237(1)	21.179(2)	2043.5(3)
Kapunda (AU)	8.5703(5)	11.2454(8)	21.175(1)	2040.8(1)
Cornwall [16]	8.546(8)	11.222(5)	21.158(5)	2029.12

standards. The raw intensities were converted to the concentrations using automatic *PAP* matrix correction software package. The calculation of theoretical content of H₂O corresponding to ideal formula provided the totals near 120 wt.% (Table 2); it indicates a strong dehydratation, corresponding to loss of two H₂O molecules during sample coating in vacuum and analysis. This loss is indicated by irregular fracturing of the analyzes samples [2]. On the basis of 14 (O,OH,F), empirical formula of fluellite from Krásno may be expressed as Al_{1.98}(PO₄)_{1.07}F_{1.99}(OH)_{0.75}·7H₂O. Chemical composition of Kapunda fluellite (AU) was check by ED spectrum at the same EMPA, obtained ED spectra for both samples (*CZ*,AU) are practically identical, only very minor Fe content was found at AU sample.

Raman and infrared spectroscopy

Fragments of single crystals of fluellite were placed on a polished metal surface on the stage of an Olympus BHSM microscope, which is equipped with $10\times$, $20\times$, and $50\times$ 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 polarised 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 acquisition on the crystals using the highest magnification (50×) were accumulated to improve the signal to noise ratio in the spectra. Spectra were calibrated using the 520.5 cm⁻¹ line of a silicon wafer. Previous studies by the authors provide more details of the experimental technique. Alignment of all crystals in a similar orientation has been attempted and achieved. However, differences in intensity may be observed due to minor differences in the crystal orientation.

Infrared spectrum of fluellite sample from Kapunda was recorded by microdiffuse reflectance method (DRIFTS) on a Nicolet Magna 760 FTIR spectrometer (range 4000–600 cm⁻¹, resolution 4 cm⁻¹, 128 scans, 2 level zero-filtering, Happ–Genzel apodization), equipped with Spectra Tech InspectIR micro FTIR accessory. Sample of amount less than 0.050 mg was mixed without using pressure with KBr. Samples were immediately recorded together with the same KBr as a reference.

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

Table 2Chemical composition of fluellite from Krásno (CZ).

wt.%	Mean 1-4	1	2	3	4	Ideal ^b
Al_2O_3	36.40	36.98	36.32	36.23	36.06	30.89
P_2O_5	27.29	27.10	27.39	26.93	27.74	21.51
F	13.65	13.54	13.37	13.96	13.75	11.51
H_2O^a	47.95	48.95	47.94	47.58	47.30	40.95
-F = O	5.75	5.70	5.63	5.88	5.79	4.86
total	119.54	120.87	119.38	118.82	119.06	100.00

Additional elements (Na, K, Mn, Sr, Ba, Ca, Cu, Zn, Fe, Bi, Si, As, S and Cl) were analyzed; the analysis confirmed their absence or contents below detection limits (ca 0.01-0.05 wt.%).

^a H₂O content was calculated on the basis of ideal composition.

b Ideal composition calculated from formula Al₂(PO₄)F₂(OH)·7H₂O.

undertaken until reproducible results were obtained with squared correlations of r^2 greater than 0.995.

Results and discussion

Crystal symmetry and vibrational spectra of fluellite

Fluellite, Al₂PO₄F₂(OH)·7H₂O, is orthorhombic, space group Fddd – D_{2h}^{24} , Z = 8. The structure consists of AlF₂O₄·H_{3.5} octahedra linked through (PO₄) tetrahedra forming channels which contain the remaining water molecules. The water molecules and hydroxyl ions are hydrogen bonded. Two oxygen atoms in the Al³⁺ octahedra are shared with the (PO₄) tetrahedra. The other two are statistically one quater that of a hydroxyl ion and three quaters that of a water molecule [16,17]. According to Nakamoto [29], octahedral units XY_6 exhibit six normal vibrations v_1 (A_{1g}) and v_2 (E_g) stretching and v_5 (F_{2g}) bending vibrations are Raman active, while only v_3 (F_{1u}) stretching and v_4 (F_{1u}) bending vibrations are infrared active. Symmetry lowering in the case of XY₄Z₂ may cause RA and IR activation of corresponding vibrations and also splitting of degenerate vibrations. Free $(PO_4)^{3-}$ anion exhibits tetrahedral T_d symmetry. In the case of a free ion of T_d symmetry, there are 9 normal vibrations characterized by four fundamental distinguishable modes of vibrations: $v_1(A_1)$ symmetric stretching vibration, Raman active, $v_2(\delta)$ (E) doubly degenerate bending vibration, Raman active, v_3 (F_2) triply degenerate antisymmetric stretching vibration, Raman and infrared active, $v_4(\delta)(F_2)$ triply degenerate bending vibration, Raman and infrared active. T_d symmetry lowering may cause IR activation of the v_1 and v_2 vibrations and splitting of the doubly degenerate v_2 and triply degenerate v_3 and v_4 vibrations. [29,30]. An overlap of stretching and bending vibrations of AlO(OH)F₂ octahedra with stretching and especially with bending vibrations of (PO₄)³⁻ tetrahedra vibrations are supposed. Two fluellite samples were investigated, one sample from the Krásno, Czech Republic (CZ) and one from Kapunda, Australia (AU). RRUFF Raman spectrum of fluellite (specimen R070473 - Gold Quarry mine, Maggie Creek District, Eureka County, Nevada, USA) (cm⁻¹): 1120, 1096, 1038, 910, 651, 585, 524, 462, 406, 313, 276, 211, 173 (Fig. S1). As usually, no interpretation of this spectrum was presented. Tentative assignment and interpretation of the Raman and infrared spectra of fluellite (Table 3) is realized with special regard to [29-33].

Raman and infrared spectroscopy

The Raman spectra of fluellite samples in the 100–4000 cm⁻¹ spectral range are illustrated in Figs. 1a and 1b. These spectra show the position of the Raman bands and their relative intensities. It is obvious that there are large parts of the spectrum where little or no intensity is observed. Therefore, the spectrum is subdivided into sections according to the type of vibration is being investigated. In this way the precise position of the bands can be detailed. The infrared spectrum of fluellite (AU) in the 500–4000 cm⁻¹ spectral range is shown in Fig. 1c. As for the Raman spectrum, the infrared spectrum is subdivided into sections depending upon the type of vibration being examined. The complete infrared spectrum displays the position of the infrared bands and their relative intensity.

Raman and infrared region of v OH stretching vibrations is presented in Figs. 2a–2c. Raman band at $3667 \, \mathrm{cm}^{-1}$ (CZ) and infrared bands $3629 \, \mathrm{and} \, 3559 \, \mathrm{cm}^{-1}$ (AU) are assigned to the v OH stretching vibrations of weakly hydrogen bonded hydroxyls, (OH)⁻. Raman bands at 3396, 3314 and $3124 \, \mathrm{cm}^{-1}$ (CZ) and 3411, 3356, $3222 \, \mathrm{and} \, 3113 \, \mathrm{cm}^{-1}$ (AU) and infrared bands at 3441, $3221 \, \mathrm{and} \, 3047 \, \mathrm{cm}^{-1}$ (AU) are attributed to the v OH stretching vibrations of hydrogen bonded, structurally (symmetrically) distinct water

Table 3Tentative assignment of fluellite spectra.

CZ	AU	AU	USA*		Tentative assignment	
Raman l	Raman	IR	Raman			
3667		3629		{	v OH stretch of (OH)	
	2411	3559		(
2206	3411 3356	3441				
3396 3314	3222	3221		{	v OH stretch of water molecules	
3124	3113	3047				
1670	1675	1660		1		
1070	1073	1624		Į	δH_2O bend	
	1603	1024		1	o 1120 octid	
	1005	1575		(
1583	1503	1537		1	overtones of combination bands	
		1224		1		
		1175		1		
1122	1123	_	1120	1	v ₃ (PO ₄) ³ - antisymmetric stretch	
1096	1083	1102	1096			
	1061	1061		(
1036	1037	1026	1038		ν ₁ (PO ₄) ³ - symmetric stretch	
	1003			1		
	963	965)	S A1 OI I hand	
897	926	920	910)	δ Al-OH bend	
835			880	(
646	638		651	(
	614			{	$v_4(\delta) (PO_4)^{3-}$ out-of-plane bend; $v Al(O(OH)F)_6$ bend	
588	588		585	(
557				(
525	522		524	1	$v_4(\delta) (PO_4)^{3-}$ out-of-plane bend	
513	510			(2	
459			462	{	$v_2(\delta) (PO_4)^{3-}$ bend	
410	410		406	(
	399				ν Al(O(OH)F) ₆ stretch	
2.42	360			{		
342			212			
311 295	297		313	/	O-Al-O skeletal vibrations	
295	297		276		O-AI-O skeietai viorations	
219	279		270	<		
251	249					
220	ムサブ			1		
208			211			
199			211	1		
191	194					
173			173		4.04	
151	153			1	lattice vibrations	
139	141					
123				1		
116	113					
108						

^a USA RRUFF spectrum (R070473) of fluellite from Gold Quarry mine, Maggie Creek District, Eureka County, Nevada (USA).

molecules. Hydrogen bond lengths, $O-H\cdots O$, vary approximately in the range from ~ 3.2 to ~ 2.67 Å [34].

Raman bands at $1670~\text{cm}^{-1}$ (CZ) and $1675~\text{and}~1603~\text{cm}^{-1}$ (AU) and infrared bands at $1660~\text{and}~1624~\text{cm}^{-1}$ (AU) (Figs. 3a–3c) are connected with v_2 (δ) bending vibrations of structurally nonequivalent water molecules. Raman bands at $1583~\text{cm}^{-1}$ (CZ) and $1503~\text{cm}^{-1}$ (AU) and infrared bands at $1575~\text{cm}^{-1}$ and $1537~\text{cm}^{-1}$ (AU) may probably be assigned to overtones or combination bands.

The Raman spectra of fluellite in the $800(900)-1200~\rm cm^{-1}$ spectral range is reported in Figs. 4a and 4b. The Raman spectra are dominated an intense band at $1036~\rm cm^{-1}$ (CZ) and $1037~\rm cm^{-1}$ (AU) and the infrared spectrum (Fig. 4c) an weak band at $1026~\rm cm^{-1}$ (AU) assigned to the $v_1~\rm PO_4^{3-}$ symmetric stretching vibration. The Raman spectrum reported in this paper is in harmony with the spectrum provided in the RRUFF data base. The RRUFF spectrum shows an intense sharp band at $1038~\rm cm^{-1}$. The Raman spectra (Figs. 4a and 4b) show resolved component bands at $1122~\rm and~1096~\rm cm^{-1}$ (CZ) and $1123~\rm 1083~\rm and~1061~\rm cm^{-1}$ (AU) together with the infrared spectrum (AU) those at $1224~\rm 1175$,

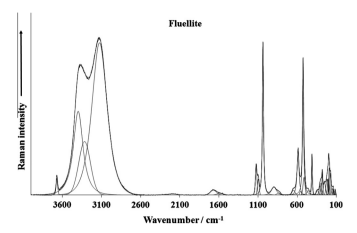


Fig. 1a. Raman spectrum of fluellite sample CZ over the $100-4000~{\rm cm}^{-1}$ spectral range.

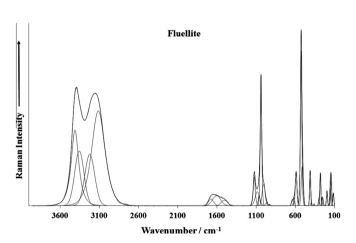


Fig. 1b. Raman spectrum of fluellite sample AU over the $100-4000~{\rm cm}^{-1}$ spectral range.

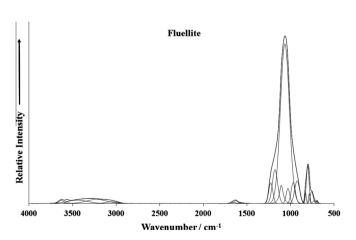


Fig. 1c. Infrared spectra of fluellite AU over the $500\text{--}4000~\text{cm}^{-1}$ spectral range.

1102 and a strong band at 1061 cm $^{-1}$. RRUFF Raman spectrum exhibits bands at 1120 and 1096 cm $^{-1}$. All these bands are assigned to the ν_3 PO $_4^{3-}$ antisymmetric stretching vibrations. There is also a weak Raman band at 1003 cm $^{-1}$ (AU), which may probably be a shoulder to the very intensive band of the ν_1 (PO $_4$) $^3-$ vibration or the δ Al–OH bending vibration. In the Raman spectrum of fluellite (AU) two low intensity component bands are observed at 963

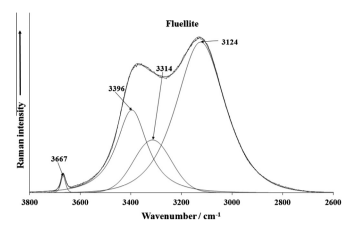


Fig. 2a. Raman spectrum of fluellite CZ over the $2600-3800 \text{ cm}^{-1}$ spectral range.

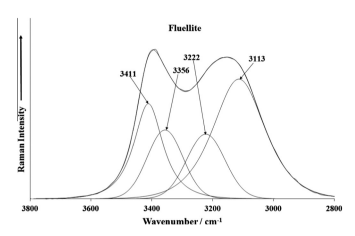


Fig. 2b. Raman spectrum of fluellite AU over the $2600-3800~\text{cm}^{-1}$ spectral range.

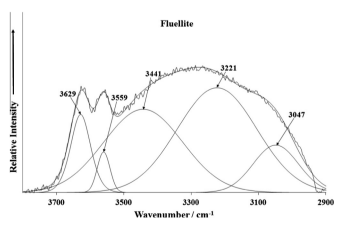


Fig. 2c. Infrared spectrum of fluellite AU over the 2900–3800 cm⁻¹ range.

and 926 cm⁻¹, which are related to the infrared bands (AU) at 965 and 920 cm⁻¹. These bands together with the Raman bands (CZ) at 897 and 835 cm⁻¹ may be attributed to the Al–OH bending modes or to libration modes of water molecules. Normally the intensity of hydroxyl deformation modes are of a quite low intensity in the Raman spectrum but may show significantly greater intensity in the infrared spectrum. In the RRUFF Raman spectrum two broadish weak bands were observed at around 880 and 910 cm⁻¹. It is suggested that these two bands may be due to hydroxyl deformation modes of the AlOH units. However, weak

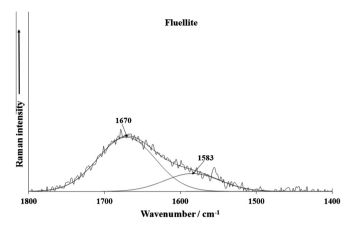


Fig. 3a. Raman spectrum of fluellite CZ over the 1400–1800 cm⁻¹ spectral range.

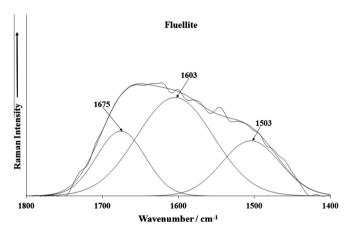


Fig. 3b. Raman spectrum of fluellite AU over the 1400–1800 cm⁻¹ spectral range.

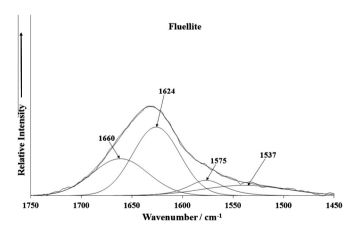
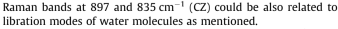


Fig. 3c. Infrared spectrum of fluellite AU over the 1450–1750 cm⁻¹ range.



The infrared spectrum of fluellite in the $850-1300~cm^{-1}$ spectral range is shown in Fig. 4c and shows much greater complexity than the Raman spectra. It is noted that the two infrared bands at 920 and 965 cm⁻¹ attributed to the water librational modes show much greater intensity. Weak infrared bands was found at $1026~cm^{-1}$, which is attributed to the v_1 (PO₄) symmetric stretching vibrations and a strong band at $1061~cm^{-1}$ accompanied with

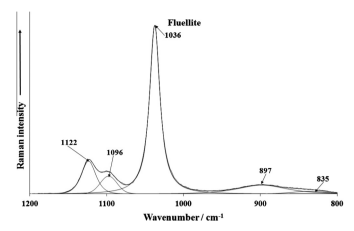


Fig. 4a. Raman spectrum of fluellite CZ over the 800–1200 cm⁻¹ range.

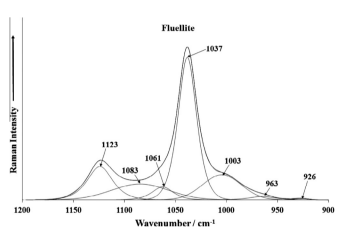


Fig. 4b. Raman spectrum of fluellite AU over the 900–1200 cm⁻¹ range.

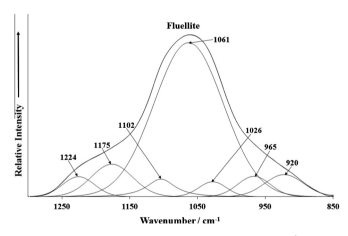


Fig. 4c. Infrared spectrum of fluellite AU over the 850–1300 cm⁻¹ range.

some related weaker bands/shoulders at 1102, 1175 and $1224~cm^{-1}$ may be assigned to the split triply degenerate ν_3 $\left(PO_4\right)^{3-}$ antisymmetric stretching modes.

The Raman spectra of fluellite in the 350(400)– $700 \, \mathrm{cm^{-1}}$ spectral range and in the 100– $350(400) \, \mathrm{cm^{-1}}$ spectral range are displayed in Figs. 5a and 5b and 6a and 6b. Raman bands are observed at 646, 588, 557, 525 and 513 $\mathrm{cm^{-1}}$ (CZ) and 638, 614, 588, 522 and 510 $\mathrm{cm^{-1}}$ (AU) [RRUFF 651, 585, 524 $\mathrm{cm^{-1}}$] are assigned to the v_4 out of the plane bending modes of the $(PO_4)^{3-1}$

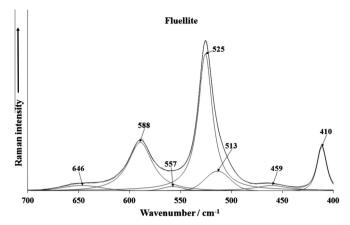


Fig. 5a. Raman spectrum of fluellite CZ over the 400-700 cm⁻¹ range.

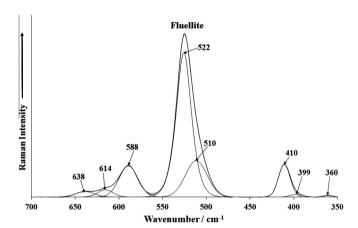


Fig. 5b. Raman spectrum of fluellite AU over the 350-700 cm⁻¹ range.

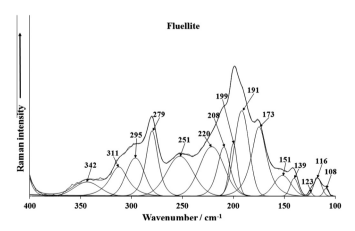


Fig. 6a. Raman spectrum of fluellite CZ over the $100-400 \text{ cm}^{-1}$ range.

units. Some overlap of these bands especially in the range from 585 to 646 cm $^{-1}$ with the v Al(O,OH,F) $_6$ octahedra stretching vibrations may be expected [32]. Raman bands 459 and 410 cm $^{-1}$ (CZ) and 410 cm $^{-1}$ (AU) [RRUFF 462 and 406 cm $^{-1}$] are observed. These bands are attributed to the v $_2$ (PO $_4$) 3 bending modes. Raman bands at 342 and 311 cm $^{-1}$ (CZ) and 399 and 360 cm $^{-1}$ (AU) [RRUFF 313 cm $^{-1}$] may be assigned to the v Al(O(OH)F) $_6$ stretching vibrations. Strong Raman bands are observed at 295, 279 and 251 cm $^{-1}$ (CZ), 297, 279, 258 and 249 cm $^{-1}$ (AU) [RRUFF

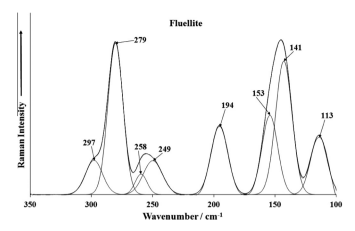


Fig. 6b. Raman spectrum of fluellite AU over the 100-350 cm⁻¹ range.

276 cm⁻¹] are related to the O–Al–O skeletal stretching vibrations. Other Raman bands for fluellite samples studied are observed at 220, 208, 199, 191, 173, 151, 139, 123, 116, 108 cm⁻¹ (CZ) and 194, 153, 141 and 113 cm⁻¹ [RRUFF 211 and 173 cm⁻¹]. These bands are described as lattice vibrations.

Conclusions

- Raman and infrared spectra of two well defined fluellite samples were recorded.
- 2. Observed Raman and infrared bands are tentatively interpreted and assigned to the stretching and bending vibrations of $(PO_4)^{3-}$ tetrahedra and (AlO_4F_2) octahedra, and of vibrations of hydrogen bonded water molecules and hydroxyl ions.
- 3. Approximate O–H···O hydrogen bond lengths are inferred from observed Raman and infrared bands connected with the v OH stretching vibrations of water molecules and hydroxyl ions.

Acknowledgements

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Appendix A. Supplementary materials

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.saa.2014.01.116.

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