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THE CRYSTAL STRUCTURE OF NEAR-END-MEMBER FERROAXINITE FROM AN IRON-CONTAMINATED PEGMATITE AT MALEŠOV, CZECH REPUBLIC

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Abstract

We have characterized ferroaxinite (up to 95 mol.% of end-member composition) from a contaminated granitic pegmatite cutting the magnetite-rich portion of an Fe-skarn body near Malešov, Czech Republic, by means of electron-microprobe analysis, single-crystal X-ray diffraction, infrared spectroscopy and temperature-dependent ⁵⁷Fe Mössbauer spectroscopy. The following crystal-chemical formula was obtained: ^{VI}[Ca_{2.00} Ca_{2.00} (Fe²⁺_{1.83}Mn_{0.10}Mg_{0.06})_{Σ1.99} (Al_{1.97}Fe³⁺_{0.04})_{Σ2.01} Al_{2.00}] ^{IV}[B_{2.00}Si_{8.00}] O₃₀ OH_{2.00}. The crystal structure was refined to R(F) = 0.0184 for 4667 observed reflections with $F_0 > 4\sigma(F_0)$. We obtained the following unit-cell parameters: *a* 7.152(1), *b* 9.206(2), *c* 8.962(2) Å, α 91.84(3), β 98.17(3), γ 77.33(3)°, *V* 569.86(19) Å³. Relative to the other end-members of axinite group, the only significant differences of structural parameters occur at the *Y* site, where Fe²⁺ atoms predominate; the mean *Y*-O bond length is 2.229 Å. The unusually long Fe–O(14) bond (2.704 Å) of the distorted *Y* site suggests a [5 + 1] coordination rather than the previously reported distorted [6] coordination for the group. The scarcity of near-end-member ferroaxinite in nature is due to preferred incorporation of Mn into the structure. The formation of near-end-member ferroaxinite in the late stage of pegmatite evolution at Malešov was favored by high activity of Fe, and lack of Mn and Mg in parent low-temperature fluids.

Keywords: ferroaxinite, axinite group, electron-microprobe analysis, structure refinement, ⁵⁷Fe Mössbauer spectroscopy, granitic pegmatite, Malešov, Czech Republic.

Sommaire

Nous avons caractérisé la ferroaxinite contenant jusqu'à 95% du pôle, découverte dans une pegmatite granitique contaminée recoupant la portion riche en magnétite d'un skarn près de Malešov, en République Tchèque, au moyen d'analyses à la microsonde électronique, diffraction X sur monocristal, spectroscopie infra-rouge et spectroscopie de Mössbauer ⁵⁷Fe à température variable. Nous avons obtenu la formule structurale suivante: ^{VI}[Ca_{2.00} Ca_{2.00} (Fe²⁺_{1.83}Mn_{0.10}Mg_{0.06})_{Σ1.99} (Al_{1.97}Fe³⁺_{0.04})_{Σ2.01} Al_{2.00}] ^{IV}[B_{2.00}Si_{8.00}] O₃₀ OH_{2.00}. Nous en avons affiné la structure jusqu'à un résidu *R*(*F*) de 0.0184 en utilisant 4667 réflexions observées [*F*₀ > 4σ(*F*₀)]. Nous obtenons les paramètres réticulaires suivants: *a* 7.152(1), *b* 9.206(2), *c* 8.962(2) Å, α 91.84(3), β 98.17(3), γ 77.33(3)°, V 569.86(19) Å³. Par rapport aux autres membres du groupe de l'axinite, les seules différences importantes portent sur les paramètres structuraux propres au site *Y*, dans lequel les atomes de Fe²⁺ prédominent; la longueur de liaison *Y*–O moyenne est 2.229 Å. La présence d'une liaison Fe–O(14) anormalement. La rareté de feroaxinite si rapprochée du pôle serait due à la préférence de cette structure pour le Mn. La formation d'une telle composition de ferroaxinite au stade tardif d'évolution

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de la pegmatite à Malešov est favorisée par l'activité élevée de Fe, et le manque de Mn et de Mg dans la phase fluide circulant à faibles températures.

(Traduit par la Rédaction)

Mots-clés: ferroaxinite, groupe de l'axinite, données de microsonde électronique, affinement de la structure, spectroscopie de Mössbauer ⁵⁷Fe, pegmatite granitique, Malešov, République Tchèque.

INTRODUCTION

Near-end-member ferroaxinite was found in a granitic pegmatite cutting an iron-rich skarn body at Malešov, near Kutná Hora, Czech Republic. As the previously published descriptions of the structure of ferroaxinite were obtained from samples with only ~80 mol.% of the ferroaxinite component (Swinnea *et al.* 1981, Andreozzi *et al.* 2004), we report here on the result of a crystal-structure refinement of this near-end-member ferroaxinite, supplemented by electron-micro-probe analyses, infrared and temperature-dependent Mössbauer spectroscopic studies. We describe the crystal chemistry, crystal structure and probable conditions of formation of ferroaxinite, as well as relation-ships among end-members of the axinite group.

BACKGROUND INFORMATION

Minerals of the axinite group typically occur in Ca- and B-rich but relatively Al-poor environments (Pringle & Kawachi 1980, Grew 1996). They have been extensively studied in the past decade (Pieczka & Kraczka 1994, Belokoneva et al. 1997, 2001, Fuchs et al. 1997, Andreozzi et al. 2000a, b, 2004, Salviulo et al. 2000, Novák & Filip 2002, Zabinski et al. 2002); for a comprehensive list of references to earlier work, see the review of Grew (1996). The general formula, (CaMn)₄ (FeMnMg)₂ Al₄ B₂ Si₈ O₃₀ (OH)₂, was proposed by Sanero & Gottardi (1968) and revised by Lumpkin & Ribbe (1979). We use, throughout this work, the site nomenclature according to the generalized structural formula improved by Andreozzi *et al.* (2004): VI[X(1)] $X(2) Y Z(1) Z(2)]_2 IV[T(1) T(2) T(3) T(4) T(5)] O_{30}$ $(OH)_2$, where VI and IV are coordination numbers; X(1)= Ca (Na), X(2) = Ca (Mn²⁺), Y = Fe²⁺, Mn²⁺, Mg (Zn, Fe³⁺, Al), Z(1) = Al (Fe³⁺), Z(2) = Al, T(1)-T(4) = Siand T(5) = B (Si).

The four presently known end-members of the axinite group are classified according to the dominant divalent cation located at the *X* and *Y* sites: ferroaxinite, manganaxinite and magnesioaxinite, all with four Ca atoms per formula unit (*apfu*) and dominant Fe, Mn or Mg at the *Y* site, respectively, and tinzenite with $2 \le Ca \le 4 apfu$ and dominant Mn at both the *X*(2) and *Y* sites (Milton *et al.* 1953, Sanero & Gottardi 1968, Basso *et al.* 1973, Lumpkin & Ribbe 1979, Belokoneva *et al.* 2001, Andreozzi *et al.* 2004). Ferroaxinite and Mg-poor

manganaxinite are the most common compositions in nature (Grew 1996, Andreozzi *et al.* 2000b); however, no previously reported axinite approached closely the ferroaxinite end-member in composition (Grew 1996, Andreozzi *et al.* 2004, Henry *et al.* 2005).

Manganaxinite and tinzenite occur in metamorphosed Mn-deposits and related rocks, as well as in hydrothermal veins and in some occurrences of granitic pegmatite (Ozaki 1972, Grew 1996, Novák et al. 1999). Rare magnesioaxinite is found solely in metamorphic rocks with a high Mg:Fe ratio (Jobbins et al. 1975, Andreozzi et al. 2000a, Novák & Filip 2002). Ferroaxinite is typical in metamorphic rocks and Alpine veins (Ozaki 1972, Pringle & Kawachi 1980, Grew 1996), but it also occurs in granitic pegmatites (Ozaki 1972, Pieczka & Kraczka 1994, Filip 2002). Ferroaxinite with relatively low Mn and Mg contents is a typical accessory mineral in contaminated pegmatites cutting Fe-skarn bodies in the Moldanubicum, Czech Republic, where it is typically associated with plagioclase (An₀₋₅₂), hastingsite to ferro-edenite, clinozoisite-epidote, Ca-rich schorl to uvite, prehnite, titanite and hedenbergite-diopside (Filip 2002).

CRYSTAL STRUCTURE AND CRYSTAL CHEMISTRY OF AXINITE-GROUP MINERALS

The crystal structure of axinite-group minerals (intermediate composition between ferroaxinite and manganaxinite) was originally proposed by Ito & Takéuchi (1952) and subsequently revised by Ito et al. (1969) and Takéuchi et al. (1974). The structural details of a member of the manganaxinite-tinzenite solid-solution series were presented by Basso et al. (1973), and more recently, the structures of relatively pure manganaxinite and of tinzenite were investigated by Belokoneva et al. (1997, 2001), respectively. Ferroaxinite with ~80 mol.% of the end-member component was examined by singlecrystal X-ray diffraction by Swinnea et al. (1981). Andreozzi et al. (2000a) described the structure of near-end-member magnesioaxinite. New structural data for nearly end-member manganaxinite and ferroaxinite with ~80 mol.% of the end-member component were also given by Andreozzi et al. (2004).

The crystal structure of axinite (space group $P\overline{1}$) is described as a sequence of layers roughly parallel to ($\overline{121}$), containing tetrahedrally and octahedrally coordinated cations. The layer of tetrahedra consist of isolated B₂ Si₈ O₃₀ planar clusters (see Takéuchi *et*

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al. 1974, Andreozzi *et al.* 2000a), whereas the layer of octahedra is a continuous framework of six-fold Fe(Mn,Mg)–Al–Al–Al–Al–Fe(Mn,Mg) finite chains [*i.e.*, Y-Z(1)-Z(2)-Z(2)-Z(1)-Y] laterally connected by highly distorted CaO₆ octahedra. The existence of the six-membered ring consisting of two disilicate groups [T(1) and T(2) sites] connected by two BO₄ [T(5)] tetrahedra is typical of the layer of tetrahedra (Takéuchi *et al.* 1974). The hydrogen atom is located between donor O(16) and acceptor O(13) (Swinnea *et al.* 1981, Belokoneva *et al.* 1997, Andreozzi *et al.* 2000a).

Divalent iron enters exclusively the *Y* site, whereas Fe^{3+} is ordered at both *Z*(1) and, to a lesser extent, also at the *Y* site. The *Z*(2) site, markedly smaller than the *Z*(1) site, is invariably fully occupied by Al (Andreozzi *et al.* 2000a, 2004). In ferroaxinite–manganaxinite solid solutions, an increasing Mn content causes enlargement of the *Y* site and subsequently also of the neighboring *Z*(1) site (Andreozzi *et al.* 2004), thus enabling progressive incorporation of Fe³⁺ into the *Z*(1) site mainly in manganaxinite. In manganaxinite–tinzenite solid-solutions, Mn also enters the smaller *X*(2) site (Basso *et al.* 1973, Andreozzi *et al.* 2000a, 2004), whereas the reported presence of Mn in the larger *X*(1) site (Belokoneva *et al.* 1997, 2001) is rather questionable.

The existence of compositional gaps within the axinite group (Pringle & Kawachi 1980, Grew 1996, Andreozzi *et al.* 2000b, 2004) is believed to be caused by both petrological (scarcity of rocks of appropriate composition, Pringle & Kawachi 1980) and structural factors (Andreozzi *et al.* 2000a, b, 2004, Novák & Filip 2002). Nevertheless, there is an almost linear increase of both unit-cell volume and <*Y*–O> mean distance from magnesioaxinite through ferroaxinite to manganaxinite (Andreozzi *et al.* 2000a, 2004, Salviulo *et al.* 2000).

ANALYTICAL METHODS

Sample

The sample of ferroaxinite examined was found in a contaminated granitic pegmatite cutting the magnetiterich portion of a body of Fe-skarn (grossular-andradite, diopside-hedenbergite and magnetite) at Malešov, near Kutná Hora, Czech Republic (the sample studied is from the mineralogical collection of the Institute of Geochemistry, Mineralogy and Mineral Resources, Faculty of Science, Charles University in Prague, Czech Republic; sample no. 19288). The skarn body is located in the Malín Formation of the Gföhl unit, Kutná Hora crystalline unit, Bohemian Massif (Synek & Oliveriová 1993), representing HP-HT rocks metamorphosed during the polyphase Variscan deformation (Beard et al. 1991, Kachlík 1999). The dike of granitic pegmatite, up to 1 m wide, shows a homogeneous internal structure; it is highly contaminated by surrounding iron-rich rocks (magnetite-rich accumulations). The primary minerals of the pegmatite include dominant oligoclase, minor quartz and Fe-rich amphibole (mostly hastingsite). The mineral assemblage with ferroaxinite is developed in small pockets. These formed at the hydrothermal stage, and the following sequence of crystallization was observed: albite \rightarrow chlorite (Al-bearing chamosite) + calcite I \rightarrow ferroaxinite + epidote \rightarrow Fe-rich prehnite + calcite II (Fig. 1). Violet, mostly transparent ferroaxinite forms well-developed isolated axe-shaped crystals (Fig. 1), up to 5 mm in size, with evident striae. Indices of refraction, measured by the immersion method, are: $\alpha 1.677(2)$, $\gamma 1.687(2)$ and $\gamma - \alpha 0.010$.

Electron-microprobe analysis and ⁵⁷Fe Mössbauer and infrared spectroscopy

Electron-microprobe analyses (EMPA) were done using a CAMECA SX100 instrument at the Joint Laboratory of Electron Microscopy and Microanalysis, Institute of Geological Sciences, Masaryk University, Brno, and the Czech Geological Survey, Praha, Czech Republic. The following conditions were used: accelerating voltage of 15 keV, sample current of 10 nA, and beam diameter of 5 μ m. Augite (Si, Mg), andradite (Ca, Fe), jadeite (Na), almandine (Al), rhodonite (Mn), synthetic TiO₂ (Ti) and chromite (Cr) were used as standards. Data were reduced using the X-phi routine (Merlet 1994). Chemical formulae of ferroaxinite were calculated on the basis of 32 anions and from the assumed stoichiometry B = 2, OH = 2 *apfu*.

The ⁵⁷Fe transmission Mössbauer spectra of a powdered sample of ferroaxinite were accumulated in constant-acceleration mode using a ⁵⁷Co-in-Rh source and a 1024-channel detector at room temperature (RT) and at low temperatures (LT), at both 80 K and 25 K. With respect to the strongly overlapped doublets in axinite spectra (*cf.* Fuchs *et al.* 1997, Andreozzi *et al.*



FIG. 1. BSE image of near-end-member ferroaxinite (Ax) from Malešov (Prh: Fe-rich prehnite, Ep: Fe-rich epidote).

2004), we also applied RT ⁵⁷Fe Mössbauer spectroscopy with resonant detection of gamma rays using the moving sample to get narrower spectral lines and a better signal-to-noise ratio than in conventional ⁵⁷Fe Mössbauer spectrometers (see Mashlan *et al.* 2004). Spectra were fitted by Lorentz functions using the computer program CONFIT2000, v. 4.2.3 (Žák 2001). The isomer shift was calibrated with regard to an α -Fe foil. The percentages of the Fe²⁺ and Fe³⁺ contents were established by measuring the integrated area of the respective doublets in the 25 K spectrum (see McCammon 2004).

A Fourier-transform infrared (FTIR) absorption spectrum of ferroaxinite was recorded in the 4000–400 cm⁻¹ spectral region using a Perkin Elmer FTIR 1760X spectrometer equipped with a TGS detector. The sample was prepared by mixing 1 mg of powdered sample with 300 mg of KBr (dried before hand at 120°C) and pressing in an evacuated die at 10 tonnes. A total of 128 scans were done in air with a resolution of 4 cm⁻¹.

TABLE I. SINGLE-CRYSTAL X-RAY DATA, DATA COLLECTION AND DETAILS OF STRUCTURE-REFINEMENT FOR NEAR-END-MEMBER FERROAXINITE FROM MALEŠOV

	Crys	tal data	
$a (Å) b (Å) c (Å) a (°) \beta (°) \gamma (°) V (Å^3)$	7.152(1) 9.206(2) 8.962(2) 91.84(3) 98.17(3) 77.33(3) 569.86(19)	Space group Z F (000) ρ_{cate} (g cm ⁻¹) μ (mm ⁻¹)	PT 2 562 3.312 2.841
Crystal dim	ensions (mm)	0.27 × 0.20 × 0).13

Data collection and processing

Single-crystal four-circle diffractometer Nonius Kappa CCD, Mo tube, graphite monochromator, capillary-optics collimator 300 µm in diameter, program COLLECT (Nonius 2004), scan mode: φ and ω scans, frame width = 2°, crystal-to-detector distance 30 nm, 20_{max} = 70°, T = 293 K, Nonius program suite DENZO-SMN (Otwinowski & Minor 1997), corrections for Lorentz, polarization and background; absorption was corrected according to the multi-scan method (Otwinowski & Minor 1997), Otwinowski *et al.* 2003).

Total number of frames	603
Collection time per degree (s)	10
h, k, l ranges	-11→11, -14→14, -14→14
Total reflections collected	9923
Unique reflections	5012
$R_{\rm int} = \Sigma F_o^2 - F_o^2(\rm mean) / \Sigma F_o^2$	0.0148
"Observed" reflections, $F_o \ge 4\sigma(F_o)$	4667

Refinement

 Full-matrix least-squares on F^2 , program SHELXL-97 (Sheldrick 1997); 242 variable parameters (p), $\Delta/\sigma \le 0.001$

 R1(F) for "observed" reflections

 0.0184

$wR2(F^{*})^{o}$ for all reflections	0.0513
Extinction coefficient	0.0348(10)
a, b ^c	0.02, 0.33
GooF = $\{\Sigma [w(F_0^2 - F_c^2)^2] / (n-p)\}^{0.5}$	1.093
$\Delta \rho_{\min}, \Delta \rho_{\max} (e/Å^3)$	-0.47, 0.53

$$\begin{split} ^{b} & \mathcal{R}1(F) = \Sigma(||F_{o}| \cdots |F_{c}||) / \Sigma F_{o}; \quad wR2(F^{2}) = [\Sigma w(F_{o}^{2} \cdots F_{c}^{2})^{2} / \Sigma wF_{o}^{4}]^{1/2} \\ ^{c} & w = 1 / \{\sigma^{2}(F_{o}^{2}) + [a^{*}\mathsf{P}]^{2} + [b^{*}\mathsf{P}]\}, \quad \mathcal{P} = [\max(0,F_{o}^{2})] + 2^{*}F_{c}^{2} / / 3]. \end{split}$$

X-ray-diffraction experiments and crystal-structure refinement

Data for structure refinement (SREF) were collected with an automatic four-circle single-crystal X-ray diffractometer equipped with a CCD area detector (Nonius Kappa CCD; MoK α radiation, $\lambda = 0.71073$ Å) and with a 300- μ m diameter capillary-optics collimator to provide better resolution. For the collection of intensity data, a suitable fragment of a very good quality and with the approximate dimensions $0.13 \times 0.20 \times 0.27$ mm was cut and mounted on a glass fiber. A full sphere of diffraction data up to $2\theta = 70^{\circ}$ was collected (see Table 1 for details). The intensity data were processed with the Nonius program suite DENZO–SMN, corrected for Lorentz, polarization, absorption and background effects.

The positional parameters for magnesioaxinite given by Andreozzi et al. (2000a) were used as starting parameters for a full-matrix least-squares refinement using SHELXL97 (Sheldrick 1997). For easier comparison with the previously published data on axinite-group minerals, we adopted the cell orientation recommended by Peacock (1937), best representing the unique set defined by the Eisenstein-reduced lattice (see Andreozzi et al. 2000a, Salviulo et al. 2000). The final steps of the refinement resulted in $wR2(F^2) = 0.0513$ for the 5012 unique reflections, and R1(F) = 0.0184 for the 4667 observed reflections. Relevant information on crystal data, data collection, and refinements is given in Table 1. Polyhedron-distortion parameters and polyhedron volumes were calculated by using of the Geometri software (kindly provided by G.B. Andreozzi).

RESULTS

Crystal chemistry: EMPA, ⁵⁷Fe Mössbauer and FTIR spectroscopy

The ferroaxinite from Malešov is chemically homogeneous in the BSE image (Fig. 1). Its chemical composition is very close to the ideal stoichiometry: 7.95 < Si < 8.05 (8.01), 3.92 < Al < 4.01 (3.97) and 3.95 < Ca < 4.04 (3.99) (all in *apfu*; Table 2). The Y site is occupied by dominant Fe²⁺, between 1.77 and 1.95 (1.83) apfu (from a combination of EMPA and Mössbauer spectroscopy; Table 2) and by minor to negligible amounts of Mn, between 0.06 and 0.14 (0.10) apfu and between 0.03 and 0.09 Mg (0.06) apfu. These proportions are relatively constant in all analyzed crystals (Fig. 2). Hence, this sample represents nearly ideal ferroaxinite, (Ca)₄ (Fe)₂ Al₄ B₂ Si₈ O₃₀ (OH)₂ (up to 95 mol.%, Fig. 2), and it is the purest sample described to date (cf. Grew 1996, Andreozzi et al. 2000b, Henry et al. 2005). Minor to trace amounts of Fe^{3+} (0.04 apfu, established from a combination of EMPA, SREF and Mössbauer spectroscopy), and negligible Ti, Cr and Na (less than 0.01 apfu) were found. The stoichiometric



FIG. 2. Composition of ferroaxinite from Malešov in the Mn-Mg-Fe²⁺tot triangle.

content of boron was confirmed from the T(5)-site volume calculated from SREF results, in agreement with results of Andreozzi *et al.* (2000b) (see the next section). Chemical compositions of selected associated minerals also are given in Table 2.

The ⁵⁷Fe Mössbauer spectra of ferroaxinite from Malešov (Fig. 3) are very similar to those of ferroaxinite published in the literature (cf. Fuchs et al. 1997, Andreozzi et al. 2004). They are simpler than the spectrum published by Pieczka & Kraczka (1994) and Zabinski et al. (2002) for ferroaxinite from Strzegom, Poland, where rather unusual substitutions were described. However, their sample obviously contained impurities of associated epidote (unpublished data of J. Filip). Our Mössbauer spectra were fitted with two strongly overlapped doublets of quadrupole splitting (Fig. 3). The Mössbauer parameters refined at RT are: isomer shifts (δ): 1.14 and 0.22 mm/s, and quadrupole splitting (ΔE_0): 2.11 and 0.38 mm/s (0.56 mm/s in case of resonant spectra), respectively. The Mössbauer parameters at LT are: $\delta = 1.27$ and 0.32 mm/s, and ΔE_0 = 2.58 and 0.44 mm/s, respectively. According to the existing literature and results from SREF and EMPA, the two doublets were attributed to ${}^{Y}Fe^{2+}$ and ${}^{Z1}Fe^{3+}$. The δ values are typical of high-spin Fe^{2+} and Fe^{3+} and are within the range for octahedrally coordinated iron in common silicates (cf. Gonser 1975, Dyar et al. 1998, McCammon 2004). The relative areas of absorption in the LT (25 K) spectra are 98% for Fe²⁺ and 2%for Fe³⁺.

The FTIR spectrum (Fig. 4) reflects the great complexity of the axinite structure. The large number

of sharp bands in the 1200 to 400 cm⁻¹ spectral region, corresponding to fundamental Si-O, B-O, Al-O, and Fe-O vibrations, cannot be assigned without uncertainty. Nevertheless, this part of the spectrum is comparable to FTIR spectra of other axinite-group minerals, with no significant differences (e.g., Fuchs et al. 1997). The spectrum in the OH-stretching frequency region consists of a sharp (FWHM ~40 cm⁻¹) absorption band at 3382 cm⁻¹ (Fig. 4) caused by the stretching vibration of the O(16)-H group bonded to O(13). The evident asymmetry of this band (Fig. 4) may imply a slight disorder of the hydrogen bond within the ferroaxinite structure, reflecting substitutions at the coordinating sites (e.g., Fe^{3+} -for-Al). The adjacent small and broad band at around 3485 cm⁻¹ indicates a stretching mode of another O-H bond in the ferroaxinite structure. The high background in the 3000 to 3700 cm⁻¹ spectral region and the weakly pronounced band at ca. 1600 cm⁻¹, typical of the H₂O-bending mode, indicate the presence of free water in fluid inclusions (if water adsorbed on the powdered sample is excluded).

Crystal structure

The unit-cell parameters obtained from SREF (Table 1) are close to the published values for most samples of ferroaxinite. The final coordinates (Table 3) are close to those previously reported for other axinite structures (*cf.* Swinnea *et al.* 1981, Belokoneva *et al.* 1997, Andreozzi *et al.* 2000a, 2004). The complete table of structure factors is available from the Depository of Unpublished Data, CISTI, National Research

	1	2	3	4	5	6	7	8	9	10	
SiO, wt%	42.24	42.18	41.66	42.21	42.24	42.13	38.02	22.31	42 74	69.18	
FiO ₂	0.04	0.01	0.04	0.01	0.00	0.00	0.02	0.07	0.02	0.02	
ALO.	17.70	17.61	17.38	17.93	17.97	17.81	25.21	20.16	20.61	19.76	
Cr ₂ O ₂	0.02	0.01	0.00	0.00	0.00	0.01	0.01	0.02	0.00	0.02	
CaO	19.52	19.70	19.67	19.52	19.81	19.49	22.89	0.00	26.27	0.06	
MgO	0.33	0.11	0.19	0.23	0.15	0.14	0.00	1.12	0.00	0.01	
FeO *	11.58	11.63	12.42	11.59	11.91	11.77	n.d.	45.69	n.d.	n.d.	
$Fe_2O_3 **$	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.	11.56	n.d.	5.43	n.d.	
MnO	0.65	0.78	0.43	0.68	0.47	0.53	0.04	0.31	0.01	n.d.	
Na ₂ O	0.01	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.01	12.20	
B ₂ O ₃ #	6.10	6.09	6.05	6.11	6.13	6.09	-	-	-	-	
H ₂ O "	1.58	1.58	1.57	1.58	1.59	1.58	1.90	10.33	4.26	-	
Fotal	99.77	99.70	99.43	99.88	100.26	99.54	99.65	100.01	99.34	101.25	
FeO ##	11.35	11.40	12.17	11.36	11.67	11.53	-	-	-	-	
Ee ₂ O ₃ ##	0.26	0.26	0.28	0.26	0.26	0.26	~	-	-	-	
Si apfu	8.018	8.025	7.977	8.003	7.988	8.017	3.003	2.591	3.009	2.987	
Гi	0.006	0.001	0.006	0.001	0.000	0.000	0.001	0.006	0.001	0.001	
41	3.960	3.949	3.922	4.007	4.005	3.994	2.347	2.760	1.709	1.006	
Cr	0.003	0.002	0.000	0.000	0.000	0.002	0.001	0.002	0.000	0.001	
Ca	3.970	4.016	4.035	3.966	4.014	3.974	1.937	0.000	1.982	0.003	
Mg	0.093	0.031	0.054	0.065	0.042	0.040	0.000	0.194	0.000	0.001	
Fe ^{2 ; ##}	1.801	1.813	1.949	1.801	1.846	1.836	-	4.438	-	-	
Fe ^{3+ ht}	0.037	0.037	0.040	0.037	0.038	0.037	0.687	-	0.289	-	
Mn ²⁺	0.105	0.126	0.070	0.109	0.075	0.085	0.003	0.030	0.000	-	
Na	0.004	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.002	1.021	
В	2.000	2.000	2.000	2.000	2.000	2.000	-	-	-	-	
Н	2.000	2.000	2.000	2.000	2.000	2.000	1.000	8.000	2.000	-	
E cations	19.997	19.999	20.058	19.994	20.009	19.985	7.979	10.022	6.992	5.019	

TABLE 2. REPRESENTATIVE CHEMICAL COMPOSITIONS OF FERROAXINITE AND OF ACCOMPANYING MINERALS FROM PEGMATITE AT MALEŠOV

* All Fe expressed as FeO. ** All Fe expressed as Fe₂O₃. ² Calculated from stoichiometry. ⁴⁹ Determined by Mössbauer spectroscopy, n.d.: not determined. Columns: 1–6 axinite, 7 Fe-rich epidote, 8 chamosite,

9 Fe-rich prehnite, 10 albite.

Council, Ottawa, Ontario K1A 0S2, Canada. Selected bond-distances for tetrahedral and octahedral sites are presented in Table 4, along with calculated sitevolumes and values of indices of deformation of the coordination polyhedra, for example the mean quadratic elongation (λ) and the angular variance (σ^2), as defined by Robinson *et al.* (1971). As the overall structural features of ferroaxinite (Table 4) are comparable to those of magnesioaxinite and manganaxinite (tinzenite), described in detail by Andreozzi *et al.* (2000a) and Belokoneva *et al.* (1997, 2001), our discussion will focus on the interesting differences (see Discussion).

The SREF indicates complete occupancy of the T(1) to T(4) sites by Si, of the X(1) and X(2) by Ca, and of Z(2) by Al. Hence, Fe³⁺-for-Si, Al-for-Si and B-for-Si substitutions (*cf.* Andreozzi *et al.* 2000a, 2004) are unlikely, and the tinzenite component, according to the vector MnCa₋₁, is present to a negligible extent only. At the Y site, the longest Y–O(14) bond amounts to

2.704 Å, and the second longest *Y*–O(6) bond is 2.372 Å (Table 4). The mean distance of the remaining four bonds is 2.075 Å. The *Z*(1) site, with refined occupancy Al_{0.9837(19)}Fe_{0.0163(19)} (Table 3), is the center of an almost regular octahedron ($\lambda = 1.006$ and $\sigma^2 = 19.4$), with mean <Z(1)–O> = 1.911 Å. The volume of this site (9.216 Å³) is 0.170 Å³ larger than that of the *Z*(2) site (Table 4), occupied solely by Al.

The volume of the T(5) site (1.671 Å³) as well as the values of both distortion parameters ($\lambda = 1.006$, $\sigma^2 = 25.15$) suggest full occupancy of the T(5) site by boron in the ferroaxinite from Malešov. In fact, these values compare well with the respective values of the T(5) site in magnesioaxinite (*cf.* Andreozzi *et al.* 2000a). Moreover, for the axinite group, a T(5)-site volume of *ca.* 1.675 Å³ was found to be characteristic, and indicative of full occupancy by boron (*cf.* Andreozzi *et al.* 2000b).



FIG. 3. Mössbauer spectra of ferroaxinite from Malešov. The two doublets indicate major Fe^{2+} and minor Fe^{3+} (see text).

DISCUSSION

Chemical composition of ferroaxinite and mechanisms of substitution

The chemical composition of ferroaxinite from Malešov is very close to end-member ferroaxinite. It has the following crystal-chemical formula derived from EMPA, ⁵⁷Fe Mössbauer spectroscopy and SREF data: ^{VI}[Ca_{2.00} Ca_{2.00} (Fe²⁺_{1.83}Mn_{0.10}Mg_{0.06})_{Σ1.99} (Al_{1.97}Fe³⁺_{0.04})_{Σ2.01} Al_{2.00}] ^{IV}[B_{2.00}Si_{8.00}] O₃₀ OH_{2.00}. The structurally refined Al:Fe ratio, Al_{0.9837(19})Fe_{0.0163(19}) at the *Z*(1) site, is in good agreement with the small Fe³⁺ content deduced from the 25 K Mössbauer spectrum (2% Fe_{tot}, Al:Fe ratio of Al_{0.981}Fe_{0.019}; *cf*. Table 2). Consequently, the homovalent exchange-vectors observed are: minor MnFe²⁺₋₁ and MgFe²⁺₋₁, and negligible Fe³⁺Al³⁺₋₁. No Fe³⁺ was detected at the *Z*(2) and *Y* sites or in any of tetrahedral sites.

Structure refinement

The previously published refinements of the ferroaxinite structure are based on samples containing relatively high concentrations of Mg (0.21 and 0.30 apfu), but relatively low Mn (0.15 and 0.06 apfu; Swinnea et al. 1981, Andreozzi et al. 2004, respectively). Thus, structural parameters derived from those samples are clearly controlled by the presence of these cations, invariably located at the Y site. The unit-cell parameters of near-end-member ferroaxinite from Malešov (Table 1) are intermediate between those of magnesioaxinite (smaller parameters; cf. Andreozzi et al. 2000a) and manganaxinite (larger parameters; cf. Belokoneva et al. 1997). This is in agreement with the results of Salviulo et al. (2000), who demonstrated the differences in powder-diffraction data among end-members of the axinite group; according to Vegard's law, these reflect the different ionic radii of the octahedrally coordinated divalent cations present at the Y site (Mg: 0.72 Å, Fe: 0.78 Å, Mn: 0.83 Å; Shannon 1976).



FIG. 4. FTIR spectrum of ferroaxinite from Malešov. The ripples at $\sim 2900 \text{ cm}^{-1}$ are due to impurities of organic matter.

Compared to the magnesioaxinite end-member (Andreozzi *et al.* 2000a), the mean *Y*–O distance (2.229 Å) and polyhedron volume at the *Y* site (13.305 Å³) of near-end-member ferroaxinite are markedly larger than those of the Mg end-member (Andreozzi *et al.* 2000a). Similarly, there is markedly increasing deformation of the *Y*O₆ polyhedron from magnesioaxinite to ferroaxinite, reflected by an increase in the σ^2 value from 177.7 (Andreozzi *et al.* 2000a) to 224 (this work). The negligible changes in the bond distances and distortion parameters of the remaining octahedra, and particularly the tetrahedra, are comparable to those of axinite-group phases of various compositions (*cf.* Andreozzi *et al.* 2000a *versus* Table 4).

In accordance with previous studies of the axinite structure (*e.g.*, Belokoneva *et al.* 1997, Andreozzi *et al.* 2000a), the indices of deformation of the Z(1) and Z(2) coordination polyhedra, *i.e.*, λ and σ^2 (Table 4), indicate almost regular octahedra. In excellent agreement with SREF, the Mössbauer parameters of Fe³⁺ (δ and ΔE_Q , see previous section), interpreted above as due to substitution of Fe³⁺ for Al at the Z(1) site, indicate a low distortion of the Z(1) site (*cf.* McCammon 2004).

TABLE 3. ATOM COORDINATES AND EQUIVALENT ISOTROPIC ($U_{\rm eq}$) AND ANISOTROPIC DISPLACEMENT PARAMETERS (Ų) FOR NEAR-END-MEMBER FERROAXINITE FROM MALEŠOV

Atom	x	у	Ζ.	$U_{ m eq}$	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
<i>T</i> (1)	0.21024(4)	0.44997(3)	0.23430(3)	0.00447(5)	0.00501(11)	0.00399(11)	0.00438(12)	0.00039(8)	0.00035(9)	-0.00106(9)
T(2)	0.21917(4)	0.27475(3)	0.52333(3)	0.00379(5)	0.00349(11)	0.00387(11)	0.00406(11)	0.00019(8)	0.00036(8)	-0.00101(8)
T(3)	0.69820(4)	0.25644(3)	0.01151(3)	0.00459(6)	0.00519(11)	0.00459(11)	0.00374(11)	0.00040(8)	0.00070(8)	-0.00041(9)
T(4)	0.64140(4)	0.01946(3)	0.23044(3)	0.00413(5)	0.00392(11)	0.00491(11)	0.00372(11)	0.00019(8)	0.00077(8)	-0.00115(9)
T(5)	0.46146(16)	0.63436(12)	0.28684(12)	0.00474(17)	0.0046(4)	0.0049(4)	0.0047(4)	-0.0004(3)	0.0008(3)	-0.0009(3)
$Z(1)^{*}$	0.05241(4)	0.80071(3)	0.25398(2)	0.00379(9)	0.00382(14)	0.00396(14)	0.00362(15)	0.00019(9)	0.00061(9)	-0.00084(9)
Z(2)	0.35207(4)	0.93593(3)	0.42115(4)	0.00415(6)	0.00418(12)	0.00429(13)	0.00414(13)	-0.00033(10)	0.00059(10)	-0.00133(10)
Y^*	0.76793(2)	0.590928(19) 0.111915(19)	0.00825(5)	0.00543(8)	0.01147(8)	0.00813(8)	0.00305(5)	0.00081(5)	-0.00209(5)
X(1)	0.74654(3)	0.34815(2)	0.39531(2)	0.00780(5)	0.00888(9)	0.00599(8)	0.00774(9)	0.00133(6)	-0.00066(6)	-0.00090(6)
X(2)	0.18294(3)	0.10007(2)	0.08383(2)	0.00845(5)	0.00941(9)	0.00784(9)	0.00767(9)	0.00229(6)	-0.00257(7)	-0.00303(7)
O(1)	0.05360(11)	0.60321(8)	0.18957(9)	0.00670(13)	0.0068(3)	0.0049(3)	0.0078(3)	0.0002(2)	-0.0002(2)	-0.0006(2)
O(2)	0.23119(12)	0.33833(9)	0.09578(9)	0.00882(14)	0.0116(3)	0.0072(3)	0.0077(3)	-0.0021(2)	0.0018(3)	-0.0019(3)
O(3)	0.41844(11)	0.48704(8)	0.31215(9)	0.00670(13)	0.0055(3)	0.0053(3)	0.0095(3)	0.0013(2)	-0.0005(2)	-0.0023(2)
O(4)	0.13537(12)	0.37210(9)	0.36980(9)	0.00996(14)	0.0083(3)	0.0132(4)	0.0087(3)	0.0062(3)	-0.0003(3)	-0.0032(3)
O(5)	0.02163(11)	0.24270(9)	0.56436(9)	0.00626(13)	0.0051(3)	0.0087(3)	0.0057(3)	0.0009(2)	0.0014(2)	-0.0026(2)
O(6)	0.32702(11)	0.38060(8)	0.64393(9)	0.00649(13)	0.0048(3)	0.0061(3)	0.0081(3)	-0.0019(2)	-0.0005(2)	-0.0015(2)
O(7)	0.38056(11)	0.12747(8)	0.49578(9)	0.00539(12)	0.0046(3)	0.0045(3)	0.0067(3)	-0.0014(2)	0.0005(2)	-0.0006(2)
O(8)	0.53476(11)	0.34395(9)	0.87689(9)	0.00692(13)	0.0072(3)	0.0080(3)	0.0048(3)	0.0014(2)	0.0014(2)	0.0007(2)
O(9)	0.87563(11)	0.15555(8)	0.93367(9)	0.00590(12)	0.0057(3)	0.0061(3)	0.0055(3)	-0.0002(2)	0.0017(2)	-0.0001(2)
O(10)	0.76780(12)	0.36672(9)	0.13914(9)	0.00874(14)	0.0113(3)	0.0081(3)	0.0068(3)	-0.0017(2)	0.0006(3)	-0.0028(3)
O(11)	0.60268(12)	0.13512(9)	0.08713(9)	0.00901(14)	0.0100(3)	0.0099(3)	0.0078(3)	0.0044(3)	0.0009(3)	-0.0032(3)
O(12)	0.43624(11)	0.98117(9)	0.24413(8)	0.00590(12)	0.0048(3)	0.0086(3)	0.0051(3)	0.0007(2)	0.0011(2)	-0.0027(2)
O(13)	0.72081(11)	0.10006(9)	0.38469(8)	0.00604(12)	0.0070(3)	0.0071(3)	0.0045(3)	-0.0010(2)	0.0008(2)	-0.0027(2)
O(14)	0.79432(11)	0.87507(9)	0.17743(9)	0.00700(13)	0.0050(3)	0.0081(3)	0.0067(3)	-0.0011(2)	0.0003(2)	0.0008(2)
O(15)	0.32502(11)	0.74622(8)	0.35434(8)	0.00497(12)	0.0046(3)	0.0047(3)	0.0057(3)	-0.0011(2)	0.0011(2)	-0.0008(2)
O(16)	0.09682(11)	0.99539(8)	0.32260(9)	0.00578(13)	0.0047(3)	0.0062(3)	0.0064(3)	-0.0009(2)	0.0014(2)	-0.0007(2)
Н	0.993(3)	0.963(2)	0.631(2)	0.021(5)**					. ,	

* Refined occupancy ratios for Z(1): Al_{0.9837(19)}Fe_{0.0163(19)}; for Y: Fe_{0.9413(19)}Mg_{0.0587(19)}. ** Isotropic refinement U_{iso} .

TABLE 4. BOND LENGTHS (Å) FOR THE OCTAHEDRALLY AND TETRAHEDRALLY COORDINATED SITES OF FERROAXINITE FROM MALEŠOV

$X(1) (Ca) \rightarrow X(1) = O^{>}$ $M \text{ volume}$ λ σ^{2}	$\begin{array}{cccc} O(3) & 2.4382(11)\\ O(5) & 2.3490(12)\\ O(6) & 2.4625(10)\\ O(10) & 2.3364(10)\\ O(13) & 2.3291(9)\\ O(15) & 2.5879(10)\\ & 2.417\\ & 15.545(4)\\ & 1.138\\ & 474.6 \end{array}$	$Z(1) (A1) - C = \frac{Z(1) - C}{M \text{ volume}}$ $\frac{\lambda}{\sigma^2}$	$\begin{array}{llllllllllllllllllllllllllllllllllll$	$T(1) (Si) - $ $< T(1) - O > $ $T \text{ volume} $ $\lambda $ $\sigma^{2} $ $T(2) (Si) - $	$\begin{array}{cccc} O(1) & 1.6190(10)\\ O(2) & 1.5885(10)\\ O(3) & 1.6532(9)\\ O(4) & 1.6385(10)\\ & 1.625\\ & 2.193(1)\\ & 1.003\\ & 10.5\\ \end{array}$
X(2) (Ca)	O(2) 2.2909(10) O(9) 2.3693(11) O(9) 2.4754(10) O(12) 2.2439(12) O(14) 2.3916(10) O(14) 2.3916(10) O(16) 2.5797(10)	Z(2) (Al) -	$\begin{array}{ccc} O(7) & 1.9083(9) \\ O(7) & 1.9197(10) \\ O(12) & 1.8645(9) \\ O(13) & 1.9442(9) \\ O(15) & 1.8692(9) \\ O(16) & 1.8833(10) \end{array}$	< T(2) - O > T volume λ σ^2	$\begin{array}{c} 0(5) & 1.536(6)\\ 0(6) & 1.6546(10)\\ 0(7) & 1.6139(10)\\ & 1.625\\ & 2.170(1)\\ & 1.010\\ \end{array}$
< X(2) - O >	2.392	<z(2) o="" –=""></z(2)>	1.898		37.0
M volume	15.467(4)	M volume	9.049(2)	T(3) (Si) –	O(8) 1.6491(10)
λ	1.119	λ	1.005		O(9) 1.6286(10)
σ^2	375.5	σ^2	18.5		O(10) 1.6038(10)
					O(11) 1.6396(9)
Y (Fe) –	O(1) 2.0889(9)	T(5) (B) –	O(3) 1.4859(14)	<t(3) o="" –=""></t(3)>	1.63
	O(2) 1.9929(10)		O(6) 1.5296(14)	T volume	2.215(1)
	O(6) 2.3717(10)		O(8) 1.4921(14)	λ	1.003
	O(8) 2.1318(9)		O(15) 1.4372(14)	σ	11.1
	O(10) 2.0864(9)	< T(5) - O >	1.486		
	O(14) 2.7036(10)	T volume	1.671(1)	T(4) (Si) –	O(11) 1.6485(10)
<y -="" o=""></y>	2.229	۸ _.	1.006		O(12) 1.6041(8)
M volume	13.305(3)	σ-	25.2		O(13) 1.6371(10)
۸ _2	1.085			<m(a) (d)<="" td=""><td>0(14) 1.6304(10)</td></m(a)>	0(14) 1.6304(10)
σ	224.1			< I(4) - 0 >	1.03
) volume	2.209(1)
				~ ~ ²	1.004
				U	10.2

The distortion parameters λ (the mean quadratic elongation) and σ^2 (the angular variance) are as defined by Robinson *et al.* (1971).

Similarly, relatively large Mössbauer parameters for Fe²⁺ located at the Y site (δ and ΔE_0 , see previous section), indicating a highly irregular site, are in a good accordance with structural evidence of significant distortion of this octahedron (λ and σ^2 , Table 4). Moreover, the Fe-O(14) bond in the YO_6 octahedron (the longest one at 2.704 Å) is not typical for octahedral coordination of Fe²⁺. In the previous studies, this coordination polyhedron was interpreted as a "significantly distorted" or "irregular" octahedron (see Takéuchi et al. 1974, Belokoneva et al. 1997, 2001, Andreozzi et al. 2000a). However, considering the unusually long Y-O(14) bond in all end-members [Mg-O(14) = 2.593]Å, manganaxinite: Mn–O(14) = 2.68 Å, tinzenite: Mn– O(14) = 2.61 Å; Andreozzi et al. 2000a, Belokoneva et al. 1997, 2001, respectively], the distorted octahedral environment of the Y-site cations might be considered as a [5 + 1] coordination rather than the previously reported irregular [6] coordination. Such [5 + 1] coordination has only rarely been reported in the structure of minerals (e.g., in some phosphates, amphiboles and in iron oxides; Chopin *et al.* 2001, Cámara & Oberti 2005, Kelm & Mader 2005, respectively).

On the basis of the stretching frequency of the main OH band in the FTIR spectrum ($\nu = 3382 \text{ cm}^{-1}$), a bond length of about 2.8 Å from the hydrogen donor oxygen O(16) to the acceptor O(13) can be deduced from the distance-frequency correlation of Libowitzky (1999). This length is in very good agreement with the refined O(16)-H...O(13) distance of 2.7794(13) Å. The refined angle O(16)-H...O(13) is $150.7(19)^\circ$, and the (freely) refined O(16)-H bond length is 0.83(2)Å. In order to tentatively explain the additional lowintensity O–H stretching band at *ca*. 3485 cm^{-1} in the FTIR spectrum (Fig. 4), a second possible location of the H atom bonded to O(16) is assumed to be close to the apparent acceptor atom O(6). This position, not previously described from axinite-group minerals, is characterized by the following refined bond-parameters: O(16)-H...O(6) = 3.106 Å, O(16)-H = 0.83 Å, and angle O(16)-H... $O(6) = 125.72^{\circ}$.



FIG. 5. The distorted YO₆ octahedron in the crystal structure of near-end-member ferroaxinite from Malešov (thermal ellipsoids at 99% probability; bond lengths in Å); note the long *Fe*–O(14) bond, suggesting a [5+1] coordination.

Crystal-chemical constraints within the axinite group and origin of the ferroaxinite from Malešov

As was inferred for magnesioaxinite (Novák & Filip 2002), the scarcity of near-end-member ferroaxinite in nature is controlled by the preferred incorporation of the large Mn^{2+} cation at the Y site; hence, axinite generally acts as a sink for Mn (Novák & Filip 2002). The presence of smaller divalent cations, Fe and chiefly Mg, at the Y site causes the contraction of the whole framework (cf. Andreozzi et al. 2000a, 2004), and it could control the scarcity of both near-end-member ferroaxinite and magnesioaxinite compositions in nature. At Malešov, a highly iron-contaminated dike of granitic pegmatite cuts magnetite-rich parts of an iron-rich skarn body, where nearly no Mn was available in parental fluids; thus, near-end-member ferroaxinite formed. In addition, the tendency of axinite to form close to end-member compositions at low temperatures, as well as increased miscibility among all end-members at elevated temperatures (Pringle & Kawachi 1980, Grew 1996), probably also played an important role. Our near-end-member ferroaxinite, associated with chlorite and prehnite, formed at relatively low temperature (well below 300°C, Filip 2002) during the late (hydrothermal) stages of pegmatite evolution at Malešov.

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