Fetiasite (Fe²⁺, Fe³⁺, Ti)₃O₂ [As₂O₅], a new arsenite mineral: Its description and structure determination

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ABSTRACT

Fetiasite, (Fe²⁺, Fe³⁺, Ti⁴⁺)₃O₂As³⁺O₅, is a new mineral found at Pizzo Cervandone, Italy, and Binntal, Switzerland, in the Central Alps. Fetiasite is monoclinic, $P2_1/m$, Z = 2, a = 10.614(2), b = 3.252(1), c = 8.945(1) Å, $\beta = 108.95(2)^\circ$; V = 291.9(2) Å³. The strongest powder-diffraction lines [d_{obs} (Å), (hkl), I/I_0] are as follows: 2.749, (210, $\overline{2}11$),100; 2.814, (202,301),94; 2.391,(110),85; 2.985,($\overline{1}03$),67; 1.779,($\overline{5}04,\overline{5}11$),48; 1.709,($510,\overline{6}03$),35; 1.754,(204, $\overline{6}01$),32; 3.526,(102),31. Fetiasite is brown to black, has a metallic to semimetallic luster, and shows perfect cleavage on {100}. Microhardness VHN₅₀ = 440–490 kg/mm² corresponding to a Mohs hardness of ≈ 5 ; $D_{obs} = 4.6$, $D_{calc} = 4.74-4.80$ g/cm³. In polished section, the mineral appears creamy white; anisotropy is visible along the grain boundaries. Reflectance values (in air) are 15.4–16.0% (470 nm), 15.3–15.8% (546 nm), 14.8–15.2% (589 nm), 14.2–15.0% (650 nm).

Fetiasite and other associated arsenic minerals (e.g., asbecasite, cafarsite, cervandonite) are products of As-bearing solutions remobilized from pre-Alpine ore concentrations (tennantite, chalcopyrite) by Alpine metamorphism.

The crystal structure was solved and refined to r = 0.029 for 1168 observed reflections (Pizzo Cervandone) and to r = 0.023 for 1200 observed reflections (Binntal). The structure contains two As₂O₅ groups, with the As³⁺ cations in trigonal pyramidal coordination, and three nonequivalent octahedral sites, two essentially occupied by Fe and one statistically occupied by Fe and Ti. The octahedra are joined together by sharing edges and corners to form a framework, characterized by the presence of channels running along [010], in which the As atoms are situated.

INTRODUCTION

The new mineral fetiasite was first found by the Italian mineral collector Claudio Albertini in 1986 in the mountain region of Pizzo Cervandone (3155 m above sea level) that forms the border between Italy (Alpe Devero) and Switzerland (Binntal, Canton Valais). This region has become famous since the 1960s for a remarkable number of rare and new minerals, mainly arsenites and arsenates (e.g., asbecasite, cafarsite: Graeser, 1966; Graeser and Roggiani, 1976; gasparite: Graeser and Schwander, 1987; cervandonite: Armbruster et al., 1988). The first specimen of fetiasite was found at about 3000 m above sea level at the eastern slope of Pizzo Cervandone in Italy. Shortly thereafter, the same mineral was detected by Swiss collectors (A. Skrapits, A. Gorsatt, R. Lüssi) about 4 km to the north in two mineral fissures at Gorb in Binntal (formerly well known for the occurrence of spectacular anatase crystals). Material from this locality, though completely different in size and morphology, was essentially identical in other features to that from the Italian occurrence. Therefore, it seems reasonable to give the status of type locality to both occurrences.

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TABLE 1. Physical and optical data of fe	tiasite
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$ \begin{array}{c c} \hline Color: brown to black & Luster: metallic to semimetallic \\ \hline Cleavage: \{100\} perfect & Fracture: uneven to conchoidal \\ \hline Hardness: VNH_{s0} = 438 kg/mm^2 (430-447) \\ VHN_{s0} = 490 kg/mm^2 (487-494) \\ \hline Mohs = \approx 5 (according to VHN) \\ \hline D_{pycn} = 4.6(1) g/cm^3 \rightarrow low because of alteration (locality C) \\ \hline D_{calc} = 4.74 g/cm^3 (locality A); \\ \hline D_{calc} = 4.80 g/cm^3 (locality B) \end{array} $				
Obse	rvations in reflected	I light (Leitz MPV compact)		
	In air	In oil (N = 1.515)		
Color Internal refl. Anisotropy	creamy white none observed visible along grain boundaries	white, with weak brown tint none observed clearly visible		
Bireflectance	not visible	 very weak along grain bound- aries 		
Reflectance values for two isolated crystals at COM wavelengths (SiC standard, minimum and maximum values)				
470 546 589 650	nm: 15.4–16.0% nm: 15.3–15.8% nm: 14.8–15.2% nm: 14.2–15.0%	4.17–5.12% 3.69–4.68% 3.70–4.53% 3.79–4.26%		

The name fetiasite was derived from the mineral's chemical composition Fe-Ti-As oxide. The type material is preserved at the Department of Mineralogy, Natural History Museum, Basel, and at the Mineralogical Institute, University of Basel. The mineral and the mineral name have been accepted by the Commission on New Minerals and Mineral Names, IMA.

GEOLOGICAL SITUATION

The whole mountain range between Alpe Devero and Binntal is built up partly by serpentinites and mainly by gneisses belonging to the Monte Leone nappe, one of the lower units of the Pennine Simplon nappes. The host rocks at the two occurrences are gneisses of variable mineral composition that were influenced by Alpine metamorphism in upper greenschist to lower amphibolite facies; the accompanying Mesozoic calcareous metasediments (Bündnerschiefer) contain abundant kyanite of Alpine origin. The whole region is characterized by a remarkable positive anomaly of the element As. This is manifested by the occurrence of a large number of very special arsenic oxide minerals (arsenites and arsenates), a number of which have been determined as new minerals within the last 25 yr. Presumably, this As anomaly has been caused by a remobilization process during Alpine metamorphism of primary Cu- and As-bearing sulfide ores (tennantite) of Hercynian age, which, in a later state, was responsible for the formation of arsenic sulphosalts in the Triassic dolomite at the famous locality Lengenbach, Binntal (Graeser, 1966; Graeser and Roggiani, 1976). It is not yet clear if there is an age difference between the formation of the arsenites, which were probably formed at an earlier stage, and the arsenates, which are probably the product of subsequent oxidation under atmospheric conditions.

The three occurrences (subsequently designated as A, B, and C) of fetiasite are clearly distinguishable by the



Fig. 1. Fetiasite aggregates (2 mm across) from locality A. Photograph by W. Gabriel.

morphology of the respective crystals. That fetiasite from the three localities is chemically and structurally identical, whereas the host rocks are different, suggests deposition by migrating As-bearing solutions.

Locality A is at Pizzo Cervandone, Italy, about 3000 m above sea level. The first small fetiasite aggregates (up to 2 mm) were found in an Alpine mineral fissure in a light-colored, feldspar-rich, two-mica gneiss of presumably orthogneiss character. The sample contained abundant chlorite (ripidolite), small colorless quartz crystals, and some anatase.

Localities B and C are at Binntal, Switzerland. Fetiasite was detected here in two fissures in dark crystalline rocks ranging from biotite-rich gneisses to mica schists. This is the region that, under the name Lercheltini-Zone, had a worldwide reputation for its spectacular anatase crystals early in this century. The fetiasite crystals from here were collected in the old abandoned clefts that had been mined for anatase about 100 yr ago; surprisingly, these professional mineral collectors ("Strahler") overlooked the new mineral for many years, as well as asbecasite, cafarsite, and cervandonite. The fissures containing fetiasite and these other As minerals are located at about 2000 m above sea level.

PHYSICAL AND OPTICAL PROPERTIES

All fetiasite crystals are covered by a thin brown-red alteration layer that renders the mineral difficult to recognize. Beneath this alteration, fetiasite exhibits the physical properties given in Table 1.

Although the chemical and structural data are identical, the crystal morphology is remarkably different at the three localities: At locality A, only a single hand specimen exists; it consisted of several minute tabular crystals about 0.2 mm in diameter, forming globular aggregates of pinecone shape, rarely exceeding 2 mm in diameter (Fig. 1). On splitting up these aggregates to select single crystals, a perfect cleavage was always observed, on {100}, according to single-crystal X-ray study. At locality B, fe-



Fig. 2. Idealized crystal drawings (Shape) of fetiasite from locality B and locality C; crystal at right is viewed along **b**.

tiasite occurs as radial aggregates (up to 40 mm across) of shingle-shaped crystals of 20 mm maximum length. By analogy with the crystals from locality C (measured by an optical goniometer), fetiasite at locality B is elongated along the **b** axis, with the following forms present: a {100}, c {001}, m {011}. The perfect cleavage on {100} is easily recognizable (see Fig. 2a). At locality C, about 5 m above locality B, characteristic isolated fetiasite crystals reach 1 cm in length and have a peculiar boat-like shape. It was possible to measure a fragment of one of these crystals with an optical goniometer (see Table 2). In agreement with the single-crystal X-ray study, the following forms were identified: $a \{100\}, c \{001\}, m \{011\}$. In contrast to the crystals of the other localities mentioned above, fetiasite from locality C is slightly elongated along the c axis and is platy on {100} (see Fig. 2b). These crystals, like the corresponding ones from localities A and B, are superficially altered to an X-ray amorphous powder of brown-red color.

CHEMICAL COMPOSITION

Preliminary qualitative analyses of the mineral were obtained with SEM-EDS techniques (Cambridge Stereoscan Mark 2A). The EDS analyses revealed the presence of Fe, Ti, and As only, with no substantial quantities of other elements. Comparison of the IR spectrum (Perkin Elmer 883) of fetiasite with the spectra of magnussonite (Långban, Sweden) and mimetite (Tsumeb mine, Namibia) showed fetiasite to be an arsenite mineral, with As as As³⁺ (see Fig. 3).

The chemical analyses were carried out by means of an ARL electron microprobe using the following standards: quartz (Si), corundum (Al), rutile (Ti), FeOOH and fayalite (Fe), MnOOH and spessartine (Mn), AlAsO₄ (As), wollastonite (Ca), and NaF and albite (Na). The results

TABLE 2. Measurements from the optical goniometer

Form	ф _{оbs} (°)	ρ _{obs} (°)	ф _{саю} (°)	P _{calc} (°)
001	90	19.1	90	19.1
100.	90	90	90	90
011	7	70	7.2	70.03

Note: measurements were made on fetiasite, locality C; $a:1:c = \infty:1:$ 2.73, $\beta = 109.1^{\circ}$ (from goniometry); a:1:c = 3.26:1:2.75, $\beta = 108.95^{\circ}$ (X-ray study).



Fig. 3. Infrared spectra of arsenite and arsenate minerals (IR-Laboratorium, MPI, Basel). (a) (IR 049) fetiasite, Cervandone, Italy; (b) (IR 142) mimetite, Tsumeb mine, Namibia; (c) (IR 298) magnussonite, Långban, Sweden.

of the crystal structure determination (see the next section) showed Fe to be present as Fe^{2+} and Fe^{3+} , with a Fe^{2+}/Fe^{3+} ratio of 1.5. The final electron microprobe results are presented in Table 3. Fetiasite crystals from the two localities, Pizzo Cervandone, Italy (A), and Binntal, Switzerland (B), are virtually identical; there is good agreement between the microprobe data and the results of the structure determination.

X-RAY DIFFRACTION STUDY

Single-crystal photographs (Weissenberg and precession techniques) showed monoclinic symmetry, with conditions for nonextinction (0k0: k = 2n) consistent with the possible space groups $P2_1/m$ or $P2_1$; the subsequent structure determination showed $P2_1/m$ to be correct. This is in agreement with the evident centrosymmetric crystal habit of the idiomorphic fetiasite crystals found later (locality C). The powder diffraction pattern is reported in Table 4, and the unit-cell data are reported in Table 5.

The crystal from locality A used for structure determination was a cleavage plate with largest dimensions 0.10×0.09 mm; the crystal from locality B was $0.08 \times$ 0.07×0.10 mm. Both crystals were mounted on an Enraf-Nonius CAD-4 diffractometer, and 25 intense reflections having a θ value in the range 12.6-20.1° were centered using graphite-monochromated MoK α radiation (λ = 0.71073 Å). Least-squares refinement of their setting angles resulted in the unit-cell parameters reported in Table 5, together with an orientation matrix relating the crystal axes to the diffractometer axes. A total of 2940 diffracted intensities for sample A and 2925 for sample B were collected at room temperature with variable scan speed (the maximum scan time for each reflection was 90 s), with $-17 \le h < 17, 0 \le k \le 5$, and $-15 \le l \le 15$, to a maximum 2θ angle of 70°. An ω scan was used in the collection of the intensities of sample A, whereas a $\theta/2\theta$ scan was used for sample B. The diffracted intensities were corrected for Lorentz, polarization, and background

TABLE 3. Chemical composition of fetiasite

TABLE 4.	X-ray	powder	diffraction	data	for	fetiasite
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	Loca	ality A	Loca	lity B	
	1	П	Ш	IV	
FeO	23.12	23.69	24.23	23.84	
Fe ₂ O ₃ *	17.13	17.58	17.93	17.64	
MnO	1.25	1.27	0.89	0.84	
TiO ₂	10.09	10.37	11.17	10.95	
As ₂ O ₃	46.95	47.09	46.76	46.73	
Total	98.54	100.00	100.98	100.00	
Fe ²⁺	1	.38	1.40		
Mn	0	.08	0.05		
Fe ³⁺	0	.92	0.93		
Ti	0	.54	0.58		
Total	2	.92	2.96		
As	2	.0	2.0		
0	7	.0	7.	0	

Note: I = mean of seven individual analyses of fetiasite (locality A); II = $(Fe_{136}^{+}Fe_{032}^{-2}Ti_{0.54}Mn_{0.08})_{\Sigma=2.92}[O_2|As_2O_5]$; III = mean of ten individual analyses of fetiasite (locality B); IV = $(Fe_{140}^{+}Fe_{033}^{+}Ti_{0.58}Mn_{0.05})_{\Sigma=2.96}[O_2|As_2O_5]$. * The ratio $Fe^{2+}/Fe^{3+} = 1.5$ is from structure determination.

effects. An empirical absorption correction was applied according to the procedure already described in Demartin et al. (1992); the minimum transmission factor is 0.42 and 0.63 for A and B, respectively. A total of 1470 independent reflections for A ($R_{av} = 2.4\%$) and 1462 for B ($R_{av} = 1.7\%$) were obtained, and only those with $I > 2\sigma_I$ were considered for the structure solution and refinement.

Scattering factors for neutral atoms, including anomalous dispersion corrections, were taken from Cromer and Waber (1974) and Cromer (1974), respectively. The structure was solved by direct methods using Multan (Germain et al., 1971) and refined by full-matrix least squares, minimizing the function $\Sigma w(F_o - k|F_c|)^2$. Weights assigned to individual observations were $1/\sigma^2(F_c)$. where $\sigma(F_o) = [\sigma_I^2 + (kI)^2]^{\frac{1}{2}}/2F_oLp$, σ_I^2 is the standard deviation for each reflection as derived from counting statistics, k = 0.04 is a coefficient for improving the goodness of fit, and Lp is the Lorentz-polarization factor. Anisotropic displacement parameters were assigned to all the atoms. The refined extinction coefficients were 3.03 \times 10⁻⁷ for A and 1.35 \times 10⁻⁶ for B. The final R and R_w indices were 0.029 and 0.033 for 1168 reflections (sample A) and 0.023 and 0.030 for 1200 reflections (sample B), respectively. The maximum residual in the final difference-Fourier synthesis was 1.7 and 1.2 e/Å³ for A and B, respectively. The fractional atomic coordinates and their estimated standard deviations are reported in Table 6. The anisotropic displacement parameters are reported in Table 7. Selected interatomic distances and angles are reported in Table 8. Observed and calculated structure factor amplitudes for samples A and B are reported in Tables 9A and 9B,1 respectively.

	Lo	ocality A*		Locali	Locality B**		ty C†
hkl	d _{calc}	$d_{\rm obs}$	// I _o	d _{calc}	d _{obs}	d _{calc}	d _{obs}
001	8.493	8.5	12	8.465	8.43	8.45	8.41
102	4.472	4.48	21	4.453	4.45	4.442	4.441
002	4.246	4.25	10	4.233	_	4.223	4.220
301	3.533	2 506	21	3.529	2 5 2 0	3.522	0 505
102	3.518	3.520	31	3.512	3.520	3.506	3.520
300	3.346	3.340	15	3.346	3.344	3.340	3.342
302	3.184	3.183	12	3.175	3.174	3.168	3.171
110	3.104	3.107	21	3.104	3.088	3.086	3.085
103	2.993	2.985	67	2.981	2.981	2.974	2.976
202	2.818	0.014	04	2.816	0.010	2.811	0.010
301	2.814	2.011	94	2.814	2.013	2.816	2.013
211	2.752	0.740	100	2.750	0 740	2.737	0 707
210	2.736	2.749	100	2.736	2.740	2.723	2.131
112	2.637	0.000	45	2.633		2.622	
303	2.627	2.030	15	2.617	_	2.617	_
402	2.559	2.563	27	2.553	2.554	2.548	2.548
212	2.514	0 510	0E	2.510		2.499	0.400
400	2.509	2.510	25	2.509	_	2.505	2.499
112	2.393	2.391	85	2.391	2.383	2.381	2.381
114	1.842	1 926	12	1.837	1 020	1.830	1 922
411	1.834	1.030	13	1.834	1.032	1.829	1.000
504	1.779 🏾	1 770	40	1.773	1 771	1.768	1 760
511	1.775	1.//5	40	1.774	1.771	1.769	1.709
204	1.759	1 754	20	1.756	1 755	1.753	1 754
501	1.754	1.7.54	52	1.753	1.755	1.750	1.704
510	1.710	1 700	25	1.710	1 709	1.705	1 705
503	1.706	1.709	35	1.702	1.700	1.699	1.705
)20	1.632			1.632		1.622	
313	1.628	1.629	20	1.627	1.621	1.622	1.622
502	1.621			1.621		1.618	
220	1.552			1.552		1.543	
214	1.549	1.549	20	1.547	1.541	1.542	1.543
501	1.548			1.548		1.546	
		1.459	32				1.455
		1.431	27				1.423
		1.412	16				1.404
a =	10.625(4)Å		10.615(3	3) Å	10.595(3	3) Å
b =	3.264(2	Å (3.264(1) Å	3.244(1) Å
C =	8.990(6)Å		8.952(3	3) Å	8.931(2	2) Å
$\beta =$	= 109.15(4)°			108.98(2)	0	108.95(5)	0
a:1:c =	3.255:1:2.	754		3.252:1:2	.743	3.266:1:2	.753

Note: lattice parameters were refined from powder diagrams.

* Debye-Scherrer camera, 114.6 mm, FeKα.

** Guinier-IV camera, FeKα1.

† Guinier-Hägg camera, CuKα1.

DESCRIPTION OF THE STRUCTURE

The structure (Fig. 4) can be considered as belonging to the so-called "3-Å fiber axis wallpaper structures" (Moore and Araki, 1974; Konnert et al., 1976). It is characterized by three independent sites occupied by Fe and Ti atoms in octahedral coordination. The octahedra are joined together by sharing edges and corners to form a three-dimensional framework with channels extending along [010]. In these channels, As atoms are bound with trigonal pyramidal coordination to adjacent O atoms.

The As atoms are aligned along [010] at intervals corresponding to the value of the unit-cell parameter b, so that a sequence of atomic coordinates, $x, -\frac{3}{4}, z; x, \frac{1}{4}, z;$ $x, \frac{1}{4}, z$, etc., is observed. Each As atom is bound to two O atoms lying in the same plane normal to **b** (i.e., O5 and O6 for As1 and O1 and O2 for As2) and to a bridging O atom (i.e., O8 connecting two As1 atoms and O7 connecting two As2 atoms, respectively).

¹ A copy of Table 9A and 9B may be ordered as Document AM-94-564 from the Business Office, Mineralogical Society of America, 1130 Seventeenth Street NW, Suite 330, Washington, DC 20036, U.S.A. Please remit \$5.00 in advance for the microfiche.

	Cervandone (A)	Binntal (B)	Binntal (C)
a (Å)	10.614(2)*	10.616(1)*	10.595(3)**
b (Å)	3.252(1)	3.242(1)	3.244(1)
c (Å)	8.945(1)	8.932(1)	8.931(2)
B (°)	108.95(2)	108.89(2)	108.95(5)
V (Å3)	291.9(2)	290.9(2)	290.3(1)
Space group $Z = 2$	P2,/m	P2,/m	P2,/m
* Single-crys ** Refined fro	stal data. om powder data (Gui	nier-Hägg).	

TABLE 5. Unit-cell data for fetiasite, $(Fe^{2+}, Fe^{3+}, Ti)_3O_2[As_2O_5]$

Since the occupation factor for both O7 and O8 is 0.5, we clearly have a case of statistical disorder between empty and occupied sites, thereby excluding the presence of an infinite polyarsenate chain. This disorder is confirmed by other details; for instance, full occupancy of O7 and O8 would result in an O-O distance of 2.687(12) Å. However, such a short distance would be compatible only with the existence of a strong H bond, in this case contradicted by the absence of frequencies due to OH stretching in the infrared spectrum and of residual peaks in the expected position in the final difference-Fourier synthesis. The only way of avoiding this short distance is by short-range ordering of atoms and vacancies, leading to the average occupation of 0.5 that we have observed.

As this model of disorder involving the O atoms bound to the As atoms implies a strong correlation along the same channel, there must be a lack of correlation among different channels; otherwise, the b parameter would be a multiple of the value reported here. However, no additional reflections have been recorded, even from overexposed films in Weissenberg or precession cameras.

For the reasons discussed above, the mineral is clearly an arsenite, with two AsO_3^3 -groups sharing a corner, forming an $As_2O_3^4$ -dimeric group. The geometry matches the corresponding data reported by Hawthorne (1985) for a series of natural arsenites.

DISCUSSION

The chemical formula for fetiasite can be written as Me₃As₂O₇, or Me₃O₂[As₂O₅] if the presence of As₂O₅⁴⁻ groups is emphasized. Unlike the AsO_4^3 ion, AsO_3^3 even in nature easily forms polymeric clusters. For instance, a tetrameric As₄O₈⁴⁻ group occurs in stenhuggarite, Ca-FeSbAs₂O₇ (Coda et al., 1977), whereas indefinite arsenite chains can be observed in trippkeite, CuAs₂O₄ (Zemann, 1951; Pertlik, 1975), and in the corresponding (not isostructural) zinc compound leiteite, ZnAs₂O₄ (Ghose et al., 1987). Similar to fetiasite, dimeric As₂O₅⁻ groups also occur in paulmooreite, Pb₂[As₂O₅] (Araki et al., 1980), and in schneiderhöhnite, Fe²⁺Fe³⁺₃As₅O₁₃ (Hawthorne, 1985). In these minerals, the average As-O distances in the dimeric groups are 1.778(4) and 1.784(2) Å, respectively, which are significantly longer than the corresponding average value in fetiasite [1.767(1) Å].

TABLE 6.	Fractional	atomic	coordinates	and	displacement	pa-
	rameters					

Atom	Mult	x	у	z	$B_{\rm eq}$ (Å ²)
As1	0.50	0.40970(4)	1/4	0.66474(5)	0.717(7)
		0.41010(3)	1/4	0.66453(4)	0.792(5)
As2	0.50	0.24978(4)	1/4	0.24797(5)	0.782(7)
		0.24954(3)	1/4	0.24816(4)	0.831(5)
Fe1	0.50	0.04922(6)	1/4	0.86648(7)	0.89(1)
		0.04912(5)	1/4	0.86648(6)	0.902(8)
Fe2	0.50	0.64241(6)	1/4	0.02199(7)	0.67(1)
		0.64280(4)	1/4	0.02237(5)	0.719(7)
(Fe,Ti)	0.53*	0.88301(9)	1/4	0.39371(9)	1.24(1)
	0.55*	0.88267(6)	1/4	0.39356(7)	1.26(1)
01	0.50	0.0815(3)	1/4	0.1156(4)	1.41(7)
		0.0816(3)	1/4	0.1159(3)	1.41(5)
02	0.50	0.3292(4)	1/4	0.1035(4)	1.50(7)
		0.3287(3)	1/4	0.1036(3)	1.49(5)
03	0.50	0.8322(3)	1/4	0.1581(4)	1.09(6)
		0.8324(2)	1/4	0.1586(3)	1.02(4)
04	0.50	0.9930(3)	1/4	0.6193(4)	1.76(8)
		0.9928(3)	1/4	0.6192(3)	1.67(5)
05	0.50	0.4332(3)	1/4	0.8726(4)	1.26(6)
		0.4335(2)	1/4	0.8722(3)	1.20(5)
06	0.50	0.2348(4)	1⁄4	0.5993(5)	3.2(1)
		0.2347(3)	1/4	0.5990(3)	2.69(8)
07	0.25	0.7279(7)	1/4	0.6769(8)	1.4(1)
		0.7275(5)	1/4	0.6763(6)	1.31(9)
08	0.25	0.5520(9)	1/4	0.3827(9)	2.1(2)
		0.5518(6)	1⁄4	0.3834(7)	1.9(1)

Note: estimated standard deviations are in parentheses. First rows: sample A; second rows: sample B. Isotropic equivalent displacement parameter (B_{eq}) defined as $\frac{4}{3}(a^2B_{11} + b^2B_{22} + c^2B_{33} + ac(\cos\beta)B_{33})$. * Refined as Ti.

In fetiasite, Me essentially corresponds to a mixture of Fe and Ti, and, for that reason, the formula could be alternatively written as $(Fe,Ti)_3O_2[As_2O_5]$. Since these two metals are present in three crystallographic sites, the composition of each of these sites is not necessarily the same as for the others, even in the same sample. Refinement of the occupation parameter of these sites confirmed by the value of the relative bond-length averages (see below) indicates the virtual absence of Ti in the sites indicated as Fe1 and Fe2.

A complication arises from considering the oxidation state of Fe and Ti in the mineral. Since, as we have seen, As is in the +3 state, a total of eight formal electron charges should be assigned to Fe, Ti, and Mn in the three sites. If Ti is all in the +4 state and Mn is in the +2state, this would correspond to a total of $0.55 \times 4 = 2.20$ formal charges for the sample from Cervandone, and the remaining 8 - 2.20 - 0.14 = 5.66 formal charges should be attributed to the Fe atoms, leading to a fractional average oxidation state of 5.66/2.30 = 2.46. This value implies the presence of both Fe2+ and Fe3+; similar cases are encountered in a number of complex oxides of transition elements, both natural and artificial. In the same way, an average oxidation state of 2.39 can be deduced for the sample from Binntal. From these considerations, a Fe²⁺-Fe³⁺ ratio of about 1.5 can be assumed in our specimens.

This estimate for the average oxidation state for Fe may change noticeably if Ti is considered to be in the +3

TABLE 7. Anisotropic displacement factors

Atom	U11	U_{22}	U ₃₃	U_{12}	U ₁₃	U_{23}
As1	0.0088(1)	0.0095(2)	0.0088(1)	0	0.0027(1)	0
	0.0094(1)	0.0109(2)	0.0101(1)	0	0.00359(9)	0
As2	0.0093(1)	0.0100(2)	0.0104(1)	0	0.0032(1)	0
	0.0101(1)	0.0106(2)	0.0114(1)	0	0.00416(9)	0
Fe1	0.0113(2)	0.0102(3)	0.0123(2)	0	0.0039(2)	0
	0.0115(2)	0.0101(2)	0.0131(2)	0	0.0045(1)	0
Fe2	0.0072(2)	0.0084(3)	0.0096(2)	0	0.0021(2)	0
	0.0080(2)	0.0087(2)	0.0107(2)	0	0.0031(2)	0
(Fe, Ti)	0.0174(3)	0.0159(4)	0.0116(3)	0	0.0016(2)	0
	0.0191(2)	0.0144(3)	0.0121(2)	0	0.0022(2)	0
01	0.008(1)	0.024(2)	0.018(1)	0	0.000(1)	0
	0.0100(9)	0.022(1)	0.020(1)	0	0.0027(8)	0
02	0.020(1)	0.023(2)	0.017(1)	0	0.010(1)	0
	0.021(1)	0.021(1)	0.0194(9)	0	0.0135(7)	0
03	0.009(1)	0.022(2)	0.009(1)	0	0.0000(9)	0
	0.0081(8)	0.020(1)	0.0094(8)	0	0.0010(7)	0
04	0.010(1)	0.047(3)	0.009(1)	0	0.0016(9)	0
	0.0100(9)	0.043(2)	0.0100(9)	0	0.0021(7)	0
05	0.012(1)	0.027(2)	0.009(1)	0	0.0031(9)	0
	0.0082(8)	0.027(1)	0.0103(8)	0	0.0025(7)	0
06	0.013(1)	0.091(5)	0.014(1)	0	0.002(1)	0
	0.010(1)	0.077(3)	0.015(1)	0	0.0034(8)	0
07	0.023(3)	0.012(3)	0.017(2)	0	0.005(2)	0
	0.021(2)	0.015(3)	0.014(2)	0	0.005(2)	0
08	0.033(4)	0.026(5)	0.020(3)	0	0.008(3)	0
	0.028(2)	0.024(3)	0.026(2)	0	0.014(2)	0

Note: first rows: sample A; second rows: sample B. The form of the anisotropic displacement parameter is $exp[-2\pi^2(h^2a^{*2}U_{11} + k^2b^{*2}U_{22} +$ $l^2 c^{*2} U_{33} + 2hka^* b^* U_{12} + 2hla^* c^* U_{13} + 2klb^* c^* U_{23}$ where a^* , b^* , and c^* are reciprocal lattice parameters.

state. However, such a situation is very unlikely, since in general the redox equilibrium $Fe^{3+} + Ti^{3+} \Rightarrow Fe^{2+} + Ti^{4+}$ is definitely shifted to the right, and no well-documented occurrence of Ti³⁺ in a terrestrial mineral has been reported.

From the above data and considerations, the theoretical composition (wt%) in terms of the constituent oxides can be deduced for both samples and is reported in Table 3. In spite of all the difficulties encountered, the agreement between the theoretical and the experimental composition is remarkable.

For the (Fe,Ti) position, the average metal to O distance (2.056 and 2.053 Å for the samples A and B, respectively) is not too different from 2.061 and 2.022 Å, the corresponding average distance for ilmenite (Ohgaki et al., 1989) and for taramellite (Mazzi and Rossi, 1980), respectively. Upon applying an empirical calculation (Brown, 1981) to the mean (Fe, Ti)-O distance, the bondvalence estimates range from 2.64 to 3.12, depending on the sample and especially on the kind of atom used as reference. Accordingly, we might reasonably assume that the average oxidation state for the atoms in this site is about +3. This implies that the oxidation state for the Fe atom in this position does not exceed +2.

The same empirical bond-valence calculation applied to the mean Fe1-O distance leads to a value of about 2.11 for the oxidation state of the metal in this site for both samples; for Fe2, the average metal to O bond length is smaller, corresponding to a bond-valence estimate of about 2.46 for both samples. In the latter case, the pres-

Sample	A	В
As1-05	1.793(3)	1.790(2)
As1-06	1.756(4)	1.762(3)
As1-O8°	1.761(3)	1.757(2)
(As1-0)	1.770	1.770
As2-01	1.794(4)	1.792(3)
As2-02	1.757(4)	1.754(3)
As2-07ª	1.746(3)	1.742(2)
(As2-O)	1.766	1.763
Fe1-O1ª	2.142(4)	2.141(3)
Fe1-O1 ^b	2.177(2)	2.173(2)
Fe1-01°	2.177(2)	2.173(2)
Fe1-03 ^d	2.111(2)	2.108(2)
Fe1-O3 ^e	2.111(2)	2.108(2)
Fe1-O4º	2.095(4)	2.093(3)
(Fe1-O)	2.136	2.133
Fe2-029	2.053(2)	2.050(2)
Fe2-02 ^h	2.053(2)	2.050(2)
Fe2-03	1.987(3)	1.987(2)
Fe2-05	2.189(4)	2.193(2)
Fe2-05 ^d	2.160(2)	2.159(2)
Fe2-05°	2.053(2)	2.050(2)
(Fe2-O)	2.083	2.082
(Ti,Fe)-O3	1.998(3)	1.991(2)
(Ti,Fe)-O4	1.976(4)	1.976(3)
(Ti,Fe)-O4	2.118(2)	2.118(2)
(Ti,Fe)-O4'	2.118(2)	2.118(2)
(Ti,Fe)-06 ^a	2.064(3)	2.058(2)
(Ti,Fe)-O6°	2.064(3)	2.058(2)
((Ti,Fe)-O)	2.056	2.053
Note: a = x, y, 1	+ z; b = -x, -y, 1 - z; c	= -x, 1 - y, 1 - z; d =
1 – x, –y, 1 – z; e	x = 1 - x, 1 - y, 1 - z; t =	= 2 - x, 1 - y, 1 - z; g
$= 1 - x_1 - y_2 - z_1$	= 1 - X, 1 - V, -Z; = 1	x, y, -1 + z; = 2 - x,

TABLE 8. Selected interatomic distances (Å) and angles (°)

-y, 1 - z.

ence of an oxidation state intermediate between +2 and +3 is shown even more clearly. For simplicity, the oxidation state in Fe1 can be assumed to be about +2, and the state in Fe2 to be a mixture of +2 and +3.

The presence of atoms of different nature in a certain site can be verified by checking the mean-square displacement along the bond direction (Hirshfeld, 1976). Owing to the particularly rigid nature of most bonds, this displacement should be essentially the same for directly bonded atoms. Deviations in this sense are evidence for the presence of atoms differing either in nature or in the electron configuration; for ligand-metal bonds, these differences are significant if they exceed a limiting value of about 3×10^{-3} Å² (see for instance Bürgi, 1984; Chandrasekhar and Bürgi, 1984). Here, using data from the sample B, for either As1 or As2 the difference in meansquare displacement along each bond with the adjoining O atoms does not exceed four times the estimated standard deviation (σ). In our opinion this difference is hardly significant, since the estimated standard deviation is usually underestimated for various reasons. On the other hand, the corresponding difference for Fe1 and Fe2 amounts to 8 \times 10⁻³ Å², i.e., to 7 σ and to 8 σ , respectively; for the (Fe,Ti) site, it is up to $35 \times 10^{-3} \text{ Å}^2$, i.e., to 19o. This essentially confirms the results obtained from the bond lengths and the occupation parameters: i.e., atoms of different kinds are located in the octahedral sites. The most reasonable assumption is that Ti replaces a



Fig. 4. Poliedri (Pilati, 1990) drawing of the structural framework seen almost along the channel direction.

substantial amount of Fe in the (Fe,Ti) site, whereas the other two metal sites are occupied essentially by Fe atoms in two oxidation states, particularly in the site Fe2.

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