Kuratite, Ca₄(Fe₁₀²⁺Ti₂)O₄[Si₈Al₄O₃₆], the Fe²⁺-analogue of rhönite, a new mineral from the D'Orbigny angrite meteorite

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ABSTRACT

Kuratite, ideally $Ca_4(Fe_{10}^{2+}Ti_2)O_4[Si_8Al_4O_{36}]$, the Fe^{2+} -analogue of rhönite and a new member of the sapphirine supergroup, was identified from the D'Orbigny angrite meteorite by electron microscopy and micro-Raman spectroscopy. Based on the least-squares refinement of 25 d-spacings measured from selected-area electron diffraction patterns of 11 zone axes, the symmetry of kuratite was shown to be triclinic (space group $P\overline{1}$ by analogy to rhönite) with a=10.513(7), b=10.887(7), c=9.004(18) Å, $\alpha=105.97(13)$, $\beta=96.00(12)$, $\gamma=124.82(04)^\circ$, $V=767\pm2$ Å 3 and Z=1 for the 40 oxygen formula. The empirical formula based on eight electron microprobe analyses is $(Ca_{3.88}Na_{0.02}REE_{0.03}^{2+}Mn_{0.03}Mg_{0.01}Ni_{0.02}Zn_{0.01}Sr_{0.01})_{\Sigma4.01}$ $(Fe_{2.98}^{2+}9.9Ti_{2.00})_{\Sigma11.98}(Si_{7.80}Al_{3.52}Fe_{0.64}^{2+}P_{0.05}S_{0.02})_{\Sigma12.03}O_{39.98}F_{0.01}Cl_{0.01}$. The simplified formula is $Ca_4(Fe_{10}^{2+}Ti_2)O_4[Si_8Al_4O_{36}]$. Micro-Raman spectroscopy showed four main bands resembling those of lunar rhönite but with higher frequencies due to different chemical composition. Analogous to the occurrence of kuratite in terrestrial basaltic rocks, kuratite coexisting with Al, Ti-bearing hedenbergite, ulvöspinel, iron-sulfide, tsangpoite, Ca-rich fayalite and kirschsteinite in D'Orbigny angrite most probably was formed at >1000°C by rapid cooling of an interstitial melt, which is subsilicic, almost Mg-free but enriched in Al-P-Ca-Ti-Fe.

KEYWORDS: kuratite, new mineral, angrite, D'Orbigny, rhönite, sapphirine supergroup.

Introduction

THE sapphirine supergroup comprises the sapphirine and surinamite groups, which constitute a polysomatic series (Grew *et al.*, 2008). The sapphirine group is further divided into three subgroups according to the occupancy of the two largest *M* sites at boundaries of octahedral walls in the structure (Bonaccorsi *et al.*, 1990), namely, sapphirine (Mg), aenigmatite (Na) and rhönite (Ca). The rhönite subgroup comprises rhönite

 $Ca_4(Mg_8Fe_3^{2+}Ti_2)O_4[Si_6Al_6O_{36}], dorrite $Ca_4(Mg_6Al_6)$ O_4[Si_8Al_8Fe_3^{3+}O_{36}], serendibite $Ca_4(Mg_6Al_6)$ O_4[Si_6B_3Al_3O_{36}], høgtuvaite $Ca_4(Fe_6^{2+}Fe_6^{3+})O_4$ [Si_8Be_2Al_2O_{36}], makarochkinite $Ca_4(Fe_8^{2+}Fe_2^{3+}Ti_2)O_4[Si_8Be_2Al_2O_{36}], welshite $Ca_4(Mg_9Sb_3^{5+})O_4$ [Si_6Be_3AlFe_2^{3+}O_{36}], the unnamed Ti^{3+}-bearing Mg analogue of rhönite $Ca_4(Mg_7AlTi_2^{3+}Ti_2^{4+})O_4[Si_5Al_7O_{36}]$ and the unnamed Fe^{2+} analogue of rhönite $Ca_4(Fe_{10}^{2+}Ti_2)O_4[Si_8Al_4O_{36}]$ (Grew et $al.$, 2008, and references therein). The Fe^{2+} analogue of rhönite currently listed as a valid unnamed mineral with IMA code $UM1972-18-SiO:AlCaFeMgTi$ (Smith and Nickel, 2007, and updates) has now been formally named kuratite in honour of $Professor$$

* E-mail: slhwang@mail.ndhu.edu.tw DOI: 10.1180/minmag.2016.080.043 Dr Gero Kurat (1938-2009), former Head of the Mineralogical-Petrographical Department and Curator of the Meteorite Collection at the Natural History Museum in Vienna, Austria. He was a pioneer in meteorite research and a gifted mineralogist and petrologist. He identified the unusual D'Orbigny meteorite as an angrite. The new mineral and its name were approved by the IMA-CNMNC (IMA 013-109). Type material is deposited in the collections of the Naturhistorisches Museum Wien, Vienna, Austria, inventory number Section D'Orbigny C-N1172-NH Wien. Here we report the description and characterization of the new mineral kuratite, ideally $Ca_4(Fe_{10}^{2+}Ti_2)$ $O_4[Si_8Al_4O_{36}]$, the Fe^{2+} analogue of rhönite. The origin and formation conditions of kuratite in the angrite meteorite are also addressed, based on the available information on terrestrial kuratite occurrences.

Occurrence and appearance

Kuratite occurs in D'Orbigny, a 16.5 kg angrite meteorite found in July 1979 in Argentina. The polished angrite slab sample c. 1.5 cm \times 1.5 cm \times 0.3 cm in size used for the present study was kindly loaned by the Naturhistorisches Museum Wien, Austria. Petrography of the D'Orbigny angrite has been described in detail by Mittlefehldt et al. (2002), Kurat et al. (2004) and Varela et al. (2003). In addition to the presence of some hollow shells and open vesicles of various sizes and shapes with occasional glass filling, D'Orbigny angrite consists of the following silicate phase assemblage: tabular crystals of nearly pure anorthite, euhedral to anhedral augite crystals with Mg-free and Al, Ti-bearing hedenbergite rims, euhedral to anhedral olivine crystals with Mg-rich cores (Mg ~64 mol.%) and complex Ca-rich fayalite + kirschsteinite overgrowths toward the rims (Fig. 1a,b). The whole microstructure can be characteristically described as olivine + augite aggregates with highly irregular outlines suggestive of brittle fracture (as circled in Fig. 1a) surrounded by anorthite crystals. Pockets consisting of several phases with an overall brighter contrast in scanning electron microscope (SEM) back-scattered electron (BSE) imaging due to Fe sulfides and ulvöspinel are common along grain boundaries within the olivine + augite aggregates (Fig. 1a,b). Detailed SEM observations indicate that the multiple phase pockets sit mainly at olivine-augite triple junctions, and consist of: (1) Fe-sulfide spherules

and euhedral ulvöspinel; (2) fibrous Na-bearing anorthite; (3) fibrous whitlockite, which was described as Ca-phosphate (Mittlefehldt et al., 2002); (4) the new mineral tsangpoite, approved recently by IMA-CNMNC and to be published in a separate paper (IMA 2014-110, Ca₅(PO₄)₂SiO₄, hexagonal; Hwang et al., 2015), and previously described as silico-phosphate (Mittlefehldt et al., 2002; Kurat et al., 2004); (5) the new mineral kuratite, previously described as Fe-Ca-Al-Ti silicate (Mittlefehldt et al., 2002), or Fe-Al-Ti silicate (Kurat et al., 2004), or 'rhönite' (Jambon and Boudouma, 2011); and (6) Al, Ti-bearing hedenbergite filling vesicles (Fig. 1). Kuratite crystals are typically euhedral to anhedral, < 20 µm in size, and occur, together with whitlockite and submicrometre-sized Fe sulfide, within a rim of olivine of Ca-rich favalite composition $(Ca \approx 20 \text{ mol.\%}) + \text{kirschsteinite} (Ca \approx 40 \text{ mol.\%})$ intergrowths at the contact between Fe-sulfide spherules and the Al, Ti-bearing hedenbergite filling vesicles (Fig. 1c-e). Such intergrowths were not observed at the contacts without hedenbergite between Fe-sulfide spherules and the olivine overgrowth rim enriched with Fe/Ca (Fig. 1f). Whitlockite inclusions in kuratite are common (Fig. 1d,e).

Chemical data

Chemical analyses were carried out using a JEOL JXA8500-F FE-electron microprobe at Academia Sinica, Taiwan (wavelength-dispersive spectroscopy mode, 12 kV, 5 nA, 2 µm beam diameter). Natural and synthetic standard minerals and diffracting crystals (in parenthesis) used for calibration were diopside for $SiK\alpha$ (TAP), rutile for $TiK\alpha$ (PET), corundum for Al $K\alpha$ (TAP), chromium oxide for $CrK\alpha$ (PET), hematite for $FeK\alpha$ (LiF), tephroite for MnK α (PET), periclase for MgK α (TAP), Ni oxide for NiKα (LiF), wollastonite for $CaK\alpha$ (PET), albite for $NaK\alpha$ (TAP), adularia for $KK\alpha$ (PET), apatite for $PK\alpha$ (PET), fluorite for $FK\alpha$ (TAP), tugtupite for $ClK\alpha$ (PET), celestite for $SK\alpha$ (PET), Zn oxide for $ZnK\alpha$ (LiF), celestite for $SrL\alpha$ (PET), YAl garnet (YAG) for YL α (PET), LaP₅O₁₄ for LaLα (LiF), PrP₅O₁₄ for PrLβ (LiF), NdP₅O₁₄ for $NdL\beta$ (LiF). Counting times for the peak of each element and backgrounds both above and below the peak were 20 and 10 s, respectively. Eight analyses of kuratite and their average are given in Table 1a. The empirical formula (atoms per formula unit) of the average, based on 28 cations and 40 oxygens, is

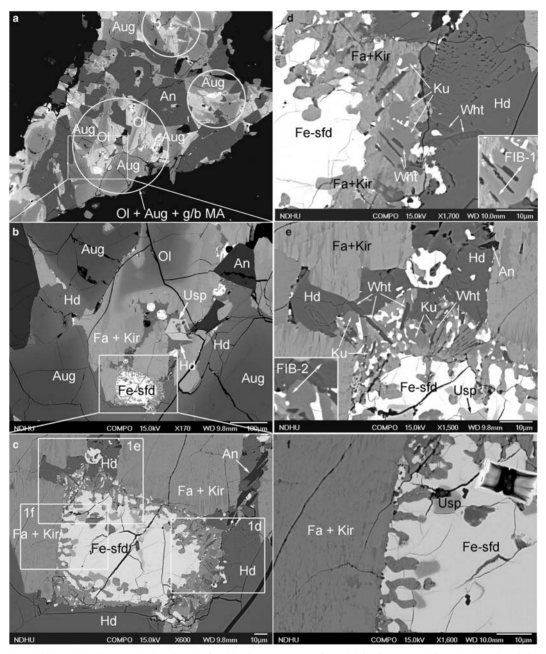


Fig. 1. SEM-BSE micrographs showing (a) the ubiquitous presence of a multiple phase assemblage (MA) at the triple-junction of an olivine (OI) + augite (Aug) grain aggregate (circled) with a highly irregular outline; (b) the main constituents of the multiple-phase assemblage, including Fe sulfide (Fe-sfd), ulvöspinel (Usp) and the Al, Ti-bearing hedenbergite (Hd), with complex Ca-rich fayalite (Fa) + kirschsteinite (Kir) overgrowths toward the olivine rims; (c), (d) and (e) kuratite (Ku) and whitlockite (Wht) crystals surrounded by Fe/Ca-olivine intergrowth at the periphery of Fesulfide spherule facing hedenbergite; (c) and (f) the absence of Ku + Wht at the Fe-sulfide boundary facing olivine. Note that fibrous whitlockite crystals are frequently enclosed by kuratite (d). Two FIB thin sections for electron diffraction study of kuratite structure are shown as inserts in (d) and (e).

TABLE 1a. Chemical composition (wt.%) of kuratite from D'Orbigny angrite.

	B5-a-3	B5-a-4	B5-a-5	B5-a-6	B5-a-7	B5-a-8	B5-a-9	B5-a-10	Average
SiO ₂	25.37	25.16	25.95	25.06	26.39	25.21	25.50	25.80	25.55
TiO_2	8.43	9.17	8.52	8.51	9.78	8.68	8.33	8.15	8.70
$Al_2\tilde{O}_3$	10.32	10.18	9.53	9.35	8.86	10.29	10.01	9.90	9.80
Cr_2O_3	0.00	0.00	0.00	0.00	0.00	0.00	0.11	0.00	0.01
Y_2O_3	0.08	0.00	0.05	0.04	0.03	0.00	0.00	0.00	0.03
La_2O_3	0.23	0.00	0.00	0.00	0.00	0.00	0.03	0.07	0.04
Pr_2O_3	0.15	0.00	0.22	0.00	0.18	0.12	0.06	0.00	0.09
Nd_2O_3	0.18	0.01	0.00	0.00	0.13	0.21	0.00	0.00	0.07
FeO Total	42.01	41.09	40.98	41.76	41.95	41.86	41.60	41.57	41.60
MnO	0.20	0.02	0.04	0.13	0.06	0.03	0.24	0.15	0.11
NiO	0.01	0.00	0.15	0.00	0.00	0.19	0.00	0.18	0.07
SrO	0.06	0.08	0.04	0.04	0.05	0.03	0.00	0.06	0.05
ZnO	0.00	0.00	0.00	0.00	0.22	0.00	0.03	0.10	0.04
MgO	0.00	0.02	0.00	0.06	0.00	0.02	0.00	0.01	0.01
CaO	11.92	11.86	11.84	11.78	11.56	12.10	11.96	11.82	11.86
Na ₂ O	0.00	0.02	0.13	0.03	0.01	0.00	0.07	0.04	0.04
K_2O	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.01	0.00
P_2O_5	0.27	0.16	0.20	0.23	0.08	0.17	0.19	0.27	0.20
F	0.00	0.00	0.00	0.00	0.00	0.02	0.03	0.00	0.01
C1	0.03	0.00	0.02	0.04	0.03	0.04	0.00	0.01	0.02
SO_3	0.09	0.07	0.06	0.13	0.08	0.05	0.09	0.05	0.08
Total	99.36	97.82	97.72	97.16	99.42	99.01	98.23	98.19	98.38

 $\begin{array}{l} (Ca_{3.88}Na_{0.02}REE_{0.03}^{3+}Mn_{0.03}Mg_{0.01}Ni_{0.02}Zn_{0.01}\\ Sr_{0.01})_{\Sigma4.01}(Fe_{9.98}^{2+}Ti_{2.00})_{\Sigma11.98}(Si_{7.80}Al_{3.52}Fe_{0.64}^{3+}P_{0.05}\\ S_{0.02})_{\Sigma12.03}O_{39.98}F_{0.01}Cl_{0.01} \ (Table\ 1b;\ REE=\ rare-earth\ minerals).\ Most\ analyses\ show\ the\ presence\ of\ 5-8\%\ Fe^{3+}\ among\ all\ Fe\ analysed.\ The\ simplified\ formula\ is\ Ca_4(Fe_{11}^{2+}Ti_2)O_4[Si_8Al_4O_{36}]. \end{array}$

Crystallography

Two transmission electron microscopy (TEM) samples of dimensions $5 \mu m \times 5 \mu m \times 100 nm$ were prepared by the focused ion beam (FIB) technique from two kuratite grains (inserts in Fig. 1d,e). A JEOL 3010 analytical electron microscope (AEM) was used for imaging and energy-dispersive X-ray (EDX) analyses. Selectedarea electron diffraction (SAED) patterns of kuratite were taken at various tilt angles for phase identification. A representative TEM bright field micrograph showing a kuratite crystal with abundant polysynthetic nano-twins and whitlockite inclusions in thin section FIB-1 is shown in Fig. 2a. Surrounding Fa+Kir intergrowths can also be clearly seen. The twin plane is $(0\overline{1}1)$, similar to that reported for other rhönite subgroup minerals.

In contrast, the kuratite crystal in thin section FIB-2 is twin-free and inclusion-free, and is associated with euhedral whitlockite and a rounded Fe sulfide within a polycrystalline hedenbergite matrix (Fig. 2b). Kuratite is bounded by small faces, such as $(0\overline{1}1)$, $(0\overline{2}1)$, $(0\overline{3}1)$ etc., as indicated in Fig. 2c.

Unit-cell parameters of kuratite were determined from SAED patterns. The error of the d-spacing measurements on electron diffraction patterns taken at a camera length of 120 cm and calibrated by an Al standard was estimated to be ± 0.002 nm. Based on the least-squares refinement of 25 d-spacings (Table 2) measured from eleven diffraction patterns of the twin-free kuratite crystal in section FIB-2 (insert in Fig. 1e; Fig. 2c), including the six shown in Fig. 3a-f and five not shown ($[0\overline{1}\ \overline{1}]$, $[0\overline{1}\ \overline{2}]$, $[0\overline{1}\,\overline{3}]$, $[10\overline{2}]$, $[\overline{1}\,\overline{2}\,\overline{2}]$), unit-cell parameters of kuratite were determined to be very similar to those of other rhönite subgroup minerals (see unitcell data below; Table 3). In comparison to rhönite with abundant Mg2+ (Table 3), the slightly larger unit-cell parameters of kuratite must be due to Fe²⁺ replacing Mg²⁺ at the octahedral sites. Note that the double diffraction effect in electron diffraction usually prohibits quantitative determination of the

Table 1b. Calculated formula (atoms per formula unit) for the averaged composition of kuratite.

Constituent	wt.%	Symbol	No. ions	No. O atoms
SiO ₂	25.55	Si ⁴⁺	7.796	15.593
TiO ₂	8.70	Ti ⁴⁺	1.997	3.993
$Al_2\tilde{O}_3$	9.80	$A1^{3+}$	3.524	5.286
Cr_2O_3	0.01	Cr^{3+}	0.002	0.004
Y_2O_3	0.03	Y^{3+}	0.005	0.007
La_2O_3	0.04	La ³⁺	0.005	0.007
Pr_2O_3	0.09	Pr^{3+}	0.010	0.015
Nd_2O_3	0.07	Nd^{3+}	0.008	0.011
ΣFeO	41.60	ΣFe	10.615	10.933
FeO	39.10	Fe^{2+}	9.978	9.978
Fe ₂ O ₃	2.77	Fe ³⁺	0.637	0.955
MnO	0.11	Mn^{2+}	0.028	0.028
NiO	0.07	Ni ²⁺	0.017	0.017
SrO	0.05	Sr^{2+}	0.009	0.009
ZnO	0.04	Zn^{2+}	0.009	0.009
MgO	0.01	Mg^{2+}	0.005	0.005
CaO	11.86	Ca^{2+}	3.877	3.908
Na ₂ O	0.04	Na ⁺	0.024	0.012
K_2O	0.00	K^{+}	0.000	0.000
P_2O_5	0.20	P^{5+}	0.052	0.129
F	0.01	F^{-}	0.010	
Cl	0.02	Cl ⁻	0.010	
SO_3	0.08	S^{6+}	0.018	0.055
O = Cl, F	-0.0087			
Total wt.%	98.37			
Σ				39.991
O = Cl, F				-0.010
Σ (O atoms)				39.981
Σ (cations)			27.999	
Σ (anions)				40.001
-				

relative intensities among the reflections. This is especially true for the low-order zone axis patterns such as $[00\overline{1}]$ and $[\overline{1}\ \overline{1}\ \overline{2}]$, in which many reflections are present and double diffraction effects prevail (Fig. 3a,b). However, for the high-order zone-axis patterns with fewer reflections, the strong reflections can be nonetheless ascertained by careful examination, such as $22\overline{1}$ in the $[\overline{1}\ \overline{1}\ \overline{4}]$ zone, $03\overline{1}$ and $\overline{2}5\overline{1}$ in the $[\overline{1}\ \overline{1}\ \overline{3}]$ zone, $\overline{2}4\overline{1}$ in the $[0\overline{1}\ \overline{4}]$ zone, $\overline{1}1\overline{1}$ in the $[11\overline{3}]$ zone from section FIB-2 (Fig. 3c-f), as well as $4\overline{2}0$, $20\overline{3}$ and $21\overline{3}$ in the [362] and [364] zone axis patterns from section FIB-1 (Fig. 3g,h). It is no surprise that these strong reflections of kuratite are among the set of strong reflections in the X-ray diffraction data of makarochkinite, $Ca_4(Fe_8^2+Fe_2^3+Ti_2)O_4[Si_8Be_2Al_2O_{36}]$

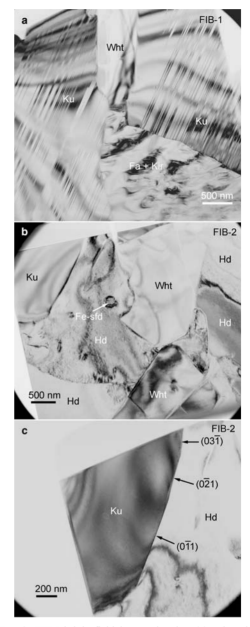


Fig. 2. TEM bright-field image showing (a) a kuratite crystal in thin section FIB-1 with abundant polysynthetic (011) nano-twins, which is associated with a whitlockite inclusion in a matrix of Fe/Ca-olivine intergrowths; (b) and (c) the twin-free kuratite crystal with (011), (011) and (011) facets in thin section FIB-2, with the associated euhedral whitlockite crystals, Fe sulfide, and the polycrystalline hedenbergite matrix.

Table 2. Observed (obs) and refined (ref) *d*-spacings (Å) for kuratite from section FIB-2.

hkl	$d_{ m obs}$	$d_{\rm ref}$	hkl	$d_{ m obs}$	d_{ref}
$1\overline{1}0$	9.43	9.436	$2\overline{3}1$	3.54	3.528
010	8.15	8.150	$1\overline{3}1$	3.50	3.496
100	8.14	8.143	111	3.41	3.404
$01\overline{1}$	7.51	7.512	$21\overline{1}$	3.22	3.229
$1\overline{1}1$	6.47	6.473	$21\overline{2}$	3.04	3.034
$2\overline{1}0$	5.17	5.159	$03\overline{1}$	2.97*	2.968
$1\overline{2}1$	5.14	5.144	$2\overline{4}1$	2.72*	2.716
$11\overline{1}$	4.90	4.895	$4\overline{2}0$	2.58*	2.580
110	4.51	4.516	211	2.50	2.497
$02\overline{1}$	4.45	4.448	$22\overline{1}$	2.44*	2.450
$2\overline{2}1$	4.25	4.256	$\overline{2}51$	2.13*	2.128
201	4.25	4.250	$\overline{4}1\overline{1}$	2.12*	2.116
$2\overline{1}1$	4.08	4.076			

*Strong reflections. Note that three strong reflections, 2.71 $(20\overline{3})$, 2.58 $(4\overline{2}0)$ and 2.57 $(21\overline{3})$, were ascertained from diffraction patterns of section FIB-1. Reflections 2.71 $(20\overline{3})$ and 2.57 $(21\overline{3})$ were not recorded in diffraction patterns of section FIB-2 and hence not listed for unit-cell refinement.

(Grew *et al.*, 2005), which has a composition very similar to kuratite except for the presence of Be (Table 3). Unit-cell data thus derived for kuratite show it is triclinic and in space group $P\overline{1}$ (by analogy to rhönite), a = 10.513(7), b = 10.887(7), c = 9.004(18) Å, $\alpha = 105.97(13)$, $\beta = 96.00(12)$, $\gamma = 124.82(04)^{\circ}$, $V = 767 \pm 2$ Å³, Z = 1 for the formula with 40 oxygens. The *a:b:c* ratio calculated from the unit-cell parameters is 0.966:1:0.827.

Powder X-ray diffraction data are not available for kuratite. Measured and refined d-spacings from TEM SAED patterns of section FIB-2 are listed in Table 2. The eight strongest reflections in electron diffraction, as ascertained from high-order zone-axis electron diffraction patterns of sections FIB-1 and FIB-2, are (in the order of decreasing d spacing, Å) 2.97 $(03\overline{1})$, 2.72 $(\overline{241})$, 2.71 $(20\overline{3})$, 2.58 $(4\overline{20})$, 2.57 $(21\overline{3})$, 2.45 $(22\overline{1})$, 2.13 $(\overline{251})$ and 2.12 $(\overline{411})$.

Raman spectroscopy

The Raman spectrum of kuratite, measured using a LABRAM HR confocal micro-Raman spectrometer equipped with an Ar+ laser with 514.5 nm excitation at Academia Sinica, Taiwan, shows four main scattering peaks near 563–571, 697–699 (strongest), 852–856 and 986–996 cm⁻¹ (Fig. 4)

resembling that of lunar rhönite (Treiman, 2008) but with higher frequencies due to different chemical composition. A weak shoulder on the strongest peak at 720 cm⁻¹ was recognized in lunar rhönite (Treiman, 2008) but was not present in kuratite.

Physical and optical properties

Calculated density is 3.906 g cm⁻³ using the empirical formula. Other physical properties and optical properties are not available because of the minute size of the kuratite crystals in the sample. The optical data available from the literature for Fe²⁺-dominant rhönite, i.e. kuratite, are $\alpha=1.805\pm0.007,~\beta=1.815\pm0.007,~\gamma=1.845\pm0.007,~2V=50\pm3^{\circ}$ for Puy de St.-Sandoux rhönite (X $_{\rm Mg}=0.49$); $Z\wedge c=38$ –43° for Kaiserstuhl rhönite (X $_{\rm Mg}=0.41$); $Z\wedge c=58^{\circ}$ (X $_{\rm Mg}=0.02$) for Luna 24 rhönite (Grapes and Keller, 2010; Grünhagen and Seck, 1972; Olsson, 1983; Treiman, 2008).

Discussion

Relation to other species

Rhönite is the only sapphirine-group mineral reported to have compositions with both Mg²⁺/ $(Mg^{2+} + Fe^{2+}) < 0.5$ and $Mg^{2+}/(Mg^{2+} + Fe^{2+}) > 0.5$. As plotted in figure 4 of Grapes and Keller (2010), there is a continuous range of Fe2+ and Mg substitution with $Mg^{2+}/(Mg^{2+} + Fe^{2+}) = 0.16-0.49$ for several terrestrial Fe2+-rich rhönites; e.g. 'rhönite' in tephrite glass and limburgite of basaltic rocks from the Kaiserstuhl volcanic complex (Grapes and Keller, 2010), 'rhönite' in the unusual fassaite-melilite-rhönite paragenesis from an alkaline basalt contaminated by Ca from coral, Saint-Leu, Réunion Island (Havette et al., 1982), 'rhönite' in the Mesozoic basanites from Skane (Scania) (Olsson, 1983), rhönite from a melaphonolite of Puy de Saint-Sandoux (Auvergne) (Grünhagen and Seck, 1972) and 'rhönite' in glass from kaersutite-bearing spinel-wehrlite xenoliths from Foster Crater (Gamble and Kyle, 1987). According to the 50% rule (Nickel and Grice, 1998), these so-called 'rhönite' phases could well be categorized as kuratite with the characteristic end-member composition Ca₄(Fe₁₀²⁺Ti₂) O₄[Si₈Al₄O₃₆]. Note that kuratite was first anticipated in the paper reporting the nomenclature of the sapphirine supergroup (Grew et al., 2008), in which the rhönites from Réunion, Skane, Puy de Saint-

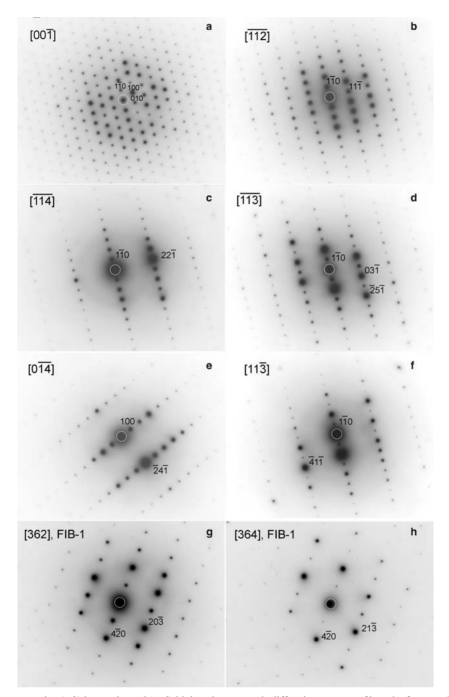


Fig. 3. Representative (a,b) low-order and (c-f) high-order zone-axis diffraction patterns of kuratite from section FIB-2 and (g,h) high-order zone-axis diffraction patterns from section FIB-1. The transmitted beam is circled in each diffraction pattern.

TABLE 3. Comparison of unit-cell parameters between kuratite and other rhönite subgroup minerals.

	a (Å)	b (Å)	c (Å)	α (°)	β (°)	γ (°)	sample type
Kuratite	10.5	10.9	9.0	106.0	0.96	124.8	D'Orbigny angrite
$(Ca_{3.88}Na_{0.02}REE_{0.13}^{3+}Nn_{0.03}Ma_{0.01}Ni_{0.02}Zn_{0.01}Sr_{0.01})_{\Sigma 4.01}(Fe_{9.98}^{2+} Ti_{2.00})_{\Sigma 11.98}(Si_{7.80}Al_{3.52}Fe_{9.64}^{3+}P_{0.05}S_{0.02})_{\Sigma 12.03}O_{39.98}F_{0.01}Cl_{0.$	$Mg_{0.01}Ni_{0.02}Zn_{0.01}Sr$	$(F_{0.01}^{2+})_{\Sigma 4.01} (F_{9.98}^{2+} T_{12.0}^{2})$	$_{10})_{\Sigma 11.98}(\mathrm{Si}_{7.80}\mathrm{Al}_{3.52}\mathrm{I}$	$Fe_{0.64}^{3+}P_{0.05}S_{0.02})_{\Sigma12.0}$	$_{3}O_{39.98}F_{0.01}Cl_{0.01}$		
Rhönite ¹ 27 8 47 40	10.4	10.8	8.9	105.9	96.1	124.8	volcanic breccias
$Ca_{3.8}Na_{0.2}(Mg_{6.6}Fe_{1.2}^{2+}Fe_{2.2}^{3+}Ti_2)(Si_6AI_6)O_{40}\\ Ca_4(Mg_8Fe_2^{3+}Ti_2)(Si_6AI_6)O_{40}$	$^{12}_{2})(Si_{6}Al_{6})O_{40}$						
Rhönite	10.4	10.8	6.8	106.0	0.96	124.7	Allende meteorite
$\begin{array}{l} Ca_4(Mg_{5.8}Fe_{0.6}^{2+}Ti_{1.6}^{3+}AI_{1.0}V_{0.6}Ti_{2.4})(Si_4AI_8)O_{40} \\ Ca_4(Mg_7Ti_3^{3+}Ti_3)(Si_4AI_8)O_{40} \end{array}$	$_{6}$ Ti _{2.4})(Si ₄ Al ₈)O ₄₀						
Makarochkinite ²	10.4	10.8	6.8	105.7	96.2	124.9	granitic pegmatite
$ \begin{array}{l} (Ca_{3.28}Na_{0.50}Mn_{0.22})(Fe_{7.12}^{2+}Fe_{2.92}^{3+}Ti_{1.2} \\ Ca_4(Fe_8^2+Fe_2^2+Ti_2)(Si_8Be_2Al_2)O_{40} \end{array} $	${ m Fe}_{2.92}^{3+}{ m Ti}_{1.22}{ m Mg}_{0.50}{ m Mm}_{2){ m O}_{40}}$	10.04Nb _{0.076} Ta _{0.014})(\$	$_{.22}Mg_{0.50}Mn_{0.04}Nb_{0.076}Ta_{0.014})(Si_{8.96}Be_{1.82}Al_{1.08}Fe_{0.14}^{3+})O_{40}$	$^{3+}_{0.14})O_{40}$			

Sandoux and Foster Crater were recognized as a distinct species, the Fe²⁺-analogue of rhönite [see table 1 and figures 9 to 11 in Grew *et al.* (2008)].

As for the extraterrestrial occurrences, the first description of kuratite from a meteorite was made by Mittlefehldt et al. (2002) reporting the discovery of an unknown Fe-Al-Ti-silicate with a cation/Si ratio of ~2.65. Kurat et al. (2004) found the unknown Fe-Al-Ti silicate in multi-phase inclusions in anorthite, forming euhedral crystals up to 20 um in diameter. This then unknown phase was also reported to be enclosed by augite and associated with magnetite, troilite and kirschsteinite. Its major chemical composition is homogeneous (Mittlefehldt et al., 2002; Kurat et al., 2004) and is characterized by high and unfractionated REE (~100 × CI) contents (Varela et al., 2005). More recently, kuratite with Mg²⁺/(Mg²⁺+ Fe^{2+}) = 0-0.17 was also found in the plutonic igneous angrite NWA 4590 (Kuehner and Irving, 2007), volcanic angrite SAHARA 99555 (Jambon and Boudouma, 2011), as well as magmatic inclusions in augite grains of the Luna 24 regolith (Treiman, 2008).

The ordering scheme in kuratite

For the rhönite subgroup minerals, it is well documented that in half of the unit cell, there are six different tetrahedral sites T1-T6, seven octahedral sites M1-M7, and two additional eightfold sites M8 and M9 (Bonaccorsi *et al.*, 1990; Grew *et al.*, 2008). The structure of rhönite minerals can be described as octahedral strips (M3-M7 octahedral sites and M8 and M9 eightfold sites) connected via tetrahedral chains with chain tetrahedra (T1-T4) and wings (T5 and T6 sites) and additional insular octahedra (M1 and M2). The octahedral strips are three to four octahedra wide

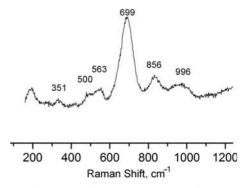


Fig. 4. Representative Raman spectrum of kuratite.

Bonaccorsi et al. (1990), ²Grew et al. (2005)

and run parallel to the crystallographic a axis. The tetrahedral chains are pyroxene-like and aligned also along the a axis. The M8 and M9 sites are occupied by Ca. The Fe^{2+} and Mg^{2+} ions prefer M1to M6 sites. Smaller Ti⁴⁺ ions are concentrated at the M7 site. As for the T sites, Si and Al probably occupy T1-T4 sites in equal proportion, and T5 and T6 are dominantly Si (Grew et al., 2008, and references therein). Although it is not possible to determine the structure of kuratite directly due to its minute crystal size, the close similarity in crystal chemistry, unit-cell type and parameters between kuratite and rhönite subgroup minerals, e.g. rhönite and makarochkinite (Table 3), do indicate that kuratite should be similar to rhönite and makarochkinite in every aspect; Ca at M8 and M9, Fe^{2+} at M1-M6, Ti at M7, Si and Al at T1-T6. By comparison with the site occupancies in makar- $(Ca_4(Fe_8^{2+}Fe_2^{3+}Ti_2)(Si_8Be_2Al_2)O_{40})$ ochkinite with similar Fe/Ti ratio at M sites and the Si/(Be + Al) ratio at T related to kuratite, Ti⁴⁺ is probably ordered at the M7 site in kuratite, whereas Fe^{2+} are disordered over the M1-M6 sites (Grew et al., 2005).

Origin of kuratite in the angrite meteorite

The formation of the angrite meteorite is still under debate, although most studies favour a basaltic igneous origin (Keil, 2012). Following the igneous scenario, it is not surprising that kuratite was found in angrite D'Orbigny, plutonic igneous angrite NWA 4590 (Kuehner and Irving, 2007), volcanic angrite SAHARA 99555 (Jambon and Boudouma 2011) and Luna 24 regolith (Treiman 2008), similar to the common occurrences of other Fe²⁺-dominant rhönites in terrestrial undersaturated basaltic rocks such as tephrite and limburgite from volcanic complexes (Grapes and Keller, 2010), basanite glass in spinel-wehrlite xenoliths (Gamble and Kyle, 1987), alkaline basalt within coral (Havette et al., 1982) and basanite (Olsson, 1983). In fact, texture observations indicate that terrestrial kuratite in basaltic rocks was the last phase crystallized. The frequent presence of glass groundmass further indicates that kuratite was most likely a quench phase either on eruption (e.g. undersaturated alkaline basaltic rocks; Grapes and Keller, 2010) or from rapid cooling of an infiltrated melt into the host rocks (e.g. kaersutite-bearing wehrlite xenoliths; Gamble and Kyle, 1987). In accordance with the above, the occurrence of kuratite + whitlockite embedded in Ca-rich fayalite + kirschsteinite intergrowths (Fig. 1d,e) at the contact between Fe-sulfide spherules and hedenbergite suggests that these phases could have crystallized from an interstitial melt at a very late stage of rock formation. Although it is not clear regarding the origin of this interstitial melt phase, i.e. residue from fractional crystallization or a fresh, infiltrating melt (Mittlefehldt et al., 2002; Kurat et al., 2004; Varela et al., 2003), this interstitial melt must be subsilicic, almost Mg-free but enriched in Al-P-Ca-Ti-Fe. Alternatively, the multiple phase assemblage at olivine-augite triple junctions might represent metasomatic products resulting from reactions between an intruding metasomatic agent, which could be either gas, liquid or melt, and the porous and fractured olivine + augite + Fe-sulfide aggregates (see Fig. 1a). In either case, the crystallization temperatures of kuratite must be >1000°C, considering the co-presence of Ca-rich favalite and kirschsteinite (Mukhopadhyay and Lindsley, 1983), the 950-1180°C stability field of rhönites based on experimental phase relations for various undersaturated alkaline basaltic rocks (Kunzmann, 1999), and the environment for the formation of Fe²⁺-dominant rhönite as a quenched phase in glass with liquidus temperatures of 1038-1137°C in tephrite and limburgite bulk compositions (Grapes and Keller, 2010).

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