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Revision 2

**Crystal structure of richetite revisited: crystallographic evidence for the presence of
pentavalent uranium**

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10 **ABSTRACT:**

11 Revision of crystal structure of the rare U-oxide mineral richetite provided crystallographic
12 evidence for the presence of pentavalent U. The structure of richetite, space group $P\bar{1}$, $a =$
13 $12.0919(2)$, $b = 16.3364(4)$, $c = 20.2881(4)$ Å, $\alpha = 68.800(2)^\circ$, $\beta = 78.679(2)^\circ$, $\gamma = 76.118(2)^\circ$,
14 with $V = 3600.65(14)$ Å³ and $Z = 1$, was solved by charge-flipping algorithm and refined to an
15 agreement index (R) of 5.6% for 9955 unique reflections collected using microfocus X-ray
16 source. The refined structure, in line with the previous structure determination, contains U-O-
17 OH sheets of the α -U₃O₈ type (protasite topology) and an interstitial complex comprising
18 Pb²⁺, Fe²⁺, Mg²⁺ cations and molecular H₂O. However, the polyhedral geometry, the bond-
19 valence sum incident at one U site within the sheet (U17) together with charge-balance
20 requirements, indicate that U17 site is occupied by U⁵⁺. The U17Φ₇ (Φ: O, OH) polyhedra is
21 rather distorted, with two shorter U–O bond-lengths (~2.01 Å), four longer U–O bond-lengths
22 (~2.2 Å) and one, very long U–O bond (2.9 Å). The color of richetite also supports the
23 presence of U⁵⁺ in the structure. The current results show that α -U₃O₈ type of sheet can
24 incorporate U⁵⁺. Richetite is the second mineral containing pentavalent uranium in Nature.

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26 **Keywords:** Richetite, uranyl oxide hydroxy-hydrate, crystal structure, pentavalent uranium,
27 weathering

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29 Running title: Pentavalent U in richetite

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INTRODUCTION

32 Uranyl-oxide hydroxy-hydrate minerals (further labelled as UOH) are important
33 products of supergene weathering of primary U^{4+} minerals, predominantly uraninite. They
34 form in the initial alteration stages and are common constituents of the oxidized parts of
35 uranium deposits, and usually replace uraninite *in-situ*, forming massive aggregates called
36 “*gummites*” (Finch and Ewing 1992; Finch and Murakami 1999; Krivovichev and Plášil
37 2013; Plášil 2014). Weathering of uraninite, also called hydration-oxidation weathering, is of
38 further relevance because of the analogy between the alteration of uraninite and UO_{2+x} in
39 spent nuclear fuel (Janeczek et al. 1996). The crystallography and crystal chemistry of this
40 mineral group has attracted a lot of attention and this group is nowadays extensive (see e.g.,
41 Plášil et al. 2016). These minerals and the synthetic UOH phases have been studied
42 intensively by X-ray diffraction and only a few of their structures are unknown.

43 Richetite is a rare UOH mineral, occurring at few localities in the world. It was
44 originally described by Vaes (1947) from Shinkolobwe mine, Haut-Katanga province,
45 Democratic Republic of Congo, Africa, and later studied by Piret and Deliens (1984).
46 Richetite has a large triclinic unit-cell (Burns 1998), which is in line with results of Piret and
47 Deliens (1984). However, the structure has several issues prompting reexamination of the
48 structure.

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REDETERMINATION OF THE CRYSTAL STRUCTURE

Single-crystal X-ray diffraction

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52 The crystal used in this study was obtained from a sample provided by Jean-Claude
53 Leydet (Brest, France) and originates from the type locality, Shinkolobwe mine (Haut-
54 Katanga province, DRC, Africa).

55 A tabular greyish brown fragment of richetite was selected under an optical
56 microscope and used for X-ray study. Data were collected using a Rigaku (Oxford diffraction)
57 SuperNova diffractometer, using MoK α radiation ($\lambda = 0.71073 \text{ \AA}$) from a micro-focus X-ray
58 tube collimated and monochromatized by mirror optics and detected by an Atlas S2 CCD
59 detector. From 39599 collected reflections, 13469 were independent and 9955 were unique
60 observed with the criterion $I_{\text{obs}} > 3\sigma(I)$. Integration of the diffraction data, including
61 corrections for background, polarization and Lorentz effects, was done using the CrysAlis
62 RED program. The absorption correction combining empirical scaling and spherical-
63 absorption correction was done with CrysAlis program; SCALE3 Abspack algorithm.

64 The structure of richetite was solved by the charge-flipping algorithm using the Shelxt
65 program (Sheldrick 2015). The structure model was refined by full-matrix least-squares in the
66 Jana2006 program (Petříček et al. 2014) based on F^2 . The reflection conditions were
67 consistent with the space-group $P-1$, which was further confirmed by the successful
68 refinement. The possibility for twinning by reticular merohedry was tested by Jana2006
69 (Petříček et al. 2016) (transformation matrix 1 -2 -1/1 0 0/0 -1 1), however it was negative.
70 The crystal used for the experiment was found to be a split crystal; the contribution of the
71 second fragment to the dataset was corrected by detecting fully separated, fully overlapped
72 and partially separated reflections in Jana2006 (Petříček et al. 2016). The structure solution
73 provided nearly complete structure sheets and missing atoms (mostly O atoms) were located
74 from the difference-Fourier maps. Anisotropic displacement parameters were used for U, Pb
75 and Fe atoms. Unconstrained and unrestrained refinement converged smoothly to final $R =$
76 0.056 for 9955 unique observed reflections (Table 1). Final atom coordinates and
77 displacement parameters are listed in Tables 2 and 1S (Supplementary file), selected
78 interatomic distances are in Tables 3 and 4, and the bond-valence sums (calculated by the
79 procedure of Brown, 1981, 2002) are listed in Table 2. The original crystallographic

80 information file (cif) is provided as Supplementary material and can be downloaded from
81 XXXX.

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DESCRIPTION OF THE CRYSTAL STRUCTURE

84 The structure of richetite contains 18 unique U sites, 8 unique Pb sites, 1 mixed Fe/Mg site,
85 and 85 O sites (of which 18 correspond to H₂O groups) (Fig. 1). The U sites are coordinated
86 by seven ligands (O or OH⁻) in two classes of distances: ~1.8 Å and 2.1 to ~2.8 Å. as it is
87 characteristic for the UO₂²⁺ ion (Evans 1963; Burns et al. 1997a; Lussier et al. 2016). The
88 structure contains 8 Pb sites; none of them is fully occupied, site-scattering refinement
89 showed occupancies ranging from 0.14 to 0.95. The coordination polyhedra around Pb atoms
90 are irregular; ligands are represented by O_{Ur} atoms and H₂O groups, with Pb-Φ bond-lengths
91 ranging from 2.4 to 3.1 Å (Table 5). Richetite structure contains also one symmetrically
92 unique octahedrally coordinated site, occupied by divalent cations Fe²⁺ and Mg²⁺, with <M²⁺-
93 Φ> bond-length of 2.07 Å. The M²⁺ octahedron is quite regular and is formed by two O_{Ur}
94 atoms (of the U2) and four H₂O groups. Site-scattering refinement gave M²⁺ = 0.62 Fe²⁺ +
95 0.38 Mg²⁺.

96

The U17 site

97
98 Several U sites in the structure of richetite exhibit rather irregular coordination. In case of
99 U17 (Fig. 2; Table 4) O62 and O72 atoms (OH groups), which usually should be O_{Ur} atoms
100 with U-O bond-lengths of 1.8 Å, have U-O distances of 2.006(19) and 2.01(2) Å. Moreover,
101 O62-U17-O72 bond-angle is 171.82°, different from the usually linear UO₂²⁺ ion. Bond-
102 valence analysis (Table 2) indicates that the U17 site is occupied by pentavalent U.

103

The sheets of polyhedra

105 The sheet of polyhedra found in richetite has the protasite uranyl–anion topology, i.e.
106 the α - U_3O_8 sheet (Fig. 3) (Burns 1998, 2005; Lussier et al., 2016). All pentagons are occupied
107 by U atoms, the U17 site by U^{5+} , and the rest by U^{6+} . Based upon distribution of $(\text{OH})^-$ within
108 the equatorial ligands (excluding two OH groups associated with U17), we can distinguish
109 several structures of protasite topology. In richetite (Burns 1998), we have the AABAAB...
110 sequence, where considering type-A triangles (which have (OH) groups at all corners), and
111 type-B triangles (which contain only O^{2-} anions), richetite has twice as many A triangles as B
112 triangles. The O:OH ratio in richetite is 3:2, however, there are also two OH groups linked to
113 U17 (in case of other U sites in richetite, they are O_{Ur} atoms), linking U17 to Pb1 through
114 O62, and to Pb4 and Pb7 through O72. In case of protasite (Pagoaga et al. 1987), all (OH)
115 groups are located at the corners of triangles of the topology, such that all triangles in the
116 sheet contain two (OH) groups. The sheets in the structures of becquerelite (Burns and Li
117 2002) and billietite (Finch et al. 2006) (considering there the α - U_3O_8 sheet only) have the
118 composition $[(\text{UO}_2)_6\text{O}_4(\text{OH})_6]^{2-}$ with O:OH = 2:3. In the becquerelite and billietite sheet
119 anion-topologies, all (OH) groups are located at the corners of triangles, and all triangles
120 contain three (OH) groups.

121

122 *The interlayer complex*

123 As indicated by Burns (1998), the structure of richetite contains two different
124 interlayer complexes containing Pb^{2+} , M^{2+} and H_2O groups. Adjacent sheets of protasite
125 topology are linked through an extensive network of $\text{Pb}-\text{O}$, $M^{2+}-\text{O}$ and $\text{O}-\text{H}\dots\text{O}$ bonds. The
126 interstitial complex comprising M^{2+} octahedra occurs at $\mathbf{b} = 0$, $\mathbf{c} = 0$ and is built from two
127 tetramers (Pb2, Pb4, Pb7 and Pb8) linked by M^{2+} octahedra (Fig. 4). Four of the ligands
128 coordinated to M^{2+} site are H_2O groups, and two are O_{Ur} atoms. There is an additional O site

129 (O84) that is occupied by an H₂O group that links to the structure through H-bonds only. The
130 coordination environment of the corresponding Pb sites is shown in Fig. 4.

131 The interlayer complex that does not contain the M^{2+} site occurs at $b \sim 0.5$, $c \sim 0.5$. It
132 contains a dimer of Pb1 Φ_8 and Pb3 Φ_9 ($\Phi = O, OH, H_2O$) polyhedra and a tetramer of Pb5 Φ_8
133 and Pb6 Φ_8 ($\Phi = O, H_2O$) polyhedra (Fig. 5). Between these clusters of polyhedra there are
134 three independent O sites that belong to H₂O groups that are not coordinated directly to any
135 metal cation site.

136

137 *The structural formula*

138 The structural formula of the studied richetite crystal is therefore
139 $M^{2+}_{0.50}Pb_{4.86}[U^{5+}(U^{6+}O_2)_{17}O_{18}(OH)_{14}](H_2O)_{-19.5}$, $Z = 2$. This formula is in line with the color
140 of richetite (Fig. 6). Most UOH minerals are orange or yellow, whereas those minerals
141 containing U^{5+} , wyartite (Burns and Finch 1999) and dehydrated wyartite (Hawthorne et al.
142 2006) are similar in color to richetite.

143

144 *α -U₃O₈ topology and the presence of U^{5+}*

145 Incorporation of U^{5+} into α -U₃O₈ type sheets has not been considered; all minerals
146 with U^{5+} or U^{4+} present adopt β -U₃O₈ sheets (Burns and Finch 1999; Burns et al. 1997b;
147 Hawthorne et al. 2006). For shinkolobweite, $Pb_{1.25}[U^{5+}(H_2O)_2(UO_2)_5O_8(OH)_2](H_2O)_5$ (Olds et
148 al. 2017), there are sheets resembling β -U₃O₈ topology. Geometrically, the U17 site in
149 richetite is consonant with the idealized α -U₃O₈ topology and bond-valence analysis clearly
150 indicates incorporation of U^{5+} . Thus, incorporation of U^{5+} into α -U₃O₈ topology is possible.

151

152 **THE ROLE OF RICHTITE IN URANINITE ALTERATION**

153 UOH minerals are important products of oxidation-hydration weathering of uraninite,
154 or UO_2 in spent nuclear fuel (Finch and Ewing 1992; Janeczek et al. 1996; Wronkiewicz et al.
155 1992, 1996; Krivovichev and Plášil 2013; Plášil 2014). Based on field as well as laboratory
156 observations (for references, see above cited papers), a weathering sequence for UOH has
157 been established (after Finch and Ewing 1992; Fig. 7). At very early stages of uraninite
158 alteration under oxidizing conditions, result in minerals with a high-proportion of molecular
159 H_2O and low content of metal cations such as schoepite, $[(\text{UO}_2)_8\text{O}_2(\text{OH})_{12}](\text{H}_2\text{O})_{12}$ (Finch et
160 al. 1996, 1998). With increasing time UOH structures will incorporate metal cations released
161 from the gradually weathering uraninite (such as radiogenic Pb, and others) or from host-
162 rocks (Na, K, Ca etc.). The position of richetite in the alteration sequence is determined by the
163 molar proportion of H_2O and *Me* close to fourmarierite, masuyite and protasite. Also the
164 value of charge deficiency per anion (CDA; defined by Schindler and Hawthorne 2008),
165 ~ 0.21 v.u., is close to that of fourmarierite (0.19 v.u.) and masuyite (0.22 v.u.). It has been
166 shown that the CDA value correlates closely with increase of *Me* and decrease of H_2O in the
167 structures, therefore higher CDA corresponds to older products during the weathering
168 sequence. There are two or three other UOH minerals that contain reduced forms of U. They
169 are ianthinite (with U^{4+}), wyartite and dehydrated wyartite (with U^{5+} in addition to U^{6+}). The
170 presence of the reduced form of an easily oxidized species indicates high gradients of redox
171 conditions within the systems where these phases form. All of these phases form during initial
172 stages of uraninite weathering. Ianthinite is related to the schoepite family of minerals (Fig.
173 7); there is a solid-state spontaneous phase transition from ianthinite to schoepite. The
174 position of wyartite is not clear; the usual mineral association comprises uranophane,
175 schoepite and fourmarierite. Samples of ianthinite, besides those from Shinkolobwe, also
176 come from the well-known uranium deposit Menzenschwand (Krunkebachtal, Baden-
177 Württemberg, Germany), where ianthinite is usually associated with pyrite and often fills

178 vugs in altered uraninite-pyrite aggregates in a quartz matrix. Richetite forms during later
179 stages of uraninite weathering. The presence of Fe^{2+} in richetite also indicates special
180 geochemical conditions. It seems likely that most of sulfides (as a source of Fe) had
181 undergone complete dissolution prior to the formation of richetite. It is clear that partial
182 reduction of U^{6+} to U^{5+} is most probably connected to the $\text{Fe}^{2+}/\text{Fe}^{3+}$ pair, which is also the
183 most frequent redox agent in Nature. To assess the role of Fe in the formation of richetite,
184 more detailed textural work is needed. To conclude, minerals where U is present in a reduced
185 valence state, such as in ianthinite, wyartite and richetite, may play an important role during
186 the alteration of uraninite or SNF under less-oxidizing conditions, or at places with reduced
187 $f\text{O}_2$ or where the redox conditions are characterized by high gradients, as in roll-front,
188 environments with extremely low pH etc. Such phases also might play role during long-term
189 storage of SNF in geological repositories under reducing conditions (Ewing 2015; Ewing et
190 al. 2016).

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321

322 Caption to Figures

323 FIGURE 1. Crystal structure of richetite viewed down **a**. Uranyl polyhedra are drawn in yellow
324 color, except of $U^{5+}17$ polyhedra (red); M^{2+} octahedra are green, Pb atom dark grey and
325 O atoms are displayed as red balls. Unit-cell edges are outlined in solid black line.

326 FIGURE 2. Coordination environment around U17 site, occupied by U^{5+} , with displayed bond-
327 lengths and O–U–O bonding-angle.

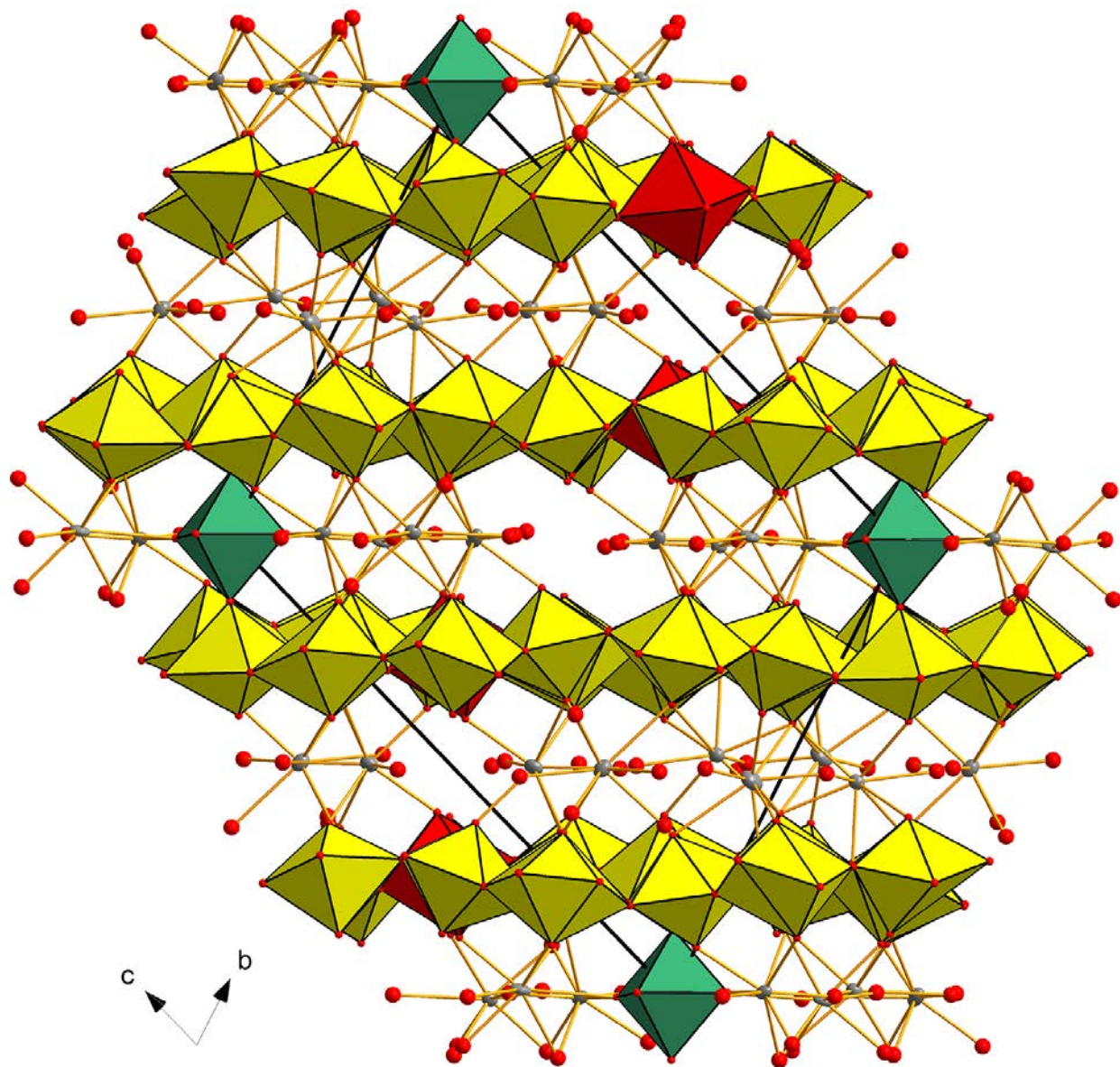
328 FIGURE 3. Sheet of uranyl polyhedra of the α - U_3O_8 type (or protasite topology) found in the
329 structure of richetite. In red is displayed U17 polyhedron, occupied by U^{5+} ; the
330 distribution of OH groups within the sheet is shown by blue balls.

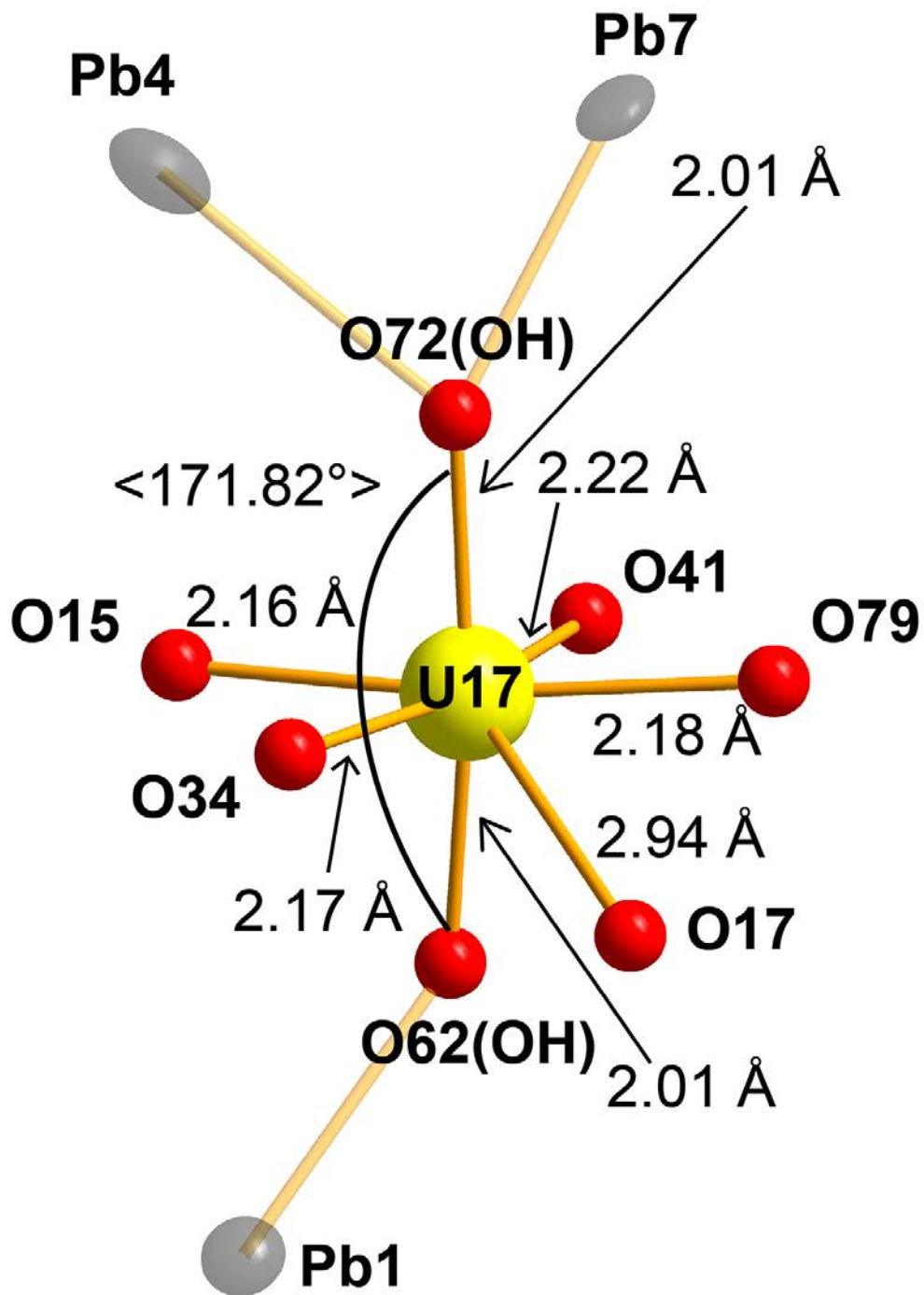
331 FIGURE 4. Interstitial constituents at **b**~0, **c**~0 (extended to for about four unit-cell content).
332 The M^{2+} (mixed Fe1/Mg1 site) octahedra are shown in green color, Pb^{2+} -sites are dark
333 grey, O atoms are represented by red balls. H_2O groups are labelled (W).

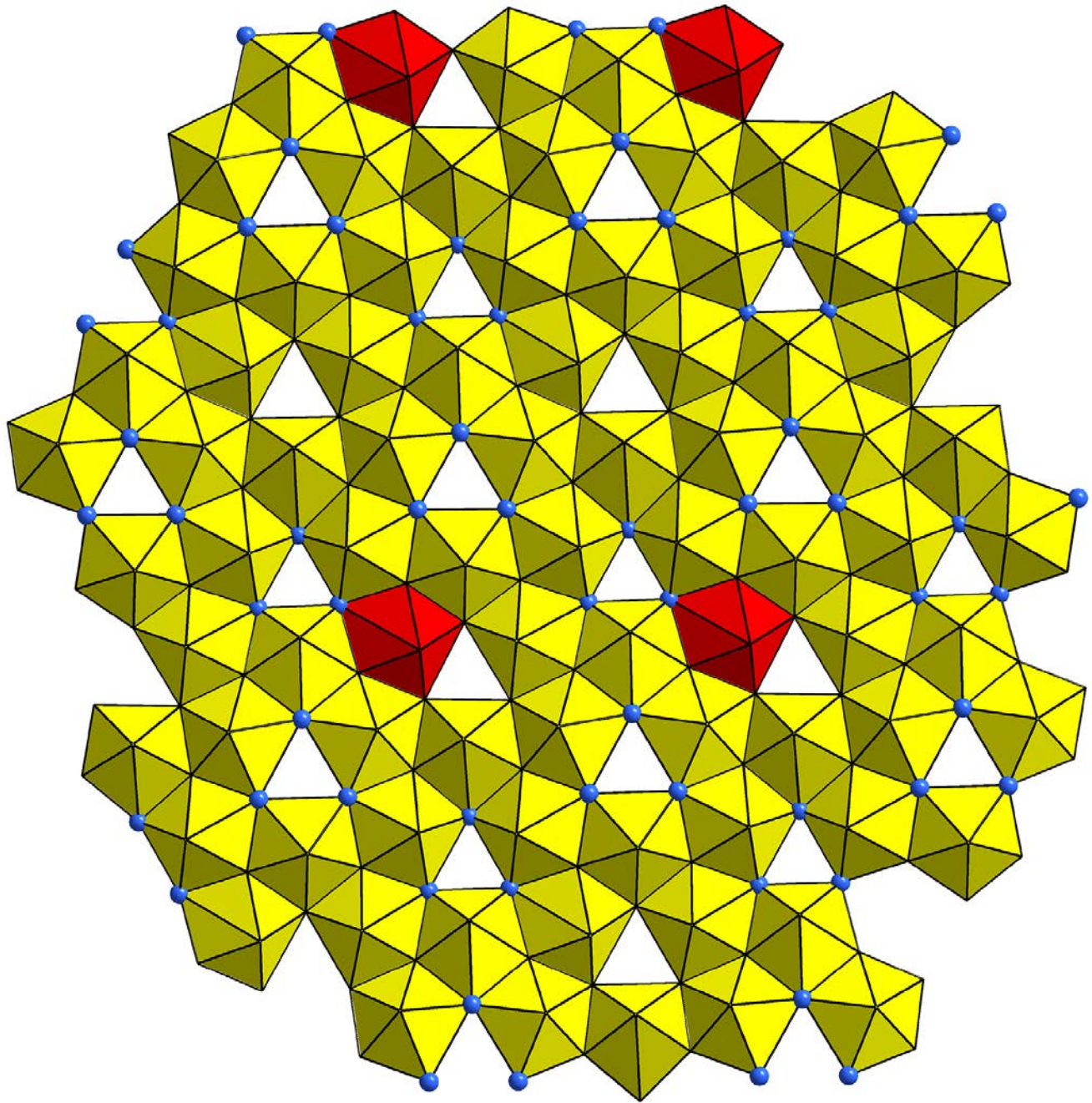
334 FIGURE 5. Interstitial constituents at **b**~0.5, **c**~0.5 (extended to for about four unit-cell
335 content). Pb^{2+} -sites are dark grey, O atoms are represented by red balls. H_2O groups are
336 labelled (W).

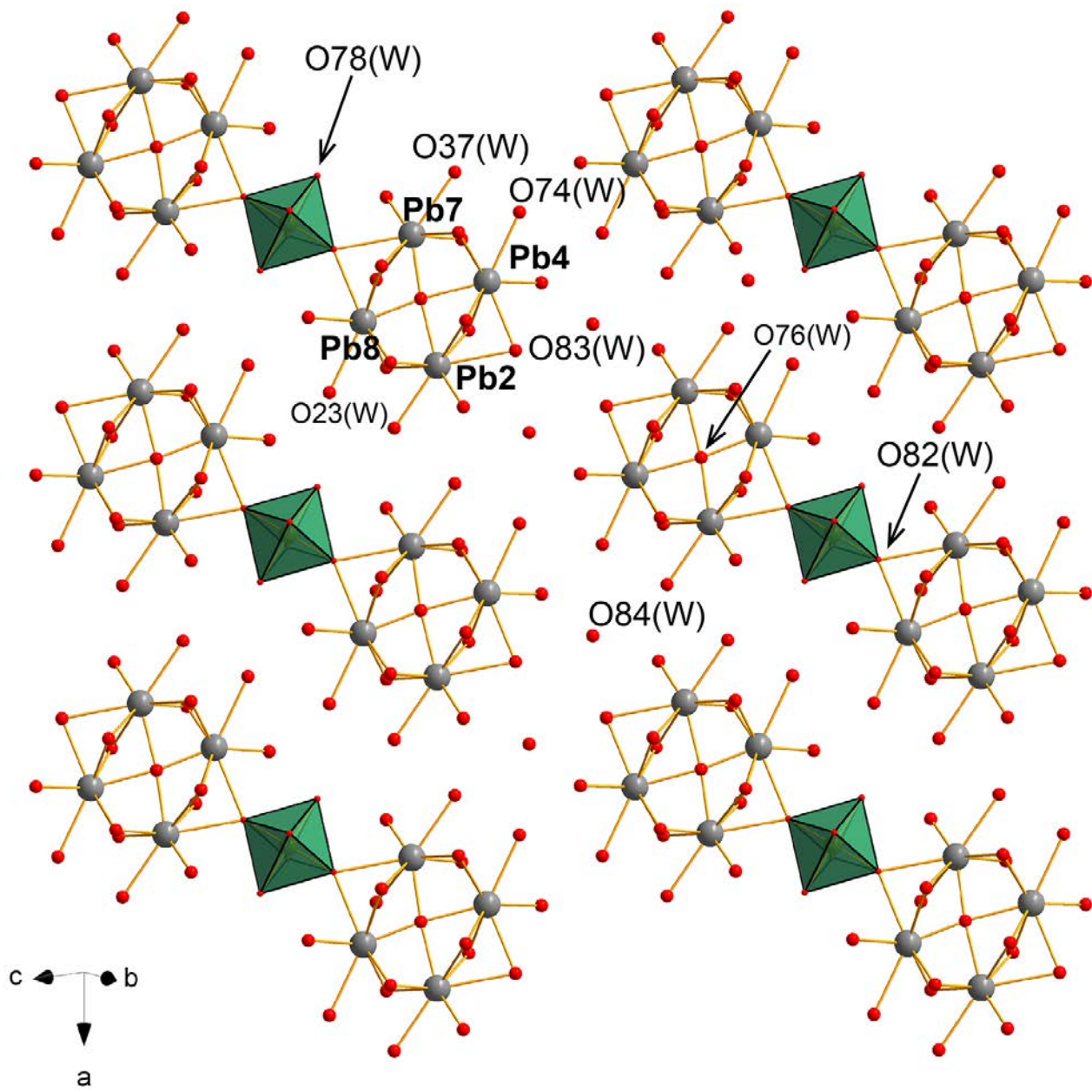
337 FIGURE 6. Richetite crystals (olive brown) among masuyite (orange). Shinkolobwe mine (type
338 locality), Haute-Katanga province, DRC, Africa. FOV 2 mm, photo S. Wolfsried.

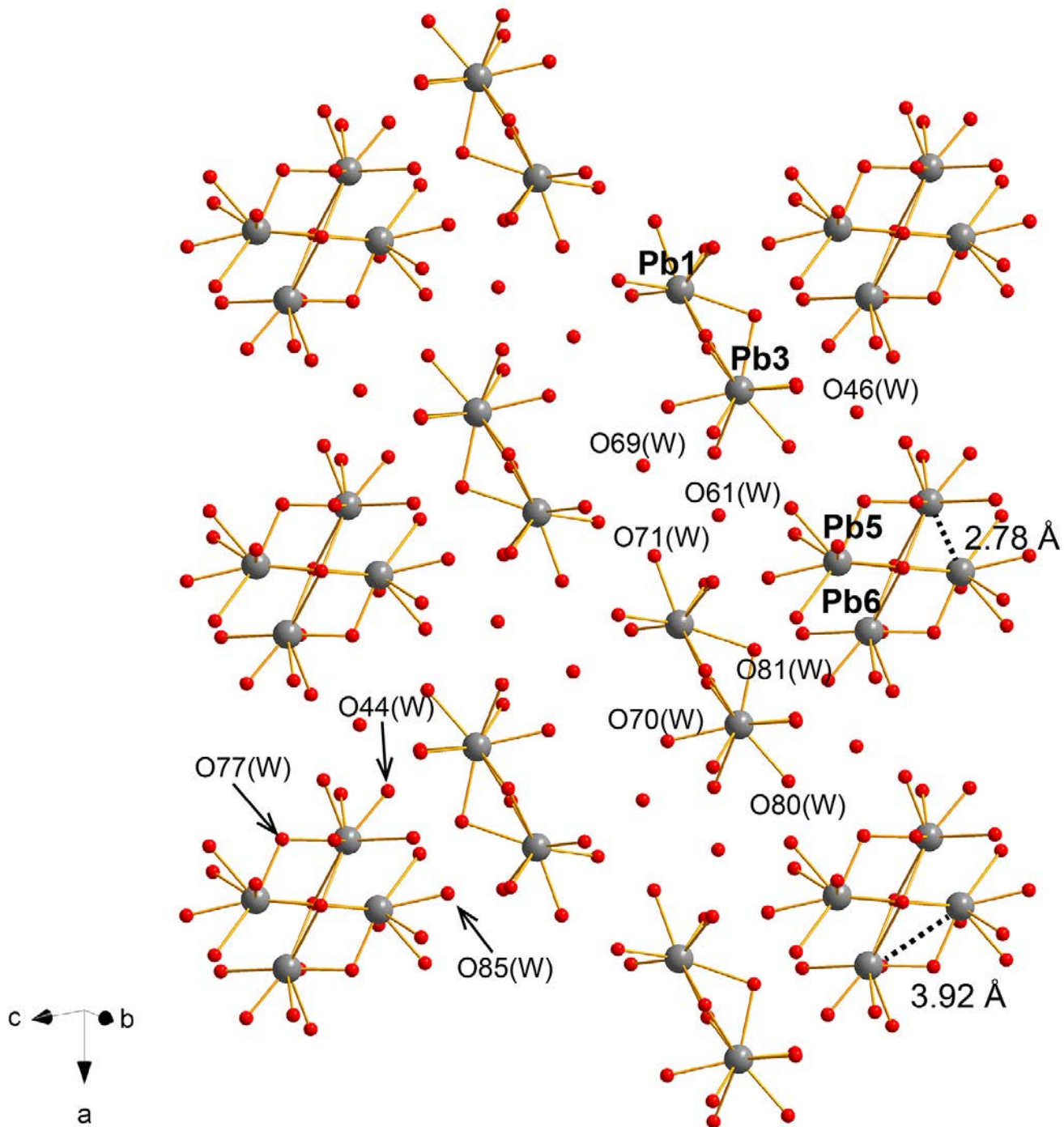
339 FIGURE 7. Composition of uranyl-oxide hydroxy-hydrate minerals as a function of proportion
340 of molecular H_2O and a content of metal cations (Me).













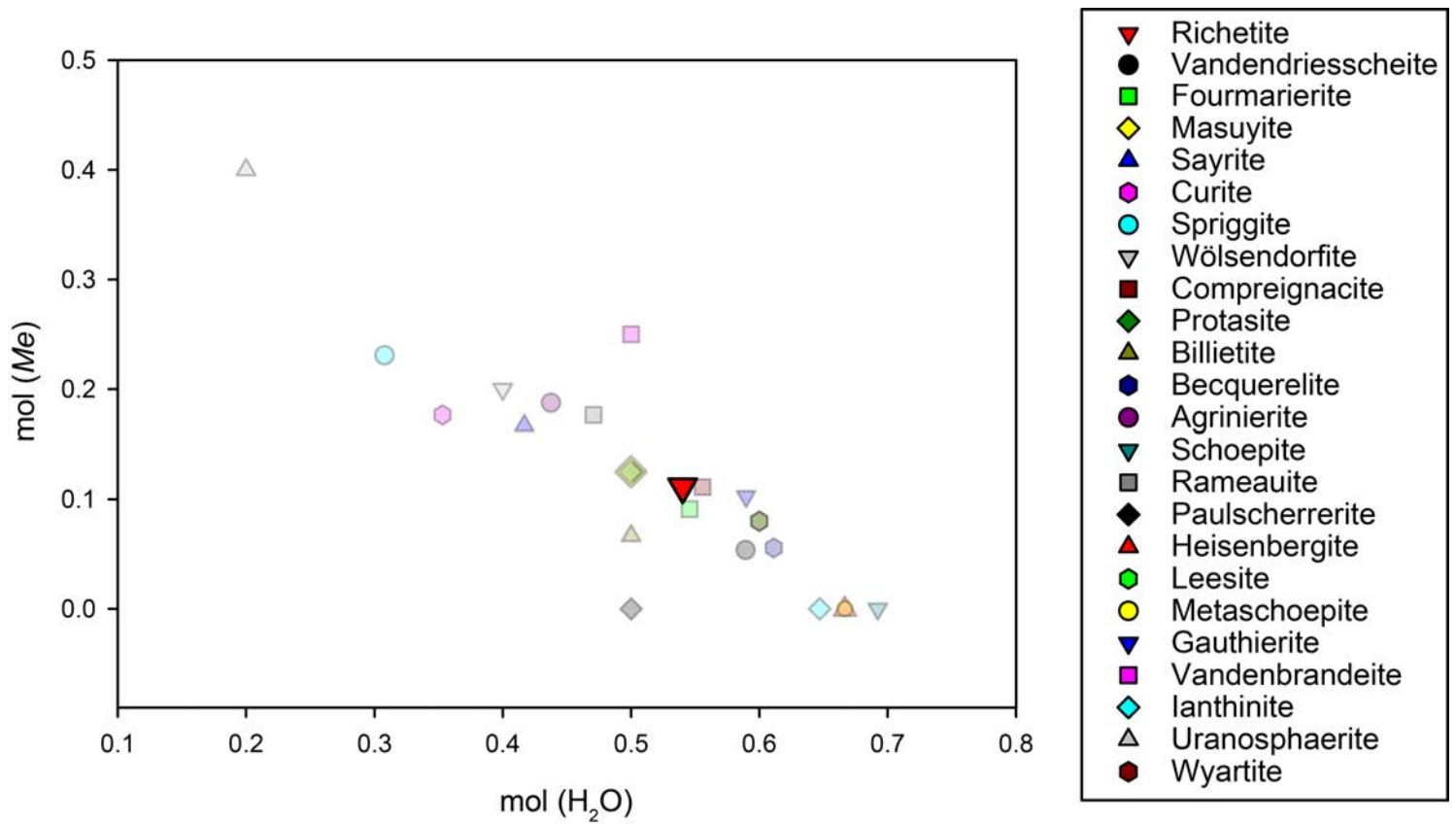


Table 1. Summary of data collection conditions and refinement parameters for richetite.

Structural formula	$(\text{Fe}^{2+}_{0.31}\text{Mg}_{0.19})\text{Pb}_{4.86}[\text{U}^{5+}(\text{U}^{6+}\text{O}_2)_{17}\text{O}_{18}(\text{OH})_{14}](\text{H}_2\text{O})_{\sim 19.5}$
Unit cell parameters	$a = 12.0919(2), b = 16.3364(4), c = 20.2881(4) \text{ \AA}$ $\alpha = 68.800(2), \beta = 78.6794(18), \gamma = 76.1181(19)$
V	$3600.68(14) \text{ \AA}^3$
Z	2
Space group	$P-1$
D_{calc} (g cm^{-3})	6.194 (for the formula given above)
Temperature	298 K
Diffractometer	Rigaku SuperNova, Atlas S2 CCD
Radiation	$\text{MoK}\alpha$ (0.7107 \AA)
(wavelength)	
Crystal dimensions	$0.174 \times 0.135 \times 0.029 \text{ mm}$
Collection mode	ω scans to cover the Ewald sphere
Frame width, counting time	$1.0^\circ, 300 \text{ s}$
Limiting θ angles	$3.40\text{--}28.10^\circ$
Limiting Miller indices	$-15 < h < 15, -21 < k < 21, -26 < l < 26$
No. of reflections	39599
No. of unique reflections	13469
No. of observed reflections (criterion)	9955 [$I_{\text{obs}} > 3\sigma(I)$]
μ (mm^{-1})	51.76
$T_{\text{min}}/T_{\text{max}}$	0.055/0.079
Coverage, R_{int}	0.98, 0.043
F_{000}	5479
Refinement	Full matrix least-squares by Jana2006 on F^2
Parameters refined	593
R, wR (obs)	0.0560, 0.1142
R, wR (all)	0.0771, 0.1210
GOF (obs, all)	1.99, 1.80
Weighting scheme	$1/(\sigma^2(I) + 0.0004I^2)$
$\Delta\sigma_{\text{min}}, \Delta\sigma_{\text{max}}$ ($\text{e}/\text{\AA}^3$)	$-4.12, 6.56$ (1.29 \AA from O70 atom)
Twin ratio; twin matrix	$0.8482(17)/0.15118(17); \begin{pmatrix} -1 & 0 & 0 \\ 0 & -0.328 & -0.671 \\ 0 & -1.328 & 0.328 \end{pmatrix}$

Table 2. Atom positions, occupation factors, displacement parameters (equivalent and isotropic, in Å²) and bond-valence sums (in valence units) for the crystal structure of richetite.

Atom	Occ. (<1)	<i>x/a</i>	<i>y/b</i>	<i>z/c</i>	<i>U</i> _{eq} / <i>U</i> _{iso} (Å ²)	ΣBV
U1		0.56884(6)	0.32974(5)	0.40750(4)	0.0139(3)	5.93(8)
U2		1.06978(6)	0.82861(5)	-0.09742(4)	0.0137(3)	5.81(8)
U3		0.26120(6)	-0.02209(6)	0.75313(4)	0.0151(3)	6.07(8)
U4		0.42599(6)	0.16277(6)	0.57945(4)	0.0153(3)	5.95(7)
U5		0.43851(6)	0.83810(6)	-0.08721(4)	0.0160(3)	5.92(7)
U6		0.75344(6)	0.48241(5)	0.25073(4)	0.0136(3)	5.97(8)
U7		0.05207(6)	0.51722(6)	0.24384(4)	0.0150(3)	5.88(7)
U8		0.93510(6)	0.34134(6)	0.41004(4)	0.0152(3)	5.89(7)
U9		0.58292(6)	0.66312(6)	0.08266(4)	0.0160(3)	5.81(8)
U10		0.93432(6)	0.65868(6)	0.07252(4)	0.0143(3)	5.93(7)
U11		0.25508(6)	0.64896(6)	0.09384(4)	0.0165(3)	5.91(8)
U12		0.43717(7)	0.51334(6)	0.24642(4)	0.0177(3)	5.86(9)
U13		0.75112(6)	0.15270(6)	0.59592(4)	0.0158(3)	6.01(8)
U14		0.74315(6)	0.82801(5)	-0.05630(4)	0.0149(3)	5.95(8)
U15		0.54990(6)	0.02117(6)	0.74772(4)	0.0150(3)	5.91(8)
U16		0.24484(6)	0.32640(5)	0.44176(4)	0.0140(3)	5.98(8)
U17		0.08254(7)	0.16447(6)	0.57735(4)	0.0186(3)	5.37(7)
U18		-0.06322(7)	0.01441(6)	0.74493(4)	0.0197(3)	5.93(9)
Pb1	0.945(3)	0.83195(8)	-0.12157(7)	0.62331(5)	0.0317(4)	1.739(18)
Pb2	0.857(3)	0.54386(8)	0.71947(7)	0.26790(5)	0.0260(4)	1.90(2)
Pb3	0.862(3)	0.47019(9)	0.22291(8)	0.27319(5)	0.0331(5)	1.829(19)
Pb4	0.865(3)	0.18826(9)	0.36382(8)	0.63576(6)	0.0400(5)	1.772(19)
Pb5	0.776(3)	0.97785(9)	0.41404(8)	0.11436(6)	0.0260(5)	1.82(2)
Fe1/Mg1	0.62(2)/0.38(2)	1	1	0	0.018(2)	2.24(3)
Pb6	0.242(3)	0.1971(4)	0.4301(3)	0.0430(2)	0.047(2)	1.93(2)
Pb7	0.146(3)	0.03-99(5)	0.2196(4)	0.7709(3)	0.030(3)	1.89(2)
Pb8	0.142(3)	0.3150(5)	0.1421(4)	0.8660(3)	0.030(3)	1.89(2)
O1		0.0913(11)	-0.0665(9)	0.7974(7)	0.023(3)*	2.08(3)
O2		0.0981(10)	0.4323(9)	0.3675(6)	0.018(3)*	1.206(16)
O3		0.4133(13)	-0.0543(12)	0.8074(9)	0.042(5)*	2.18(4)
O4		0.4194(12)	0.2765(10)	0.4770(8)	0.027(4)*	2.12(3)
O5		0.9215(11)	0.7776(10)	-0.0263(8)	0.025(4)*	2.12(3)
O6		1.2686(10)	0.8320(8)	-0.1317(6)	0.014(3)*	1.283(19)
O7		0.5713(10)	0.2598(9)	0.3542(6)	0.018(3)*	1.80(5)
O8		0.9032(12)	0.4544(10)	0.3083(8)	0.026(4)*	2.09(3)
O9		0.3690(10)	0.4284(8)	0.3672(6)	0.016(3)*	1.261(16)
O10		0.6419(11)	0.2013(9)	0.4986(7)	0.020(3)*	1.261(19)
O11		0.7685(10)	0.3286(9)	0.3719(6)	0.019(3)*	1.27(2)
O12		0.6265(11)	0.5420(9)	0.1730(7)	0.025(3)*	1.86(3)
O13		0.5863(11)	0.4432(9)	0.3063(7)	0.023(3)*	2.11(3)
O14		0.0834(12)	0.6081(10)	0.1337(7)	0.027(4)*	2.13(3)
O15		0.2550(11)	0.1902(9)	0.5404(7)	0.020(3)*	1.98(3)
O16		0.7605(11)	0.6852(9)	0.0397(7)	0.023(3)*	1.92(3)
O17		0.8854(11)	0.2093(9)	0.4979(7)	0.025(3)*	1.243(18)

O18	0.7116(12)	0.2575(10)	0.6124(7)	0.031(4)*	1.65(6)
O19	0.5745(11)	0.1264(10)	0.6393(7)	0.023(3)*	2.08(3)
O20	1.0637(11)	0.9043(9)	-0.0475(7)	0.022(3)*	1.73(6)
O21	0.3867(11)	0.7066(9)	-0.0031(7)	0.024(3)*	1.275(19)
O22	0.8698(10)	0.9315(9)	-0.1340(6)	0.019(3)*	1.219(16)
O23	0.5446(12)	1.0730(10)	-0.0756(8)	0.035(4)*	0.0158(5)
O24	0.5206(11)	0.7666(9)	-0.1382(7)	0.022(3)*	1.74(4)
O25	0.2096(10)	0.3929(9)	0.4993(6)	0.017(3)*	1.87(5)
O26	0.2083(12)	0.7539(10)	0.1094(7)	0.032(4)*	1.63(5)
O27	0.8848(11)	0.7363(9)	0.1224(7)	0.026(3)*	1.60(5)
O28	1.0774(11)	0.7525(9)	-0.1464(7)	0.023(3)*	1.67(5)
O29	1.1420(11)	0.6983(9)	-0.0045(7)	0.019(3)*	1.227(18)
O30	0.5726(11)	0.7295(9)	0.1433(7)	0.022(3)*	1.74(5)
O31	-0.0129(11)	0.6115(9)	0.2723(7)	0.024(3)*	1.93(5)
O32	0.5986(10)	0.9344(9)	-0.1298(6)	0.017(3)*	1.211(16)
O33	1.0159(11)	0.2690(9)	0.3601(7)	0.023(3)*	1.81(5)
O34	0.0688(12)	0.2863(10)	0.4863(7)	0.029(4)*	2.06(3)
O35	0.4782(11)	0.0771(9)	0.5359(7)	0.023(3)*	1.49(5)
O36	0.7707(11)	0.7686(9)	-0.1192(7)	0.023(3)*	1.63(5)
O37	0.1637(13)	0.6894(11)	0.3020(8)	0.044(4)*	0.0204(6)
O38	0.3895(11)	0.0552(9)	0.6809(7)	0.023(3)*	1.92(3)
O39	0.8560(11)	0.4142(10)	0.4595(7)	0.027(3)*	1.59(5)
O40	0.8945(11)	0.5407(9)	0.1709(7)	0.025(3)*	1.91(3)
O41	0.1386(12)	0.0414(10)	0.6659(8)	0.035(4)*	1.57(3)
O42	0.7904(11)	0.0467(9)	0.5790(7)	0.024(3)*	1.86(5)
O43	0.2763(11)	0.2681(9)	0.3779(7)	0.023(3)*	1.82(5)
O44	0.3383(9)	0.3771(8)	0.1210(6)	0.009(3)*	0.110(3)
O45	0.8009(11)	0.3875(10)	0.2179(7)	0.027(3)*	1.68(5)
O46	0.5375(12)	0.4237(10)	0.0715(7)	0.029(4)*	0
O47	0.7347(10)	0.0708(9)	0.7308(6)	0.019(3)*	1.259(18)
O48	0.2334(11)	0.5642(9)	0.2324(7)	0.023(3)*	1.237(18)
O49	0.4195(12)	0.6214(11)	0.1394(8)	0.032(4)*	2.07(3)
O50	0.4822(11)	0.1089(9)	0.7840(7)	0.022(3)*	1.93(5)
O51	0.3543(12)	0.9081(10)	-0.0377(7)	0.027(3)*	1.63(5)
O52	0.9824(11)	0.5753(9)	0.0243(7)	0.020(3)*	1.84(4)
O53	0.5674(10)	0.3984(9)	0.4601(6)	0.017(3)*	1.62(5)
O54	0.3051(11)	0.5431(9)	0.0778(7)	0.022(3)*	1.61(5)
O55	0.6269(13)	-0.0711(11)	0.7168(8)	0.040(4)*	1.80(5)
O56	0.3661(11)	0.2460(9)	0.6258(7)	0.025(3)*	1.85(5)
O57	0.3234(11)	-0.1116(10)	0.7142(7)	0.027(3)*	1.66(5)
O58	0.7074(11)	0.5843(10)	0.2783(7)	0.026(3)*	1.83(5)
O59	0.6034(12)	0.5921(10)	0.0254(7)	0.029(4)*	1.49(5)
O60	0.1217(12)	0.4186(10)	0.2184(7)	0.032(4)*	1.63(5)
O61	0.1581(12)	-0.1940(10)	0.6882(7)	0.029(4)*	0
O62	0.1175(12)	0.0890(10)	0.5142(8)	0.036(4)*	1.33(4)
O63	0.5676(10)	0.7897(9)	-0.0118(7)	0.018(3)*	2.03(4)
O64	-0.0776(12)	-0.0694(10)	0.7115(7)	0.035(4)*	1.87(6)
O65	0.4315(13)	0.4201(11)	0.2199(8)	0.038(4)*	1.69(6)
O66	-0.0643(11)	0.0997(9)	0.7798(7)	0.024(3)*	1.75(6)
O67	0.7132(10)	0.8952(9)	0.0000(6)	0.017(3)*	1.70(5)

O68		0.2040(12)	0.0675(10)	0.7909(8)	0.035(4)*	1.61(5)
O69		0.3070(14)	-0.0599(12)	0.5604(9)	0.048(5)*	0
O70		0.4776(18)	-0.0868(15)	0.6098(11)	0.081(7)*	0.210(7)
O71		1.0388(11)	-0.0868(10)	0.5766(7)	0.025(3)*	0.249(6)
O72		0.0718(15)	0.2366(13)	0.6408(9)	0.058(5)*	1.30(4)
O73		0.4279(11)	0.6030(10)	0.2808(7)	0.029(3)*	1.85(6)
O74		0.0351(14)	0.5711(12)	0.4209(9)	0.051(5)*	0.116(3)
O75		0.6599(12)	0.1895(10)	0.8039(8)	0.034(4)*	0.123(3)
O76		0.2442(11)	0.2423(10)	0.7573(7)	0.031(4)*	0.516(8)
O77		0.1987(12)	0.5527(10)	-0.0634(8)	0.035(4)*	0.415(9)
O78		1.1511(12)	0.9506(10)	0.0539(7)	0.032(4)*	0.184(6)
O79		-0.0779(13)	0.1232(11)	0.6358(8)	0.035(4)*	2.05(3)
O80		1.3589(11)	0.6965(9)	-0.1864(7)	0.024(3)*	0.191(5)
O81		0.2462(13)	0.2491(11)	0.2473(8)	0.048(5)*	0.322(6)
O82		0.9164(12)	0.9135(10)	0.0806(7)	0.029(4)*	0.297(9)
O83		0.4094(13)	0.4197(11)	0.5856(8)	0.043(4)*	0.214(4)
O84		0.3286(15)	0.5479(12)	0.4356(9)	0.056(5)*	0
O85	0.5	1.030(2)	0.278(2)	0.2254(15)	0.032(8)*	0.221(13)

*Refined with isotropic atomic displacement parameters.

Table 3. Interatomic distances among U atoms (in Å) in the structure of richetite.

U1–O4	2.216(12)	U2–O1 ⁱ	2.215(12)	U3–O1	2.256(13)
U1–O7	1.829(16)	U2–O5	2.189(12)	U3–O3	2.185(15)
U1–O9	2.657(11)	U2–O6	2.381(12)	U3–O6 ⁱⁱ	2.666(10)
U1–O10	2.355(11)	U2–O20	1.843(17)	U3–O38	2.208(12)
U1–O11	2.372(12)	U2–O22	2.658(11)	U3–O41	2.314(14)
U1–O13	2.227(12)	U2–O28	1.830(17)	U3–O57	1.849(17)
U1–O53	1.801(16)	U2–O29	2.388(11)	U3–O68	1.816(18)
<U1–O _{Ur} >	1.82	<U2–O _{Ur} >	1.84	<U3–O _{Ur} >	1.83
<U1–O _{eq} >	2.37	<U2–O _{eq} >	2.37	<U3–O _{eq} >	2.33
U4–O4	2.226(13)	U5–O3 ⁱⁱⁱ	2.246(14)	U6–O8	2.210(15)
U4–O10	2.876(12)	U5–O6 ^{iv}	2.431(14)	U6–O11	2.807(11)
U4–O15	2.246(13)	U5–O21	2.342(12)	U6–O12	2.212(13)
U4–O19	2.211(14)	U5–O24	1.827(15)	U6–O13	2.233(12)
U4–O35	1.842(16)	U5–O32	2.610(14)	U6–O40	2.236(12)
U4–O38	2.217(12)	U5–O51	1.795(15)	U6–O45	1.837(17)
U4–O56	1.854(16)	U5–O63	2.211(14)	U6–O58	1.866(17)
<U4–O _{Ur} >	1.85	<U5–O _{Ur} >	1.81	<U6–O _{Ur} >	1.85
<U4–O _{eq} >	2.36	<U5–O _{eq} >	2.37	<U6–O _{eq} >	2.34
U7–O2	2.487(11)	U8–O2 ^v	2.577(14)	U9–O12	2.194(11)
U7–O8 ^{iv}	2.240(13)	U8–O8	2.241(13)	U9–O16	2.210(12)
U7–O14	2.208(12)	U8–O11	2.378(14)	U9–O21	3.011(14)
U7–O31	1.791(15)	U8–O17	2.371(12)	U9–O30	1.882(17)
U7–O40 ^{iv}	2.518(14)	U8–O33	1.817(15)	U9–O49	2.206(12)
U7–O48	2.430(14)	U8–O34 ^v	2.259(13)	U9–O59	1.868(18)
U7–O60	1.826(16)	U8–O39	1.808(16)	U9–O63	2.252(12)
<U7–O _{Ur} >	1.81	<U8–O _{Ur} >	1.81	<U9–O _{Ur} >	1.88
<U7–O _{eq} >	2.38	<U8–O _{eq} >	2.37	<U9–O _{eq} >	2.38
U10–O5	2.234(13)	U11–O14	2.247(13)	U12–O9	2.430(10)
U10–O14 ^v	2.208(13)	U11–O21	2.335(12)	U12–O12	2.525(12)
U10–O16	2.231(13)	U11–O26	1.795(17)	U12–O13	2.211(12)
U10–O27	1.821(16)	U11–O29 ^{iv}	2.425(14)	U12–O48	2.447(12)
U10–O29	2.770(12)	U11–O48	2.638(12)	U12–O49	2.256(13)
U10–O40	2.285(12)	U11–O49	2.241(14)	U12–O65	1.812(19)
U10–O52	1.874(16)	U11–O54	1.815(15)	U12–O73	1.810(18)
<U10–O _{Ur} >	1.85	<U11–O _{Ur} >	1.81	<U12–O _{Ur} >	1.81
<U10–O _{eq} >	2.35	<U11–O _{eq} >	2.38	<U12–O _{eq} >	2.37
U13–O10	2.378(14)	U14–O5	2.241(12)	U15–O3	2.197(13)
U13–O17	2.357(12)	U14–O16	2.433(12)	U15–O19	2.265(12)
U13–O18	1.789(17)	U14–O22	2.463(12)	U15–O32 ^{vi}	2.483(11)
U13–O19	2.221(12)	U14–O32	2.444(11)	U15–O38	2.419(14)
U13–O42	1.820(16)	U14–O36	1.804(17)	U15–O47	2.480(14)
U13–O47	2.566(11)	U14–O63	2.274(12)	U15–O50	1.796(15)
U13–O79 ^v	2.247(14)	U14–O67	1.787(15)	U15–O55	1.819(16)
<U13–O _{Ur} >	1.80	<U14–O _{Ur} >	1.80	<U15–O _{Ur} >	1.81
<U13–O _{eq} >	2.35	<U14–O _{eq} >	2.37	<U15–O _{eq} >	2.37
U16–O2	2.471(11)	U17–O15	2.161(12)	U18–O1	2.222(12)
U16–O4	2.227(13)	U17–O17 ^{iv}	2.943(14)	U18–O22 ⁱⁱ	2.422(10)
U16–O9	2.409(12)	U17–O34	2.166(12)	U18–O41	2.684(12)

U16–O15	2.390(12)	U17–O41	2.228(13)	U18—O47iv	2.427(12)
U16–O25	1.794(15)	U17–O62	2.005(19)	U18—O64	1.784(18)
U16–O34	2.294(13)	U17–O72	2.01(2)	U18—O66	1.774(17)
U16–O43	1.805(16)	U17–O79	2.186(12)	U18—O79	2.290(13)
<U16–O _{Ur} >	1.80	<U17–O>	2.24	<U16–O _{Ur} >	1.78
<U16–O _{eq} >	2.36			<U16–O _{eq} >	2.41

Symmetry codes: (i) $x+1, y+1, z-1$; (ii) $x-1, y-1, z+1$; (iii) $x, y+1, z-1$; (iv) $x-1, y, z$; (v) $x+1, y, z$; (vi) $x, y-1, z+1$; (vii) $-x+2, -y, -z+1$; (viii) $-x+1, -y, -z+1$; (ix) $-x+1, -y+1, -z+1$; (x) $-x+1, -y+1, -z$; (xi) $-x+2, -y+1, -z$; (xii) $-x, -y+1, -z+1$; (xiii) $-x+2, -y+2, -z$; (xiv) $-x+1, -y+2, -z$.

Table 4. Interatomic distances among Pb and M^{2+} sites (in Å) in the structure of richetite.

Pb1–O33 ^{vii}	2.613(12)	Pb2–O30	2.434(15)	Pb3–O7	2.531(16)
Pb1–O42	2.519(13)	Pb2–O50 ^{ix}	2.578(13)	Pb3–O24 ^x	2.664(15)
Pb1–O43 ^{viii}	2.998(16)	Pb2–O56 ^{ix}	2.874(17)	Pb3–O43	2.986(13)
Pb1–O55	2.954(13)	Pb2–O58	2.562(13)	Pb3–O55 ^{viii}	2.919(18)
Pb1–O62 ^{viii}	2.617(16)	Pb2–O73	2.535(17)	Pb3–O57 ^{viii}	2.710(12)
Pb1–O64 ^v	2.729(18)	Pb2–O75 ^{ix}	2.915(13)	Pb3–O65	2.952(15)
Pb1–O71	2.616(13)	Pb2–O76 ^{ix}	2.695(13)	Pb3–O70 ^{viii}	2.655(16)
Pb1–O81 ^{viii}	2.851(13)	Pb2–O83 ^{ix}	3.081(13)	Pb3–O80 ^{xi}	2.703(12)
<Pb1–O>	2.74	<Pb2–O>	2.71	Pb3–O81	2.766(13)
				<Pb3–O>	2.77
Pb4–O25	2.606(12)	Pb5–O28 ^{xi}	2.772(15)	Pb6–O16 ^x	2.847(17)
Pb4–O31 ^{xii}	2.591(12)	Pb5–O40	2.623(16)	Pb6–O36 ^x	3.013(14)
Pb4–O56	2.548(12)	Pb5–O45	2.683(12)	Pb6–O44	2.350(13)
Pb4–O58 ^{ix}	2.798(18)	Pb5–O52	2.613(12)	Pb6–O52 ^{iv}	3.051(12)
Pb4–O72	2.73(2)	Pb5–O52 ^{xi}	2.707(15)	Pb6–O52 ^x	2.811(16)
Pb4–O74 ^{xii}	2.949(14)	Pb5–O60 ^v	3.023(16)	Pb6–O54	2.839(18)
Pb4–O76	2.637(12)	Pb5–O77 ^x	2.417(14)	Pb6–O59 ^x	2.554(12)
Pb4–O83	2.905(13)	Pb5–O85	2.58(3)	Pb6–O77	2.359(14)
<Pb4–O>	2.72	<Pb5–O>	2.68	<Pb6–O>	2.72
Pb7–O27 ^{ix}	2.871(18)	Pb8–O23 ^{vi}	3.039(14)	M^{2+} –O20	2.046(16)
Pb7–O31 ^{xii}	2.534(15)	Pb8–O27 ^{ix}	2.768(13)	M^{2+} –O20 ^{xiii}	2.046(16)
Pb7–O37 ^{xii}	2.924(14)	Pb8–O30 ^{ix}	2.693(17)	M^{2+} –O78	2.148(13)
Pb7–O66	2.507(17)	Pb8–O50	2.439(13)	M^{2+} –O78 ^{xiii}	2.148(13)
Pb7–O68	2.744(14)	Pb8–O67 ^{ix}	2.533(13)	M^{2+} –O82	2.013(12)
Pb7–O72	2.511(19)	Pb8–O68	2.93(2)	M^{2+} –O82 ^{xiii}	2.013(12)
Pb7–O76	2.529(15)	Pb8–O76	2.394(13)	< M^{2+} –O>	2.07
Pb7–O82 ^{ix}	3.073(13)	Pb8–O82 ^{ix}	3.023(14)		
<Pb7–O>	2.71	<Pb8–O>	2.73		

Symmetry codes: (i) $x+1, y+1, z-1$; (ii) $x-1, y-1, z+1$; (iii) $x, y+1, z-1$; (iv) $x-1, y, z$; (v) $x+1, y, z$; (vi) $x, y-1, z+1$; (vii) $-x+2, -y, -z+1$; (viii) $-x+1, -y, -z+1$; (ix) $-x+1, -y+1, -z+1$; (x) $-x+1, -y+1, -z$; (xi) $-x+2, -y+1, -z$; (xii) $-x, -y+1, -z+1$; (xiii) $-x+2, -y+2, -z$; (xiv) $-x+1, -y+2, -z$.

Table 5. Comparison of the $U17\Phi_7$ polyhedral geometry in richetite with other compounds.

	U- Φ (Å)							O-U-O (°)	BV (v.u.)	Ref.
Richetite (U17)	2.161	2.943	2.166	2.228	2.01	2.01	2.186	171.8	5.37	this work
Wyartite (U3)	2.07	2.09	2.06	2.14	2.44	2.47	2.480	167.0	5.07	1
Dehyd. wyartite (U2)	2.095	2.095	2.092	2.092	2.476	2.512	2.301	163.6	5.10	2
U ₂ MoO ₈	2.06	2.06	2.11	2.18	2.36	2.46	2.73	178.1	4.92	3
	2.08	2.08	2.13	2.15	2.32	2.35	2.58	164.1	5.12	
USbO ₃	1.93	2.02	2.13	2.30	2.35	2.43	2.50	173.0	5.23	4
UVO ₅	2.05	2.07	2.21	2.21	2.30	2.30	2.32	179.9	5.26	5
U ₅ O ₁₂ Cl	2.06	2.06	2.25	2.25	2.30	2.30	2.54	178.9	4.95	6

1 = Burns and Finch (1999); 2 = Hawthorne et al. (2006); 3 = Serezhkin et al. (1973); 4 = Dickens and Stuttard (1992); 5 = Dickens et al. (1992); 6 = Cordfunke et al. (1985).