1 Revision 1

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3	Carlsonite, (NH ₄) ₅ Fe ³⁺ ₃ O(SO ₄) ₆ ·7H ₂ O, and huizingite-(Al), (NH ₄) ₉ Al ₃ (SO ₄) ₈ (OH) ₂ ·4H ₂ O,
4	two new minerals from a natural fire in an oil-bearing shale near Milan, Ohio
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20	Abstract
21	The new minerals carlsonite (IMA2014-067), $(NH_4)_5 Fe^{3+}_3O(SO_4)_6 \cdot 7H_2O$, and
22	huizingite-(Al) (IMA2015-014), (NH ₄) ₉ Al ₃ (SO ₄) ₈ (OH) ₂ ·4H ₂ O, formed from a natural fire in
23	an oil-bearing shale near Milan, Ohio. Carlsonite crystals are yellow to orange-brown thick
24	tablets, flattened on {001}, or stout prisms, elongated on [110], up to about 0.5 mm in size.
25	The mineral has a tan streak, vitreous luster, Mohs hardness of 2, brittle tenacity, irregular

26	fracture, perfect $\{001\}$ cleavage, calculated density of 2.167 g·cm ⁻³ and is easily soluble in
27	H ₂ O. Carlsonite is optically biaxial (–), $\alpha = 1.576(1)$, $\beta = 1.585(1)$ and $\gamma = 1.591(1)$ (white
28	light). Huizingite-(Al) crystals, typically intergrown in light greenish yellow drusy aggregates,
29	are tabular to bladed, flattened on $\{100\}$, up to about 0.25 mm in maximum dimension. The
30	mineral has a white streak, vitreous luster, Mohs hardness of 21/2, brittle tenacity, irregular
31	fracture, no cleavage, calculated density of 2.026 g·cm ⁻³ and is easily soluble in H_2O .
32	Huizingite-(Al) is optically biaxial (+) with $\alpha = 1.543(1)$, $\beta = 1.545(1)$ and $\gamma = 1.563(1)$
33	(589.6 nm light). Raman and infrared spectroscopy was conducted on both minerals. Electron
34	microprobe analyses provided the empirical formulas
35	$[(NH_4)_{4.64}Na_{0.24}K_{0.12}]_{\Sigma 5.00}Fe^{3+}{}_{3.05}O(SO_4)_6{}^{\cdot}6.93H_2O \text{ and }$
36	$[(NH_4)_{8.76}Na_{0.22}K_{0.02}]_{\Sigma 9.00}(Al_{1.65}Fe^{3+}_{1.34})_{\Sigma 2.99}(OH)_{1.98}(H_2O)_{4.02}(SO_4)_{8.00} \text{ for carlsonite and } B_{1.01}(SO_4)_{1.01}($
37	huizingite-(Al), respectively. Huizingite compositions with Fe>Al were noted. Carlsonite is
38	triclinic, <i>P</i> -1, $a = 9.5927(2)$, $b = 9.7679(3)$ $c = 18.3995(13)$ Å, $\alpha = 93.250(7)^{\circ}$, $\beta = 95.258(7)^{\circ}$,
39	$\gamma = 117.993(8)^{\circ}$, $V = 1506.15(16)$ Å ³ , and $Z = 2$. Huizingite-(Al) is triclinic, <i>P</i> -1, <i>a</i> =
40	9.7093(3), $b = 10.4341(3)$, $c = 10.7027(8)$ Å, $\alpha = 77.231(5)^{\circ}$, $\beta = 74.860(5)^{\circ}$, $\gamma = 66.104(5)^{\circ}$,
41	$V = 948.73(9)$ Å ³ , and $Z = 1$. The crystal structures of carlsonite ($R_1 = 0.030$) and huizingite
42	$(R_1 = 0.040)$ are bipartite, each consisting of a structural unit and an interstitial unit. For
43	carlsonite, the structural unit is a $[Fe^{3+}_{3}O(H_2O)_3(SO_4)_6]^{5-}$ cluster and the interstitial complex is
44	$[(NH_4)_5(H_2O)_4]^{5+}$. For huizingite-(Al), the structural unit is a $[(Al,Fe^{3+})_3(OH)_2(H_2O)_4(SO_4)_6]^{5-}$
45	cluster and the interstitial complex is $[(NH_4)_9(SO_4)_2]^{5+}$. In the carlsonite cluster, three FeO ₆
46	octahedra share a common vertex, while in the huizingite-(Al) cluster, three (Al,Fe)O ₆
47	octahedra form an abbreviated corner-linked chain. The cluster in carlsonite is the same as
48	that in metavoltine, while the huizingite-(Al) cluster is unique. The range of Lewis basicity of
49	the structural unit in carlsonite is $0.23-0.11$ valence units (vu) and in huizingite-(Al) it is
50	0.20–0.12 vu; the corresponding Lewis acidities of the interstitial complexes in these

51	structures are 0.13 and 0.14 vu, respectively. A characteristic Lewis acid strength of 0.13 vu is
52	suggested for NH_4^+ when it is in its most typical coordinations of 7 to 8. Carlsonite is named
53	for the late Ernest H. Carlson (1933–2010), former Professor of Mineralogy at Kent State
54	University. Huizingite is named for Terry E. Huizing (born 1938), Adjunct Curator of
55	Mineralogy for the Cincinnati Museum Center, and Marie E. Huizing (born 1939), Editor of
56	Rocks & Minerals since 1978.
57	
58	Keywords: carlsonite; huizingite-(Al); new mineral; crystal structure; Raman spectroscopy;
59	Infrared spectroscopy; Lewis acidity-basicity; Huron Shale burn site, Milan, Ohio.
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61	INTRODUCTION
62	This paper reports the descriptions of the first two terrestrial (non-meteoritic) minerals
63	to have been first discovered in the state of Ohio. These minerals, carlsonite and huizingite-
64	(Al), formed as the result of a non-anthropogenic fire in an oil-bearing shale along the Huron
65	River.
66	Carlsonite is named for Ernest H. Carlson (1933-2010). Dr. Carlson (Ph.D., McGill
67	University, 1966) was professor of mineralogy at Kent State University in Kent, Ohio, from
68	1966 until his retirement in 2009. He was a Fellow of both the Society of Economic
69	Geologists and the Association of Exploration Geochemists. Dr Carlson is perhaps best
70	known for his studies of Ohio minerals and his authorship of Ohio Geological Survey Bulletin
71	69, Minerals of Ohio (1991). At the time of his death, Dr. Carlson was engaged in initial
72	studies of the minerals of the Huron Shale burn site at which carlsonite and huizingite-(Al)
73	were found, and he performed some of the early identifications of the minerals from this
74	occurrence.

75 Huizingite is named for Terry E. Huizing (born 1938) and Marie E. Huizing (born 76 1939) of Cincinnati, Ohio. Terry has been an avid mineral collector since childhood. He and 77 Marie met in college and married in 1961 following graduation, he with a B.S. in chemical 78 engineering and she with a degree in secondary education with a major in English. Marie 79 caught the mineral-collecting bug from Terry and both became (and still are) very active in 80 the Cincinnati Mineral Society. Marie wrote and edited the society's newsletter, *The Quarry*, 81 for more than 15 years. She was asked to assume the duties of Editor of *Rocks & Minerals* in 82 1978, a job that she has very effectively executed ever since, with Terry at her side serving as 83 a Consulting Editor and helping with the promotion of the magazine. Terry has also served as 84 North American representative for several other mineralogical publications, has published a 85 number of mineralogical articles and, since the late 1970s, has served as the Adjunct Curator 86 of Mineralogy for the Cincinnati Museum Center (formerly the Cincinnati Museum of Natural 87 History). In recognition of these and many other contributions to the earth sciences, both 88 Terry and Marie have received numerous honors. Terry received the Cincinnati Mineral 89 Society Educational Foundation Award in 1984 and the American Federation Scholarship 90 Foundation Award in 1991 for "distinguished achievement in the field of earth sciences". 91 Marie received the Cincinnati Mineral Society Educational Foundation Award in 1978, the 92 Carnegie Mineralogical Award for 1995 and the Mineralogical Society of America's 93 Distinguished Public Service Award for 2007. Terry and Marie Huizing have agreed to the 94 naming of this mineral in their honor. The -(Al) suffix is used to indicate that this mineral is 95 the Al-dominant member of a series with its not-yet-described Fe-dominant counterpart, for 96 which the name huizingite-(Fe) is proposed. 97 The new minerals and their names were approved by the Commission on New

Minerals, Nomenclature and Classification of the International Mineralogical Association

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99 (carlsonite: IMA2014-067; huizingite-(Al): IMA2015-014). The carlsonite description is

based on one holotype and one cotype specimen, both of which are deposited in the
collections of the Natural History Museum of Los Angeles County, catalogue numbers 65544
and 65545. The description of huizingite-(Al) is based on one holotype specimen, which is
deposited in the collections of the Natural History Museum of Los Angeles County, catalogue
number 65576.

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OCCURRENCE

107 Carlsonite and huizingite-(Al) occur in a suite of minerals that resulted from a natural 108 fire in an oil-bearing shale exposed at the interface between an eroded stream cliff (up to 7.5 109 m high) and its talus pile (~4.5 m thick) along the Huron River in north-central Ohio, 110 approximately 6.1 km WSW of Milan, Ohio, U.S.A. (41°16'41.4"N, 82°40'27.0"W). The fire 111 started in September of 2009, as the result of spontaneous combustion, and burned until 112 March of 2011 (Figure 1). The occurrence is referred to as the Huron Shale burn site. 113 The rock unit exposed is the late Devonian Huron Shale member of the Ohio Shale 114 formation, a dark grey to black marine, carbon-rich shale containing ironstone concretions, 115 finely divided pyrite and pyrite nodules, and >10% organic matter. A sample taken from a 116 four-meter cliff exposure near Milan, about six kilometers to the northeast of the fire site, 117 yielded 5.2 gallons of oil per ton of shale (Hoover 1960). The geometry of the talus slope 118 favors access of oxygen to pyrite, but also the sequestration of heat, which led to the 119 spontaneous combustion. The fire was concentrated at the interface between talus slope and 120 cliff, but burned away from the cliff into the talus pile in some areas. It burned to a depth of 121 about 2 m. The mineral suite formed by sublimation and condensation on the surfaces of 122 rocks within the talus pile as gases from the fire cooled (Figure 2). Surface temperatures at the 123 vent were between 204°C and 260°C. A temperature probe operated by the Huron County 124 Engineers Office registered a 426°C reading at a depth of half a meter. These temperatures

125 were measured during a moderately intense stage of the fire. The cliff and talus slope formed 126 as the result of natural erosion; the fire occurred without any form of human intervention and 127 was sustained naturally by the oil in the shale. 128 Minerals directly associated with carlsonite include anhydrite, boussingaultite, 129 gypsum, and lonecreekite. Minerals directly associated with huizingite-(Al) include 130 adranosite-(Al), anhydrite, boussingaultite, mascagnite, and salammoniac. Other minerals in 131 the assemblage include adranosite-(Fe), alunogen, anhydrite, boussingaultite, clairite, 132 ferrinatrite, gypsum, halotrichite, kremersite, letovicite, lonecreekite, mascagnite, metavoltine, 133 pyracmonite, sabieite (2*H* and 3*R* polytypes; Kampf et al. 2014), salammoniac, sulfur, 134 tschermigite, and voltaite. 135 Carlsonite and huizingite-(Al) are exceedingly rare minerals, thus far known only 136 from a few specimens at a single occurrence, and neither has previously been reported as a 137 synthetic phase. They formed at ambient pressure, but in an extreme and ephemeral 138 environment characterized by a steep temperature gradient, indicating that they probably have 139 very narrow stability ranges. 140 141 **APPEARANCE AND PROPERTIES** 142 Carlsonite 143 Carlsonite crystals are thick tablets, flattened on $\{001\}$, or stout prisms, elongated on 144 [110], up to about 0.5 mm in size (Fig. 3). The crystals forms observed are $\{100\}$, $\{001\}$, $\{1-$ 10}, {111}, (1-1-1) and {01-2} (Fig. 4). Cross-hatched twinning was rarely observed under 145 146 crossed polars. Carlsonite is yellow to orange-brown, with a tan streak. Crystals are 147 transparent with vitreous luster and display no fluorescence. The Mohs hardness based upon 148 scratch tests is 2. Tenacity is brittle, fracture is irregular and cleavage is perfect on {001}. The 149 density calculated based on the empirical formula using single-crystal unit-cell data is 2.167

150	$g \cdot cm^{-3}$. Crystals are easily soluble in room temperature H ₂ O. The mineral is optically biaxial
151	(-) with indices of refraction $\alpha = 1.576(1)$, $\beta = 1.585(1)$, and $\gamma = 1.591(1)$ determined in white
152	light. The 2V measured directly using a spindle stage is $80(1)^\circ$. The calculated 2V is 78°.
153	Strong $r > v$ dispersion was observed. The incompletely determined optical orientation is X
154	$\approx \perp \{001\}, Z \approx [110]$. The pleochroism is X yellow, Y and Z orange; $X < Y \approx Z$. The Gladstone-
155	Dale compatibility index $1 - (K_P/K_C)$ is 0.001 for the empirical formula, in the range of
156	superior compatibility (Mandarino 2007).
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158 Huizingite-(Al)

159 Huizingite-(Al) crystals are tabular, on {100}, to bladed, up to about 0.25 mm in

160 maximum dimension. Crystals are typically intergrown in drusy aggregates (Figs. 5 and 6).

161 The crystal forms observed are $\{100\}$, $\{010\}$, $\{001\}$ and $\{110\}$ (Fig. 7). No twinning was

162 observed. Huizingite-(Al) is light greenish yellow, with a white streak. Crystals are

transparent with vitreous luster and display no fluorescence. The Mohs hardness based upon

164 scratch tests is $2\frac{1}{2}$. Tenacity is brittle, fracture is irregular and the mineral displays no

165 cleavage. The density calculated based on the empirical formula using single-crystal cell data

166 is 2.026 g·cm⁻³. Crystals are easily soluble in room temperature H_2O . The mineral is optically

167 biaxial (+) with indices of refraction $\alpha = 1.543(1)$, $\beta = 1.545(1)$, $\gamma = 1.563(1)$ determined

using 589.6 nm light. The 2V measured directly using a spindle stage is $40(3)^\circ$. The calculated

169 2*V* is 37°. Strong r > v dispersion was observed. The optical orientation is $X^{\wedge} \mathbf{a} = 24^{\circ}$, $Y^{\wedge} \mathbf{b} =$

170 25°, $Z \wedge c = 27°$. The pleochroism is X pale yellow, Y and Z nearly colorless; $X > Y \approx Z$.

171 Crystals exhibit irregularly zoned extinction, apparently related to Al:Fe compositional

172 zonation. During conoscopic observation, the isogyres were quite diffuse, presumably due to

the combination of zoned optical properties and strong dispersion. The Gladstone-Dale

compatibility index $1 - (K_P/K_C)$ is -0.015 for the empirical formula, in the range of superior compatibility (Mandarino 2007).

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RAMAN AND INFRARED SPECTROSCOPY

178 Raman analysis of single crystals of huizingite-(Al), within a 0.28 mm cluster of 179 crystals, and a single crystal of carlsonite in a similar sized cluster were conducted with a 180 Renishaw InVia Raman Microscope. The spectrometer interfaced to the microscope employed 181 a 1800 groove/mm grating and a charge-coupled detector. The samples were excited using a 182 HeNe laser (632 nm) that was focused onto the sample using a $20 \times (0.40 \text{ N.A.})$ objective for 183 the huizingite and a $50 \times (0.85 \text{ N.A.})$ objective for carlsonite. The same objectives were 184 employed to collect the back-scattered Raman radiation. The spectra were collected with a spectral resolution of 4 cm⁻¹, over the range of 100–4000 cm⁻¹, using an integration time of 185 186 120 s per scan, and samples in unknown orientations. The collected spectra were essentially featureless above 1600 cm⁻¹. Five individual scans were averaged to produce the final Raman 187 188 spectrum (Figs. 8 and 9). Abscissa values were calibrated using the phonon band of single-189 crystal silicon located at 520.7 ± 0.3 wavenumber. 190 The huizingite spectrum was collected at full laser power (10.0 mW) with no harm to the sample. The following intense bands are observed at (centroid positions in cm^{-1}): 223, 191 192 263, 448, 468, 478, 618, 641, 673, 980, 1003, 1010, 1027, 1064, 1123, 1151 and 1205. Most of the observed bands in the Raman spectra relate to the four structurally distinct SO_4^{-2} 193 194 groups. Based on comparison to other sulfate minerals the following (speculative) band assignments can be made: 1) The most intense bands 980, 1003, 1010, and 1027 cm⁻¹ are from 195 the symmetric stretch, v_1 , of the four sulfate groups; 2) bands 1064, 1123, 1151, and 1205 cm⁻ 196 ¹ are from the v_3 modes of the sulfate groups; 3) bands in the 618-673 cm⁻¹ region are from the 197 198 v_4 modes of the sulfate groups; 4) bands in the 448–478 cm⁻¹ region are from the v_2 modes of

the sulfate groups; and 5) bands at 223 and 263 cm⁻¹ are uncertain but possibly due to ^[VI]Fe-O
modes.

201 The carlsonite spectrum was collected at 10% laser power (1.0 mW) because the full 202 10 mW beam caused sample burning and the formation of hematite, for which a sharp Raman 203 spectrum was produced. After data collection, a 5 mW beam was used for further analysis and 204 did not affect the sample. The following intense bands are observed at (centroid positions in cm⁻¹): 245, 275, 436, 487, 514, 552, 576, 617, 629, 670, 1015, 1066, 1104, 1140, 1160, 1188 205 206 and 1219. Band assignments are similar to those of huizingite-(Al). Raman signals from structural ammonium and water, in the vicinity of 3300 cm⁻¹, 207 208 were also investigated in both samples. For huizingite, there were no observable peaks in this 209 region. With the 10.0 mW beam, a very weak but discernible envelope of peaks was observed 210 around 3300 cm⁻¹ in carlsonite; however, at lower powers these peaks were not detectable. 211 Although the presence of ammonium in huizingite was not detected in Raman and 212 only a weak broad signal was observed in carlsonite, FTIR clearly shows the presence of 213 ammonia in these two minerals. Attenuated total internal reflection (ATR) spectra were 214 collected with a Perkin-Elmer Spotlight 400 infrared microscope interfaced to a Perkin-Elmer 215 Spectrum One Fourier transform infrared spectrometer (FTIR). The system employed a $100 \times$ 216 100 µm, liquid nitrogen cooled, mercury cadmium telluride (HgCdTe) detector. The standard 217 drop-down germanium internal reflection element was employed in conjunction with a $50 \times$ 218 50 μ m aperture. Each spectrum collected (Figs. 10 and 11) represents the average of 128 individual scans possessing a spectral resolution of 4 cm⁻¹. 219 The normal vibrational modes of unbound ammonium have frequencies of 3033 =220 $v_1(A_1)$, 1680 = $v_2(E)$, 3137 = $v_3(F_2)$ and 1400 = $v_4(F_2)$ cm⁻¹ (Nakamoto 1986). Of these only v_3 221 222 and v_4 are IR active; however, the lower symmetry sites of a crystal structure may allow the v_1

and v_2 modes to become active. The frequencies of these modes are also shifted as a result of

224	bonding effects in carlsonite and huizingite. FTIR spectra clearly show the presence of
225	ammonium in these minerals by the 1416 and 1413 cm ⁻¹ bands, respectively. Other modes are
226	observable but their IR absorptions can be difficult to distinguish from those of structural
227	water. For example, the 3044 and 3022 cm ⁻¹ bands, in these minerals respectively, are likely
228	the v_1 mode in ammonium. The ammonium v_3 mode appears at 3176 cm ⁻¹ in carlsonite and at
229	3192 cm ⁻¹ in huizingite. The band at 1667 cm ⁻¹ in huizingite is likely from the ammonium v_2
230	vibrational mode but a band in this region for carlsonite is not observed. In carlsonite the
231	weak band with a maximum at 1624 is most likely the bending mode of the water molecules.
232	This feature is not observed in the huizingite, possibly due to the lower water concentration
233	and broadening of the signal due to hydrogen bonding. Finally, the broad shoulder around
234	3500 cm ⁻¹ in carlsonite is likely from the OH stretching modes in water which may in part
235	also contribute to the band associated with ammonium in the vicinity of 3300 cm ⁻¹ (Szakall et
236	al. 2012). Thus, there is an absence of or weak contribution by typical H_2O absorptions,
237	especially in the OH stretching region. Similar results were reported by Knop et al (1985) for
238	$(NH_4)_2[AlF_5(H_2O)]$. They concluded that the OH stretching absorption in that phase underlies
239	the group of peaks in the $3100-2900 \text{ cm}^{-1}$ region that result from ammonium absorptions.
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241	CHEMICAL COMPOSITION
242	Analyses of carlsonite (7 on six crystals) and huizingite-(Al) (10 on four crystals) were
243	performed at the University of Utah on a Cameca SX-50 electron microprobe with four
244	wavelength-dispersive spectrometers utilizing Probe for EPMA software (Probe Software,
245	Inc.). A 15 KeV accelerating voltage and a 20 μ m beam diameter were used for both minerals;
246	a 20 nA beam current was used for carlsonite and a 10 nA beam current was used for
247	huizingite-(Al). Counting times were 40 s for N and 20 s for other elements. Nitrogen was
248	analyzed with a 60Å W/Si multilayer pseudocrystal (Cameca PC-1). The sample and nitrogen

standard (synthetic AIN) were carbon-coated at the same time to assure an equivalent

thickness of the carbon layer. Other standards employed are albite (Na), sanidine (K),

251 hematite (Fe) and celestine (S). Raw X-ray intensities were corrected for matrix effects with a

 $\phi(\rho z)$ (PAP) algorithm (Pouchou and Pichoir 1991).

253 Electron microprobe analysis of low atomic number elements such as nitrogen is 254 complicated by a low cross-section for ionization and high absorption of the soft X-rays. Our 255 analyses confirmed the presence of significant N; however, our $(NH_4)_2O$ values are less than 256 those predicted by the structures (50% of predicted for carlsonite and 86% of predicted for 257 huizingite-(Al). The nature of the structure suggests that NH_4^+ is weakly bonded and some is 258 probably quickly lost, along with H_2O , under vacuum, either initially during carbon coating of 259 the sample or subsequently in the microprobe chamber. Monitoring of the NK α intensity 260 during each spot analysis showed only a slight decrease in N concentration with time under 261 the electron beam for which a correction was applied, but as noted above, it is likely that NH_3 262 was lost under vacuum prior to the analyses. Unfortunately, there is insufficient material for 263 direct determinations of N or H by CHN analysis. Consequently, the (NH₄)₂O and H₂O 264 contents were calculated by stoichiometry based upon the crystal-structure determinations. 265 For carlsonite the high EPMA total after addition of calculated (NH₄)₂O and H₂O is attributed 266 to the loss of these constituents under vacuum, which results in higher concentrations for the 267 remaining constituents. While the loss of (NH₄)₂O and H₂O under vacuum should be expected 268 to provide a high EPMA total for huizingite as well, in fact the analytical total was low. We 269 attribute this to huizingite being very susceptible to beam damage, even when using low beam 270 current and a 20 µm diameter defocused beam. The results for carlsonite are presented in 271 Table 1a and those for huizingite in Table 1b. 272 The empirical formula for carlsonite (based on 6 S *apfu*) is

273 $[(NH_4)_{4.64}Na_{0.24}K_{0.12}]_{\Sigma 5.00}Fe^{3+}_{3.05}O(SO_4)_6 \cdot 6.93H_2O$. The simplified formula is

	274	(NH ₄) ₅ Fe ³⁺ ₃ O(SO ₄) ₆ .	7H ₂ O, which requires	(NH ₄) ₂ O 13.34, F	e ₂ O ₃ 24.54, SO ₃ 49.21, H ₂
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275 12.92, total 100%. The empirical formula for huizingite-(Al) (based on 38 O apfu) is

276 $[(NH_4)_{8.76}Na_{0.22}K_{0.02}]_{\Sigma 9.00}(Al_{1.65}Fe^{3+}_{1.34})_{\Sigma 2.99}(SO_4)_{8.00}(OH)_{1.98}\cdot 4.02H_2O$. The simplified end-

277 member formula is $(NH_4)_9Al_3(SO_4)_8(OH)_2 \cdot 4H_2O$, which for the Al end member requires

278 (NH₄)₂O 20.96, Al₂O₃ 13.68, SO₃ 57.30, H₂O 8.06, total 100 wt%, and for the intermediate

279 member with Al:Fe = 1 requires $(NH_4)_2O$ 20.18, Al_2O_3 6.59, Fe_2O_3 10.31, SO_3 55.16, H_2O

280 7.76, total 100 wt%.

281 Among the 10 analyses of huizingite on which the results in Table 1b are based, the 282 Al:Fe ratio ranged from 0.86 to 2.03 or, in terms of the empirical formula, from $(Al_{1,39}Fe_{1,61})$ 283 to $(Al_{2,03}Fe_{0,99})$. Both Al- and Fe-dominant regions were found in every crystal analyzed and 284 no spatial core-to-rim variation was found in a survey of Al and Fe contents over these and 285 additional crystals. Our findings indicate that, on average, the crystals are significantly higher 286 in Al than Fe and, in crystal regions where Fe is greater than Al, it is only slightly so. Of the 287 10 analyses included in Table 1, five are Al dominant (Al:Fe = 1.43-2.03) and five are Fe dominant (Al:Fe = 0.86–0.97). Furthermore, Al significantly dominates over Fe (on average) 288 289 in the crystal used for the structure refinement $(Al_{1,72}Fe_{1,28})$. Because available material does 290 not allow the characterization of the Fe-dominant phase and the material characterized here is, 291 on average, Al dominant, we are describing only the Al-dominant phase. However, because 292 we have demonstrated the existence of the Fe-dominant phase, we recommend the use of a 293 suffix-based nomenclature for the end-members (Al- and Fe-dominant) of this series, i.e. 294 huizingite-(Al) and huizingite-(Fe). 295 296 X-RAY CRYSTALLOGRAPHY AND STRUCTURE DETERMINATION 297

Both powder and single-crystal X-ray diffraction studies were carried out using a
 Rigaku R-Axis Rapid II curved imaging plate microdiffractometer, with monochromatized

299 MoK α radiation. For the powder-diffraction studies, a Gandolfi-like motion on the φ and ω 300 axes was used to randomize the sample. Observed d values and intensities were derived by 301 profile fitting using JADE 2010 software (Materials Data, Inc.). Data (in Å for MoK α) are 302 given in Table 2 along with the calculated pattern. 303 The Rigaku CrystalClear software package was used for processing intensity data, 304 including the application of empirical multi-scan absorption corrections using ABSCOR 305 (Higashi, 2001). The structures were solved by direct methods using SIR2004 (Burla et al. 306 2005). SHELXL-2013 (Sheldrick 2008) was used for the refinements of the structures. For 307 both structures, the location of all non-hydrogen sites was straightforward. After refinement 308 of these sites with anisotropic displacement parameters, difference-Fourier maps provided the 309 locations of all H atom sites associated with the H₂O and NH₄ groups in carlsonite and the 310 OH, H₂O, and NH₄ groups in huizingite-(Al); however, the H sites associated with N5 in 311 huizingite-(Al) are disordered and partially occupied, as can be expected for a NH_4 group on a 312 center of symmetry $(0, \frac{1}{2}, 0)$. The H sites were refined using soft distance restraints with N–H 313 = 0.90(3), O-H = 0.82(3), H-H of NH₄ = 1.45(3) and H-H of H₂O = 1.30(3) Å. The H-H 314 restraint was not used for the disordered NH_4 group mentioned above. U_{eq} of each H was set 315 to 1.2 times that of the N or O atom to which it is bonded. The refinement of the huizingite 316 structure converged well with these restraints; however, the carlsonite refinement exhibited 317 relatively large shifts in the final refinement cycle: maximum shift/su = 1.323 and mean 318 shift/su = 0.036. The removal of the distance restraints allowed the refinement to converge 319 nicely (maximum shift/su = 0.002 and mean shift/su = 0.000); however, some of the resulting 320 N-H and O-H distances were significantly too high or too low. Consequently, we have 321 elected to report the refinement of the carlsonite structure using the aforementioned restraints. 322 Details of the sample, data collection, and structure refinement for each mineral are provided

323	in Table 3, final atom coordinates and displacement parameters in Table 4, selected bond
324	distances in Table 5 and a bond-valence analysis in Table 6.
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326	DISCUSSION OF THE STRUCTURES
327	Carlsonite
328	Carlsonite is a bipartite structure that consists of a structural unit and an interstitial
329	complex, as elucidated by Schindler and Hawthorne (2001). The $[Fe^{3+}_{3}O(H_2O)_3(SO_4)_6]^{5-}$
330	cluster defines the structural unit. The interstitial complex, ideally $[(NH_4)_5(H_2O)_4]^{5+}$, balances
331	the charge of the structural unit and links the structural units together. The atomic
332	arrangement of carlsonite is depicted in Figure 12 and the structural unit is shown in Figure
333	13.
334	A polytypic relationship may exist between carlsonite and clairite,
335	$(NH_4)_2Fe_3(SO_4)_4(OH)_3\cdot 3H_2O$; however, clairite crystals are generally of too poor quality for
336	definitive single-crystal study. Note that clairite is reported to be slowly soluble in $\mathrm{H_2O}$
337	(Martini 1983). Our examination of clairite crystals from the Huron Shale burn site confirmed
338	them to be very slowly soluble, requiring several hours to dissolve. By contrast, carlsonite
339	crystals are easily soluble in H ₂ O, arguing against a simple polytypic relationship between the
340	species.
341	The $[Fe^{3+}_{3}O(H_2O)_3(SO_4)_6]^{5-}$ cluster in the structure of carlsonite (Figure 6) is identical
342	to that found in the structure of metavoltine, $Na_6K_2Fe^{2+}Fe^{3+}_6O_2(SO_4)_{12}$ ·18H ₂ O (Giacovazzo et al. 1997)
343	al. 1976). In fact, the structures of carlsonite and metavoltine are remarkably similar, as
344	shown in Figure 5. Note that the placements of the NH ₄ groups in the structure of carlsonite
345	are similar to the placements of Na and K in the structure of metavoltine, providing further

346 support for our assignments of the NH₄ sites. The same cluster is also found in the structure of

Maus' salt, $K_5Fe^{3+}_3(SO_4)_6(OH)_2 \cdot nH_2O$, and several related compounds (cf. Scordari et al. 1994).

- 349
- 350 Huizingite-(Al)
- 351 Huizingite-(Al) is also a bipartite structure. The $[(Al,Fe^{3+})_3(OH)_2(H_2O)_4(SO_4)_6]^{5-}$
- 352 cluster defines the structural unit and the interstitial complex has the formula

353 $[(NH_4)_9(SO_4)_2]^{5+}$. The atomic arrangement of huizingite-(Al) is depicted in Figure 14 and the

354 structural unit is compared to that of carlsonite in Figure 13.

355 The structural unit in the structure of huizingite-Al is a cluster containing the same

356 polyhedral components, three $M^{3+}O_6$ octahedra (M^{3+} = Fe or Al) and six SO₄ groups, that

357 make up the cluster in the structure of carlsonite; however, the polyhedra are linked quite

differently. In the carlsonite cluster, three FeO₆ octahedra share a common vertex, while in the

359 huizingite-(Al) cluster, three (Al,Fe)O₆ octahedra form an abbreviated corner-linked chain.

360 The polyhedral cluster in the huizingite-(Al) structure is unique, but has the same topology as

361 a segment of the $[Fe^{3+}(SO_4)_3]^{3-}$ polyhedral chain in the structure of sideronatrite (Scordari and

362 Ventruti 2009).

363

364 NH₄–O bonding

The NH₄–O bond lengths in the structure of carlsonite vary from 2.796 to 3.380 Å, corresponding to coordinations of 7, 6, 8, 7, and 8 for NH₄1, NH₄2, NH₄3, NH₄4, and NH₄5, respectively. Those in huizingite-(Al) vary from 2.729 to 3.281 Å, corresponding to

coordinations of 7, 7, 7, 8, and 6 for NH_41 , NH_42 , NH_43 , NH_44 , and NH_45 , respectively.

- 369 Khan and Baur (1972) surveyed NH₄-containing structures and noted that NH₄-O
- 370 coordinations vary from 4 to 9; for small (4 or 5) coordinations, the NH₄ group behaves more
- 371 like a conventional hydrogen-bond donor, forming nearly linear N–H…O bonds; for higher

372	coordinations, the NH ₄ group behaves more like an alkali cation, with either the NH ₄ group
373	exhibiting orientational disorder or the H bonds being polyfurcated. In a study of the crystal
374	structure of hannayite, Mg ₃ (NH ₄) ₂ (HPO ₄) ₄ ·8H ₂ O, Catti and Franchini-Angels (1976)
375	described the hybrid (or dual) bonding behavior of the NH_4^+ group between an ordered
376	hydrogen-bond donor and a strongly electropositive large alkali-like cation. They observed
377	that this dual behavior for NH_4^+ is apparently quite common. The dual-bonding behavior of
378	NH ₄ is clearly observed in the structures of carlsonite and huizingite-(Al) (Table 5); with the
379	exceptions of HN5a in carlsonite and HN3d in huizingite-(Al), each of the H atoms associated
380	with the NH ₄ groups forms a single short, nearly linear hydrogen bond to an O atom, while
381	other NH ₄ –O bonds are more appropriately regarded as electrostatic in nature. It is
382	noteworthy that the dual-bonding behavior of NH4 was also reported in pyracmonite,
383	$(NH_4)_3 Fe^{3+}(SO_4)_3$ (Demartin <i>et al.</i> , 2010), a mineral that is also found in the Huron River burn
384	site mineral suite.
385	

386 Lewis acidity-basicity

387 Hawthorne and Schindler (2008) noted that the strengths of the relatively weak bonds 388 between the strongly bonded structural unit (SU) and the interstitial complex (IC) usually 389 control the stability of a structure. Furthermore, they pointed out that the interaction between 390 the typically anionic SU and the typically cationic IC can be examined using the Principle of 391 Correspondence of Lewis acidity-basicity. The Lewis basicity (LB) of the SU reflects its 392 capacity to donate electron density and the Lewis acidity (LA) of the IC reflects its capacity to 393 accept electron density; these quantities are essentially equivalent to bond valence and are 394 expressed in valence units (vu). The Principle of Correspondence states that, when the LA 395 closely matches the LB, a stable structure will form.

396	Using the approach detailed by Hawthorne and Schindler (2008), the characteristic
397	range in LB for the SU in carlsonite can be calculated as follows: The effective charge (EC) =
398	the formal charge of the SU less the charge transferred by hydrogen bonding (6 H-bonds) = 5^{-}
399	$-6 \times 0.2 = 6.2^{-}$. The charge deficiency per anion (CDA) = EC per O atom (28 O atoms) =
400	$6.2^{-}/28 = 0.22^{-}$. The range in the average number of bonds (<nb<sub>in>) from the IC to O atoms</nb<sub>
401	in the SU obtained from Figure 8c in Hawthorne and Schindler (2008) $\langle NB_{in} \rangle = 0.95 - 1.95$.
402	The range in total number of bonds (RB) to the SU = $\langle NB_{in} \rangle \times number of O atoms = (0.95 - 100)$
403	1.95 × 28 = 26.6–54.46. The range in LB of the SU of carlsonite = EC / RB = 6.2 / (26.6–
404	54.46) = 0.23–0.11 vu. For huizingite-(Al), the same procedure was used; however, it is worth
405	noting that the unconnected SO ₄ group, included as part of the IC, must be considered as part
406	of the SU for the purpose of computing the LB because of the unconnected SO_4 group's
407	strong internal bonding. The resulting range in LB for the SU in huizingite-(Al) was
408	calculated to be 0.20–0.12 vu.
409	Previous studies of Lewis acidity have largely overlooked the NH_4^+ group. In fact,
410	prior listings of Lewis acid strengths for cations (cf. Hawthorne and Schindler 2008;
411	Hawthorne 2012) do not include the NH_4^+ group. For purposes of calculating the LA of the IC
412	in NH_4 phases, the distinction between hydrogen bonds and electrostatic bonds, discussed
413	earlier, does not seem pertinent and the NH_4^+ group is probably best treated as a normal
414	cation. The LA of the IC in carlsonite and huizingite-(Al) is computed as the formal charge of
415	an NH_4^+ group (+1) divided by the average number of bonds emanating from each of the
416	$\mathrm{NH_4}^+$ groups, modified by any transformer H ₂ O groups. [For an explanation of transformer
417	$\rm H_2O$ groups, the reader is referred to Hawthorne and Schindler (2008)]. The calculated LA of
418	the IC in carlsonite is 0.13 vu and in huizingite-(Al) is 0.14 vu. In both cases, the principle of
419	correspondence of Lewis acidity-basicity holds. Furthermore, the fact that in both cases the

420 match occurs at the lower range of Lewis basicity is an indication that the environment in

421 which these phases formed was not highly acidic.

422 Finally, it is worth considering the typical or average value of Lewis acid strength that should be attributed to the NH_4^+ cation, in general. The 0.13–0.14 vu values noted above for 423 424 carlsonite and huizingite-(Al) correspond to average coordination numbers (CN) in the 7 to 8 425 range. It is well known that NH₄⁺ forms many oxysalts that are isostructural with corresponding K⁺ salts (cf. Khan and Baur, 1972). The most common CN for K⁺ is 8, which 426 427 corresponds to a Lewis acid strength of 0.125 vu. Indeed, Brown (1981) provided a Lewis acid strength of 0.13 vu for K^+ and this has been used by subsequent workers. Consequently, 428 we suggest that 0.13 vu be used as the characteristic Lewis acid strength for NH_4^+ , in most 429 430 cases (7 to 8 CN); however, this value is probably not appropriate for those structures in which NH_4^+ has small (4 or 5) CN. For example, NH_4^+ has an average CN of 4 in 431 $(NH_4)_3PO_4 \cdot 3H_2O$ (Mootz and Wunderlich 1970), with each of three distinct NH_4^+ groups 432 forming four hydrogen bonds to O atoms of PO_4^{3-} groups (which constitute the structural 433 unit). The PO₄³⁻ oxyanion has a characteristic Lewis basicity of 0.25 vu (cf. Hawthorne and 434 Schindler, 2008). In this structure, and others in which NH_4^+ has a CN of 4, it seems more 435 436 appropriate to assign it a Lewis acid strength of 0.25 vu (charge/CN). 437

438 **Relation of the structures to paragenesis**

In a series of papers, Scordari and coworkers examined the structures and stabilities of some hydrated alkali iron sulfates (see Scordari et al. 1994, and references therein). These featured, in particular, phases containing the same $[Fe^{3+}_{3}O(H_2O)_3(SO_4)_6]^{5-}$ cluster that is found in carlsonite and metavoltine. They showed that these compounds gradually alter through dehydration to form phases with the ferrinatrite, Na₃Fe(SO₄)₃·3H₂O, structure and that the transformation involves the rearrangement of the $[Fe^{3+}_{3}O(H_2O)_3(SO_4)_6]^{5-}$ clusters into

chains of composition $[Fe^{3+}(SO_4)_3]^{3-}$. Compounds containing Na⁺, K⁺, and H₃O⁺ were examined; however, phases containing NH₄⁺ were not considered. Interestingly, as noted above, the huizingite-(Al) cluster is essentially a segment of a sideronatrite polyhedral chain.

449

IMPLICATIONS

450 The close structural relationship between carlsonite and metavoltine is intriguing 451 considering that metavoltine is a rather widely distributed mineral occurring as an alteration 452 product of pyrite in arid climates, as a fumarolic sublimate and solfataric precipitate, as a 453 post-mining product, and as a coal-fire sublimate. The last of these modes of occurrence, 454 whether of natural or anthropogenic origin, is rather similar to that at the Huron River burn 455 site. Ammonium-bearing phases are typical of coal-fire occurrences [e.g. the burning coal 456 dumps of the Upper Silesian Coal Basin (Parafiniuk and Kruszewski 2009) and the burning 457 Anna I coal mine dump, Alsdorf, Germany (Witzke et al. 2015)] and can occur, as well, in 458 other types of deposits; yet an ammonium analogue of metavoltine has not been reported, 459 either as a natural or synthetic phase; although Wendlandt and Harrison (2006) did report a 460 NH₄-bearing metavoltine as a precipitate associated with uranium mill tailings disposal cells 461 at the White Mesa Mill in Utah. It seems remarkable that the new mineral carlsonite has not 462 previously been reported from other NH₄ mineral occurrences and particularly from those in 463 which these phases have formed by sublimation from high-temperature gases. There are 464 sufficient similarities in the powder X-ray diffraction patterns of carlsonite and metavoltine to 465 suspect that carlsonite may have been mistaken for metavoltine in some previous studies of 466 NH₄-rich mineral assemblages.

The new heteropolyhedral cluster in the structure of huizingite-(Al) is of interest
simply because of its uniqueness, but more so because of insight that its existence may
provide into the structural and paragenetic relations among the various hydrated ferric sulfate

470 minerals. In particular, it may exist as a complex in aqueous solutions or in solid-state 471 transformations involving the formation and/or breakdown of sideronatrite-style 472 $[Fe^{3+}(SO_4)_3]^{3-}$ chains. The fact that it has thus far only been found in a rare phase formed 473 under extreme and very ephemeral conditions suggests that it has a very narrow stability 474 range and its existence is normally transitory.

- 475
- 476

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574	FIGURE CAPTIONS
575	Figure 1. Huron Shale burn site along the Huron River near Milan, Ohio, in early 2011.
576	
577	Figure 2. Recently formed minerals near the interface between the cliff face and talus slope.
578	The white dendritic crystals are salammoniac. (FOV = 22 cm).
579	
580	Figure 3. Carlsonite crystal. (FOV = 1 mm).
581	
582	Figure 4. Crystal drawing of carlsonite tablet; clinographic projection in standard orientation.
583	
584	Figure 5. Drusy intergrowths of huizingite-(Al) crystals with anhydrite. (FOV = 2.2 mm
585	across).
586	
587	Figure 6. Huizingite-(Al) crystals on anhydrite. (FOV = 0.6 mm across).
588	
589	Figure 7. Crystal drawing of huizingite-(Al) tablet; clinographic projection in standard
590	orientation.
591	
592	Figure 8. Raman spectrum of carlsonite.
593	
594	Figure 9. Raman spectrum of huizingite-(Al).
595	
596	Figure 10. FTIR spectrum of carlsonite.
597	
598	Figure 11. FTIR spectrum of huizingite-(Al).

600	Figure 12. Atomic arrangements of carlsonite and metavoltine, viewed down [100]. $Fe^{3+}O_6$
601	octahedra are orange, $Fe^{2+}O_6$ octahedra are green, SO_4 tetrahedra are yellow, K atoms are
602	light blue spheres, Na atoms are dark blue spheres, N atoms are red spheres, O atoms of
603	isolated H ₂ O groups are large white spheres and H atoms are small white spheres. N–H and
604	O-H bonds are shown as sticks. Unit cells are shown with dashed lines.
605	
606	Figure 13. The structural units in carlsonite and huizingite-(Al). $Fe^{3+}O_6$ and AlO_6 octahedra
607	are orange, SO ₄ tetrahedra are yellow.
608	
609	Figure 14. Atomic arrangement of huizingite-(Al) viewed down [100]. The structural
610	components are as described in figures 12 and 13. Note that the disordered NH ₄ 5 group on the
611	center of symmetry at $(0,\frac{1}{2},0)$.
612	
613	

Const.	wt%	Range	SD	Normalized
$(NH_4)_2O$	6.30	6.03-6.48	0.15	
$(NH_4)_2O*$	12.75			12.30
Na ₂ O	0.79	0.42 - 1.13	0.24	0.76
K ₂ O	0.59	0.47 - 0.72	0.09	0.57
Fe ₂ O ₃	25.70	25.17 - 26.42	0.51	24.79
SO_3	50.67	49.87 - 51.73	0.79	48.88
H_2O^*	13.16			12.70
Total	103.66			100.00

614 Table 1a. Electron microprobe analytical results for carlsonite.

615 616

6	Table 1b.	Electron	micror	orobe a	analyt	ical	results	for	huiz	ingite-	(Aľ)
											`	

Const.	wt%	Range	SD	Normalized
$(NH_4)_2O$	15.88	14.71 – 17.27	0.85	
$(NH_4)_2O^*$	18.57			19.70
Na ₂ O	0.55	0.30 - 1.05	0.27	0.58
K ₂ O	0.09	0.06 - 0.12	0.02	0.10
Al ₂ O ₃	6.85	5.53 - 8.59	1.20	7.27
Fe ₂ O ₃	8.71	6.40 - 10.17	1.36	9.24
SO ₃	52.14	49.91 - 53.66	1.11	55.32
H_2O^*	7.35			7.80
Total	94.26			100.00

617 * based upon the crystal structure determination.

619 Table 2a. Powder X-ray diffraction data for carlsonite.*

1	1	1	100	1 1 1		1	1	T	1 1 1
I _{obs}	$d_{\rm obs}$	$d_{\rm calc}$	Icalc	hKl	I _{obs}	$d_{\rm obs}$	$d_{\rm calc}$	Icalc	h k l
100	9.23	9.1020	100	002			3.0717	8	-312
		8.5695	4	010			3.0376	4	-1-23
		(8 3 9 3 9	25	100	16	3 035	2 0289	7	-215
40	8 76	8 2026	23	110	10	5.055	2 0142	2	213
40	8.20	0.2950	~~~	-110			3.0142	5	2-32
		(8.1161	22	0-11			2.9907	2	1-2.5
		8.0439	6	-101			2.9687	4	-232
		(7.6129	13	-111	10	2 0 4 2	2.9590	3	-132
43	7.57	{ 7.4831	17	1-11	12	2.945	2.9285	3	122
		7 4353	22	011			2 9203	4	-313
		7 2612		101			2 8007	2	212
0	667	6.6206	-	101			2.077	2	212
0	0.07	0.0290	9	-102			2.0944	3	3-2 Z
		6.2011	2	-112			2.84/6	6	-125
4	5.83	5.7953	4	102	10	2.849	{ 2.8363	2	2-33
5	5 28	∫ 5.2642	2	-103			(2.8303	3	-1-2 4
5	3.20	5.2403	4	0-13			(2.7799)	3	-233
		(49453	11	-1-11	5	2 758	2 7465	3	115
23	4.93	1 4 8726	11	-120	-		2 7237	2	1_3 4
		1 9/19	2	-120 112			2.7257	2	225
		4.0440	3	1-1 5			2.7055	2	2-2.5
		4.//5/	9	-210			2.6823	3	213
10	4.74	{ 4.7682	2	1-21	6	2.666	{ 2.6713	2	3-13
		(4.7069	2	013			(2.6592	3	-324
		4.3937	2	-212			2.6311	2	025
8	4 302	4 2847	4	020			2 6286	2	3-32
Ũ		4 2153	5	_2 0 1	7	2 463	2 4668	6	220
		(4.1469)	2	201	1	2.405	2.4000	2	220 241
8	4.136	4.1408	2	-220			2.4312	2	2-41
		(4.1305	2	-1-1-3	6	2 389	2.4010	3	-1-3 1
		4.0814	2	2-12	Ũ	_	(2.3878	2	-420
		4.0580	2	0-22			2.3621	3	-422
		4.0234	2	2-21			2.3534	2	026
		(3.9746	2	201			(2.3442)	3	0-27
6	3.957	3 8978	6	1-23			2 3245	3	-411
14	2 776	2 7080	10	123	10	2.332	2.5215	2	1 2 2
14	5.720	2 7020	10	-123			2.5220	2	-1-5 5
		3.7030	2	0-2.3			2.3105	3	-340
13	3 597	3.5996	2	2-13			2.2947	2	-341
15	5.071	(3.5836	12	113	8	2 268	€ 2.2755	3	008
		3.5168	3	-105	0	2.208	2.2525	3	304
0	2 4 4 1	(3.4524	7	-214			2.2388	2	-342
9	3.441	3.4141	3	1-24	_		(2.2263)	2	-335
		3 3879	3	2_23	5	2.210	2 2085	2	0-3.6
		(2 2260	2	022			2 1020	້າ	120
20	3.328	3.3209	17	023			2.1938	2	4-32
		(3.3148	1/	-204	2	2 1 2 8	2.1402	2	0-4.2
		3.3061	2	1-1 5	-	0	(2.1254	2	108
		3.2693	4	-1-2 1			1.9354	1	3-27
		(3.2536	5	203			1.9277	1	-350
15	3 246	3 2443	8	-124	4	1 9225	1 9241	1	3-51
10	5.210	3 2409	3	120	•	1.7220	1 9195	1	4-25
		2 1566	7	120 214			1 0170	1	
		3.1300	21	2-14			1.91/9	1	223
		3.1382	4	-310	5	1 8628	1.8706	2	1-5 1
		3.1332	4	2-3 1	5	1.0020	(1.8623	2	-4-11
41	3.144	3.1280	3	114	0	1 8216	∫ 1.8204	4	0 0 10
		3.1176	8	121	8	1.0210	1.8039	2	-541
		3.1074	3	-231					
		3 1006	4	-224					
		15.1000	т	-447					

620 *Calculated lines with I < 2 are omitted except those needed to index the observed line at

621 1.9225 Å.

Id	d	1	<u> </u>	I			d	<u>1</u>	<u> </u>	I	d		d	I	h k l
$I_{\rm obs}$ $a_{\rm obs}$	$u_{\rm calc}$	Icalc	пкі	I _{obs}	$a_{\rm obs}$		$u_{\rm calc}$	Icalc	пкі	I _{obs}	$a_{\rm obs}$		$u_{\rm calc}$	Icalc	пкі
30 10.33	10.2428	45	$0\ 0\ 1$				2.8049	5	331	7	1.8706		1.8745	6	520
	9.4581	16	010				2.7976	5	-2-22			1	1.8480	3	-4-32
60 8 82	9 7110	70	100	0	2 772		2 7616	10	202				1 9/25	2	1 / 2
00 8.82	0./110	19	100	2	2.112		2.7040	10	502	-	1 0 0 5 0		1.0455	2	-145
11 8.07	8.0581	39	110				2.7545	2	-212	1	1.83/8	1	1.8385	3	510
AC 7.47	7.4504	8	011				2.7250	3	332				1.8350	2	0-43
26 /.4/	7 3731	44	101			1	2 7 1 9 4	7	233			1	1 8259	2	4-13
20 (55	(52(4	50	0 1 1			1	2.7174	, 7	122			- 4	1.0257	2	2 2 1
28 0.55	0.5304	59	0-1 1				2./100	/	133			,	1.8122	2	3-3 1
5 6.14	6.0830	17	-101	20	2.697	<	2.6973	5	-3-21			1	1.8057	2	254
	5.5952	47	-1-11				2.6860	7	330				1.8012	2	1-34
32 5.60	5 1788	16	110			1	2 6760	5	221	0	1 7080	1	1 7005	3	513
,	5.4788	10	-110			7	2.0709	5	2-2 1	3	1./909	1	1.7995	5	515
(5.1214	20	002				2.6664	2	-130				1.7962	2	0-25
(0 5 0 2 7	5.0312	9	112				2.6623	2	-301			1	1.7926	2	450
69 5.037 Y	5 0122	66	121				2 6558	2	313			1	1 7734	2	-3-14
	4.0251	10	1 1 1				2.0330	2	0 2 2			1	1 7705	2	1 4 5
N N	4.9251	10	1-1 1				2.0111	3	0-2.3			1	1.//05	3	145
1	4.8408	15	102	9	2.571		2.5764	13	141	10	1 7656)	1.7692	2	1-2 5
	4.7573	24	211				2.5574	2	014	10	1.7050		1.7647	2	-5-21
32 4.776 <u>{</u>	1 7122	7	1 1 1				2 5 4 1 0	2	1 2 2				1 7509	2	204
	4.7423	~ ~	-111				2.5410	4	-1-2.5				1./390	2	-304
	4.7290	22	020				2.5324	4	3-11			1	1.7549	5	415
25 4.521	4.5242	45	021	8	2.521		2.5129	10	-123			(1.7429	8	154
(4 3555	11	200				2 4664	2	-203	11	1 7411	2	1 7417	3	-5-11
22 4.337	4 2107	20	200			(2.1001	2	202		1.7 111		1 7250	Š	412
	4.518/	20	201	18	2 4 5 8	- 2	2.4020	9	Z-Z Z			5	1./339	2	-4-1 3
	4.2759	7	0-12	10	2.100	(2.4563	12	-2-13			(1.7209	4	-4-23
1	4.1844	10	221				2.4313	4	-3-31	8	1.7141	2	1.7166	3	3-24
	4 1276	14	1 2 2				2 4204	2	204				1 7124	4	442
	4.1270	14	122				2.4204	4	204			(1./134	4	-4-4 2
41 4.122 {	4.1094	5	-1-21				2.4090	4	-3-12				1.6960	2	260
	4.0949	18	0-21				2.3941	3	0-14			(1.6899	2	-314
	4 0847	32	-102				2 3860	2	024	7	1 6817	2	1 6852	2	4-14
2	4.0047	52	220				2.3000	2	0 4 1	,	1.0017		1.0052	2	000
	4.0290	2	220			- (2.3728	2	041			1	1.6//2	2	026
	3.8477	3	-1-12	11	2 260)	2.3645	4	040			1	1.6689	2	263
10 3.831	3.8292	16	1-12	11	2.309)	2.3614	6	1-14				1.6671	4	360
	3 7252	2	022			(2 3515	3	1 / 1	5	1 6550	1	1 6623	2	160
	3.7232	2	022				2.3313	5	-1-4 1	5	1.0550	1	1.0023	2	100
	3.6866	2	202				2.3289	3	-3-22				1.6429	2	0-16
	3.6593	2	-112			1	2.3225	3	-213			1	1.6417	2	3-33
	3 6320	4	-120			1	2 3 1 6 7	3	1-3.2			1	1 6116	2	550
20 2 524	2 5221	50	1120	12	2.316	<	2.2107	5	242			1	1 6060	2	412
38 3.334	5.5521	20	115			- 1	2.3090	5	243				1.0008	2	-415
	3.4493	3	131			(2.3053	3	431	0	1 6072	1	1.6032	2	-422
	3.4449	19	-121				2.2953	2	134	0	1.00/5		1.5997	2	346
	3 1260	100	2 2 1	5	2 212		2 2108	5	3/3				1 5074	2	161
	3.4209	100	-2-2 1	5	2.212		2.2100	5	242				1.59/4	2	-1-0 1
100 3 427	3.4219	6	2-11				2.1857	2	2-2-3			1	1.5907	3	036
100 5.427	3.4143	7	003	10	2.175		2.1735	8	3-13			1	1.5898	9	504
	3 4018	2	1-2.1				2 1 5 6 0	3	-133				1 5867	2	155
	2 2010	14	120				2.1512	2	1 2 1	12	1.5870	<	1 5011	-	160
	5.5919	14	150				2.1312	4	-4-2 1				1.3044	2	400
	3.3543	33	013	11	2 1 3 8	J	2.1379	10	-3-32			7	1.5800	2	-1-16
1	3.2898	17	231	11	2.150	1	2.1367	3	3-21				1.5700	2	545
	3 2682	2	0 - 2.2				2 1128	3	-124			(1 5586	2	620
	2 2167	12	1 2 2			1	2.0040	0	140	0	1 5565)	1 5572	2	241
	3.2107	12	125		• • • •)	2.0949	0	-140	9	1.5505)	1.5575	5	5-41
	3.2140	3	311	16	2.087	5	2.0922	2	442			(1.5564	3	611
68 3.204 (3.2063	56	-211			(2.0861	8	225			1	1.5457	3	-4-14
	3 1979	12	213				20734	2	1-24				1 5441	5	-3-15
	2 1690	12	210				2.0701	2	212	8	1.5419	<	1 5 4 2 7	Š	0 2 6
	5.1089	2	230			1	2.0001	2	-3-1 3				1.5427	2	0-20
	3.1585	7	132			- (2.0562	12	252			1	1.5324	2	562
	3.1527	31	030			1	2.0513	2	-401	-	1 5110	ſ	1.5155	2	-2-62
	3 1172	4	310	15	2 048	1	2.0474	2	0-42	5	1.5110	5	1 5052	2	-1-26
	2 0067	11	222	10	2.010		2.0171	2	151			2	1 4040	2	555
	3.0907	11	232			- 1	2.0429	3	151	8	1 4888	2	1.4940	3	222
	3.0854	5	0-13			1	2.0253	7	-2-1 4	0	1	C	1.4900	4	406
	3.0813	3	223			1	2.0169	3	424			1	1.4778	2	-4-3 4
	3 0420	76	2_12				2 0086	8	205				1 4760	3	3 2 7
04 2042	2.0210	/0	1 2 1	16	2.0097	<	2.0000	4	203	8	1.4698	<	1.4(02	2	551
94 3.043	5.0310	9	-1-5 1				2.0040	4	352			1	1.4093	2	560
	3.0184	43	312			l	1.9985	5	-3-23			(1.4685	2	-520
	3,0010	8	322				1.9879	2	-142			7	1.4608	4	373
	2 0800	ñ	203			(1 02/6	-	521			1	1 /502	5	2.50
	2.2000	47	203	8	1.9338	3	1.0211	7	214	8	1.4543	<	1.4500	3	452
	2.9586	1	1-13			(1.9311	3	5-14			1	1.4588	4	456
	2.9464	5	301				1.9095	2	-2-5 1			(1.4559	2	-334
	2,9350	2	1-2.2			(1.9086	2	532				1.4370	2	156
	2 8605	-	_1 1 2	6	1 8044)	1 0051		051	7	1 / 207		1 1224	2	625
5 0 0 40	2.0003	4	-1-1-3	0	1.0700		1.7031	4	117	/	1.420/		1.4234	3	023
5 2.840	2.8564	6	032			l	1.8908	2	-115						
							1.8889	2	153						

622	Table 2b.	Powder X-ray	diffraction da	ta for	huizingite-(A	41).*
						/-

623 *Calculated lines with I < 2 are omitted.

624	Table 3a. Sample and crystal data	a for carlsonite.*
625	Diffractometer	Rigaku R-Axis Rapid II
626	X-ray radiation/power	MoK α ($\lambda = 0.71075$ Å)/50 kV, 40 mA
627	Temperature	298(2) K
628	Structural formula	$(NH_4)_5 Fe^{3+}_{3}O(SO_4)_6 \cdot 7H_2O$
629	Space group	<i>P</i> -1
630	Unit-cell dimensions	$a = 9.5927(2)$ Å $\alpha = 93.250(7)^{\circ}$
631		$b = 9.7679(3)$ Å $\beta = 95.258(7)^{\circ}$
632		$c = 18.3995(13)$ Å $\gamma = 117.993(8)^{\circ}$
633	V	$1506.15(16) \text{ Å}^3$
634	Ζ	2
635	Density (for above formula)	2.153 g cm^{-3}
636	Absorption coefficient	1.968 mm ⁻¹
637	<i>F</i> (000)	998
638	Crystal size	200 x 130 x 110 μm
639	θ range	3.07 to 27.45°
640	Index ranges	$-12 \le h \le 12, -12 \le k \le 12, -23 \le l \le 23$
641	Reflections collected / unique	$28480 / 6869; R_{\text{int}} = 0.017$
642	Reflections with $F_0 > 4\sigma(F)$	6430
643	Completeness to $\theta = 27.45^{\circ}$	99.7%
644	Max. and min. transmission	0.813 and 0.694
645	Refinement method	Full-matrix least-squares on F^2
646	Parameters refined/restrained	71/517
647	GoF	1.124
648	Final <i>R</i> indices $[F_0 > 4\sigma(F)]$	$R_1 = 0.0297, wR_2 = 0.0812$
649	R indices (all data)	$R_1 = 0.0314, wR_2 = 0.0826$
650	Largest diff. peak / hole	$+0.94 / -0.77 e/A^3$
651	$R_{int} = \Sigma F_0^2 - F_0^2 (mean) / \Sigma [F_0^2].$	GoF = S = { $\Sigma[w(F_o^2 - F_c^2)^2]/(n-p)$ } ^{1/2} . $R_1 = \Sigma F_o - F_c /\Sigma F_o $.
652	$wR_2 = \{\Sigma[w(F_0^2 - F_c^2)^2] / \Sigma[w(F_o^2 - F_c^2)^2] \}$	$[F_0^2]^2$ $[V_0^2]^{1/2}$; $w = 1/[\sigma^2(F_0^2) + (aP)^2 + bP]$ where <i>a</i> is 0.0505, <i>b</i> is
653	1.0203 and P is $[2F_c^2 + Max]$	$(F_0^2, 0)]/3$
654	-	

655	Table 3b. Sample and crystal data	a for huizingite-(Al).*
656	Diffractometer	Rigaku R-Axis Rapid II
657	X-ray radiation/power	MoK α ($\lambda = 0.71075$ Å)/50 kV, 40 mA
658	Temperature	298(2) K
659	Structural formula	$[(NH_4)_9(SO_4)_2][(Al_{1.72}Fe_{1.28})_3(SO_4)_6(OH)_2(H_2O)_4]$
660	Space group	<i>P</i> -1
661	Unit-cell dimensions	$a = 9.7093(3)$ Å $\alpha = 77.231(5)^{\circ}$
662		$b = 10.4341(3)$ Å $\beta = 74.860(5)^{\circ}$
663		$c = 10.7027(8)$ Å $\gamma = 66.104(5)^{\circ}$
664	V	948.73(9) Å ³
665	Ζ	1
666	Density (for above formula)	2.021 g cm^{-3}
667	Absorption coefficient	1.113 mm ⁻¹
668	<i>F</i> (000)	597
669	Crystal size	90 x 60 x 30 μm
670	θ range	3.12 to 27.45°
671	Index ranges	$-12 \le h \le 12, -13 \le k \le 13, -13 \le l \le 13$
672	Reflections collected / unique	$18529 / 4323; R_{\text{int}} = 0.046$
673	Reflections with $F_0 > 4\sigma(F)$	3543
674	Completeness to $\theta = 27.45^{\circ}$	99.6%
675	Max. and min. transmission	0.967 and 0.906
676	Refinement method	Full-matrix least-squares on F^2
677	Parameters refined/restrained	342 / 50
678	GoF	1.059
679	Final <i>R</i> indices $[F_0 > 4\sigma(F)]$	$R_1 = 0.0399, wR_2 = 0.0874$
680	R indices (all data)	$R_1 = 0.0519, wR_2 = 0.0925$
681	Largest diff. peak / hole	$+0.40 / -0.50 e/A^{3}$
682	$*R_{int} = \Sigma F_o^2 - F_o^2(mean) / \Sigma [F_o^2].$	GoF = S = { $\Sigma[w(F_o^2 - F_c^2)^2]/(n-p)$ } ^{1/2} . $R_1 = \Sigma F_o - F_c /\Sigma F_o $.
683	$wR_2 = \{\Sigma[w(F_0^2 - F_c^2)^2] / \Sigma[w(F_0^2 - F_c^2)^2] \}$	$[F_o^2]_2^2]^{1/2}; w = 1/[\sigma^2(F_o^2) + (aP)^2 + bP]$ where <i>a</i> is 0.0375, <i>b</i> is

684 1.0704 and P is $[2F_c^2 + Max(F_o^2, 0)]/3$

686		x/a	y/b	z/c	U _{eq}	U^{11}	U^{22}	U^{33}	U^{23}	U^{13}	U^{12}
687	Fe1	0.62121(3)	0.25279(3)	0.24651(2)	0.01520(7)	0.01411(13)	0.01468(14)	0.01811(14)	0.00311(10)	0.00349(10)	0.00755(11)
688	Fe2	0.36807(3)	0.39286(3)	0.26517(2)	0.01689(7)	0.01465(14)	0.01465(14)	0.02217(14)	0.00177(10)	0.00219(10)	0.00774(11)
689	Fe3	0.24002(3)	0.00884(3)	0.25780(2)	0.01645(7)	0.01478(14)	0.01498(14)	0.02002(14)	0.00208(10)	0.00407(10)	0.00715(11)
690	S 1	0.42868(6)	-0.06325(6)	0.13719(3)	0.01942(11)	0.0201(2)	0.0189(2)	0.0188(2)	-0.00139(17)	0.00331(17)	0.00918(19)
691	S2	0.71360(6)	0.54664(6)	0.36759(3)	0.01832(10)	0.0169(2)	0.0167(2)	0.0190(2)	0.00004(17)	0.00171(17)	0.00638(18)
692	S3	0.51278(6)	-0.00006(6)	0.36580(3)	0.01964(11)	0.0211(2)	0.0206(2)	0.0191(2)	0.00663(18)	0.00275(18)	0.01097(19)
693	S4	0.06836(6)	0.13994(6)	0.14676(3)	0.01828(10)	0.0176(2)	0.0207(2)	0.0179(2)	0.00111(17)	0.00119(17)	0.01052(19)
694	S5	0.59911(6)	0.49178(6)	0.13715(3)	0.02132(11)	0.0215(2)	0.0221(2)	0.0214(2)	0.00860(19)	0.00461(18)	0.0104(2)
695	S6	0.12297(6)	0.18539(6)	0.37145(3)	0.01986(11)	0.0205(2)	0.0248(2)	0.0188(2)	0.00278(18)	0.00495(17)	0.0140(2)
696	01	0.4974(2)	-0.1674(2)	0.13588(11)	0.0371(4)	0.0406(10)	0.0300(9)	0.0522(11)	0.0044(8)	0.0200(8)	0.0239(8)
697	O2	0.3344(2)	-0.0787(3)	0.06780(10)	0.0440(5)	0.0403(10)	0.0527(12)	0.0252(8)	0.0022(8)	-0.0062(7)	0.0130(9)
698	O3	0.55703(19)	0.10241(18)	0.15437(8)	0.0263(3)	0.0268(8)	0.0218(8)	0.0243(7)	-0.0015(6)	0.0095(6)	0.0058(6)
699	O4	0.32513(18)	-0.10129(17)	0.19731(8)	0.0226(3)	0.0218(7)	0.0192(7)	0.0282(7)	0.0017(6)	0.0093(6)	0.0099(6)
700	O5	0.7032(3)	0.5393(2)	0.44527(10)	0.0418(4)	0.0596(12)	0.0411(10)	0.0236(8)	0.0061(7)	0.0136(8)	0.0216(10)
701	O6	0.8492(2)	0.6918(2)	0.35524(10)	0.0317(4)	0.0224(8)	0.0250(8)	0.0379(9)	0.0040(7)	0.0068(7)	0.0027(7)
702	O7	0.7256(2)	0.40985(19)	0.33607(9)	0.0298(4)	0.0290(8)	0.0281(8)	0.0333(8)	-0.0112(7)	-0.0089(6)	0.0182(7)
703	08	0.5675(2)	0.54338(19)	0.32936(11)	0.0351(4)	0.0231(8)	0.0224(8)	0.0565(11)	-0.0096(8)	-0.0120(7)	0.0126(7)
704	09	0.5916(2)	0.0827(2)	0.43761(9)	0.0355(4)	0.0345(9)	0.0451(10)	0.0225(8)	-0.0019(7)	-0.0020(7)	0.0171(8)
705	O10	0.4965(2)	-0.15715(19)	0.35909(9)	0.0313(4)	0.0373(9)	0.0232(8)	0.0369(9)	0.0121(7)	0.0061(7)	0.0162(7)
706	011	0.60687(18)	0.08306(18)	0.30713(8)	0.0225(3)	0.0214(7)	0.0243(7)	0.0266(7)	0.0106(6)	0.0075(6)	0.0133(6)
707	O12	0.35299(19)	-0.0077(2)	0.35345(8)	0.0267(3)	0.0226(7)	0.0374(9)	0.0251(7)	0.0133(7)	0.0079(6)	0.0166(7)
708	013	0.0973(3)	0.1789(2)	0.07287(9)	0.0405(4)	0.0583(12)	0.0505(11)	0.0245(8)	0.0128(8)	0.0146(8)	0.0332(10)
709	O14	-0.10093(18)	0.0754(2)	0.15429(10)	0.0295(3)	0.0172(7)	0.0282(8)	0.0389(9)	-0.0047(7)	0.0006(6)	0.0089(6)
710	015	0.1212(2)	0.02310(19)	0.16453(9)	0.0269(3)	0.0307(8)	0.0247(8)	0.0281(8)	-0.0049(6)	-0.0063(6)	0.0181(7)
711	016	0.15765(18)	0.28191(17)	0.20106(9)	0.0244(3)	0.0203(7)	0.0192(7)	0.0325(8)	-0.0032(6)	-0.0042(6)	0.0105(6)
712	O17	0.5322(3)	0.4022(3)	0.06609(11)	0.0589(6)	0.0678(15)	0.0686(16)	0.0312(10)	-0.0096(10)	-0.0107(10)	0.0299(13)
713	O18	0.7109(2)	0.6528(2)	0.13069(12)	0.0401(4)	0.0342(9)	0.0261(9)	0.0648(12)	0.0245(9)	0.0180(9)	0.0143(8)
714	019	0.4691(2)	0.4878(2)	0.17653(11)	0.0390(4)	0.0364(10)	0.0509(11)	0.0501(11)	0.0331(9)	0.0238(8)	0.0316(9)
715	O20	0.68350(19)	0.42422(19)	0.18223(9)	0.0274(3)	0.0233(7)	0.0277(8)	0.0376(9)	0.0173(7)	0.0103(6)	0.0148(7)
716	O21	0.1458(2)	0.1434(2)	0.44455(9)	0.0317(4)	0.0362(9)	0.0367(9)	0.0227(8)	0.0091(7)	0.0041(7)	0.0172(8)
717	O22	-0.0144(2)	0.2120(2)	0.36220(9)	0.0349(4)	0.0331(9)	0.0534(11)	0.0330(9)	0.0053(8)	0.0072(7)	0.0323(9)
718	O23	0.2705(2)	0.32705(19)	0.35946(8)	0.0279(3)	0.0315(8)	0.0230(8)	0.0241(7)	-0.0007(6)	0.0077(6)	0.0086(7)
719	O24	0.09649(18)	0.05547(18)	0.31503(8)	0.0248(3)	0.0195(7)	0.0240(8)	0.0291(8)	-0.0028(6)	0.0065(6)	0.0089(6)

685 <u>Table 4a. Atom coordinates and displacement parameters ($Å^2$) for carlsonite.</u>

720	O25	0.41093(16)	0.21995(16)	0.25570(7)	0.0161(3)	0.0141(6)	0.0146(6)	0.0205(6)	0.0024(5)	0.0033(5)	0.0075(5)
721	OW1	0.85062(18)	0.28090(18)	0.23993(9)	0.0228(3)	0.0183(7)	0.0251(8)	0.0286(8)	0.0043(6)	0.0048(6)	0.0129(6)
722	HW1a	0.904(3)	0.290(3)	0.2794(12)	0.027						
723	HW1b	0.857(3)	0.219(3)	0.2110(13)	0.027						
724	OW2	0.3064(2)	0.5696(2)	0.27352(10)	0.0319(4)	0.0327(9)	0.0234(8)	0.0420(9)	-0.0049(7)	-0.0087(7)	0.0186(7)
725	HW2a	0.249(3)	0.578(4)	0.2395(14)	0.038						
726	HW2b	0.353(3)	0.655(3)	0.2965(15)	0.038						
727	OW3	0.0645(2)	-0.2161(2)	0.26112(11)	0.0363(4)	0.0333(9)	0.0187(8)	0.0495(11)	0.0008(7)	0.0231(8)	0.0036(7)
728	HW3a	0.054(4)	-0.296(3)	0.2366(16)	0.044						
729	HW3b	0.003(4)	-0.254(4)	0.2902(15)	0.044						
730	OW4	0.0689(2)	0.5530(2)	0.17521(13)	0.0443(5)	0.0374(11)	0.0252(9)	0.0635(13)	0.0037(9)	-0.0073(9)	0.0120(8)
731	HW4a	0.042(4)	0.587(4)	0.1395(16)	0.053						
732	HW4b	0.003(4)	0.468(3)	0.1749(19)	0.053						
733	OW5	0.8216(5)	0.3194(4)	0.05751(17)	0.0793(9)	0.106(3)	0.0638(18)	0.0733(19)	-0.0013(14)	0.0157(17)	0.0453(18)
734	HW5a	0.894(5)	0.323(6)	0.024(2)	0.095						
735	HW5b	0.746(4)	0.237(5)	0.025(2)	0.095						
736	OW6	0.1324(8)	0.5631(6)	0.4345(4)	0.171(3)	0.215(6)	0.120(4)	0.202(5)	-0.087(4)	-0.095(5)	0.132(4)
737	HW6a	0.037(9)	0.507(10)	0.454(4)	0.205						
738	HW6b	0.100(9)	0.548(10)	0.3879(19)	0.205						
739	OW7	0.0086(3)	0.2942(3)	-0.07344(12)	0.0488(5)	0.0386(11)	0.0510(13)	0.0443(11)	0.0053(10)	0.0016(9)	0.0120(10)
740	HW7a	0.079(3)	0.283(4)	-0.089(2)	0.059						
741	HW7b	-0.069(3)	0.211(3)	-0.076(2)	0.059						
742	N1	0.4767(3)	0.2788(3)	0.50578(12)	0.0367(5)	0.0395(12)	0.0447(13)	0.0306(11)	0.0137(9)	0.0080(9)	0.0223(11)
743	HN1a	0.380(2)	0.263(4)	0.4955(16)	0.044						
744	HN1b	0.538(3)	0.370(3)	0.4912(16)	0.044						
745	HN1c	0.497(4)	0.279(3)	0.5524(11)	0.044						
746	HN1d	0.497(4)	0.211(3)	0.4809(15)	0.044						
747	N2	0.2935(3)	0.1274(3)	-0.02010(11)	0.0337(4)	0.0356(11)	0.0352(12)	0.0266(10)	-0.0011(9)	0.0064(9)	0.0139(10)
748	HN2a	0.314(3)	0.065(3)	0.0086(14)	0.040						
749	HN2b	0.223(3)	0.142(3)	0.0058(15)	0.040						
750	HN2c	0.375(3)	0.211(3)	-0.0159(16)	0.040						
751	HN2d	0.242(3)	0.072(3)	-0.0594(12)	0.040						
752	N3	0.2574(3)	0.5122(3)	0.05027(14)	0.0462(6)	0.0539(16)	0.0478(15)	0.0372(12)	0.0040(11)	-0.0012(11)	0.0259(13)
753	HN3a	0.210(4)	0.559(4)	0.0251(16)	0.055						
754	HN3b	0.326(3)	0.508(4)	0.0219(16)	0.055						

755 756 757 758 759 760 761	HN3c HN3d N4 HN4a HN4b HN4c HN4d	$\begin{array}{c} 0.317(3) \\ 0.188(3) \\ 0.1104(3) \\ 0.197(3) \\ 0.101(4) \\ 0.131(4) \\ 0.029(3) \end{array}$	0.587(3) 0.432(3) -0.1644(3) -0.131(3) -0.249(3) -0.092(3) -0.185(3)	$\begin{array}{c} 0.0872(14)\\ 0.0637(18)\\ 0.47709(12)\\ 0.5133(13)\\ 0.4560(15)\\ 0.4508(14)\\ 0.5018(15) \end{array}$	$\begin{array}{c} 0.055\\ 0.055\\) & 0.0364(5)\\ 0.044\\ 0.044\\ 0.044\\ 0.044\\ 0.044\end{array}$	0.0345(12)	0.0410(13)	0.0329(11)	0.0024(9)	0.0005(9)	0.0183(10)
762 763	N5 HN5a	0.7340(2) 0.818(2)	-0.1376(2) -0.075(3)	0.25778(11) 0.2489(14)) 0.0249(4) 0.030	0.0269(10)	0.0240(9)	0.0268(9)	0.0098(8)	0.0084(8)	0.0130(8)
764	HN5b	0.010(2) 0.733(3)	-0.199(3)	0.2487(14) 0.2887(13)	0.030						
765	HN5c	0.665(3)	-0.186(3)	0.2213(11)	0.030						
766	HN5d	0.692(3)	-0.088(3)	0.2820(13)	0.030						
767											
768						-					
769	Table 4	b. Atom coor	dinates and	displacement	parameters (Å	²) for huizingit	te-(Al).				
770		x/a	y/b	z/c	$U_{ m eq}$	U^{11}	U^{22}	U^{33}	U^{23}	U^{13}	U^{12}
771	Al1	0.5	0.5	0.0	0.0184(3)	0.0193(4)	0.0182(4)	0.0178(4)	-0.0042(3)	-0.0023(3)	-0.0069(3)
772	Al2	0.40551(6)	0.74296(5)	0.21230(5)	0.01774(18)	0.0188(3)	0.0186(3)	0.0167(3)	-0.0022(2)	-0.0046(2)	-0.0070(2)
773	S 1	0.37237(7)	0.24458(7)	0.06816(6)	0.02067(16)	0.0203(3)	0.0226(3)	0.0197(3)	-0.0017(3)	-0.0034(3)	-0.0094(3)
774	S2	0.19631(8)	0.76866(7)	0.01591(7)	0.02286(16)	0.0212(3)	0.0237(3)	0.0241(3)	-0.0041(3)	-0.0083(3)	-0.0059(3)
775	S3	0.69632(8)	0.54010(7)	0.34623(7)	0.02388(16)	0.0233(3)	0.0260(4)	0.0229(3)	-0.0004(3)	-0.0087(3)	-0.0084(3)
776	S4	0.82969(8)	0.87469(8)	0.46665(7)	0.02857(18)	0.0290(4)	0.0314(4)	0.0258(4)	-0.0049(3)	-0.0064(3)	-0.0104(3)
777	01	0.2184(2)	0.2426(2)	0.0949(2)	0.0316(5)	0.0237(10)	0.0381(12)	0.0351(12)	-0.0009(10)	-0.0041(9)	-0.0163(9)
778	O2	0.4652(2)	0.1410(2)	0.15867(19)	0.0298(5)	0.0316(11)	0.0315(11)	0.0268(10)	0.0050(9)	-0.0129(9)	-0.0121(9)
779	O3	0.3605(2)	0.3897(2)	0.07501(19)	0.0273(4)	0.0303(11)	0.0242(10)	0.0275(10)	-0.0066(9)	0.0025(8)	-0.0135(8)
780	O4	0.5546(2)	0.7897(2)	0.06772(18)	0.0270(4)	0.0332(11)	0.0302(11)	0.0219(10)	-0.0067(9)	-0.0005(8)	-0.0174(9)
781	05	0.1315(3)	0.8847(2)	-0.0795(2)	0.0426(6)	0.0561(15)	0.0300(12)	0.0404(13)	-0.0019(10)	-0.0321(12)	-0.0020(11)
782	06	0.0895(2)	0.7015(3)	0.0888(2)	0.0421(6)	0.0299(12)	0.0458(14)	0.0546(15)	-0.0169(12)	0.0065(11)	-0.0218(10)
783	O7	0.3338(2)	0.6619(2)	-0.05358(19)	0.0277(4)	0.0259(10)	0.0286(11)	0.0261(10)	-0.0075(9)	-0.0053(8)	-0.0055(8)
784	08	0.2471(2)	0.8229(2)	0.1065(2)	0.0300(5)	0.0313(11)	0.0289(11)	0.0316(11)	-0.0073(9)	-0.0158(9)	-0.0052(9)
785	O9	0.8258(3)	0.5652(3)	0.2550(3)	0.0496(6)	0.0284(12)	0.0644(17)	0.0488(15)	0.0017(13)	-0.0001(11)	-0.0190(12)
786	O10	0.7139(3)	0.5238(2)	0.4811(2)	0.0409(6)	0.0540(15)	0.0393(13)	0.0305(12)	-0.0002(10)	-0.0237(11)	-0.0112(11)
787	011	0.6796(3)	0.4144(2)	0.3211(2)	0.0402(6)	0.0463(14)	0.0262(11)	0.0528(15)	-0.0047(11)	-0.0229(11)	-0.0101(10)

788	012	0.5535(2)	0.6654(2)	0.3284(2)	0.0289(5)	0.0285(11)	0.0280(11)	0.0294(11)	-0.0049(9)	-0.0129(9)	-0.0046(8)
789	O13	0.8548(4)	0.7691(3)	0.5804(3)	0.0744(10)	0.130(3)	0.0511(17)	0.0597(18)	0.0213(15)	-0.0568(19)	-0.0426(19)
790	O14	0.6914(3)	0.8920(3)	0.4237(2)	0.0486(6)	0.0343(13)	0.0790(19)	0.0398(13)	-0.0089(13)	-0.0069(11)	-0.0279(13)
791	015	0.9596(3)	0.8390(4)	0.3573(3)	0.0640(9)	0.0256(13)	0.110(3)	0.0517(16)	-0.0445(17)	-0.0001(12)	-0.0073(14)
792	016	0.8101(3)	0.0103(3)	0.5023(2)	0.0485(6)	0.0687(17)	0.0422(14)	0.0407(14)	-0.0108(12)	0.0045(12)	-0.0332(13)
793	OH1	0.4586(2)	0.5597(2)	0.16627(19)	0.0228(4)	0.0241(10)	0.0224(10)	0.0230(10)	-0.0026(8)	-0.0092(8)	-0.0069(8)
794	H1	0.523(3)	0.503(3)	0.203(3)	0.027						
795	OW1	0.3499(3)	0.9294(2)	0.2612(2)	0.0303(5)	0.0384(12)	0.0282(11)	0.0246(11)	-0.0031(9)	0.0000(9)	-0.0164(9)
796	HW1a	0.295(3)	0.946(4)	0.333(3)	0.036						
797	HW1b	0.386(4)	0.988(3)	0.235(3)	0.036						
798	OW2	0.2507(2)	0.7242(2)	0.3647(2)	0.0322(5)	0.0260(11)	0.0301(12)	0.0363(12)	0.0034(10)	-0.0061(10)	-0.0098(9)
799	HW2a	0.276(4)	0.664(3)	0.422(3)	0.039						
800	HW2b	0.159(3)	0.756(3)	0.366(3)	0.039						
801	N1	0.4800(3)	0.7877(3)	0.6057(3)	0.0382(6)	0.0377(16)	0.0347(15)	0.0448(17)	-0.0049(14)	-0.0088(13)	-0.0155(13)
802	HN1a	0.555(3)	0.798(3)	0.547(3)	0.046						
803	HN1b	0.486(4)	0.794(3)	0.686(2)	0.046						
804	HN1c	0.393(3)	0.859(3)	0.586(3)	0.046						
805	HN1d	0.465(4)	0.709(3)	0.609(3)	0.046						
806	N2	0.0275(4)	0.8436(4)	0.7176(3)	0.0474(8)	0.054(2)	0.063(2)	0.0396(17)	0.0096(16)	-0.0211(15)	-0.0368(17)
807	HN2a	0.107(3)	0.765(3)	0.716(4)	0.057						
808	HN2b	0.008(4)	0.890(3)	0.783(3)	0.057						
809	HN2c	-0.056(3)	0.835(4)	0.712(3)	0.057						
810	HN2d	0.054(4)	0.903(3)	0.646(3)	0.057						
811	N3	0.1115(4)	0.5045(4)	0.3225(3)	0.0516(8)	0.063(2)	0.052(2)	0.055(2)	0.0019(17)	-0.0280(18)	-0.0311(18)
812	HN3a	0.142(4)	0.517(4)	0.389(3)	0.062						
813	HN3b	0.127(4)	0.409(2)	0.338(4)	0.062						
814	HN3c	0.019(3)	0.552(3)	0.318(4)	0.062						
815	HN3d	0.178(4)	0.515(4)	0.251(3)	0.062						
816	N4	0.2297(3)	0.1194(3)	0.8502(3)	0.0363(6)	0.0359(15)	0.0386(16)	0.0396(16)	-0.0118(13)	-0.0132(13)	-0.0121(13)
817	HN4a	0.261(4)	0.126(3)	0.765(2)	0.044						
818	HN4b	0.230(4)	0.039(2)	0.888(3)	0.044						
819	HN4c	0.134(2)	0.182(3)	0.865(3)	0.044						

820	HN4d	0.286(3)	0.150(3)	0.881(3)	0.044						
821	N5	0.0000	0.5000	0.0000	0.0552(13)	0.072(4)	0.037(3)	0.050(3)	-0.014(2)	-0.030(3)	0.002(3)
822	HN5a	0.044(10)	0.555(9)	0.013(10)	0.066						
823	HN5b	0.061(7)	0.419(5)	0.039(6)	0.066						
824	HN5c	0.059(8)	0.494(9)	-0.081(4)	0.066						
825	Site occu	upancies: Al	1: Al/Fe 0.5	97/0.403(5), A	l2: Al/Fe 0.562	2/0.438(4), HN	15A: 0.50(9), I	HN5B: 0.69(8)), HN5C: 0.58	(9)	

827	Table 5a.	Selected l	bond	distances	(Å)) for carlsonite.
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828	NH ₄ 1–O5	2.830(3)	S1-01	1.4492(17)	Fe1–O25	1.9136(13)
829	NH ₄ 1–O10	2.858(3)	S1O2	1.4522(18)	Fe1–O20	1.9937(15)
830	NH ₄ 1–O21	2.886(3)	S1-O3	1.4949(16)	Fe1-011	2.0087(14)
831	NH ₄ 1–O9	2.902(3)	S104	1.5000(15)	Fe1–O7	2.0100(16)
832	NH ₄ 1–O5	3.151(3)	<s1–o></s1–o>	1.4741	Fe1–O3	2.0220(15)
833	NH ₄ 1–OW6	3.355(6)			Fe1–OW1	2.1019(15)
834	NH ₄ 1–O23	3.380(3)	S2–O5	1.4452(17)	<fe1–o></fe1–o>	2.0083
835	<nh41-o></nh41-o>	3.052	S2–O6	1.4537(17)		
836			S2–O7	1.4844(16)	Fe2025	1.9192(13)
837	NH ₄ 2–O2	2.796(3)	S2–O8	1.4952(16)	Fe2–O8	1.9911(17)
838	NH ₄ 2–O17	2.836(3)	<s2–o></s2–o>	1.4696	Fe2016	2.0014(15)
839	NH ₄ 2–O13	2.847(3)			Fe2019	2.0191(17)
840	NH ₄ 2–O14	2.934(3)	S309	1.4467(17)	Fe2–O23	2.0390(16)
841	NH ₄ 2–O1	2.976(3)	S3010	1.4643(17)	Fe2–OW2	2.0735(16)
842	NH ₄ 2–O18	3.052(3)	S3-011	1.4930(15)	<fe2–o></fe2–o>	2.0072
843	<nh<sub>42–O></nh<sub>	2.907	S3012	1.4944(16)		
844	·		<\$3–O>	1.4746	Fe3025	1.9465(14)
845	NH ₄ 3–OW5	2.911(4)			Fe3–O4	1.9793(14)
846	NH ₄ 3–O13	2.951(3)	S4013	1.4489(17)	Fe3024	2.0030(15)
847	NH ₄ 3–O17	2.961(4)	S4014	1.4642(16)	Fe3015	2.0200(15)
848	NH ₄ 3–OW7	3.016(3)	S4-015	1.4907(15)	Fe3012	2.0360(15)
849	NH ₄ 3–O19	3.038(3)	S4016	1.4920(15)	Fe3–OW3	2.0560(17)
850	NH ₄ 3–O1	3.096(3)	<s4–o></s4–o>	1.4740	<fe2–o></fe2–o>	2.0068
851	NH ₄ 3–OW4	3.170(4)				
852	NH ₄ 3–O17	3.278(4)	S5017	1.4410(21)	Hydrogen bor	nds
853	<nh<sub>43–O></nh<sub>	3.053	S5–O18	1.4491(18)	OW1022	2.765(2)
854			S5019	1.4846(17)	OW1014	2.717(2)
855	NH ₄ 4–OW6	2.838(5)	S5–O20	1.4897(16)	OW2····OW4	2.711(3)
856	NH ₄ 4–O9	2.858(3)	<\$5–O>	1.4661	OW2…O10	2.713(3)
857	NH ₄ 4–O6	2.930(3)			OW3…OW4	2.699(3)
858	NH ₄ 4–O21	2.966(3)	S6-O21	1.4555(16)	OW3…O6	2.676(2)
859	NH ₄ 4–O21	3.042(3)	S6–O22	1.4557(16)	OW4…OW7	2.719(3)
860	NH ₄ 4–O22	3.172(3)	S6-O23	1.4902(17)	OW4…OW1	2.908(3)
861	NH ₄ 4–O12	3.310(3)	S6024	1.5007(15)	OW5…OW7	3.198(4)
862	<nh44o></nh44o>	3.017	<\$6–O>	1.4755	OW5…O2	2.921(4)
863					OW6…none	
864	NH ₄ 5–O14	2.894(3)			OW6…none	
865	NH ₄ 5–O1	2.939(3)			OW7…O18	2.808(3)
866	NH ₄ 5–O18	2.940(3)			OW7…O2	2.976(3)
867	NH ₄ 5–O6	2.986(3)				
868	NH ₄ 5–O10	3.018(3)				
869	NH ₄ 5–O11	3.067(2)				
870	NIL 5 024	2 116(2)				
071	NTI43-024	5.110(5)				
8/1	NH ₄ 5–024 NH ₄ 5–08	3.189(3)				
867 868 869 870	NH ₄ 5–O10 NH ₄ 5–O10 NH ₄ 5–O11	2.940(3) 2.986(3) 3.018(3) 3.067(2) 2.116(3)			0 w / 02	2.770(3)

	873	Table 5b.	Selected bond	distances (Å) for	huizingite-(A
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374	NH ₄ 1–O14	2.824(4)	NH ₄ 4–O14	2.847(4)	S101	1.455()
375	NH ₄ 1–O11	2.957(4)	NH_44-O5	2.860(4)	S1-O2	1.457
376	NH ₄ 1–O2	3.008(4)	NH ₄ 4–O6	2.883(4)	S1–O3	1.488(2
377	NH41-014	3.056(4)	NH4404	2.999(3)	S104	1.4938
78	NH41-O10	3.081(4)	NH ₄ 4–O15	3.092(4)	<si1-0></si1-0>	1.473
379	NH41-016	3.086(4)	NH ₄ 4–O2	3.100(3)		
80	NH ₄ 1–O12	3.281(4)	NH ₄ 4–O9	3.103(4)	S2–O5	1.441(2
81	<nh41-o></nh41-o>	3.042	NH ₄ 4–O1	3.122(3)	S2-O6	1.449(2
82			<nh<sub>44–O></nh<sub>	3.001	S2-O7	1.490(2
883	NH4205	2.797(4)			S2–O8	1.491(2
84	NH ₄ 2–O13	2.902(4)	$NH_{4}5-O1(\times 2)$	2.837(2)	<si2–o></si2–o>	1.468
85	NH ₄ 2–O16	2.962(4)	$NH_{4}5-O9(\times 2)$	2.872(3)		
886	NH ₄ 2–O1	2.996(4)	$NH_{4}5-O6(\times 2)$	2.979(2)	S3–O9	1.454(2
87	NH ₄ 2–O11	3.030(4)	<nh<sub>45–O></nh<sub>	2.896	S3-O10	1.462(2
888	NH ₄ 2–O16	3.254(4)			S3011	1.473(2
89	NH ₄ 2–O15	3.277(5)	$All-OH1(\times 2)$	1.9051(19)	S3012	1.493(2
90	<nh42–o></nh42–o>	3.031	Al1–07(×2)	1.906(2)	<si3–o></si3–o>	1.471
91			$A11-O3(\times 2)$	2.0225(19)		
92	NH₄3–O13	2.729(4)	< <u>A</u> 11–O>	1.945	S4013	1.4490
393	NH ₄ 3–O9	2.836(4)	-		S4014	1.462(2
94	NH ₄ 3–O6	2.860(4)	Al2–OH1	1.9107(19)	S4015	1.463(
95	NH ₄ 3–O10	2.926(4)	Al2–OW2	1.943(2)	S4016	1.472
96	NH ₄ 3–O3	3.163(4)	Al2–04	1.9480(19)	<si4–o></si4–o>	1.462
97	NH ₄ 3–OW2	3.251(4)	A12–O8	1.948(2)		
398	NH ₄ 3–O15	3.258(5)	Al2-OW1	1.950(2)	Hydrogen bo	nds
99	<nh<sub>43–O></nh<sub>	3.003	Al2-012	1.952(2)	OH1…O11	2.786(.
00			<al2–o></al2–o>	1.942	OW1…O16	2.660(.
01					OW1…O2	2.761(.
002					OW2…O10	2.688(
003					OW2…O15	2.601

Table 6a. Bond distances and angles for N–H–O bonds in carlsonite.

900						
907	N–H…O	d(N-H)	<i>d</i> (H···O)	d(N-O)	<n-h…o< td=""><td>NH₄…O bonding*</td></n-h…o<>	NH ₄ …O bonding*
908	N1-HN1a…O21	0.87(2)	2.08(2)	2.886(3)	155(3)	hydrogen bond
909	N1-HN1a…O5	0.87(2)	2.63(3)	3.151(3)	120(2)	electrostatic bond
910	N1-HN1a···O23	0.87(2)	2.84(3)	3.380(3)	122(2)	electrostatic bond
911	N1-HN1b…O5	0.88(2)	1.97(2)	2.830(3)	163(3)	hydrogen bond
912	N1-HN1b…OW6	0.88(2)	3.07(3)	3.357(6)	101(2)	electrostatic bond
913	N1-HN1c…O10	0.86(2)	2.09(2)	2.858(3)	149(3)	hydrogen bond
914	N1-HN1d…O9	0.89(2)	2.02(2)	2.902(3)	168(3)	hydrogen bond
915	N2-HN2a…O2	0.91(2)	1.89(2)	2.796(3)	174(3)	hydrogen bond
916	N2-HN2b…O13	0.92(2)	1.93(2)	2.847(3)	172(3)	hydrogen bond
917	N2-HN2c…O17	0.82(2)	2.16(2)	2.836(3)	140(3)	hydrogen bond
918	N2-HN2c…O1	0.82(2)	2.72(3)	2.976(3)	100(2)	electrostatic bond
919	N2-HN2c…O18	0.82(2)	2.82(3)	3.052(3)	99(2)	electrostatic bond
920	N2-HN2d…O14	0.84(2)	2.10(2)	2.934(3)	176(3)	hydrogen bond
921	N3-HN3a…OW5	0.90(2)	2.07(2)	2.911(4)	156(3)	hydrogen bond
922	N3-HN3b…O17	0.89(2)	2.16(3)	2.961(4)	148(3)	hydrogen bond
923	N3-HN3b…O17	0.89(2)	2.71(3)	3.278(4)	123(3)	electrostatic bond
924	N3-HN3c…O1	0.90(2)	2.26(2)	3.096(3)	154(3)	hydrogen bond
925	N3-HN3c…O19	0.90(2)	2.61(3)	3.038(3)	110(2)	electrostatic bond
926	N3-HN3d…O13	0.82(2)	2.23(2)	2.951(3)	147(3)	hydrogen bond
927	N3-HN3d…OW7	0.82(2)	2.79(3)	3.016(3)	98(2)	electrostatic bond
928	N3-HN3d…OW4	0.82(2)	2.88(3)	3.170(4)	103(2)	electrostatic bond
929	N4-HN4a…O9	0.93(2)	1.97(2)	2.858(3)	160(3)	hydrogen bond
930	N4-HN4b…OW6	0.86(2)	2.01(2)	2.839(5)	161(3)	hydrogen bond
931	N4-HN4b…O6	0.86(2)	2.71(3)	2.930(3)	96(2)	electrostatic bond
932	N4-HN4c…O21	0.84(2)	2.24(2)	2.966(3)	144(3)	hydrogen bond
933	N4-HN4c…O12	0.84(2)	2.76(3)	3.310(3)	124(2)	electrostatic bond
934	N4-HN4d…O21	0.89(2)	2.20(2)	3.042(3)	158(3)	hydrogen bond
935	N4-HN4d····O22	0.89(2)	2.54(3)	3.172(3)	129(2)	electrostatic bond
936	N5-HN5a…O14	0.79(2)	2.30(2)	2.894(3)	133(2)	electrostatic bond
937	N5-HN5a···O24	0.79(2)	2.51(2)	3.116(3)	135(2)	electrostatic bond
938	N5-HN5b…O6	0.85(2)	2.21(2)	2.986(3)	152(2)	hydrogen bond
939	N5-HN5b…O8	0.85(2)	2.47(2)	3.189(3)	143(2)	electrostatic bond
940	N5-HN5c…O1	0.84(2)	2.22(2)	2.939(3)	144(2)	hydrogen bond
941	N5-HN5c…O18	0.84(2)	2.43(2)	2.940(3)	120(2)	electrostatic bond
942	N5-HN5d…O11	0.89(2)	2.22(2)	3.067(2)	159(2)	hydrogen bond
943	N5-HN5d…O10	0.89(2)	2.32(2)	3.018(3)	136(2)	electrostatic bond

944	* The NH ₄ O bond is interpreted as being predominantly an ordered hydrogen bond if	C
945	$d(\text{H} \cdots \text{O})$ is short (< 2.27 Å) and <n–h···o (="" is="" large=""> 140°).</n–h···o>	

/ – /						
948	N–H…O	d(N-H)	<i>d</i> (H···O)	d(N-O)	<n–h···o< td=""><td>NH4…O bonding*</td></n–h···o<>	NH4…O bonding*
949	N1-HN1a…O14	0.86(2)	2.00(2)	2.824(4)	159(3)	hydrogen bond
950	N1-HN1b…O2	0.90(2)	2.15(2)	3.008(4)	161(3)	hydrogen bond
951	N1-HN1c…O14	0.91(2)	2.37(3)	3.056(4)	132(3)	electrostatic bond
952	N1-HN1c…O16	0.91(2)	2.20(2)	3.086(4)	163(3)	hydrogen bond
953	N1-HN1d…O10	0.88(2)	2.66(3)	3.081(4)	110(2)	electrostatic bond
954	N1-HN1d…O11	0.88(2)	2.16(2)	2.957(4)	149(3)	hydrogen bond
955	N2-HN2a…O11	0.87(2)	2.17(2)	3.030(4)	170(3)	hydrogen bond
956	N2-HN2b…O5	0.88(2)	2.10(3)	2.797(4)	135(3)	hydrogen bond
957	N2-HN2c····O1	0.87(2)	2.45(3)	2.996(4)	121(3)	electrostatic bond
958	N2-HN2c…O13	0.87(2)	2.18(3)	2.902(4)	140(3)	hydrogen bond
959	N2-HN2c…O16	0.87(2)	2.80(3)	3.254(4)	114(3)	electrostatic bond
960	N2-HN2d…O15	0.93(2)	2.63(3)	3.277(5)	127(3)	electrostatic bond
961	N2-HN2d…O16	0.93(2)	2.08(2)	2.962(4)	160(3)	hydrogen bond
962	N3-HN3a…O10	0.89(2)	2.09(2)	2.926(4)	157(3)	hydrogen bond
963	N3-HN3a…OW2	0.89(2)	2.71(3)	3.251(4)	121(3)	electrostatic bond
964	N3-HN3b…O13	0.93(2)	1.83(2)	2.729(4)	163(3)	hydrogen bond
965	N3-HN3c…O9	0.84(2)	2.10(3)	2.836(4)	146(3)	hydrogen bond
966	N3-HN3c…O15	0.84(2)	2.91(4)	3.258(5)	107(3)	electrostatic bond
967	N3-HN3d…O3	0.89(2)	2.42(3)	3.163(4)	142(3)	electrostatic bond
968	N3-HN3d…O6	0.89(2)	2.34(3)	2.860(4)	118(3)	electrostatic bond
969	N4-HN4a…O14	0.88(2)	1.98(2)	2.847(4)	166(3)	hydrogen bond
970	N4-HN4a…O15	0.88(2)	2.66(3)	3.092(4)	112(2)	electrostatic bond
971	N4-HN4b…O5	0.85(2)	2.11(2)	2.860(4)	148(3)	hydrogen bond
972	N4-HN4b…O2	0.85(2)	2.77(3)	3.100(3)	105(2)	electrostatic bond
973	N4-HN4c···O6	0.89(2)	2.00(2)	2.883(4)	171(3)	hydrogen bond
974	N4-HN4c···O9	0.89(2)	2.80(3)	3.103(4)	102(2)	electrostatic bond
975	N4-HN4d…O1	0.89(2)	2.51(3)	3.122(3)	126(2)	electrostatic bond
976	N4-HN4d…O4	0.89(2)	2.11(2)	2.999(3)	172(3)	hydrogen bond
977	N5-HN5a…O6	0.90(3)	2.11(4)	2.979(2)	163(9)	hydrogen bond
978	N5-HN5b…O1	0.90(3)	1.95(3)	2.837(2)	170(7)	hydrogen bond
979	N5-HN5c…O9	0.90(3)	1.99(4)	2.872(3)	164(8)	hydrogen bond
980	* The NH ₄ …O bon	d is interpre	eted as bein	g predomina	ntly an ordere	d hydrogen bond if
					-	

946 947 Table 6b. Bond distances and angles for N-H-O bonds in huizingite-(Al).

* The NH₄...O bond is interpreted as being predominantly at $d(\text{H} \cdots \text{O})$ is short (< 2.27 Å) and <N-H \cdots O is large (> 140°). гy 981

	01	02	03	04	05	06	07	08	09	O10	011	012	013	014	015	016	017	O18	019	O20	O21	O22	O23	O24	O25	OW1	OW2	OW3	OW4	OW5	OW6	OW7	$\Sigma_{\rm c}$
NH_41					0.20 0.08				0.16	0.18											0.17		0.04								0.05		0.68
NH ₄ 2	0.13	0.21											0.19	0.15			0.19	0.11															0.98
NH ₄ 3	0.10												0.14				0.14 0.06		0.11										0.08	0.16		0.12	0.61
NH ₄ 4						0.15			0.18			0.05									0.14 0.11	0.08									0.19		0.70
NH ₄ 5	0.15					0.13		0.07		0.12	0.10			0.16				0.15						0.09									0.67
Fe1			0.49				0.51				0.51									0.53					0.66	0.40							3.10
Fe2								0.53								0.52			0.49				0.47		0.65		0.43						3.09
Fe3				0.55								0.47			0.49									0.52	0.60			0.45					3.08
S1	1.60	1.59	1.42	1.40																													6.01
S2					1.62	1.58	1.46	1.42																									6.08
S3									1.61	1.54	1.42	1.42																					5.99
S4													1.61	1.54	1.43	1.43																	6.01
S5																	1.64	1.60	1.46	1.44				1 10									6.14
S6																					1.58	1.58	1.44	1.40									6.00
HW1a						-						-										0.19				0.81							1.00
HW1b						-						-		0.20												0.80							1.00
HW2a						-						-															0.80		0.20				1.00
HW2b						-				0.20		-															0.80						1.00
HW3a						-						-																0.80	0.20				1.00
HW3b						0.21						-																0.79					1.00
HW4a						-						-																	0.80			0.20	1.00
HW4b																										0.14			0.86				1.00
HW5a						-						-																		1.00			1.00
HW5b		0.14																												0.86			1.00
HW6a																					ļ	ļ									1.00		1.00
HW6b																					ļ	ļ									1.00		1.00
HW7a																		0.18			ļ	ļ										0.82	1.00
HW7b		0.11																			ļ	ļ										0.89	1.00
Σ_{a}	1.98	2.05	1.91	1.95	1.90	2.07	1.97	2.02	1.95	2.04	2.03	1.94	1.94	2.05	1.92	1.95	2.03	2.04	2.06	1.97	2.00	1.85	1.95	2.01	1.91	2.15	2.03	2.04	2.14	2.02	2.24	2.03	

Table 7a. Bond-valence analysis for carlsonite. Values are expressed in valence units.*

*NH₄⁺–O bond strengths from Garcia-Rodriguez *et al.* (2000); Fe³⁺–O bond strengths from Brown and Altermatt (1985); Si⁴⁺–O bond strengths from Brese and O'Keeffe (1991).

	01	02	03	04	05	06	07	08	09	010	011	012	013	014	015	016	OH	OW1	OW2	$\Sigma_{\rm c}$
NH ₄ 1		0.12								0.10	0.14	0.06		0.20 0.11		0.10				0.83
NH ₄ 2	0.12				0.21						0.11		0.16		0.06	0.06 0.14				0.86
NH ₄ 3			0.08			0.18			0.19	0.15			0.26		0.06					0.92
NH ₄ 4	0.09	0.09		0.12	0.18	0.17			0.09					0.19	0.10					1.03
NH ₄ 5	0.19 ×2→					0.13 ×2→			0.17 ×2→											0.98
A11			0.40 ×2→				0.55 ×2→										0.55 ×2→			2.99
Al2				0.49				0.49				0.49					0.55	0.49	0.50	3.01
S1	1.58	1.57	1.44	1.42																6.01
S2					1.64	1.60	1.44	1.43												6.11
S3									1.58	1.55	1.50	1.42								6.05
S4													1.60	1.55	1.55	1.51				6.21
H1											0.18						0.82			1.00
HW1a																0.22		0.78		1.00
HW1b		0.19																0.81		1.00
HW2a										0.21									0.79	1.00
HW2b															0.24				0.76	1.00
Σ_{a}	1.98	1.97	1.92	2.03	2.03	2.08	1.99	1.92	2.03	2.01	1.93	1.97	2.02	2.05	2.01	2.03	1.92	2.08	2.05	

Table 7b. Bond-valence analysis for huizingite-(Al). Values are expressed in valence units.*

* NH_4^+ -O bond-valence parameters from Garcia-Rodriguez *et al.* (2000); all other bond-valence parameters from Brown and Altermatt (1985). The All-O and Al2-O bond strengths are based upon the refined Al/Fe site occupancies.





























carlsonite [Fe³⁺₃O(H₂O)₃(SO₄)₆]⁵⁻ cluster huizingite $\label{eq:constraint} \begin{array}{l} \mbox{huizingite} \\ \mbox{[(AI,Fe^{3+})_3(OH)_2(H_2O)_4(SO_4)_6]^{5-}} \\ \mbox{cluster} \end{array}$



