Revision 4 1 2 A new biogenic, struvite-related phosphate, the ammonium-analogue of 3 hazenite, (NH₄)NaMg₂(PO₄)₂·14H₂O 4 5 Hexiong Yang¹, Livia Martinelli^{2,3}, Flavia Tasso³, Anna Rosa Sprocati³, Flavia Pinzari^{2,4}, 6 Zhenxian Liu⁵, Robert T. Downs¹, and Henry J. Sun⁶ 7 ¹Department of Geosciences, University of Arizona, 1040 E. 4th Street, Tucson, Arizona 85721, USA 8 9 ²Istituto Centrale per il Restauro e la Conservazione del Patrimonio Archivistico e Librario, Rome, Italy 10 ³Unità Tecnica Caratterizzazione, Prevenzione e Risanamento Ambientale, ENEA-CASACCIA, Rome, 11 Via Anguillarese 301, 00123 Rome, Italy 12 ⁴Consiglio per la Ricerca e la sperimentazione in Agricoltura Centro di ricerca per lo studio delle relazioni 13 tra pianta e suolo. Via della Navicella 2-4, 00184 Rome, Italy 14 ⁵Geophysical Laboratory, Carnegie Institution of Washington, Washington, D.C. 20015, USA 15 ⁶Desert Research Institute, 755 Flamingo Road, Las Vegas, Nevada 89119, USA 16 17 Abstract 18 A new biogenic, struvite-related phosphate, the ammonium analogue of hazenite 19 (AAH), ideally $(NH_4)NaMg_2(PO_4)_2 \cdot 14H_2O$, has been found in cultures containing the 20 bacterial strain Virgibacillus sp.NOT1 (GenBank Accession Number: JX417495.1) 21 isolated from an XVII Century document made of parchment. The chemical composition 22 of AAH, determined from the combination of electron microprobe and X-ray structural 23 analyses, is [(NH₄)_{0.78}K_{0.22}]NaMg₂(PO₄)₂·14H₂O. Single-crystal X-ray diffraction shows 24 that AAH is orthorhombic with space group *Pmnb* and unit-cell parameters a =6.9661(6), b = 25.236(3), c = 11.292(1) Å, and V = 1985.0(3) Å³. Compared to hazenite, 25 the substitution of NH_4^+ for K⁺ results in a noticeable increase of the average A-O (A = 26 27 $NH_4^+ + K^+$) bond length and the unit-cell volume for AAH, as also observed for struvite 28 vs. struvite-K. Both infrared and Raman spectra of AAH resemble those of hazenite, as 29 well as struvite. Our study reveals that AAH forms only in cultures with Na-bearing 30 solutions and pH below 10.0. No AAH or hazenite was found in experiments with 31

the K-bearing solutions, suggesting the necessity of a Na-bearing solution for AAHformation.

34 *Key words*: ammonium phosphate, hazenite, struvite-type materials, biomineral, crystal

- 35 structure, X-ray diffraction, infrared and Raman spectra
- 36
- 37

Introduction

38 Phosphate formation through microbial activities is one of the most common 39 mechanisms for the biological transformation of inorganic phosphates (Gibson 1974; 40 Kamnev et al. 1999; Desmidt et al. 2013). Among all biogenic phosphates, struvite, 41 $(NH_4)MgPO_4 \cdot 6H_2O_4$, is the most widespread in a variety of environments, such as bat 42 droppings, decomposing foods, infection (e.g., urinary tract) stones in humans, water 43 treatment facilities, and in a range of bacterial cultures (Sánchez-Román et al., 2007; 44 Weil 2008; Desmidt et al. 2013 and references therein). The specific roles that 45 microorganisms play in struvite formation are not well understood. It has been speculated 46 that bacterial cell surfaces may serve as nucleation sites and biological activities provide 47 a steady supply of phosphate and ammonia as the crystals grow (e.g., Ben Omar 1994, 48 1995, 1998; Chen et al. 2010).

49 A number of compounds are isotypic with, or structurally analogous to, struvite 50 (Dickens and Brown 1976; Weil 2008; Yang et al. 2011). A general chemical formula for struvite-type materials can be expressed as $A^+M^{2+}(XO_4) \cdot nH_2O$, where n = 6 - 8, X = P or 51 52 As, $A = NH_4$, K, Rb, Cs, and Tl, and M = Mg, Fe, Co, Ni, Zn, Mn. A common structural 53 feature of struvite-type compounds is that all M cations are octahedrally coordinated by 54 six H₂O molecules and no H₂O molecule is shared between $M(H_2O)_6$ octahedra. The XO₄ 55 tetrahedra and $M(H_2O)_6$ octahedra are interlinked through hydrogen bonding. The 56 struvite-type structure was once thought unable to accommodate A cations smaller than 57 K^+ (Banks et al. 1975). Nevertheless, Mathew et al. (1982) synthesized a Na-analogue of struvite, NaMg(PO₄)·7H₂O, in which the small Na⁺ (relative to K⁺) is compensated by an 58

59	additional H ₂ O molecule. More intriguingly, Yang and Sun (2004) and Yang et al. (2011)
60	described the new biomineral hazenite, KNaMg ₂ (PO ₄) ₂ ·14H ₂ O, which possesses many
61	structural features similar to those for both struvite-(K), KMg(PO ₄)·6H ₂ O (Mathew and
62	Schroeder 1979; Graeser et al. 2008) and synthetic NaMg(PO ₄)·7H ₂ O (Mathew et al.
63	1982). Hazenite represents the first struvite-type phosphate that contains both K and Na
64	as the A ions. In this paper, we report a new biologically-formed phosphate, an
65	ammonium analogue of hazenite, ideally (NH ₄)NaMg ₂ (PO ₄) ₂ ·14H ₂ O.
66	
67	Experimental Methods
68	Formation of the ammonium analogue of hazenite (AAH)
69	The AAH crystals used in this study were formed in cultures containing the
70	bacterial strain Virgibacillus sp.NOT1 (GenBank Accession Number: JX417495.1),
71	which was isolated from an XVII Century document made of parchment and identified
72	through 16S rDNA sequencing. The growth medium was prepared in Blood Agar Base
73	N.2 (Oxoid, Code: CM0271) with the following components: Proteose peptone 15.0 g/L;
74	liver digest 2.5 g/L; yeast extract 5.0 g/L; sodium chloride 5.0 g/L, agar 12.0 g/L. The
75	final pH was 7.4(2) at 25°C. Four different salts at two or three different concentrations
76	were added to the growth media (Table 1). A 10-ml aliquot of each salt-supplemented
77	agar medium was used for the pH measurements before solidification using a glass
78	electrode specific for high viscosity samples (Metrhom 6.0239.100 Viscotrode). The
79	plates were then inoculated by surface streaking and incubated aerobically at 25 °C.
80	Cultures were checked for crystal formation periodically for up to 60 days. As the growth
81	media began to dry, elongated tabular or prismatic AAH crystals (up to $0.50 \times 0.12 \times$
82	0.08 mm) appeared on or in the bacterial colonies in runs #1, 2, and 4 (Fig. 1).
83	

84 Characterization of AAH crystals

The AAH crystals were first examined using a Variable Pressure EVO 50
Scanning Electron Microscope. Qualitative chemical analysis of AAH was performed
with electron-dispersive spectroscopy following the procedure given by Gazulla et al.
(2013), which revealed the major elements P, Mg, Na, K, and N, plus trace Ca. The
presence of N in AAH was further confirmed by the infrared (IR) spectroscopy and X-ray
structure determination (see below).

91 The quantitative analysis of the AAH chemical composition was conducted with a 92 CAMECA SX100 electron microprobe at 10 kV and 5 nA, with a beam size of 20 µm to 93 minimize the sample damage by the electron beam. The average composition of five 94 analysis points is (wt.%): P₂O₅ 42.2(4), MgO 23.7(3), Na₂O 9.0(1), K₂O 2.5(2), and CaO 95 0.16(8), with a sum of 77.5(6) wt.%. Due to the high degree of hydration and the rapid 96 deterioration of the sample during the microprobe analysis, this composition was used 97 only for the estimation of cation ratios. By assuming two P cations per formula, the 98 relative ratio of P:Mg:Na:K is 2.00:1.98:0.98:0.18. The actual composition of the crystal, 99 $[(NH_4)_{0.78}K_{0.22}]$ NaMg₂(PO₄)₂·14H₂O, was determined by the combination of the electron 100 microprobe and X-ray structural analyses (see below), which can be idealized as 101 $(NH_4)NaMg_2(PO_4)_2 \cdot 14H_2O.$

The IR and Raman spectra of AAH were acquired at the U2A beam line of the
National Synchrotron Light Source at Brookhaven National Laboratory. The details of
experimental procedures and the optical layout of the beamline have been described by
Liu et al. (2002).

106 Single-crystal X-ray diffraction data of AAH (from run 2) were collected at 107 ambient temperature on a Bruker X8 APEX2 CCD X-ray diffractometer equipped with 108 graphite-monochromatized Mo K_{α} radiation. All reflections were indexed on the basis of 109 an orthorhombic unit-cell (Table 2). The intensity data were corrected for X-ray 110 absorption using the Bruker program SADABS. The systematic absences of reflections 111 suggest possible space group *Pmnb* (#62) or *P2*₁*nb* (#33). The structure model of hazenite

112	(Yang et al. 2011) was adopted and refined using SHELX97 (Sheldrick 2008) based on
113	the space group <i>Pmnb</i> , because it yielded the better refinement statistics in terms of bond
114	lengths and angles, atomic displacement parameters, and R factors. The H atoms in all
115	H_2O molecules were located, but not those in NH_4^+ . The positions of all atoms were
116	refined with anisotropic displacement parameters, except for H atoms, which were
117	refined with a fixed isotropic displacement parameter ($U_{eq} = 0.04$). The site occupancy of
118	K was refined against N, yielding a relative ratio of $0.22(1) : 0.78(1)$. This ratio was
119	adopted in our empirical formula for AAH. Final coordinates and displacement
120	parameters of atoms are listed in Table 3, and selected bond-distances in Table 4.
121	

122

Discussion

123 Crystal Structure

124 AAH is isostructural with hazenite, which exhibits many structural features 125 resembling those of struvite-type materials (Yang and Sun 2004; Yang et al. 2011). The 126 crystal structure of AAH is characterized by six distinct non-hydrogen cation sites, including two octahedral sites for Mg^{2+} (Mg1 and Mg2), two tetrahedral sites for P^{5+} (P1 127 and P2), one trigonal prismatic site for Na⁺, and one six-coordinated, very irregular site 128 129 for A^+ . These cation sites form three types of layers that are stacked along the *b*-axis in a repeating sequence of ABCBABCB... (Fig. 2), where layer A consists of $Mg1(H_2O)_6$ 130 131 octahedra and NaO₆ trigonal prisms, layer B of P1O₄ and P2O₄ tetrahedra, and layer C of 132 $Mg2(H_2O)_6$ octahedra and AO_6 polyhedra. These layers are linked together by hydrogen 133 bonds, plus the A-O bonds between layers B and C (A-O5-P2). As noted by Yang et al. 134 (2011), the struvite-type structure, which displays a layer stacking sequence of 135 BCBCBC..., can be readily derived from the hazenite-type structure by replacing its layer 136 A with layer C. In general, because the effective radius of NH_4^+ (1.48 Å) is larger than that of K^+ 137

138 (1.38 Å for six-coordination) (Shannon 1976), the substitution of NH_4^+ for K^+ in a crystal

will result in the increase in both the average *A*-O bond length and the unit-cell volume,
as have been observed in numerous compounds (e.g. Abu El-Fadl et al. 2006; Bogdanov
et al. 2011; Shin 2011; Lim and Lee 2013). For AAH and hazenite, the average *A*-O bond
lengths are 2.972 and 2.919 Å (Yang et al. 2011), respectively (Table 4), and the unit-cell
volumes are 1985.0(5) and 1958.7(2) Å³ (Table 5). Similar results are also found for
struvite vs. struvite-K (Table 5).

145

146 Infrared and Raman spectra

147 Figures 3 and 4 show the IR and Raman spectra of AAH. There have been 148 copious IR and Raman spectroscopic studies on struvite-related materials (Banks et al. 149 1975; Angoni et al. 1998; Stefov et al. 2004, 2005; Frost et al. 2005; Koleva 2007; Cahil 150 et al. 2008). Yang and Sun (2004) and Yang et al. (2011) showed that the IR and Raman 151 spectra of hazenite are similar to those of struvite and made the tentative assignments of 152 major bands for both spectra. These assignments should be applicable to AAH as well 153 because of their isotypism. Nonetheless, as in the case for struvite vs. struvite-K (e.g., 154 Stefov et al. 2005; Cahil et al. 2008), the IR spectrum of AAH is expected to be more 155 complicated than that of hazenite owing to the presence of the significant amount of NH_4^+ . The presence of NH_4^+ in crystals generally gives rise to two discernible groups of 156 bands: one at ~ 1430 cm⁻¹ attributable to the H-N-H bending vibrations and the other at 157 \sim 3300 cm⁻¹ originating from the N-H stretching vibrations, although the exact numbers 158 159 and positions of bands in each group may vary, depending on the bonding environments 160 and the local symmetry of NH_4^+ . For AAH, the bands ascribable to the H-N-H bending vibrations are observed at \sim 1379 and 1388 cm⁻¹. However, it is difficult to 161 162 unambiguously assign which bands arise from the N-H stretching vibrations due to overlap with the O-H stretching vibrations in the range of 2700 to 3700 cm⁻¹. 163 164 Compared to the band positions for the H-N-H bending vibrations for AAH, those for struvite are at much greater wavenumbers (\sim 1432 and 1468 cm⁻¹) (Figure 3). This 165

166	evident difference is related to the bonding environments around NH_4^+ in the two
167	compounds. In struvite, NH_4^+ is coordinated by six O atoms, with a wide range of the N-
168	O distances, from 2.800 to 3.498 Å, and an average N-O distance of 3.136 Å (Ferraris et
169	al. 1986), whereas all six O atoms coordinated to A in AAH fall between 2.720 and 3.095
170	Å, with an average A-O distance of 2.972 Å (Table 4) [see Yang and Sun (2004) for a
171	detailed discussion on the bonding differences around the A site between the hazenite-
172	and struvite-type structures]. While the large separation between the two H-N-H bending
173	modes $(1468 - 1432 = 36 \text{ cm}^{-1})$ for struvite is primarily a consequence of the marked
174	distortion of its A site, the greater wavenumbers of the two bending bands in struvite
175	(relative to those in AAH) may be explained by its longer average N-O distance. For a N-
176	HO bond, the longer N-O distance means a stronger N-H and weaker HO bonding,
177	which makes the H-N-H bending more difficult and the corresponding bands appear at
178	higher wavenumbers.

179

180 Implications of AAH

181 As shown in Table 1, AAH appears to form only in cultures with the Na-bearing solutions and pH below 10.0 (runs #1, 2, and 4). No AAH or hazenite was found in 182 183 experiments with the K-bearing solutions (runs 7-10). Hence, a Na-rich environment 184 seems to be essential for the formation of both AAH and hazenite (Yang and Sun 2004). 185 In addition, we observed both AAH and struvite in run #1, suggesting that they are 186 overlapping in formation environments. In nature, Mono Lake in California, which is 187 known for its unique biological and geochemical features, currently consists of a 188 hypersaline (84–92 g/L), alkaline (pH = 9.8) Na-CO₃-Cl-SO₄ brine (Yang et al. 2011 and 189 references therein). This environment is obviously analogous to that of our experimental 190 run #1 or #2. Hazenite was discovered on the south shore of Mono Lake (Yang et al. 191 2011), where no struvite has been documented. However, on the north shore of the lake, 192 especially on Paoha Island, where guano is relatively well preserved, struvite is quite

abundant (Cooper and Dunning 1969; Walker 1988). These places, therefore, could serveas candidates for the formation and discovery of AAH in nature.

195 Research on the precipitation of struvite from sewage has been an attractive 196 subject as it may offer a potential route for dephosphorization of wastewater from 197 industries and recovery of phosphates for fertilizers (e.g., Doyle and Parsons 2002; Shu et 198 al. 2006; Forrest et al. 2008; Machnicha et al. 2008; Muster et al. 2013). Given the strong 199 resemblances in both chemistry and structure between struvite and AAH, a better 200 understanding of the formation mechanism of AAH, especially in terms of the extent of 201 bacterial involvements, will unquestionably provide additional knowledge of 202 biomineralization of struvite-type phosphate materials and might lead to another route 203 for dephosphorization of wastewater. Furthermore, the struvite-type structure allows a complete substitution of AsO_4^{3+} for PO_4^{3+} (Weil 2008 and references therein). Thus, it 204 205 would be intriguing to explore whether the hazenite-type structure can also have the As-206 analogues, synthetic or natural. 207 208 Acknowledgements 209 This study was supported by the Science Foundation Arizona. The authors are 210 grateful to Piero Colaizzi from the Istituto Centrale per il Restauro e la Conservazione del Patrimonio Archivistico e Librario of Rome for his kind technical assistance with SEM-211 212 EDS. The careful and constructive reviews by Drs. W. Kolitsch and J. Pasteris are greatly

appreciated.

214

- 215
- 216

References Cited

217 Abu El-Fadl, A., Soltan, A.S., and Shaalan, N.M. (2006) Influence of cationic

218	substitution on lattice constants and optical characterization in solution grown
219	mixed crystals of potassium-ammonium zinc chloride. Crystal Research and
220	Technology, 41, 1013-1019.
221	Angoni, K., Popp, J., and Kiefe, W. (1998) A vibrational spectroscopy study of "urinary
222	Sand". Spectroscopy Letters, 31, 1771-1782.
223	Abbona, F., Calleri, M., and Ivaldi, G. (1984) Synthetic struvite, MgNH ₄ PO ₄ ·6H ₂ O:
224	Correct polarity and surface features of some complementary forms, Acta
225	Crystallographica, B40, 223-227.
226	Banks, E., Chianelli, R., and Korenstein, R. (1975) Crystal chemistry of struvite analogs
227	of the type MgMPO ₄ · $6H_2O$ (M ⁺ = K ⁺ , Rb ⁺ , Cs ⁺ , Tl ⁺ , NH ₄ ⁺). Inorganic Chemistry,
228	14, 1634-1639.
229	Ben Omar, N., Entrena, M., Gonzalez-Munoz, M.T., Arias, J.M., and Huetas, F. (1994)
230	The effects of pH and phosphate on the production of struvite by Myxococcus
231	Xanthus. Geomicrobiology Journal, 12, 81-90.
232	Ben Omar, N., Matinez-Canamero, M., Gonzalez-Munoz, M.T., Arias, J.M., and Huetas,
233	F. (1995) Myxocuccus xanthus' killed cells as inducers of struvite crystallization:
234	Its possible role in the biomineralization processes. Chemosphere, 30, 2387-2396.
235	Ben Omar, N., Gonzalez-Munoz, M.T., Penalver, J.M.A. (1998) Struvite crystallization
236	on Myxocuccus cells. Chemosphere, 36, 475-481.
237	Bogdanov, E.V., Vasil'ev, A.D., Flerov, I.N., and Laptash, N.M. (2011) Effects of cation
238	substitution in fluorine-oxygen molybdates $(NH_4)_{2-x}A_xMoO_2F_4$. Physics of the
239	Solid State, 53, 303-308.
240	Cahil, A, Soptrajanov, B., Najdoski, M, Lutz, H.D., Engelen, B., and Stefov, V. (2008)
241	Infrared and Raman spectra of magnesium ammonium phosphate hexahydrate
242	(struvite) and its isomorphous analogues. Part VI: FT-IR spectra of isomorphous
243	isolated species. NH_4^+ ions isolated in MKPO ₄ ·6H ₂ O (M = Mg, Ni) and PO ₄ ³⁻

4/9

244	ions isolated in MgNH ₄ AsO ₄ ·6H ₂ O. Journal of Molecular Structure, 876, 255-
245	259.
246	Chen, L., Shen, Y., Xie, A., Huang, F., Zhang, W., and Liu, S. (2010) Seed-mediated
247	synthesis of unusual struvite hierarchical superstructures using bacterium. Crystal
248	Growth & Design, 10, 2073-2082.
249	Cooper, J.F. and Dunning, G.E. (1969) Struvite found at Mono Lake. Mineral
250	Information Service, 22, 44-45.
251	Desmidt, E., Ghyselbrecht, K., Monballiu, A., Rabaey, K., Verstraete, W., and
252	Meesschaert, B. D. (2013) Factors influencing urease driven struvite precipitation.
253	Separation and Purification Technology, 110, 150-157.
254	Dickens, B. and Brown, W.E. (1972) The crystal structure of CaKAsO ₄ ·8H ₂ O. Acta
255	Crystallographica, B28, 3056-3065.
256	Doyle, J.D. and Parsons, S.A. (2002) Struvite formation, control, and recovery. Water
257	Research, 36, 3925-3940.
258	Ferraris, G., Fuess, H., and Joswig, W. (1986) Neutron diffraction study of
259	$Mg(NH_4)(PO_4).6H_2O$ (struvite) and survey of water molecules donating short
260	hydrogen bonds. Acta Crystallographica B, 42, 253-258.
261	Forrest, A.L., Fattah, K.P., Mavinic, D.S. and Koch, F.A. (2008) Optimizing struvite
262	production for phosphate recovery in WWTP. Journal of Environmental
263	Engineering, 134, 395-402.
264	Frost, R.L., Weier, M.L., Martens, W.N., Henry, D.A., and Mills, S.J. (2005) Raman
265	spectroscopy of newberyite, hannayite, and struvite. Spectroschimica Acta, A62,
266	181-188.
267	Gazulla, M.F., Rodrigo, M., Blasco, E., and Orduna, M. (2013) Nitrogen determination
268	by SEM-EDS and elemental analysis. X-ray Spectrometry, 42 (in press. DOI
269	10.1002/xrs.2490).
270	Gibson, R.I. (1974) Descriptive human pathological mineralogy. American Mineralogist,

59, 1177-1182.

272	Graeser, S., Postl, W., Bojar, HP., Berlepsch, P., Armbruster, T., Raber, T., Ettinger, K.,
273	Walter, F. (2008): Struvite-(K), KMgPO ₄ ·6H ₂ O, the potassium equivalent of
274	struvite – a new mineral. European Journal of Mineralogy, 20, 629-633.
275	Kamnev, A.A., Antonyuk, L.P., Colina, M., Chernyshev, A.V., Ignatov, V. (1999)
276	Investigation of a microbially produced structural modification of magnesium-
277 278	ammonium orthophosphate. Monatshefte für Chemie, 130, 1431-1442. Koleva, V.G. (2007) Vibrational behavior of the phosphates ions in dittmarite-type
279	compounds M'M"PO4·H ₂ O (M'=K ⁺ , NH ₄ ⁺ ; M"=Mn ²⁺ , Co ²⁺ , Ni ²⁺).
280	Spectrochimica Acta, A66, 413-418.
281	Lim, A.R. and Lee, M. (2013) Structural properties in mixed $LiK_{1-x}(NH_4)_xSO_4$ (x = 0,
282	0.06, and 1) crystals by nuclear magnetic resonance. Journal of Molecular
283	Structure, 1033, 113-120.
284	Liu, Z., Hu, J., Mao, H.K., and Hemley, R.J. (2002) High-pressure synchrotron x-ray
285	diffraction and infrared microspectroscopy: applications to dense hydrous phases.
286	Journal of Physics: Condensed Matter, 14, 10641–10646.
287	Machnicha, A., Grubel, K., and Suschka, J. (2008) Enhanced biological phosphorus
288	removal and recovery. Water Environment Research, 80, 617-623.
289	Mathew, M., and Schroeder, L.W. (1979) Crystal structure of a struvite analogue,
290	MgKPO ₄ ·6H ₂ O. Acta Crystallographica, B35, 11-13.
291	Mathew, M., Kingsbury, P., Takagi, S., and Brown, W.E. (1982) A new struvite-type
292	compound, magnesium sodium phosphate heptahydrate. Acta Crystallographica,
293	B38, 40-44.
294	Muster, T.H., Douglas, G.B., Sherman, N., Seeber, A., Wright, N., and Guezuekara, Y.
295	(2013) Towards effective phosphorus recycling from wastewater: Quantity and
296	quality. Chemosphere, 91, 676-684.
297	Sánchez-Román, M., Rivadeneyra, M.A., Vasconcelos, C., and McKenzie, J.A. (2007)
298	Biomineralization of carbonate and phosphate by moderately halophilic bacteria.
299	FEMS Microbiology Ecology, 61, 273-84.
300	Shannon, R.D. (1976) Revised effective ionic radii and systematic studies of interatomic

301	distances in balides and chalcogenides. Acta Crystallographica, A32, 751, 767
301	distances in nandes and chalcogendes. Acta Crystanographica, A52, 751–707.
302	Sheldrick, G. M. (2008) A short history of SHELX. Acta Crystallographica, A64, 112-
303	122.
304	Shin, H.K. (2011) Protonic conductivity and phase transition in $K_{1-x}(NH_4)_xHSO_4$. Solid
305	State Ionics, 189, 29-32.
306	Shu, L., Schneider, P., Jegatheesan, V., and Johnsom, J. (2006) An economic evaluation
307	of phosphorous recovery as struvite from digester supernatant. Bioresource
308	Technology, 97, 2211-2216.
309	Stefov, V., Soptrajanov, B., Spirovski, F., Kuzmanovski, I., Lutz, H.D., and Engelen, B.
310	(2004) Infrared and Raman spectra of magnesium ammonium phosphate
311	hexahydrite (struvite) and its isomorphous analogues. I. Spectra of protiated and
312	partially deuterated magnesium potassium phosphate hexahydrite. Journal of
313	Molecular Structure, 689, 1-10.
314	Stefov, V., Soptrajanov, B., Kuzmanovski, I., Lutz, H.D., and Engelen, B. (2005)
315	Infrared and Raman spectra of magnesium ammonium phosphate hexahydrite
316	(struvite) and its isomorphous analogues. III. Spectra of protiated and partially
317	deuterated magnesium ammonium phosphate hexahydrite. Journal of Molecular
318	Structure, 752, 60-67.
319	Walker, J. (1988) Paoha Island phosphates. Quarterly of San Bernardino County Museum
320	Association, 35, 46-47.
321	Weil, M. (2008) The struvite-type compounds $M[Mg(H_2O)_6](XO_4)$, where $M = Rb$, Tl
322	and $X = P$, As. Crystal Research and Technology, 43, 1286-1291.
323	Yang, H. and Sun, H.J. (2004) Crystal structure of a new phosphate compound,
324	Mg ₂ KNa(PO ₄) ₂ ·14H ₂ O. Journal of Solid State Chemistry, 177, 2991-2997.
325	Yang, H., Sun, H. J., and Downs, R.T. (2011) Hazenite, KNaMg ₂ (PO ₄) ₂ ·14H ₂ O, a new
326	biologically related phosphate mineral, from Mono Lake, California, U.S.A.
327	American Mineralogist, 96, 675-681.
328	
329	
330	

331	
332	
333	
334	
335	List of Tables
336	
337	Table 1. Agar media prepared for the biomineralization experiments.
338	
339 340	Table 2. Crystallographic data and refinement results for the ammonium-analogue of hazenite.
341	
342	Table 3. Coordinates and displacement parameters of atoms for the ammonium-analogue
343	of hazenite.
344	
345	Table 4. Selected non-hydrogen bond distances (Å) in hazenite and its ammonium-
346	analogue.
347	
348	Table 5. Comparison of unit-cell data for hazenite vs. its ammonium-analogue and
349	struvite vs. struvite-K.
350	
351	
352	
353	
354	
355	
356	List of Figure Captions
337 250	Figure 1 Crystels of the emmonium englosus of herenite (a) Crystels taken from growth
338 250	Figure 1. Crystais of the annionium-analogue of nazenite. (a) Crystais taken from growth
339	and (d) are back sectored electron images of three colonies on the top of Figure
300	1a (in order from left to right)
362	ra (in order nom ien to right).
363	Figure 2 Crystal structure of the ammonium analogue of hazenite. Tetrahedra – PO^{3}
364	arouns and octahedra – $Mg(H_2\Omega)_c$. The largest medium and smallest spheres
365	represent $A (- NH_4 + K)$ Ow9 (the H ₂ O molecule bonded to Na only) and Na
366	respectively. See the text for the definition of layers A B and C
367	respectively. See the text for the definition of hayers <i>H</i> , <i>D</i> , and <i>C</i> .
368	Figure 3. Infrared spectrum of the ammonium-analogue of hazenite, along with that of
369	struvite taken from the RRUFF project (rruff.info/R050511) for comparison
370	
371	Figure 4. Raman spectrum of the ammonium-analogue of hazenite. along with that of
372	hazenite taken from the RRUFF project (rruff.info/R100029) for comparison. The
373	spectra are shown with vertical offset for more clarity.

AAH Crystals







Mag = 147 X File Name = Na2CO3 5% new 39.tif Chamber = 31 Pa Signal A = CZ BSD Brightness = 33.6 % | Probe = 253 pA EHT = 20.00 kV Contrast = 78.3 % Spot Size = 442 WD = 12.5 mm











Run #	Supplemented salt	Concentration	Final agar pH at 25°C	Products identified
1	Na ₂ CO ₃	2%	9.5(3)	ammonium hazenite, struvite
2	Na ₂ CO ₃	5%	9.9(2)	ammonium hazenite
3	Na ₂ CO ₃	10%	10.0(2)	No visible crystals
4	NaCl	2%	6.9(1)	ammonium hazenite (very few)
5	NaCl	5%	6.9(4)	halite
6	NaCl	10%	6.7(3)	halite
7	KCl	5%	7.0(2)	sylvite
8	KCl	15%	6.6(1)	sylvite
9	K_2CO_3	5%	9.9(5)	struvite-K
10	K_2CO_3	15%	10.5(6)	kalincinite

Table 1. Agar media prepared for the biomineralization experiments.

Ideal structural formula	$(NH_4)NaMg_2(PO_4)_2 \cdot 14H_2O$
Space group	<i>Pmnb</i> (No. 62)
<i>a</i> (Å)	6.9661(6)
<i>b</i> (Å)	25.236(3)
c(Å)	11.2919(11)
$V(\text{\AA}^3)$	1985.0(3)
Ζ	4
$\rho_{\rm calc} ({\rm g/cm}^3)$	1.795
λ (Å)	0.71069
$\mu (\text{mm}^{-1})$	0.445
θ range for data collection	1.56 to 28.28
No. of reflections collected	21864
No. of independent reflections	2646
No. of reflections with $I > 2\sigma(I)$	1962
No. of parameters refined	198
R(int)	0.068
Final R factors $[I > 2\sigma(I)]$	$R_1 = 0.042, wR_2 = 0.091$
Final R factors (all data)	$R_1 = 0.066, wR_2 = 0.099$
Goodness-of-fit	1.09
R(int) Final R factors $[I > 2\sigma(I)]$ Final R factors (all data) Goodness-of-fit	$0.068 R_1 = 0.042, wR_2 = 0.091 R_1 = 0.066, wR_2 = 0.099 1.09$

Table 2. Summary of crystallographic data and refinement results for ammonium analogue of hazenite

Atom	x	У	Z	Uiso	U11	U22	U33	U23	U13	U12
 А	3⁄4	0.22146(6)	0.4983(1)	0.0386(6)	0.0505(11)	0.0295(9)	0.0356(10)	-0.0080(6)	0	0
Na	3/4	0.00057(7)	0.9266(2)	0.0367(6)	0.0280(10)	0.0420(11)	0.0401(11)	0.0115(8)	0	0
Mg1	3/4	0.95466(4)	0.6394(1)	0.0143(4)	0.0139(6)	0.0142(6)	0.0149(6)	0.0021(4)	0	0
Mg2	3⁄4	0.22385(4)	0.8370(1)	0.0160(4)	0.0185(6)	0.0152(6)	0.0141(6)	0.0007(4)	0	0
P1	1/4	0.12865(3)	0.7211(1)	0.0149(2)	0.0157(4)	0.0152(4)	0.0139(4)	-0.0009(3)	0	0
P2	3/4	0.12589(3)	0.2172(1)	0.0142(2)	0.0132(4)	0.0147(4)	0.0145(4)	0.0023(3)	0	0
01	1/4	0.13574(10)	0.8572(2)	0.0215(5)	0.0230(12)	0.0289(13)	0.0127(12)	-0.0019(10)	0	0
02	1/4	0.18372(9)	0.6614(2)	0.0222(5)	0.0269(13)	0.0179(11)	0.0218(12)	0.0019(10)	0	0
03	0.4317(2)	0.09817(6)	0.6845(1)	0.0201(4)	0.0172(8)	0.0206(8)	0.0224(9)	-0.0029(6)	0.0002(7)	0.0024(7)
04	3/4	0.06616(9)	0.1907(2)	0.0241(6)	0.0229(13)	0.0153(12)	0.0341(14)	-0.0001(10)	0	0
05	3/4	0.13598(10)	0.3515(2)	0.0219(5)	0.0224(12)	0.0281(13)	0.0151(12)	0.0007(10)	0	0
06	0.9311(2)	0.15090(6)	0.1616(1)	0.0192(4)	0.0162(8)	0.0216(8)	0.0198(8)	0.0063(7)	0.0019(7)	-0.0014(7)
OW7	3/4	0.00987(12)	0.5072(3)	0.0292(7)	0.0207(14)	0.0318(16)	0.0352(16)	0.0180(12)	0	0
OW8	3/4	0.89480(13)	0.7615(3)	0.0366(8)	0.0145(13)	0.0469(18)	0.0484(19)	0.0336(14)	0	0
OW9	0.5359(3)	0.91630(8)	0.5449(2)	0.0253(4)	0.0237(10)	0.0349(11)	0.0173(9)	-0.0010(7)	0.0028(8)	-0.0107(8)
OW10	0.9586(3)	0.99535(7)	0.7373(2)	0.0233(4)	0.0203(9)	0.0194(9)	0.0302(10)	0.0001(7)	-0.0064(8)	-0.0016(8)
OW11	0.5341(3)	0.25019(8)	0.7262(2)	0.0292(4)	0.0306(10)	0.0182(9)	0.0387(11)	0.0070(9)	-0.0124(9)	0.0004(9)
OW12	0.9644(3)	0.19572(7)	0.9484(2)	0.0238(4)	0.0273(10)	0.0267(10)	0.0174(9)	0.0022(7)	0.0000(8)	0.0048(8)
OW13	3/4	0.15560(11)	0.7295(3)	0.0281(6)	0.0171(13)	0.0255(14)	0.0417(16)	-0.0129(12)	0	0
OW14	3/4	0.29694(11)	0.9275(3)	0.0376(8)	0.0681(22)	0.0171(14)	0.0276(16)	-0.0019(12)	0	0
OW15	0.0114(3)	0.05941(8)	0.9812(2)	0.0329(5)	0.0421(13)	0.0272(11)	0.0293(11)	-0.0019(8)	0.0093(10)	-0.0003(9)
H11	0.834(4)	0.026(1)	0.487(3)							
H21	0.843(4)	0.880(1)	0.789(3)							
H31	0.453(5)	0.899(1)	0.573(3)							

Table 3. Coordinates and displacement parameters of atoms for ammonium analogue of hazenite

H32	0.546(5)	0.909(1)	0.473(3)
H41	0.066(5)	0.976(1)	0.757(3)
H42	0.994(4)	0.025(1)	0.727(3)
H51	0.446(5)	0.228(1)	0.715(3)
H52	0.512(5)	0.277(1)	0.717(3)
H61	0.932(5)	0.182(1)	0.016(3)
H62	0.053(5)	0.174(1)	0.918(3)
H71	0.841(4)	0.136(1)	0.726(3)
H81	3⁄4	0.317(2)	0.896(5)
H82	3⁄4	0.300(2)	0.011(4)
H91	0.053(5)	0.081(1)	0.946(3)
H92	0.960(5)	0.076(1)	0.038(3)

Note: $A = 0.78(NH_4) + 0.22K$; The Uiso parameters for all H atoms were fixed at 0.04.

hazenite		NH₄-analogue of Hazenite		
K-O5	2.672(3)	A-O5	2.720(3)	
-OW11 (x2)	2.980(2)	-OW11 (x2)	3.068(2)	
-OW12 (x2)	2.908(2)	-OW12 (x2)	2.940(2)	
-OW13	3.064(3)	-OW13	3.095(3)	
Avg.	2.919	Avg.	2.972	
Na-OW10 (x2)	2.563(3)	Na-OW10 (x2)	2.589(3)	
-OW15 (x2)	2.425(3)	-OW15 (x2)	2.429(2)	
-OW15 (x2)	2.472(3)	-OW15 (x2)	2.478(2)	
Avg.	2.487	Avg.	2.498	
Mg1-OW7	2.032(3)	Mg1-OW7	2.042(3)	
-OW8	2.040(3)	-OW8	2.046(3)	
-OW9 (x2)	2.066(2)	-OW9 (x2)	2.074(2)	
-OW10 (x2)	2.083(2)	-OW10 (x2)	2.095(2)	
Avg.	2.062	Avg.	2.071	
Mg2-OW11 (x2)	2.060(2)	Mg2-OW11 (x2)	2.066(2)	
-OW12 (x2)	2.074(2)	-OW12 (x2)	2.077(2)	
-OW13	2.093(3)	-OW13	2.107(3)	
-OW14	2.109(3)	-OW14	2.109(3)	
Avg.	2.078	Avg.	2.084	
P1-O1	1.544(3)	P1-O1	1.548(3)	
-O2	1.540(3)	-O2	1.544(3)	
-O3 (x2)	1.530(2)	-O3 (x2)	1.538(2)	
Avg.	1.536	Avg.	1.542	
P2-O4	1.535(3)	P2-O4	1.537(3)	
-O5	1.534(3)	-O5	1.538(3)	
-O6 (x2)	1.543(2)	-O6 (x2)	1.544(2)	
Avg.	1.539	Avg.	1.541	

Table 4. Selected non-hydrogen bond distances (Å) in hazenite and its ammonium analogue

Name	Locality	a (Å)	<i>b</i> (Å)	c (Å)	$V(\text{\AA}^3)$	References
Hazenite Hazenite NH ₄ -Hazenite	Synthetic Mono Lake Run #1 Pup #2	6.9316(5) 6.9349(4) 6.962(1) 6.9661(6)	25.1754(18) 25.1737(15) 25.223(7) 25.226(3)	11.2189(10) 11.2195(8) 11.293(2) 11.292(1)	1957.8(3) 1958.7(2) 1983(1) 1985.0(3)	Yang and Sun (2004) Yang et al. (2011) This study
NH ₄ -Hazenne	Kull #2	6.055(1)	23.230(3)	11.292(1)	1985.0(5)	Exercise at al. (1086)
Struvite	Synthetic	6.966(1) 6.966(2)	6.142(1) 6.142(1)	11.217(2) 11.217(2)	479.2(2) 479.9(1)	Abbona et al. (1986) This study
Struvite-K	Synthetic	6.873(2) 6.892(2)	6.145(1) 6.160(2) 6.166(2)	11.226(4) 11.087(3) 11.139(4)	479.9(4) 469.4(3) 473.4(3)	Mathew (1979) Graeser et al. (2008)
Struvite-K	Run #9	6.8741(4)	6.1481(4)	11.1094(6)	469.51(4)	This study

Table 5. Comparison of unit-cell data for hazenite vs. NH₄-analogue of hazenite and struvite vs. struvite-K.