1	The Crystal Structure and Chemistry of Natural Giniite and
2	Implications for Mars
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12 13	REVISION 1 Abstract
14	Investigations of planetary processes using phosphate minerals often focus on igneous,
15	recrystallized, or potentially metasomatized minerals, likely as a result of the minerals commonly
16	available for study within meteorites and lunar samples. However, Mars is a relatively
17	phosphorus-rich planet and possesses abundant evidence of past aqueous surface interactions.
18	Therefore, secondary phosphate phases may be important on the Martian surface. Giniite
19	$[Fe^{2+}Fe_4^{3+}(PO_4)_4(OH)_2 \cdot 2H_2O]$ is a secondary phosphate mineral that has been suggested as a
20	potentially significant phase at locations in Gusev Crater and Meridiani Planum on Mars.
21	Although relatively rare as a natural mineral on Earth, giniite has gained attention as an
22	important mineral in industry and technology, especially the lithium battery industry, and the
23	ferrian version of the mineral is often synthesized. This suggests giniite may be important as an
24	in-situ resource utilization (ISRU) target for future extended human missions to Mars. Despite
25	this, there are few data available on the natural mineral and the last characterization of the
26	structure was over 40 years ago. There has also been confusion in the literature as to whether
27	giniite is orthorhombic or monoclinic. In this work we revisit and document the chemistry and
28	crystal structure of natural giniite from the type locality at the Sandamab pegmatite in Namibia

29	using updated techniques. Our results refine and update what was previously known regarding
30	the structure and chemistry of giniite and support the potential of the mineral as a possible
31	Martian scientific and resource target for further study to aid future missions.
32	
33	Keywords
34	Giniite, Fe-phosphate, ferrous giniite, ferric giniite, ferrian, phosphate,
35	hydroxyphosphate, Martian habitability, Mars, ISRU, XRD,
36	
37	Introduction
38	The study of phosphorus minerals yields insight into planetary interior and surface
39	processes. For instance, primary or igneous phosphate minerals have been used to investigate
40	volatile abundances in the interiors of Earth, Mars, the Moon, and asteroidal bodies (e.g.
41	McCubbin et al., 2011; Patiño Douce et al., 2011; Filiberto et al., 2016; Jones et al., 2014;
42	McCubbin et al., 2014), as potential indicators of oxygen fugacity during late stage magma
43	crystallization (Shearer et al., 2015), and even as recorders of past aqueous surface environments
44	on Mars (Mojzsis and Arrhenius, 1998; Hurowitz et al., 2006; Ming et al., 2006; Adcock and
45	Hausrath, 2015). Phosphorus, as phosphate or a more reduced species, is also an essential
46	nutrient for all known life, and considered important in prebiotic reactions that led to life on
47	Earth (Wald, 1964; Westheimer, 1987; Powner et al., 2009; Pasek and Kee, 2011; Benner and
48	Kim, 2015; Burcar et al., 2016). Consequently, P-bearing minerals have important implications
49	for past and present habitability and the potential for life on other planetary bodies (Weckwerth
50	and Schidlowski, 1995; Mojzsis et al., 1996; Yang et al., 2011; Adcock et al., 2013; Adcock and
51	Hausrath, 2015).

52	Investigations of planetary processes using phosphate minerals often focus on igneous,
53	recrystallized, or potentially metasomatized minerals (e.g. Brearley and Jones, 1998; Jones et al.,
54	2014; McCubbin and Jones, 2015; Adcock et al., 2017). For extraterrestrial studies, this focus is
55	likely the result of the minerals commonly available for study within meteorites and lunar
56	samples. Though various secondary phosphate minerals are present in some meteorites (Dyar et
57	al., 2014), the most common phosphate minerals in most meteorites and lunar samples are
58	merrillite [Ca ₉ NaMg(PO ₄) ₇] and apatite [Ca ₅ (PO ₄) ₃ (F,Cl,OH)], often of igneous origin (Shearer
59	et al., 2006; McCubbin et al., 2014; Adcock and Hausrath, 2015). This is especially the case for
60	Martian meteorites (McSween and Treiman, 1998).
61	However, Mars is a relatively phosphorus-rich planet (Wänke and Dreibus, 1988; Taylor,
62	2013) and analyses from the surface of Mars indicate relatively high P-concentrations in soils,
63	rocks, and dust compared to Earth (Goetz et al., 2005; Gellert et al., 2006; Yang and Post, 2011;
64	Rampe et al., 2017; Yen et al., 2017; Rampe et al., 2020). Mars also possesses abundant
65	evidence of past aqueous surface interactions (Carr and Head III, 2003; Hurowitz et al., 2006;
66	Grotzinger et al., 2014; Adcock and Hausrath, 2015; Rampe et al., 2016; McCollom et al., 2018).
67	Reactive transport modeling of measured rock profiles at Gusev Crater indicate dissolution of
68	primary phosphate minerals, the product of which would be secondary phosphates (Adcock and
69	Hausrath, 2015). Thermodynamic modeling by Berger et al., (2016) indicates the formation of
70	the secondary phosphate mineral strengite (FePO ₄ \cdot 2H ₂ O) at Gale Crater, Mars. Recent data from
71	the Alpha Particle X-ray Spectrometer (APXS) on board Curiosity at Gale Crater indicate
72	enrichments of P and Mn in nodules, veins, and other surface features there (Berger et al., 2021).
73	ChemMin diffraction data from Gale Crater also show the potential presence of secondary
74	flourapatite (Rampe et al., 2017) and a secondary manganese-bearing phosphate of the jahnsite-

75	whiteite group (Treiman et al., 2021). These models and observations suggest aqueously altered
76	or precipitated secondary phosphate minerals may not be uncommon phases on the Martian
77	surface and thus warrant more in-depth consideration.
78	Among the potential secondary minerals that may occur on Mars is giniite
79	$[Fe^{2+}Fe^{3+}_4(PO_4)_4(OH)_2 \cdot 2H_2O]$. Although rare on Earth, giniite has gained attention in industry,
80	including as a potential component in Li-ion battery production (Hong et al., 2003; Whittingham,
81	2004; Lv et al., 2017), wastewater processing (Duan et al., 2013; Han et al., 2017; Priambodo et
82	al., 2017), biomedical materials, and as a deprotonation catalyst (Chen et al., 2014; Nedkov et
83	al., 2016).
84	In comparison to Earth, Mars has notably higher Fe and P content, and Hausrath et al.,
85	(2013) first suggested that giniite may be a significant secondary phase on that planet based on
86	terrestrial hydrothermal experiments that produced giniite, and Mössbauer data acquired from
87	Gusev Crater on Mars. In later work, alteration experiments using P-enriched basalts as analogs
88	of rocks at Meridiani Planum also produced giniite, suggesting the mineral may likewise be of
89	importance at that location on Mars (McCollom et al., 2018). The confirmation of giniite on
90	Mars would be scientifically important. Like primary minerals, secondary minerals are products
91	of their formation environments, and are therefore potentially useful indicators of past Martian
92	environmental conditions (e.g. Klingelhöfer et al., 2004; Wiseman et al., 2008). Minerals like
93	giniite also have implications for the availability of bio-essential phosphate in these past
94	environments, and thus are astrobiologically relevant. Giniite may further represent a potential
95	resource for extended human missions to Mars. For instance, as a hydrated mineral, giniite
96	contains approximately 48 liters of H ₂ O per metric ton (not including the OH ⁻ component) that
97	can be driven off between 175° and 300° C (Jambor and Dutrizac, 1988; Rouzies et al., 1994;

98	Gonçalve et al., 2017; Liu et al., 2017). McCollom et al., (2018) suggest as much as 8% of the								
99	Fe in the Burns Formation at Meridiani Planum on Mars may be held in giniite, with another								
100	29% in jarosite (a hydrated Fe-sulfate holding \sim 10% H ₂ O) (Morris et al., 2006). Therefore,								
101	giniite, along with other hydroxides in Martian soils or rocks, could be a useful source of water.								
102	The mineral could also be processed into fertilizer, and oxidation of the Fe^{2+} component in								
103	minerals has been shown to produce H ₂ or CH ₄ , potential fuel components (e.g. Miller et al.,								
104	2017; Adcock et al., 2021). In Situ Resource Utilization (ISRU) is the practice of using materials								
105	or energy sources derived at space destinations (such as Mars) to replace or supplement								
106	resources that would otherwise have to be transported from Earth (Sanders and Larson, 2011;								
107	Sridhar et al., 2000; Starr and Muscatello, 2020). The above qualities, including the potential								
108	availability, suggest that giniite may be a good subject for further investigation as a possible								
109	Martian ISRU target.								
110	Despite the potential importance of giniite both terrestrially and on Mars, most of the								
111	research and data currently available focus upon the synthetic, generally ferrian, analog of the								
112	mineral, rather than the natural, typically more reduced form. To our knowledge, there are few								

data on the natural mineral and the only characterization was over 40 years ago by Keller (1980a;

114 1980b), who originally documented the mineral with X-ray diffraction, electron microprobe and

thermogravimetric analysis (TGA). The original structure was reported as orthorhombic

116 (Fleischer et al., 1980; Keller, 1980a), but following work suggested this assessment was the

result of twinning and the structure was revised to monoclinic (Keller, 1980b). Despite this,

giniite is still sometimes reported as orthorhombic (e.g. Liu et al., 2017; Zhang et al., 2013). In

this work we use modern single-crystal X-ray diffraction with an area detector on natural ginite

120 from the type locality, to reassess the structure in the correct monoclinic system. Additionally, Raman

spectroscopy and electron microprobe techniques are employed to revisit and document thechemistry.

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Background

125 Minerals exhibit characteristics of their formation environments. In the case of secondary phosphate minerals, these characteristics may include temperature, pH, chemistry, and oxidation 126 127 state of any interacting fluids (Moore, 1973; Hawthorne, 1998; Dill et al., 2008). Vivianite $[Fe_3^{2+}(PO_4)_2 \cdot 8H_2O]$ for instance, indicates a near neutral pH, anoxic, reducing, low temperature, 128 high water:rock ratio aqueous formation environment (Hawthorne, 1998; McGowan and 129 Prangnell, 2006). Brushite $[Ca(HPO_4) \cdot 2H_2O]$ and monetite $[Ca(HPO_4)]$ are indicators of acidic, 130 131 and potentially biogenic formation environments on Earth (Dumitras et al., 2004; Dosen and Giese, 2011; Frost and Palmer, 2011; Frost et al., 2013b). Although not confirmed on Mars, 132 brushite has been suggested as a possible acidic weathering product on that planet (Hurowitz et 133 134 al., 2006; Ming et al., 2006). Secondary Al- and Fe-phosphates are also probable components in the P-enriched amorphous fraction of soils documented at Gale Crater by the Mars Science 135 Laboratory *Curiosity*, again with implications for past aqueous Martian weathering environments 136 (Morris et al., 2013; Tu et al., 2014; Vaniman et al., 2014; Rampe et al., 2016). All of these 137 secondary phosphate minerals have the potential to inform us about past and present formation 138 environments, including those on Mars. 139 Giniite is no exception and its presence and paragenesis can provide detailed insights 140 141 regarding its formation environment. On Earth, the mineral typically occurs within pegmatites or iron-bearing ore bodies (Keller, 1980a; Jambor and Dutrizac, 1988; Nunes et al., 2009). In 142 pegmatite settings, the mineral is a product of hydrothermal alteration of primary triphylite 143

[LiFePO₄]. Keller (1980a) identified it as part of a paragenetic or age sequence of triphylite \rightarrow 144 145 hureaulite \rightarrow [an unidentified dark green mineral + giniite] \rightarrow tavorite \rightarrow leukophosphite. Keller (1980a) speculated that giniite took the place of barbosalite $[Fe^{2+}Fe^{3+}_{2}(PO_{4})_{2}(OH)_{2}]$ in a similar sequence 146 147 at other pegmatites in the region (e.g. Keller and Knorring, 1989), suggesting giniite is the 148 product of mid- to late-stage hydrothermal alteration (<250 °C) (Hawthorne, 1998). In metal ore bodies the genetic history of giniite is less clear. Phosphorus is a common 149 150 impurity in iron-bearing metal ore bodies (Cheng et al., 1999; Delvasto et al., 2008; Ofoegbu, 151 2019), and at the Silver Coin mine in Nevada, USA, phosphate minerals, including giniite, are products of acidic hydrothermal precipitation associated with the ore body (Adams et al., 2015). 152 However, at mines near Saalfeld, Germany, giniite appears to occur as a low temperature 153 weathering product of the ore body associated with a hydrous mineral gel formed in open natural 154 caves (Ullrich, 2017). 155

Recent industrial, technologic, and scientific interests in the mineral (noted above) have
led to several synthesis methods and studies. Most of these synthesis methods produce a fully

158 ferric form of the mineral we refer to as ferrian giniite $[Fe^{3+}_{5}(PO_{4})_{4}(OH)_{3} \cdot H_{2}O]$ (e.g. Jambor and

159 Dutrizac, 1988; Rouzies et al., 1994; Frost et al., 2007; Duan et al., 2013; Zhang et al., 2013;

160 Nedkov et al., 2016; Han et al., 2017; Liu et al., 2017). However, mixed Fe valence giniite more

similar to natural giniite has also been synthesized with ferrous/ferric content up to $Fe^{2+}_{1.7}/Fe^{3+}_{3.3}$

162 (Rouzies et al., 1994). To account for different Fe^{2+}/Fe^{3+} ratios, charge balance is maintained by

163 changes in the OH^{-/} H₂O ratio through hydrating/dehydrating OH⁻ sites (Keller, 1980a; Rouzies

tet al., 1994) and can be expressed as

165
$$[Fe^{2+}_{x} Fe^{3+}_{(5-x)}(PO_4)_4(OH)_{(3-x)} \cdot (1+x)H_2O].$$

166 It is of note, however, that in the absence of measured data, giniite is often reported with $2 H_2O$

units regardless of Fe valence or OH⁻ content (e.g. Roncal-Herrero et al., 2009; Zhang et al.,

168 2015; Priambodo et al., 2017). In addition, x=1.7 is the highest Fe²⁺ content currently observed.

169 Corbin et al., (1986) synthesized a fully ferrous phase $[Fe_5^{2+} P_4O_{20}H_{10}, \text{ or } Fe_5^{2+} H_2(PO_4)_4 \cdot 4H_2O]$

170 (i.e., x=5) referred to as giniite (e.g. Dyar et al., 2014; Gonça ves et al., 2017). However, a fully

171 ferrous giniite structure is not stable, and instead this chemistry adopts the hureaulite structure

172 $[Mn_5(PO_3OH)_2(PO_4)_2 \cdot 4H_2O]$ (Corbin et al., 1986).

Although the rare natural occurrence of giniite on Earth somewhat limits paragenetic and 173 174 minerogenetic data, the conditions of giniite synthesis can yield further insight into the environments where the mineral may form. Based on synthesis methods, giniite forms under 175 relatively high water:rock ratios (>10:1) and formation temperatures from 25° to 250°C (>150°C 176 appearing optimum for highest yields) (Rouzies et al., 1994; Roncal-Herrero et al., 2009; 177 Hausrath et al., 2013; Nedkov et al., 2016; Gonca ves et al., 2017). Highly acidic conditions 178 appear to be optimum (pH 0.6 to 2) (Jambor and Dutrizac, 1988; Hausrath et al., 2013). 179 However, in the presence of other monovalent cations like Na and K, giniite can form at as high 180 as pH 6 (Gonça ves et al., 2017). While these conditions represent a range of potential settings 181 for giniite formation, the mineral can exhibit variable chemistry and morphology while 182 maintaining the giniite structure (Jambor and Dutrizac, 1988; Zhang et al., 2013) and these 183 184 aspects may reflect specific details of formation conditions. For instance, Roncal-Herrero (2009) found that crystal morphology of giniite synthesized at 150°C was spheroidal while giniite 185 synthesized at 200°C had a bi-pyramidal morphology. Other studies have noted morphology 186 dependencies based on solution Fe and P concentrations (dendritic, spherical, and octahedral) 187 (Liu et al., 2017), and the presence of different organic compounds in synthesis solutions 188 (spheres and star-like) (Duan et al., 2013; Han et al., 2017). Tubular morphologies have been 189

reported as products of biogenic formation at 25°C (Nedkov et al., 2016). The pH and presence
of Li, Na, or K in solution also appear to act on morphology (asterisk- or flower-like, and
dendritic morphologies) (Gonça ves et al., 2017). It is not clear if the Li, Na, or K substitute into
giniite in that study, however SO₄²⁻ substitution for PO₄³⁻ has been noted (Jambor and Dutrizac,
1988) and divalent cations including Mn and Mg are known to incorporate into the mineral
(Keller, 1980a). These conditions, though potentially broader, are not inconsistent with what is
known about natural giniite formation environments.

197 Although giniite has not been definitively identified on Mars, several lines of evidence suggest that it may be present. Experiments that mimic Martian conditions of acidic solutions 198 placed in contact with a mixture of fluorapatite, olivine, and basaltic glass at 150°C by Hausrath 199 et al. (2013) produced ferrian giniite [Fe₅(PO₄)₄(OH)₃·2H₂O]. Follow-up Mössbauer 200 measurements of the ferrian giniite alteration products were consistent with analyses performed 201 by the Mars Exploration Rover Spirit on Paso Robles soil at Gusev Crater. Hausrath et al. (2013) 202 could not confirm the phase on Mars based on Mössbauer alone, as measurements were also 203 204 consistent with ferric sulfate phases (Hausrath et al., 2013; Dyar et al., 2014). However, $Fe_2O_3 +$ FeO concentrations in PasoRobles and PasoLight1 soils on Mars are too high to be accounted for 205 solely by sulfate phases and indicate acid fluid transport in high fluid:rock ratios to the location 206 207 of the Paso Robles soils, general conditions shown to precipitate giniite. McCollom et al. (2018) also suggested giniite may help explain phosphate immobility in the Burns formation at 208 Endurance Crater, as well as the potentially high P measured in alteration halos around fractures 209 previously documented in the Stimson and Murray formations at Gale Crater suggesting the 210 presence of phosphate-rich fluids (Yen et al., 2017). The P-containing fluids that formed these 211 alteration haloes have also been quantitatively modeled (Hausrath et al., 2018). Rampe et al. 212

213	(2017), suggest acidic phosphate-rich fluids in the lower Murray at Gale Crater. This indicates
214	conditions suitable for the natural formation of giniite, especially considering giniite genesis at
215	the Silver Coin Mine, Nevada. Therefore, formation of giniite or other ferric phosphate phases
216	may have occurred at multiple locations on Mars. Updating and refining our fundamental
217	knowledge of these minerals, and investigating them more deeply, will help us to detect and
218	identify them on the Martian surface.
219	
220	Experimental
221	Materials
222	A natural sample of giniite was acquired from the Sandamab pegmatite, Namibia by the
223	RRUFF project (Lafuente et al., 2016). Sandamab (sometimes spelt Sandamap) pegmatite is the
224	type locality for the mineral as originally identified by Keller (1980a; 1980b). The sample was a
225	black fragment within a more massive sample of associated triphylite, hureaulite, yellow-
226	greenish tavorite, and black heterosite and was preliminarily identified as giniite based on color
227	and rough habit. RRUFF reference number is R060765 (Supplementary Figure S1).
228	
229	Analytical Methods
230	The broad scan Raman spectrum of giniite was collected from a randomly oriented
231	crystal with a Thermo Almega microRaman system using a solid-state laser with a frequency of
232	532 nm at 150 mW and a thermoelectrically cooled CCD detector. The laser is partially polarized
233	with 4 cm ^{-1} resolution and a spot size of 1 μ m.
234	Chemical analyses were carried out on a Cameca SX-100 electron microprobe (EMP).
235	The giniite sample was mounted in epoxy, polished and carbon coated. Samples were analyzed

236 using wavelength-dispersive spectroscopy (WDS) X-ray analysis. Analysis conditions were 15 237 keV and 10 nA using a 2 μ m beam. Standards and configuration package are detailed in Supplementary Table S1. Multiple analyses (n=15, Figure 1) of the giniite crystal were taken and 238 239 averaged. EMP does not detect OH-, H₂O, or Fe valence. To estimate OH⁻ and H₂O, ideal stoichiometry as determined by Keller (1980a) of $Fe^{2+}Fe^{3+}_4(PO_4)_4(OH)_2 \cdot 2H_2O$ was used. To 240 estimate the Fe valence ratio, the EMP data were fit to the ideal formula with $Fe^{3+} = 4$ formula 241 242 units. Synthetic giniite is known to have variable Fe valence and this may be possible for natural 243 giniite as well. Therefore, the EMP data were further fit to a range of stoichiometries based on $Fe^{2+}_{x}Fe^{3+}_{(5-x)}(PO_{4})_{4}(OH)_{(3-x)}$ · (1+x)H₂O (Keller, 1980; Rouzies et al., 1994) by varying x to 244 determine the Fe^{3+} formula units and calculating the resulting Fe^{2+} formula units and the 245 estimated OH and H₂O. A "best fit" was determined based on total Fe and PO₄ formula units 246 being closest to the ideal of 5 and 4 respectively, in the same fit. 247 Single-crystal X-ray diffraction (SC-XRD) of giniite was carried out using MoKa 248 radiation on a Rigaku XtaLAB Synergy diffractometer and radiation at 50 kV and 1mA. All 249 250 reflections were indexed on the basis of a monoclinic unit-cell (Table 1, Supplementary Table S2). The systematic absences of reflections suggested the possible space group Pn, or P2/n. The 251 crystal structure was solved and refined using SHELX2018 (Sheldrick, 2015a; Sheldrick, 2015b) 252 253 based on space group P2/n, because it yielded better refinement statistics in terms of bond lengths and angles, atomic displacement parameters, and R factors. All H atoms were located 254 from the difference Fourier maps. The ideal chemistry was assumed during the refinements. The 255 positions of all atoms were refined with anisotropic displacement parameters, except those for 256 the H atoms, which were refined only with isotropic parameters. 257

259

Results

260 Structure

X-ray diffraction data collected in this study allowed calculation of a powder X-ray 261 262 diffraction pattern that is a very close match to published patterns of both a synthetic ferric 263 giniite (Frost et al., 2007) and Keller's (1980a) natural giniite sample (Figure 2). The close match suggests the study sample is indeed giniite. Final coordinates and displacement parameters of 264 265 atoms in giniite are listed in Supplementary Table S3, and selected bond distances in Table 2. 266 Calculated bond-valence sums using the parameters of Brese and O'Keeffe (1991) are given in Table 3. Crystallographic results of the single crystal data indicate giniite is monoclinic P2/n, a =267 10.3472(6), b = 5.1497(2), c = 14.2338(7) Å, $\beta = 111.175(6)^{\circ}$ and V = 707.24(7) Å³ (Table 1 and 268 Supplementary Table S2). Atomic positions in Supplementary Table S3 show three unique Fe 269 sites (here labeled Fe1, Fe2 and Fe3) and two unique PO₄ sites (P1 and P2), as well as O and H 270 positions. Determined Fe-O and P-O bond length values fall within expected ranges (Gagné and 271 Hawthorne, 2018; Kanowitz and Palenik, 1998) with average (Fe1-O) bond lengths (2.14 Å) 272 longer than the other two (2.01 Å) indicating that Fe1 contains Fe^{2+} while Fe2 and Fe3 contain 273 Fe^{3+} (Table 2). Bond valence calculations are consistent with this assignment and are generally as 274 expected (Table 3) with the potential exception of the O7 site which is ~0.45 deficient. 275 276 Chemistry Results of 15 EMP analyses (Figure 1 and Supplementary Table S4) were averaged into a 277

single analysis which appears in Table 4 along with Keller's (1980a) original EMP analysis. Low
analysis totals are the result of unaccounted for OH⁻ and molecular water which the technique
cannot directly detect. EMP can also not detect Fe valence state. In order to estimate wt% OH⁻,

H₂O, and Fe²⁺/Fe³⁺ for comparison to Keller (1980a), the analyses from this study and Keller

(1980a) were fit to the ideal giniite formula of $Fe^{2+}Fe^{3+}_4(PO_4)_4(OH)_2 \cdot 2H_2O$ with $Fe^{3+}=4.00$.

283 This fitting produced an average stoichiometry of

284 $(Fe^{2+}_{0.80}Mn_{0.11}Mg_{0.02})_{\Sigma=0.93}Fe^{3+}_{4.00}(PO_4)_{4.03}(OH)_{2.00} \cdot 2H_2O$ for this study and

285 $(Fe^{2+}_{0.67}Mn_{0.07}Mg_{0.13})_{\Sigma=0.87}Fe^{3+}_{4.00}(PO_4)_{4.05}(OH)_{2.00} \cdot 2H_2O$ for Keller (1980a) data. Incorporating

the estimates as wt.% into the EMP data improved the analysis totals for this study and Keller

287 (1980a) to ~97.15 and 91.83 wt% respectively (Table 4).

EMP data fit to a range of Fe valence ratios based on $[Fe^{2+}_{x} Fe^{3+}_{(5-x)}(PO_4)_4(OH)_{(3-x)}$.

289 $(1+x)H_2O$ (Keller, 1980; Rouzies et al., 1994), with x used to determine the Fe³⁺ valence (i.e.

290 Fe^{2+} was calculated), produced a "best fit" stoichiometry of

291 $(Fe^{2+}_{0.90}Mn_{0.11}Mg_{0.02})_{\Sigma=1.03}Fe^{3+}_{3.92}(PO_4)_{4.05}(OH)_{1.92} \cdot 2.08H_2O$ (Supplementary Tables S5 and S6).

This fit is based on x = 1.08 to determine Fe³⁺, however, the calculated Fe²⁺ is 1.03 formula units.

A broad scan Raman spectrum in Figure 3 is consistent with previously reported giniite 294 spectra. There is one non-equivalent hydroxyl group and one water molecule with two non-295 equivalent O-H pairs in giniite, consistent with the single sharp peak at 3324 cm⁻¹ that overlaps 296 with the broad peak at \sim 3250 cm⁻¹, typical of overlapping O-H stretching modes for OH and H₂O 297 (e.g. Frost et al., 2011; Kolesov, 2006; Weber et al., 2018). PO₄ symmetric stretching modes are 298 in the 800 - 1100 cm⁻¹ range with asymmetric v_3 stretching in the 1100 to 1200 cm⁻¹ range. In 299 phosphates, bands in the 400 - 650 cm⁻¹ range are generally representative of O-P-O angle 300 bending (both v_2 and v_4) (Frost et al., 2007; Hausrath and Tschauner, 2013). The sharp bands in 301 the 200 – 440 cm⁻¹ range are typical of Fe-O stretches (Aatig et al., 2016; Frost et al., 2007, 302 2013a,c). For a plot of raw data, see Supplementary Figure S2. 303

305

Discussion

306 Minerals like giniite are a challenge to chemically characterize. They contain both ferric 307 and ferrous iron as well as OH^{-} and H_2O . These components are often undetectable or 308 undistinguishable by common analytical techniques, including electron microprobe. As a result, 309 Keller (1980a) noted low EMP totals for giniite (Table 4) in that study. Using EMP and X-ray data, Keller was able to produce a generalized formula for giniite accounting for the OH/H₂O 310 which can be expressed as 311 $Fe^{2+x} Fe^{3+}_{(5-x)} (PO)_4 (OH)_{(3-x)} \cdot (1+x)(H_2O)$ 312 Thermo-gravimetric analysis allowed Keller (1980a) to estimate water content and thereby 313 conclude "x" in the expression to be 1 which produced an Fe^{3+} : $Fe^{2+}+Fe^{3+}$ formula unit ratio of 314 0.80 (i.e., Fe^{2+}/Fe^{3+}_4). Thus, $Fe^{2+}Fe^{3+}_4(PO)_4(OH)_2 \cdot 2H_2O$ was accepted as the ideal formula for 315 giniite by the International Mineralogical Association (IMA). 316 In the present study, Raman data, which are generally consistent with data from a 317 318 previously characterized synthetic ferric giniite sample (Frost et al., 2007), confirm OH/H₂O in the study sample. EMP analyses from this study, fit to the ideal formula by fixing $Fe^{3+} = 4$ 319 formula units, generate an Fe^{3+} : $Fe^{2+}+Fe^{3+}$ formula unit ratio of ~0.80 and calculated 320 stoichiometry even closer to the ideal formula than Keller's original work fit in the same way 321 (Supplementary Table S5). Like Keller (1980a), the original EMP analyses here produced low 322 totals. However, accounting for the Fe valence and OH/H₂O produces totals of ~97% (Table 4). 323 An additional stoichiometric best fit with the EMP data using the more generalized formula of 324 giniite $[Fe^{2+}x Fe^{3+}(5-x) (PO)_4(OH)_{(3-x)} \cdot (1+x)(H_2O)]$ also produces an Fe^{3+} : $Fe^{2+}+Fe^{3+}$ formula unit 325 ratio of ~0.80 (i.e., $Fe^{2+}_{1,03}/Fe^{3+}_{3,92}$ or $x \sim 1$) (Supplementary Table S6). Thus, chemical data in 326 this study are consistent with Keller (1980a) and the ideal giniite formula. 327

328 It is of note, however, that in determining the ideal formula for giniite, Keller (1980a) 329 assumed the divalent site stoichiometry to be a fixed integer. A number of studies have documented synthetic giniite with Fe^{3+} : $Fe^{2+}+Fe^{3+}$ ratios ranging from 1 (ferrian giniite) to as 330 low as 0.65 (e.g. Rouzies et al., 1994; Roncal-Herrero et al., 2009; Duan et al., 2013; Goncalve 331 et al., 2017; Priambodo et al., 2017). Consequently, Fe^{2+} is observed to range from 0 to 1.7 332 formula units while still maintaining the giniite structure. In fact, one of the interesting aspects of 333 giniite is the range of chemistry, and even morphology, that the mineral can possess while still 334 335 maintaining the giniite structure (e.g. Jambor and Dutrizac, 1988; Zhang et al., 2013; Gonca ves et al., 2017; Martins et al., 2020). Some of this range can be attributed to analytical uncertainty. 336 Data in this study, for instance, allow for variation in the Fe^{3+} : $Fe^{2+}+Fe^{3+}$ ratio of 0.81 to 0.76 337 (Fe²⁺ content of 1.0 to 1.2) before the summed Fe or PO₄ stoichiometry begin to deviate too far 338 from giniite. However, the observed range in the literature is too large to be explained by this 339 uncertainty alone. While these variations occur in synthetic forms of giniite, and this study 340 focuses on a natural sample as would be found on Mars, the synthesis conditions of at least some 341 342 of the giniite discussed here are similar to the known petrogenetic/minerogenetic conditions of natural giniite in both pegmatite and ore body settings. Therefore, though the data in this study 343 are consistent with the ideal Fe^{3+} : $Fe^{2+}+Fe^{3+}$ ratio of 0.80 inferred by Keller (1980a), this ratio 344 may only be nominally true in natural specimens. 345

Crystallographically, Keller (1980a) originally reported giniite as orthorhombic and this was how it was initially documented as a new mineral (Fleischer et al., 1980). Follow-up work by Keller (1980b) showed that the orthorhombic determination was a probable consequence of unrecognized twinning, and the natural mineral was actually monoclinic. However, synthetic giniite has still been reported as orthorhombic (e.g. Liu et al., 2017; Zhang et al., 2013), possibly

351	as a product of confusion from the literature. Single crystal data/refinement results in this study							
352	of natural giniite indicate our sample to be monoclinic, consistent with Keller (1980b).							
353	The giniite structure can be described as short chains (Fe1-Fe2-Fe1 trimers) of face-							
354	sharing irregular FeO ₆ octahedra oriented along [100], corner-linked by both 6-coordinated							
355	evenly spaced FeO ₆ octahedra associated with Fe3 arranged in columns along [010], and							
356	alternating (P1)O ₄ and (P2)O ₄ tetrahedra (Figure 4). In the context of this description the Fe1 site							
357	houses the ferrous iron component. The O atom of the H_2O molecule is bonded to ferric Fe3 and							
358	ferrous Fe1 atoms. The O atom in the OH group is bonded to all three Fe atoms and, with H, it is							
359	tetrahedrally coordinated (Figure 4). The structure shares some similarities with that of							
360	barbosalite $[Fe^{2+}Fe^{3+}_2(PO_4)_2(OH)_2]$ (Poienar et al., 2020; Redhammer et al., 2000) which also							
361	has Fe trimers joined to phosphate and an additional Fe octahedral site, although in that mineral							
362	the trimer valences are ordered Fe^{3+} , Fe^{2+} , Fe^{3+} , whereas in giniite the trimers are sequenced							
363	Fe^{2+} , Fe^{3+} , Fe^{2+} . These similarities are in-line with speculation by Keller (1980a) that giniite							
364	shares certain structural components with other Fe-hydroxy-phosphates and that in more							
365	generalized pegmatitic settings giniite may take the place of barbosalite in the evolution of							
366	secondary minerals associated with the decomposition of triphylite.							
367	Of further interest in this study is the "O7" site associated with the Fe1 and P2 sites. Fe-O							

and P-O bond lengths and bond valence calculations are generally as expected (Tables 2 & 3). However, bond valence sums for the O7 site are deficient (~1.5 e). Non-ideal bond valence sums can result for many reasons, including vacancies or undetected substitutions/components in the structure. In this study, the configuration of the O7 sites results in them being adjacent across an opening in the giniite structure (Figure 4, Supplementary Figure S3). The Fe1-O7-P2 bond angle is 127.33° and the O7-O7 distance is approximately 2.46 Å. It may be possible that H⁺ protons

374 occupy some of these "O7 voids" oscillating between O7 sites, and a shared H⁺ proton would 375 account for the valence value. Proton oscillation or "hopping" between sites has previously been indicated in the Fe-phosphate minerals barbosalite $[Fe^{2+}Fe^{3+}_{2}(PO_{4})_{2}(OH)_{2}]$ and ludlamite 376 377 $[(Fe,Mn,Mg)_3(PO4)_2 \cdot 4H_2O]$ where H⁺ ions have been suggested to oscillate between OH sites 378 and PO₄ (Frost et al., 2013a,c). Many analytical techniques cannot detect H⁺, and for techniques that can, because there is already significant OH⁻ or H₂O in giniite, it is unlikely such additional 379 H⁺ would be easily identified. If undetected H⁺ resides in the "O7 void", this may explain the 380 381 bond valence deficiency and add to the interesting nature of the mineral. Giniite has gained recent attention in industry and as a potential mineral occurring on 382 Mars. The characteristic of the mineral to maintain the giniite structure and chemistry while 383 384 exhibiting different morphologies based on formation environment mean it could be a powerful 385 indicator of past environments and potentially of life, on Earth or Mars. Overall, the results of this study refine and better detail the structure of giniite and confirm that giniite is indeed a 386 monoclinic mineral rather than orthorhombic. The results here are also consistent with the ideal 387 formula, including the Fe valence ratio, for natural giniite. However, based on synthetic giniite, 388 variations in the Fe valence ratio cannot be ruled out in the natural mineral, and this should be 389 further investigated in the future. Along with morphology, the Fe valence ratio may be an 390 391 indicator of formation conditions. In addition, if considering giniite as a potential Martian resource, valence can influence how, and how much, water and H⁺ can be evolved from the 392 mineral. 393 394 Raman spectrometers, like the spectrometers on the SuperCam and Scanning Habitable

Environments with Raman & Luminescence for Organics & Chemicals (SHERLOC) instrument
 suites on *Perseverance*, X-ray diffractometers, like the CheMin instrument on *Curiosity*, and the

397	Mössbauer instruments carried by Spirit and Opportunity all have the potential to identify giniite							
398	in situ. This is especially the case if a combination of these instruments is deployed. We							
399	encourage deployment of these mineralogical instruments on future Mars missions with data							
400	libraries that include secondary phosphates to help identify minerals like giniite that can							
401	elucidate habitability and potentially be used as a resource.							
402	Implications							
403	Giniite has become a recent mineral of interest as a potential component in applications							
404	including water purification, energy storage, and bio-medical materials on Earth. However,							

secondary phosphate minerals, like giniite, are also likely to hold a wealth of information

406 regarding past Martian surface processes and past environments. If confirmed on Mars, the

407 variable morphologies and broad formation conditions of giniite may make the mineral

408 especially important as an indicator of past environments and habitably. The discovery of giniite

409 with tubular morphologies on Mars, or in samples returned from Mars, might also be an indicator

410 of past life on the planet. In addition, terrestrial investigations of giniite as a resource in

411 technology and industry have potential implications for Mars. If substantial amounts of giniite

412 are present on Mars, long-term human exploration missions to the planet may be able to utilize

the mineral in applications developed on Earth (e.g., water purification, energy storage, bio-

414 medical materials). Beyond this, giniite is also potentially a resource for phosphate, water, and

415 fuel generation on Mars.

Deepening our knowledge base of secondary minerals that are possible or probable on the Martian surface, like giniite, will enhance our resource flexibility during long-term missions on Mars while also yielding insight into the Martian past. Secondary phosphate minerals in samples potentially collected by the *Perseverance* rover and returned by future missions will further our

420	knowledge of ancient aqueous environments on Mars, their habitability, and potential resources
421	for future human missions. However, if minerals like giniite are to be fully explored as scientific
422	and practical resources, up to date fundamental data, like those provided in this study, are
423	essential.
424	
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428	

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Figure Captions
Figure 1. BSE image at 15 kV of giniite. Dots are locations of EMP analysis on the crystal.
Figure 2. Calculated comparison of XRD patterns of natural (top and bottom patterns) and
synthetic ferrian (middle pattern) giniite. Patterns are calculated for $CuK\alpha$ radiation.
Figure 3. Broad Scan Raman spectra of giniite showing band consistent with giniite chemistry
and structure. Typical PO4, Fe-O, and O-H ranges are shown at bottom of figure and are
consistent with Raman of synthetic giniite and other minerals with structure similarities. 2nd-
order Savitzky-Golay filter applied (interval of 10). See Supplementary Figure S2 for plot of raw
data.
Figure 4. Giniite structure viewed down b (top) and a (bottom) axes. Four unit cells pictured.
Spheres on tetrahedral and octahedral corners are oxygen. Smaller spheres are H ⁺ , O7 oxygen
atoms discussed in the text are labeled in lower right cell of the top panel. Supplementary Figure
S3 is a close up of this "void" area.

762		Tables
763		
764	Table 1. Summary of cry	stallographic data for giniite
	Empirical chemical formula	$(Fe^{2+}_{0.80}Mn_{0.11}Mg_{0.02})_{\Sigma=0.93}Fe^{3+}_{4}(PO_{4})_{4.03}(OH)_{2}\cdot 2H_{2}O$
	Ideal chemical formula	$Fe^{2+}Fe^{3+}_{4}(PO_{4})_{4}(OH)_{2}\cdot 2H_{2}O$
	Crystal symmetry	Monoclinic
	Space group	P2/n
	a (Å)	10.3472(6)
	<i>b</i> (Å)	5.1497(2)
	<i>c</i> (Å)	14.2338(7)
	β (°)	111.175(6)
	$V(Å^3)$	707.24(7)
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Table 2. Select bolid distances (A) for glinte								
Fe1—O8	1.986 (2)	P1—O3	1.518 (2)					
Fe1—O7	2.069 (2)	P101	1.526 (2)					
Fe1—O2	2.082 (2)	P1—O4	1.544 (2)					
Fe1—O10W	2.183 (2)	P1—O2	1.553 (2)					
Fe1—O4	2.253 (2)	Ave.	1.54 (2)					
Fe1—O9H	2.279 (2)							
Ave.	2.14 (12)							
		P2—O6	1.513 (2)					
Fe2—O4 x 2	1.995 (2)	P2—O5	1.521 (2)					
Fe2—O2 x 2	2.013 (2)	P2—O8	1.537 (2)					
Fe2—O9H x 2	2.024 (2)	P2—O7	1.563 (2)					
Ave.	2.01 (2)	Ave.	1.53 (2)					
Fe3—O6	1.938 (2)							
Fe3—O5	1.954 (2)							
Fe3—O3	1.969 (2)							
Fe3—O1	1.985 (2)							
Fe3—O9H	2.024 (2)							
Fe3—O10W	2.226 (2)							
Ave.	2.02 (11)							

Table 2. Select bond distances (Å) for giniite

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	Fe1	Fe2	Fe3	P1	P2	Sum
01			0.543	1.323		1.866
02	0.356	0.503 x 2↓		1.233		2.092
03			0.567	1.354		1.920
O4	0.224	0.529 x 2↓		1.263		2.016
05			0.591		1.341	1.932
06			0.617		1.373	1.990
07	0.369				1.199	1.568
08	0.462				1.287	1.749
09	0.209	0.489 x 2↓	0.489			1.187
O10	0.271		0.283			0.554
Sum	1.891	3.042	3.090	5.173	5.200	

771	Table 3. Bond	Valence	calculations	for Fe-O	and P-O	bonds.
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Analysis					Fitted ^d		
					This Study	This Study	
(wt%)	Keller	This Study ^b		Keller	Ideal	Best Fit	Ideal
Fe ₂ O ₃	46.07	51.25	(0.34)	39.21	42.74	41.73	43.80
FeO	-	-	-	6.17	7.66	8.57	9.85
Al ₂ O ₃	1.20	-	-	1.20			
MgO	0.68	0.10	(0.02)	0.68	0.10	0.10	
P_2O_5	36.99	38.34	(0.25)	36.99	38.34	38.34	38.94
MnO	0.63	1.07	(0.20)	0.63	1.07	1.07	
H ₂ O _(Total) ^a	-	-	-	6.95	7.24	7.31	7.41
Total	85.57	90.76	(0.37)	91.83	97.15	97.11	100.00

Table 4. Comparison of chemical analyses of giniite by EMP in oxide wt%. This study
compared to Keller (1980a).

777NOTE: Parenthetical values are 1 standard deviation. ${}^{a}H_{2}O_{(Total)}$ is the wt% sum of OH and molecular H2O. ${}^{b}Based$ 778on 15 analyses. ${}^{d}Values$ in table are based on a calculated fit. Fits for Keller and "This Study Ideal" based on ideal779formula and fully occupied Fe3+ sites. "This Study Best Fit" represent best fit result from a range of calculated

780 stoichiometries using EMP data from this study.

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Figures



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783 784

Figure 1. BSE image at 15 kV of giniite. Dots as locations of EMP analysis on the crystal.

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Figure 2. Calculated comparison of XRD patterns of natural (top and bottom patterns) and synthetic ferrian (middle pattern) giniite. Patterns are calculated for $CuK\alpha$ radiation.



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Figure 4. Giniite structure viewed down *b* (top) and *a* (bottom) axes. Four unit cells pictured. Spheres on tetrahedral and octahedral corners are oxygen. Smaller spheres are H^+ . O7 oxygen atoms discussed in the text are labeled in lower right cell of the top panel. Supplementary Figure S3 is a close up of this "void" area.