

1 **Revision 1**

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3 **Maruyamaite, $K(\text{MgAl}_2)(\text{Al}_5\text{Mg})\text{Si}_6\text{O}_{18}(\text{BO}_3)_3(\text{OH})_3\text{O}$, a potassium-dominant tourmaline**

4 **from the ultrahigh-pressure Kokchetav massif, northern Kazakhstan:**

5 **Description and crystal structure**

6
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ABSTRACT

25 Maruyamaite, ideally $K(MgAl_2)(Al_5Mg)Si_6O_{18}(BO_3)_3(OH)_3O$, was recently approved as

26 the first K-dominant mineral-species of the tourmaline supergroup. It occurs in ultra-high

27 pressure quartzofeldspathic gneisses of the Kumdy-Kol area of the Kokchetav Massif, northern

28 Kazakhstan. Maruyamaite contains inclusions of microdiamonds, and probably crystallized near

29 the peak pressure conditions of UHP metamorphism in the stability field of diamond. Crystals

30 occur as anhedral to euhedral grains up to 2 mm across, embedded in a matrix of anhedral quartz

31 and K-feldspar. Maruyamaite is pale brown to brown with a white to very pale-brown streak and

32 has a vitreous luster. It is brittle and has a Mohs hardness of ~ 7 ; it is non-fluorescent, has no

33 observable cleavage or parting, and has a calculated density of 3.081 g cm^{-3} . In plane-polarized

34 transmitted light, it is pleochroic, O = darkish brown, E = pale brown. Maruyamaite is uniaxial

35 negative, $\omega = 1.634$, $\epsilon = 1.652$, both ± 0.002 . It is rhombohedral, space group $R3m$, $a = 15.955(1)$,

36 $c = 7.227(1) \text{ \AA}$, $V = 1593(3) \text{ \AA}^3$, $Z = 3$. The strongest ten X-ray diffraction lines in the powder

37 pattern are [d in $\text{\AA}(I)(hkl)$]: {note to typesetting: minus signs are overbars on top of the following

38 digit} 2.581(100)(051), 2.974(85)(-132), 3.995 (69)(-240), 4.237(59)(-231), 2.046(54)(-162),

39 3.498(42)(012), 1.923(36)(-372), 6.415(23)(-111), 1.595(22)(-5.10.0), 5.002(21)(021) and

40 4.610(20)(030). The crystal structure of maruyamaite was refined to an R_1 index of 1.58% using

41 1149 unique reflections measured with MoK α X-radiation. Analysis by a combination of electron

42 microprobe and crystal-structure refinement gave SiO $_2$ 36.37, Al $_2$ O $_3$ 31.50, TiO $_2$ 1.09, Cr $_2$ O $_3$

43 0.04, Fe $_2$ O $_3$ 0.33, FeO 4.01, MgO 9.00, CaO 1.47, Na $_2$ O 0.60, K $_2$ O 2.54, F 0.30, B $_2$ O $_3$ (calc)

44 10.58, H $_2$ O(calc) 2.96, sum 100.67 wt%. The formula unit, calculated on the basis of 31 anions

45 pfu with B = 3, OH = 3.24 apfu (derived from the crystal structure) and the site populations

46 assigned to reflect the mean interatomic distances, is $(K_{0.53}Na_{0.19}Ca_{0.26}\square_{0.02})_{\Sigma X=1.00}$
47 $(Mg_{1.19}Fe^{2+}_{0.55}Fe^{3+}_{0.05}Ti_{0.14}Al_{1.07})_{\Sigma Y=3.00}(Al_{5.00}Mg_{1.00})(Si_{5.97}Al_{0.03}O_{18})(BO_3)_3(OH)_3(O^{2-}_{0.60}$
48 $F_{0.16}OH_{0.24})$. Maruyamaite, ideally $K(MgAl_2)(Al_5Mg)(BO_3)_3(Si_6O_{18})(OH)_3O$, is related to oxy-
49 dravite: ideally $Na(MgAl_2)(Al_5Mg)(BO_3)_3(Si_6O_{18})(OH)_3O$, by the substitution ${}^XK \rightarrow {}^XNa$.

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51 *Keywords:* Maruyamaite, tourmaline, new mineral, electron-microprobe analysis, optical
52 properties, crystal-structure refinement, Kokchetav Massif, northern Kazakhstan, ultrahigh-
53 pressure, microdiamond inclusions.

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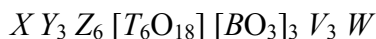
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INTRODUCTION

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The general formula of the minerals of the tourmaline supergroup can be written as

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where $X = \text{Na, K, Ca, Pb}^{2+}, \text{Bi, } \square$ (vacancy); $Y = \text{Li, Mg, Fe}^{2+}, \text{Mn}^{2+}, \text{Al, Cr}^{3+}, \text{V}^{3+}, \text{Fe}^{3+}, \text{Ti}^{4+}$; Z

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$= \text{Mg, Fe}^{2+}, \text{Al, Fe}^{3+}, \text{V}^{3+}, \text{Cr}^{3+}$; $T = \text{Si, Al, B}$; $B = \text{B}$; $V = \text{OH, O}$; $W = \text{OH, F, O}$. The tourmaline

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supergroup minerals are important indicator minerals, providing both chemical (e.g., Selway et

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al. 1998a,b, 1999, 2000a, b, 2002; Novák et al. 2004, 2011; Agrosi et al. 2006; Lussier and

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Hawthorne 2011; Lussier et al. 2008, 2011a,b; Hawthorne and Dirlam 2011; Henry and Dutrow

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1990, 1996; Dutrow and Henry 2011; Bačík et al. 2011; van Hinsberg and Schumacher 2009;

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van Hinsberg et al. 2011a) and isotopic (e.g., Marschall et al. 2006; van Hinsberg and Marschall

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2007; Ludwig et al. 2011; Marschall and Jiang 2011; van Hinsberg et al. 2011b; Shabaga et al.

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2010; Hezel et al. 2011) information on the evolution of their host rocks. However, the chemical

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composition of tourmalines is also controlled by short-range and long-range constraints (e.g.,

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Hawthorne 1996, 2002; Hawthorne and Henry 1999; Bosi and Lucchesi 2007; Bosi 2010, 2011,

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2013; Bosi et al. 2010; Henry and Dutrow 2011; Skogby et al. 2012). The dominant constituents

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at the X site are Na, Ca and \square , and until now, the remaining constituents, K, Pb^{2+} and Bi, have

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been found only in subordinate amounts. However, Ota et al. (2008a,b) and Shimizu and

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Ogasawara (2013) reported K-rich and K-dominant tourmaline from diamondiferous Kokchetav

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UHP metamorphic rocks in northern Kazakhstan (Kaneko et al. 2000). Here we describe a new

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^xK -dominant tourmaline from these rocks. The new species and name have been approved by the

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International Mineralogical Association Commission on New Minerals, Nomenclature and

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Classification (2013-123). Maruyamaite is named in honour of Professor Shigenori Maruyama

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(Earth-Life Science Institute, Tokyo Institute of Technology, Japan), born 24th December, 1949,

78 in Tokushima, Japan. Professor Maruyama is a prominent figure in the field of regional tectonics
79 and led the project which unraveled the mechanism of continental subduction and exhumation
80 (from a depth greater than 120 km) of the Kokchetav massif of northern Kazakhstan, and
81 discovered the diamond-bearing UHP rocks that contain the new K-dominant tourmaline.
82 Holotype material is deposited in the collections of the National Museum of Nature and Science,
83 Tsukuba, Japan, registered number NSM- MF15696.

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OCCURRENCE

86 Maruyamaite occurs as core domains of chemically zoned tourmaline in a
87 quartzofeldspathic rock collected from the Kundry-Kol area of the Kokchetav Massif (Ota et al.
88 2008a,b; Shimizu and Ogasawara 2013). The rock occurs as thin layers of variable thickness
89 (less than several cm) in diamond-bearing pelitic gneiss, and consists mainly of quartz, K-
90 feldspar and tourmaline (up to 20 vol%) with minor amounts of goethite, titanite, zircon,
91 phengite, phlogopite, apatite, chlorite, zoisite, pumpellyite, graphite and diamond. Tourmaline
92 occurs as anhedral to euhedral crystals up to 2 mm across with strong chemical zoning; K
93 decreases from core to rim (Fig. 1). Although the mantle and rim also contain high amounts of K
94 for magnesian tourmaline, their K-contents in atoms per formula unit (apfu) do not exceed those
95 of Na or Ca (i.e., the mantle and the rim compositions correspond to K-bearing dravite or oxy-
96 dravite). Microdiamond inclusions occur only in the K-dominant core of tourmaline
97 (maruyamaite) and in zircon. On the other hand, flake graphite and quartz occur in the mantle
98 and rim. More detailed descriptions of the chemical zoning and inclusion content of tourmaline
99 in these rocks are given by Shimizu and Ogasawara (2013).

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PHYSICAL PROPERTIES

102 Maruyamaite is pale brown to brown with a white to very pale-brown streak, has a
103 vitreous luster and does not fluoresce in ultraviolet light. It has a Mohs hardness of ~7, and is
104 brittle with a splintery fracture; the calculated density is 3.081 g cm^{-3} . A spindle stage was used
105 to orient a crystal for measurement of optical properties. The optical orientation was determined
106 by transferring the crystal from the spindle stage to a single-crystal diffractometer and measuring
107 the relative axial relations by X-ray diffraction. In transmitted polarized light, maruyamaite is
108 pleochroic with O = darkish brown, E = pale brown. It is uniaxial negative, with indices of
109 refraction $\omega = 1.634$, $\varepsilon = 1.652$, both ± 0.002 , measured with gel-filtered Na light ($\lambda = 589.9 \text{ nm}$).

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RAMAN SPECTROSCOPY

112 Raman spectra of maruyamaite were obtained using an Ar^+ laser with a wavelength of
113 514.5nm. The Raman spectra (Fig. 2) show a strong single band at 3572 cm^{-1} . The peak may be
114 assigned to an O-H stretching band involving (OH) at the O3 site. Below 1500 cm^{-1} , there is a
115 very weak band at 1230 cm^{-1} (internal modes of the BO_3 group), a group of bands (977 m, 1051
116 m, 1106 sh) centered on $\sim 1000 \text{ cm}^{-1}$ (internal modes of the SiO_4 group), another group of bands
117 (669 m, 703 s, 789 vs) centered on $\sim 720 \text{ cm}^{-1}$ (delocalised Si_6O_{18} ring motions and AlO_6
118 deformations), and lower-frequency bands at 538 vw, 500 w, 367 vs, 242 m, 212 s and 155 w
119 cm^{-1} (vs = very strong, s = strong, m = medium, w = weak, vw = very weak, sh = shoulder)
120 (various motions involving the more weakly bonded cations K, Na, Mg). Assignments follow
121 those of McKeown (2008).

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MÖSSBAUER SPECTROSCOPY

125 The Mössbauer spectrum was collected in transmission geometry at room temperature
126 (RT) using a ^{57}Co point source. The spectrometer was calibrated using the RT spectrum of $\alpha\text{-Fe}$.
127 For preparing the Mössbauer absorber, several K-tourmaline grains were attached to adhesive
128 tape and mounted on a Pb disk with a 500 μm aperture, and the collection time was one month.
129 The spectrum was analyzed using a Voigt-based quadrupole-splitting distribution (QSD) method
130 (Rancourt and Ping 1991). To account for thickness and/or instrumental broadening, the
131 Lorentzian linewidth of the symmetrical elemental doublets of the QSD was allowed to vary
132 during spectrum fitting. There are two symmetrical doublets with Mössbauer parameters typical
133 of octahedrally coordinated Fe^{2+} and one doublet with parameters typical of octahedrally
134 coordinated Fe^{3+} (Fig. 3, Table 1). The area ratio of the Fe^{3+} doublet to the Fe^{2+} doublets is 0.076
135 : 0.924, and this value was also taken as the $\text{Fe}^{3+} : \text{Fe}^{2+}$ ratio (i.e., assuming equal recoil-free
136 fractions for Fe^{2+} and Fe^{3+}).

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CHEMICAL COMPOSITION

139 The crystal used for the collection of the X-ray intensity data was embedded in epoxy,
140 polished, carbon coated and analyzed with a Cameca SX-100 electron microprobe operating in
141 wavelength-dispersion mode with an excitation voltage of 15 kV, a specimen current of 10 nA, a
142 beam diameter of 10 μm , peak count-times of 20 s and background count-times of 10 s. The
143 following standards and crystals were used: Si, Mg, Ca: diopside, TAP, LTAP, LPET; Ti:
144 magnesio-hornblende, LLiF; Fe: fayalite, LLiF; Mn: rhodonite, LLiF; Na: albite, TAP; Al:
145 andalusite, TAP; K: orthoclase, LPET; F: fluoro-riebeckite, LTAP; Zn: willemite, LLiF; Cr:
146 chromite, LLiF. Data reduction was done using the $\phi(\rho Z)$ procedure of Pouchou and Pichoir

147 (1985). The average of eleven analyses on a single grain is given in Table 2.

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X-RAY POWDER DIFFRACTION

150 X-ray powder-diffraction data were collected with a Bruker D8 Discover SuperSpeed
151 micro-powder diffractometer with a multi-wire 2D detector using a modified Gandolfi
152 attachment ($\text{CuK}\alpha$, $\lambda = 1.54178 \text{ \AA}$); 50kV/60 mA; two 30 min. frames, merged; no internal
153 standard used. Data were indexed on the following refined cell-dimensions: $a = 15.915(3)$, $c =$
154 $7.120(2) \text{ \AA}$, $V = 1561.8(7) \text{ \AA}^3$, and the pattern is given in Table 3.

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CRYSTAL-STRUCTURE REFINEMENT

157 A fragment was extracted from the center of a strongly zoned crystal of maruyamaite in
158 thin section, attached to a tapered glass fiber and mounted on a Bruker APEX II ULTRA three-
159 circle diffractometer equipped with a rotating-anode generator (MoK α), multilayer optics and an
160 APEX II 4K CCD detector. A total of 6230 intensities was collected to $60^\circ 2\theta$ using 2 s per 0.2°
161 frame and a crystal-to-detector distance of 5 cm. Empirical absorption corrections (SADABS;
162 Sheldrick 2008) were applied and equivalent reflections were corrected for Lorentz, polarization
163 and background effects, averaged and reduced to structure factors. The unit-cell dimensions were
164 obtained by least-squares refinement of the positions of 4678 reflections with $I > 10\sigma I$ and are
165 given in Table 4, together with other information pertaining to data collection and structure
166 refinement.

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All calculations were done with the SHELXTL PC (Plus) system of programs. The structure was refined by full-matrix least-squares methods with anisotropic-displacement parameters for all atoms, and converged to a final R1 index of 1.58%. Refined atom coordinates

170 and anisotropic-displacement parameters are listed in Table 5, selected interatomic distances are
171 given in Table 6, and refined site-scattering values (Hawthorne et al. 1995) and assigned site-
172 populations are given in Table 7. The CIF is on deposit and available as listed below.¹

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174

CHEMICAL FORMULA

175 In the absence of a determination of H₂O content, the chemical composition (Table 2) of
176 tourmaline was initially reduced to a chemical formula on the basis of 31 anions per formula unit
177 with B = 3 apfu and (OH) + F = 4 apfu, and the resulting formula provided a starting point for a
178 more accurate normalization. For maruyamaite, the Y- and Z-site contents, together with the
179 observed <Y-O> and <Z-O> distances (Table 6), indicate that there is significant disorder of Mg
180 and Al over the Y and Z sites. Taylor et al. (1995) refined the structure of an OH-deficient uvite
181 with significant disorder of this type, and Hawthorne (1996) has discussed how the bond-valence
182 requirements for O at the O1 site can drive the order-disorder reaction ${}^Y\text{Al} + {}^Z\text{Mg} = {}^Y\text{Mg} + {}^Z\text{Al}$.
183 This issue has been examined in more detail by Bosi (2013) who produced the expression
184 ${}^{O1}(\text{OH}) = 2 - 1.01\Sigma s^{O(1)} - 0.21 - F$ apfu to estimate the amount of (OH) at the O1 site where
185 $\Sigma s^{O(1)}$ is the bond valence incident at the O1 anion (exclusive of H). We used this expression to
186 estimate the amount of (OH) at O1 and then renormalized the chemical formula on this basis. We
187 again recalculated the ${}^{O1}(\text{OH})$ content on the basis of the new formula, and iterated this process
188 to convergence. There is the possibility of ${}^X\text{K}$ bonding to O1, but the observed distance of 3.218
189 Å results in an additional 0.03 v.u. incident at O1 and a negligible change (0.02 apfu) in the (OH)
190 content. The final calculated (OH) content of the crystal is 3.24 apfu, and calculation of the

¹ Deposit item AM-XX-XXX, CIF, is stored on the MSA web site and available via the American Mineralogist Table of Contents. Find the article in the table of contents at GSW (ammin.geoscienceworld.org) or MSA (www.minsocam.org), and then click on the deposit link.

191 empirical chemical formula with this OH content gave the following result:

192 $(K_{0.53}Na_{0.19}Ca_{0.26}\square_{0.02})_{\Sigma X=1.00}(Mg_{2.20}Fe^{2+}_{0.55}Fe^{3+}_{0.05}Ti_{0.14}Al_{0.07})_{\Sigma Y=3.01}(Al_6)(Si_{5.97}Al_{0.03}O_{18})$

193 $(BO_3)_3(OH)_3(O^{2-}_{0.60}F_{0.16}OH_{0.24})$.

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195

SITE POPULATIONS

196 The T site is almost completely occupied by Si according to the formula derived from the
197 electron-microprobe analysis, and the <T-O> distance (Table 6) is 1.621 Å, close to the value of
198 1.620 Å proposed as the <Si-O> distance in the tourmaline structure by MacDonald and
199 Hawthorne (1995). Although the Z site in the tourmaline structure is dominated by Al,
200 Hawthorne et al. (1993) showed that there is significant disorder of Al and Mg over the Y and Z
201 sites in tourmaline, and the <Z-O> distance is sensitive to occupancy by other cations
202 (Hawthorne et al. 1993; Burns et al. 1994; Bosi 2008, 2011; Bosi and Lucchesi 2004; Bosi and
203 Skoby 2013; Bosi et al. 2004; Clark et al. 2011; Ertl et al. 2010a,b; Novák et al. 2013). Where Z
204 is completely occupied by Al, the <Z-O> distances are in the range 1.904-1.910 Å (e.g.,
205 Hawthorne et al. 1993; Bosi 2008; Lussier et al. 2011a; MacDonald et al. 1993; Selway et al.
206 1998a,b). Bosi and Lucchesi (2007) showed that <Z-O> distances are also inductively affected
207 by different occupancies at the Y site. The <Z-O> distance varies linearly with the mean radius
208 of the constituent Z cations, and the <Z-O> distance in the holotype maruyamaite structure
209 (1.932 Å, Table 6) is significantly longer than the values for tourmaline structures where $Z = Al_6$
210 (1.904-1.911 Å, see above), indicating that maruyamaite has significant disorder of Mg and Al
211 over the Y- and Z-sites, as discussed above. In Figure 4, the <Z-O> distance in maruyamaite
212 (1.932 Å) intersects the line through the rest of the data at 5.00 Al pfu (marked by the pink
213 diamond in Fig. 4), and hence the site population of Z was set at $Al_{5.00}Mg_{1.00}$ apfu and the site

214 population of Y was modified accordingly.

215 The resultant site-populations for maruyamaite are given in Table 7. Maruyamaite,
216 ideally $\text{K}(\text{MgAl}_2)(\text{Al}_5\text{Mg})(\text{BO}_3)_3(\text{Si}_6\text{O}_{18})(\text{OH})_3\text{O}$, is the K analogue of oxy-dravite, ideally
217 $\text{Na}(\text{MgAl}_2)(\text{Al}_5\text{Mg})(\text{BO}_3)_3(\text{Si}_6\text{O}_{18})(\text{OH})_3\text{O}$ (Bosi and Skogby 2013), and the new rootname is in
218 accord with Henry et al. (2011), the current IMA-approved nomenclature for the tourmaline
219 supergroup minerals.

220

221 DISCUSSION

222 The direct association of maruyamaite with diamond inclusions at the Kokchetav locality
223 suggests that high K contents in magnesian tourmaline may be an indicator of UHP conditions.
224 However, tourmaline developed in other UHP terrains typically do not exhibit comparable K
225 contents, and generally contain <0.05 apfu K (cf. Ertl et al. 2010b). This apparent inconsistency
226 was examined in the experimental study of Berryman et al. (2014) who found that K contents in
227 synthetic dravitic tourmalines in K-bearing aqueous fluids increase to K-dominant contents at
228 UHP conditions. However, with significant amounts of Na and K in the aqueous fluids, the
229 amount of K in the synthetic tourmaline was greatly reduced under UHP conditions.

230 Significant amounts of K occur in povondraite (Grice et al. 1993; Záček et al. 2000), a
231 tourmaline in which Fe^{3+} dominates over Al and which forms at relatively low pressure,
232 suggesting that tourmalines with high contents of Fe^{3+} can accommodate far more K than Al-rich
233 tourmaline at low pressure. The implication is that K-dominant magnesian tourmalines such as
234 maruyamaite are not only a function of UHP conditions, but also the presence of highly potassic
235 fluids.

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IMPLICATIONS

238 The crystallization of maruyamaite requires both UHP conditions and K-dominated
239 compositions, and these conditions could occur by partial melting of subducted continental crust
240 (including sedimentary rocks) at high pressure in the stability field of diamond (Ota et al.
241 2008a,b). This conclusion is consistent with previous studies (e.g., Hwang et al. 2005) which
242 have shown that K-rich fluid was present at the Kokchetav UHP stages and played an important
243 role in the formation of metamorphic diamond and other UHP minerals such as K-rich
244 clinopyroxene.

245 As neither maruyamaite nor other tourmalines have been found in diamond-bearing
246 zircon (Shimizu and Ogasawara 2013), and both K-tourmaline and dravite are likely to be
247 unstable at peak metamorphic conditions of the Kokchetav Massif (> 6Gpa, e.g., Ogasawara et
248 al. 2002) according to Ota et al. (2008a,b) and Berryman et al. (2014), it is probable that
249 maruyamaite formed during exhumation after peak metamorphism (but still at UHP conditions).
250 The discovery of maruyamaite and related K-rich tourmaline in the Kokchetav UHP rocks
251 highlights the potential of tourmaline as a recorder of metamorphic history.

252 Wunder et al. (2015) showed that tourmaline can incorporate small amount of nitrogen as
253 NH_4^+ (which has an ionic radius similar to that of K) at high pressure. High-pressure
254 tourmalines, including maruyamaite, can be a container of volatile elements such as H, B and N
255 in the deep mantle, and play an important role in the recycling of crustal material.

256

257

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REFERENCES

- 268 Agrosi, G., Bosi, F., Lucchesi, S., Melchiorre, G., and Scandale, E. (2006) Mn-tourmaline
269 crystals from island of Elba (Italy): Growth history and growth marks. American
270 Mineralogist, 91, 944–952.
- 271 Bačík, P., Méres Š., and Uher, P. (2011) Vanadium-bearing tourmaline in metacherts from
272 Chvojnica, Slovak Republic: crystal chemistry and multistage evolution. Canadian
273 Mineralogist, 49, 195–206.
- 274 Berryman, E., Wunder, B., and Rhede, D. (2014) Synthesis of K-dominant tourmaline. American
275 Mineralogist, 99, 539–542.
- 276 Bosi, F. (2008) Disordering of Fe²⁺ over octahedrally coordinated sites of tourmaline. American
277 Mineralogist, 93, 1647–1653.
- 278 ——— (2010) Octahedrally coordinated vacancies in tourmaline: a theoretical approach.
279 Mineralogical Magazine, 74, 1037–1044.
- 280 ——— (2011) Stereochemical constraints in tourmaline: from a short-range to a long-range
281 structure. Canadian Mineralogist, 49, 17–27.
- 282 ——— (2013) Bond-valence constraints around the O1 site of tourmaline. Mineralogical
283 Magazine, 77, 343–351.
- 284 Bosi, F., and Lucchesi, S. (2004) Crystal chemistry of the schorl-dravite series. European Journal
285 of Mineralogy, 16, 335–344.
- 286 ——— (2007) Crystal chemical relationships in the tourmaline group: Structural constraints on
287 chemical variability. American Mineralogist, 92, 1054–1063.
- 288 Bosi, F., and Skogby, H. (2013) Oxy-dravite, Na(Al₂Mg)(Al₅Mg)(Si₆O₁₈)(BO₃)₃(OH)₃O, a new
289 mineral species of the tourmaline supergroup. American Mineralogist, 98, 1442–1448.

- 290 Bosi, F., Lucchesi, S., and Reznitskii, L. (2004) Crystal chemistry of the dravite-chromdravite
291 series. *European Journal of Mineralogy*, 16, 345–352.
- 292 Bosi, F., Balić-Žunić, T., and Surour, A.A. (2010) Crystal structure analysis of four tourmalines
293 from the Cleopatra’s Mines (Egypt) and Jabal Zalm (Saudi Arabia), and the role of Al in
294 the tourmaline group. *American Mineralogist*, 95, 510–518.
- 295 Burns, P.C., MacDonald, D.J., and Hawthorne, F.C. (1994) The crystal chemistry of manganese-
296 bearing elbaite. *Canadian Mineralogist*, 32, 31–41.
- 297 Clark, C.M., Hawthorne, F.C, and Ottolini, L. (2011) Fluor-dravite,
298 $\text{NaMg}_3\text{Al}_6\text{Si}_6\text{O}_{18}(\text{BO}_3)_3(\text{OH})_3\text{F}$, a new mineral of the tourmaline group from the Crabtree
299 emerald mine, Mitchell county, North Carolina: Description and crystal structure.
300 *Canadian Mineralogist*, 49, 57–62.
- 301 Dutrow, B.L., and Henry, D.J. (2011) Tourmaline: A geologic DVD. *Elements*, 7, 301–306.
- 302 Ertl, A., Rossman, G.R., Hughes, J.M., London, D., Wang, Y., O’Leary, J.A., Dyar, M.D.,
303 Prowatke, S., Ludwig, T., and Tillmanns, E. (2010a) Tourmaline of the elbaite-schorl
304 series from the Himalaya Mine, Mesa Grande, California: A detailed investigation.
305 *American Mineralogist*, 95, 24–40.
- 306 Ertl, A., Marschall, H.R., Giester, G., Henry, D.J., Schertl, H.P., Ntaflos, T., Luvizotto, G.L.,
307 Nasdala, L., and Tillmanns, E. (2010b) Metamorphic ultrahigh-pressure tourmaline:
308 Structure, chemistry, and correlations to P-T conditions. *American Mineralogist*, 95, 1–
309 10.
- 310 Grice, J.D., Ercit, T.S., and Hawthorne, F.C. (1993) Povondraite, a redefinition of the tourmaline
311 ferridravite. *American Mineralogist*, 78, 433–436.
- 312 Hawthorne, F.C. (1996) Structural mechanisms for light-element variations in tourmaline.

- 313 Canadian Mineralogist, 34, 123–132.
- 314 ——— (2002) Bond-valence constraints on the chemical composition of tourmaline. Canadian
315 Mineralogist, 40, 789–797.
- 316 Hawthorne, F.C., and Dirlam, D.M. (2011) Tourmaline, the indicator mineral: From atomic
317 arrangement to Viking navigation. Elements, 7, 307–312.
- 318 Hawthorne, F.C., and Henry, D.J. (1999) Classification of the minerals of the tourmaline group.
319 European Journal of Mineralogy, 11, 201–215.
- 320 Hawthorne, F.C., MacDonald, D.J., and Burns, P.C. (1993) Reassignment of cation site-
321 occupancies in tourmaline: Al/Mg disorder in the crystal structure of dravite. American
322 Mineralogist, 78, 265–270.
- 323 Hawthorne, F.C., Ungaretti, L., and Oberti, R. (1995) Site populations in minerals: terminology
324 and presentation of results of crystal-structure refinement. Canadian Mineralogist, 33,
325 907–911.
- 326 Henry, D.J., and Dutrow, B.L. (1990) Ca substitution in Li-poor aluminous tourmaline. Canadian
327 Mineralogist, 28, 111–124.
- 328 Henry, D.J., and Dutrow, B.L. (1996) Metamorphic tourmaline and its petrologic applications.
329 Reviews in Mineralogy, 33, 503–557.
- 330 Henry, D.J., and Dutrow, B.L. (2011) The incorporation of fluorine in tourmaline: Internal
331 crystallographic controls or external environmental influences? Canadian Mineralogist,
332 49, 41–56.
- 333 Henry, D.J., Novák, M., Hawthorne, F.C., Ertl, A., Dutrow, B.L., Uher, P., and Pezzotta, F.
334 (2011) Nomenclature of the tourmaline super-group minerals. American Mineralogist,
335 96, 895–913.

- 336 Hezel, D.C., Kalt, A., Marschall, H.R., Ludwig, T., and Meyer, H.-P. (2011) Major-element and
337 Li, Be compositional evolution of tourmaline in an S-type granite–pegmatite system and
338 its country rocks: an example from Ikaria, Aegean Sea, Greece. *Canadian Mineralogist*,
339 49, 321–340.
- 340 Hwang, S.L., Shen, P., Chu, H.T., Yui, T.F., Liou, J.G., Sobolev, N.V., and Shatsky, V.S. (2005)
341 Crust-derived potassic fluid in metamorphic microdiamond. *Earth and Planetary Science*
342 *Letters*, 231, 295–306.
- 343 Kaneko, Y., Maruyama, S., Terabayashi, M., Yamamoto, H., Ishikawa, M., Anma, R.,
344 Parkinson, C.D., Ota, T., Nakajima, Y., Katayama, I., Yamamoto, J., and Yamauchi, K.
345 (2000) Geology of the Kokchetav UHP-HP metamorphic belt, Northern Kazakhstan. *The*
346 *Island Arc*, 9, 264–283.
- 347 Ludwig, T., Marschall, H.R., Pogge von Strandmann, P.A.E., Shabaga, B.M., Fayek, M., and
348 Hawthorne, F.C. (2011) A secondary ion mass spectrometry (SIMS) re-evaluation of B
349 and Li isotopic compositions of Cu-bearing elbaite from three global localities.
350 *Mineralogical Magazine*, 75, 2485–2494.
- 351 Lussier, A.J., and Hawthorne, F.C. (2011) Oscillatory zoned liddicoatite from central
352 Madagascar. II. Compositional variations and substitution mechanisms. *Canadian*
353 *Mineralogist*, 49, 89–104.
- 354 Lussier, A.J., Aguiar, P.M., Michaelis, V.K., Kroeker, S., Herwig, S., Abdu, Y., and Hawthorne,
355 F.C. (2008) Mushroom elbaite from the Kat Chay mine, Momeik, near Mogok,
356 Myanmar: I. Crystal chemistry by SREF, EMPA, MAS NMR and Mössbauer
357 spectroscopy. *Mineralogical Magazine*, 72, 747–761.
- 358 Lussier, A.J., Hawthorne, F.C., Aguiar, P.M., Michaelis, V.K., and Kroeker, S. (2011a) Elbaite-

- 359 liddicoatite from Black Rapids glacier, Alaska. *Periodico di Mineralogia*, 80, 57–73.
- 360 Lussier, A.J., Abdu, Y. Hawthorne, F.C., Michaelis, V.K., Aguiar, P.M., and Kroeker, S. (2011b)
- 361 Oscillatory zoned liddicoatite from Anjanabonoina, central Madagascar. I. Crystal
- 362 chemistry and structure by SREF and ¹¹B and ²⁷Al MAS NMR spectroscopy. *Canadian*
- 363 *Mineralogist*, 49, 63–88.
- 364 MacDonald, D.J., and Hawthorne, F.C. (1995) The crystal chemistry of Si ↔ Al substitution in
- 365 tourmaline. *Canadian Mineralogist*, 33, 849–858.
- 366 MacDonald, D.J., Hawthorne, F.C., and Grice, J.D. (1993) Foitite, $\square[\text{Fe}^{2+}_2(\text{Al},\text{Fe}^{3+})]\text{Al}_6\text{Si}_6\text{O}_{18}$
- 367 $(\text{BO}_3)_3(\text{OH})_4$, a new alkali-deficient tourmaline: description and crystal structure.
- 368 *American Mineralogist*, 78, 1299–1303.
- 369 Marschall, H.R., and Jiang, S.-Y. (2011) Tourmaline Isotopes: No element left behind. *Elements*,
- 370 7, 313–319.
- 371 Marschall, H.R., Ludwig, T., Altherr, R., Kalt, A., and Tonarini, S. (2006) Syros metasomatic
- 372 tourmaline: Evidence for very high- $\delta^{11}\text{B}$ fluids in subduction zones. *Journal of*
- 373 *Petrology*, 47, 1915–1942.
- 374 McKeown, D.A. (2008) Raman spectroscopy, vibrational analysis and heating of buergerite
- 375 tourmaline. *Physics and Chemistry of Minerals*, 35, 259–270.
- 376 Novák, M., Povondra, P., and Selway, J.B. (2004) Schorl-oxy-schorl to dravite-oxy-dravite
- 377 tourmaline from granitic pegmatites; examples from the Moldanubicum, Czech Republic.
- 378 *European Journal of Mineralogy*, 16, 323–333.
- 379 Novák, M., Škoda, P., Filip, J., Macek, I., and Vaculovič, T. (2011) Compositional trends in
- 380 tourmaline from intragranitic NYF pegmatites of the Třebíč Pluton, Czech Republic;
- 381 electron microprobe, Mössbauer and LA-ICP-MS study. *Canadian Mineralogist*, 49, 359–

- 382 380.
- 383 Novák, M., Ertl, A., Povondra, P., Galiová, M.V., Rossman, G.R., Pristacz, H., Prem, M.,
384 Giester, G., Gadas, P., and Škoda, R. (2013) Darrellhenryite, $\text{Na}(\text{LiAl}_2)\text{Al}_6(\text{BO}_3)_3\text{Si}_6\text{O}_{18}$
385 $(\text{OH})_3\text{O}$, a new mineral from the tourmaline supergroup. *American Mineralogist*, 98,
386 1886–1892.
- 387 Ogasawara, Y., Fukasawa, K., and Maruyama, S. (2002) Coesite exsolution from supersilicic
388 titanite in UHP marble from the Kokchetav Massif, northern Kazakhstan. *American*
389 *Mineralogist*, 87, 454–461.
- 390 Ota, T., Kobayashi, K., Katsura, T., and Nakamura, E. (2008a) Tourmaline breakdown in a
391 pelitic system: implications for boron cycling through subduction zones. *Contributions to*
392 *Mineralogy and Petrology*, 155, 19–32.
- 393 ——— (2008b) Boron cycling by subducted lithosphere; insights from diamondiferous
394 tourmaline from the Kokchetav ultrahigh-pressure metamorphic belt. *Geochimica et*
395 *Cosmochimica Acta*, 72, 3531–3541.
- 396 Pertlik, F., Ertl, A., Körner, W., Brandstätter, F., and Schuster, R. (2003) Na-rich dravite in the
397 marbles from Friesach, Carinthia, Austria: Chemistry and crystal structure. *Neues*
398 *Jahrbuch für Mineralogie Monatshefte*, 2003, 277–288.
- 399 Pouchou, J.L., and Pichoir, F. (1985) ‘PAP’ $\phi(\rho Z)$ procedure for improved quantitative
400 microanalysis. In J.T. Armstrong, Ed., *Microbeam Analysis*, p. 104–106. San Francisco
401 Press, San Francisco, California.
- 402 Rancourt D.G., and Ping J.Y. (1991) Voigt-based methods for arbitrary-shape static hyperfine
403 parameter distributions in Mössbauer spectroscopy. *Nuclear Instruments and Methods in*
404 *Physics Research*, B58, 85–97.

- 405 Selway, J.B., Černý, P., and Hawthorne, F.C. (1998a) Feruvite from lepidolite pegmatites at Red
406 Cross lake, Manitoba. *Canadian Mineralogist*, 36, 433–439.
- 407 Selway, J.B., Novák, M., Hawthorne, F.C., Černý, P., Ottolini, L., and Kyser, T.K. (1998b)
408 Rossmanite, $\square[\text{LiAl}_2]\text{Al}_6\text{Si}_6\text{O}_{18}(\text{BO}_3)_3(\text{OH})$, a new alkali-deficient tourmaline:
409 Description and crystal structure. *American Mineralogist*, 83, 896–900.
- 410 Selway, J.B., Novák, M., Černý, P., and Hawthorne, F.C. (1999) Compositional evolution of
411 tourmaline in lepidolite-subtype pegmatites. *European Journal of Mineralogy*, 11, 569–
412 584.
- 413 Selway, J.B., Černý, P., Hawthorne, F.C., and Novák, M. (2000a) The Tanco pegmatite at Bernic
414 Lake, Manitoba. XIV. Internal tourmaline. *Canadian Mineralogist*, 38, 877–891.
- 415 Selway, J.B., Novák, M., Černý, P., and Hawthorne, F.C. (2000b) The Tanco pegmatite at Bernic
416 Lake, Manitoba. XIII. Exocontact tourmaline. *Canadian Mineralogist*, 38, 869–976.
- 417 Selway, J.B., Smeds, S-A., Černý, P., and Hawthorne, F.C. (2002) Compositional evolution of
418 tourmaline in the petalite-subtype Nyköpingsgruvan pegmatites, Utö, Stockholm
419 Archipelago, Sweden. *GFF*, 124, 93–102.
- 420 Shabaga, B.M., Fayek, M., and Hawthorne, F.C. (2010) Boron and lithium isotopic compositions
421 as provenance indicators of Cu-bearing tourmalines. *Mineralogical Magazine*, 74, 241–
422 255.
- 423 Sheldrick, G.M. (2008) A short History of SHELX. *Acta Crystallographica*, A64, 112–122.
- 424 Shimizu, R., and Ogasawara, Y. (2013) Diversity of potassium-bearing tourmalines in
425 diamondiferous Kokchetav UHP metamorphic rocks: a geochemical recorder from peak
426 to retrograde metamorphic stages. *Journal of Asian Earth Science*, 63, 39–55.
- 427 Skogby, H., Bosi, F., and Lazor, P. (2012) Short-range order in tourmaline: a vibrational

- 428 spectroscopic approach to elbaite. *Physics and Chemistry of Minerals*, 39, 811–816.
- 429 Taylor, M.C., Cooper, M.A., and Hawthorne, F.C. (1995) Local charge-compensation in
430 hydroxyl-deficient uvite. *Canadian Mineralogist*, 33, 1215–1221.
- 431 van Hinsberg, V.J., and Marschall, H.R. (2007) Boron isotope and light element sector zoning in
432 tourmaline: Implications for the formation of B-isotopic signatures. *Chemical Geology*,
433 238, 141–148.
- 434 van Hinsberg, V.J., and Schumacher, J.C. (2009) The geothermobarometric potential of
435 tourmaline, based on experimental and natural data. *American Mineralogist*, 94, 761–
436 770.
- 437 van Hinsberg, V.J., Henry, D.J., and Dutrow, B.L. (2011a) Tourmaline as a petrologic forensic
438 mineral: A unique recorder of its geologic past. *Elements*, 7, 327–332.
- 439 van Hinsberg, V.J., Henry, D.J., and Marschall, H.R. (2011b) Tourmaline: an ideal indicator of
440 its host environment. *Canadian Mineralogist*, 49, 1–16.
- 441 Wunder, B., Berryman, E., Plessen, B., Rhede, D., Koch-Muller, M., and Heinrich, W. (2015)
442 Synthetic and natural ammonium-bearing tourmaline. *American Mineralogist*, 100, 250–
443 256.
- 444 Záček, V., Jirá, F., Petrov, A., and Hyršl, J. (2000) Tourmalines of the povondraite-(oxy) dravite
445 series from the cap rock of meta-evaporite in Alto Chapare, Cochabamba, Bolivia.
446 *Journal of the Czech Geological Society*, 45, 3–12.

447

FIGURE CAPTIONS

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Figure 1. (A) Photomicrograph and (B) characteristic X-ray K-K α map of

diamondiferous K-bearing tourmaline. Tourmaline shows discontinuous chemical zoning with

representative K-contents in apfu of 0.58 (core), 0.15 (mantle), and 0.04 (rim). Diamond

inclusions occur only in the K-dominant tourmaline core, i.e., maruyamaite (shown bounded by

red broken lines). Abbreviations in (A) are: Dia: diamond, Gr: graphite, Kfs: K-feldspar, Qz:

quartz, and Tur: tourmaline.

Figure 2. Raman spectra of maruyamaite with the laser (a) perpendicular, and (b) parallel

to the *c*-axis.

Figure 3. Mössbauer spectrum of maruyamaite; Full lines: Fe²⁺ doublets, dashed line:

Fe³⁺ doublet.

Figure 4. Variation in $\langle Z-O \rangle$ as a function of the ^ZAl* (= Al + Fe³⁺) content at the Z site

for selected tourmalines; uvites from McDonald and Hawthorne (1995) (green circles) and Clark

et al. (unpublished) (yellow circle); dravites from Bosi and Lucchesi (2004) (samples 235a,

235b, 65e, red circles) and Pertlik et al. (2003) (blue circle). The dashed line shows extrapolation

of holotype maruyamaite (pink diamond) to the regression line defined by the rest of the data and

the corresponding value of ^ZAl indicated by the dashed arrow: 5.00 apfu.

TABLE 1. Mössbauer parameters (mm/s) for maruyamaite

Cation	Centre shift (mm/s)	Quadrupole splitting	Area ratio
Fe ²⁺	1.086(8)	2.46(15)	44(5)
Fe ²⁺	1.099(20)	1.89(5)	49(5)
Fe ³⁺	0.35(10)	0.66(28)	2(1.8)

TABLE 2. Chemical composition (wt%) for maruyamaite

Constituent	Wt%	Range	Standard deviation
SiO ₂	36.37	35.83–36.74	0.30
TiO ₂	1.09	0.90–1.21	0.03
Al ₂ O ₃	31.50	31.25–32.06	0.11
B ₂ O ₃	10.58	–	–
Cr ₂ O ₃	0.04	0.01–0.06	0.01
FeO*	4.28	4.07–4.55	0.02
Fe ₂ O ₃	0.33	–	–
FeO	4.01	–	–
MgO	9.00	8.78–9.19	0.10
CaO	1.47	1.39–1.55	0.01
Na ₂ O	0.60	0.52–0.65	0.03
K ₂ O	2.54	2.43–2.64	0.05
F	0.30	0.20–0.37	0.05
H ₂ O	2.96	–	–
O = F	<u>-0.13</u>	–	–
Total	100.67	–	–

* FeO determined by electron-microprobe analysis.

TABLE 3. Powder-diffraction pattern for maruyamaite

I_{rel}	$d_{(meas.)}$ Å	$d_{(calc.)}$ Å	h	k	l	I_{rel}	$d_{(meas.)}$ Å	$d_{(calc.)}$ Å	h	k	l
23	6.415	6.404	-1	1	1	15 B	2.132*	2.135	-3	3	3
21	5.002	4.994	0	2	1			2.135	0	3	3
20	4.610	4.605	0	3	0	54	2.046	2.046	-1	6	2
59	4.237	4.233	-2	3	1	7	1.994	1.994	-4	8	0
69	3.995	3.988	-2	4	0	36	1.923	1.923	-3	7	2
42	3.498	3.496	0	1	2	6	1.883	1.882	-1	5	3
12	3.389	3.386	-1	4	1	10	1.852	1.852	-6	8	1
2	3.204	3.202	-2	2	2	2 B	1.825*	1.820	-6	7	2
3	3.113	3.116	-4	4	1	6 B	1.787*	1.792	-1	1	4
85	2.974	2.972	-1	3	2			1.785	-3	6	3
12	2.904	2.903	-3	5	1	3 B	1.746	1.748	0	2	4
10	2.634	2.629	-3	4	2	5	1.692	1.693	-2	8	2
100	2.581	2.581	0	5	1	21	1.664	1.665	-6	6	3
2	2.495	2.497	0	4	2			1.665	0	6	3
3	2.456	2.456	-2	6	1	18	1.644	1.644	-2	9	1
14 B	2.386*	2.409	0	0	3	22	1.595	1.595	-5	10	0
		2.383	-2	5	2	5	1.547	1.548	4	-10	1
19	2.347	2.347	-5	6	1	11	1.530*	1.535	0	9	0
4 B	2.305*	2.306	-1	2	3			1.529	-7	9	2
		2.303	0	6	0	19	1.511	1.512	0	5	4
11	2.196	2.195	-5	5	2	4	1.485	1.486	-2	6	4
18	2.168	2.167	-4	7	1	15	1.458*	1.461	-5	6	4

{note to typesetting: minus signs are overbars in above table}

TABLE 4. Miscellaneous information for maruyamaite

<i>a</i> (Å)	15.955(10)	crystal size (μm)	30 x 60 x 80
<i>c</i>	7.227(4)	radiation	MoKα
<i>V</i> (Å ³)	1593(2)	No. unique reflections	1149
		No. $ F_o > 5\sigma F$	1149
Space group	<i>R3m</i>	R_{int} %	1.02
<i>Z</i>	3	R_{obs} %	1.58
$D_{calc.}$ (gcm ⁻³)	3.081	wR_2 %	3.98
		GOF	1.131

TABLE 5. Atom coordinates and displacement parameters for maruyamaite

Atom	<i>x</i>	<i>y</i>	<i>z</i>	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}	U_{eq}
X	0	0	0.2206(2)	0.0154(6)	0.0154(6)	0.0227(8)	0	0	0.0077(3)	0.0178(5)
Y	0.12420(5)	0.06210(3)	0.63605(12)	0.0105(4)	0.0079(3)	0.0176(4)	-0.00237(12)	-0.0048(2)	0.00526(19)	0.0117(2)
Z	0.29818(4)	0.26169(4)	0.61244(11)	0.0068(2)	0.0066(3)	0.0071(2)	0.00016(18)	-0.00038(18)	0.00330(19)	0.00684(13)
T	0.19192(3)	0.19010(3)	1.00	0.0055(2)	0.0055(2)	0.0077(2)	-0.00091(16)	-0.00071(17)	0.00256(16)	0.00633(12)
B	0.10985(11)	0.2197(2)	0.4561(4)	0.0078(8)	0.0079(12)	0.0082(11)	0.0006(9)	0.0003(4)	0.0040(6)	0.0079(5)
O1	0	0	0.7753(5)	0.0151(10)	0.0151(10)	0.0106(15)	0	0	0.0075(5)	0.0136(7)
O2	0.06089(8)	0.12178(15)	0.4875(3)	0.0138(7)	0.0075(9)	0.0175(10)	0.0014(7)	0.0007(3)	0.0037(5)	0.0136(4)
O3	0.26395(18)	0.13197(9)	0.5127(3)	0.0282(12)	0.0148(7)	0.0069(8)	-0.0000(4)	-0.0000(8)	0.0141(6)	0.0151(4)
O4	0.09278(8)	0.18556(15)	0.0708(3)	0.0083(6)	0.0145(10)	0.0114(8)	-0.0016(7)	-0.0008(4)	0.0073(5)	0.0107(4)
O5	0.18438(16)	0.09219(8)	0.0920(3)	0.0153(10)	0.0077(6)	0.0119(8)	0.0002(3)	0.0003(7)	0.0077(5)	0.0108(4)
O6	0.19555(10)	0.18571(10)	0.7784(2)	0.0114(6)	0.0112(6)	0.0073(6)	-0.0013(5)	-0.0008(5)	0.0062(5)	0.0098(3)
O7	0.28470(10)	0.28492(9)	0.07961(19)	0.0073(6)	0.0069(6)	0.0104(6)	-0.0014(5)	-0.0015(5)	0.0014(5)	0.0091(3)
O8	0.20934(10)	0.26999(11)	0.4420(2)	0.0056(6)	0.0101(6)	0.0161(6)	0.0022(5)	0.0005(5)	0.0039(5)	0.0106(3)

TABLE 6. Selected interatomic distances (Å) in maruyamaite

X-O2	2.560(2)	Z-O3	1.993(1)
X-O4	2.783(2)	Z-O6	1.898(1)
X-O5	<u>2.712(2)</u>	Z-O7	1.907(1)
<X-O>	2.685	Z-O7	1.963(1)
		Z-O8	1.900(1)
T-O4	1.629(1)	Z-O8	<u>1.932(1)</u>
T-O5	1.646(1)	<Z-O>	1.932
T-O6	1.606(1)		
T-O7	<u>1.604(1)</u>	Y-O1	1.990(2)
<T-O>	1.621	Y-O2 x2	2.011(1)
		Y-O3	2.127(2)
B-O2	1.372(3)	Y-O6 x2	<u>1.999(1)</u>
B-O8 x2	<u>1.379(2)</u>	<Y-O>	2.023
<B-O>	<u>1.377</u>		

TABLE 7. Site-scattering values (*epfu*) and assigned site occupancies in maruyamaite

Site	Site population (<i>apfu</i>) from EMPA	Site scattering (<i>epfu</i>)	
		refined	calculated
X	0.53 K + 0.19 Na + 0.26 Ca	16.6(1)	17.3
Y	1.19 Mg + 0.55 Fe ²⁺ + 0.05 Fe ³⁺ + 1.07 Al + 0.14 Ti	46.9(6)	46.4
Z	5.00 Al + 1.00 Mg	78*	77
T	5.97 Si + 0.03 Al	84*	84

* fixed.

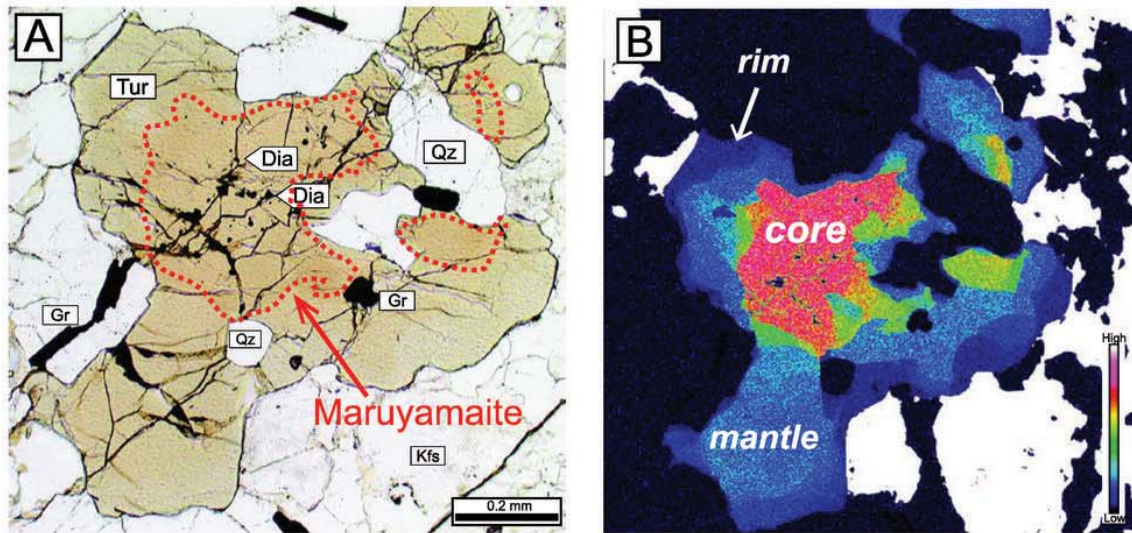


FIGURE 1

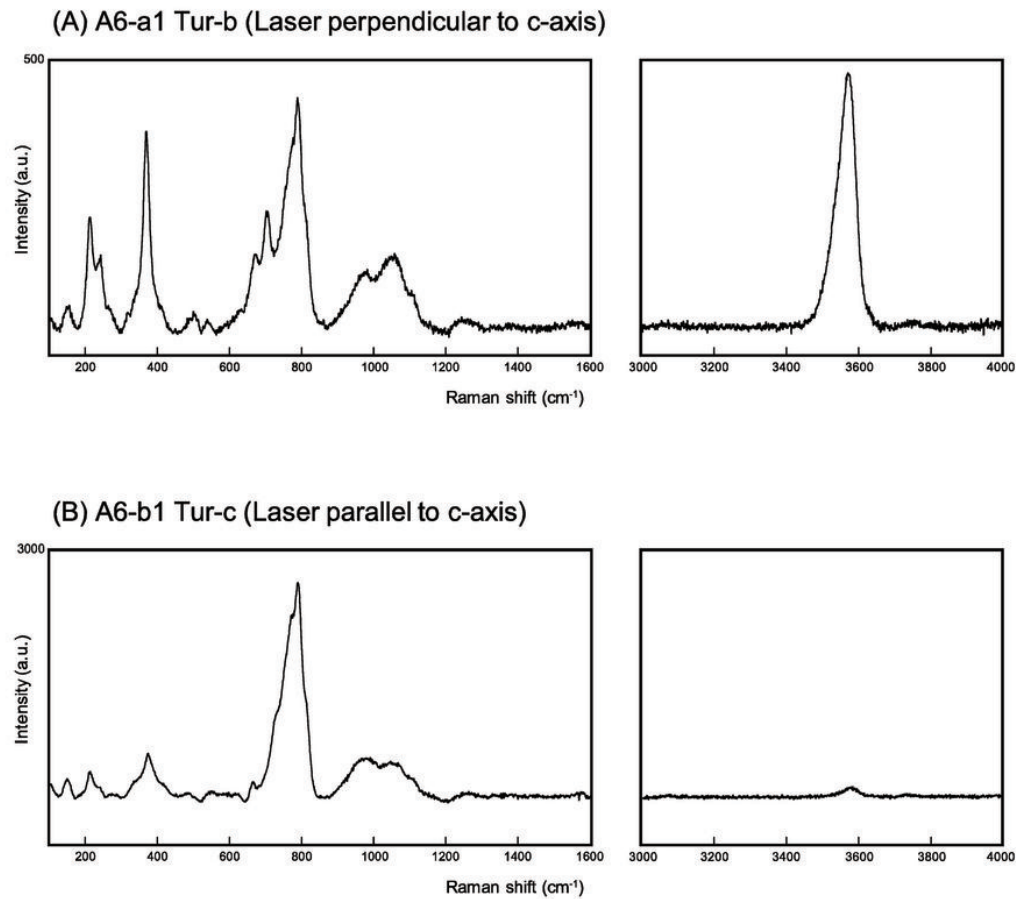


FIGURE 2

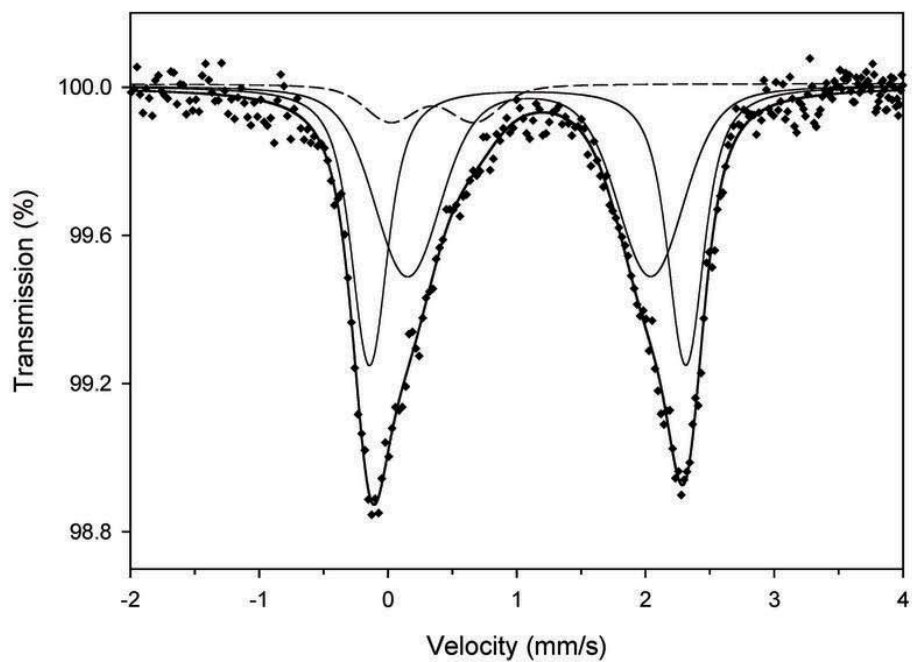


FIGURE 3

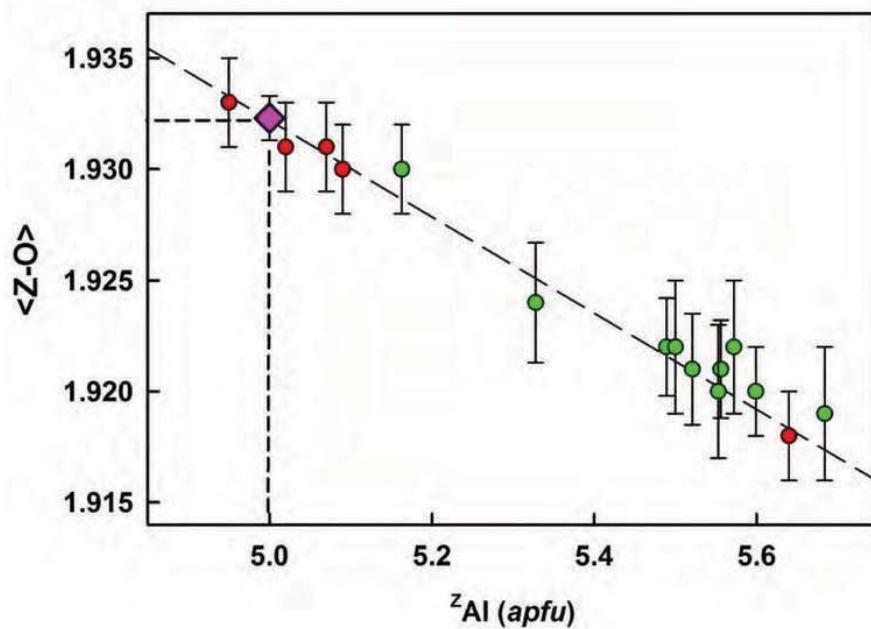


FIGURE 4