

1 **Investigation of the crystal structure of low water content hydrous olivine to 29.9 GPa: a high-**
2 **pressure single-crystal X-ray diffraction study**

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12

13 **Abstract**

14 Olivine is the most abundant mineral in the Earth's upper mantle and subducting slabs. Studying
15 the structural evolution and equation of state of olivine at high-pressure is of fundamental
16 importance in constraining the composition and structure of these regions. Hydrogen can be
17 incorporated into olivine and significantly influence its physical and chemical properties.
18 Previous infrared and Raman spectroscopic studies indicated that local structural changes occur
19 in Mg-rich hydrous olivine ($Fo \geq 95$; 4883-9000 ppmw water) at high-pressure. Water contents
20 of natural olivine are commonly less than 1000 ppmw, it is thus inevitable to investigate the
21 effects of such water contents on the equation of state (EoS) and structure of olivine at high-
22 pressure. Here we synthesized low water content hydrous olivine (Fo_{95} ; 1538 ppmw water) at
23 low SiO_2 activity and identified that the incorporated hydrogens are predominantly associated
24 with the Si sites. We performed high-pressure single-crystal X-ray diffraction experiments on
25 this olivine to 29.9 GPa. A third-order Birch-Murnaghan equation of state (BM3 EoS) was fit to
26 the pressure-volume data, yielding the following EoS parameters: $V_{T0} = 290.182(1) \text{ \AA}^3$, $K_{T0} =$
27 $130.8(9) \text{ GPa}$, and $K'_{T0} = 4.16(8)$. The K_{T0} is consistent with those of anhydrous Mg-rich olivine,
28 which indicates that such low water content has negligible effects on the bulk modulus of
29 olivine. Furthermore, we carried out the structural refinement of this hydrous olivine as a
30 function of pressure to 29.9 GPa. The results indicate that, similar to the anhydrous olivine, the
31 compression of the M1-O and M2-O bonds are comparable, which are larger than that of the Si-O
32 bonds. The compression of M1-O and M2-O bonds of this hydrous olivine are comparable with
33 those of anhydrous olivine, while the Si-O1 and Si-O2 bonds in the hydrous olivine are more
34 compressible than those in the anhydrous olivine. Therefore, this study suggests that low water

35 content has negligible effects on the EoS of olivine, though the incorporation of water softens the
36 Si-O1 and Si-O2 bond.

37 **1. Introduction**

38 The incorporation of water into mantle nominally anhydrous minerals (NAMs) has been a hot
39 topic of geophysics and geochemistry, as it strongly influences the physical and chemical
40 properties of NAMs. Among the main mineral phases in the upper mantle, olivine is undoubtedly
41 the center of the topic, due to its large abundance in the mantle (Ita and Stixrude 1992). Natural
42 mantle-derived olivine can contain a few to hundreds of ppmw of water (e.g., Bell and Rossman
43 1992; Beran and Libowitzky 2006; Peslier 2010). Several factors affect the water solubility of
44 olivine, including pressure, temperature, oxygen fugacity, water fugacity, and silica activity (e.g.,
45 Fei et al. 2018; Kohlstedt et al. 1996; Mosenfelder et al. 2006; Qin et al. 2018; Smyth et al.
46 2006). Experimental studies have suggested that the maximum is as high as 8900 ppmw (Smyth
47 et al. 2006).

48 The incorporation of hydrogen in olivine and its effects on the various physical and chemical
49 properties have long been investigated (e.g., Jacobsen et al. 2008; Mao et al. 2010; Chen et al.
50 2011; Ghosh et al. 2013; Manghnani et al. 2013; Wang et al. 2019). Vibrational spectroscopy
51 (such as infrared spectroscopy) has been mostly employed to qualitatively and quantitatively
52 determine the extent of incorporation of hydrogen. In a Fourier transform infrared spectroscopy
53 (FTIR) spectrum, the O-H vibrational bands are within the region of 3000-3700 cm^{-1} . However,
54 the mechanisms of hydrogen incorporation, as constrained by the locations of these vibrational
55 bands have long been controversial. The most significant controversy concerns which
56 substitution mechanism accounts for the high-frequency O-H bands above 3450 cm^{-1} . Some of

57 the previous studies have interpreted these bands in terms of the 2H^+ for Mg^{2+} substitution
58 associated with the Mg vacancies, based on the polyhedral O-O edge lengths (e.g., Kudoh et al.
59 2006; Smyth et al. 2006; Hushur et al. 2009; Manghnani et al. 2013). However, other studies
60 have attributed these bands to the 4H^+ for Si^{4+} substitution associated with the Si vacancies,
61 based on the compositional effects on the incorporation of water (e.g., Matveev et al. 2001;
62 Berry et al. 2005; Berry et al. 2007; Kovács et al. 2010).

63 Recently, combined nuclear magnetic resonance (NMR), vibrational spectroscopy and first-
64 principles calculation studies have led to a conclusion that the high-frequency O-H bands above
65 3450 cm^{-1} in olivine are due to the 4H^+ for Si^{4+} substitution associated with the Si vacancies,
66 which is the predominant hydration mechanism in olivine, and the 2H^+ for Mg^{2+} substitution
67 associated with the Mg vacancies is responsible for the low-frequency ($< 3400\text{ cm}^{-1}$) O-H bands
68 (e.g., Balan et al. 2011; Umemoto et al. 2011; Balan et al. 2017; Xue et al. 2017).

69 However, most of these studies on the hydrogen incorporation mechanism were performed at
70 ambient conditions. Very recently, an in situ High-pressure FTIR study on hydrous olivine has
71 revealed hydrogen transfer between the Si storage sites with pressure (Yang et al. 2019).

72 Therefore, it is important to investigate the effects of the hydrogen transfer on the structure and
73 equation of state (EoS) of olivine at high-pressure. However, compared to anhydrous olivine
74 (Mao et al. 1970; Liu 1975; Durben et al. 1993; Liu and Mernagh 1993; Andrault et al. 1995;
75 Downs et al. 1996; Zha et al. 1998; Zhang 1998; Rouquette et al. 2008; Nestola et al. 2011;
76 Finkelstein et al. 2014; Zhang et al. 2017b; Angel et al. 2018; Zhang et al. 2019), high-pressure
77 studies on hydrous olivine are relatively limited (e.g., Manghnani et al. 2013). At ambient
78 temperature, high-pressure powder X-ray diffraction (PXRD) experiments revealed that olivine
79 containing several thousand ppmw water retains its structure to $\sim 34\text{ GPa}$. However, high-

80 pressure Raman spectroscopy (RS) has detected subtle discontinuous changes around 20 GPa
81 (Hushur et al. 2009; Manghnani et al. 2013).

82 Previous studies suggested that the incorporation of several thousand ppmw water (≥ 4883
83 ppmw) has effects on the elasticity and EoS of olivine (e.g., Jacobsen et al. 2008; Hushur et al.
84 2009; Mao et al. 2010; Manghnani et al. 2013); however, the water contents of natural mantle
85 olivine are commonly less than 1000 ppmw (e.g., Peslier 2010). Therefore, it is inevitable to
86 investigate the effects of relatively low water content on the EoS and structure of olivine at high-
87 pressure. High-pressure single-crystal X-ray diffraction (SCXRD) is a powerful tool for
88 investigating the pressure-induced changes in the crystal structure and provides the most reliable
89 unit-cell parameter data to determine the EoS (Angel et al. 2000; Dubrovinsky et al. 2010). With
90 high-pressure SCXRD, one can determine the effects of pressure on the compression of the
91 individual coordination polyhedron, and thus examine the effects of incorporation of hydrogen
92 associated with cationic vacancies. As a result, SCXRD may be able to investigate the effects of
93 minor water content on the olivine structure at high-pressure. Here we report a high-pressure
94 SCXRD study to investigate the EoS and structural evolution of low water content hydrous Mg-
95 rich olivine (Fo₉₅; 1538 ppmw water). Furthermore, the effects of water on the EoS and high-
96 pressure structure of olivine will be discussed.

97 **2. Materials and Methods**

98 **2.1. Synthesis of hydrous olivine**

99 The olivine used in this study was synthesized by the method of high-pressure solid-solid
100 reactions using a multi-anvil pressure apparatus at the Institute of Geochemistry, Chinese
101 Academy of Sciences, Guiyang, China. The sample assembly was similar to the authors'

102 previous study (Fan et al. 2017; Xu et al. 2018). To obtain hydrous olivine, we used an
103 omphacite + brucite ($\text{Mg}(\text{OH})_2$) mixture as the starting material. Several natural omphacite
104 crystals with grain sizes of $\sim 200\text{-}400\ \mu\text{m}$, were selected from a crushed large eclogitic omphacite.
105 The brucite powder was used as the water source surrounding omphacite crystals in the
106 experimental platinum capsule. We used a Ni foil as the oxygen buffer in the synthesis (Rauch
107 and Keppler 2002; Xu et al. 2018). This sample assembly allowed olivine to grow at low SiO_2
108 activity. The sample was first compressed to 4.0 GPa over 35 minutes and then heated to $1200\ ^\circ\text{C}$
109 in 30 minutes. After a run duration of 24 hours, the olivine crystals ($100\text{-}400\ \mu\text{m}$ size) were
110 obtained from the quenched run product.

111 **2.2. Chemical and FTIR Analysis**

112 Selected crystals with sizes larger than $\sim 100\ \mu\text{m}$ were used for electron microprobe analysis
113 (EMPA). Analyses were performed with a JEOL Hyperprobe JXA-8500F microscope, operating
114 at a 15 kV accelerating voltage, 20 nA beam current, and the beam size of $10\ \mu\text{m}$. The empirical
115 chemical formula was calculated as $\text{Mg}_{1.904\pm 9}\text{Ni}_{0.089\pm 9}\text{Fe}_{0.015\pm 3}\text{Si}_{0.991\pm 1}\text{O}_4$ based on the EMPA data
116 (Table S1). The composition of the sample expressed in end-member molar percentages is
117 $\text{Fo}_{95}\text{Lie}_5$, where Fo and Lie are forsterite (Mg_2SiO_4) and liebenbergite (Ni_2SiO_4), respectively.
118 Unpolarized FTIR was employed to analyze the incorporation of water in the olivine. We
119 obtained two crystals without any inclusions or fractures from the quenched products for the
120 FTIR analysis, and the grain size was $\sim 400\ \mu\text{m}$. The experimental details can be seen in Xu et al.
121 (2018). The obtained FTIR spectra (Figure 1) were very similar to previous studies on synthetic
122 olivine (Lemaire et al. 2004; Smyth et al. 2006) and indicated that the sample has four strong
123 peaks at 3612 , 3577 , 3565 and $3555\ \text{cm}^{-1}$, and a weaker peak at $3477\ \text{cm}^{-1}$. These peaks are

124 associated with the Si site (Kohlstedt et al. 1996; Lemaire et al. 2004; Berry et al. 2005; Berry et
125 al. 2007; Walker et al. 2007; Xue et al. 2017). Additionally, a broad peak located between 3100
126 and 3400 cm^{-1} confirmed that the 2H^+ for Mg^{2+} substitution also takes place in the synthetic
127 olivine, but its extent is rather insignificant, relative to that at Si sites (Lemaire et al. 2004; Xue
128 et al. 2017). The water content was estimated from integrated absorbances using the calibration
129 of Bell et al. (2003), and the result was 1538(37) ppmw.

130 **2.3. Single-crystal X-ray diffraction at high-pressure**

131 An olivine crystal with size ca. $40 \times 35 \times 7 \mu\text{m}$ was selected from the crushed FTIR sample,
132 which was mounted onto a polymer holder for ambient SCXRD measurement. Then, the same
133 sample was loaded into a short symmetric diamond anvil cell (DAC) with two Type-I diamonds
134 ($300 \mu\text{m}$ culets) mounted on Boehler-Almax-type WC seats, and this sample assembly allowed a
135 $\pm 32^\circ$ opening angle. A rhenium gasket was indented to a thickness of $\sim 40 \mu\text{m}$ by the diamond
136 anvils, and a $180\text{-}\mu\text{m}$ sample chamber was cut using a laser drilling following indentation. Gold
137 powder was loaded as pressure marker (Fei et al. 2007). A small ruby sphere was loaded as the
138 pressure indicator for the gas-loading with neon (Rivers et al. 2008).

139 Ambient and high-pressure SCXRD experiments were carried out with a six-circle
140 diffractometer at the experimental station 13-BM-C of the Advanced Photon Source, Argonne
141 National Laboratory. The experimental details can be seen in previous studies (e.g., Xu et al.
142 2017a; Zhang et al. 2017a). To obtain precise and sufficient data to constrain the unit-cell
143 evolution with pressure, we collected the diffraction data at 40 different pressures over 0-29.9
144 GPa (Table S2), and at least 150 reflections (Figure 2) were used to refine the unit-cell
145 parameters at each pressure point. In addition, we collected diffraction data with increased

146 coverage/more reflections (at least 550) with multiple detector positions at 11 different pressures
147 for full structure determination (Table S3). The refinement of the unit-cell parameters and the
148 data reduction were completed with the GSE_ADA/RSV software package (Dera et al. 2013).
149 Structure refinements at various pressures were carried out with SHELXL, Olex2, and VESTA
150 software packages (Sheldrick 2008; Dolomanov et al. 2009; Momma and Izumi 2011). We
151 employed a previously reported olivine structure (Nord et al. 1982) as the initial model of the
152 structure refinement. In the olivine structure (M1M2TO₄), there are two non-equivalent
153 octahedral sites M1 and M2. In the ambient structural refinement, the T sites were fully occupied
154 by Si, while the M1 and M2 sites are fully occupied by a mixture of Mg and Ni with a refinable
155 ratio. We did not include Fe in the refinement as it has a similar electron number with Ni and its
156 insignificant content compared to Mg and Ni. Atoms sharing the same site were set to share the
157 same fractional coordinates and atomic displacement parameters (ADP). Anisotropic ADP was
158 applied to all atoms. Ambient structure refinement led to the Mg/Ni ratios of 91/9 and 98/2 for
159 M1 and M2, respectively, which was very consistent with the results of the chemical analysis. In
160 the high-pressure structural refinements, the Mg/Ni ratios of M1 and M2 were fixed at 91/9 and
161 98/2, respectively, and isotropic ADP was applied to all atoms, due to the limited opening angle
162 of the DAC. Unit-cell parameters, refinement details, atomic coordinates, as well as calculated
163 polyhedral parameters including bond length and volume are listed in Tables S(2-6). The cif files
164 can be obtained in the supplementary material.

165 **3. Results and discussion**

166 **3.1. Equation of state and compressional behavior of coordination polyhedral geometry**

167 Upon compression, olivine retained its initial structure to the maximum pressure of 29.9(2) GPa.
168 The unit-cell parameters of olivine decreased continuously with increasing pressure over the
169 experimental range of 0-29.9 GPa, as shown in Figure 3. The pressure-volume (P - V) data were
170 fit without any constraints, using a third-order Birch-Murnaghan (BM3) EoS (Birch 1947) using
171 the program EoSFit7c (Angel et al. 2014; Gonzalez-Platas et al. 2016). The obtained EoS
172 parameters including zero- P unit-cell volume (V_0), isothermal bulk modulus (K_{T0}), and its
173 pressure derivative (K'_{T0}) were $V_0 = 290.182(1) \text{ \AA}^3$, $K_{T0} = 130.8(9) \text{ GPa}$, and $K'_{T0} = 4.16(8)$,
174 respectively. We also determined the EoS for each unit-cell edge using a BM3 EoS. The results
175 are reported in Table S7, which shows that the b direction is the softest, while a is the stiffest; the
176 axial compressibility scheme is $\beta_b > \beta_c > \beta_a$. The F_E - f_E plot (Birch 1978; Angel 2000) is shown
177 in Figure 4, which shows that data for a , b , c , and V lie on inclined straight lines, and a weighted
178 linear fit of the F_E - f_E data yielded K_{T0} and K'_{T0} , which are in excellent agreement with those
179 indicated by the BM3 EoS fit (Table S7), indicating that the BM3 EoS is a reasonable choice to
180 fit the high-pressure data.

181 To better understand the compression mechanism, we analyzed the polyhedral evolution with
182 pressure using the structure refinements at high-pressure. As in previous high-pressure studies on
183 olivine and other mantle major minerals such as pyroxene (Zhang et al. 1997; Periotto et al.
184 2012; Xu et al. 2017b), in the hydrous olivine, the SiO_4 tetrahedron was much more
185 incompressible than the M1O_6 and M2O_6 octahedra (Figure 5). As shown in Figure 5, polyhedral
186 volumes of M1O_6 and M2O_6 underwent nearly linear compression, accompanying the decrease
187 of the unit-cell volume. M2O_6 is slightly more compressible than M1O_6 . The SiO_4 tetrahedron is
188 much stiffer than M1O_6 and M2O_6 . The compressional behavior of the M1O_6 and M2O_6 is very
189 close to that of the unit-cell volume (Figure 5), thus control the bulk compression of the olivine

190 structure. The bond lengths of M1-O, M2-O, and Si-O as a function of pressure are shown in
191 Figures (6-8). During compression from 0 to 29.9 GPa, the mean bond length of M1-O and M2-
192 O shortened by 5 and 7%, respectively. By comparison, the Si-O bonds were much stiffer, only
193 shortened by 2% over the experimental pressure range.

194 **3.2. Effects of water on compression behavior of olivine**

195 We compared the EoS results of the hydrous olivine in this study to those of Mg-rich ($Fo \geq 95$)
196 olivine from previous studies. To make the comparison reasonable, only the data collected using
197 high-pressure SCXRD from the previous studies were considered. The EoS determination of
198 Hazen (1976) was based on only four pressure measurements to a maximum pressure of 5 GPa,
199 and such data coverage should be insufficient to allow robust refinements of three EoS
200 parameters (V_{T0} , K_{T0} , and K'_{T0}), thus we did not consider it. The EoS study of Kudoh and
201 Takéuchi (1985) was based on 7 pressure measurements within 3.1-14.9 GPa, as they used a 4:1
202 mixture of methanol and ethanol as the pressure medium, and only five data (3.1-8.6 GPa) were
203 collected under hydrostatic conditions (Angel et al. 2007). Therefore, for the same reason
204 mentioned above, we did not consider the EoS results of Kudoh and Takéuchi (1985) as well.
205 Finally, the EoS parameters of anhydrous Fo_{100} obtained by Downs et al. (1996), Zhang (1998),
206 Poe et al. (2010) and Finkelstein et al. (2014) were considered. Due to the effects of the trade-off
207 between the K_{T0} and K'_{T0} , we refitted the P - V data of the previous studies (Downs et al. 1996;
208 Zhang 1998; Poe et al. 2010; Finkelstein et al. 2014) and this study with fixed $K'_{T0} = 4.2$ that is
209 the value for the anhydrous Fo_{100} suggested by a high-pressure Brillouin Light Scattering study
210 (Zha et al. 1996). By doing so, the fitting of the P - V data of the previous studies obtained K_{T0} of
211 124(3)-130.4(4) GPa, which are comparable to the K_{T0} derived from the fitting of the P - V data in
212 this study (130.5(2) GPa). Therefore, low water content (1538 ppmw) could have negligible

213 effects on the K_{70} of olivine. However, previous studies on hydrous olivine with water contents
214 larger than or equal to 4883 ppmw suggested that such water contents reduce the bulk modulus
215 of olivine (Jacobsen et al. 2008; Mao et al. 2010; Manghnani et al. 2013).

216 We also compared the compressions of bond lengths and polyhedral volumes of hydrous olivine
217 in this study to those of anhydrous olivine from previous studies. As shown in Figures (6-9), the
218 hydrous olivine Fo₉₅ in this study and anhydrous Fo₁₀₀ from Pamato et al. (2019) are in good
219 agreement in the compression trends of M1-O, M2-O, and Si-O bonds. However, the
220 compressions of bond lengths of anhydrous Fo₁₀₀ from earlier high-pressure SCXRD studies
221 (Hazen 1976; Kudoh and Takéuchi 1985) deviate the trends determined by this study and Pamato
222 et al. (2019), which is possibly caused by the lower accuracy of diffractometers decades ago, as
223 indicated by the larger uncertainties of the unit-cell parameters and bond lengths (Figures 3 and
224 (6-8)). Therefore, we only further compared our results of the bond lengths to those of anhydrous
225 Fo₁₀₀ from Pamato et al. (2019).

226 We analyzed the pressure-bond length and pressure-polyhedral volume data of olivine in this
227 study and Pamato et al. (2019) by using weighted linear regression (Figures (6-9) and Table S8).
228 Due to the data range of Pamato et al. (2019) was 0-7.66 GPa, only the data within the range of
229 0-10 GPa of this study were used, to make the comparisons reasonable. As shown in Figures (6-
230 7) and Table S8, the compressions of M1-O and M2-O between the anhydrous and hydrous
231 olivine are similar. The M1-O bonds of the anhydrous Fo₁₀₀ decrease near linearly with pressure
232 and the rates are 0.0044(3)-0.0054(2) Å/GPa, which are comparable with the shrinking rates of
233 the M1-O bonds for the hydrous Fo₉₅ (0.0036(3)-0.0048(2) Å/GPa; Figure 6). Similarly, the M2-
234 O bonds of these two olivines decrease with pressure at comparable rates of 0.0049(5)-0.0077(6)
235 and 0.0053(3)-0.0069(6) Å/GPa for the anhydrous Fo₁₀₀ and hydrous Fo₉₅ (Figure 7),

236 respectively. The similarity in the compressions of M1-O and M2-O bond lengths between the
237 anhydrous and hydrous olivine results in the comparable polyhedral ($M1O_6$ and $M2O_6$)
238 compressibilities between the hydrous and anhydrous olivine (Figure 9).

239 Compared with M1-O and M2-O, the Si-O bonds behave quite differently under pressure
240 between the anhydrous and hydrous olivine (Figure 8). As shown in Figure 8 and Table S8, for
241 the anhydrous Fo_{100} , the Si-O1 and Si-O2 bonds decrease at very low rates of 0.0006(7) and
242 0.0010(5) Å/GPa, respectively. Additionally, the small values of the correlation coefficients (R^2
243 = 0.394 and 0.084 for the Si-O1 and Si-O2, respectively) indicate the variation of bond length is
244 insensitive to the pressure change, while the compression of the Si-O3 is more significant as
245 indicated by the higher shrinking rate (0.0015(1) Å/GPa) and R^2 (0.936). The Si-O1 and Si-O2
246 bonds in the hydrous Fo_{95} are more compressible than those in the anhydrous Fo_{100} . As shown in
247 Figure 8 and Table S8, the bond lengths of the Si-O1 and Si-O2 of hydrous Fo_{95} decrease at rates
248 of 0.0025(2) and 0.0013(6) Å/GPa, respectively, and the higher R^2 values (0.992 and 0.731 for
249 the Si-O1 and Si-O2, respectively) also indicate that the bond length is more sensitive to the
250 pressure change than that of the anhydrous Fo_{100} . On the other hand, the shrinking rate of the Si-
251 O3 for the hydrous Fo_{95} (0.0011(1) Å/GPa) is lower than that for the anhydrous Fo_{100} (0.0015(1)
252 Å/GPa). In a view of the polyhedral compression, the SiO_4 tetrahedron in the hydrous Fo_{95} is
253 more compressible than that in the anhydrous Fo_{100} olivine (Figure 9).

254 **4. Implications**

255 Previous high-pressure spectroscopic and XRD studies suggested that the incorporation of
256 several thousand ppmw of water (≥ 4883 ppmw) causes some local structural changes at high-
257 pressure and affects the elasticity and EoS of olivine (e.g., Jacobsen et al. 2008; Hushur et al.

258 2009; Mao et al. 2010; Manghnani et al. 2013; Yang et al. 2019). However, the water contents of
259 natural olivine are commonly less than 1000 ppmw (e.g., Peslier et al. 2010; Novella et al. 2015),
260 and petrological experiments also suggested that the water storage capability of olivine in
261 peridotite in the upper mantle is lower than 2000 ppmw (Ferot and Bolfan-Casanova 2012;
262 Tenner et al. 2012). Therefore, the water contents of mantle olivine, in most cases, should be less
263 than 2000 ppmw. In this study, the hydrous olivine was with moderate water content (1538
264 ppmw) that could be more expected in real mantle olivine than those in hydrous olivine from the
265 previous studies. The EoS determination indicated that such low water content has negligible
266 effects on the bulk modulus. In addition, the detailed structural refinements at high-pressure
267 suggested that the compressions of the M1-O and M2-O bonds of the hydrous olivine in this
268 study are similar to those of anhydrous olivine from the previous study. However, the
269 compressions of the Si-O bonds were changed by the incorporation of water, and the Si-O1 and
270 Si-O2 bond in the hydrous olivine are significantly more compressible than those in the
271 anhydrous olivine. Our FTIR measurements indicated that the incorporation of hydrogen is
272 predominantly associated with the vacancies in Si sites (Figure 1), which is consistent with the
273 fact that the incorporation of water softens the Si-O bonds instead of the M1-O and M2-O bonds
274 (Figures 6-9). However, the water-induced change in the compressional behavior of the Si-O
275 bonds does not affect the bulk compression of olivine significantly. The incorporation of ~1500
276 ppmw does not significantly reduce the bulk modulus of olivine, which can be interpreted in
277 terms of that such low water content has negligible effects on the compressional wave velocity,
278 according to the Newton-Laplace equation approximation. Therefore, the results of this study
279 indicate that the effects of incorporation of water on the elasticity and EoS of olivine could be
280 significant only when the water content is at least higher than 1538 ppmw.

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477

478

479 **Figures**

480 **Figure 1.** Selected unpolarized FTIR spectra of hydrous olivine synthesized at 4 GPa and 1250
481 °C. Integration of the spectrum indicates an H₂O content of 1538 ppmw.

482 **Figure 2.** Single crystal X-ray diffraction patterns of hydrous olivine at (a). $P = 1.2(1)$ GPa and
483 (b) $P = 29.9(2)$ GPa.

484 **Figure 3.** Evolution of unit-cell parameters as a function of pressure for the hydrous olivine in
485 this study: (a) a , (b) b , (c) c , and (d) V . Mg-rich ($Fo \geq 95$) olivine from previous studies are
486 also shown for comparison (Hazen 1976; Kudoh and Takéuchi 1985; Will et al. 1986;
487 Downs et al. 1996; Zhang 1998; Poe et al. 2010; Manghnani et al. 2013; Finkelstein et al.
488 2014; Pamato et al. 2019). Solid lines correspond to the BM3 EoS of this study.

489 **Figure 4.** Eulerian strain-normalized pressure (F_E - f_E) plot (Angel 2000) ($F_E = P/3f_E (1 + 2f_E)^{5/2}$;
490 $f_E = [(V_0/V)^{2/3} - 1]/2$) of a (a) b (b), c (c), and V (d) for the hydrous olivine in this study.

491 **Figure 5.** Normalized polyhedral volume as a function of a normalized unit-cell volume at
492 different pressures. The solid line represents the $y = x$ line. The error bars were calculated
493 using the method described by Zhang et al. (2019).

494 **Figure 6.** Pressure dependence of M1-O bond lengths for the hydrous olivine in this study: (a)
495 M1-O1, (b) M1-O2, (c) M1-O3 and (d) M1-O average. Mg-rich ($Fo \geq 95$) olivine from
496 previous studies are also shown for comparison (Hazen 1976; Kudoh and Takéuchi 1985;
497 Pamato et al. 2019). The solid lines represent weighted linear regression results (Table S8)
498 for the data of Pamato et al. (2019) (red lines) and this study (black lines).

499 **Figure 7.** Pressure dependence of M2-O bond lengths for the hydrous olivine in this study: (a)
500 M2-O1, (b) M2-O2, (c) M2-O3a and M2-O3b, and (d) M2-O average. Mg-rich ($Fo \geq 95$)

501 olivine from previous studies are also shown for comparison (Hazen 1976; Kudoh and
502 Takéuchi 1985; Pamato et al. 2019). The solid lines represent weighted linear regression
503 results (Table S8) for the data of Pamato et al. (2019) (red lines) and this study (black lines)

504 **Figure 8.** Pressure dependence of Si-O bond lengths for the hydrous olivine in this study: (a) Si-
505 O1, (b) Si-O2, (c) Si-O3 and (d) Si-O average. Mg-rich ($Fo \geq 95$) olivine from previous
506 studies are also shown for comparison (Hazen 1976; Kudoh and Takéuchi 1985; Pamato et
507 al. 2019). The solid lines represent weighted linear regression results (Table S8) for the data
508 of Pamato et al. (2019) (red lines) and this study (black lines)

509 **Figure 9.** Pressure dependence of polyhedral volumes for the hydrous olivine in this study: (a)
510 $M1O_6$, (b) $M2O_6$ and (c) SiO_4 . Mg-rich ($Fo \geq 95$) olivine from the previous study is also
511 shown for comparison (Pamato et al. 2019). The solid lines represent weighted linear
512 regression results (Table S8) for the data of Pamato et al. (2019) (red lines) and this study
513 (black lines)

Figure 1

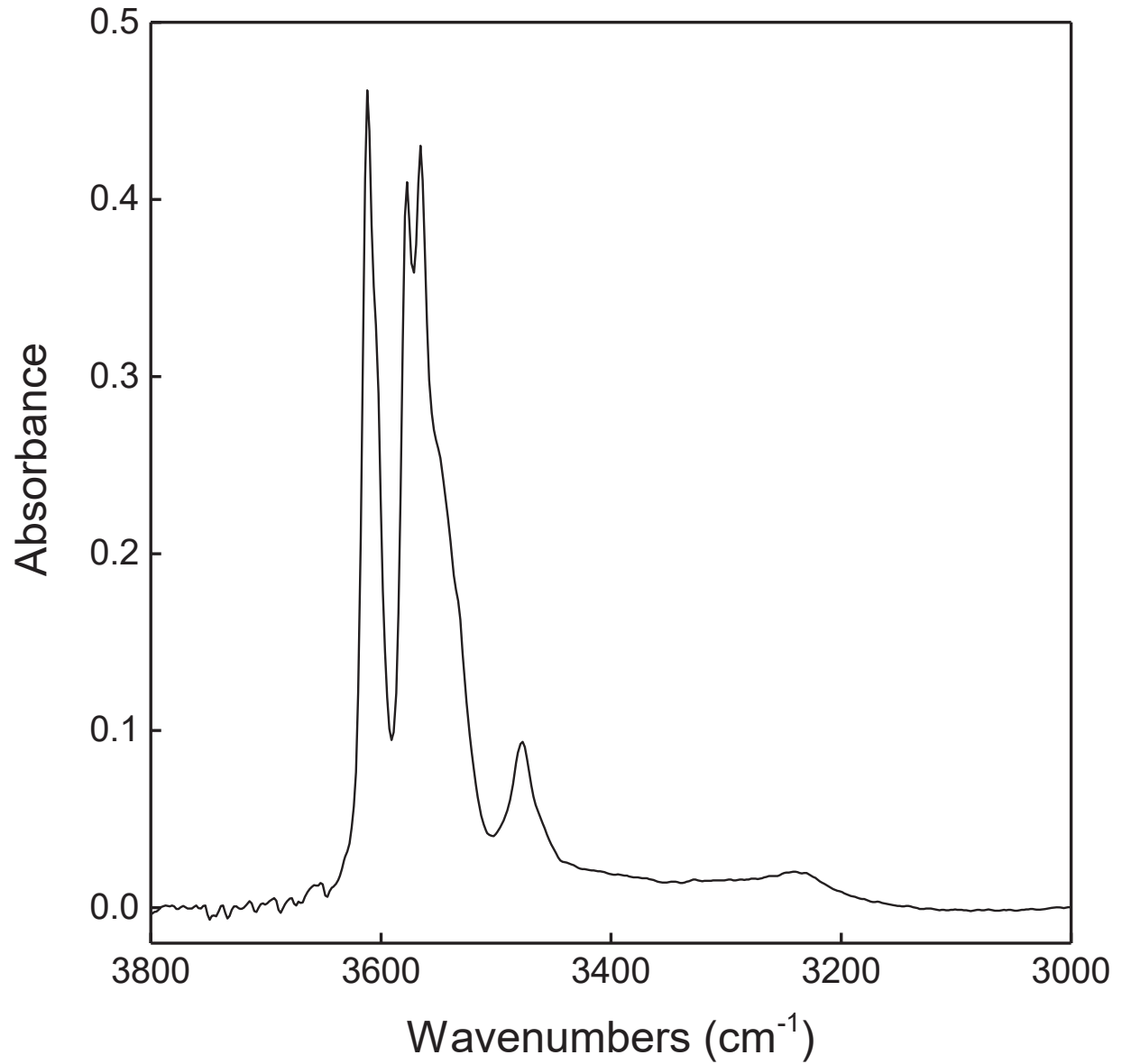


Figure 2

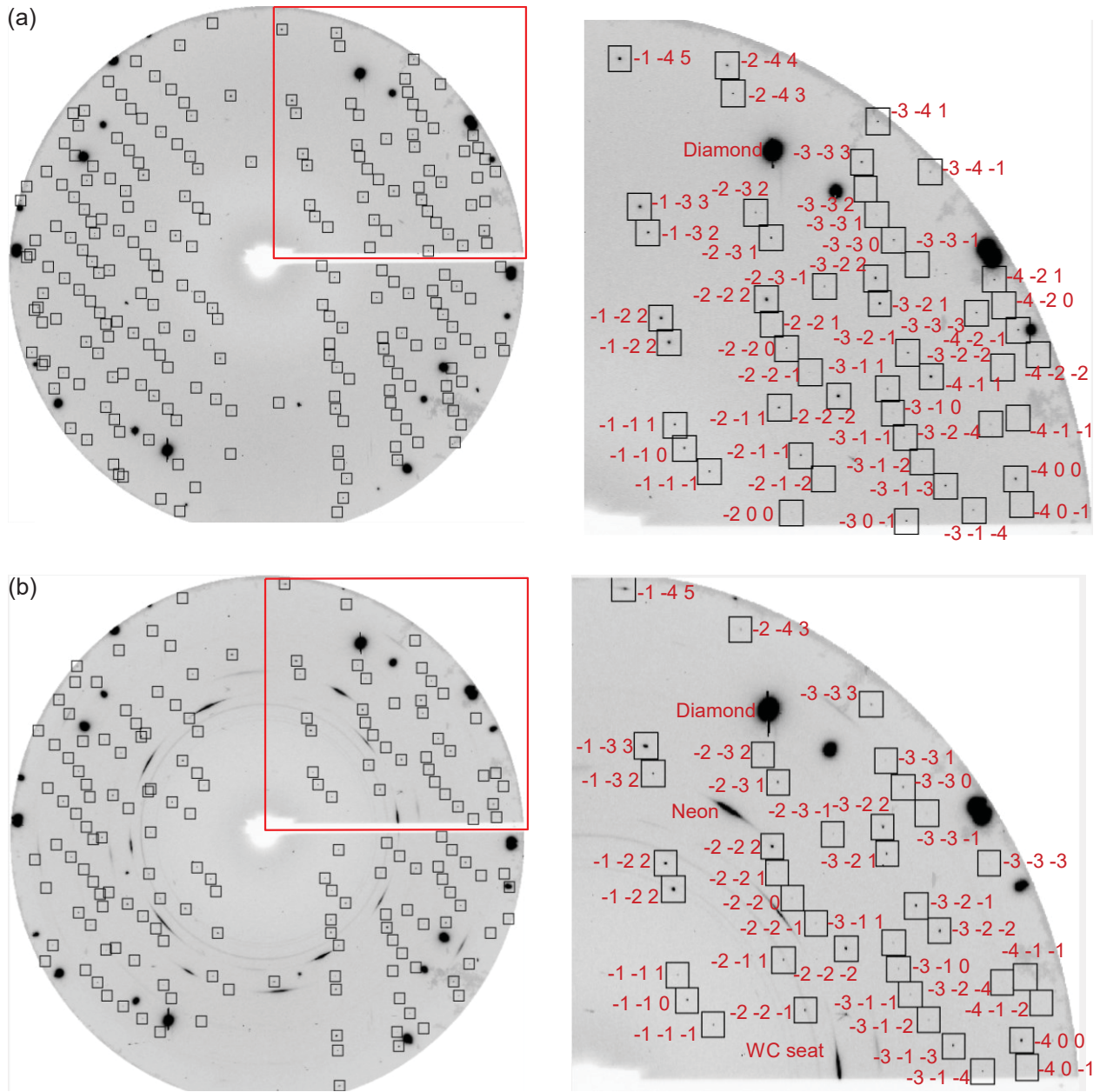


Figure 3

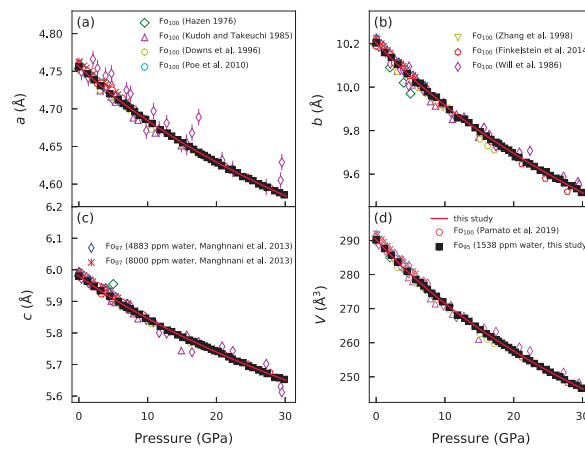


Figure 4

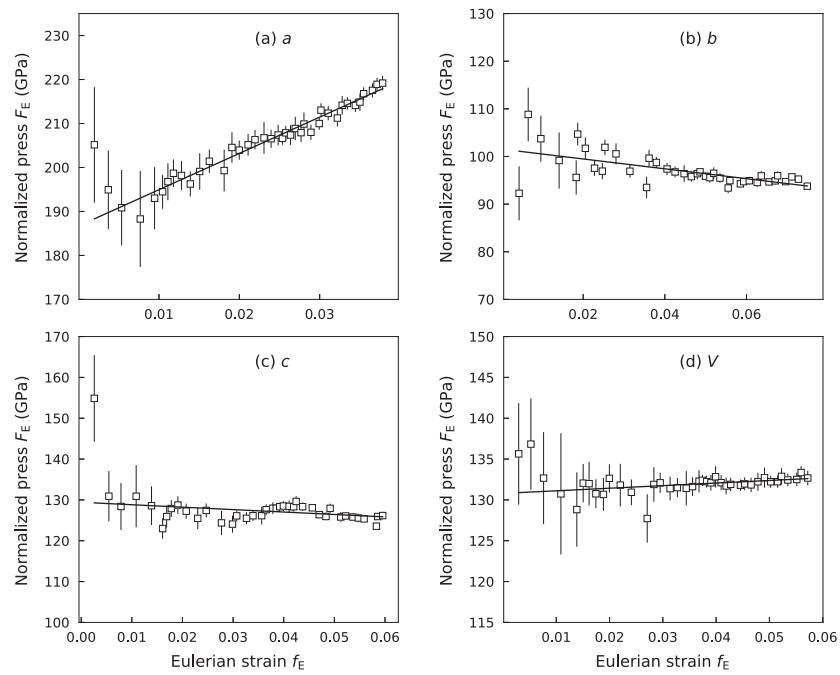


Figure 5

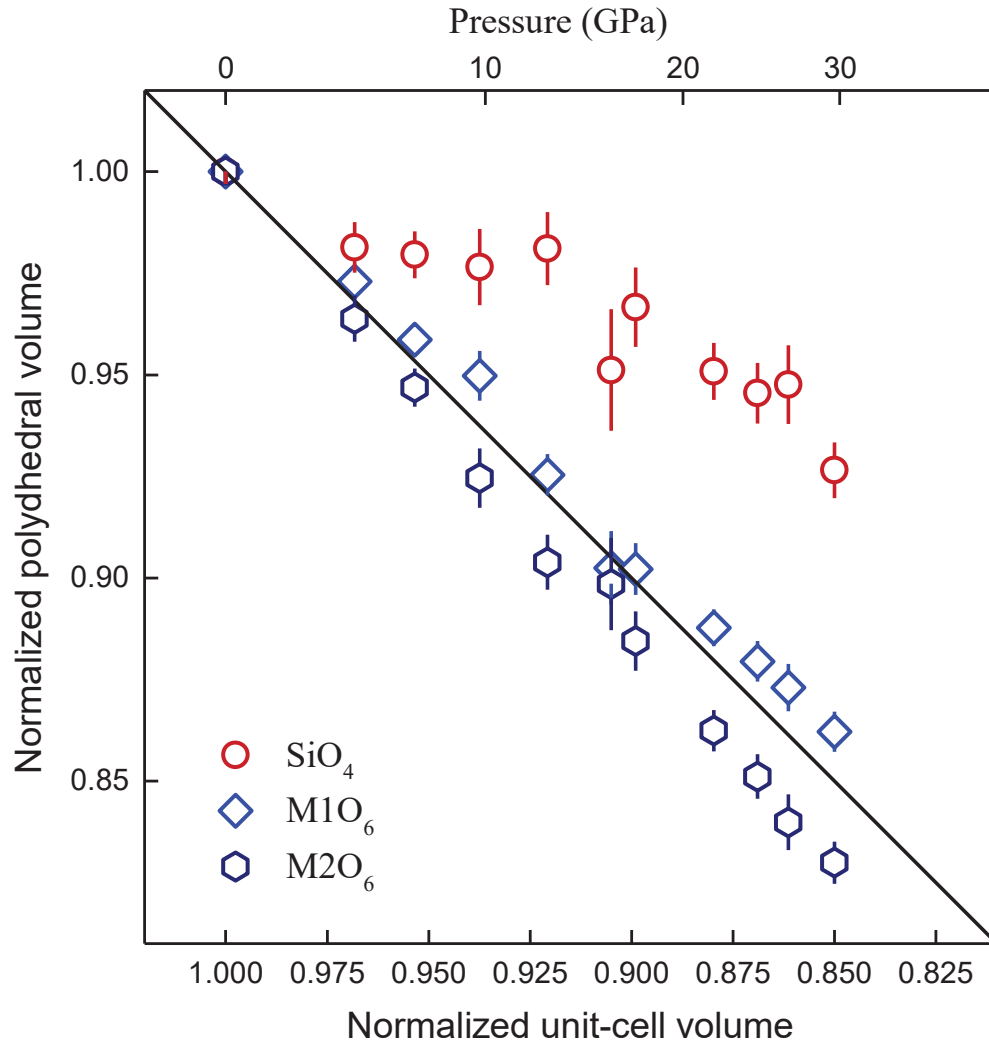


Figure 6

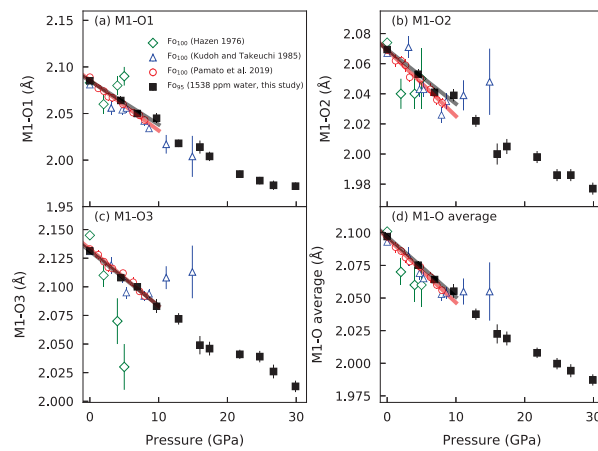


Figure 7

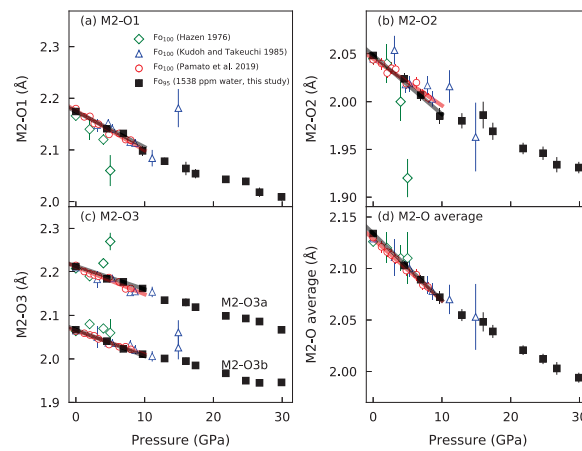


Figure 8

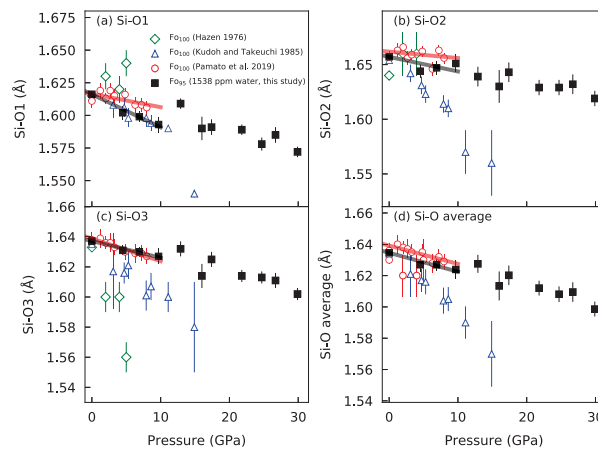


Figure 9

