

1 **Revision 2:**

2 ***P–V–T* equation of state of hydrous phase A up to 10.5 GPa**

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15 **ABSTRACT**

16 Pressure–volume–temperature (*P–V–T*) data of synthetic Mg₇Si₂O₈(OH)₆ phase A,
17 were collected under *P–T* conditions up to approximately 10.5 GPa and 900 K by
18 energy dispersive X-ray diffraction using a cubic type multi-anvil apparatus, MAX80,
19 located at the Photon Factory – Advanced Ring (PF–AR) at the High Energy
20 Accelerator Research Organization (KEK). *P–V* EoS using only room temperature
21 data yielded $V_0=511.6$ (2) Å³, $K_{T0}=106.8$ (18) GPa, and pressure derivative $K_T'=3.88$
22 (38). These properties were consistent with the subsequent equation of state (EoS)
23 analysis. The compressibility of phase A was very anisotropic along the axes, with the
24 *a*-axis as approx. 26% more compressible than the *c*-axis, which is normal to the

25 plane of the distorted close-packed layers. A fit of the present data to the
26 high-temperature Birch–Murnaghan EoS yielded $V_0=511.7$ (3) \AA^3 , $K_0=104.4$ (24) GPa,
27 $K''=4.39$ (48), $(\partial K_T/\partial T)_P = -0.027$ (5) GPa K^{-1} , and thermal expansion $\alpha=a+bT$ with
28 values of $a=2.88$ (27) $\times 10^{-5}$ K^{-1} and $b=3.54$ (68) $\times 10^{-8}$ K^{-2} . The lattice dynamical
29 approach by the Mie–Grüneisen–Debye EoS yielded $\theta_0=928$ (114) K, $q=2.9$ (10), and
30 $\gamma_0=1.19$ (8). The isobaric heat capacity C_P of phase A at 1 atm. was calculated based
31 on the Mie–Grüneisen–Debye EoS fit of present P – V – T data. In addition, the density
32 of subducting slab with different degree of serpentinization was also calculated along
33 the cold geotherm up to ~ 13 GPa. The serpentinization of subducting slab will
34 significantly lower the density of slab at shallower depth, however, this effect
35 becomes negligible when antigorite dehydrated to phase A. Because the phase A
36 bearing subducting slab is supposed to be denser than the surrounding mantle, the
37 water can transport into deeper parts of the upper mantle and the mantle transition
38 zone.

39 **Keywords:** bulk modulus, phase A, dense hydrous magnesium silicate (DHMS),
40 equation of state (EoS), heat capacity, high pressure and high temperature (HPHT),
41 thermal expansion

42 INTRODUCTION

43 Dense hydrous magnesium silicate minerals (DHMS) are potential water carriers in
44 the deep mantle. Phase A is one group of these phases that lies on the
45 forsterite–brucite interface in the $\text{MgO-SiO}_2\text{-H}_2\text{O}$ (MSH) system with an ideal

46 formula of $\text{Mg}_7\text{Si}_2\text{O}_8(\text{OH})_6$. Its water content is about 12 wt%. In a water-saturated
47 peridotite system, serpentine decomposes to phase A, clinoenstatite and water at 6.2
48 GPa, 853 K (Schmidt & Poli 1998). Ohtani et al. (2004) suggested that the serpentine
49 transformed to phase A at depth greater than 180 km (~6 GPa) in a cool subducting
50 slab, and that phase A was stable up to 11 GPa before hydrous phase E becomes the
51 main water carrier in the slab. Komabayashi et al. (2005a) conducted experiments to
52 examine antigorite bulk composition and pointed out that antigorite breaks down to
53 phase A + enstatite + water at ~5 GPa, 823 K. Komabayashi et al. (2005b) also
54 conducted experimental and theoretical study of stability of phase A and derived the
55 thermochemical parameters. Therefore, phase A is an important water carrier in the
56 subduction zone after antigorite is dehydrated. Its EoS has been of particularly
57 interest.

58 The EoS of Fe-free phase A was reported in several studies. A range of values for
59 bulk moduli (K_0) and the pressure derivative (K') has also been reported. Pawley et al.
60 (1995) investigated the thermal expansivity and compressibility of phase A in the
61 MSH system. They reported a value of $K_0 = 145$ (5) GPa (K' fixed to 4), thermal
62 expansion $\alpha_0 = 4.9$ (2) $\times 10^{-5}/\text{K}$, and obtained the temperature derivative of bulk
63 modulus $(\partial K_T/\partial T)_P = -0.049$ GPa/K using the empirical equation $(\partial K_T/\partial T)_P =$
64 $-7\alpha K_0$. Kuribayashi et al. (2003) found that $K_0 = 105$ (4) GPa and $K' = 3.9$ (8) from
65 single-crystal X-ray diffraction (XRD) up to 11.2 GPa. The Kuribayashi's bulk
66 modulus was markedly lower than the Pawley's one. Crichton and Ross (2002) also
67 measured the lattice parameters using single crystal and obtained $K_0 = 97.5$ (4) GPa,

68 $K' = 5.97$ (14). The datasets from Crichton and Ross (2002) and from Kuribayashi et
69 al. (2003) were mutually consistent when considering the tradeoff between K and K' .
70 Recently, Holl et al. (2006) studied the Fe-bearing phase A, $\text{Mg}_{6.85}\text{Fe}_{0.14}\text{Si}_2\text{O}_8(\text{OH})_6$
71 using single-crystal XRD up to 33 GPa at room temperature. They reported the value
72 of $K_0 = 102.9$ (28) GPa and $K' = 6.4$ (3). In addition, Sanchez-Valle et al. (2006, 2008)
73 determined the single crystal elastic properties of phase A by Brillouin spectroscopy,
74 and obtained the adiabatic bulk and shear moduli and the derivatives, $K_s = 106$ (1)
75 GPa, $K' = 5.8$ (3) and $\mu = 61$ (1) GPa and $\mu' = 1.8$ (1). Nevertheless, few experiments
76 were done under high-temperature conditions. No report of the relevant literature
77 describes an experiment conducted under conditions of simultaneous high pressure
78 and high temperature (HPHT).

79 To investigate the thermoelastic properties (e.g., temperature dependence of bulk
80 modulus, thermal expansion, etc.) of phase A, we conducted P - V - T experiments
81 under conditions of simultaneous HPHT on synthetic phase A by synchrotron powder
82 XRD up to ~ 10.5 GPa and 900 K. Results are compared with those of earlier studies.
83 The Mie-Grüneisen-Debye EoS of phase A was also reported.

84 **EXPERIMENTAL METHODS**

85 **Synthesis of Phase A**

86 Phase A was synthesized using powder mixture of MgO, $\text{Mg}(\text{OH})_2$ and SiO_2 , in
87 the Kawai-type (MA8-type) high pressure apparatus, ORANGE 1000, in the
88 Geodynamics Research Center (GRC), Ehime University, Japan. 14/8 type cell

89 assembly was used. That is, the edge length of the octahedron pressure medium is 14
90 mm. The truncation edge length (TEL) of the second stage tungsten carbide is 8 mm.
91 Semi-sintered MgO doped with 17% CoO was used as the pressure medium. For this
92 study, LaCrO₃ was used as a thermal insulator; Pt foil of 30 μm thickness was used as
93 both a heater and a sample container. The cell was compressed to 11 GPa and heated
94 to 1123 K for 30 minutes, and the charge was quenched by turning off the electric
95 power supply. Then the pressure was released gradually for ~10 hours.

96 The recovered sample was not well sintered. It was easily crushed to powder by
97 agate mortar to be submicron grain size. The powder XRD pattern of the recovered
98 sample showed that it is pure phase A.

99 *In-situ* X-ray diffraction

100 HPHT synchrotron XRD experiments were conducted using a 6-6 type multi-anvil
101 (6-6 type MA) assembly operated in a cubic anvil (DIA) apparatus, MAX80, at
102 beam-line AR-NE5C at the PF-AR, Tsukuba, Japan. Details of 6-6 type MA were
103 described by Nishiyama et al. (2008). Figure 1 shows the cell assemblage used for
104 *in-situ* X-ray diffraction experiments, which was the same as that used by Yang et al.
105 (2014). The powdered sample and pressure standard were placed into NaCl sleeves.
106 Cylindrical graphite was used as a furnace, whereas ZrO₂ sleeve with h-BN X-ray
107 window was used as a thermal insulator. Cubic-shaped boron epoxy was used as the
108 pressure medium. The temperature was monitored with a W_{97%}Re_{3%}-W_{75%}Re_{25%}
109 thermocouple. The junction of thermocouple was sandwiched in the center of the

110 furnace between the pressure standard and the sample. Pressure effects on the
111 electromotive force (EMF) of the thermocouple were neglected during temperature
112 measurements. A mixture of NaCl and Au with a weight ratio of 10:1 was used as a
113 pressure standard.

114 The energy dispersive X-ray diffraction method was used in the current study. The
115 incident beam was collimated to a rectangular shape with 100 μm in vertical and 200
116 μm in horizontal by slits. Data were collected using a solid-state detector (SSD)
117 connected to a multi-channel analyzer, which was calibrated with the characteristic
118 X-rays of several metal standards: Cu, Ag, Pr, Dy, W, Er, Au, Bi, Pb, Ta, and La. The
119 X-rays diffracted from the sample and pressure marker were collected at a fixed 2θ
120 angle of $\sim 6^\circ$ in the vertical direction. The 2θ angle was calibrated by the unit cell
121 volume of NaCl at ambient conditions. To minimize the pressure and temperature
122 uncertainty, XRD data of both the sample and pressure marker were collected at
123 positions as close as possible to the thermocouple junction.

124 The pressures were calculated from the EoS of NaCl proposed by Decker (1971).
125 XRD patterns were analyzed using a program, *XRayAna*, by which the unit cell
126 parameters were calculated from the peak positions using least-squares method. To
127 calculate the unit cell parameters, 5–7 diffraction peaks from NaCl and 6–11 peaks
128 from phase A were used.

129 In the P – V – T experiments, the cell was compressed to a desired load after the
130 diffraction patterns were collected at ambient conditions. Then it was heated to a

131 desired temperature to anneal the sample. The first cycle was heated to 773 K. It was
132 efficient to remove the deviatoric stress because the diffraction peaks became sharp
133 after heating. Diffraction patterns were collected at every 100 K interval during
134 cooling. After data collection at room temperature, the cell was compressed further to
135 the next desired pressure. Similar cycles for data collection were repeated. Figure 2
136 portrays the pressure–temperature paths of the P – V – T experiments. As shown in
137 Table 1, we collected 49 data points.

138 RESULTS AND DISCUSSION

139 Room temperature equation of state

140 The unit cell parameters of phase A obtained at ambient conditions were $a=7.858$
141 (4) Å, $c=9.566$ (8) Å, which were consistent with results reported from earlier studies
142 (Horiuchi et al. 1979; Pawley et al. 1995; Kagi et al. 2000; Crichton and Ross 2002).
143 Figure 3 presents unit cell volumes at room temperature against pressure. Those room
144 temperature data were fitted with the Birch–Murnaghan EoS, which is

$$145 \quad P = \frac{3K_{T0}}{2} \left[\left(\frac{V}{V_0} \right)^{-7/3} - \left(\frac{V}{V_0} \right)^{-5/3} \right] \left\{ 1 + \frac{3}{4} (K'_{T0} - 4) \left[\left(\frac{V}{V_0} \right)^{-2/3} - 1 \right] \right\}, \quad (1)$$

146 where V_0 , K_0 , and K_0' respectively denote the unit cell volume, the isothermal bulk
147 modulus, and its pressure derivative. The fitted curve is shown as the solid line in
148 Figure 3, which yields values of $V_0=511.6$ (1) Å³, $K_0=106.3$ (6) GPa, with K_0' fixed as
149 4. Comparison of the value of K_0 revealed that phase A is more compressible than
150 forsterite ($K_0= 125$ (2) GPa, Downs et al. 1996) and the hydroxyl chondrodite

151 ($K_0=115.0$ (8) GPa, Ross and Crichton 2001).

152 In Figure 3, data reported from earlier studies (Pawley et al. 1995; Crichton and
153 Ross 2002; Kuribayashi et al. 2003; Holl et al. 2006) are shown for comparison. Our
154 data show good agreement with the single-crystal diffraction data reported by
155 Crichton and Ross (2002) and by Kuribayashi et al. (2003). Although Pawley et al.
156 (1995) gave larger volume under high pressure, which might be attributable to the
157 non-hydrostatic pressure in their cell assembly, as reported by Crichton and Ross
158 (2002). Holl et al. (2006) used Fe-bearing phase A. Their data show slightly greater
159 volume and are more compressible than the Mg end-member phase A under high
160 pressure.

161 The linear compressibilities of phase A along the a -axis and c -axis are very
162 anisotropic (Fig. 4). Linear compressibility of the c -axis is $\beta_c=2.21(1) \times 10^{-3}/\text{GPa}$, and
163 is approximately 26% less compressible than the a -axis ($\beta_a=2.96(3) \times 10^{-3}/\text{GPa}$)
164 because the c -direction is normal to the closest packed oxygen array in the structure.

165 High-temperature Birch–Murnaghan equation of state

166 We used two thermal EoS to analyze the P – V – T data and to extract thermoelastic
167 parameters of phase A: high-temperature Birch–Murnaghan (HTBM) EoS and
168 Mie–Grüneisen–Debye (MGD) EoS.

169 The HTBM EoS is described again by Eq. (1), but the V_0 , K_0 , and K_0' are replaced
170 respectively by parameters at atmospheric pressure and at a constant temperature T :
171 V_{0T} , K_T , and K_T' . Temperature effects for K_T and V_{0T} are given as

172
$$K_T = K_{T_0} + (\partial K_T / \partial T)_P (T - T_0) \quad (2)$$

173
$$V_{0T} = V_0 \exp\left[\int_{T_0}^T \alpha dT\right] \quad (3)$$

174 where $(\partial K_T / \partial T)_P$, T_0 , and α respectively represent the temperature derivative of the
175 bulk modulus, the reference temperature (300 K), and the volumetric thermal
176 expansion at atmospheric pressure. Thermal expansion α is represented empirically
177 (e.g. Nishihara et al. 2005) as

178
$$\alpha(0, T) = a + bT. \quad (4)$$

179 The K_T' was assumed to be constant against temperature. Six parameters must be
180 fitted in this approach: V_0 , K_{T_0} , K_T' , $(\partial K_T / \partial T)_P$, a , and b . Table 2 presents the
181 thermoelastic parameters derived by fitting our P - V - T data to the HTBM EoS and
182 MGD EoS in addition to parameters reported in earlier studies. The data points below
183 873 K were used for the fitting because of a few data set at 973 K (See Fig. 2). The
184 root mean square (RMS) misfit in pressure in the EoS fitting is also shown in Table 2,
185 which is very low (0.075 GPa) and which is similar to the uncertainty of the
186 experimental pressure. This low misfit is related to the very good internal consistency
187 of our P - V - T data. Figure 5 presents the isothermal compression curves calculated
188 from the obtained thermoelastic parameters. Apparently, our data are well fitted.
189 Thermoelastic parameters derived from this study show good agreement with results
190 reported by Kuribayashi et al. (2003) and by Crichton and Ross (2002). The
191 temperature derivative of the bulk modulus $(\partial K_T / \partial T)_P$ was determined for the first

192 time based on a series of direct measurements. The smaller value means that the bulk
193 modulus is less dependent on the temperature than in the result reported by Pawley et
194 al. (1995), who calculated that value empirically. The volumetric thermal expansion at
195 atmospheric pressure and room temperature is $3.9 \times 10^{-5}/\text{K}$, which is slightly lower
196 than the value reported by Pawley et al. (1995).

197 Mie–Grüneisen–Debye equation of state

198 The P – V – T data were also analyzed using the Mie–Grüneisen–Debye (MGD) EoS
199 (e.g., Jackson and Rigden 1996; Tange et al. 2012). The pressure can be expressed as
200 the sum of the static pressure at room temperature $P(V, T_0)$ and thermal pressure ΔP_{th}
201 at any temperature T and given volume V (e.g. Anderson 1984). Eq. (1) was used as
202 the expression of the static pressure $P(V, T_0)$. The Debye model was used for the
203 thermal pressure part ΔP_{th} (MGD model) (e.g., Jackson and Rigden 1996; Poirier
204 2000),

$$205 \quad \Delta P_{th}(V, T) = \frac{\gamma}{V} [E_{th}(V, T) - E_{th}(V, T_0)] \quad , \quad (5)$$

206 where $E_{th}(V, T)$ and γ respectively represent the internal thermal energy and the
207 Grüneisen parameter. The internal thermal energy is calculable based on the Debye
208 model as

$$209 \quad E_{th}(V, T) = 9nRT \left[\frac{\theta}{T} \right]^{-3} \int_0^{\frac{\theta}{T}} \frac{x^3}{e^x - 1} dx, \quad (6)$$

210 where n , R , and θ respectively denote the number of atoms per formula unit, the gas

211 constant, and the Debye temperature. The Grüneisen parameter γ and the volume
212 dependence of the Debye temperature are expressed below (e.g., Nishihara et al. 2005;
213 Tange et al. 2009).

$$214 \quad \gamma(V) = \gamma_0 \left(\frac{V}{V_0} \right)^q \quad (7)$$

$$215 \quad \theta = \theta_0 \exp\left(\frac{\gamma_0 - \gamma}{q}\right) \quad (8)$$

216 Therein, q is a dimensionless parameter. For this approach, six parameters must be
217 fitted: V_0 , K_{T0} , K'_T , θ_0 , q , and γ_0 .

218 Table 2 presents results of MGD EoS fit. The V_0 , K_{T0} , and K'_T values are consistent
219 with the HTBM analyses within uncertainty. The thermal parameters are found as $\theta_0 =$
220 929 (114) K, $q = 2.9$ (10), and $\gamma_0 = 1.19$ (8).

221 The MGD EoS includes the expression of thermal energy (E_{th}). Therefore, we were
222 able to calculate the heat capacity of phase A from the MGD EoS parameters in Table
223 2 (Jackson and Rigden 1996; Poirier 2000). The isochoric heat capacity C_V is
224 calculated by taking the partial derivative of thermal energy E_{th} with respect to
225 temperature T ($C_V = (\partial E_{th} / \partial T)_V$), then the isobaric heat capacity C_P is obtainable by
226 conversion of the isochoric heat capacity C_V using the thermodynamic relation of
227 $C_P = C_V(1 + \alpha \kappa T)$. Figure 6 presents the calculated C_P of phase A at atmospheric
228 pressure. The value estimated by Holland and Powell (1998) is also shown for
229 comparison. Data obtained under lower temperatures are mutually consistent within
230 uncertainty, although our results obtained under higher temperatures have a slightly

231 higher value than those reported from an earlier study. This report is the first to
232 describe the heat capacity of phase A based on experimental data.

233

234

IMPLICATIONS

235 Oceanic lithosphere consists of a thin layer oceanic crust (MORB), harzburgite,
236 residual lherzolite and depleted pyrolite (Ringwood, 1982) for the schematic profile of
237 oceanic lithosphere. Considering the water saturated upper part of the oceanic
238 lithosphere, i.e, the oceanic crust and harzburgite part, the subducting slab was
239 modeled by a combination of 80% harzburgite + 20% MORB, in which, the
240 harzburgite consists of 80% olivine + 20% pyroxene. The addition of H₂O to this
241 ultramafic bulk composition may generate serpentine by serpentinization of the
242 ultramafic rocks. Antigorite is the stable serpentine mineral in ultramafic rocks
243 metamorphosed under the moderate temperatures of blue schist and green schist
244 facies conditions (Evans, 1976; Evans, 1977; Evans, 2004), and the antigorite will be
245 stable to ~200 km before dehydrated to hydrous phase A (Bromiley and Pawley, 2003;
246 Komabayashi et al., 2005a; Ulmer and Trommsdorff, 1995; Wunder and Schreyer,
247 1997). Here, we calculated the density profile of subducting slab with different degree
248 of serpentinization by using the new thermoelastic parameters of antigorite (Yang et al,
249 2014) and phase A, and discussed the effect of serpentinization on the density of
250 subducting slab.

251 Thermoelastic parameters of mineral phases used for the density calculation were

252 listed in Table 3. The density of each mineral phase was calculated along the cold
253 geotherm (Komabayashi et al., 2005b), and then the density of harzburgite was
254 modeled by 80% olivine and 20% pyroxene. Effect of degree of serpentinization on
255 the density was calculated by using the antigorite/phase A instead of olivine, e.g. ,
256 harzburgite with 10% serpentinization means 10% olivine changes to antigorite at
257 shallow depth; the antigorite dehydrates to hydrous phase A + Enstatite +water
258 ($5\text{Atg}=2\text{PhA}+8\text{En}+14\text{H}_2\text{O}$) at 6GPa (e.g. Ohtani et al., 2004), and we assumed that
259 the water from this reaction was escaped from the system by migrating upwards. See
260 Figure 7 (a) for the density profiles of harzburgites. The density profile of MORB
261 from Irifune and Ringwood (1987) was employed to calculate the density of
262 subducting slab. The density of subducting slab was modeled by 80% harzburgite and
263 20% MORB (Figure 7 (b)). The density of surrounding mantle was indicated by the
264 profile of AK135 (Kennett et al., 1995). The model calculation shows that the
265 serpentinization of subducted slab decreases the density significantly at shallower
266 depth, however, this effect become negligible when antigorite dehydrated to phase A
267 at ~200km depth. Because the phase A bearing subducting slab is supposed to be
268 denser than the surrounding mantle, the water can transport into deeper parts of the
269 upper mantle and the mantle transition zone.

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- 381

382 **TABLE 1.** Lattice parameters and the unit-cell volume of phase A at each P-T
 383 condition

No.	Pressure (GPa)	<i>T</i> (K)	<i>a</i> (Å)	<i>c</i> (Å)	<i>V</i> (Å ³)
AR198001	0.01 (1)	300	7.858 (4)	9.566 (8)	511.6 (7)
AR198010	6.12 (6)	300	7.713 (2)	9.433 (5)	485.9 (4)
AR198018	7.10 (8)	300	7.690 (2)	9.416 (5)	482.3 (3)
AR198031	8.07 (8)	300	7.670 (1)	9.393 (2)	478.6 (1)
AR198040	9.56 (3)	300	7.642 (1)	9.366 (2)	473.7 (1)
AR198049	9.14 (6)	300	7.650 (1)	9.374 (2)	475.1 (1)
AR198055	7.08 (7)	300	7.690 (1)	9.415 (3)	482.1 (2)
AR198056	5.31 (6)	300	7.726 (1)	9.453 (3)	488.7 (3)
AR198009	6.17 (4)	373	7.717 (2)	9.436 (5)	486.7 (4)
AR198017	7.12 (6)	373	7.695 (2)	9.420 (5)	483.1 (4)
AR198020	8.31 (3)	373	7.669 (2)	9.393 (4)	478.5 (3)
AR198030	8.09 (5)	373	7.675 (1)	9.397 (2)	479.4 (1)
AR198039	9.62 (3)	373	7.646 (1)	9.369 (2)	474.3 (1)
AR198048	9.27 (6)	373	7.652 (1)	9.376 (2)	475.5 (1)
AR198054	7.21 (7)	373	7.693 (1)	9.419 (3)	482.7 (2)
AR198008	6.12 (7)	473	7.725 (2)	9.446 (6)	488.2 (4)
AR198016	7.16 (6)	473	7.702 (2)	9.425 (5)	484.1 (3)
AR198021	8.44 (4)	473	7.674 (2)	9.398 (4)	479.3 (3)
AR198029	8.14 (4)	473	7.680 (1)	9.404 (2)	480.4 (2)
AR198038	9.70 (3)	473	7.652 (1)	9.374 (2)	475.3 (1)
AR198047	9.45 (6)	473	7.657 (1)	9.381 (2)	476.3 (1)
AR198053	7.39 (7)	473	7.697 (1)	9.423 (2)	483.5 (2)
AR198003	6.51 (1)	573	7.728 (3)	9.447 (7)	488.7 (6)
AR198007	6.19 (5)	573	7.732 (2)	9.453 (5)	489.5 (4)
AR198015	7.24 (6)	573	7.708 (2)	9.432 (5)	485.3 (4)
AR198022	8.48 (3)	573	7.681 (2)	9.406 (4)	480.5 (3)
AR198028	8.23 (4)	573	7.687 (1)	9.410 (2)	481.5 (2)
AR198037	9.82 (2)	573	7.658 (1)	9.380 (3)	476.4 (2)
AR198046	9.63 (6)	573	7.661 (1)	9.384 (2)	477.0 (1)
AR198052	7.61 (7)	573	7.702 (1)	9.428 (2)	484.3 (2)
AR198006	6.34 (3)	673	7.741 (2)	9.462 (5)	491.0 (4)
AR198014	7.41 (9)	673	7.716 (2)	9.439 (5)	486.7 (4)
AR198023	8.53 (4)	673	7.689 (2)	9.415 (5)	482.1 (4)

AR198027	8.37 (3)	673	7.694 (1)	9.418 (2)	482.8 (2)
AR198036	9.96 (2)	673	7.664 (1)	9.385 (3)	477.4 (2)
AR198045	9.86 (5)	673	7.666 (1)	9.390 (2)	477.9 (1)
AR198051	7.80 (7)	673	7.708 (1)	9.434 (2)	485.4 (2)
AR198005	6.44 (2)	773	7.748 (2)	9.472 (5)	492.4 (4)
AR198013	7.53 (7)	773	7.723 (2)	9.447 (5)	487.9 (4)
AR198024	8.61 (4)	773	7.697 (3)	9.424 (6)	483.5 (4)
AR198026	8.63 (4)	773	7.695 (3)	9.427 (6)	483.5 (5)
AR198035	10.12 (2)	773	7.669 (1)	9.391 (3)	478.3 (2)
AR198044	10.07 (6)	773	7.671 (1)	9.394 (2)	478.7 (1)
AR198012	7.61 (2)	873	7.733 (2)	9.457 (5)	489.7 (4)
AR198025	8.71 (4)	873	7.703 (3)	9.436 (7)	484.9 (5)
AR198034	10.22 (2)	873	7.675 (1)	9.397 (3)	479.3 (2)
AR198043	10.25 (7)	873	7.676 (1)	9.399 (2)	479.7 (1)
AR198033	10.32 (6)	973	7.681 (1)	9.405 (3)	480.5 (2)
AR198042	10.47 (7)	973	7.683 (1)	9.402 (3)	480.6 (3)

384 Parenthesis shows one standard deviation.

385

386 **TABLE 2.** Thermoelastic parameters of phase A from this study and earlier studies

	RT BM EoS	HT BM EoS	MGD EoS	K03	C&R02	P95	H06 [#]	S-V06 [#] s
V_0 (Å ³)	511.6 (2)	511.7 (3)	511.7 (3)	512.2 (6)	512.56 (3)	510.8	512.3 (3)	513.67
K_{T0} (GPa)	106.8 (18)	104.4 (24)	104.9 (23)	105 (4)	97.5 (4)	145 (5)	102.9 (28)	106 (1)
K_T'	3.88 (38)	4.39 (48)	4.29 (45)	3.9 (8)	5.97 (14)	4 (fixed)	6.4 (3)	5.8 (3)
dK_T/dT (GPa/K)	-	-0.027 (5)	-	-	-	-0.049*	-	-
$\alpha=a+bT$	-	-	-	-	-	-	-	-
a (10 ⁻⁵ /K)	-	2.88 (27)	-	-	-	4.9 (2)	-	-
b (10 ⁻⁸ /K ²)	-	3.54 (68)	-	-	-	-	-	-
θ_0 (K)	-	-	928 (114)	-	-	-	-	-
γ_0	-	-	1.19 (8)	-	-	-	-	-
q	-	-	2.9 (10)	-	-	-	-	-
RMS misfit (GPa)		0.075	0.152					

387 K03:Kuribayashi et al. (2003), C&R02: Crichton and Ross (2002), P95: Pawley et al.
 388 (1995), H06: Holl et al. (2006), and S-V06: Sanchez-Valle et al. (2006). #: Fe-bearing
 389 phase A, \$: adiabatic property by Brillouin spectroscopy, * calculated using empirical
 390 equation $dK/dT=-7\alpha K_{T0}$ (Holland et al., 1996)

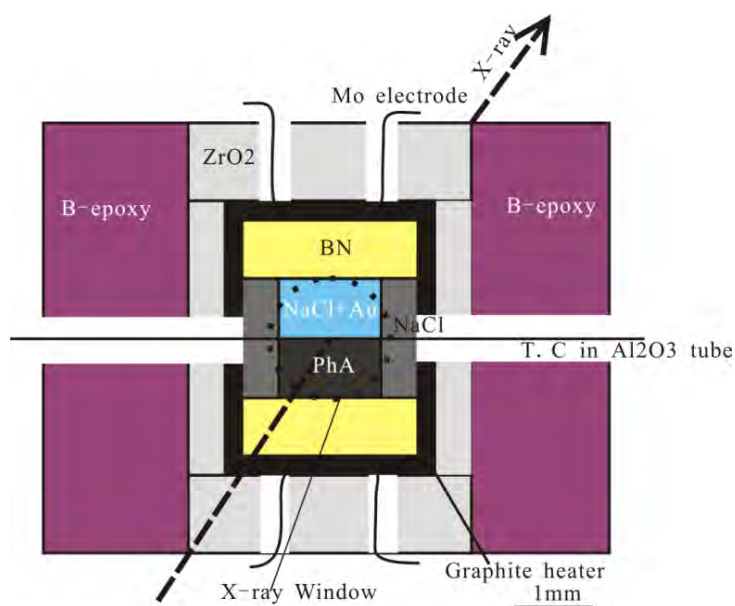
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392 **TABLE 3.** Thermoelastic parameters used in the density calculation

Mineral	V_0 (\AA^3)	K_0 (GPa)	K_0'	$(dK/dT)_P$ (GPa/K)	$a \times 10^{-5}$ (K^{-1})	$b \times 10^{-8}$ (K^{-2})	Ref.
olivine	292.13 (10)	129	4.61	-0.019 (2)	2.73 (34)	2.22 (81)	Liu and Li (2006)
enstatite	832.68 (9)	102.8 (2)	10.2 (12)	-0.037 (5)	2.86 (29)	0.72 (16)	Zhao et al. (1995)
antigorite	367.3 (2)	62.9	6.1	-0.027 (4)	3.92 (49)	-	Yang et al. (2014)
phase A	511.7 (3)	104.4 (24)	4.39 (48)	-0.027 (5)	2.88 (27)	3.54 (68)	This study

393 Parenthesis shows one standard deviation.

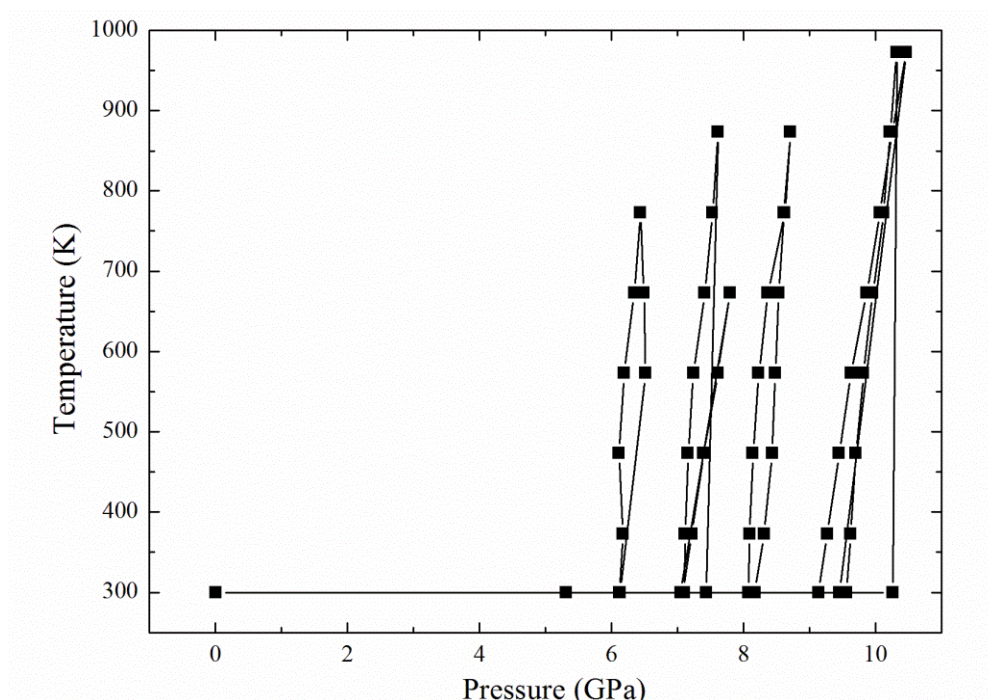
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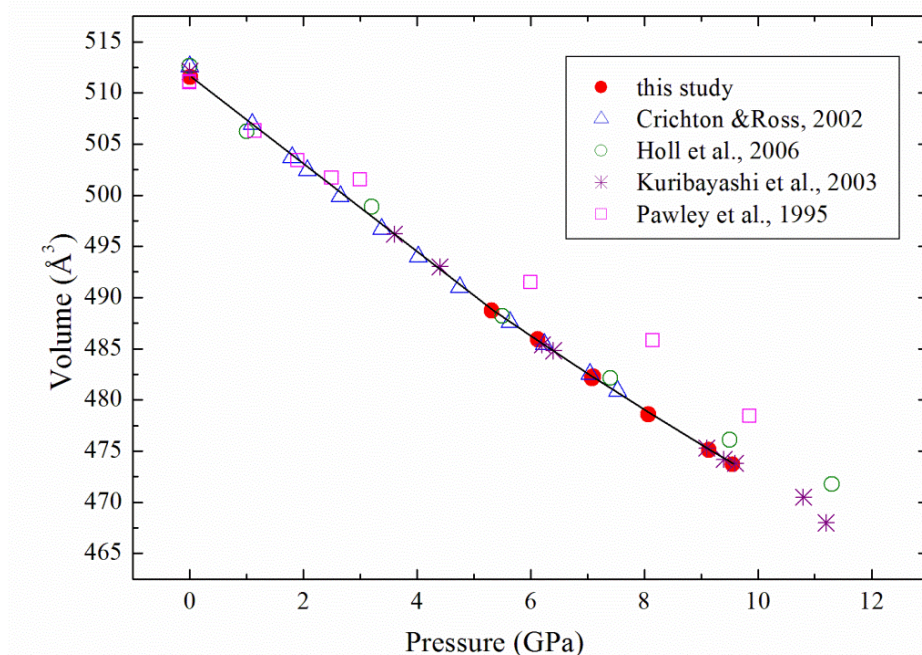
396 **FIGURE 1.** Cell assembly used for the present *in-situ* X-ray diffraction experiments.
 397 The X-ray path is shown by the arrow and the dotted circle, representing the “X-ray
 398 window” covered by h-BN in the ZrO₂ sleeve.

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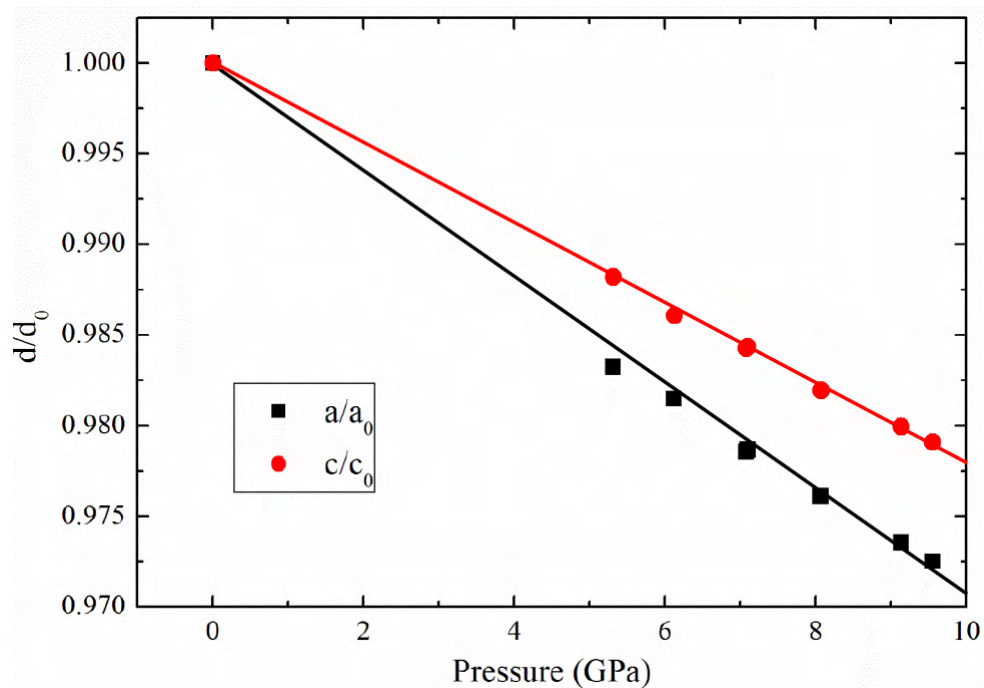
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FIGURE 2. Pressure–temperature pathway of the present experiment. Most P–V–T data used for EoS analysis were collected during cooling.



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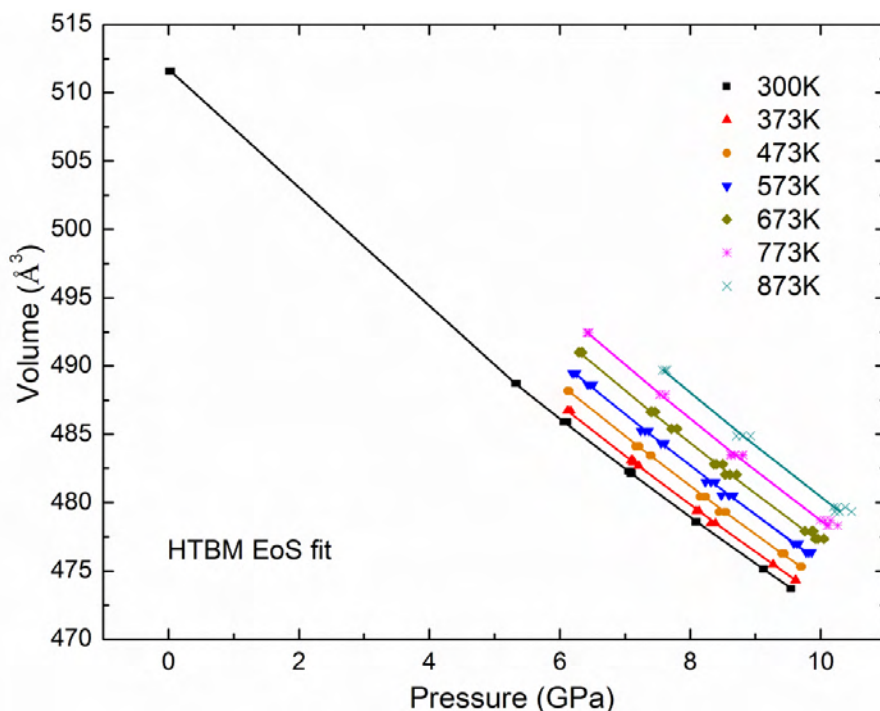
FIGURE 3. Room temperature compression data of phase A. The solid curve is the fit of room temperature data using the Birch–Murnaghan EoS. Data from earlier studies are also shown for comparison. Error bars are comparable with data markers. Data from Holl et al. (2006) are Fe-bearing phase A, whereas the others are of pure Mg end members.



411

412 **FIGURE 4.** Variation of unit-cell axis length with increasing pressure at room
413 temperature. Linear compression (β) of the a-axis and c-axis are, respectively,
414 $\beta_a=2.96(3) \times 10^{-3}/\text{GPa}$ and $\beta_c=2.21(1) \times 10^{-3}/\text{GPa}$. The a-axis is approximately 26%
415 more compressible than the c-axis. Error bars are within the data markers.

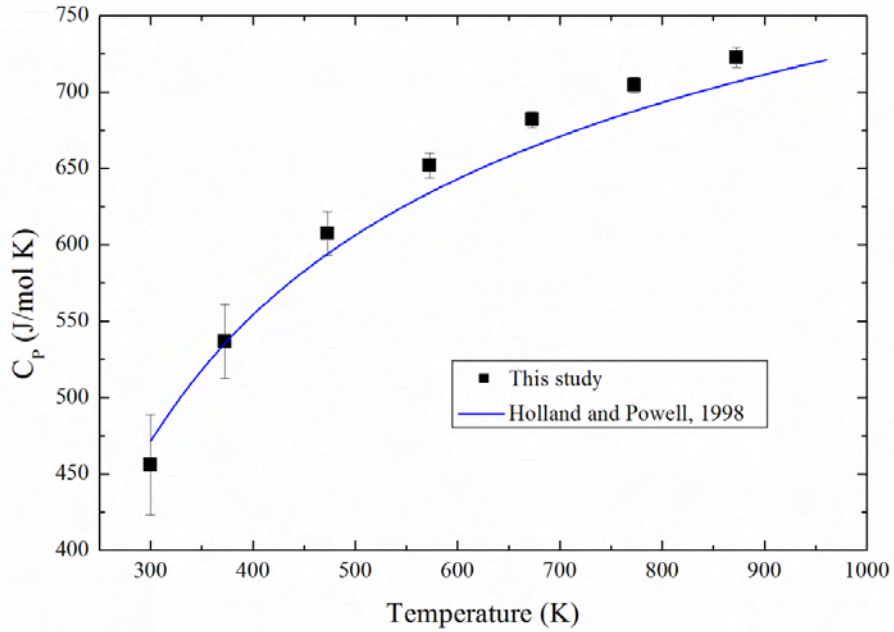
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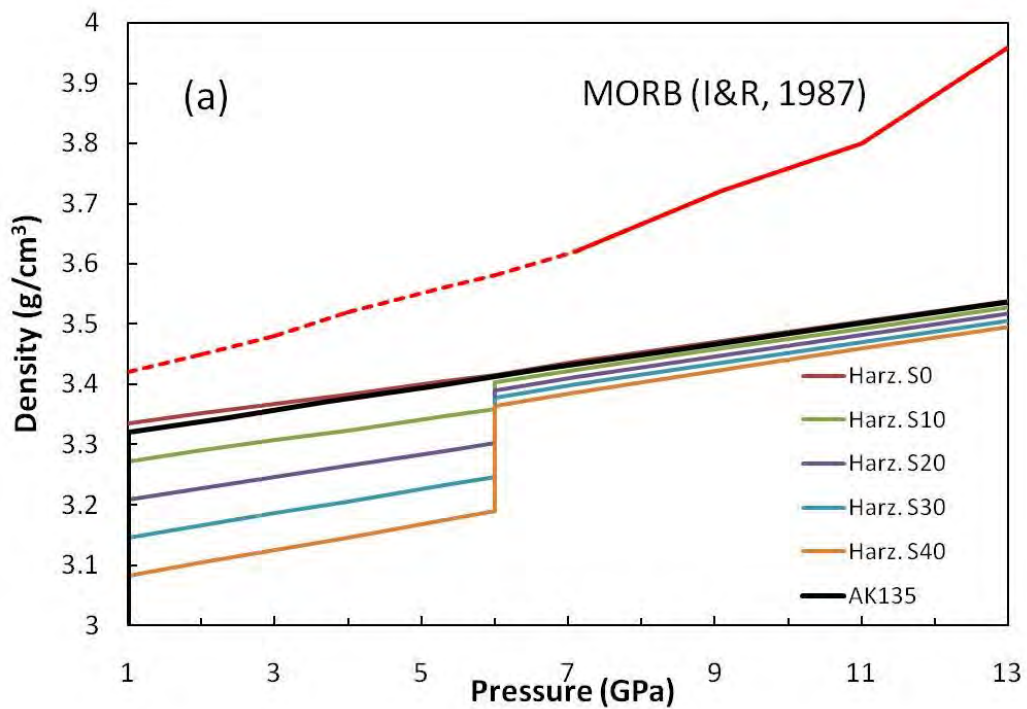
418 **FIGURE 5.** P–V–T data of phase A with the HTBM EoS fit. Solid lines are
419 isothermal compression curves at 300 K, 373 K, 473 K, 573 K, 673 K, 773 K, and
420 873 K. Error bars are within the data markers.

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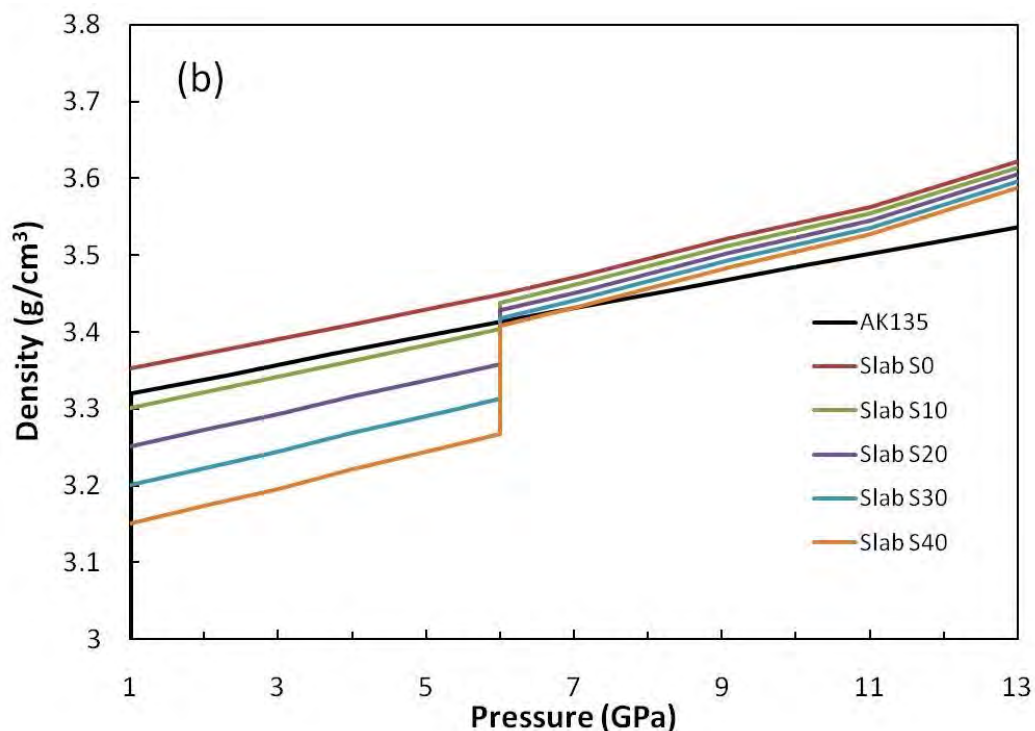


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FIGURE 6. Isobaric heat capacity C_p of phase A at 1 atm. Solid squares are based on MGD EoS fit of present P–V–T data. Their error bars correspond to the error of Debye temperature θ . Values of C_p estimated by Holland and Powell (1998) are shown for comparison.



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429

430 **FIGURE 7.** (a) Density of harzburgite and (b) subducting slab with different degree
431 of serpentinization (S0, S10, S20, S30, S40 and S50 show the serpentinization of 0,
432 10, 20, 30, 40 and 50%, respectively), along cold subduction zone geotherm
433 (Komabayashi et al., 2005b). Red line in (a) indicates the density of MORB
434 (Irfune and Ringwood, 1987). The density of MORB at lower pressure range (the
435 dashed line) was estimated from the density at high pressure range. The density
436 profile of AK135 (Kennett et al., 1995) is shown for comparison.

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