1	Crystal structure, thermal expansivity and elasticity of OH-
2	chondrodite: Trends among dense hydrous magnesium silicates
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15	Abstract: We report the structure and thermoelastic properties of OH-chondrodite. The sample was
16	synthesized at 12 GPa and 1523 K, coexisting with hydroxyl-clinohumite and hydrous olivine. The Fe
17	content Fe/(Fe+Mg) is 1.1 mol%, and the monoclinic unit-cell parameters are: $a = 4.7459(2)$ Å, $b =$
18	10.3480(7) Å, $c = 7.9002(6)$ Å, $\alpha = 108.702(7)^{\circ}$, and $V = 367.50(4)$ Å ³ . At ambient conditions the crystal
19	structure was refined in space group $P2_1/b$ from 1915 unique reflection intensities measured by single-
20	crystal x-ray diffraction. The volume thermal expansion coefficient was measured between 150 and 800 K,
21	resulting in $\alpha_V = 2.8(5) \times 10^{-9} (K^{-2}) \times T + 40.9(7) \times 10^{-6} (K^{-1}) - 0.81(3)(K)/T^2$, with an average value of
22	38.0(9)×10 ⁻⁶ K ⁻¹ . Brillouin spectroscopy was used to measure a set of acoustic velocities from which all
23	thirteen components (C_{ij}) of the elastic tensor were determined. The Voigt-Reuss-Hill average of the

25 GPa. The Reuss bound on the isothermal bulk modulus (K_{T0}) is 114.2(14) GPa. From the measured

moduli yield for the adiabatic bulk modulus, $K_{s0} = 117.9(12)$ GPa, and for shear modulus, $G_0 = 70.1(5)$

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26 thermodynamic properties, the Grüneisen parameter (γ) is calculated to be 1.66(4). Fitting previous static

27	compression data using our independently measured bulk modulus (isothermal Reuss bound) as a fixed
28	parameter, we refined the first pressure derivative of the bulk modulus, $K_{\rm T}' = 5.5(1)$. Systematic trends
29	between H ₂ O content and physical properties are evaluated among dense hydrous magnesium silicate
30	(DHMS) phases along the forsterite-brucite join.
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32	Keywords: chondrodite, dense hydrous magnesium silicates, crystal structure, thermal expansivity, bulk
33	modulus, Grüneisen parameter

35 Introduction:

36 The dense hydrous magnesium silicate (DHMS) minerals along the brucite-forsterite join include phase A, 37 norbergite, chondrodite, humite, and clinohumite (Fig. 1) as well as other DHMS minerals in the peridotite 38 system. Among these phases, phase A, chondrodite, and clinohumite have been observed as dehydration 39 products of serpentine as well as in experiments on MgO-SiO₂-H₂O systems at pressures of 5~12 GPa 40 (Berry and James 2001; Smyth et al. 2006; Wunder 1998). These phases could serve as H₂O carriers in the 41 Earth's upper mantle and transition zone. Due to the critical role of water in controlling silicate phase 42 relations and physical properties at upper mantle conditions, the crystal structures and physical properties 43 of DHMS have been extensively studied (e.g. Angel et al. 2001; McGetchin et al. 1970; Kanzaki 1991; 44 Faust and Knittle 1994; Irifune et al. 1998; Kawamoto 2006; Komabayashi and Omori 2006; Ohtani et al. 45 2000; Sanchez-Valle et al. 2006; Shieh et al. 2000; Wunder et al. 1995; Ye et al. 2013). The composition 46 and various thermoelastic properties of phase A, chondrodite, and clinohumite are summarized in Table 1.

47 The humite minerals are found in metamorphosed limestones and dolomites and contain variable 48 proportions of F and OH (Friedrich et al. 2002). Variable Ti and F/OH humites are also known in high-49 pressure ultramafic rocks (Evans and Trommsdorff 1983) as well as being recently described in a 50 serpentinized wehrlite (Shen et al. 2014). Within kimberlites, mineral associations including humites have 51 indicated equilibrium pressures in excess of 6 GPa (Taskaev and Ilupin 1990). Synthetic hydroxyl-52 chondrodite (nominally $Mg_5Si_2O_{10}H_2$) is stable up to at least 12 GPa at temperatures below 1473 K 53 (Burnley and Navrotsky 1996), and Fe-Ti-bearing compositions bracket the stability fields of chondrodite 54 and clinohumite in typical mantle compositions (Khodyrev et al. 1992). Fluorine may significantly increase 55 its thermal stability (Engi and Lindsley 1980) and incompressibility (Friedrich et al. 2002), but the 56 estimated concentration of fluorine in the upper mantle is limited (e.g. McDonough and Sun 1995). Hence, 57 the geological occurrence of chondrodite may be restricted to low-temperature regimes such as cold, 58 subducting slabs and the mantle wedge above them.

The crystal structure of chondrodite is illustrated in *Fig.* 2. The structure is monoclinic (space group $P2_1/b$, *a*-unique). The non-conventional setting of space group $P2_1/c$ can be chosen to preserve the association with the *a*- and *b*-axes of olivine. In the crystal structure of OH-chondrodite, there are three distinct (Mg,Fe)-O₆ octahedra (M1, M2 and M3) and one Si-O₄ tetrahedron (Si). The ratio of M1 : M2 : M3 63 : Si equals 1:2:2:2, since the M1 sites are fixed at the special position of (1/2, 0, 1/2). Hydrogen atoms 64 are bonded to O5 sites, which are also the positions for F anions in natural F-bearing samples. 65 Thermoelastic properties of DHMS phases along the olivine-brucite join have been reported for a range of 66 compositions (with variable H₂O, Fe, Ti, F, etc.) including thermal expansivity (Pawley et al. 1995; Ye et 67 al. 2013; etc), isothermal compressibility (Crichton and Ross 2002; Kuribayashi et al. 2004; Freidrich et al. 68 2002; Holl et al. 2006; Ross and Crichton 2001), and adiabatic elastic moduli (Fritzel and Bass 1997; 69 Jacobsen et al. 2008; Jiang et al. 2006; Sanchez-Valle et al. 2006; Sinogeikin and Bass 1999; Phan 2009; 70 Xia et al. 1998). In this study, we examined the crystal structure of a Fe-bearing, OH-chondrodite sample 71 by single-crystal X-ray diffraction and determined the lattice coefficients of thermal expansivity between 72 150 and 800 K. Brillouin spectroscopy was used to measure a set of velocities from three different platelets 73 that constrain the thirteen independent C_{ij} of the elasticity tensor. By comparing the current results with 74 physical properties of other DHMS phases along the forsterite-brucite join, we establish trends as a 75 function water content that can be used to model the physical properties of mineral assemblages found in 76 hydrated subducting slabs.

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78 Experiments

79 Sample synthesis and composition

80 The Fe-bearing OH-chondrodite crystals were synthesized in the 5000-ton multi-anvil press at 81 Bayerisches Geoinstitut, Bayreuth, Germany, run SZ0407B (Smyth et al. 2006). An outer 3.5-mm welded 82 Pt capsule contained two separate 1.6 mm-diameter welded Pt inner capsules in a matrix of brucite to 83 maintain H activity. The inner capsule contained a 1-mm single-crystal cube of San Carlos olivine, plus a 84 mixture of brucite and forsterite. The inner capsules also contained small amount of a Re-ReO₂ oxygen 85 buffer to maintain the iron in the ferrous state. The run was made in an 18-mm sintered MgO octahedron 86 using 54-mm WC cubes with 8-mm corner truncations. The nominal pressure was 12 GPa at 1250°C with a 87 heating duration of 180 min. The original objective of the experiment was to hydrate the San Carlos 88 olivine, but rupture of the inner capsule exposed the San Carlos olivine to excess brucite, with which it 89 reacted to form Phase A, chondrodite, and clinohumite. The chondrodite crystals were up to 0.25 mm in 90 diameter and light tan in color.

91 A single crystal about 120 µm in size was selected and polished for chemical analysis by electron 92 microprobe following the experimental procedure described previously for the coexisting clinohumite (Ye 93 et al. 2013). Six points at different locations of the crystal were chosen for analysis, and the average values 94 together with their standard deviations are listed in Table 2. The water content was calculated to be 5.26 95 wt.%, assuming the molar ratio of H:O to be 2:10. The formula is expressed as 96 97 1.1 % which is lower than those of the coexisting OH-clinohumite (4 % Fe) (Ye et al. 2013) and hydrous 98 olivine (3 % Fe) (Jacobsen et al. 2008, hy-Fo₉₇) in the run products.

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100 Single-crystal X-ray diffraction (XRD)

101 A single crystal of OH-chondrodite measuring $100 \times 85 \times 80 \ \mu\text{m}^3$ was chosen for X-ray diffraction 102 at ambient conditions. Unit-cell parameters were refined on a Bruker P4 four-circle diffractometer with a 103 dual-scintillation point detector system using an 18-kW rotating Mo-anode X-ray source operating at 50 kV 104 and 250 mA. The average wavelength for Mo $K\alpha_1$ - $K\alpha_2$ mixed characteristic wavelengths was determined 105 to be 0.71073 Å by measuring a spherical single crystal of anhydrous forsterite (Fo₁₀₀). Least squares fitting was performed on 48 centered reflections within the following classes: (120), $(1\overline{1}2)$, $(1\overline{2}2)$, $(1\overline{3}1)$, (130), 106 $(112), (1\overline{3}2), (131), (210), (1\overline{1}3), (1\overline{2}3), (122), (1\overline{4}1), (211), (2\overline{1}2), (2\overline{2}3), (222), and (2\overline{4}1), and the$ 107 108 refined unit-cell parameters at ambient condition are: a = 4.7459(2) Å, b = 10.3480(7) Å, c = 7.9002(6) Å, 109 $\alpha = 108.702(7)^{\circ}$, and V = 367.50(4) Å³. Intensity data were collected using a Bruker APEX II CCD 110 detector mounted on a P4 diffractometer up to $2\theta = 75^{\circ}$. Refinement of atomic positions and anisotropic 111 displacement parameters was carried out using SHELXL-97 (Sheldrick 2008) in the software package 112 WinGX (Farrugia 2012), with the default neutral atom scattering factors in SHELXL-97. In total, 10333 113 reflections were collected with 1915 unique and 1833 with $I > 4\sigma$ used in the refinement. Model fit 114 parameters are: $R_1 = 0.0213$ for $I > 4\sigma$, and 0.0223 for all the unique reflections; goodness of fit (GooF) = 115 1.674; and for merging of equivalent reflections, $R_{int} = 0.039$.

116 The refined atomic position coordinates and occupancy factors are listed in *Table 3*, and the 117 anisotropic displacement parameters are given in *Appendix 1*. In this study, Fe and Mg occupancies in the 118 M1, M2 and M3 sites were refined, as well as Si occupancy, and the O occupancies were fixed at full 119 occupancy. In addition, the H position was refined with an isotropic displacement parameter (U_{eq}) fixed at 120 0.054.

121 The same single crystal of OH-chondrodite was then used for thermal expansion measurements 122 below and above room temperature. At each temperature step, the refinement of unit-cell parameters was 123 performed using the same experimental procedure as the room-temperature measurement described above. 124 Low-temperature measurements were carried out at 253 K, 203 K and 153 K. Temperatures were measured 125 and controlled by a Bruker LT-2A controller and N₂ gas stream nozzle. For high-temperature 126 measurements, the crystal was transferred to a silica glass capillary, sealed at both ends. Ten high-127 temperature steps were carried out from 350 K to 787 K using a Bruker high-temperature device with a 128 two-prong ceramic-coated Pt wire radiant heater and an Omega temperature control unit (Ye et al. 2009). 129 The refined unit-cell parameters at low and high temperatures are given in Appendix 2.

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131 Brillouin Spectroscopy

Three crystals of OH-chondrodite (from run SZ0407B) were prepared for Brillouin scattering experiments by double-sided polishing to produce parallel plates measuring about 100 μm across and 30-50 μm thick. Polishing was carried out with 3 μm diamond lapping film and finishing with 1 μm and then finally 0.5 μm diamond lapping film to produce an optical-quality surface free of visible scratches.

Prior to Brillouin spectroscopy, Raman spectroscopy was carried out on each of the three platelets. Unpolarized Raman spectra were obtained using a 200 mW Ar-ion laser ($\lambda = 514.532$ nm) with a confocal optical setup in back-scattering geometry using holographic optics and a single-grating 0.5 m spectrometer with a 1100 x 330 pixel CCD camera. Raman spectra of the three OH-chondrodite crystals used in the Brillouin scattering experiments are shown in *Fig. 3 (A-D)*.

141 Brillouin Spectra were obtained in a forward symmetric scattering geometry (Speziale and Duffy 142 2002; Speziale et al. 2014) with a six-pass Sandercock tandem Fabry-Perot interferometer, using a solid-143 state laser ($\lambda = 532.15$ nm, 150 mW at the sample). Acoustic velocities (V_i) were determined as:

144
$$V_i = \frac{\Delta \omega_i \lambda}{2\sin(\frac{\theta}{2})}, \qquad (1)$$

where $\Delta \omega_i$ is the Brillouin scattering frequency shift of the acoustic modes, and θ is the scattering angle in the sample (70° in this study). For each platelet, measurements were performed in ~5-10° steps with the total number of spectra in a given platelet depending on its size and the quality of the signal. Measured acoustic velocities are plotted as functions of azimuthal angle in *Fig. 4 (A-C)*. Further details about the Brillouin scattering measurements can be found in Speziale and Duffy (2002) and Speziale et al. (2014).

Three platelets were used for Brillouin scattering, with the orientation of each platelet defined with three Eulerian angles (θ , ϕ , χ). Chondrodite has a monoclinic structure with thirteen elastic constants: C_{11} , C_{22} , C_{33} , C_{44} , C_{55} , C_{66} , C_{12} , C_{13} , C_{23} , C_{15} , C_{25} , C_{35} , and C_{46} . The elastic constants and acoustic velocities are related by Christoffel's equation (Every 1980):

154
$$\det |C_{ijkl}n_{j}n_{l} - \rho V_{P,S}^{2}\delta_{ik}| = 0$$
 (2)

155 where C_{ijkl} is the elastic tensor in full suffix notation, n_i is the direction cosine of the phonon propagation 156 direction, density $\rho = 3.099$ g/cm³ (according to the composition determined by EPMA and unit-cell 157 volume from single-crystal XRD at ambient condition), and $V_{P,S}$ is the acoustic velocity of either 158 longitudinal (V_P) or transverse (V_S) in the Brillouin spectra. The inversion was carried out by non-linear 159 lest-squares using Levenberg-Marquardt method (Press et al. 1988) with closed-form expressions for the 160 Christoffel equation given by Every (1980). The inversion process was iterative. The elastic constants of 161 Sinogeikin and Bass (1999) were used as the initial model and held fixed while the inversion was 162 performed for the three Eulerian angles defining the orientation of each plane. The relationship between the 163 Eulerian angles and the direction cosines of the phonon propagation direction is given by Shimizu (1995). 164 In subsequent inversions, the elastic constants and orientations were alternately varied or allowed to vary 165 together, and the process was continued until convergence was achieved. Different sets of initial elastic 166 constants were tested to confirm that the final results were insensitive to the starting model. The results are 167 shown in Fig. 4(A-C). The good quality of the fit is demonstrated by the small root-mean-square difference 168 between the measured and calculated velocities which was $V_{\text{RMS}} = 39$ m/s.

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170 Results and Discussion

171 Raman spectroscopy

172 Unpolarized Raman spectra of the three OH-chondrodite platelets used for Brillouin scattering are 173 shown in Fig. 3. The spectra are generally similar to previous Raman studies of OH-chondrodite (Lin et al. 174 1999; Mernagh et al. 1999). In Fig. 3, the polished plane orientation is expressed as (hkl), fitted from the 175 velocity dispersion curves in the Brillouin scattering study. Band assignments for some of the modes shown 176 in Fig. 3 are given elsewhere (Cynn et al. 1996; Lin et al. 1999). In the O-H stretching region, we find a 177 similar strong doublet as was found for F-free synthetic OH-chondrodite (Lin et al. 1999). The main band 178 positions shown in Fig. 3 are 3525 and 3565 cm⁻¹, which are shifted by about 10 cm⁻¹ from the positions 179 shown by Lin et al. (1999) at 3515 and 3554 cm⁻¹. Also in Fig. 3 is shown a deconvolution of the O-H 180 stretching doublet (dashed lines), which is composed of at least four separate bands at 3525, 3560, 3565, 181 and 3580 cm⁻¹. The 3525 cm⁻¹ band may be further deconvolved into three separate bands to best fit the 182 observed spectrum.

183

184 Crystal Structure

185 According to the refined Mg and Fe occupancies in *Table 3*, the total Fe mol% is 0.9(4)%, consistent 186 within uncertainty with that of (1.1 mol%) obtained from electron microprobe analyses. The Fe content 187 from single-crystal XRD (site occupancy) is an approximation as minor elements (Ti, Ni, Mn, Ca, Al, < 188 0.04 mol%) were ignored in the structural refinement. The Si occupancy refinement from X-ray diffraction 189 is consistent with that from microprobe analyses. The bond lengths and polyhedral volumes were calculated 190 using the software package XTALDRAW (Downs et al. 1993), and listed in Table 4. The octahedral 191 volumes shows the order of M2 > M3> M1, and the Si tetrahedral volume is 2.21(1) Å³. The hydrogen 192 bond geometry shows an O5-H distance of 1.06(2) Å and O5...H of 2.12(2) Å with a hydrogen bond angle 193 of 122.7°. The O5-H bond length (O/F-H bonds in some chondrodites) has been refined by neutron 194 diffraction (Friedrich et al. 2001 and Kunz et al. 2006), as well as X-ray diffraction (Ottolini et al. 2000 and 195 Friedrich et al. 2002). Neutron diffraction better reflects the OF5-H interatomic distances than single-196 crystal X-ray diffraction. Friedrich et al. (2001) reported a bond length of 1.029(6) Å for OF-H and 197 1.872(6) Å for OF5...H, while Kunz et al. (2006) reported 0.976(7) Å and 1.940(7) Å, respectively. Our 198 O5-H bond length is close to those from neutron experiments, whereas the O5...H interatomic distance is 199 10 % longer than previous results from neutron diffraction.

200 To compare the compositional effects of Fe, F, and Ti on the crystal structure of chondrodite, the 201 unit-cell volumes and M-OF bond lengths are summarized in Table 5 for this study (SZ0407B) and 202 literature values with the following abbreviations used throughout: Berry and James 2002, B02; Friedrich 203 et al. 2001 and 2002, F01 and F02; Fujino and Takéuchi 1978, F78; Gibbs et al. 1970, G70; Kuribayashi et 204 al. 2004, K04; Kunz et al. 2006, K06; Ottolini et al. 2000, O00; Yamamoto 1977, Y77. The α angle in 205 Sample F78 is significantly larger than other samples in *Table 5*, which is likely due to the effect of Ti^{4+} 206 incorporation (Robinson et al. 1973). In addition, α angle values in SZ0407B and Y77 are 0.35~0.4° 207 smaller than those in Y77, B02, G70, O00, F01, F02, K06 and K04, implying that F⁻ might increase the 208 angle. The unit cell volumes decrease in the order of SZ0407B and Y77 > F78 > F01, F02, K06 and K04 > B02 and G70 > Y77 consistent with the order of decreasing ionic radii ($Fe^{2+} > Mg^{2+} > Ti^{4+}$) and anions (O^{2-} 209 210 > F⁻). Samples SZ0407B, Y77 and F78 are F-free, while F78 has a high Ti⁴⁺ concentration of 8.4 mol%. 211 Samples F01, F02, K06 and K04 have higher Fe²⁺ concentrations than B02, G70 and Y77, and B02 has the 212 highest F^- content among these samples. The most significant effect of F^- on the crystal structure is that F^- 213 decreases distances between OFOH and surrounding cations, i.e. one M2-OF bond and two M3-OFOH 214 bonds. Samples SZ0407B and Y77 typically have larger M-OF bond lengths than other samples, as shown 215 in Table 5, while sample F78 has significantly smaller M3-OF bond lengths, due to the occupancies of smaller cations Ti⁴⁺ in M3 site. Fe²⁺ should increase the M-O bond lengths but such effect is not well 216 217 observed here, perhaps because of the low concentration of Fe^{2+} in the reported crystal structures and, 218 possible technical discrepancies among different measurements.

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220 Thermal Expansivity

To relate the anisotropy of thermal expansion to the crystal structure, the principal axes of thermal expansion are determined for this monoclinic structure, reducing the number of thermal expansion coefficients from 4 to 3: X_1 , X_2 , and X_3 (Cliffe and Goodwin 2012). The variations with temperature of the principle expansion axes are listed in *Appendix 2*, and the average axial thermal expansion coefficients are listed in *Table 6*, as well as the orientations of the principle expansion axes. There is no difference between *a* and X_1 for a-unique monoclinic structure, while the thermal expansion coefficient of X_2 is 8% smaller than that of *b* axis for the compensation of α angle (>90°) expansion. 228 The thermal expansion coefficient is defined as:

229
$$\alpha(T) = \frac{1}{V} \left(\frac{\partial V}{\partial T} \right)_P = \left(\frac{\partial \ln(V)}{\partial T} \right)_P.$$
(3)

230 Fei (1995) proposed a useful formula for modeling $\alpha(T)$ above room temperature:

231
$$\alpha(T) = a_0 + a_1 T + \frac{a_2}{T^2}$$
. (Fei equation) (4)

232 On the basis of the Mie-Grüneisen-Debye Equation of Stae, Suzuki (1975), Suzuki et al. (1979) and Kumar

233 (1995, 1996, 2000) developed expression for the unit-cell volume as function of temperature:

234
$$V_0(T) = V_0(0) \cdot \left[1 + \frac{1 - (1 - 4k \cdot E_{th}(T) / Q_0)^{1/2}}{2k}\right]$$
 (Suzuki equation) (5)

235
$$V_0(T) = V_0(0) \cdot \left[1 - \frac{\ln[1 - (2k+2) \cdot E_{th}(T) / Q_0]^{1/2}}{2k+2}\right] \text{ (Kumar equation)}$$
(6)

where $k = (K_T'-1)/2 = 2.25$ ($K_T' = 5.5$ as in Table 1), $Q_0 = V_0(0)K_{T0}(0)/\gamma_{MGD}$. $V_0(0)$ and $K_{T0}(0)$ are the volume and isothermal bulk modulus taken at 0 K and 0 GPa. γ_{MGD} is a parameter related to the thermodynamic Grüneisen parameter γ_{th} (Barron et al. 1980). The thermal energy $E_{th}(T)$ is expressed by the Debye model:

240
$$E_{th}(T) = \frac{9nRT}{(\Theta_D / T)^3} \int_0^{\Theta_D / T} \frac{x^3}{e^x - 1} \cdot dx$$
(7)

241 where *R* is the gas constant, *n* is the number of atoms in the formula unit (19 for OH-chondrodite), and $\Theta_{\rm D}$

242 is the Debye temperature, and the acoustic Debye temperature Θ_{ac} is given by:

243
$$\Theta_{ac} = \frac{h}{k} \cdot \left(\frac{3N}{4\pi}\right)^{1/3} \cdot \left(\frac{n \cdot \rho}{M}\right)^{1/3} \cdot v_m \tag{8}$$

244
$$\frac{3}{v_m^3} = \frac{2}{v_s^3} + \frac{1}{v_p^3}$$
(9)

where *h*, *k*, *N* are Boltzmann, Plank and Avagadro's constants, respectively, and M is the molar mass. The calculated Θ_{ac} values for phase A, chondrodite and clinohumite are listed in *Table 1*. The Θ_{ac} values for 247 these DHMS phases are close to that (729 K) for San Carlos olivine (Liu and Li 2006), and we adopt $\Theta_D =$

248 741 K to calculate the thermal expansion coefficient of chondrodite in this study.

249 The unit-cell volume and principal expansion axes are plotted in Fig. 5(A) as a function of 250 temperature, and fitted by the curves calculated from Fei equation (Eq. 4). The principal expansion axes 251 show an order of $\alpha_{x3} > \alpha_{x1} > \alpha_{x2}$. We derive the thermal expansion coefficients for unit-cell volume as: a_0 252 $= 40.9(7) \times 10^{-6} \text{ K}^{-1}$, $a_1 = 2.8(5) \times 10^{-9} \text{ K}^{-2}$, $a_2 = -0.81(3) \text{ K}$ according to Fei equation. Next, the V(T) dataset 253 from this study is also fitted by Suzuki and Kumar equations (Eq. 5 and 6). We used $K_{T0}(298K) = 114$ GPa 254 (Table 1), and assuming that OH-chondrodite has the same dK_{T0}/dT as forsterite (-0.023 GPa/K, Kroll et al. 255 2012), we obtain $K_{T0}(0K) = 121$ GPa. Then the fittings result in $\gamma_{MGD} = 1.19(6)$ from Kumar equation; 256 1.21(8) from Suzuki equation; and $V_0(0K) = 365.8(3) \text{ Å}^3$ for both Kumar and Suzuki equations.

257 The calculated $\alpha_V(T)$, calculated from Fei, Kumar and Suzuki equations, are plotted in Fig. 5(B), 258 respectively, and extrapolated to 1500 K where the breakdown of OH-chondrodite was observed (Burnley 259 and Navrotsky 1996). The Fei equation generates negative α_V below 140 K, whereas the Kumar and Suzuki 260 equations give larger and more reasonable α_V at such low temperatures. From 200 K to 800 K, these three 261 equations agree with each other well, with discrepancy of 5% at 800 K. When extrapolated to 1500 K, the 262 Kumar and Suzuki equations indicate α_V values 20% and 13% larger, respectively, compared with that 263 from Fei equation. In addition, for $\alpha(T)$ relation as proposed by Fei (1995), the sum of the thermal 264 expansion coefficients for principal axes at the measured temperatures is also plotted in Fig. 5(B), which 265 consistent with the solid curve of $\alpha_V(T)$ from Fei equation.

266 The axial thermal expansivity for the present Fe-bearing OH-chondrodite sample shows anisotropy 267 in the order of $\alpha_c > \alpha_a > \alpha_b$, whereas axial compressibilities determined from single-crystal XRD data at 268 high pressure yield an anisotropy of $\beta_b \approx \beta_c > \beta_a$ for Fe-free OH-chondrodite (Ross and Crichton 2001; 269 Kuribayashi et al. 2004), and $\beta_b > \beta_c > \beta_a$ for Fe,F-bearing samples (Friedrich et al. 2002; Kuribayashi et al. 270 2004). The axial compressibilities are also consistent with the elasticity studies: $C_{11} > C_{33} \approx C_{22}$ for Fe-free 271 OH-chondrodite (this study, in the following discussion), and $C_{11} > C_{33} > C_{22}$ for Fe,F-bearing sample 272 (Sinogeikin and Bass 1999). Here, the elastic moduli are transferred to a-unique monoclinic symmetry to 273 be consistent with the studies of crystal structure and isothermal compression. In this study, we find that the

274 α angle increases with temperature, which is consistent with the observations that α decreases with 275 pressure (Ross and Crichton 2001; Friedrich et al. 2002; Kuribayashi et al. 2004). The angle α increases as 276 temperature increasing with a thermal expansion coefficient of 1.43(10) x10⁻⁶ K, consistent with that for 277 clinohumite (1.41~1.53 x10⁻⁶ K from Ye et al. 2013).

278 The average volume thermal expansion coefficients (α_0) for the DHMS phases along the forsterite-279 brucite join are plotted in Fig. 6, as a function of water content ($C_{\rm H2O}$, expressed as wt.% H₂O). The $\alpha_0(V)$ 280 values are recalculated over temperature range of 300 - 787 K for forsterite (Ye et al. 2009), clinohumite 281 (Ye et al. 2013) and chondrodite (this study) to allow for consistent comparison with previous experiments 282 on phase A (Pawley et al. 1995) and brucite (Fukui et al. 2003). There have been many previous studies on 283 thermal expansivity of anhydrous forsterite (e.g. Ye et al. 2009; Kroll et al. 2012; and Trots et al. 2012). To 284 simplify Fig. 6, we plotted the anhydrous forsterite thermal expansivity data from Ye et al. (2009) as representative. The α_0 for hydrous forsterite (0.89 wt% H₂O) is ~ 0.8×10⁻⁶ K⁻¹ larger than that for 285 anhydrous forsterite. The Fe-bearing clinohumite (Fe % = 4.1 mol%) shows $\alpha_0 \sim 2.2 \times 10^{-6} \text{ K}^{-1}$ smaller than 286 287 Fe-free clinohumite (Ye et al. 2013). The α_0 for the current OH-chondrodite sample is 40.8(4) ×10⁻⁶ K⁻¹ in

288 the range of 300 - 787 K, whereas it is $38.0(9) \times 10^{-6}$ K⁻¹ in 153 - 787 K.

A linear regression for the data of *Fig.* 6 yields a systematic relation between α_0 and C_{H2O} as:

290 a

$$\alpha_0 (10^{-6} \,\mathrm{K}^{-1}) = 35.3(4) + 1.02(3) \times C_{\mathrm{H2O}} \,(\mathrm{wt\%}) \qquad (\mathrm{R}^2 = 0.9947) \tag{10}$$

Each additional 1 wt% H₂O content increases α_0 by $1.02(3) \times 10^{-6}$ K⁻¹ (or about 3 %). Thus, the thermal expansion coefficient increases with H₂O content for DHMS along the forsterite-brucite join, which is consistent with the observation that the bulk moduli decrease as H₂O content increases (Ross and Crichton 2001; Holl et al. 2006; Jacobsen et al. 2008; Sanchez-Valle et al. 2006).

After heating to 787 K, the single crystal of OH-chondrodite was quenched to room temperature, and the unit cell parameters after heating are: a = 4.7455(4) Å, b = 10.3475(8) Å, c = 7.9008(9) Å, $\alpha = 108.706(8)^{\circ}$, and V = 367.47(7) Å³, consistent with the initial ones before heating. We conclude that this Fe-bearing OH-chondrodite sample persisted to 787 K at ambient pressure without any observable phase transition or dehydration.

301 Elastic moduli

302 The calculated elastic moduli for OH-chondrodite (Fe/(Mg+Fe) = 1.1 %) are listed in *Table 7*, with 303 comparison to F-bearing chondrodite (Fe/(Mg+Fe) = 5.5 %, F/(F+OH) = 32%, Sinogeikin and Bass 1999). 304 Nearly all of the longitudinal and shear elastic constants are lower than those reported by Sinogeikin and 305 Bass (1999) for a F-rich natural chondrodite. The Brillouin inversion was performed in standard monoclinic 306 symmetry (b-unique, $P2_1/c$) where zigzag chains of edge-sharing octahedra are aligned along the a axis and 307 lie in the plane perpendicular to the b axis. The elastic indices are also reported in the a-unique coordinate 308 system $(P2_1/b)$ (values given in parentheses in *Table 7*) to facilitate comparison with the studies of the 309 crystal structure. The longitudinal constants C_{22} and C_{33} (in standard monoclinic symmetry hereafter) are 310 close to the values from Sinogeikin and Bass (1999), whereas C_{11} is 12% smaller. The shear constants C_{55} 311 and C₆₆ are about 5 % smaller, while C₄₄ is 8% smaller. The off-diagonal moduli are not constrained as well 312 in either study. The Voigt / Reuss bounds, and Voigt-Reuss-Hill (VRH) average of the aggregate moduli 313 (Watt 1980) for OH-chondrodite are summarized in Table 8. The bulk modulus found here is identical 314 within uncertainty to that obtained by Sinogeikin and Bass (1999), and consistent with those reported in 315 static compression studies (Ross and Crichton, 2001, Kuribayashi et al., 2004; Friedrich et al., 2002). The 316 shear modulus, G_0 , is however about 6% lower than that reported by Sinogeikin and Bass (1999) for a 317 natural F-bearing sample, indicating that the shear modulus may increase with F content.

318 Using the density (ρ), isothermal bulk modulus (K_{S0}), thermal expansion coefficient (α_0) and heat 319 capacity (C_P) listed in *Table 1*, we calculated the Grüneisen parameter (γ) for phase A, OH-chondrodite and 320 OH-clinohumite, from the formula:

$$\gamma = \frac{\alpha \cdot K_s}{\rho \cdot C_p} \tag{11}$$

The calculated γ values are also listed in *Table 1*. The Grüneisen parameter for phase A (1.69(8)), chondrodite (1.66(4)), and clinohumite (1.50(4)) is larger than that for forsterite (1.28, Gillet 1991), since phase A, chondrodite and clinohumite have smaller C_P and ρ values than forsterite, whereas variation in the product $\alpha \cdot K_S$ is relatively less significant among these phases. Because the Reuss bound on the bulk modulus, K_{S0} , corresponds to constant stress across grain boundaries, which is relevant for static compression studies, we calculate the isothermal Reuss bound bulk modulus for OH-chondrodite: $K_{T0} = 114.2(14)$ GPa, with $K_{S0} = 116.4(12)$ GPa and the formula:

$$K_T = \frac{K_S}{1 + \alpha \cdot \gamma \cdot T} \tag{12}$$

330 Next, on the basis of reported compression datasets for Fe-free OH-chondrodite (Ross and Crichton 2001; 331 Kuribayashi et al. 2004), we refined the pressure derivative of the bulk modulus, $K_{\rm T}$, and obtain $K_{\rm T}$ ' = 332 5.5(1) (Fig. 7) by fixing K_{T0} at 114.2 GPa, the Reuss bound from Brillouin scattering. Fig. 7 also shows 333 that both these high-pressure datasets are consistent with each other. It should be noted that our OH-334 chondrodite has 1.1 mol% Fe, but such a low Fe concentration should probably be expected to have little 335 effect on the bulk modulus. The refined results are also included in Table 8. Our results are consistent with 336 the chondrodite sample from Friedrich et al. (2002) (Fe-free, F/(F+OH) = 0.32) which further indicates that 337 F content does not strongly affect the bulk modulus. It should also be noted that the study by Faust and 338 Knittle (1994) could be affected by non-hydrostatic stress for compression to 42 GPa in the pressure 339 medium of methanol, ethanol and water mixture.

Over the past three decades, extensive studies have been carried out on the elasticity of DHMS phases along the forsterite-brucite join (Suzuki et al. 1983; Webb 1989; Zha et al. 1996 and 1998; Fritzel and Bass 1997; Sinogeikin and Bass 1999; Sanchez-Valle et al. 2006; Jacobsen 2008 and 2009; Phan 2009; as well as this study). The density (ρ), elastic moduli (K_{S0} and G_0 , Voigt-Reuss-Hill averages) and acoustic velocities (V_P and V_S) are summarized in *Fig.* 8(*A*), (*B*) and (*C*), respectively, with linear regression fittings listed below:

346	ρ (g/cm ³) = 3.27(2) – 0.029(2)× $C_{\rm H2O}$	$(R^2 = 0.9767)$	(13)
347	$K_{\rm S0}$ (GPa) = 129.6(9) – 2.69(8)× $C_{\rm H20}$	$(R^2 = 0.9898)$	(14)
348	G_0 (GPa) = 79.3(6) - 1.45(5)× $C_{\rm H2O}$	$(R^2 = 0.9866)$	(15)
349	$V_{\rm P}$ (km/s) = 8.54(4) - 0.073(3)× $C_{\rm H2O}$	$(R^2 = 0.9724)$	(16)
350	$V_{\rm S}$ (km/s) = 4.95(3) – 0.036(2)× $C_{\rm H2O}$	$(R^2 = 0.9565)$	(17)

The liner fits show that 1 wt% H₂O reduces K_{S0} and G_0 by 2.69(8) GPa (2%) and 1.45(5) GPa (2%), respectively, We also find that the isothermal bulk modulus (K_{T0}) decreases by 2.71 GPa for every 1 wt% H₂O added along the forsterite-brucite join (Ross and Crichton 2001; Holl et al. 2006). On the other hand, a linear relation for density versus water content is plotted in *Fig.* 8(A). Addition of 1wt% H₂O reduces ρ by 0.029(2) g/cm³, and linear regressions for moduli and velocities versus density (Birch's law) could be fulfilled on basis of Eq. (13).

357 In summary, we have presented new thermoelastic properties of OH-chondrodite, which was 358 synthesized at 12 GPa and 1523 K, and coexists with OH-clinohumite and hydrous olivine. The synthesis 359 conditions correspond to the pressure-temperature condition around 360-km depth in a hydrous subducting 360 slab. The chondrodite sample was characterized by multiple experimental methods: electron microbe, 361 Raman spectroscopy, single-crystal X-ray diffraction (153–787 K), and finally by Brillouin spectroscopy. 362 For DHMS phases along the forsterite-brucite joint, every 1 wt% increase in C_{H2O} increases the mean 363 thermal expansion coefficient (α_0) by 3 %, decreases the moduli (K_{T0} , K_{S0} and G_0) by 2 %, and decreases 364 the seismic velocities (V_P and V_S) by ~ 8 %. Hydration also enlarges the Grüneisen parameters for phase A, 365 chondrodite and clinohumite by 17~32 %, compared with anhydrous forsterite. The elastic properties OH-366 chondrodite found in this study are consistent with the previous isothermal compression studies of Fe,F-367 free samples. The thermal expansion coefficient $\alpha(T)$ at ambient pressure is constrained by different models. 368 The Fei equation yields: $\alpha_V = 2.8(5) \times 10^{-9} (K^{-2}) \times T + 40.9(7) \times 10^{-6} (K^{-1}) - 0.81(3) (K)/T^2$ (150–800 K); 369 Suzuki and Kumar equations fit: $V_0(0 \text{ K}) = 365.8(3) \text{ Å}^3$, $\Theta = 741 \text{ K}$, $\gamma_{MGD} = 1.20(9)$ ($\gamma = 1.66(4)$ calculated 370 from Eq. 11). The thermoelastic properties reported in this study are important to anchoring more complete 371 P-V-T equations of state for chondrodite and other DHMS phases in future studies at simultaneous high 372 pressure and high temperature.

373

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375

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546 Figure Captions

547 Fig. 1 Composition of hydrous magnesium silicates in the hydrous peridotite system (MgO-SiO₂-

- 548 Mg(OH)₂). A: phase A; Nrb: Norbergite; Chn: chondrodite; Hu: humite; Chu: clinohumite; Fo: Forsterite;
- En: enstatite; B: phase B; Ahy B: anhydrous phase B; Shy B: supper hydrous phase B; Tlc: talc; Ant:
- antigorite; E: Phase E (non-stoichiometric, Si/Mg 1.8~2.1, C_{H20} 6~18 wt%); D: Phase D (Si/Mg 0.5~0.8,
- 551 C_{H2O} 13~19 wt%); 10 Å (C_{H2O} 7~13 wt%); Hy Wad: hydrous wadsleyite (C_{H2O} up to 3.3wt%).
- Fig. 2. Crystal structure of OH-chondrodite viewed down the *a* axis, with hydrogen positions shown as
 black spheres. M1-M3 refer to (Mg,Fe) octahedra, and the silicate tetrahedron is labeled Si.
- **Fig. 3** Unpolarized Raman spectra of the three OH-chondrodite crystals used in the Brillouin scattering experiments: (**A**) low-wavenumber region, (**B**) mid-wavenumber region, (**C**) high-wavenumber region showing O-H stretching bands, (**D**) high-wavenumber region of one sample shown with the two main O-H stretching bands deconvolved into six different components (dashed lines), sum of the fitted peaks (solid line) and spectral data points (grey dots). The unpolarized Raman spectra were collected in backscatter geometry with the incident laser perpendicular to each plane expressed as (*hkl*), fitted by Brillouin spectroscopy.
- Fig. 4. (A-C). Longitudinal acoustic (LA) and transverse acoustic (TA) velocities in three different planes
 (*hkl*) of OH-chondrodite as a function of crystallographic direction. The azimuthal angle is relative to an
 arbitrary starting angle. The solid curves were calculated using the best fit elastic constants for comparison
 with the measured velocities (filled circles).
- **Fig. 5.** (A) Unit-cell volume and principal expansion axes of OH-chondrodite as a function of temperature between 153 and 787 K. The solid curves are fitting to the data normalized to 153 K, using the Fei equation. Experimental uncertainties are smaller than the size of the symbols. (B) Volumetric thermal expansion coefficient as a function of temperature from Fei, Suzuki and Kumar equations, which are extrapolated to 1500 K.
- 570 Fig. 6. The average thermal expansion coefficients as a function of water content for DHMS phases along
- 571 the forsterite-brucite join. The temperature ranges are 300 787 K for forsterite (Fo, Ye et al. 2009),
- 572 clinohumite (Chu, Ye et al. 2013), and chondrodite (Chn, this study); 293 873 K for phase A (PhA,
- 573 Pawley et al. 1995); 299 593 K for brucite (Bru, Fukui et al. 2003).

- 574 Fig. 7. Unit-cell volumes of Fe-free OH-Chondrodite samples (Ross and Crichton 2001; Kuribayashi et al.
- 575 2001) as a function of pressure, with 3rd-order Birch-Murnaghan Equation of State (BM EoS) fitting. The
- 576 volumes are normalized to their respective V_0 values at ambient conditions.
- 577 Fig. 8. Elastic properties of DHMS along the forsterite-brucite join: (A) density as a function water content,
- **578** (B) bulk modulus (K_0) and shear modulus (G_0) as a function of water content, (C) compressional (V_P) and
- 579 shear-wave (V_S) velocities as a function of water content. Linear fits to the data (dashed lines) are given in
- the text (equations 13-17). References are given in Table 9.
- 581

		Pha	ase A	Chond	rodite	C	linohumi	te	
Formul	la	Mg ₇ S	$i_2O_{14}H_6$	Mg ₅ Si	$_{2}O_{10}H_{2}$	Ν	Ig ₉ Si ₄ O ₁₈ I	H_2	
$C_{\rm H2O}$ (v	vt%)	1	1.8	5.	.4		2.9		
ρ (g/cn	n ³)	2.9	$49(1)^{a}$	3.09	$9(1)^{b}$		3.134(1)	с	
α_0 (10-	⁶ K ⁻¹)	4	$6(2)^{d}$	38.0	$(9)^{b}$		36.8(6) ^c		
$K_{\rm S0}$ (G)	Pa)	10	$0(1)^{e}$	118	$(1)^{b}$		$119(2)^{e}$		
G_0 (GP	'a)	6	$1(1)^{e}$	70.1	$(5)^{b}$		$77(1)^{e}$		
V _P (km	√s)	7.8	$4(8)^{e}$	8.26	$(8)^{b}$		8.41(9) ^e		
Vs (km	√s)	4.5	$5(7)^{e}$	4.76	$(4)^{b}$		$4.96(8)^{e}$		
$K_{\rm T0}$ (G	Pa)	97.	$5(4)^{a}$	114($(1)^{b}$		119.4(7) ^f		
K _T '		6.0	$(1)^{a}$	5.5($5.5(1)^f$ $4.8(2)$		$4.8(2)^{f}$		
$C_{\mathrm{P}}\left(\mathbf{J}\cdot\mathbf{k}\right)$	$(g^{-1} \cdot K^{-1})$	924	$(1)^{g}$	874($(1)^{g}$		$884(1)^{h}$		
$\Theta_{ac}(K)$	$)^i$	726	(13)	741((10)		760(15)		
γ^i		1.0	59(8)	1.6	6(4)		1.50(4)		
a: Cric	hton and Ros	ss (2002):	b: This stu	dy; <i>c</i> : Ye et :	al. (2013	3); <i>d</i> : Pawle	ey et al. (1	995); e: Phan (2009);
refined	on the data f	from Ros	s and Crich	ton (2001) at	nd Kurit	bayashi et a	il. (2004);	g: Cynn et al. (1996);
Hollan	d and Powell	1998. i;	the Debye t	emperatures	and Gri	ineisen par	ameters a	re calculated in	this stu
Table 2	2. Compositio	on of OH	-chondrodit	e SZ0407B 1	from ele	ctron micro	oprobe an	alyses.	
	wt%		apfu [#]			wt%		apfu	
SiO2	34 09(6)	Si 1	945(3)		NiO	0.04(1)	Ni	0.0018(5)	
TiO	0.003(3)	Ti 0	0001(1)		MnO	0.008(3)	Mn	0.0010(3)	
Al_2O_3	0.005(3)	A1 0	0017(2)		CaO	0.000(3) 0.014(3)	Ca	0.0001(1) 0.0008(2)	
MgO	59.34(5)	Mg 5	5.047(4)		$H_2O^{\$}$	5.26	Н	2	
FeO	1.22(3)	Fe (0.058(2)		Total	100.00(6)		-	
#: aton	ns per formul	la unit.							
\$: The	wt% and apf	u values	for H ₂ O/H a	are calculated	d from s	toichiomet	ry, assum	ing that the mol	ar ratic
110'	2:10.						•		
H:O 18							-		
H:O 18							-		
H:O 18									
H:O IS Table 3	3. Atomic pos	sitions an	d occupanc	y factors fro	m X-ray	intensity s	scan (spac	e group: <i>P</i> 2 ₁ /b)	
H:O is Table 3 Atom	3. Atomic pos x/a	sitions an	d occupanc y/b	y factors fro $\frac{z}{c}$	m X-ray	intensity s	can (spac	e group: <i>P</i> 2 ₁ /b)	
Table 3 Atom	3. Atomic pos x/a 0.5	sitions an	d occupanc y/b 0	y factors fro z/c 0.5	m X-ray Mg	r intensity s oc : 0.991(2)	can (spac cupancy / Fe: 0.00	e group: <i>P</i> 2 ₁ /b) 9(2)	
Table 3 Atom M1 M2	3. Atomic pos $\frac{x/a}{0.5}$	<u>sitions an</u> (6) 0.1	<u>d occupanc</u> y/b 0 7542(3)	$\frac{y \text{ factors fro}}{z/c}$ 0.5 0.30892(4)	m X-ray Mg Mg	<u>oc</u> : 0.991(2) : 0.992(2)	<u>ccan (spac</u> cupancy / Fe: 0.00 / Fe: 0.00	e group: P2 ₁ /b) 9(2) 8(2)	
Table 3 Atom M1 M2 M3	3. Atomic pos $\frac{x/a}{0.5}$ 0.00761(0.48780)	<u>sitions an</u> (6) 0.1 (6) 0.8	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3)	y factors fro z/c 0.5 0.30892(4) 0.07707(4)	m X-ray Mg Mg Mg	intensity s oc : 0.991(2) : 0.992(2) : 0.989(2)	ccan (spac cupancy / Fe: 0.009 / Fe: 0.003 / Fe: 0.01	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	
H:O is Table 3 Atom M1 M2 M3 Si	3. Atomic pos x/a 0.5 0.00761(0.48780(0.077690	<u>sitions an</u> (6) 0.1 (6) 0.8 (4) 0.1	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2)	y factors fro z/c 0.5 0.30892(4) 0.07707(4) 0.70194(2)	m X-ray Mg Mg Mg	intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	ccan (spac cupancy / Fe: 0.009 / Fe: 0.001 / Fe: 0.01 3(2)	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	
Table 3 Atom M1 M2 M3 Si O1	3. Atomic pos x/a 0.5 0.00761(0.48780(0.07769(0.77577(sitions an (6) 0.1 (6) 0.8 (4) 0.1 11) 0.0	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2) 0294(5)	y factors fro z/c 0.5 0.30892(4) 0.07707(4) 0.70194(2) 0.29661(7)	m X-ray Mg Mg Mg	v intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	ccan (spac cupancy / Fe: 0.009 / Fe: 0.001 / Fe: 0.01 3(2) 1	e group: <i>P</i> 2 ₁ /b) 9(2) 8(2) 1(2)	
Table 3AtomM1M2M3SiO1O2	3. Atomic pos x/a 0.5 0.00761(0.48780(0.07769(0.77577(0.72759(sitions an (6) 0.1 (6) 0.8 (4) 0.1 11) 0.0 11) 0.2	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2) 0294(5) 4411(5)	y factors fro <u>z/c</u> 0.5 0.30892(4) 0.07707(4) 0.70194(2) 0.29661(7) 0.12853(7)	m X-ray Mg Mg Mg	v intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	ccan (spac cupancy / Fe: 0.009 / Fe: 0.003 / Fe: 0.01 3(2) 1	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	
Table 3AtomM1M2M3SiO1O2O3	3. Atomic pos x/a 0.5 0.00761(0.48780(0.07769(0.77577(0.72759(0.22561(<u>sitions an</u> (6) 0.1 (6) 0.8 (4) 0.1 11) 0.0 11) 0.2 11) 0.1	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2) 0294(5) 4411(5) 6797(5)	y factors fro z/c 0.5 0.30892(4) 0.07707(4) 0.70194(2) 0.29661(7) 0.12853(7) 0.52799(7)	<u>m X-ray</u> Mg Mg Mg	r intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	ccan (spac cupancy / Fe: 0.009 / Fe: 0.001 / Fe: 0.01 3(2) 1 1	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	
H:O is Table 3 Atom M1 M2 M3 Si O1 O2 O3 O4	3. Atomic pos x/a 0.5 0.00761(0.48780(0.07769(0.77577(0.72759(0.22561(0.26251(sitions an (6) 0.1 (6) 0.8 (4) 0.1 11) 0.0 11) 0.2 11) 0.1 11) 0.8	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2) 10294(5) 4411(5) 6797(5) 5683(5)	y factors fro z/c 0.5 0.30892(4) 0.70194(2) 0.29661(7) 0.12853(7) 0.52799(7) 0.29738(7)	m X-ray Mg Mg Mg	r intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	acan (spac cupancy / Fe: 0.009 / Fe: 0.01 3(2) 1 1 1 1	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	
H:O is Table 3 Atom M1 M2 M3 Si O1 O2 O3 O4 O5	3. Atomic pop x/a 0.5 0.00761(0.48780(0.077690 0.77577(0.72759(0.22561(0.26251(0.26372(sitions an (6) 0.1 (6) 0.8 (4) 0.1 11) 0.0 11) 0.2 11) 0.1 11) 0.8 14) 0.0	<u>d occupanc</u> <u>y/b</u> 0 7542(3) 8398(3) 4073(2) 10294(5) 4411(5) 6797(5) 5683(5) 6017(6)	y factors fro z/c 0.5 0.30892(4) 0.07707(4) 0.29661(7) 0.12853(7) 0.52799(7) 0.29738(7) 0.10540(9)	m X-ray Mg Mg Mg	r intensity s oc : 0.991(2) : 0.992(2) : 0.989(2) 0.98	acan (spac cupancy / Fe: 0.009 / Fe: 0.01 3(2) 1 1 1 1 1 1	e group: P2 ₁ /b) 9(2) 8(2) 1(2)	

Table 1. Thermoelastic properties of Phase A, chondrodite and clinohumite (F-free) at ambient conditions.

629	Table 4. Bond lengths (Å) and polyhedral volumes (Å ³).									
630	M1		M2		M3					
631	O1×2:	2.0805(5)	O1:	2.0713(6)	01:	2.2413(6)				
632	O3×2:	2.1253(5)	O2:	2.2257(6)	O2:	2.0140(6)				
633	O4×2:	2.1227(5)	O3:	2.0387(6)	O2:	2.1123(6)				
634	Avg. bond:	2.1095(5)	O3:	2.1772(6)	O4:	2.1363(6)				
635	Poly. $V(Å^3)$:	12.04(1)	O4:	2.1983(6)	O5:	2.0604(7)				
636			O5:	2.0630(7)	O5:	2.0819(7)				
637	Si		Avg. bond:	2.1290(7)	Avg. bond:	2.1077(7)				
638	01:	1.6461(6)	Poly. $V(Å^3)$:	12.47(1)	Poly. $V(Å^3)$:	12.13(1)				
639	O2:	1.6427(6)								
640	O3:	1.6463(6)	Н							
641	O4:	1.6151(6)	O5:	1.06(2)						
642	Avg. bond:	1.6376(6)								
643	Poly. $V(Å^3)$:	2.22(1)								

645
646 Table 5. Selected bond length (Å) for chondrodite samples from this and previous studies.

647		Fe $\%^*$	$F \%^+$	Ti %*	α (°)	$V(\text{\AA}^3)$	М	2-OF5	M3-OF51	M3-OF5 ₂
648	SZ0407B ^{\$}	1.1	0	0	108.702(7)	367.50(4) 2.0)628(5)	2.0599(6)	2.0825(6)
649	Y77 ^{\$}	0	0	0	108.71(5)	368.7(2)	2.	133(3)	2.056(3)	2.091(3)
650	B02	1.4	77	0	109.11(1)	357.26(3) 2.	.054(8)	2.011(8)	2.015(9)
651	G70	1	65	0	109.059(2)	359.30(1) 2	.045(2)	2.016(1)	2.042(2)
652	O00	0.8	62	0	109.08(2)	359.47(1) 2	.051(3)	2.019(3)	2.045(3)
653	F01 ^{&}	5.6	51	0.46	109.097(2)	363.14(4) 2	.060(1)	2.033(1)	2.059(1)
654	F02*	5.6	58	0.46	109.065(1)	361.98(2) 2	.057(8)	2.01(1)	2.055(9)
655	K06 ^{&}	5.6	58	0.46	109.065(3)	361.98(2) 2	.068(5)	2.048(5)	2.067(5)
656	K04 ^{&}	4.4	37	0.4	109.07(1)	361.0(2)	2	.057(1)	2.024(2)	2.053(1)
657	F78	12.5	0	8.4	109.33(1)	363.83(5) 2.	051(2)	1.976(2)	1.987(2)
658 659 660 661 662 663 664 665 666 667	F/812.508.4109.33(1)363.83(5)2.051(2)1.976(2)1.987SZ0407B: this study; Y77: Yamamoto (1977); B02: Berry and James (2002); G70: Gibbs et al. (1970); Ottolini et al. (2000); F01: Friedrich et al. (2001); F02: Friedrich et al. (2002); K06: Kunz et al. (2006); K04: Kuribayashi et al. (2004): F78: Fujino and Takéuchi (1978). *: Atomic percentage in the total metallic cations. +: Molar ratio of F/(F+OH). \$: synthesized from experiments, while others are from natural sources. &: data measured at ambient condition.Table 6. Average thermal expansion coefficients for the unit-cell parameters and the principal expansion								al. (1970); O00: al. (2006); l expansion	
668	Unit-cell p	aramete	ers	Pri	ncipal expansio	n axes	0	rientations	3	
669	α ₀ (10 ⁻⁶ K ⁻¹)			α ₀ (10 ⁻⁶ K	-1)	a	b	с	
670	a 12	.7(3)		X_1	12.7(3)		1	0	0	
671	b 11	.5(2)		X_2	10.6(2)		0	0.8353	0.5499	
672	c 14	4.6(4)		X_3	14.7(4)		0 .	0.1636	0.9865	
673	α 1.4	43(10)								
674	V 38	3.0(9)								
675 676 677										

678 Table 7. Single-crystal elastic constants (in GPa) for chondrodite at ambient conditions.

		0 ,		/			
679		This study	SB89		This study	SB89	
680	longitudina	al		off-diagona	al		
681	$C_{11}(C_{33})$	188.0(7)	213.4(15)	$C_{12}(C_{13})$	61(10)	70(3)	
682	$C_{22}(C_{11})$	278.0(10)	275.3(15)	$C_{13}(C_{23})$	71.4(7)	59(2)	
683	$C_{33}(C_{22})$	195.7(7)	198.4(12)	$C_{23}(C_{12})$	74(3)	67(3)	
684	shear			$C_{15}(C_{14})$	3.9(4)	7.2(10)	
685	$C_{44}(C_{66})$	64.0(10)	69.7(6)	$C_{25}(C_{24})$	2.6(10)	-1.7(12)	
686	$C_{55}(C_{44})$	68.3(4)	72.1(9)	$C_{35}(C_{34})$	2.5(4)	-2.6(8)	
687	$C_{66}(C_{55})$	71.3(5)	75.2(7)	$C_{46}(C_{56})$	4.1(6)	-0.7(4)	

688 *SB89*:: Sinogeikin and Bass (1999).

689 The elastic constants are given in the standard monoclinic coordinate system (*b*-unique, $P2_1/c$), while the 690 values inside parentheses are those for an *a*-unique coordinate system ($P2_1/b$).

Table 8. Aggregate elastic moduli and pressure derivative of bulk modulus for chondrodite

694		F/(F+OH)	Fe (mol%)	$K_{\rm T0}~({\rm GPa})$	K_{T} '	G_0 (GPa)
695	This study	0	1.1	119.4 ^{&}		71.1 (Voigt bound)
696				116.4 ^{&}		69.2 (Reuss bound)
697				117.9(12) ^{&}		70.1(5) (VRH average)
698				114.2^{+}	$5.5(1)^{\#}$	
699	Sinogeikin and Bass (1999)	0.32	5.5	118.4(16)&		75.6(7)
700	Ross and Crichton (2001)	0	0	115.7(8)	4.9(2)	
701	Kuribayashi et al. (2004)	0	0	117(2)	4^{*}	
702	Kuribayashi et al. (2004)	0.37	4.4	124.1(4)	4^{*}	
703	Friedrich et al. (2002)	0.55	0	117(4)	5.6(1)	
704	Faust and Knittle (1994)	0.69	0	136.2	3.7^{*}	
705						

705 &: adiabatic bulk modulus.

706 +: K_{T0} calculated from Reuss bound K_{S0} .

#: Refined on the datasets of Ross and Crichton (2001) and Kuriyabashi et al. (2004), with $K_{T0} = 114.2$ GPa.

*: fixed without refinement.

709

691 692

	H_2O	density	Fe	$K_{ m S0}$	G_0	$V_{ m P}$	$V_{\rm S}$
	(wt%)	(g/cm^3)	(mol%)	(GPa)	(GPa)	(km/s)	(km/s)
Olivine:							
Suzuki et al. (1983)	0	3.225(1)	0	128.9(2)	81.4(2)	8.58(1)	5.02(1)
Zha et al. (1996)	0	3.221(1)	0	128.8(5)	81.6(2)	8.59(3)	5.03(1)
Webb (1989)	0	3.350(1)	10	129.5(2)	77.5(2)	8.34(1)	4.81(1
Zha et al. (1998)	0	3.343(1)	10	131.1(19)	79.4(8)	8.42(9)	4.87(4)
Jacobsen et al. (2008, 2009)	0.8	3.240(1)	3	125.2(2)	77.7(1)	8.40(1)	4.90(1)
Jacobsen et al. (2008, 2009)	0.9	3.180(1)	0	125.4(4)	79.6(3)	8.53(2)	5.00(2
Clinohumite:							
Fritzel and Bass (1997)	2.0^{a}	3.261(1)	6.5	125(2)	73(5)	8.26(9)	4.7(3)
Phan (2009)	2.9	3.134(1)	0	119(2)	77(1)	8.41(9)	4.96(8
Chondrodite:							
Sinogeikin and Bass (1999)	3.3 ^b	3.227(1)	9.4	118(2)	75.6(7)	8.24(8)	4.84(5)
This study	5.3	3.099(1)	1.1	118(1)	70.6(5)	8.26(8)	4.76(4
Phase A:							
Sanchez-Valle et al. (2008)	11.7	2.976(1)	1.9	106(1)	61(1)	7.93(8)	4.53(8
Phan (2009)	11.8	2.949(1)	0	100(2)	61(1)	7.84(9)	4.55(8)
Brucite:							
Jiang et al. (2006)	30.9	2.380(1)	0	43.8(8)	35.2(3)	6.2(1)	3.85(3
Xia et al. (1998)	30.9	2.380(1)	0	46(1)	34.9(5)	6.2(1)	3.83(5

Table 9. Studies of elasticity for DHMS along the olivine-brucite join, listed in the order of H₂O content increasing 713

739 Appendix 1. Anisotropic displacement parameters (Å²).

Ator	m U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}	$U_{ m eq}$
M1	0.0058(2)	0.0069(2)	0.0061(2)	0.0014(1)	0.0009(1)	0.0005(1)	0.0064(1)
M2	0.0085(2)	0.0065(2)	0.0071(2)	0.0029(1)	-0.0004(1)	0.0004(1)	0.0072(1)
M3	0.0072(1)	0.0094(2)	0.0074(1)	0.0040(1)	-0.0004(1)	0.0004(1)	0.0077(1)
Si	0.0042(1)	0.0056(1)	0.0054(1)	0.0022(1)	0.0000(0)	0.0000(0)	0.0050(1)
01	0.0074(2)	0.0065(2)	0.0084(2)	0.0030(1)	0.0005(2)	-0.0001(1)	0.0073(1)
O2	0.0068(2)	0.0074(2)	0.0070(2)	0.0015(2)	0.0002(1)	-0.0001(1)	0.0073(1)
O3	0.0066(2)	0.0082(2)	0.0065(2)	0.0035(1)	0.0002(1)	-0.0001(1)	0.0068(1)
O4	0.0043(2)	0.0084(2)	0.0075(2)	0.0027(2)	0.0001(1)	0.0000(1)	0.0067(1)
05	0.0189(3)	0.0100(2)	0.0202(3)	0.0080(2)	0.0127(2)	0.0059(2)	0.0156(1)
Н							0.054 (fixed)

751
752
753 Appendix 2. Unit-cell parameters of chondrodite as functions of temperature, as well as the principle expansion axes, which are normalized to the ones at 153 K

		,								_
755	<i>T</i> (K)	<i>a</i> (Å)	<i>b</i> (Å)	<i>c</i> (Å)	lpha (°)	$V(\text{\AA}^3)$	X_1/X_{10}	X_2/X_{20}	X_3/X_{30}	
756	153(2)	4.7402(3)	10.3341(8)	7.8899(8)	108.687(8)	366.12(4)	1	1	1	
757	203(2)	4.7419(3)	10.3387(8)	7.8926(8)	108.693(8)	366.53(4)	1.00036	1.00030	1.00045	
758	253(2)	4.7438(3)	10.3439(7)	7.8969(7)	108.698(7)	367.04(4)	1.00078	1.00076	1.00099	
759	300(2)	4.7459(2)	10.3480(7)	7.9002(7)	108.702(7)	367.50(4)	1.00120	1.00114	1.00142	
760	350(3)	4.7489(2)	10.3530(7)	7.9052(7)	108.706(8)	368.13(4)	1.00183	1.00164	1.00201	
761	396(3)	4.7516(2)	10.3583(7)	7.9105(8)	108.708(9)	368.77(4)	1.00240	1.00217	1.00266	
762	443(3)	4.7546(2)	10.3637(7)	7.9168(7)	108.715(9)	369.47(4)	1.00304	1.00267	1.00344	
763	495(4)	4.7578(2)	10.3695(7)	7.9226(8)	108.722(9)	370.18(4)	1.00371	1.00318	1.00418	
764	544(4)	4.7610(2)	10.3771(7)	7.9286(7)	108.735(9)	370.95(5)	1.00439	1.00380	1.00498	
765	587(4)	4.7640(2)	10.3827(8)	7.9345(8)	108.737(8)	371.67(5)	1.00502	1.00435	1.00571	
766	635(5)	4.7674(3)	10.3892(8)	7.9403(9)	108.750(9)	372.41(6)	1.00574	1.00486	1.00648	
767	685(5)	4.7704(3)	10.3951(8)	7.9470(9)	108.764(9)	373.14(6)	1.00637	1.00534	1.00734	
768	736(6)	4.7738(3)	10.4022(8)	7.9545(9)	108.771(9)	373.99(6)	1.00709	1.00599	1.00829	
769	787(6)	4.7773(3)	10.4099(8)	7.9619(9)	108.795(9)	374.84(6)	1.00783	1.00653	1.00929	
770										-





784 Figure 3.







797 Figure 7.

