



Article Calculated Elasticity of Al-Bearing Phase D

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Abstract: Using first-principles calculations, this study evaluates the structure, equation of state, and elasticity of three compositions of phase D up to 75 GPa: (1) the magnesium endmember $[MgSi_2O_4(OH)_2]$, (2) the aluminum endmember $[Al_2SiO_4(OH)_2]$, and (3) phase D with 50% Alsubstitution $[AlMg_{0.5}Si_{1.5}O_4(OH)_2]$. We find that the Mg-endmember undergoes hydrogen-bond symmetrization and that this symmetrization is linked to a 22% increase in the bulk modulus of phase D, in agreement with previous studies. $Al_2SiO_4(OH)_2$ also undergoes hydrogen-bond symmetrization, but the concomitant increase in bulk modulus is only 13%—a significant departure from the 22% increase of the Mg-end member. Additionally, Al-endmember phase D is denser (2%–6%), less compressible (6%–25%), and has faster compressional (6%–12%) and shear velocities (12%–15%) relative to its Mg-endmember counterpart. Finally, we investigated the properties of phase D with 50% Al-substitution $[AlMg_{0.5}Si_{1.5}O_4(OH)_2]$, and found that the hydrogen-bond symmetrization, equation of state parameters, and elastic constants of this tie-line composition cannot be accurately modeled by interpolating the properties of the Mg- and Al-endmembers.

Keywords: phase D; dense hydrous magnesium silicate; hydrogen bond symmetrization; elasticity



Citation: Thompson, E.C.; Campbell, A.J.; Tsuchiya, J. Calculated Elasticity of Al-Bearing Phase D. *Minerals* 2022, 12, 922. https://doi.org/10.3390/ min12080922

Academic Editor: Jordi Ibanez-Insa

Received: 25 May 2022 Accepted: 20 July 2022 Published: 22 July 2022

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1. Introduction

Mineral physics experiments and first-principles calculations have identified several mechanisms for water storage inside the Earth, including nominally anhydrous and hydrous phases, but few of these phases are stable at the extreme pressure and temperature conditions of the Earth's lower mantle. As serpentine-bearing lithospheric plates subduct, serpentine exposed to the increasing pressures and temperatures of the geotherm decomposes into a series of dense hydrous magnesium silicates (DHMSs) [1]. These DHMSs contain wt.% quantities of water (OH⁻) and are important carriers of water in subduction zones [2–6]. Of the known DHMSs, phase D, (Mg,Al)(Si,Al)₂O₄(OH)₂, has the second highest pressure stability, rendering phase D capable of transporting water through the transition zone and into the lower mantle [7–11].

Recent studies indicate that aluminum substitution into DHMSs increases the thermodynamic stability of these phases [7,12–18], and that Al-bearing DHMSs may host more water than their magnesium endmember counterparts [18,19]. Additionally, Al-bearing phase D is a likely precursor to the solid solution formed by phase H [MgAlO₂(OH)₂] and δ -(Al,Fe)OOH—a solid solution with *P*-*T* stability that extends to the core-mantle boundary [13,14,20,21]. Owing to the important role that Al-bearing phase D may play in the storage and cycling of hydrogen in the Earth's lower mantle, this study evaluates the influence of Al-substitution on the structure, equation of state, and elasticity of phase D using first-principles density functional theory (DFT) calculations.

Magnesium endmember phase D $[MgSi_2O_4(OH)_2]$ has trigonal symmetry and is in the $P\bar{3}1m$ space group [22]. The crystal structure is based on a hexagonal closest packed array of O atoms, with non-hydrogen cations occupying two different octahedrally coordinated

sites. In Mg-endmember phase D, the SiO₆ and MgO₆ octahedra occur in two separate layers stacked along *c*-axis, leading these sites to be referred to as the S-site and M-site, respectively (Figure 1a). In Al-bearing phase D, the aluminum occupies both the S- and M-sites [18] (Figure 1b). Aluminum substitutes into phase D via a Tschermak Si⁴⁺ + Mg²⁺ \leftrightarrow 2Al³⁺ substitution, and experiments report a range of compositions, including the near Al-endmember composition referred to as 'super-aluminous' phase D [19]. Based on single crystal X-ray diffraction structure refinement, this near Al-endmember phase D is also in the $P\bar{3}1m$ space group, with a high degree of Si/Al disordering and decreased octahedral distortion relative to the Mg-endmember [19]. Within the S-site layer, octahedra are edge-sharing with 1/3 of the sites vacant, producing brucite-like layers, whereas in the M-site layer the octahedra do not share edges, thereby creating space for hydroxyl bonds.



Figure 1. Crystal structures of phase D at 0 GPa including: (a) Mg-endmember $[MgSi_2O_4(OH)_2]$ (one formula unit) and (b) disordered Al-endmember $[Al_2SiO_4(OH)_2]$ (two formula units). Images were generated in VESTA [23]. Aluminum atoms are aqua, magnesium atoms are orange, silicon atoms are dark blue, oxygen are red, and hydrogen are white spheres.

Mg-endmember phase D undergoes pressure-induced hydrogen bond symmetrization at approximately 40 GPa, which was predicted by first-principles calculations [24] and confirmed by high-pressure X-ray diffraction experiments [25]. Hydrogen-bond symmetrization is the phenomenon in which the hydroxyl bonds (O–H) in a material evolve with increased pressure such that they become equal in length to the hydrogen bonds (O···H) (Figure 2). Hydrogen-bond symmetrization in MgSi₂O₄(OH)₂ profoundly impacts its compressibility, increasing the bulk modulus by up to 20% [24–26]. However, as hydrogen bond symmetrization has yet to be reported in Al-bearing phase D, it is important to probe the influence of Al-substitution on this phenomenon. Using first-principles calculations, this study evaluates three compositions of phase D: (1) the magnesium endmember [MgSi₂O₄(OH)₂], (2) the aluminum endmember [Al₂SiO₄(OH)₂], and phase D with 50% Al-substitution [AlMg_{0.5}Si_{1.5}O₄(OH)₂], to evaluate the influence of Al-substitution on the structure, equation of state, and elasticity of phase D.



Figure 2. Comparison of hydrogen bonding in Mg-endmember phase D before and after pressure induced hydrogen bond symmetrization. (**a**) shows the hydrogen off-center (HOC) structure at 0 GPa with asymmetrical O–H···H bonding, where the dashed black lines indicate the longer but more compressible hydrogen bridge bonds (O···H) and the solid black lines indicate the shorter but stiffer hydroxyl bonds (O–H). (**b**) at 70 GPa, the hydrogen are now centered (i.e., symmetric) with respect to the two neighboring oxygens. Images were generated in VESTA [23].

2. Methods

Density functional theory (DFT) based calculations were used to evaluate the structure and elasticity of three compositions of phase D [MgSi₂O₄(OH)₂, AlMg_{0.5}Si_{1.5}O₄(OH)₂, $Al_2SiO_4(OH)_2$] as a function of pressure from 0 to 75 GPa in 5 GPa pressure increments. Although previous studies have calculated the structure and elasticity of Mg-endmember phase D [26,27], these calculations were repeated to enable direct comparison between all three phase D compositions using the same pseudopotentials for all calculations. The aluminum endmember composition values reported herein are based on evaluations using a 2-unit supercell to introduce a degree of disordering (Figure 1b). An ordered structure of the Al-endmember composition was also evaluated (1 unit cell), but exhibited elevated enthalpy relative to the disordered structure across the entire pressure range of this study and is therefore less stable relative to its disordered counterpart. Two different supercells (8-unit cells) of AlMg_{0.5}Si_{1.5}O₄(OH)₂ were evaluated to assess the influence of cation disordering on phase stability and material properties. These two AlMg_{0.5}Si_{1.5}O₄(OH)₂ supercells are referred to hereafter as '88-1' and '88-2'. Using two supercells enables us to probe the interplay of composition and structure on the elasticity of AlMg_{0.5}Si_{1.5}O₄(OH)₂ and helped us to better assess the reliability of modeling the properties of intermediate compositions by interpolating those of the Mg- and Al-endmembers. A full exploration of all solid solution compositions and possible atomic configurations is beyond the scope of this study. Atomic positions of the fully optimized structures of $MgSi_2O_4(OH)_2$, $Al_2SiO_4(OH)_2$, and both supercells of $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ at 0 GPa can be found in Tables A1–A4.

First-principles simulations were performed using Quantum ESPRESSO [28], in which we applied the generalized gradient approximation (GGA) to the exchange-correlation functional [29], as it more accurately describes hydrogen bonding compared to the local density approximation [30,31]. As the present study did not include temperature and quantum zero-point vibration effects, we did not employ the empirical dispersion correction of Grimme et al. 2010 [32]; however, the influence of such corrections on the van der Waals interactions and hydrogen bonding of phase D should be evaluated in the future. The effective interaction of core electrons was approximated using previously evaluated norm-conserving pseudopotentials [33] and electronic wave functions were expanded in plane-waves with an energy cutoff of 80 Ry. The irreducible Brillouin zone was sampled by Monkhorst-Pack meshes of $5 \times 5 \times 4$, $5 \times 5 \times 2$, and $3 \times 3 \times 2$ for the Mg-endmember, Al-endmember, and tie-line compositions, respectively [34]. The effects of larger energy cutoffs and k-point sampling were found to be negligible. Elastic constants were determined

by applying strains of 0.005–0.01 to the optimized (0 K) structures, maintaining linear stress-strain relations [35].

3. Results

3.1. Structure and Hydrogen Bond Symmetrization

Optimized structures of the Mg- and Al-endmember compositions, as well as both structures of the tie-line composition (50% Al-substitution), were evaluated to determine the influence of Al-substitution on the structure and hydrogen-bonding of phase D. The resultant structures of both endmember and intermediate compositions are consistent with the previously described trigonal phase D structure (Figure 1), with minor triclinic distortion (<2%) consistent with previous calculations [26]. Across the pressure range examined, the Al-endmember structures exhibit the highest degree of distortion (0.8%-1.8%) while the structures of the tie-line composition (88-2) exhibit the lowest degree of distortion (0.2%–0.4%). Structural parameters including lattice parameters, hydroxyl and hydrogen bond lengths, and O—H \cdots O bond angles as a function of pressure from 0 to 75 GPa are reported in Tables A5–A7, respectively. In agreement with previous experimental and theoretical studies [25,27,36,37], we find that in Mg-endmember phase D the *c*-axis is more compressible than the *a*-axis at low pressures (<40 GPa) but at pressures above 40 GPa the c/a ratio becomes nearly pressure independent as shown in Figure 3. This disparity in axial compression is also observed in the Al-endmember and tie-line compositions, but the degree of this disparity, i.e., the magnitude of the negative slope of the c/a ratio as a function of pressure, is significantly reduced and limited to pressures below 30 GPa in these Al-bearing compositions (Figure 3).



Figure 3. The *c*/*a* ratio of Mg-endmember phase D (red circles), Al-endmember phase D (blue squares), and the tie-line composition in the 88-1 (purple triangles) and 88-2 structures (lavender diamonds). Grey symbols (open squares, crosses, open triangles, and open circles) show literature values from [25,27,36,37], respectively.

Similar to previous studies [14,24,25], we find that Mg-endmember phase D undergoes pressure-induced hydrogen bond symmetrization at 45 GPa. In other words, at and above pressures of 45 GPa the hydroxyl bond length (r_{O-H}) is equal to the hydrogen bond length (r_{O-H}) (Table A6). We find that Al-endmember phase D also undergoes a pressure-induced hydrogen bond symmetrization, albeit at the slightly lower pressure of 40 GPa.

Conversely, neither configuration of $AIMg_{0.5}Si_{1.5}O_4(OH)_2$ underwent complete pressureinduced hydrogen bond symmetrization in the pressure range of this study (0–75 GPa), as the more complex cation disordering introduced additional non-degenerate hydrogen sites. The pressure dependence of these sites vary, likely due to differences in nearest-neighbors and next-nearest neighbor cation occupancy [38]. Tables containing information regarding the hydrogen ($r_{O...H}$) and hydroxyl (r_{O-H}) bond lengths, as well as the O—H···O bond angles for all three compositions are reported in Tables A6 and A7, respectively. Our results deviate from the VASP ab initio calculations of Panero and Caracas (2020) [14], who reported that only roughly a quarter of hydrogen bonds in Al-endmember phase D symmetrize at pressure and that symmetrization in intermediate compositions is incremental and does not involve all bonds.

Although there is agreement in the literature concerning the existence and magnitude of a pressure-dependent evolution of the *c/a* ratio in Mg-endmember phase D, as well as its eventual stabilization at high pressures, no consensus exists regarding the cause. Furthermore, the pressure at which the *c/a* ratio is reported to stabilize varies widely, with reported stabilization pressures of 14 GPa [39], 20 GPa [40], 25 GPa [36], 35 GPa [41], 40 GPa [24] and 48 GPa [25]. In experimental studies, these differences may be attributed to differences in sample compositions including non-stoichiometric Mg/Si ratios, variable water contents, and Al- and Fe-substitution, as well as the wide range of pressure transmitting media used including ZrO₂, MgO, and Ne.

The larger question is whether the pressure evolution of the lattice parameters and eventual pressure independence of the c/a ratio is directly tied to hydrogen bond symmetrization [25], hydrogen bond disordering that occurs as a precursor to symmetrization as observed in δ -AlOOH [42], or the result of the layered structure of phase D [43]. Hydrogen bond symmetrization has been linked to shifts in axial compression both pre- and post-symmetrization in other phases [44–47], and has been described as the primary driver of the aforementioned pressure-dependent evolution of c/a ratios in phase D [24]. In this study, all three compositions of phase D have *c*-axes which are more compressible than the *a*-axes at low pressure, yet hydrogen bond symmetrization only occurs in the Al- and Mg-endmembers, and occurs at pressures slightly higher that the pressure at which the c/a ratio stabilizes. Therefore, it is likely that the observed low-pressure anisotropy is tied to the layered nature of phase D, with differences in the Al- and Mg-endmembers tied to the relative stiffness of the AlO₆, MgO₆, and SiO₆ units. However, c/a ratio stabilization is seemingly a prerequisite for hydrogen bond symmetrization, such that the two phenomenon can appear coincident. Furthermore, in intermediate compositions of Al-bearing phase D, hydrogen bond symmetrization is not expected, but the pressure-dependence of the c/a ratio will likely reflect the compressibilities and configurations of the constituent cation polyhedra.

3.2. Equation of State

Optimized (0 K) structures of the endmember and tie-line compositions were used to evaluate the volume-pressure (*V*-*P*) relationship of these phases, by fitting them to third-order Birch-Murnaghan equations of state (EOSs) [48]:

$$P(V) = \frac{3K_0}{2} \left[\left(\frac{V_0}{V}\right)^{\frac{7}{3}} - \left(\frac{V_0}{V}\right)^{\frac{5}{3}} \right] \left\{ 1 + \frac{3}{4} \left(K'_0 - 4\right) \left[\frac{V_0}{V}^{\frac{2}{3}} - 1 \right] \right\}$$
(1)

in which K_0 is the bulk modulus at ambient pressure, K'_0 is the first pressure derivative of the bulk modulus, and V_0 is the reference volume and was treated as a free parameter. The Birch-Murnaghan equation of state parameters (K_0 , K'_0 , V_0) resulting from these fits are shown in Table 1. As evidenced by the positive slopes in Figure 4 which shows Eulerian strain (f) versus normalized pressure (F_E), the third-order Birch-Murnaghan equation of state better describes the compressive behavior of all examined compositions of phase D in this study. However, parameters fit to second-order equations of state (i.e., $K'_0 = 4$) are included in Table 1 to enable direct comparison with literature values. Hydrogen bond symmetrization

of Mg-endmember phase D has been previously reported to produce a significant decrease in the compressibility of the hydrogen-symmetric structure (HC) compared to that of the hydrogen off-center (HOC) structure [24]. Consistent with this phenomenon, our F_E-f plot reveals discontinuities in the compressibility of the Al- and Mg-endmembers at 35 and 40 GPa, respectively. With this in mind, HOC and HC structures of the Mg- and Al-endmember were fit separately, deriving distinct sets of equation of state parameters (Table 1, Figure 5). Phase D with 50% Al-substitution did not undergo pressure induced hydrogen bond symmetrization and the F_E-f plot revealed no discontinuities in either the 88-1 or 88-2 configuration. Therefore, optimized structures of AlMg_{0.5}Si_{1.5}O₄(OH)₂ spanning the entire pressure range (0–75 GPa) were fit to single equations of state for each configuration.

Table 1. Equations of state parameters for phase D of varying compositions. The pressure range of each study is noted, and where the authors fit hydrogen off-center (HOC) and hydrogen-centered (HC) structures independently the structure is indicated in brackets. The structure of the tie-line compositions from this study are in brackets. Values in parentheses are uncertainties on the last digit as reported by the original authors.

Study	Composition	Pressure (GPa)	V_0 (Å ³)	K ₀ (GPa)	$\mathbf{K_0}'$
This study	$MgSi_2O_4(OH)_2$	0–75	86.20 (4)	145 (1)	5.08 (4)
This study	$MgSi_2O_4(OH)_2$	0–75	85.5 (2)	173 (2)	4 (fixed)
This study	MgSi ₂ O ₄ (OH) ₂	0-35 [HOC]	86.14 (1)	149.0 (3)	4.79 (2)
This study	$MgSi_2O_4(OH)_2$	0-35 [HOC]	85.99 (4)	160 (1)	4 (fixed)
This study	$MgSi_2O_4(OH)_2$	40-75 [HC]	84.54 (9)	182 (2)	4.26 (5)
This study	$MgSi_2O_4(OH)_2$	40–75 [HC]	84.05 (3)	194.6 (5)	4 (fixed)
This study	$Al_2SiO_4(OH)_2$	0–75	82.26 (2)	186.9 (9)	4.77 (3)
This study	$Al_2SiO_4(OH)_2$	0-75	81.88 (9)	209 (2)	4 (fixed)
This study	$Al_2SiO_4(OH)_2$	0-30 [HOC]	82.25 (1)	188.0 (9)	4.67 (7)
This study	$Al_2SiO_4(OH)_2$	0-30 [HOC]	82.19 (4)	197 (1)	4 (fixed)
This study	$Al_2SiO_4(OH)_2$	35–75 [HC]	81.47 (9)	212.4(3)	4.21 (7)
This study	$Al_2SiO_4(OH)_2$	35–75 [HC]	81.20 (2)	222.5 (5)	4 (fixed)
This study	AlMg _{0.5} Si _{1.5} O ₄ (OH) ₂ [88-1]	0-75	83.81 (2)	177.2 (8)	4.61 (3)
This study	AlMg _{0.5} Si _{1.5} O ₄ (OH) ₂ [88-1]	0-75	83.47 (8)	195 (1)	4 (fixed)
This study	AlMg _{0.5} Si _{1.5} O ₄ (OH) ₂ [88-2]	0–75	84.50 (2)	172.3 (8)	4.57 (3)
This study	AlMg _{0.5} Si _{1.5} O ₄ (OH) ₂ [88-2]	0-75	84.16 (8)	189 (1)	4 (fixed)
[37]	Mg _{1.11} Si _{1.89} O ₆ (H) _{2.22}	0 to 19.8	85.46 (4)	200 (7)	4 (fixed)
[40]	Mg _{1.11} Si _{1.6} O ₆ (H) _{3.4}	0 to 30	85.66 (4)	166 (3)	4.1 (3)
[24]	$MgSi_2O_6(H)_2$	0 to 40 [HOC]	86.41	147.29	5.252
[24]	$MgSi_2O_6(H)_2$	40 to 65 [HC]	84.93	185.80	4.209
[36]	Mg _{0.89} Fe _{0.14} Al _{0.25} Si _{1.56} O ₆ (H) _{2.93}	0 to 30.5	86.10 (5)	136.5 (33)	6.32 (30)
[39]	Mg _{0.99} Fe _{0.12} Al _{0.09} Si _{1.75} O ₆ (H) _{2.51}	0 to 20.6	85.32 (2)	141.5 (30)	6.2 (4)
[41]	Mg _{1.02} Si _{1.73} O ₆ (H) _{3.03}	0 to 44	85.43 (4)	130 (1)	8.0 (2)
[25]	Mg _{1.0} Si _{1.7} O ₆ (H) _{3.0}	0 to 55.8	85.1 (2)	167.9 (86)	4.3 (5)
[25]	Mg _{1.0} Si _{1.7} O ₆ (H) _{3.0}	0 to 30 [HOC]	85.1 (2)	173 (2)	4 (fixed)
[25]	Mg _{1.0} Si _{1.7} O ₆ (H) _{3.0}	40 to 55.8 [HC]	85.1 (2)	212 (15)	4 (fixed)
[43]	Mg _{1.1} Si _{1.8} O ₆ (H) _{2.5}	0 to 65	85.80 (5)	151.4 (12)	4.89 (8)
[49]	$Mg_{0.90}Al_{0.64}Si_{1.29}O_6(H)_{3.10}$	0 to 20.5	86.71 (fixed)	143 (5)	5.8 (7)



Figure 4. Eulerian strain (*f*) versus normalized pressure (F_E) of MgSi₂O₄(OH)₂ (red circles), Al₂SiO₄(OH)₂ (blue squares), and AlMg_{0.5}Si_{1.5}O₄(OH)₂ in the 88-1 structure (purple triangles) and 88-2 structure (lavender diamonds). In this plot, the intercept is K_0 , the slope reflects the deviation of the first derivative (K'_0) from a value of 4, and curvature reflects the behavior of K''_0 . This $f - F_E$ plot uses the V_0 values obtained from fitting a third-order Birch-Murnaghan equation of state, as indicated in lines 1, 7, 13, and 15 of Table 1. The blue and red dotted lines indicate discontinuities due to hydrogen bond symmetrization in the Al- and Mg-endmembers, respectively.



Figure 5. Formula unit (f.u.) volumes of the optimized structures of $MgSi_2O_4(OH)_2$ (red circles), $Al_2SiO_4(OH)_2$ phase D (blue squares), $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ in the 88-1 structure (purple triangles), and $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ in the 88-2 structure (lavender diamonds). Dotted lines indicate equation of states (EOSs) fit to pre-hydrogen bond symmetrization structures (lines 3, 9, 13, and 15 of Table 1). Dashed lines are EOSs fit to post-hydrogen bond symmetrization structures (lines 5 and 11 of Table 1). Grey symbols (open squares, crosses, and open circles) show literature values from [25,36,37], respectively.

Direct comparison to experimentally derived equation of state parameters can be difficult, as even in the case of Mg-endmember phase D a range of compositions have been reported [25,37,40,41,43]. Yet despite this compositional variability, our equation of state parameters are in good agreement with previously published values (Table 1). No experimental equations of state for Al-endmember phase D (Al₂SiO₄(OH)₂) or AlMg_{0.5}Si_{1.5}O₄(OH)₂ are available. However, our equation of state parameters indicate that Al-endmember phase D is slightly less compressible than the Mg-endmember and that the compressibility of

AlMg_{0.5}Si_{1.5}O₄(OH)₂ is approximately intermediate to the Mg- and Al-endmember compositions (Figure 5). As expected, pressure induced hydrogen bond symmetrization resulted in an increase in the zero-pressure bulk modulus (K_0) for both endmember compositions, accompanied by a modest reduction in (K'_0). Due to the intrinsic trade-off in these parameters, the increase in bulk modulus coincident with hydrogen bond symmetrization was determined using a fixed K'_0 value of 4. In the case of MgSi₂O₄(OH)₂, the increase in bulk modulus corresponding to hydrogen bond symmetrization is 22%, in good agreement with previous calculations [24,26] and experiments [25], while for Al₂SiO₄(OH)₂ the increase is just 13%.

3.3. Elastic Constants

The full elastic tensors of the MgSi₂O₄(OH)₂, AlMg_{0.5}Si_{1.5}O₄(OH)₂, and Al₂SiO₄(OH)₂ structures were calculated at 5 GPa intervals across the 0 to 75 GPa pressure range. Although phase D is trigonal, we calculated the 21 independent elastic constants needed to describe the slight triclinic distortion in our optimized structures. The major single crystal elastic constants (C_{11} , C_{22} , C_{33} , C_{44} , C_{55} , C_{66}) are plotted in Figure 6, and the full elastic tensors are included in tabulated form in Tables A8–A15.



Figure 6. The (a) C_{11} , (b) C_{22} , (c) C_{33} , (d) C_{44} , (e) C_{55} , and (f) C_{66} elastic constants of MgSi₂O₄(OH)₂ (red circles), AlMg_{0.5}Si_{1.5}O₄(OH)₂ (blue squares), AlMg_{0.5}Si_{1.5}O₄(OH)₂ in the 88-1 structure (dark purple triangles), and AlMg_{0.5}Si_{1.5}O₄(OH)₂ in the 88-2 structure (lavender diamonds).

The Mg-endmember constants from this study are in close agreement with the previously published calculated elastic constants of [27,50], which themselves have been extensively compared to experimental results. In the case of the C_{11} , C_{44} , and C_{55} constants, both the 88-1 and 88-2 structures of the intermediate composition are bounded by the constants of the endmember compositions, with the Al-endmember C_{44} and C_{55} significantly higher than those of the Mg-endmember. The C_{22} constant largely follows the same pattern, with elevated values for Al₂SiO₄(OH)₂ compared to MgSi₂O₄(OH)₂, but the the 88-1 structure of AlMg_{0.5}Si_{1.5}O₄(OH)₂ is virtually indistinguishable from the Al-endmember. At low pressures, the C_{33} of the intermediate composition is also bracketed by the endmember compositions, but coincident with the onset of hydrogen bond symmetrization the C_{33} of both endmembers undergo discontinuous behavior, increasing abruptly before again smoothly increasing with pressure. The O—H···O bonds within phase D are most closely aligned to the *c*-axis (Figure 1), therefore it is intuitive that the C_{33} elastic constant which indicates stiffness along the *c*-axis is the most dramatically impacted by hydrogen bond symmetrization. As neither structure of AlMg_{0.5}Si_{1.5}O₄(OH)₂ undergoes pressure induced hydrogen bond symmetrization, the C_{33} of AlMg_{0.5}Si_{1.5}O₄(OH)₂ lacks this discontinuity, increasing steadily but remaining lower than either endmember at pressures exceeding 40 GPa. Lastly, the C_{66} of the tie-line composition of both structures is slightly elevated compared to both endmember compositions. This is likely due to the fact that the triclinic distortion evident in the endmembers structures is absent in the AlMg_{0.5}Si_{1.5}O₄(OH)₂ supercells, therefore the C_{66} constant accommodates some of the strain otherwise accommodated by these lesser components (Tables A8–A15).

3.4. Moduli and Velocities

Bulk and shear moduli of the Mg-endmember, Al-endmember, and two structures of the tie-line composition were calculated from the single crystal elastic constants using the Voigt-Reuss-Hill averaging scheme as shown in Figure 7a and Table 2 [51]. The pressure dependence of the bulk modulus of both Al- and Mg-endmembers exhibit discontinuities, reflecting the decrease in compressibility which accompanies pressure-induced hydrogen bond symmetrization in these phases, particularly along the *c*-axis. A more subtle inflection is also visible in the shear modulus of the Mg-endmember although no analogous discontinuity occurs within the Al-endmember. Notably, the AlMg_{0.5}Si_{1.5}O₄(OH)₂ in both the 88-1 and 88-2 structures has a bulk modulus approximately intermediate to the Mg-and Al-endmembers prior to the onset of hydrogen bond symmetrization at low pressures, but above 45 GPa these tie-line compositions have a bulk modulus nearly indistinguishable from that of the Mg-endmember. Additionally, the shear modulus of the intermediate composition is nearly indistinguishable from that of the Al-endmember in the case of both structures evaluated across the entire pressure range investigated.

Table 2. Bulk (κ) and shear (μ) moduli of Mg-endmember phase D [MgSi₂O₄(OH)₂], Al-endmember [Al₂SiO₄(OH)₂], and tie-line composition in the 88-1 and 88-2 structures as a function of pressure from 0 to 75 GPa.

P (GPa)	E	Bulk Modul	us (GPa)		S	hear Modu	lus (GPa)	
	Mg-PhD	Al-PhD	88-1	88-2	Mg-PhD	Al-PhD	88-1	88-2
0	152.9	191.5	175.1	161.9	94.8	132.7	134.4	117.3
5	177.3	216.2	199.6	191.2	106.3	145.0	147.1	132.7
10	200.6	239.5	223.5	214.3	115.9	155.5	157.8	144.2
15	222.9	261.5	245.9	240.7	124.4	165.0	167.9	156.8
20	244.9	283.5	268.0	262.4	132.1	174.5	177.4	167.0
25	263.4	305.2	289.6	284.4	139.2	183.4	186.2	176.6
30	289.2	336.7	310.9	304.6	146.3	195.0	194.9	185.2
35	311.1	353.5	332.6	324.8	153.1	200.8	202.9	192.8
40	341.4	374.0	352.7	344.6	162.6	208.1	210.4	200.6
45	361.9	392.4	369.6	363.9	168.0	214.5	216.2	208.2
50	380.5	410.3	388.6	383.3	172.8	220.2	222.9	215.7
55	398.3	428.1	407.2	402.3	177.0	226.0	229.3	222.7
60	416.3	446.1	425.8	420.4	181.2	231.7	235.5	229.1
65	433.0	463.6	443.8	439.3	183.6	237.2	241.4	235.5
70	451.5	481.1	461.9	459.8	188.9	242.6	247.7	241.7
75	468.7	498.5	479.6	475.4	192.6	248.0	253.4	246.7



Figure 7. (a) The moduli of phase D including bulk modulus (solid line) and shear modulus (dotted line), and (b) sound velocities of phase D including compressional velocity (solid line), and shear velocity (dotted line) of Mg-endmember phase D (red circles), Al-endmember phase D (blue squares), tie-line composition in the 88-1 structure (purple triangles) and the 88-2 structure (lavender diamonds).

As phase D exhibits anisotropic compression, evident in the low-pressure evolution of the *c/a* ratio as well as hydrogen bond symmetrization, determining the influence of cation substitution on compressibility is a complex issue. Furthermore, Al-bearing compositions of phase D have physical properties that cannot be predicted by linear interpolation between the Mg- and Al-endmembers. While the low pressure anisotropy in phase D is likely dictated by differential strain accommodation due to varying cation occupancies in the layered structure, Al-substitution also suppresses hydrogen bond symmetrization, indirectly influencing the elastic behavior. Parsing these distinct but overlapping effects experimentally will likely be daunting, particularly when exploring even more complex compositions (e.g., Fe-substitution combined with Al-substitution). Therefore, first-principles calculations provide a theoretical framework and detailed structural information which complement and elucidate more compositionally complex experimental studies.

4. Discussion

Previous work has probed the influence of phase D on the velocity structure of hydrous subducting slabs, of which phase D may be a significant component [27,52,53]. These studies have focused on determining to what extent phase D, which exhibits relatively high degree of shear wave anisotropy (AV_S), contributes to observations of shear wave splitting ($S_H > S_V$) in stagnant slabs. Although Mainprice and coauthors [52] evaluated the influence of compositional variation on these properties and determined the influence of cation substitution was negligible, the extent of solid solution in that study was extremely limited [$Al_{0.03}Fe_{0.11}Mg_{1.0}Si_{1.5}O_4(OH)_2$]. Furthermore, the interpretation of the influence of either Al- or Fe-substitution on the properties of phase D in that study is complicated by the inclusion of both in a single, compositionally complex sample.

We evaluated the influence of Al-substitution on the seismic anisotropy of phase D using the Christoffel equation [54], reducing our triclinic elastic constants to the appropriate trigonal symmetry with a weighted mean to enable direct comparison to literature values. The maximum shear wave polarization anisotropy (AV_S) of Mg-endmember, Al-endmember, and the tie-line composition of phase D as a function of pressure are reported in Table A16. At 0 GPa, we find the maximum AV_S of MgSi₂O₄(OH)₂ is quite high $(AV_s = 21.86)$, in good agreement with previously reported values by [27] $(AV_s = 19.92)$ and [52] (AV_S = 17.69). However, we find a strong, negative pressure dependence of the shear wave polarization anisotropy in $MgSi_2O_4(OH)_2$, such that by the pressure of the lower mantle the magnitude is of the AV_S is halved (Figure 8a). Conversely, at 0 GPa the maximum shear wave polarization anisotropy of Al-endmember phase D ($AV_S = 10.65$) is significantly lower than that of the Mg-endmember (Table A16), but due to its strong positive pressure dependence is nearly double that of the Mg-endmember at lower mantle pressures (Figure 8b). Additionally, in Al-endmember phase D the maximum shear anisotropy exists not only along the *a*-axis, as in the Mg-endmember, but also along the b-axis.



Figure 8. Lambert equal-area upper-hemisphere projection of the shear wave polarization (AV_S) of phase D at 30 GPa, including (**a**) MgSi₂O₄(OH)₂, (**b**) AlMg_{0.5}Si_{1.5}O₄(OH)₂, (**c**) AlMg_{0.5}Si_{1.5}O₄(OH)₂ in the 88-1 structure, and (**d**) AlMg_{0.5}Si_{1.5}O₄(OH)₂ in the 88-2 structure. Images were generated using the MTEX Open Source Package [54].

Based on our observation that the elastic properties of $AIMg_{0.5}Si_{1.5}O_4(OH)_2$ could not be accurately determined by interpolating the properties of the Mg- and Al-endmembers compositions, we evaluated the maximum shear wave polarization anisotropy of the tieline composition of phase D as a function of pressure (Table A16). At 0 GPa, the maximum AV_S of both structures (88-1 and 88-2) of the tie-line compositions are intermediate to that of the Al- and Mg-endmembers (13.48 and 14.87, respectively) (Figure 8c,d). However, like the Mg-endmember, the AV_S of both the 88-1 and 88-2 structures exhibits a negative pressuredependence at pressures up to ~50 GPa. Ultimately, both structures of $AIMg_{0.5}Si_{1.5}O_4(OH)_2$ exhibit less shear wave polarization anisotropy than either the Al- or Mg-endmember to pressures up to 70 GPa (Figure 8c,d). As non-endmember, Al-bearing phase D has less shear wave anisotropy than either endmember, studies that estimate regional volume % of phase D based on matching shear-wave splitting observations based on the properties of MgSi₂O₄(OH)₂ or $AIMg_{0.5}Si_{1.5}O_4(OH)_2$ likely underestimated the amount of phase D needed to mimic observations.

5. Conclusions

By evaluating three compositions of phase D $[MgSi_2O_4(OH)_2, AlMg_{0.5}Si_1_5O_4(OH)_2, CH)_2$ $Al_2SiO_4(OH)_2$] using density functional theory based calculations, we were able to probe the extent to which Al-substitution influences the physical properties of this phase. Alendmember phase D is denser (2%–6%), less compressible (6%–25%), and has faster compressional (6%–12%) and shear velocities (12%–15%) relative to its Mg-endmember counterpart. In the complex mineralogy of a subducting slab, solid solutions of phase D are expected (Al-bearing, or even Fe-bearing) and these are the geophysical properties one would hope to incorporate into regional models. Unfortunately, based on our calculations, the properties of Al-bearing phase D cannot be determined via a simple volumetric mixing model. In evaluating the properties of two structures of phase D with 50% Al-substitution we see that these tie-line compositions exhibit properties radically different than what would be obtained by linearly interpolating between the endmembers. Furthermore, comparison of these two $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ structures reveals that not only which cations substitute into phase D, but where they substitute, can also dictate macroscale bahavior. Solid solution seemingly inhibits pressure-induced hydrogen bond symmetrization, which in turn significantly influences compressibility at the pressures of the uppermost lower mantle. Not only are the elastic tensors of the tie-line composition far from intermediate to the endmembers, but at sufficiently high pressures (>45 GPa) the shear wave velocities of the $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ are higher than those of either $MgSi_2O_4(OH)_2$ or $Al_2SiO_4(OH)_2$. Lastly, $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ has a lower maximum shear wave polarization anisotropy than either the Mg- or Al-endmember compositions, and studies which constrain the the quantity of phase D in the deep Earth by matching seismic structures to the properties of either endmember may be misleading.

Author Contributions: Conceptualization and validation, E.C.T., A.J.C. and J.T.; methodology, investigation, formal analysis and data curation, E.C.T. and J.T.; software and resources J.T.; writing, visualization, funding acquisition and project administration, E.C.T.; supervision, A.J.C. and J.T. All authors have read and agreed to the published version of the manuscript.

Funding: This work was supported by a National Science Foundation Postdoctoral Fellowship under grant EAR-1725673 and EAPSI Program grant SP-1612833, jointly funded through the U.S. National Science Foundation and the Japan Society for the Promotion of Science (JSPS) for E.C.T. This study was supported in part by NSF grant EAR-1651017 for A.J.C. This study was also partly supported by JSPS KAKENHI grants JP26400516, JP26287137, JP20K04043, and JP20K04126 for J.T.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: The data presented in this study are available in the supplemental documents.

Acknowledgments: The authors thank our anonymous reviewers who provided valuable feedback that helped to improve this manuscript.

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

Appendix A

Table A1. Atomic positions in $MgSi_2O_4(OH)_2$ at 0 GPa. Atomic positions (X, Y, Z) are in units of fractional unit cells.

Atom	x	Y	Z
Mg	0.028620	0.023768	0.975729
Si	0.347702	0.658947	0.508795
Si	0.663789	0.342861	0.508793
О	0.638624	0.007447	0.255687
О	0.012286	0.633783	0.255686
О	0.364646	0.359806	0.283017
0	0.389340	0.003503	0.722073
0	0.008345	0.384497	0.722074
0	0.661872	0.657030	0.713433
Н	0.528067	0.000198	0.049481
Н	0.005037	0.523219	0.049482

Table A2. Atomic positions in $Al_2SiO_4(OH)_2$ at 0 GPa. Positions (X, Y, Z) are in units of fractional supercell, which has been doubled along the *c*-axis (i.e., the Z direction).

Atom	Х	Y	Z
Al	0.026542	0.025246	0.488610
Al	0.348302	0.664052	0.252579
Si	0.664396	0.344483	0.255085
0	0.662522	0.020607	0.122531
О	-0.000753	0.646637	0.121026
0	0.352439	0.330242	0.134030
0	-0.005662	0.021132	0.372155
0	0.008345	0.353860	0.369841
0	0.694481	0.674300	0.361957
Н	0.530647	0.006447	0.018271
Н	0.000892	0.520245	0.018077
Al	0.029669	0.022128	0.988607
Si	0.348912	0.660008	0.755084
Al	0.668490	0.343930	0.752574
О	0.651055	-0.005148	0.621028
О	0.025027	0.658129	0.622522
О	0.334662	0.348048	0.634028
0	0.358290	-0.010052	0.869836
0	0.025561	0.690086	0.872153
О	0.678729	0.674300	0.861956
Н	0.524663	-0.003512	0.518074
Н	0.010857	0.526234	0.518265

Y	Ζ		X	Y	Ζ
0.0078	0.5008	Mg	0.0021	0.5114	0.5128
0.3360	0.2484	Si	0.1641	0.8453	0.2500
0.1742	0.2539	Al	0.3409	0.6822	0.2493
0.0093	0.1412	0	0.3274	0.5031	0.1190
0.3355	0.1528	0	0.0001	0.8297	0.1260
0.1782	0.1416	0	0.1851	0.6989	0.1355
0.0090	0.3675	0	0.1933	0.5014	0.3744
0.1844	0.3879	0	0.0017	0.6957	0.3652
0.3380	0.3570	0	0.3151	0.8294	0.3817
0.0138	-0.0099	Н	0.2643	0.4990	0.0127
0.2602	-0.0144	Н	0.0024	0.7499	-0.0212
0.0155	1.0008	Mg	0.0126	0.5009	1.0128
0.3299	0.7539	Al	0.1834	0.8397	0.7493
0.1741	0.7484	Si	0.3465	0.6629	0.7500
0.0183	0.6528	0	0.3309	0.4989	0.6260
0.3179	0.6412	0	0.0043	0.8262	0.6190
0.1722	0.6416	0	0.1969	0.5005	0.8652
0.0128	0.8879	0	0.0026	0.6921	0.8744
0.1761	0.8675	0	0.3306	0.8139	0.8817
0.3332	0.8570	0	0.2001	0.6839	0.6355
0.7684	0.4856	Н	0.2511	0.5012	0.4788
0.2475	0.4901	Н	0.0001	0.7631	0.5127
0.0114	0.5128	Al	0.5167	0.5078	0.5008
0.3453	0.2500	Si	0.6753	0.8360	0.2484

0.8311

0.8191

0.5195

0.6734

0.6773

0.5140

0.8344

0.7487

0.2696

0.5089

0.6754

0.8372

0.8367

0.5105

0.6856

0.5102

0.8392

0.6793

0.7614

0.5150

Table A3. Atomic positions in the 88-1 structure of AlMg_{0.5}Si_{1.5}O₄(OH)₂ at 0 GPa. Atomic positions (X, Y, Z) are in units of fractional s

0.3453

0.1822

0.0031

0.3297

0.1989

0.0014

0.1957

0.3294

-0.0010

0.2499

0.0009

0.3397

0.1629

-0.0011

0.3262

0.1839

0.0005

0.1921

0.3139

1.0012

0.2631

0.2493

0.1190

0.1260

0.1355

0.3744

0.3652

0.3817

0.0127

-0.0212

1.0128

0.7493

0.7500

0.6260

0.6190

0.6355

0.8652

0.8744

0.8817

0.4788

0.5127

Si

0

0

0

0

0

Ο

Η

Η

Al

Si

Si

Ο

Ο

0

0

Ο

Ο

Η

Η

Х 0.0167

0.1753

0.3311

0.3191

0.0195

0.1734

0.1773

0.0140

0.3344

0.2487

-0.2304

0.0089

0.1754

0.3372

0.3367

0.0105

0.1793

0.1856

0.0102

0.3392

0.2614

0.0150

0.5021

0.6641

0.8409

0.8274

0.5001

0.6851

0.6933

0.5017

0.8151

0.7643

0.5024

0.5126

0.6834

0.8465

0.8309

0.5043

0.7001

0.6969

0.5026

0.8306

0.7511

0.5001

Al Si

Si

0

0

0

0

0

Ο

Η

Η

Al

Si

Si

Ο

Ο

Ο

0

Ο

Ο

Η

Η

Mg

Si

Al

Ο

Ο

0

0

Ο

0

Η

Η

Mg

Al

Si

Ο

0

0

Ο

0

0

Η

Η

14 of 23

0.2539

0.1412

0.1528

0.1416

0.3675

0.3879

0.3570

-0.0099

-0.0144

1.0008

0.7539

0.7484

0.6528

0.6412

0.8879

0.8675

0.8570

0.6416

0.4856

0.4901

0.6742

0.5093

0.8355

0.6782

0.5090

0.6844

0.8380

0.5138

0.7602

0.5155

0.8299

0.6741

0.5183

0.8179

0.5128

0.6761

0.8332

0.6722

0.2684

0.7475

Η

Η

0.7465

0.5094

1.0080

0.2484

0.4853

0.4797

	х	Ŷ	Z		х	Y	Z
Al	0.0124	0.0007	0.5029	Al	0.0038	0.5037	0.5214
Al	0.1733	0.3366	0.2481	Si	0.1746	0.8521	0.2563
Si	0.3218	0.1724	0.2534	Al	0.3477	0.6795	0.2543
0	0.3223	0.0076	0.1463	0	0.3251	0.5104	0.1281
0	0.0077	0.3377	0.1511	0	0.0043	0.8163	0.1347
0	0.1839	0.1779	0.1402	0	0.1857	0.6981	0.1395
0	0.1780	0.0050	0.3699	0	0.1806	0.5113	0.3899
0	0.0065	0.1744	0.3896	0	0.0115	0.6871	0.3780
0	0.3468	0.3358	0.3663	0	0.3254	0.8380	0.3888
Н	0.2521	0.0124	-0.0239	Н	0.2664	0.5043	0.0277
Н	-0.2578	0.2492	0.0367	Н	0.0035	0.7629	0.0280
Mg	0.0090	0.0247	0.9932	Mg	0.0065	0.4886	0.9983
Si	0.1802	0.3350	0.7512	Si	0.1778	0.8361	0.7506
Si	0.3457	0.1800	0.7450	Si	0.3463	0.6705	0.7534
0	0.3364	0.0058	0.6432	0	0.3293	0.5024	0.6262
0	0.0030	0.3289	0.6287	0	0.0150	0.8291	0.6282
0	0.1816	0.1713	0.6366	0	0.1977	0.5061	0.8593
0	0.1981	0.0076	0.8720	0	0.0097	0.6971	0.8647
0	0.0113	0.1951	0.8541	0	0.3178	0.8148	0.8662
0	0.3326	0.3278	0.8530	0	0.1819	0.6817	0.6305
Н	0.2603	0.7709	0.4846	Н	0.2465	0.5080	0.4853
Н	0.0093	0.2408	0.4864	Н	0.0094	0.7484	0.4797
Al	0.5038	0.0037	0.5214	Al	0.5124	0.5007	0.5029
Si	0.6746	0.3521	0.2563	Al	0.6733	0.8366	0.2481
Al	0.8477	0.1795	0.2543	Si	0.8218	0.6724	0.2534
0	0.8251	0.0104	0.1281	0	0.8223	0.5076	0.1463
0	0.5043	0.3163	0.1347	0	0.5077	0.8377	0.1511
0	0.6857	0.1981	0.1395	0	0.6839	0.6779	0.1402
0	0.6806	0.0113	0.3899	0	0.6780	0.5050	0.3699
0	0.5115	0.1871	0.3780	0	0.5065	0.6744	0.3896
0	0.8254	0.3380	0.3888	0	0.8468	0.8358	0.3663
Н	0.7664	0.0043	0.0277	Н	0.7521	0.5124	-0.0239
Н	0.5035	0.2629	0.0280	Н	0.2422	0.7492	0.0367
Mg	0.5065	-0.0114	0.9983	Mg	0.5090	0.5247	0.9932
Si	0.6778	0.3361	0.7506	Si	0.6802	0.8350	0.7512
Si	0.8463	0.1705	0.7534	Si	0.8457	0.6800	0.7450
0	0.8293	0.0024	0.6262	0	0.8364	0.5058	0.6432
0	0.5150	0.3291	0.6282	0	0.5030	0.8289	0.6287
0	0.6819	0.1817	0.6305	0	0.6981	0.5076	0.8720
0	0.6977	0.0061	0.8593	0	0.5113	0.6951	0.8541
0	0.5097	0.1971	0.8647	0	0.8326	0.8278	0.8530
0	0.8178	0.3148	0.8662	0	0.6816	0.6713	0.6366

Η

Η

0.5093

0.7603

0.7408

0.2709

0.4864

0.4846

Table A4. Atomic positions in the 88-2 structure of $AlMg_{0.5}Si_{1.5}O_4(OH)_2$ at 0 GPa. Atomic positions (X, Y, Z) are in units of fractional supercell, which has been doubled along the *a*-, *b*-, and *c*-axis.

P (GPa)	Mg-	phD	88	3-1	88	3-2	Al-j	phD
	с	а	с	а	с	а	с	a
0	4.368	4.787	4.262	4.776	4.293	4.784	4.196	4.785
5	4.303	4.750	4.217	4.738	4.242	4.744	4.155	4.749
10	4.249	4.717	4.177	4.703	4.200	4.709	4.119	4.715
15	4.203	4.686	4.143	4.672	4.163	4.677	4.086	4.686
20	4.162	4.658	4.112	4.644	4.130	4.648	4.057	4.658
25	4.127	4.633	4.085	4.617	4.103	4.620	4.033	4.633
30	4.094	4.609	4.060	4.593	4.077	4.596	4.003	4.608
35	4.065	4.586	4.038	4.570	4.053	4.572	3.986	4.585
40	4.038	4.565	4.017	4.548	4.031	4.551	3.967	4.564
45	4.017	4.544	3.998	4.528	4.010	4.530	3.950	4.543
50	3.998	4.524	3.980	4.508	3.991	4.510	3.934	4.524
55	3.981	4.505	3.963	4.490	3.973	4.491	3.919	4.506
60	3.964	4.487	3.946	4.472	3.956	4.473	3.905	4.488
65	3.948	4.470	3.931	4.455	3.940	4.456	3.891	4.471
70	3.933	4.454	3.916	4.439	3.924	4.440	3.878	4.455
75	3.918	4.438	3.903	4.423	3.910	4.424	3.865	4.439

Table A5. Average lattice parameters *c* and *a* of Mg-endmember phase D (Mg-phD), Al-endmember phase D (Al-phD), and tie-line composition in the 88-1 and 88-2 structures as a function of pressure from 0 to 75 GPa.

Table A6. Difference between the hydroxyl bond length (r_{O-H}) and the hydrogen bond length ($r_{O...H}$) in Mg-endmember phase D (Mg-phD), Al-endmember phase D (Al-phD), and tie-line composition in the 88-1 and 88-2 structures as a function of pressure from 0 to 75 GPa.

P (GPa)	Mg-phD	Al-j	ohD		88-1			88-2			
0	0.542	0.406	0.409	0.424	0.407	0.426	0.338	0.501	0.598	0.593	0.579
5	0.475	0.357	0.365	0.395	0.368	0.373	0.292	0.445	0.533	0.527	0.538
10	0.416	0.310	0.323	0.369	0.334	0.324	0.251	0.400	0.476	0.475	0.507
15	0.363	0.263	0.283	0.348	0.305	0.280	0.219	0.364	0.422	0.431	0.482
20	0.313	0.215	0.241	0.330	0.278	0.237	0.193	0.334	0.371	0.396	0.461
25	0.263	0.184	0.211	0.314	0.254	0.198	0.174	0.315	0.333	0.370	0.453
30	0.210	0.124	0.162	0.300	0.231	0.163	0.159	0.292	0.300	0.342	0.437
35	0.151	0.038	0.095	0.287	0.211	0.134	0.147	0.274	0.271	0.319	0.423
40	0.063	0.000	0.001	0.276	0.193	0.111	0.136	0.256	0.240	0.296	0.410
45	0.002	0.000	0.000	0.266	0.177	0.094	0.127	0.242	0.213	0.278	0.399
50	0.001	0.000	0.000	0.257	0.164	0.082	0.119	0.228	0.185	0.260	0.388
55	0.000	0.000	0.000	0.249	0.152	0.073	0.111	0.217	0.161	0.245	0.379
60	0.000	0.000	0.000	0.242	0.141	0.066	0.104	0.207	0.138	0.232	0.370
65	0.000	0.000	0.000	0.235	0.132	0.061	0.098	0.198	0.120	0.220	0.363
70	0.000	0.000	0.000	0.228	0.124	0.056	0.093	0.190	0.104	0.209	0.355
75	0.000	0.000	0.000	0.223	0.117	0.053	0.088	0.183	0.093	0.200	0.349

P (GPa)	Mg-phD	Al-j	ohD	88-1			88-2				
0	175.0	175.7	177.8	179.3	173.5	174.8	174.1	175.6	175.0	174.6	172.3
5	176.0	176.5	178.2	179.4	174.1	175.5	174.5	176.0	175.8	174.9	172.8
10	176.8	177.2	178.5	179.4	174.6	176.1	174.7	176.3	176.3	175.0	173.2
15	177.5	177.7	178.8	179.3	174.9	176.6	174.9	176.6	176.7	175.0	173.5
20	178.0	178.3	179.0	179.2	175.2	177.0	175.0	176.7	176.9	174.2	173.7
25	178.5	178.9	179.7	179.1	175.4	177.3	175.0	177.0	174.8	174.2	174.7
30	178.9	179.2	179.7	179.1	175.6	177.6	175.0	177.1	174.9	174.2	174.7
35	179.3	179.8	179.8	179.0	175.8	177.7	175.0	177.2	175.0	174.2	174.6
40	179.7	180.0	180.0	178.9	175.9	177.8	175.0	177.2	175.1	174.2	174.6
45	180.0	180.0	180.0	178.9	176.0	177.9	175.0	177.2	175.2	174.1	174.5
50	180.0	180.0	180.0	178.8	176.0	177.9	175.0	177.2	175.2	174.0	174.5
55	180.0	180.0	180.0	178.8	176.1	177.9	174.9	177.2	175.2	173.9	174.4
60	180.0	180.0	180.0	178.8	176.1	177.8	174.9	177.2	175.3	173.8	174.3
65	180.0	180.0	180.0	178.8	176.1	177.8	174.8	177.1	175.2	173.7	174.2
70	180.0	180.0	180.0	178.7	176.2	177.8	174.8	177.1	175.2	173.6	174.1
75	180.0	180.0	180.0	178.7	176.2	177.7	174.8	177.1	175.2	173.4	174.0

Table A7. Bond angle of O—H···O as a function of pressure in Mg-endmember phase D (Mg-phD), Al-endmember phase D (Al-phD), and tie-line composition in the 88-1 and 88-2 structures as a function of pressure from 0 to 75 GPa.

Table A8. Elastic constants of Mg-endmember phase D as a function of pressure, in units of GPa, Part 1 of 2.

P (GPa)	<i>C</i> ₁₁	<i>C</i> ₂₂	C ₃₃	C44	C ₅₅	C ₆₆	<i>C</i> ₁₂	<i>C</i> ₁₃	<i>C</i> ₁₄	<i>C</i> ₁₅	<i>C</i> ₁₆
0.01	384	376	265	96	93	108	114	48	3	-2	71
5.01	418	411	307	111	108	122	129	64	5	-4	79
10.01	450	444	344	125	120	129	143	81	5	-7	83
15.01	481	475	381	137	132	136	158	97	6	-9	87
19.99	511	504	417	149	144	142	173	113	7	-11	91
25.00	536	531	448	161	156	145	187	127	8	-11	94
30.00	567	561	486	171	165	153	203	151	9	-14	98
35.00	594	588	525	182	176	157	218	168	9	-16	101
40.01	621	615	632	193	186	162	233	184	10	-17	104
44.98	646	639	671	201	194	166	247	201	11	-19	107
50.00	671	664	697	209	201	170	261	217	12	-21	109
54.98	695	687	720	216	208	174	275	232	12	-22	112
60.00	719	710	744	224	215	177	289	248	13	-24	114
64.97	739	732	763	230	222	179	303	263	13	-26	117
70.01	765	756	791	238	229	184	317	278	14	-27	119
74.97	787	778	813	244	235	187	331	293	14	-29	121

P (GPa)	C ₂₃	<i>C</i> ₂₄	C ₂₅	C ₂₆	C_{34}	C ₃₅	<i>C</i> ₃₆	C ₄₅	C ₄₆	C ₅₆
0.01	50	10	11	-71	27	21	5	-25	21	15
5.01	67	8	14	-77	27	22	6	-30	25	16
10.01	84	7	16	-81	27	22	8	-33	27	17
15.01	101	6	17	-85	28	22	9	-37	29	17
19.99	119	6	19	-90	28	23	10	-40	31	18
25.00	132	6	21	-93	28	23	10	-43	32	19
30.00	155	4	23	-97	29	23	10	-46	34	19
35.00	173	4	24	-100	29	23	11	-49	36	19
40.01	192	3	26	-104	30	25	14	-51	38	20
44.98	209	3	27	-106	31	25	14	-54	40	21
50.00	224	2	29	-109	31	25	14	-56	42	22
54.98	240	1	31	-112	31	25	15	-58	43	23
60.00	255	0	32	-114	31	25	15	-60	45	23
64.97	270	0	35	-115	33	27	15	-61	48	25
70.01	286	-1	35	-119	31	26	15	-63	48	25
74.97	301	$^{-2}$	37	-121	32	26	16	-65	50	26

Table A9. Elastic constants of Mg-endmember phase D as a function of pressure, in units of GPa, Part 2 of 2.

Table A10. Elastic constants of Al-endmember phase D as a function of pressure, in units of GPa, Part 1 of 2.

P (GPa)	<i>C</i> ₁₁	<i>C</i> ₂₂	C ₃₃	C44	C ₅₅	C ₆₆	<i>C</i> ₁₂	<i>C</i> ₁₃	<i>C</i> ₁₄	<i>C</i> ₁₅	<i>C</i> ₁₆
0.01	410	411	379	185	169	110	118	83	1	30	75
5.00	448	449	414	206	189	119	130	102	1	26	82
10.01	484	484	447	225	205	128	143	121	0	25	88
15.01	515	516	479	242	221	135	156	139	0	25	93
20.00	546	546	513	259	235	144	170	157	0	26	98
24.99	574	575	551	275	250	149	182	173	0	26	102
29.99	606	606	638	292	266	157	197	200	0	28	107
35.00	633	632	651	304	276	162	210	214	0	27	111
40.00	660	659	686	316	287	167	223	232	0	28	114
45.03	686	685	710	328	298	172	236	248	0	28	118
49.98	712	710	733	339	307	176	249	263	$^{-1}$	28	121
54.96	737	734	755	350	317	180	262	278	$^{-1}$	29	124
60.01	761	759	778	360	327	184	275	294	-1	29	126
64.99	786	783	801	371	336	189	288	309	$^{-2}$	29	129
69.98	809	806	823	381	345	193	301	324	-2	29	132
74.99	833	829	845	391	354	197	314	339	-2	29	134

Table A11. Elastic constants of Al-endmember phase D as a function of pressure, in units of GPa, Part 2 of 2.

P (GPa)	C ₂₃	<i>C</i> ₂₄	C ₂₅	C ₂₆	<i>C</i> ₃₄	C ₃₅	C ₃₆	C ₄₅	C ₄₆	C ₅₆
0.01	86	32	-4	-76	47	39	5	-48	5	10
5.00	106	30	$^{-2}$	-82	46	38	6	-54	7	12
10.01	125	30	$^{-2}$	-88	45	38	8	-59	9	12
15.01	143	30	$^{-2}$	-93	46	38	9	-63	9	13
20.00	162	30	$^{-2}$	-98	46	38	9	-68	9	13
24.99	180	30	-3	-102	44	37	11	-72	9	13
29.99	206	32	-3	-107	49	41	11	-76	9	13
35.00	221	32	-3	-110	48	40	11	-79	9	13
40.00	238	33	-3	-114	50	41	12	-83	9	13
45.03	254	33	-4	-117	50	42	12	-86	9	13
49.98	270	33	-4	-120	50	42	13	-88	9	13
54.96	285	33	-4	-123	51	42	13	-91	9	13
60.01	301	33	-4	-126	51	42	13	-94	10	14
64.99	316	33	-5	-129	51	43	14	-97	10	14
69.98	332	33	-5	-132	51	43	14	-99	10	14
74.99	347	34	-5	-134	52	43	14	-102	10	14

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P (GPa)	<i>C</i> ₁₁	C ₂₂	C ₃₃	C44	C ₅₅	C ₆₆	<i>C</i> ₁₂	<i>C</i> ₁₃	<i>C</i> ₁₄	<i>C</i> ₁₅	<i>C</i> ₁₆
0.01	409	409	335	121	120	125	94	62	2	-1	-1
4.98	445	445	376	136	136	136	107	81	4	-2	0
10.03	479	479	415	150	150	143	121	101	5	-4	$^{-1}$
14.97	511	511	451	163	163	152	134	119	6	-4	0
20.00	542	541	487	175	174	160	148	138	8	-5	0
25.02	571	571	522	186	186	167	161	156	8	-6	0
30.00	600	600	563	196	196	173	174	172	10	-7	0
34.98	627	627	602	206	206	179	188	191	11	-8	0
40.00	654	654	634	215	215	185	201	207	12	$^{-9}$	0
44.98	680	680	645	224	224	191	214	223	12	-10	0
50.00	707	706	672	233	233	196	228	240	13	-10	0
54.97	732	732	698	241	241	201	241	256	14	-11	0
60.01	757	757	724	249	249	206	254	272	15	-12	0
64.97	781	781	749	257	257	211	267	287	16	-13	0
70.00	806	806	773	265	265	218	281	303	17	-13	0
74.97	829	829	797	273	273	222	294	319	17	-14	0

Table A12. Elastic constants of phase D with 50% Al-substitution, in which the aluminum atoms were distributed randomly across eight unit cells, also known as configuration 88-1. Elastic constants are presented as as a function of pressure, in units of GPa. Part 1 of 2.

Table A13. Elastic constants of phase D with 50% Al-substitution, in which the aluminum atoms were distributed randomly across eight unit cells, also known as configuration 88-1. Elastic constants are presented as as a function of pressure, in units of GPa. Part 2 of 2.

P (GPa)	C ₂₃	C ₂₄	C ₂₅	C ₂₆	C ₃₄	C ₃₅	<i>C</i> ₃₆	C_{45}	C46	C ₅₆
0.01	62	-2	2	-1	1	3	1	0	5	4
4.98	81	-2	4	0	4	4	0	0	2	6
10.03	101	-4	5	0	4	4	0	0	8	8
14.97	119	-4	6	0	5	5	0	0	9	9
20.00	138	-5	7	0	6	6	0	0	10	10
25.02	156	-6	9	0	5	6	$^{-1}$	0	11	11
30.00	172	-7	10	0	6	6	0	0	12	12
34.98	191	-8	11	0	6	6	0	0	13	13
40.00	207	-9	12	0	6	6	0	0	13	13
44.98	223	-10	12	0	7	7	0	0	14	14
50.00	240	-10	13	0	7	7	0	0	15	15
54.97	256	-11	14	0	8	8	0	0	14	16
60.01	272	-12	15	0	8	8	0	0	16	16
64.97	287	-13	16	0	8	8	0	0	17	17
70.00	303	-13	17	0	8	8	0	0	17	17
74.97	319	-14	17	0	9	9	0	0	18	18

P (GPa)	<i>C</i> ₁₁	C ₂₂	C ₃₃	C ₄₄	C ₅₅	C ₆₆	<i>C</i> ₁₂	<i>C</i> ₁₃	<i>C</i> ₁₄	<i>C</i> ₁₅	C ₁₆
0.01	358	394	286	107	103	113	107	50	4	-6	0
5.02	409	433	336	122	123	126	117	79	5	-8	0
10.02	449	462	373	133	139	137	131	96	5	-11	0
15.04	485	499	424	148	153	146	143	119	7	-12	0
20.02	517	531	458	161	165	156	156	137	8	-11	0
25.03	548	559	495	176	177	162	168	157	10	-12	0
30.00	578	587	526	189	186	169	180	174	11	-12	0
34.98	607	618	556	196	196	175	193	191	12	-12	0
40.00	635	646	586	206	206	182	206	208	13	-12	0
44.98	662	672	615	216	215	188	218	224	13	-13	0
50.01	689	699	645	227	224	193	231	241	14	-13	0
54.98	713	725	681	234	232	199	244	255	16	-14	0
60.01	736	750	709	243	241	203	256	270	16	-14	0
64.98	764	775	735	251	249	208	270	287	17	-15	0
70.01	788	800	771	259	257	212	282	305	18	-15	0
74.97	812	823	782	267	264	217	295	319	18	-16	0

Table A14. Elastic constants of phase D with 50% Al-substitution, in which the eight unit cells were individually end member composition, also known as configuration 88-2. Elastic constants are presented as as a function of pressure, in units of GPa. Part 1 of 2.

Table A15. Elastic constants of phase D with 50% Al-substitution, in which the eight unit cells were individually end member composition, also known as configuration 88-2. Elastic constants are presented as as a function of pressure, in units of GPa. Part 2 of 2.

P (GPa)	C ₂₃	C ₂₄	C_{25}	C ₂₆	C ₃₄	C ₃₅	C ₃₆	C_{45}	C ₄₆	C ₅₆
0.01	67	4	7	-1	16	1	5	0	6	9
5.02	85	1	8	-1	13	6	3	0	8	11
10.02	103	2	9	0	12	2	3	1	9	12
15.04	122	$^{-1}$	11	0	13	3	3	0	11	13
20.02	139	$^{-2}$	12	0	13	5	3	0	12	12
25.03	156	-6	12	0	7	7	2	0	14	14
30.00	172	-6	13	$^{-1}$	8	7	2	$^{-1}$	15	15
34.98	189	-6	13	0	10	7	2	0	15	15
40.00	205	-6	14	0	10	7	2	0	16	16
44.98	221	-9	15	0	8	7	2	0	17	17
50.00	238	-9	15	0	9	7	3	0	17	17
54.97	253	-8	16	0	11	7	3	0	18	18
60.01	269	-9	16	0	11	7	3	0	19	18
64.97	284	-10	17	0	12	7	3	0	19	19
70.00	303	-10	18	0	10	7	3	0	20	20
74.97	317	-11	18	0	12	7	3	0	21	21

P (GPa)	Mg-PhD	Al-PhD	88-1	88-2
0	21.86	10.65	13.48	14.87
5	15.79	10.16	11.43	13.27
10	12.96	9.70	9.72	11.30
15	11.62	11.49	8.74	10.4
20	10.32	13.02	8.01	9.43
25	9.37	14.59	7.26	8.51
30	8.88	16.23	6.89	7.74
35	6.56	17.08	6.59	7.77
40	9.95	17.74	6.38	7.44
45	9.08	18.04	6.27	7.15
50	9.36	18.36	6.25	6.71
55	8.03	18.84	6.31	7.04
60	8.04	19.14	6.83	6.98
65	7.91	19.36	7.32	7.37
70	6.92	19.84	7.68	7.57
75	6.24	20.12	8.11	8.5

Fable A16. Maximum shear wave polarization anisotropy (AV_S) of Mg-endmember phase D
$MgSi_2O_4(OH)_2$], Al-endmember phase D [Al ₂ SiO ₄ (OH) ₂], and phase D with 50% Al-substitution
$AIMg_{0.5}Si_{1.5}O_4(OH)_2].$

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