Synthesis and equation of state of post-perovskites in the (Mg,Fe)₃Al₂Si₃O₁₂ system

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Abstract

The formation and properties of the post-perovskite (CaIrO3-type) phase were studied in Fe-rich compositions along the pyrope–almandine ((Mg,Fe)₃Al₂Si₃O₁₂) join. Natural and synthetic garnet starting materials with almandine fractions from 38 to 90 mol% were studied using synchrotron X-ray diffraction in the laser-heated diamond anvil cell. Single-phase post-perovskite could be successfully synthesized from garnet compositions at pressures above 148 GPa and temperatures higher than 1600 K. In some cases, evidence for a minor amount of Al₂O₃ post-perovskite was observed for Alm38 and Alm54 compositions in the perovskite + post-perovskite two-phase region. Pressure–volume data for the post-perovskite phases collected during decompression show that incorporation of Fe leads to a systematic increase of unit cell volume broadly similar to the variation observed in the (Mg,Fe)SiO₃ system. The presence of Al₂O₃ increases the stability of perovskite relative to post-perovskite, requiring higher pressures (> 148 GPa) for synthesis of pure post-perovskites. Our data together with those of Tateno et al. (2005) also suggest that in the Al-rich system the presence of Fe has no strong effect on the pressure required to synthesize the pure post-perovskite phase, but the two-phase perovskite and post-perovskite region may be broad and its width dependent on Fe content. Our results suggest that any regions highly enriched in Al₂O₃ may consist of either the perovskite phase or a mixture of perovskite and post-perovskite phases throughout the entire thickness of the D″ region. The observed synthesis pressures (> 148 GPa) for a pure post-perovskite phase are beyond that at the Earth’s core–mantle boundary (~135 GPa).

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1. Introduction

The Earth’s core–mantle boundary (the D” layer) is expected to be a complex region of structural heterogeneity and thermal variability (Hirose, 2006; Shim, 2008). The discovery and investigation of the transformation of the MgSiO₃ perovskite phase (Pv) to a post-perovskite phase (CaIrO3-type) (pPv) has dramatically altered our understanding of the region (Itatia et al., 2004; Murakami et al., 2004; Oganov and Ono, 2004; Shim et al., 2004; Wentzcovitch et al., 2006; Wookey et al., 2005). Perovskites and post-perovskites in the lower mantle are expected to be chemically complex, and the behavior of post-perovskites across a range of mantle-relevant compositions needs to be investigated in detail to properly interpret seismic data for the deep Earth.

Aluminum is an important chemical constituent of the lower mantle that may be enriched locally in particular regions such as those containing subducted basaltic remnants. In experiments using pyrope (Mg₃Al₂Si₃O₁₂) starting material, it was found that a post-perovskite phase (i.e., (Mg₀.₇₅Al₀.₂₅)(Al₀.₂₅Si₀.₇₅)O₃) was stable to 140 GPa, and a single-phase post-perovskite was formed above 170 GPa (Tateno et al., 2005). When compared with MgSiO₃, this indicates that the presence of aluminum expands the stability field of perovskite relative to post-perovskite. Other experimental studies of Al-bearing MgSiO₃ compositions (with and without Fe) also conclude that the presence of Al³⁺ will broaden and deepen the perovskite to post-perovskite boundary (Andraut et al., 2010; Catalli et al., 2009; Kubo et al., 2007; Nishio-Hamane et al., 2007). Thus, the abundance of aluminum may have important consequences for the D” region and interpretation of its seismic structure (Andraut et al., 2010; Catalli et al., 2009).

There have been a number of theoretical studies of post-perovskites in the MgSiO₃–Al₂O₃ system. Using density functional theory calculations, Tsuchiya and Tsuchiya (2008) examined the MgSiO₃–Al₂O₃ system and found that the addition of moderate amounts of Al₂O₃ slightly reduced the Pv–pPv transition pressure and produced a relatively narrow coexistence region for perovskite and post-perovskite. At higher Al₂O₃ contents, an Rh₂O₃-II-type perovskite phase was stabilized at lower pressures, and an Al₂O₃-rich post-perovskite was formed at higher pressures. Other theoretical studies have found a slight increase in the Pv–pPv transition pressure due to addition of Al⁺⁺ to MgSiO₃ (Caracas and Cohen, 2005a,b; Ono and Oganov, 2005; Zhang and Oganov, 2007). These results stand in...
contrast to the experimental studies discussed above which suggest a much stronger effect of aluminum on the phase boundary. However, one earlier theoretical study found that incorporation of $\text{Al}^{3+}$ into MgSi$_2$O$_4$ PV would lead to a large increase in transition pressure and coexistence of PV and pPV over a wide pressure range (Aberer-Knutson et al., 2005).

There has also been considerable investigation into the PV–pPV transition in Fe-containing systems. Features of the deep lower mantle such as large, low-shear velocity provinces, ultra-low velocity zones, subducted slab materials, dense primordial remnants, and core–mantle reaction products (e.g. Garnero and McNamara, 2008; Mao et al., 2005; Nakagawa et al., 2010) could potentially exhibit varying degrees of iron enrichment. In addition to applications to Earth’s mantle, the amount of Fe that post-perovskite can accommodate is also relevant to understanding the limits of core sizes in possible super-Earth extrasolar planets (Valencia et al., 2009). Some experimental studies in the (Mg,Fe)SiO$_3$ system suggest that Fe prefers post-perovskite over perovskite thereby reducing the transition pressure and producing a wide PV–pPV phase loop (Caracas and Cohen, 2005a,b; Mao et al., 2004; Stackhouse et al., 2006) whereas others report a much narrower loop more consistent with a sharp seismic discontinuity (Murakami et al., 2005; Tateno et al., 2007). Variations in Fe oxidation state and also potentially its spin state as well as coupled behavior between Fe and Al add complexity to understanding this chemical component (Catalli et al., 2010a,b; Jackson et al., 2009; Lin et al., 2008; Mao et al., 2010; Sinmyo et al., 2006).

The pyrope–almandine system (Mg,Fe)$_3$Al$_2$Si$_3$O$_{12}$ is Al$_2$O$_3$-rich (25 mol%) and exhibits complete Mg–Fe solid solution. Natural and synthetic samples with a wide range of Mg# (=Mg/(Mg + Fe)) are available (Hofmeister, 2006; Hofmeister et al., 1998). Here we report synthetic samples with a wide range of Mg# (=Mg/(Mg + Fe)) are available (Hofmeister, 2006; Hofmeister et al., 1998), Alm90 glass was synthesized in the laboratory of H. O’Neill (Australian National University).

3. Results and discussions

Samples were first compressed at room temperature to pressures in excess of 110 GPa. At these conditions, diffraction peaks for all the garnet samples were weak and broad. For each composition, several experiments using different samples (designated as A, B, C, etc.) were carried out. The samples were heated for time periods ranging from 38 to 264 min, and diffraction patterns were collected at both high and room temperatures (Table 2). Note that the pressures listed in the Table 2 represent the maximum pressure achieved during heating in each run and therefore the thermal pressures are considered. In general, due to the high iron contents of the silicate samples, the heating was quite stable and only ~10–40% of the total laser power (100 W) was adequate to heat the sample to 1300–2600 K range. After heating, additional diffraction patterns were recorded both at the heated spot and nearby locations (±6–10 μm). Depending on heating conditions, the CCD images contained relatively smooth or more spotty and heterogeneous rings. Absorption changes accompanying the phase transition sometimes led to runaway heating which resulted in spotter diffraction patterns.

We found that single-phase post-perovskite can be successfully synthesized from all studied garnet compositions at pressures above 148 GPa and temperatures higher than 1600 K (Figs. 1 and 2). Our results demonstrate that the pPV phase can simultaneously accommodate both high aluminum and high iron contents. However, a separate Al$_2$O$_3$-rich pPV phase was also observed under certain conditions (particularly with heating at 2050 K and below) during some runs for Alm54 and Alm38.

For Alm38A, the sample was compressed to 148 GPa and heated at 1490–1830 K for 60 min. Our results showed that the post-perovskite phase emerged after about 33 min heating, and after temperature quenching the pressure was determined to be ~160 GPa (Fig. 2a). In a separate run, Alm38B was compressed to 160 GPa and heated to 1760–2050 K. The pPV phase was observed during the initial heating.

### Table 1

<table>
<thead>
<tr>
<th>Composition</th>
<th>MgO (wt.%)</th>
<th>FeO (wt.%)</th>
<th>Al$_2$O$_3$ (wt.%)</th>
<th>SiO$_2$ (wt.%)</th>
<th>CaO (wt.%)</th>
<th>MnO (wt.%)</th>
<th>TiO$_2$ (wt.%)</th>
<th>Cr$_2$O$_3$ (wt.%)</th>
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<tbody>
<tr>
<td>Alm38</td>
<td>15.88</td>
<td>8.46</td>
<td>22.23</td>
<td>39.93</td>
<td>1.13</td>
<td>0.90</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>Alm54</td>
<td>11.61</td>
<td>26.14</td>
<td>22.75</td>
<td>39.37</td>
<td>0.96</td>
<td>0.28</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>Alm38B</td>
<td>5.28</td>
<td>32.84</td>
<td>20.47</td>
<td>38.86</td>
<td>1.65</td>
<td>0.61</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>Alm90</td>
<td>2.38</td>
<td>37.39</td>
<td>24.72</td>
<td>35.50</td>
<td>1.65</td>
<td>0.61</td>
<td>0.02</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Alm = almandine; Pyr = pyrope; Grs = grossular; Sps = spessartine. Compositions are given both as endmember proportions in the pyrope–almandine–grossular–spessartine quaternary as well as a recalculations based on the ABO$_3$ stoichiometry of the CaO$_2$O$_3$ type. Rels.: Alm38 (#36, Table 1, Hofmeister et al., 1998), Alm54 (#40, Table 1, Hofmeister et al., 1998), Alm73 (Al-3, Table 1, Hofmeister, 2006). The Alm90 glass was synthesized in the laboratory of H. O’Neill (Australian National University).
cycle, especially close to the heating spot. With further heating for ~10 min, peaks assignable to the perovskite phase began emerging, together with new peaks that we assigned to an Al₂O₃-rich pPv phase and some other unidentified peaks. The Al₂O₃ pPv phase could be identified mainly due to the growth of a shoulder on the 022 peak at 2.382 Å (Fig. 2b). Other weak Al₂O₃ pPv peak positions were also observed in agreement with previously reported values (Ono et al., 2006). Moreover, long heating durations at temperatures below 108 GPa with further heating. In another run, Alm54D (in a Ne medium) was decompressed to 117 GPa and heated at 1290–1780 K. After about 70 min of stable heating, only weak peaks of the pPv phase were observed. Further heating at 2110 K for 19 more minutes promoted the development of the pPv phase, as evidenced by the growth of the 131 peak and followed by the development of the 132, 113 and 042 triplet. To synthesize a better quality pPv phase for equation of state study, we then increased pressure to 154 GPa and continuously heated the sample at 1670–1880 K. Our results demonstrated that along 37 min of heating strong pPv peaks were observed and no pV peaks were present (Fig. 2a). However, scans of the quenched sample showed some unidentified peaks especially in areas where the CCD images show high degrees of preferred orientation. These regions may have been subjected to locally higher temperatures. Note that Alm54B was heated again at 1450–2110 K for another 9 min, the Al₂O₃ pPv phase was observed at initial heating but subsequently it disappeared during longer heating. In another run, Alm54C (loaded in a Ne medium) was decompressed to 113 GPa and heated to 2300 K for 30 min. During heating, weak peaks of pPv appeared, but disappeared on quench (Fig. 2c). Pressure was then increased to 117 GPa and heating to 1350–1800 K for only 8 min produced a mixture of pV and pPv that remained after quench (Fig. 2c). This indicates the lower pressure boundary of the pV–pPv transition for the Alm54 composition is close to ~117 GPa, depending on synthesis temperature. In an additional run (Alm54D), garnet was compressed to 129 GPa and heated for 30 min at 1900–2200 K and a pV + pPv mixture was confirmed.

Sample Alm73C was loaded in an argon pressure medium and compressed to ~153 GPa and heated at 2030–2340 K for about 60 min. Only weak peaks of the pPv phase were identified at 157 GPa and 2130 K. Next, the sample was compressed to 155 GPa and heated at 2160–2350 K for 120 min. Again, only single pPv phase was observed but the pattern was much stronger (Fig. 2a). Further heating at 2160–2350 K for 60 additional minutes yielded no evidence for minor peaks or the growth of any other phase. This sample was then decompressed gradually and heated at 1750–2470 K for 10–29 min at each decompression step. During this process, the diffraction peaks of the pPv phase remained and no other peaks were observed down to pressures as low as 108 GPa, most likely due to the relatively short heating durations and the slow kinetics of the pPv→pV transition. The pV phase emerged below 108 GPa with further heating. In another run, Alm73D (in a NaCl medium) was compressed to ~147 GPa and heated at 2280–2400 K for 60 min. During this run, heating was stable but only weak peaks of the pV phase were observed despite long heating. The weak pV phase pattern observed at 153 GPa and 2110 K suggested that the condition may be just above the pV phase transition boundary.

For Alm90A, sample was compressed to about 149 GPa and heated at 1600–2000 K for 67 min. However, no pPv phase was observed. After further heating for 53 min, a weak pPv phase was observed in quenched patterns with pressure of 159 GPa and no other phase was found in this study. To improve the quality of the diffraction pattern of the pV phase, we continued to heat the sample for several cycles at temperature of 1830–2420 K with pressures at 172–180 GPa. The lengthy heating resulted in strong growth of the pV phase and is evident by the quenched patterns shown in Fig. 1. Next, the sample was gradually decompressed without further heating for collecting the pV phase compression data. A comparison of calculated and measured d-spacings for the pV phase of Alm90 at 160 GPa

Table 2

Summary of experimental observations.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Calibrant</th>
<th>Max pressure (GPa)</th>
<th>Temperature (K)</th>
<th>Heating time (min.)</th>
<th>Phases observed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alm38A</td>
<td>Pt</td>
<td>160</td>
<td>1480–1830</td>
<td>60</td>
<td>pPv</td>
</tr>
<tr>
<td>Alm38B</td>
<td>Pt</td>
<td>154</td>
<td>1700–2050</td>
<td>30</td>
<td>pPv, pV, pV(tr), Al₂O₃ pPv(tr)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>153</td>
<td>1630–2030</td>
<td>13</td>
<td>pV; pPv(w); pPv(w)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>148</td>
<td>1600–2560</td>
<td>264</td>
<td>pV, pPv(w); pPv(w)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>143</td>
<td>1520–1670</td>
<td>64</td>
<td>pV, pPv(tr)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>164</td>
<td>1280–1780</td>
<td>70</td>
<td>pPv(w)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>166</td>
<td>1660–2110</td>
<td>19</td>
<td>pPv; pPv(w)</td>
</tr>
<tr>
<td>Alm38C</td>
<td>Au</td>
<td>175</td>
<td>1670–1880</td>
<td>37</td>
<td>pPv; pPv; pPv(w)</td>
</tr>
<tr>
<td>Alm54C</td>
<td>Au</td>
<td>113</td>
<td>1700–2300</td>
<td>30</td>
<td>pV</td>
</tr>
<tr>
<td>Alm54C</td>
<td>Au</td>
<td>117</td>
<td>1550–1800</td>
<td>8</td>
<td>pV + pPv</td>
</tr>
<tr>
<td>Alm54D</td>
<td>Au</td>
<td>129</td>
<td>1900–2200</td>
<td>30</td>
<td>pPv + pPv</td>
</tr>
<tr>
<td>Alm54D</td>
<td>Au</td>
<td>132</td>
<td>1800–2400</td>
<td>43</td>
<td>pV + pPv</td>
</tr>
<tr>
<td>Alm54D</td>
<td>Au</td>
<td>150</td>
<td>1900–2600</td>
<td>50</td>
<td>pPv + pPv(tr)</td>
</tr>
<tr>
<td>Alm73C</td>
<td>Au</td>
<td>157</td>
<td>2030–2340</td>
<td>60</td>
<td>pPv(w)</td>
</tr>
<tr>
<td>Alm73C</td>
<td>Au</td>
<td>176</td>
<td>2160–2350</td>
<td>180</td>
<td>pPv</td>
</tr>
<tr>
<td>Alm73D</td>
<td>Au</td>
<td>153</td>
<td>2280–2400</td>
<td>60</td>
<td>pPv(w)</td>
</tr>
<tr>
<td>Alm90A</td>
<td>Au</td>
<td>160</td>
<td>1600–2000</td>
<td>67</td>
<td>None</td>
</tr>
<tr>
<td>Alm90A</td>
<td>Au</td>
<td>180</td>
<td>1830–2420</td>
<td>116</td>
<td>pPv</td>
</tr>
</tbody>
</table>

w = weak diffraction peaks; tr = trace amount (i.e. very weak diffraction observed, usually only 1 peak); † indicates growth of a phase during heating; ↓ diminishing of a phase during heating.

Fig. 1. Representative X-ray diffraction patterns of the post-perovskite (pPv) phase synthesized from Alm90 composition. Diffraction patterns are recorded at 300 K from samples quenched at the temperature indicated in the figure. Pressure is measured at 300 K. Post-perovskite phase: pV, Re: rhenium, Au: gold, and NC: NaCl.


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The synthesis cycle of the Alm38 run shown in (a). (c) Representative X-ray diffraction patterns of the mixture of Pv+pPv phases synthesized from the Alm54 composition. The synthesis temperature conditions for each sample are indicated in the figure. Post-perovskite phase: pPv, Al₂O₃ post-perovskite phase: A, perovskite phase: pv, Re: rhenium, Pt: platinum, Au: gold, Ar: argon, NC: NaCl, Ne: neon.

Table 3

<table>
<thead>
<tr>
<th>hkl</th>
<th>d_{calc}</th>
<th>d_{act}</th>
<th>(d_{act} - d_{calc})</th>
</tr>
</thead>
<tbody>
<tr>
<td>022</td>
<td>2.4306</td>
<td>2.4305</td>
<td>0.0001</td>
</tr>
<tr>
<td>110</td>
<td>2.3484</td>
<td>2.3493</td>
<td>−0.0009</td>
</tr>
<tr>
<td>023</td>
<td>1.8150</td>
<td>1.8147</td>
<td>0.0003</td>
</tr>
<tr>
<td>131</td>
<td>1.7370</td>
<td>1.7379</td>
<td>−0.0009</td>
</tr>
<tr>
<td>042</td>
<td>1.6820</td>
<td>1.6814</td>
<td>0.0006</td>
</tr>
<tr>
<td>132</td>
<td>1.5583</td>
<td>1.5590</td>
<td>−0.0007</td>
</tr>
<tr>
<td>113</td>
<td>1.5379</td>
<td>1.5362</td>
<td>0.0017</td>
</tr>
<tr>
<td>004</td>
<td>1.5219</td>
<td>1.5229</td>
<td>−0.0010</td>
</tr>
<tr>
<td>133</td>
<td>1.3529</td>
<td>1.3533</td>
<td>−0.0004</td>
</tr>
<tr>
<td>062</td>
<td>1.2300</td>
<td>1.2311</td>
<td>−0.0011</td>
</tr>
<tr>
<td>a = 2.457(1)Å, b = 8.075(4)Å, and c = 6.094(2)Å.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

118 GPa and 1700 K, lower than the Alm38 composition. For Alm73, we successfully obtained single-phase pPv above 153 GPa. During the decompression processes, although heating has been applied on the Alm73 composition at 1750–2470 K for 10–29 min at each decompression step, we still observed the pPv phase down to about 120 GPa. The transformation to the Pv phase may have been kinetically hindered due to relatively short heating interval. For Alm90, as discussed above, the pure pPv phase has been synthesized at 159–180 GPa, in agreement with our other compositions. In general, our results on P–T synthesis conditions for post-perovskite phases are broadly similar to those reported by Tateno et al. (2005) on Mg₆Al₅Si₇O₁₆.

Pressures above 150 GPa are required for synthesis of single-phase post-perovskite in the almandine-pyrope system at ~1600 K. Thus, the presence of 20–25 wt.% Al₃O₅ in these samples increases the stability of perovskite relative to post-perovskite, requiring higher pressures (>150 GPa) for synthesis of pure post-perovskites. In the (Mg₆Fe₂)SiO₃ system, the effect of Fe on the Pv–pPv transition pressure has been controversial. Our results together with those of Tateno et al. (2005) suggest that in the Al-rich system, the presence of Fe has little effect on the pressure required to synthesize the pure post-perovskite phase. Nevertheless, the mixture of Pv + pPv found at 118 GPa in the Alm54 composition (Fig. 2c) suggests that the high iron content may lower the pressure of first appearance of pPv coexisting with Pv. Future studies will be needed to examine the phase boundaries in more detail.

These data indicate that the post-perovskite phase can be synthesized from pyrope–almandine starting materials and hence can accommodate at least 20–25 wt.% Al₃O₅ and 26–37 wt.% FeO at high P–T conditions. Some studies report significant Fe³⁺ in some aluminous post-perovskites (e.g. Catalli et al., 2009; Nishio-Hamane et al., 2007). Future work is required to assess the site occupancy and valence state of iron in our samples. However, we found no direct evidence for metallic iron that might be expected to accompany formation of large amounts of Fe³⁺ from our starting materials.

Lattice parameters for the post-perovskite phase in various compositions are given in Table 4. At least 6 peaks (022, 110, 131, 113 and 004) were used to obtain the unit cell parameters of the pPv phase in this study. Since the data were collected immediately after laser heating, we expect lattice strain effects to be minimal. We observe sharpening of peak widths during heating consistent with reduction of micro-scale differential stresses. Pressure-volume data for the pPv phases collected during decompression are plotted together with other selected data in Fig. 4. Incorporation of Fe leads to a systematic increase of unit cell volume in our samples. The volume increase is due to expansion of all three unit cell axes with Fe incorporation (Table 4). Previous limited work on a pyrope composition showed that the unit cell volume for (Mg₇.₅₆Al₂₅.₃₆)(Al₂₅.₃₆Si₂₅.₇₅)O₉ (pyrope) post-perovskite is similar to that of Mg₃SiO₇ post-perovskite (Fig. 4) (Guignot et al., 2007; Tateno et al., 2005). Pure Al₂O₃ post-perovskite exhibits a slightly lower volume than Mg₃SiO₇ post-perovskite (Ono et al., 2006) (Fig. 5). Thus, the unit cell volumes

is given in Table 3. Upon decompression, the pPv022 peak became almost undetectable at pressure below 120 GPa followed immediately by the weakening of pPv131 peak. Moreover, the pPv132 and 004 triplets merged into a broad peak and became almost indistinguishable.

Fig. 3 summarizes selected results on the P–T conditions for synthesis of perovskite and post-perovskite phases in the (Mg₆Fe₂)₅Al₂Si₅O₁₂ system. The Alm38 compositions showed a Pv+pPv mixed-phase region above 140 GPa and seems to be in agreement with the results of Tateno et al. (2005) for Mg₆Al₂Si₅O₁₂. However, our single-phase pPv was synthesized at lower pressure than theirs. During decompression, the pPv phase in Alm38 composition was found to revert back to the Pv phase at about 127 GPa and 1630–2030 K. For Alm54, the mixture phase of Pv + pPv was observed at
of post-perovskites in the pyrope–almandine system reflect the dominant role of Fe with Al playing a lesser competing role.

Fig. 5 compares data for Alm54 extending below 120 GPa to other studies. The P–V behavior is very similar to earlier reported data on an (Mg,Fe)SiO₃ composition (Mao et al., 2006a) which is expected to have a similar fraction of Fe in the octahedral site (A-site) (see below). In particular, the sharp increase in volume below 100 GPa is observed in both the Al-rich and Al-free samples with high Fe contents. However, this behavior was not observed for a very low-Al Mg90 sample (Shieh et al., 2006), suggesting this metastable behavior may be connected to high Fe contents. The Alm54 sample was not annealed during decompression below 160 GPa and we observed peak broadening during decompression, and eventually the diffraction peaks were no longer detectable below 41 GPa. For the Alm73 composition, the pPv phase was annealed during decompression and persisted to 108 GPa (without noticeable broadening) until it converted to the Pv phase upon heating. For the Alm90 composition, upon decompression below about 120 GPa, the pPv phase grew increasingly weak and became impossible to detect. Thus, the ability to metastably retain the post-perovskite phase to lower pressures within the perovskite stability field in our compositions is reduced compared to low Al₂O₃ compositions and may be affected by iron content (e.g. Mao et al., 2006a,b; Shieh et al., 2006).

Table 4

<table>
<thead>
<tr>
<th>Composition</th>
<th>P (GPa)</th>
<th>a (Å)</th>
<th>b (Å)</th>
<th>c (Å)</th>
<th>V (Å³)</th>
<th>b/a</th>
<th>c/a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alm90</td>
<td>160</td>
<td>2.457(1)</td>
<td>8.075(4)</td>
<td>6.094(2)</td>
<td>120.67(6)</td>
<td>3.285(3)</td>
<td>2.480(2)</td>
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<tr>
<td>Alm73</td>
<td>160</td>
<td>2.448(2)</td>
<td>8.017(12)</td>
<td>6.06(7)</td>
<td>119.05(7)</td>
<td>3.275(8)</td>
<td>2.478(5)</td>
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<tr>
<td>Alm54</td>
<td>159</td>
<td>2.448(8)</td>
<td>7.933(6)</td>
<td>6.061(2)</td>
<td>117.74(6)</td>
<td>3.240(13)</td>
<td>2.476(9)</td>
</tr>
<tr>
<td>Alm38</td>
<td>150</td>
<td>2.432(1)</td>
<td>7.914(9)</td>
<td>6.048(4)</td>
<td>116.40(10)</td>
<td>3.254(5)</td>
<td>2.487(3)</td>
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<tr>
<td>Alm0a</td>
<td>168</td>
<td>2.415(0)</td>
<td>7.817(2)</td>
<td>6.001(2)</td>
<td>113.29(7)</td>
<td>3.237(1)</td>
<td>2.455(1)</td>
</tr>
<tr>
<td>(Mg,Fe)₂Al₂O₄</td>
<td>172</td>
<td>2.430(1)</td>
<td>7.896(5)</td>
<td>6.045(2)</td>
<td>115.99(16)</td>
<td>3.263(3)</td>
<td>2.484(2)</td>
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<tr>
<td>MgSiO₃</td>
<td>132</td>
<td>2.450</td>
<td>7.9933</td>
<td>6.0837</td>
<td>119.12</td>
<td>3.263</td>
<td>2.484</td>
</tr>
</tbody>
</table>

* Mg₂O₃Al₂O₃SiO₃ Tateno et al., 2005.
* Mg₂O₃Fe₂O₃Al₂O₃Nishi-Hamane et al., 2007.
* Guignot et al., 2007.
et al. (2007) and correspond to 120, 140, and 160 GPa from reported volumes for these two samples are ~1% different. Open diamonds (labeled 120) data. Solid lines are linear is distributed evenly between the octahedral and 8-fold sites providing charge balance

Fig. 5. Measured unit cell volumes of the pPv phases extended to metastable conditions. The solid circles and line are from this study. The open triangles are from Mao et al., 2006a. open circles from Shieh et al. (2006); open squares from Guignot et al. (2007) and crosses from Ono et al. (2006). The error bars are smaller than the symbol size.

In summary, X-ray diffraction experiments on laser-heated samples at pressures up to ~180 GPa demonstrate that a CaO$_2$-type post-perovskite phase can be synthesized for compositions along the pyrope–almandine join up to at least 90 mol% almandine at above 150 GPa and 1600 K. The presence of 20–25 wt.% Al$_2$O$_3$ in these samples increases the stability of perovskite relative to post-perovskite, requiring higher pressures (>150 GPa) for synthesis of pure post-perovskites. Between ~120 and 150 GPa, mixtures of Pv and pPv are usually observed but the diffraction patterns are generally well even with prolonged heating, suggesting a broad miscibility phase region with sluggish kinetics as suggested by Tateno et al. (2005) for pyrope. Overall, the transformation kinetics for the Al$_2$O$_3$-rich samples appear to be more sluggish than for orthopyroxene starting compositions, and thus further work at establishing phase boundaries is required. Finally, we note that any regions highly enriched in Al$_2$O$_3$ may consist of perovskite or a mixture of Pv and pPv phases throughout the entire thickness of D' as suggested in other recent studies (Andrault et al., 2010; Catalli et al., 2009). The synthesis pressures (>150 GPa) for a pure pPv phase are greater than that encountered at Earth's core mantle boundary (~135 GPa).

Acknowledgments

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References


