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Composition dependence of spin transition in (Mg,Fe)SiO$_3$ bridgmanite

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Abstract

Spin transitions in (Mg,Fe)SiO$_3$ bridgmanite have important implications for the chemistry and dynamics of Earth’s lower mantle, but have been complex to characterize in experiments. We examine the spin state of Fe in highly Fe-enriched bridgmanite synthesized from enstatites with measured compositions (Mg$_{0.61}$Fe$_{0.38}$Ca$_{0.01}$)SiO$_3$ and (Mg$_{0.25}$Fe$_{0.74}$Ca$_{0.01}$)SiO$_3$. Bridgmanite was synthesized at 78-88 GPa and 1800-2400 K and X-ray emission spectra were measured on decompression to 1 bar (both compositions) and compression to 126 GPa.
((Mg_{0.61}Fe_{0.38}Ca_{0.01})SiO_3 only) without additional laser heating. Observed spectra confirm that Fe in these bridgmanites is dominantly high spin in the lower mantle. However, the total spin moment begins to decrease at ~50 GPa in the 74% FeSiO_3 composition. These results support density functional theory predictions of a lower spin transition pressure in highly Fe-enriched bridgmanite and potentially explain the high solubility of FeSiO_3 in bridgmanite at pressures corresponding to Earth’s deep lower mantle.

Keywords: bridgmanite; spin transition; iron-bearing silicates; lower mantle

Introduction

Iron is the Earth’s most abundant transition metal and its different electronic states have been of great interest for their possible effects on mantle seismic velocities, rheology, transport properties and chemical reactions (Lin et al. 2013). A pressure-induced spin-pairing transition has been identified in (Mg,Fe)O magnesiowüstite at lower mantle conditions by both experimental and theoretical methods (e.g. Badro et al. 2003; Speziale et al. 2005; Tsuchiya et al. 2006). The transition increases density, incompressibility, shear modulus, elastic anisotropy, electrical conductivity, and iron partitioning into (Mg,Fe)O and decreases thermal conductivity and bulk sound speed (Lin et al. 2013). Spin transitions in the dominant phase of the lower mantle, perovskite-structured (Mg,Fe,Al)(Fe,Al,Si)O_3, recently named bridgmanite, potentially have even greater impact on mantle properties. However, incorporation of Fe in bridgmanite is more complex and identification of spin transition(s) has been ambiguous.

In bridgmanite, ferrous iron (Fe^{2+}) substitutes for Mg in the larger, pseudo-dodecahedral A site, while ferric iron (Fe^{3+}) can enter both the Mg-site and the octahedral B site. At ambient conditions, both species and sites are in the high-spin (unpaired electrons) state. Pressure-
induced spin transitions in bridgmanite were first identified at 70 and 120 GPa by X-ray emission spectroscopy (XES) (Badro et al. 2004) and time-domain synchrotron Mössbauer spectroscopy (T-SMS) (Jackson et al. 2005). These transitions were suggested to represent changes of high-spin iron to low spin (all electrons paired) (e.g. Badro et al. 2004; Lin et al. 2010a; McCammon et al. 2010), mixed spin (for different Fe atoms, all or no electrons paired) (e.g. Badro et al. 2004; Catalli et al. 2010) or intermediate spin (some electrons paired) (Lin et al. 2008; McCammon et al. 2008). Studies have since reported a wide range of sharp or gradual transitions in bridgmanite for different compositions, experimental conditions and techniques (Table 1). Changes in the compressibility and electrical conductivity of Fe-bearing bridgmanite at high pressure have also been linked to these spin transitions (Catalli et al. 2010, 2011; Ohta et al. 2010; Mao et al. 2011; Boffa Ballaran et al. 2012).

Recent density functional theory (DFT) calculations (Umemoto et al. 2008; Bengtson et al. 2009; Hsu et al. 2010, 2011; Hsu and Yu 2012) have offered insights important to the interpretation of experiments on spin transitions in bridgmanite. Unlike experiments, DFT methods are not limited in accessible conditions and allow full control of parameters such as the valence state of iron. DFT calculations have determined that both ferrous and ferric iron in the pseudododecahedral site remain in the high-spin state throughout the pressure-temperature range of the Earth’s lower mantle (Hsu et al. 2010, 2011). However, ferric iron in the B site will undergo a high-to-low spin transition (Hsu et al. 2011). Theoretical calculations do not support the stability of an intermediate spin state (Hsu et al. 2010).

(Mg,Fe)SiO$_3$ bridgmanites were recently synthesized at deep mantle pressures with as much as 75% FeSiO$_3$, higher than previous estimates of the maximum solubility of the FeSiO$_3$ component (Tateno et al. 2007; Dorfman et al. 2013). Whether an increase in Fe-solubility in bridgmanite
near 60-70 GPa is related to spin transitions is unknown. Increasing Fe-content has been observed to increase the spin transition pressure in (Mg,Fe)O (Fei et al. 2007). While earlier DFT studies predicted a higher spin transition pressure with increasing Fe-content in (Mg,Fe)SiO$_3$ bridgmanite (Cohen et al. 1997; Stackhouse et al. 2007), more recent studies have predicted that the spin transition pressure decreases strongly with Fe-content for (Mg,Fe)SiO$_3$ bridgmanite with >25% FeSiO$_3$ (Bengtson et al. 2008; Umemoto et al. 2008). The key difference in the more recent studies is the structure of the low-spin bridgmanite, which is predicted (Bengtson et al. 2009; Hsu et al. 2010; Umemoto et al. 2010; Caracas et al. 2014) and recently observed (Caracas et al. 2014) to be slightly distorted relative to the high-spin structure. A decrease in spin transition pressure would suggest that Fe$^{2+}$ could adopt a low spin state in bridgmanite in highly Fe-rich heterogeneities in Earth’s deep lower mantle.

A key technique for investigating spin states of transition metals is X-ray emission spectroscopy (XES) (Rueff et al. 1999; Badro et al. 2003, 2004). XES detects exchange interactions between 3p and 3d orbitals (Peng et al. 1994; Mattila et al. 2007). These energies are sensitive to multiple aspects of the electronic environment of Fe, including bonding, coordination, valence and spin states. The energy multiplets cluster to form a main Kβ$_{1,3}$ peak and a lower energy Kβ’ satellite. Spin transitions are identified and quantified by analysis of energy shift of the Kβ$_{1,3}$ peak and intensity of the Kβ’ satellite relative to well-characterized standards (Vankó et al. 2006). In this study we examine the effects of composition on electronic spin of (Mg,Fe)SiO$_3$ bridgmanite by applying XES to bridgmanites with 38% and 74% FeSiO$_3$.

**Methods**
Starting materials were natural enstatites of compositions (Mg_{0.61}Fe_{0.38}Ca_{0.01})SiO_3 (Fe#38) and (Mg_{0.25}Fe_{0.74}Ca_{0.01})SiO_3 (Fe#74). Additional details on these samples are given by Dorfman et al. (2013). Mössbauer spectroscopy of starting materials on a ^{57}\text{Co} source at sector 3 of the Advanced Photon Source confirmed no measurable ferric iron. Enstatite powder and NaCl pressure-transmitting medium were dried in a furnace at 120°C overnight before loading. Samples were loaded in a sandwich configuration between NaCl platelets in diamond anvil cells with 150-µm culet (300-µm bevel). X-ray transparent beryllium gaskets were preindentated to 20-25-µm thickness with 50-µm diameter sample chambers. Pressures were determined from the edge of the diamond Raman peak recorded from the tip of the anvil at the sample position (Akahama and Kawamura 2006). This measurement has an estimated uncertainty of ~3%.

To transform starting materials to the bridgmanite phase, samples were pressurized to 78-88 GPa and heated from both sides to 1800-2400 K with a near-infrared laser. Complete transformation was confirmed by X-ray diffraction (XRD) at beamline 13-ID-D (GSECARS sector) of the Advanced Photon Source (APS) or beamline ID27 of the European Synchrotron Radiation Facility (ESRF). Observed diffraction patterns are consistent with previous measurements of bridgmanites synthesized from these compositions (Figure 1) (Dorfman et al. 2013). Minor amounts of CaCl_2-type SiO_2 were also observed but diffraction patterns show no (Mg,Fe)O or Fe metal.

XES was performed at beamline 16-ID-D (HPCAT sector) of the APS (Bradley et al. 2012) or beamline GALAXIES of Synchrotron SOLEIL (Rueff et al. 2015). At 16-ID-D, the beam size was 35x55 µm, incident energy was 11.3 keV, and typical count rate was 60 cts/s. Incident X-rays passed through one diamond anvil, and emission was measured through the beryllium gasket (Figure 2a). At GALAXIES, the beam size was 30x80 µm, incident energy was 10 keV
and typical count rates were 70-150 cts/s. Emission signal was maximized at GALAXIES by collecting spectra at an angle minimizing the path through the beryllium gasket (Figure 2b). For each sample, pressure was increased or decreased from the synthesis pressure without additional laser heating or annealing. Spectra were recorded from 1 bar-126 GPa for (Mg$_{0.61}$Fe$_{0.38}$Ca$_{0.01}$)SiO$_3$ composition and 1 bar-88 GPa for (Mg$_{0.25}$Fe$_{0.74}$Ca$_{0.01}$)SiO$_3$ composition.

After the XES compression experiment on the Fe#38 sample, energy-domain synchrotron Mössbauer spectroscopy was performed at beamline ID18 of the ESRF (Rüffer and Chumakov 1996). The Mössbauer source was a $^{57}$FeBO$_3$ single crystal oscillated in a sinusoidal profile (Potapkin et al. 2012). Source velocity was calibrated by a natural Fe foil. The linewidth of the source was calibrated by a K$_2$Mg$^{57}$Fe(CN)$_6$ single-line absorber. The X-ray beam was focused to 9x14 µm. A spectrum was obtained for the natural sample at 126 GPa in ~18 hours.

**Results**

For all Fe-rich bridgmanite samples at all conditions, XES profiles fall between the high- and low-spin references (Figure 3). The intensity of the Kβ’ satellite peak decreases with pressure. On decompression, the intensity of the Kβ’ satellite peak increases but remains weaker than the high-spin references at ambient conditions (Figure 3a,b). The difference between the ambient and ~80 GPa spectra is greater for Fe#74 composition than for the Fe#38 composition. On increasing pressure from 78-126 in the Fe#38 composition, Kβ’ intensity also decreases subtly but systematically (Figure 3c). At all pressures Kβ’ is observed to be greater than the low-spin reference. These observations are consistent with a gradual, incomplete spin transition and/or pressure-induced broadening (Lin et al. 2013).
To distinguish the effects of spin state and pressure, the total spin moment was determined from differences between spectra and standards. Reference spectra used in this study are XES for (Mg$_{0.75}$Fe$_{0.25}$)O at ambient conditions (high spin) and 90 GPa (low spin) observed at HPCAT, APS (Lin et al. 2010b). These spectra have been used as references in previous studies of Fe-bearing bridgmanite and post-perovskite (Mao et al. 2010, 2011). The width of the Kβ peak measured for bridgmanite at GALAXIES, SOLEIL is similar to the reference spectra. The broader Kβ peak measured for bridgmanite and high-spin reference at HPCAT, APS may be due to differences in beamline configuration or background from the Be gasket in the 90° geometry (Figure 2). To confirm that these references are comparable to our data, we also consider Fe-bearing silicates, almandine glass and enstatite, as secondary high-spin references. These samples were loaded in diamond anvil cells with Be gaskets and measured at ambient conditions at HPCAT and GALAXIES, respectively. The secondary high-spin references overlap with high-spin (Mg,Fe)O up to at least 7056 eV (Figure 3).

Although the secondary references suggest the beamline configurations in these experiments were comparable, we observe systematically lower Kβ’ intensity for bridgmanite at ambient conditions (Figure 3a,b). Fe in bridgmanite is expected to be high-spin at 1 bar (McCammon 2006; McCammon et al. 2008; Lin et al. 2012). In a previous XRD study of bridgmanite synthesized from these compositions, the bridgmanite structure was not observed at ambient conditions, possibly due to amorphization (Dorfman et al. 2013). However, the XES for high-spin amorphous almandine does not show any difference in the intensity of Kβ’ relative to other standards. If Fe-rich bridgmanite remains at ambient conditions, its distorted structure may also be responsible for the lower intensity of Kβ’. This may imply that structural differences between
(Mg,Fe)O or other standards limit their applicability as high-spin standards for Fe-rich bridgmanite.

The total spin moment is calculated from XES data using an integrated difference between spectra and high- and low-spin references from Lin et al. (2010b), assuming a linear combination of high spin (total spin moment=2) and low spin (total spin moment=0) (Vankó et al. 2006; Mao et al. 2014). Differences between references from Lin et al (2010b) and secondary references are used to estimate uncertainty. The integrated absolute difference (IAD) method has been used in multiple previous studies of bridgmanite at mantle pressures (e.g. Lin et al. 2008, 2010a; Grocholski et al. 2009; Catalli et al. 2010, 2011; Mao et al. 2011) but suffers from underestimation of spin state due to pressure-induced broadening (Mao et al. 2014). Broadening effects can be corrected by using the integrated relative difference (IRD) method, which offsets decrease in Kβ’ with increase in intensity in the trough between Kβ’ and the main Kβ peak (Mao et al. 2014). For (Mg,Fe)O, the IRD method yields a decrease in the spin moment at pressures that agree with observations from XRD and Mössbauer spectroscopy (Lin et al. 2005; Mao et al. 2014). In IAD analysis, XES spectra are aligned to a common center of mass, normalized by area, and absolute values of differences are summed over a range from 7030 to 7070 eV. In IRD analysis, XES spectra are aligned to the main Kβ peak position, normalized by area, and differences are summed over only the Kβ’ portion of the spectra (up to ~7056 eV). While the total spin moment obtained with the IAD method for Fe#38 bridgmanite decreases by ~1/3 from 80-120 GPa (Figure 5a), with the IRD method the total spin moment for this material is approximately constant with pressure from 1 bar to 126 GPa (Figure 5b). This suggests no spin transition occurs in this composition over this pressure range, and the small differences in XES data can be ascribed to pressure-induced broadening.
The synchrotron Mössbauer spectrum recorded for the Fe#38 sample at 126 GPa (Figure 4) can be used to identify the sites that contribute to the spin moment observed by XES. Fe#38 bridgmanite exhibits a main doublet with quadrupole splitting (QS) of 4.15 mm/s and center shift (CS) of 0.97 mm/s. This doublet matches previous observations of Fe$^{2+}$ in the bridgmanite A-site identified as high spin (Hsu et al. 2010) or intermediate spin (Lin et al. 2008; McCammon et al. 2008). Another peak is observed at -0.4 mm/s, which may either be a doublet with QS ~0 (McCammon et al. 2010) or one peak of a doublet with CS=1.21 mm/s and QS=3.15 mm/s, where the other peak overlaps with the main doublet (Lin et al. 2008). The former possibility would match observations and predictions for low-spin Fe$^{2+}$ (Hsu et al. 2010; McCammon et al. 2010), but this may be ruled out by XES indications of high spin and no transition. The latter doublet, shown in the fit in Figure 4, is within the range reported for high-spin Fe$^{2+}$ (Dyar et al. 2006) or low-spin Fe$^{3+}$ (Catalli et al. 2010), but low-spin Fe$^{3+}$ would be expected to exhibit a lower value for CS (e.g. Pasternak et al. 2002), and again only the high-spin option is consistent with XES. Two high-spin Fe$^{2+}$ sites are thus consistent with the synchrotron Mössbauer spectrum, no spin transition in Fe#38 bridgmanite from 1 bar to 126 GPa, and predictions from DFT (Hsu et al. 2010).

The lower total spin moment of Fe#74 bridgmanite observed at high pressure (Figure 5) indicates a spin transition does occur at lower pressures in more Fe-rich bridgmanite, as predicted by some DFT studies (Bengtson et al. 2008; Umemoto et al. 2008). At ambient conditions, the intensity of the Kβ’ peak and total spin moment for Fe#38 and Fe#74 bridgmanites are identical within uncertainty. At 78-88 GPa, immediately after synthesis, the total spin moment of Fe#74 bridgmanite is ~2/3 that of Fe#38 bridgmanite. As structures (Figure 1) and experimental conditions are similar for these three samples, the difference must be the onset of a spin
transition in Fe\textsuperscript{2+} in Fe\#74 bridgmanite. The decrease in spin moment becomes significant between 40 and 68 GPa in this composition.

Recent studies have attempted to identify structural differences in bridgmanite XRD due to spin transitions (Boffa Ballaran et al. 2012; Caracas et al. 2014). Relative to high-spin bridgmanite, low-spin bridgmanite was predicted and observed to exhibit additional diffraction peaks due to structural distortion and symmetry reduction (Caracas et al. 2014). However, all diffraction peaks for both Fe\#38 and Fe\#74 bridgmanite at 80-88 GPa match orthorhombic GdFeO\textsubscript{3}-type perovskite structure, with no extra low-symmetry peaks (Figure 1). Structural differences in lower-spin Fe\#74 bridgmanite at 88 GPa are either not present or too subtle to observe. Both the spin transition and the structural distortion may be incomplete at the maximum pressure studied.

**Discussion**

In contrast to these observations, previous XES studies (Table 1) measured spin transitions in less Fe-rich bridgmanite over this pressure range, indicated by sharper changes in Kβ position and Kβ’ intensity (Badro et al. 2004; Catalli et al. 2010, 2011; Lin et al. 2010a; Fujino et al. 2012) or much lower Kβ’ intensity (Li et al. 2004, 2006; Lin et al. 2008; Mao et al. 2011). These studies examined compositions ranging from 5-50% FeSiO\textsubscript{3} and estimated or measured Fe\textsuperscript{3+}/ΣFe from 0-100%. Differences in total spin moment observed in these studies are due in part to differences in abundances of Fe\textsuperscript{2+} and A- or B-site Fe\textsuperscript{3+}, which will each lose spin moment at different conditions. Of these sites, density functional theory predicts that only B-site Fe\textsuperscript{3+} undergoes a spin transition within a mantle pressure range (Hsu et al. 2010, 2011). In experiments, the simplest case is Fe\textsubscript{2}O\textsubscript{3}-bearing bridgmanite, which must contain Fe\textsuperscript{3+} in both A- and B-sites. Based on both XES and T-SMS analyses, bridgmanite with 10% Fe\textsubscript{2}O\textsubscript{3} was
observed to lose half its spin moment above ~55 GPa, in good agreement with density functional theory (Catalli et al. 2010). Spin transitions in B-site Fe$^{3+}$ have been reported at pressures as low as 13-24 GPa (Lin et al. 2012).

Pressure and temperature may increase the fraction of B-site Fe$^{3+}$ by migration of Fe$^{3+}$ from the A site to the B site (Grocholski et al. 2009; Caracas 2010; Catalli et al. 2011; Hsu and Yu 2012; Fujino et al. 2014). For FeAlO$_3$-bearing bridgmanite, whether the Fe$^{3+}$ resides in A- or B-site is generally unknown and has been suggested to change during heating/annealing (Caracas 2010; Hsu and Yu 2012). Diffusion of Fe$^{3+}$ to the B-site may account for the transition observed in bridgmanites with 10-15% FeAlO$_3$ by Catalli et al. (2011) and Fujino et al. (2012) at 50-70 GPa, but does not appear to have affected other similar experiments despite annealing (Nishio-Hamane et al. 2008; Potapkin et al. 2013).

For FeSiO$_3$-bearing bridgmanites, density functional theory predicts a pressure-induced distortion in the bonding of high spin A-site Fe$^{2+}$, resulting in a mixture of A1 and A2 sites with different nuclear energies, but no loss of spin moment (Hsu et al. 2010). Experimental observations of FeSiO$_3$-bearing bridgmanites are even more complex because most analyses find some Fe$^{3+}$, regardless of attempts to generate reducing conditions during synthesis (McCammon 1997; Frost and Langenhorst 2002; Frost et al. 2004; Grocholski et al. 2009). Thus spin transitions observed in bridgmanites synthesized from Fe$^{2+}$-bearing starting materials may still occur in Fe$^{3+}$ in the B-site. In combination with diffusion of A-site Fe$^{3+}$, oxidation of Fe$^{2+}$ may also increase the component of B-site Fe$^{3+}$ particularly during heating at deep lower mantle pressures (McCammon 1997; Frost et al. 2004; Zhang and Oganov 2006). A 10% FeSiO$_3$ composition examined with X-ray emission spectroscopy by Badro et al. (2004) with annealing showed loss of spin moment at 70 and 120 GPa, while unannealed samples showed only gradual
changes (Li et al. 2004, 2006; Jackson et al. 2005). A 25% FeSiO$_3$ composition exhibited X-ray emission spectra characteristic of a dominant low spin component from 80-135 GPa, but may have as much as 50% Fe$^{2+}$/ΣFe (Mao et al. 2011). The Fe$^{2+}$/ΣFe ratio is most commonly determined at high pressure by T-SMS. To our knowledge, the only study to have reported Fe$^{3+}$/ΣFe ratio below detection limit was the Lin et al. (2008) combined T-SMS-XES study which, with McCammon et al. (2008), introduced the possibility of intermediate spin state Fe$^{2+}$ in bridgmanite. These data have been reinterpreted to show either the two high-spin Fe$^{2+}$ sites or a mixture of high-spin Fe$^{2+}$ and low-spin Fe$^{3+}$, consistent with theory (Lin et al. 2013).

The lack of a transition in Fe#38 bridgmanite is consistent with high-spin Fe in our sample throughout the lower mantle pressure range. We expect that the dominant Fe species in the synthesized bridgmanites is Fe$^{2+}$ in the A site (e.g. Vanpeteghem et al. 2006), as supported by our Mössbauer spectroscopy results for the Fe#38 bridgmanite. Since typical measured Fe$^{3+}$/ΣFe ratios in bridgmanites synthesized from Al-free, Fe-rich compositions are ~15% (e.g. Frost and Langenhorst 2002), minor amounts of Fe$^{3+}$ may be present. As no annealing was performed on these samples after synthesis, changes in oxidation state or diffusion of Fe between sites on compression or decompression should be highly energetically unfavorable. Any Fe$^{3+}$ in the B site would be expected to undergo a spin transition, but a few percent low-spin Fe$^{3+}$ would not be detectable by either XES or Mössbauer spectroscopy in this work. In a similar experiment, Lin et al. (2010a) suggested a transition in Fe#40 bridgmanite to a low or intermediate spin state based on analysis using the IAD method. Their calculated values for the total spin moment are in excellent agreement with our values for Fe#38 bridgmanite using this method (Figure 5a). Consideration of pressure-induced broadening with the IRD method reveals no transition in Fe#38 bridgmanite to at least 126 GPa.
A transition in the Fe#74 bridgmanite may be consistent with phase equilibria observed in the (Mg,Fe)SiO$_3$ system. From 25-50 GPa, solubility of FeSiO$_3$ in bridgmanite increases from ~20 to ~40% FeSiO$_3$ (e.g. Tange et al. 2009). Above ~50 GPa, the bridgmanite to bridgmanite-plus-oxides phase boundary must change slope to reach the higher solubility of FeSiO$_3$ reported in recent experiments in the diamond anvil cell (Tateno et al. 2007; Dorfman et al. 2013): by ~70 GPa, the solubility of FeSiO$_3$ increases to at least 75% FeSiO$_3$. These pressures and compositions correspond well to the observed onset of the spin transition in Fe$^{2+}$.

Our observations match predictions from density functional theory studies that high Fe-content promotes the high-to-low spin transition in (Mg,Fe)SiO$_3$ bridgmanite at lower pressures (Bengtson et al. 2008; Umemoto et al. 2008). The absolute transition pressure is highly dependent on the choice of exchange-correlation functional, but the overall trend is not. For (Mg$_{0.25}$Fe$_{0.75}$)SiO$_3$ bridgmanite, the spin transition pressure was predicted to be 26-30 GPa (LDA method) or 80-106 GPa (GGA method). For (Mg$_{0.62}$Fe$_{0.38}$)SiO$_3$ bridgmanite, the transition pressure would be 55-75 GPA (LDA method) or 120-170 GPa (GGA method). The low transition pressures given by the LDA method are clearly inconsistent with our observations, but the GGA predictions are slightly higher than the pressure at which we observe the spin moment begins to decrease in our experiments. Our observations contradict the opposing prediction that the spin transition occurs at a higher pressure in FeSiO$_3$ (Cohen et al. 1997; Stackhouse et al. 2007).

**Implications**

A lack of a sharp spin transition in (Mg,Fe)SiO$_3$ bridgmanite is consistent with measured physical properties of this material. Equation of state measurements show no discontinuity in
volume compression throughout the lower mantle pressure range (Lundin et al. 2008; Mao et al. 2011; Dorfman et al. 2013). The majority of experimental measurements and density functional theory calculations find that Fe$^{2+}$-bearing bridgmanite, unlike low-spin Fe$^{3+}$-bearing bridgmanite or low-spin (Mg,Fe)O, is not significantly less compressible than MgSiO$_3$-bridgmanite (Dorfman and Duffy 2014). These results support no discontinuity in the seismic properties of Fe-bearing bridgmanite in the lower mantle.

Differences in spin transition pressure in bridgmanite between this study and previous reports reflect differences in data processing (IAD vs. IRD), synthesis conditions (pressure, temperature, hydrostaticity of stress state, stability of heating, frequency of annealing), and composition (bulk Fe-content, Fe$^{3+}$/ΣFe, Al-content). Chemical heterogeneity in the deep lower mantle may result in regional differences in Fe-spin due to differences in bulk Al-content and Fe$^{3+}$/ΣFe ratio. These regional differences would have important effects on Earth’s mantle via changes in chemical partitioning (Badro et al. 2004; Irifune et al. 2010; Fujino et al. 2014), increase in electrical conductivity (Ohta et al. 2010, 2012), thermal conductivity (Badro et al. 2004; Ohta et al. 2014) and density (Mao et al. 2011). The wide compositional range explored in this study provides improved constraints on the dependence of bridgmanite chemistry and properties on Fe-content which can be applied to chemical heterogeneities in the mantle.

Previous experiments have reached conflicting conclusions about the partitioning of iron between the bridgmanite and post-perovskite phases at the base of the mantle (Auzende et al. 2008; Hirose et al. 2008; Sinmyo et al. 2008; Sakai et al. 2009, 2010). If, as suggested by Hirose et al. (2008), iron favors the bridgmanite phase, then the beginning of the post-perovskite transition will correspond to an increase in iron content in bridgmanite, possibly increasing the post-perovskite transition pressure while decreasing the spin transition pressure in bridgmanite,
promoting lower-spin ferrous iron in deep mantle bridgmanite. Conversely, if iron partitions into the post-perovskite phase (Auzende et al. 2008; Sakai et al. 2009), the post-perovskite transition will occur at shallower depths. If increasing Fe-content decreases the spin transition in post-perovskite, the ferrous iron in the iron-rich post-perovskite may be expected to exhibit the low-spin state in Fe-rich heterogeneities.

An additional complication for deep lower mantle seismic structure is the possibility of multiple post-perovskite phases in Fe-bearing compositions (Yamanaka et al. 2012; Zhang et al. 2014). Zhang et al. (2014) have observed that at ~95 GPa, Fe partitions strongly out of bridgmanite into a new “H-phase,” of unknown structure and assumed (Mg,Fe)SiO₃ stoichiometry. This observation conflicts with a large body of experimental work and has not been replicated by other experimenters. However, if the H-phase is the stable Fe-bearing silicate in the deep mantle, only a small amount of (high-spin) Fe will remain in bridgmanite. Phase transition boundaries between other post-perovskites and H-phase are unknown. Multiple other post-perovskite structures have been reported, depending on composition (Tschauner et al. 2008; Yamanaka et al. 2012). Phase transitions and partitioning between these post-perovskite potentially provide a mechanism for multiple discontinuities in the D’’ region.

In summary, this work adds to the growing experimental consensus that supports predictions by density functional theory of no spin transition in Fe²⁺-bearing bridgmanite over the lower mantle pressure range in likely mantle compositions. X-ray emission spectroscopy data show no change from 1 bar to 126 GPa for (Mg₀.₆₁Fe₀.₃₈Ca₀.₀₁)SiO₃ bridgmanite. A transition in (Mg₀.₂₅Fe₀.₇₄Ca₀.₀₁)SiO₃ bridgmanites is consistent with predictions of a decrease in spin transition pressure at high Fe-content.
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References


**Figure captions**

Figure 1: X-ray diffraction patterns of synthesized Fe-bearing bridgmanites at 300 K and reference stick pattern for orthorhombic bridgmanite. Gray starred reference sticks are additional peaks predicted by Caracas et al. (2014) for low-spin bridgmanite. X-ray wavelength is $\lambda=0.3344$ for these measurements.

Figure 2: Experimental geometries for X-ray emission spectroscopy at a) HPCAT and b) GALAXIES beamlines.

Figure 3: X-ray emission spectra and difference spectra relative to low-spin reference observed on decompression at APS 16-ID-D (a,b) and compression at SOLEIL GALAXIES (c) for (Mg$_{0.25}$Fe$_{0.74}$Ca$_{0.01}$)SiO$_3$ (a) and (Mg$_{0.61}$Fe$_{0.38}$Ca$_{0.01}$)SiO$_3$ (b,c) compositions (colored lines). Reference spectra for (Mg$_{0.75}$Fe$_{0.25}$)O at ambient conditions (high spin, solid black line) and 90 GPa (low spin, dashed black line) (Lin et al. 2010b) and almandine-composition glass and enstatite measured at ambient conditions at HPCAT and GALAXIES, respectively, are also shown (high-spin, bold black line).

Figure 4: Energy-domain synchrotron Mössbauer spectrum recorded over ~18 hrs at beamline ID18 of the European Synchrotron Radiation Source for Fe38-bridgmanite at 126 GPa after XES experiment. The fit is a two-site model with a Lorentzian profile (Prescher et al. 2012). Site 1 has center shift of 0.98 mm/s, quadrupole splitting of 4.18 mm/s, and intensity of 55%, and site 2 has center shift of 1.21 mm/s, quadrupole splitting of 3.15 mm/s, and intensity of 45%. Sites are identified by analogy to experiments on $^{57}$Fe-enriched bridgmanite samples (Lin et al. 2008; Dorfman et al. 2014).
Figure 5: a) Total spin moment calculated from integrated absolute difference (Vankó et al. 2006) between measured spectra and reference high- and low-spin spectra (Lin et al. 2010b). Black triangles represent \((\text{Mg}_{0.25}\text{Fe}_{0.74}\text{Ca}_{0.01})\text{SiO}_3\)-composition bridgmanite while gray triangles represent \((\text{Mg}_{0.61}\text{Fe}_{0.38}\text{Ca}_{0.01})\text{SiO}_3\)-composition bridgmanite. Gray circles represent measurements of \((\text{Mg}_{0.60}\text{Fe}_{0.40})\text{SiO}_3\)-composition bridgmanite by Lin et al. (2010a). Direction of triangle points indicates compression (APS 16-ID-D) or decompression (SOLEIL GALAXIES). A loss of spin moment is observed above 40 GPa for 74\% FeSiO\textsubscript{3}. b) Total spin moment calculated from integrated relative difference method (Mao et al. 2014).
Table 1: Summary of previous observations of spin transitions in bridgmanite in multianvil or diamond anvil cell (DAC) experiments using X-ray emission spectroscopy (XES), conventional Mössbauer spectroscopy (MS), time-domain synchrotron Mössbauer spectroscopy (T-SMS), energy-domain synchrotron Mössbauer spectroscopy (E-SMS), X-ray diffraction (XRD) and electrical resistivity (ER).

<table>
<thead>
<tr>
<th>Spin transition</th>
<th>Compositions</th>
<th>Synthesis</th>
<th>Annealing</th>
<th>Methods</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>None/no effect</td>
<td>6-40% FeSiO$_3$ (0-80% Fe$^{3+}$)</td>
<td>25-88 GPa (multianvil or DAC)</td>
<td>Each step or none</td>
<td>E-SMS, ER, XES, XRD</td>
<td>Potapkin et al. (2013), This study</td>
</tr>
<tr>
<td>Gradual decrease</td>
<td>9-14% FeSiO$_3$, 10% FeAlO$_3$ (15-70% Fe$^{3+}$)</td>
<td>25-35 GPa (multianvil or DAC)</td>
<td>None or &lt;1000 K</td>
<td>MS, XES, T-SMS</td>
<td>Li et al. (2004, 2006), McCammon et al. (2008)</td>
</tr>
<tr>
<td>Transition below 50 GPa</td>
<td>10% FeSiO$_3$, 10% FeAlO$_3$ (25-70% Fe$^{3+}$)</td>
<td>23-26 GPa (multianvil)</td>
<td>None</td>
<td>T-SMS, XES, XRD</td>
<td>Li et al. (2006), Lin et al. (2012)</td>
</tr>
<tr>
<td>Gradual transition ends at 50-70 GPa</td>
<td>5-50% FeSiO$_3$ (40% Fe$^{3+}$ or unknown)</td>
<td>26-93 GPa GPa (multianvil or DAC)</td>
<td>None</td>
<td>T-SMS, XES, XRD</td>
<td>Jackson et al. (2005), Caracas et al. (2014)</td>
</tr>
<tr>
<td>Sharp transition at 50-85 GPa</td>
<td>10-40% FeSiO$_3$, 10% Fe$_2$O$_3$, 10-15% FeAlO$_3$ (10-100% Fe$^{3+}$ or unknown)</td>
<td>25-110 GPa (multianvil or DAC)</td>
<td>Each step or none</td>
<td>ER, XES, XRD, T-SMS</td>
<td>Badro et al. (2004), Grocholski et al. (2009), Catalli et al. (2010, 2011), Lin et al. (2010a), Ohta et al. (2010), Fujino et al. (2012)</td>
</tr>
<tr>
<td>Gradual transition begins at 50-70 GPa</td>
<td>74% FeSiO$_3$ (unknown Fe$^{3+}$)</td>
<td>88 GPa (DAC)</td>
<td>None</td>
<td>XES, XRD</td>
<td>This study</td>
</tr>
<tr>
<td>Sharp transition at 120-135 GPa</td>
<td>10-40% FeSiO$_3$ (10-20% Fe$^{3+}$ or unknown)</td>
<td>30-120 GPa (DAC)</td>
<td>Each step or none</td>
<td>XES, XRD, T-SMS</td>
<td>(Badro et al. 2004), Lin et al. (2010a), McCammon et al. (2010)</td>
</tr>
</tbody>
</table>
Figure 1

Intensity (arb. units)

Diffraction angle 2θ (degrees)

Orthorhombic bridgmanite reference

(Mg_{0.25}Fe_{0.74}Ca_{0.01})SiO_3 bridgmanite, 88 GPa

(Mg_{0.61}Fe_{0.38}Ca_{0.01})SiO_3 bridgmanite, 80 GPa

NaCl

SiO_2
Figure 2
(Mg$_{0.25}$Fe$_{0.74}$Ca$_{0.01}$)SiO$_3$ bridgmanite:
- Red: 88 GPa
- Orange: 60 GPa
- Green: 40 GPa
- Blue: 1 bar

References:
- (Mg,Fe)O, high spin (Lin et al. 2010)
- (Mg,Fe)O, low spin (Lin et al. 2010)
- Almandine glass, high spin

Difference relative to low spin reference

Normalized intensity (arb. units)

Energy (eV)
\[(\text{Mg}_{0.61}\text{Fe}_{0.38}\text{Ca}_{0.01})\text{SiO}_3\text{ bridgmanite:}\]

- **80 GPa**
- **66 GPa**
- **1 bar**

**References:**

- (Mg,Fe)O, high spin (Lin et al. 2010)
- (Mg,Fe)O, low spin (Lin et al. 2010)
- Almandine glass, high spin

Normalized intensity (arb. units)

Energy (eV)
Figure 3

Normalized intensity (arb. units)

Energy (eV)

(Mg$_{0.61}$Fe$_{0.39}$Ca$_{0.01}$)SiO$_3$ bridgmanite:
- 78 GPa
- 93 GPa
- 106 GPa
- 117 GPa
- 126 GPa

References:
- (Mg,Fe)O, high spin (Lin et al. 2010)
- (Mg,Fe)O, low spin (Lin et al. 2010)
- Enstatite, high spin
Figure 4
Figure 5