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Article Composition and Pressure Effects on Partitioning of Ferrous Iron in Iron-rich Lower Mantle Heterogeneities

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Abstract: Both seismic observations of dense low shear velocity regions and models of magma ocean 15 crystallization and mantle dynamics support enrichment of iron in Earth's lowermost mantle. Phys-16 ical properties of iron-rich lower mantle heterogeneities in the modern Earth depend on distribution 17 of iron between coexisting lower mantle phases (Mg,Fe)O magnesiowüstite, (Mg,Fe)SiO3 bridg-18 manite, and (Mg,Fe)SiO₃ post-perovskite. The partitioning of iron between these phases was inves-19 tigated in synthetic ferrous-iron-rich olivine compositions (Mg0.55Fe0.45)2SiO4 and (Mg0.28Fe0.72)2SiO4 20 at lower mantle conditions ranging from 33-128 GPa and 1900-3000 K in the laser-heated diamond 21 anvil cell. The resulting phase assemblages were characterized by a combination of in situ X-ray 22 diffraction and ex situ transmission electron microscopy. The exchange coefficient between bridg-23 manite and magnesiowüstite decreases with pressure and bulk Fe# and increases with temperature. 24 Thermodynamic modeling determines that incorporation and partitioning of iron in bridgmanite 25 are explained well by excess volume associated with Mg-Fe exchange. Partitioning results are used 26 to model compositions and densities of mantle phase assemblages as a function of pressure, FeO-27 content and SiO2-content. Unlike average mantle compositions, iron-rich compositions in the man-28 tle exhibit negative dependence of density on SiO2-content at all mantle depths, an important find-29 ing for interpretation of deep lower mantle structures. 30

Keywords: iron partitioning; lower mantle; mantle heterogeneities; laser-heated diamond anvil cell; 31 experimental petrology 32

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1. Introduction

Iron-enrichment has been suggested to explain seismic observations of dense regions 35 in the deep lower mantle such as large low shear velocity provinces (LLSVPs) [e.g. 1] and 36 ultra-low velocity zones (ULVZs) [e.g. 2,3]. For LLSVPs, the difference in Fe/(Mg+Fe), or 37 Fe#, relative to the surrounding mantle is inferred to be as much as 7% based on estimated 38 chemical density differences up to 2% [4]. ULVZs may be more dramatically enriched in 39 Fe to generate density differences of 10% or more. The origin of these regions is unknown, 40 but possibilities include segregation of dense material in a basal magma ocean beneath 41 the crystallizing early mantle [5], accumulation of dense material from subducted slabs 42 [6], or partial melt due to less refractory composition and/or high local temperatures [7]. 43 Understanding these geophysical observations requires dependence of physical proper-44 ties such as elasticity, rheology, and electrical and thermal conductivities on composition. 45

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Copyright: © 2021 by the authors. Submitted for possible open access publication under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/). The distribution of elements in minerals in the modern mantle is moderated by subsolidus 46 chemical partitioning. 47

Iron in Earth's lower mantle is distributed between $(Mg,Fe,Al)(Fe,Al,Si)O_3$ bridgmanite (Bdg), its polymorph $(Mg,Fe,Al)(Fe,Al,Si)O_3$ post-perovskite (pPv), and $(Mg,Fe)O_4$ ferropericlase — magnesiowüstite (Mws). Partitioning of Fe between these phases is characterized by the exchange coefficient, K_D , between coexisting silicate and oxide, related to their compositions, X, in equilibrium, where values less than 1 indicate that iron favors the oxide, and values greater than 1 the reverse: 53

$$K_{D} = \left(X_{\text{Fe}}^{\text{Silicate}} / X_{\text{Mg}}^{\text{Silicate}}\right) / \left(X_{\text{Fe}}^{\text{Mws}} / X_{\text{Mg}}^{\text{Mws}}\right)$$
(1)

This exchange coefficient is used to describe the amounts of Fe in silicate vs. oxide 54 for variable bulk compositions corresponding to hypothesized mantle rocks. Previous ex-55 periments have explored silicate-oxide partitioning behavior primarily using San Carlos 56 olivine and pyrolite compositions as analogues for the mantle [e.g. 8–15]. Due to chal-57 lenges entailed in modeling iron, computational studies using density functional theory 58 have only recently begun to address partitioning in the lower mantle [16,17]. Previous 59 studies have converged around K_D of ~0.2-0.3 in San Carlos olivine near 660-km depth, 60 but substantially higher in pyrolite. The higher K_D in pyrolite is due to incorporation of Al 61 and ferric iron in Bdg [18]. In ferrous-iron-bearing olivine compositions, KD decreases or 62 remains constant with pressure and increases with temperature and oxygen fugacity. 63

The dependence of the exchange coefficient on bulk iron content is key to the modern 64 composition of dense mantle heterogeneities. Studies of silicate-oxide iron partitioning in 65 iron-rich compositions with Fe#>16 have found that the exchange coefficient decreases 66 with increasing amounts of ferrous iron, but these studies have been limited to <50 GPa 67 [e.g. 10]. Effects of ferrous iron enrichment on partitioning have been examined to deep 68 lower mantle conditions only along experimental thermal gradients [13] or in relatively 69 iron-poor compositions [12,11,14]. However, partitioning in lower mantle minerals may 70 change at pressures >50 GPa and high ferrous iron content due to composition-dependent 71 structural and electronic changes in bridgmanite. 72

At the boundary between the transition zone and lower mantle, bridgmanite is stable 73 with no more than 12 mol% FeSiO₃, but in the deep lower mantle the solubility limit in-74 creases to at least 75% FeSiO₃ [e.g. 19 and references therein]. An increase in solubility 75 may be linked to structural distortion in Bdg and corresponding bonding changes for fer-76 rous iron predicted by density functional theory [20] and confirmed by X-ray diffraction 77 of single crystals [21]. The increase in solubility of iron in Bdg in the deep mantle may also 78 be due to composition-dependence of a high-to-low electronic spin transition in ferrous 79 iron [22]. Since the discovery of pressure-induced spin transitions in ferropericlase [23] 80 and bridgmanite [24], these electronic changes have been explored for their potential ef-81 fects on partitioning [e.g. 8] as well as other mantle properties. Differences in Fe-content 82 and Fe³⁺/ Σ Fe control the conditions at which spin transitions and partitioning changes 83 occur in the mantle [25,26]. In San Carlos olivine, low $Fe^{3+}/\Sigma Fe$ promotes ferrous iron in 84 Mws, which will undergo a high-to-low spin transition, and Fe²⁺ in Bdg would be pre-85 dicted to remain in the high-spin state throughout the lower mantle pressure range [20]. 86 However, in more iron-rich compositions, recent studies have suggested that the spin 87 transition in Fe²⁺ in Bdg may occur at ~70 GPa, in the middle of the lower mantle [22,27]. 88

At conditions corresponding to the base of the lower mantle, post-perovskite has also 89 been observed to incorporate at least 80% FeSiO3 [28]. Most studies found that Fe-rich pPv 90 is stable at lower pressures than Fe-poor pPv [e.g. 19,28], suggesting that iron prefers pPv 91 to Bdg. Partitioning of iron from Bdg into pPv is supported by observations of higher K_D 92 for pPv-Mws than for Bdg-Mws when olivine is used as a starting material [e.g. 8]. Con-93 versely, diffraction and microscopy studies of coexisting Bdg and pPv have observed the 94 reverse in pyrolitic starting materials: a preference of iron for Bdg over pPv [29,30]. The 95 conflict between these studies has been attributed to differences in oxidation state of iron. 96 Partitioning of iron in bridgmanite and post-perovskite at these higher pressures deter-97 mines whether Fe-enriched silicate phases can explain dense heterogeneities in a poten-98 tially SiO₂-rich deep lower mantle [e.g. 2]. 99

Constraints on chemistry at extreme conditions are best obtained by a combination 100 of in situ phase identification and ex situ composition measurements. In situ X-ray diffrac-101 tion (XRD) is required to determine the high-pressure phase assemblage, as mantle sili-102 cates including Fe-rich Bdg and pPv become amorphous upon quench to ambient condi-103 tions. Compositions of all phases can be measured to percent-level under best conditions 104 using a combination of focused ion beam (FIB) and analytical transmission electron mi-105 croscopy (TEM). The FIB allows extraction and polishing of samples synthesized in the 106 diamond anvil cell for nm-scale compositional measurements in the TEM. In this study, 107 we apply XRD, FIB and TEM techniques to the partitioning of Fe in Fe-rich mantle phase 108 assemblages synthesized in the laser-heated diamond anvil cell at deep lower mantle con-109 ditions. The results are combined with self-consistent thermodynamic modeling to predict 110 physical properties of Fe-rich heterogeneities in Earth's mantle. 111

2. Materials and Methods

Synthetic fayalite-rich olivines with compositions (Mg0.55Fe0.45)2SiO4 (Fa45) and 113 (Mg0.28Fe0.72)2SiO4 (Fa72) were used as starting materials. Olivines were synthesized from 114 stoichometric mixtures of MgO, SiO₂, FeO, and a small amount of Fe₂O₃ to compensate for 115 nonstoichiometry in FeO. Oxide powder mixtures were reacted in a reducing graphite 116 capsule in a piston-cylinder press (IPGP, Paris) at 1 GPa and 1200°C for 15 hours to pro-117 duce ~300-µm olivine crystals. Olivine compositions were determined with the JEOL 8200 118 Superprobe at the University of Lausanne (Table S1), and are consistent with all Fe²⁺ in 119 the starting materials. An additional Fa50 sample synthesized with ⁵⁷Fe under similarly 120 reducing conditions exhibited 100% Fe2+ based on Mössbauer spectroscopy (Supplemen-121 tary Figure S1). 122

Single olivine crystals were crushed into 3-10-µm-thick platelets and sandwiched be-123 tween insulating platelets of NaCl or MgO (the latter medium used for 1 sample which is 124 presented for comparison but not used in partitioning analysis). Materials were dried in 125 a 120° C oven overnight to remove moisture before loading. Sample assemblages were loaded in symmetric diamond anvil cells with 300-µm-diameter flat anvil culets or 100-150-µm-diameter beveled anvil culets. Gaskets were Re sheets indented to ~25-µm thickness. 129

Samples were compressed to 33-128 GPa in the diamond anvil cell and laser-heated 130 from both sides to 1900-3000 K. Pressures were determined at 300 K from the diamond 131 anvil Raman peak at the sample position [31] or the equation of state of the NaCl medium 132 [32] before heating. Pressures measured after heating differed from before heating by up 133 to ~2 GPa due to transformation and annealing of the sample and medium. Reported un-134 certainty for these pressure scales is ~1% under quasihydrostatic loading below 1 Mbar 135 [31], and nonhydrostatic stresses and relaxation may increase uncertainty in pressure to 136 ~3-5% [32]. Laser heating was performed for 15-60 min at beamline 13-ID-D of the 137 GSECARS sector of the Advanced Photon Source or at the Earth and Planetary Science 138 Laboratory at Ecole polytechnique fédérale de Lausanne (EPFL). The laser heating system 139 at GSECARS provides a flat-top beam profile for minimal temperature gradients at the 140 sample center [33] and was co-aligned with the ~3-4 micron focused synchrotron X-ray 141 beam (0.3344 Å) by visually observing X-ray-induced fluorescence on each sample. The 142 MAR-CCD imaging detector was used to record high-resolution X-ray diffraction in situ 143 at high-pressure and high-temperature conditions. In situ X-ray diffraction is not available 144 during experiments at EPFL. The EPFL system uses a single 1070 nm wavelength, 200 W 145 fiber laser in Gaussian mode divided for double-sided heating using a polarizing 146 beamsplitter. The two sides are independently controlled with rotating wave-plates and 147 focused with 60 mm achromatic lenses. For both GSECARS and EPFL systems the hot spot 148has a diameter of ~20 µm. Based on tests of the flat-top optics [33], the large (defocused) 149

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Gaussian beam spot will produce a similar temperature distribution in the heating spot150center as a flat-top spot. Temperature was determined throughout heating by spectrora-151diometry over 620-900 nm spectral range. For each sample, a single spot was heated (no152scanning of the laser across the sample), and quenched directly from lower mantle tem-153perature conditions by zeroing the laser power.154

Chemical analyses of the quench products of these experiments were obtained by 155 electron microscopy. After recovery to ambient conditions, each sample was prepared for 156 ex situ imaging and sectioning. The NaCl medium was dissolved from the surface of each 157 silicate sample with a droplet of deionized water. Each sample was then coated with 10-158 20 nm of carbon. Heated spots were identified by color (black and opaque, vs. transparent 159 unheated olivine) and radially-symmetric topography. Although the diameter of the en-160 tire laser-heated area is ~20 μ m, the diameter of the hottest part of the sample is ~4 μ m 161 (Supplementary Figures S2-S3). The hottest region exhibits voids at grain boundaries, due 162 possibly to water or release of strain, and larger grain sizes and higher modal abundance 163 of bridgmanite relative to cooler regions. 164

Thin sections of each heated spot were extracted and polished to electron transpar-165 ency using a Zeiss NVision 40 dual-beam scanning electron microscope and focused Ga+ 166 ion beam. To prevent damage by the ion beam, a protective layer of carbon 1-2-µm thick 167 was deposited over a ~2x20 µm area across the center of each heating spot. Additional 168 carbon patches were deposited to support particularly thin (~1-µm-thick samples recov-169 ered from ~1 Mbar pressure) or fragile samples (with cracks or holes). Slices were milled 170 to 1-2-µm thick at 30 kV and 1.5-27 nA before transfer to TEM grid. The center of each 171 slice was then polished to 100-200-nm thickness at 30 kV and 300 pA. Final cleaning and 172 polishing was performed at 5 kV and 80 pA at a 5° angle to the surface. As in previous 173 work [8], a few slices were lost during the lift-out procedure. 174

Sections were transferred to a FEI Tecnai Osiris analytical TEM for composition anal-175 ysis. Accelerating voltage for these measurements was 200 kV. To avoid selective migra-176 tion of elements from measurement spots [34], chemical measurements were obtained in 177 scanning mode from maps. EDX maps were acquired at 10,000-50,000 (typically ~40,000) 178 counts/second with pixel dwell time of 50 microseconds scanned over 1024x1024 pixels 179 for total scanning time ~10 min. Compositions of individual grains were summed over 180 selected ~100 nm regions within maps. In the hot spot centers, two samples had grain sizes 181 smaller than the 100-200-nm thickness of the polished thin section and due to grain over-182 lap were not suitable for composition analysis. Grain sizes decreased significantly at 183 higher pressures, requiring longer heating times. We evaluated the quality of chemical 184 measurements and magnitude of potential Soret diffusion [35] from the Si-content meas-185 ured in oxide grains due to overlap with silicate grains. All measurements included in this 186 study have less than 4% Si/(Mg+Fe+Si) measured in magnesiowüstite. Compositions 187 measured for samples obtained via EDX are listed in Table 1 and Supplementary Table 2. 188 Reported uncertainties are standard deviations of composition measurements for selected 189 grains used for composition measurements in each sample. Measurements of unheated 190 olivine regions of samples agree with compositions measured by microprobe to 1-2%, 191 supporting accuracy of EDX. 192

Table 1. Composition measurements by XRD and EDX and corresponding experimental conditions pressure (P), temper-
ature (T), and duration (t). K_D is defined as $(X_{Fe}^{Silicate}/X_{Mg}^{Silicate})/(X_{Fe}^{Mws}/X_{Mg}^{Mws})$, where X is concentration of the subscript193element in the superscript phase. Uncertainties are taken from statistical error in EDX quantification. *=Laser heating and
X-ray diffraction conducted at APS beamline 13-ID-D. Missing data: NR=Lattice parameters not resolvable from diffrac-
tion data, NA=No diffraction available, TS=grain sizes too small for EDX measurements. Phases: Bdg=bridgmanite,
pPv=post-perovskite, Mws=magnesiowüstite, and Sti=stishovite.193

Sample	P (GPa)	T (K)	t (min)	Phases	X ^{Bdg} _{Fe} (XRD)	X_{Fe}^{Bdg}	X_{Fe}^{Mws}	KD
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Fa45*	40	2000	10	Mws, Bdg, Sti (1 grain)	25±5	17.9±1.9	74.7±1.8	0.074±0.018
Fa45*	46	2050	20	Mws, Bdg	14±5			Recovery failed
Fa45*	62	2000	15	Mws, Bdg	27±5	16±3	73.6±1.1	0.07±0.03
Fa45*	66	2250	15	Mws, Bdg	21±5			Recovery failed
Fa45*	71	2440	20	Mws, Bdg	18±5		TS	
Fa45*	89	2600	15	Mws, Bdg	4±5	18±4	64±2	0.12±0.05
Fa72*	33	1980	15	Mws, Bdg, Sti	56±5	37±3	86.8±0.3	0.089±0.016
Fa72	39	2000	30	Mws, Bdg, Sti	69±5	47±7	92.4±1.2	0.07±0.04
Fa72*	48	1900	15	Mws, Bdg	NR	43±5	92.2±0.6	0.06±0.02
Fa72	52	2000	30	Mws, Bdg	NA	46.5±1.3	95.3±0.8	0.043±0.011
Fa72*	61	1850	20	Mws, Bdg	55±5		TS	
Fa72	82	2000	20	Mws, Bdg, Fe	NA	51±2	98.2±0.5	0.019±0.009
Fa72*	89	2200	35	Mws, Bdg, pPv	31±5	49±2	96.8±1.1	0.032±0.014
Fa72	128	2700	60	Mw, pPv	NA	pPv: 59.1±0.8	97±3	0.05±0.07
Fa72 in MgO	83	2200	45	Fp, Mws, Bdg	NA	4.2±0.7	21.9±0.9, 96.4±0.5	

3. Results

3.1. Phase Identification

Phases and compositions were identified ex situ for all experiments as well as in situ for experiments conducted at GSECARS with X-ray diffraction. Diffraction patterns rec-202 orded at heating spot centers after temperature quench exhibited peaks corresponding to 203 bridgmanite, magnesiowüstite, stishovite, and a single peak attributed to CaIrO3-type 204 post-perovskite (Figure 1, Supplementary Figure S4). In the Fa45 composition from 40-89 205 GPa (all pressures studied with X-ray diffraction), all observed diffraction peaks match 206 Bdg or Mws (Supplementary Figure S4). In Fa72 at 33 GPa, corresponding to the shallow 207 lower mantle, stishovite is also observed (Figure 1). The exsolution of stishovite indicates 208 that bridgmanite cannot accommodate all available iron and breaks down to a mixture of 209 bridgmanite, stishovite and magnesiowüstite. At higher pressures, the solubility of Fe in 210 Bdg increases [19,29,36], and the stishovite phase disappears. Identification of stishovite 211 and magnesiowüstite by EDX in recovered samples concurs with observations by X-ray 212 diffraction. Pure SiO2 regions were identified in composition maps of samples synthesized 213 at and below 40 GPa, but not in samples synthesized at higher pressures (Figure 2, Table 214 1). 215

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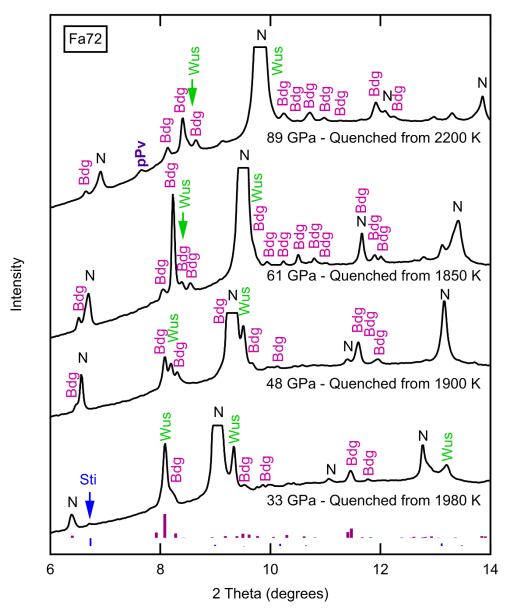


Figure 1. Diffraction patterns for phase assemblages synthesized from Fa72 at 33-89 GPa. Bdg=bridgmanite, pPv=post-perovskite, Wus=wüstite/magnesiowüstite, Sti=stishovite, N=NaCl medium. Due to overlap between oxide and NaCl peaks, we cannot resolve whether the structure 219 is B1 cubic or rhombohedral, though EDX results indicate 92-98% FeO. Reference stick patterns for 220 Bdg and Sti are provided below. For these experiments, λ =0.3344 Å. 221

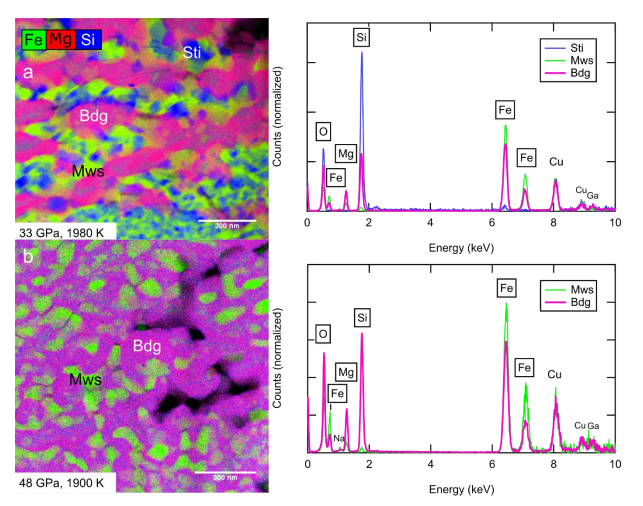


Figure 2. Energy-dispersive X-ray maps with Mg, Fe, and Si concentrations in red, green, and blue respectively and corresponding spectra obtained from selected grains for a) Fa72 heated to 1980 K at 33 GPa and b) Fa72 from 48 GPa. Bdg=bridgmanite, Mws=magnesiowüstite, Sti=stishovite. Cu signal is from the TEM grid, and minor Ga remains from ion milling.

Although Fe metal droplets have been observed in multiple previous studies of lower 227 mantle phase assemblages [e.g. 8,37], they were not typically present in samples analyzed 228 in this work. Metallic iron is detectable in recovered samples as oxygen-depleted regions 229 in EDX maps. In one sample Fe metal droplets ~10-50 nm were embedded in a large grain 230 of nearly pure FeO (Supplementary Figure S5). Unlike these droplets, Fe-rich inclusions 231 in Bdg grains were not typically depleted in oxygen (Supplementary Figure S6). In con-232 trast to previous experiments on less Fe-rich compositions, we do not observe Fe metal 233 grains within Bdg due to disproportionation. This lack of metallic Fe is evidence that iron 234 remains ferrous after synthesis of the lower mantle phase assemblage. 235

Silicates bridgmanite and post-perovskite in recovered samples could not be distin-236 guished due to amorphization. With one exception, Bdg was amorphous (e.g. Supplemen-237 tary Figure S7); Bdg with high Fe-content (Fe#≥38) is not quenchable [19]. The amorphous 238 silicate is the interconnected phase, as Bdg has been in previous studies [8]. Iron-depleted 239 Fe#4 Bdg in the sample loaded in MgO (final sample listed in Table 1) was successfully 240 quenched, and crystalline Bdg was observed before composition mapping. Bridgmanite 241 lattice planes subsequently disappeared after composition mapping due to damage by the 242 electron beam. Crystalline (Mg, Fe)SiO₃ pPv is also not quenchable to ambient conditions, 243 and we were thus unable to conclusively identify a pPv phase in recovered samples. A 244 single amorphous silicate phase quenched from synthesis at 128 GPa is assumed to have 245 been post-perovskite. 246

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3.2. Compositions of coexisting phases and partitioning

The compositions determined from EDX mapping for bridgmanite and magnesio-248 wüstite are listed in Table 1 and Supplementary Table 2. Listed uncertainties in measured 249 compositions and partitioning coefficients were obtained from a standard deviation of 250 compositions of multiple grains at the sample center, and may be an underestimate due 251 to sample topography and matrix effects. The total Fe# of the phase assemblage in the 252 center of the heating spot is ±3 Fe# of the starting materials (note that for phase assem-253 blages that include substantial stishovite, the bulk Fe# is not the average Fe# of Bdg and 254 Mws, but an average over the entire measured region), confirming that the sample center 255 did not lose or gain a significant amount Fe by Soret diffusion [35]. The outer portion of 256 the laser heating spot, where thermal gradient was significant, does exhibit compositional 257 differences due to Soret diffusion, and data from these regions were not used in further 258 analysis. Due to reaction between the sample and the MgO medium, the sample with the 259 MgO medium is not included in further analysis of compositions and partitioning. Bridg-260 manite compositions are Fe#16-18 for Fa45 and Fe#37-51 for Fa72 (Figure 3a). Measured 261 compositions of Bdg formed from Fa72 are slightly more Fe-rich than the maximum solu-262 bility of FeSiO₃ in Bdg measured at ~2000 K in multianvil press experiments to 50 GPa [36] 263 (Figure 3b). The consistently higher maximum Fe-content in Bdg in these experiments rel-264 ative to Tange et al. [36] may be due to underestimation of temperature during laser heat-265 ing, overestimation of Fe-content by EDX, or other experimental differences in run time 266 and/or redox state. Magnesiowüstites formed from Fe-rich olivines are extremely Fe-rich: 267 Fe#64-75 in Fa45 samples and Fe#87-98 in Fa72 (Figure 3b). Compositions measured by 268 EDX for Bdg and Mws determine values for K_D that are lower than previous studies of 269 less Fe-rich San Carlos olivine, decrease with pressure, and increase with temperature 270 (Figure 4). This is consistent with previous observations at lower pressures of stronger 271 preference of ferrous iron for Mws in Fe-enriched bulk compositions [e.g. 10,36]. The 272 exchange coefficient decreases with pressure for the Fa72 composition. While partitioning 273 appears flat with increasing pressure in the Fa45 composition, this is likely an effect of 274 higher temperatures during the higher pressure runs for this composition, and we may 275 expect the exchange coefficient to decrease with pressure at a constant temperature. Post-276 perovskite at 128 GPa was measured to have Fe#59, higher than any Fe# measured for 277 bridgmanite. This results in a higher post-perovskite – magnesiowüstite exchange coeffi-278 cient relative to bridgmanite-magnesiowüstite (Figure 4), in agreement with previous 279 studies of San Carlos olivine by Auzende et al. [8] and Sakai et al. [11]. 280

Lattice spacings of bridgmanite, magnesiowüstite and post-perovskite, when resolv-281 able, also provide estimates of the partitioning of iron. This analysis has previously been 282 attempted for high-pressure diffraction data [38], but more recent studies offer improved 283 constraints on the equation of state of lower mantle phases, including effects of iron on 284 the compressibility of bridgmanite and effects of the spin transition on ferropericlase. Lat-285 tice parameters of Bdg in olivine compositions in this work at pressures determined by 286 the NaCl medium are consistent with Fe#0-20 in Fa45 and Fe#40-70 in Fa72 (for more de-287 tails see Appendix A, Supplementary Figure S8, Table 1). These values are consistent with 288 EDX observations (Supplementary Figure S8) to ~20 Fe#. At all pressures, observed unit 289 cell volumes for magnesiowüstite support Fe-enrichment relative to the starting compo-290 sition (Supplementary Figure S9), also consistent with EDX observations (Figure 3b). 291

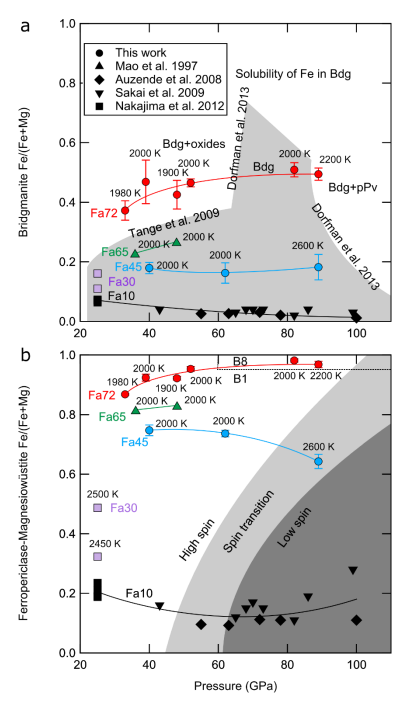


Figure 3. Composition data obtained from partitioning studies in olivine compositions plotted with relevant phase stability data. Data from this study and those from Auzende et al. [8] and Nakajima et al. [10] were obtained by EDX in the TEM. Data from Mao et al. [39] are based on X-ray diffraction at ambient conditions. a) Compositions for bridgmanite synthesized from olivine starting materials. Gray shaded region marks solubility of Fe in bridgmanite at ~2000 K from previous studies in the multianvil press [36] and laser-heated diamond anvil cell [19]. b) Compositions of (Mg,Fe)O synthesized from olivine starting materials. Shaded regions show change in the pressure of the high-to-low spin transition with composition [40,41]. The rectangle indicates the pressure/composition range of previous observations of B8 FeO [42,43]. Although two of our data indicate Fe/(Mg+Fe) for (Mg,Fe)O and synthesis conditions that could correspond to B8 structure, our XRD observations could not confirm a structural difference during or after heating.

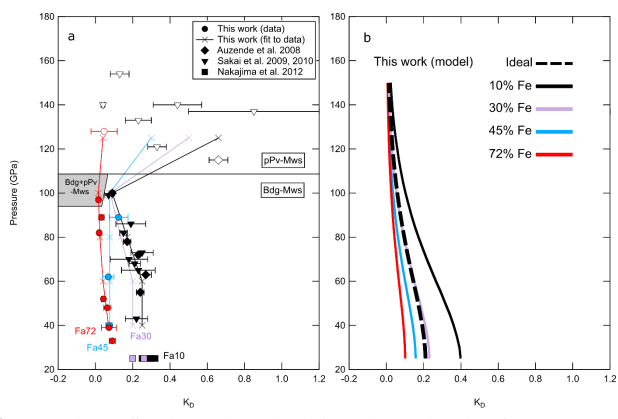


Figure 4. a) Exchange coefficient between silicate and oxide phases at lower mantle conditions from experiments in this and previous studies [8,10–12] compared to thermodynamic models. Filled/open symbols indicate a bridgmanite/postperovskite silicate phase. For most data, temperatures were in the range 1850-2200 K (temperatures for all data in this study listed in Table 1). Light gray region indicates composition/pressure space with coexisting bridgmanite and postperovskite. Values chosen for fit to compositions Fa10, Fa30, Fa45 and Fa72 at selected pressures 40, 60, 80, 100 and 125 GPa are shown in 'x' symbols and lines. b) Modeled pressure- and composition-dependence of partitioning behavior between bridgmanite and ferropericlase in olivine bulk compositions assuming ideal mixing (black dashed curve) and nonideal mixing (solid curves) using updated thermodynamic parameters based on thermal equation of state of FeO by [42] and inversion of experimental partitioning data in this study.

At 89 GPa in Fa72, lattice parameters of bridgmanite indicate only Fe# ~30, likely due 314 to partitioning of Fe into incipient pPv. The most prominent diffraction line of pPv, (022) is observed in this diffraction pattern. The lattice spacing indicated by this peak is large 316 relative to that observed for Fe#40 pPv at similar pressures, supporting very high Fe-con-317 tent [44]. The initiation of the post-perovskite transition at only 89 GPa is compatible with 318 the shallow, broad post-perovskite transition in Fe-rich (Mg, Fe)SiO3 observed in previous 319 studies [2,19,28]. 320

3.3. Thermodynamic modeling

Variation of partitioning behavior as a function of bulk iron content in the olivine 322 system implies non-ideal mixing behavior. To determine the thermodynamic properties 323 consistent with observed partitioning behavior, we carried out thermodynamic modeling 324 of partitioning in olivine bulk compositions at lower mantle conditions. Iron partitioning between Mws and Bdg when SiO₂ stishovite is not present refers to the following chemical 326 reaction, 327

$$MgSiO_3 + FeO \leftrightarrow FeSiO_3 + MgO$$

that can be rewritten in terms of chemical potential,

(3)

(2)

 $\mu_{MgSiO3} + \mu_{FeO} \leftrightarrow \mu_{FeSiO3} + \mu_{MgO}$

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where μ_i is the Gibbs free energy of species *i*. The chemical potential will vary as a function 329 of temperature: 330

$$\mu_i = \mu_i^0 + RT \log(x_i \gamma_i)$$

where x_i and γ_i are the mole fraction and coefficient of activity of species *i*, respectively. 331 We assume that mixing is ideal in the Bdg phase (i.e. $\gamma_{MgSi0_3} = \gamma_{FeSi0_3}$). However, non-ideal 332 mixing has been documented in the Mws phase [45], and can be described by a Margules 333 interaction parameter *W*. 334

$$RT\log(\gamma_{Mg0}) = W(1 - x_{Mg0})$$

$$RT\log(\gamma_{\rm Fe0}) = W(1 - x_{\rm Fe0})$$

Using equations 1-5, we obtain

$$RT \log(K_D) = \mu_{MgSi03}^0 - \mu_{FeSi03}^0 + \mu_{Mg0}^0 - \mu_{Fe0}^0 - W(2x_{Fe0} - 1)$$

In equation 6, x_{FeO} is constrained by the bulk Fe/(Fe+Mg) ratio of the system and the relative amount of Mws relative to Bdg, 337

$$x_{\rm Fe0}(1-Y) + x_{\rm FeSi03}Y = Z$$

where *Y* is the relative proportion of Bdg (i.e. Bdg/(Bdg+Mws) and *Z* is the bulk 338 Fe/(Fe+Mg) ratio of the system. It should be noted that in the ideal mixing case (i.e. *W*=0), 339 K_D is insensitive to the bulk iron content of the system that controls x_{Fe0} ; as the experimental observations described above demonstrate a dependence of K_D with bulk iron 341 content, non-ideality in either the Mws or Bdg phase is required to describe iron partition-342 ing in iron-rich olivine systems. 339

Thermodynamic parameters for Mws and Bdg phases have been constrained by previous studies, for example summarized in a database by [46]. The equation of state of ironbearing Bdg and free energies consistent with observed phase equilibria in the MgSiO₃-SeSiO₃ sytem were updated by [47]. However, an initial attempt to model iron partitioning between Mws and Bdg according to the above framework yielded K_D inconsistent with seperimental observations (Supplementary Figure S10). While experimental data indicate K_D decreasing with pressure, the modeled K_D increases with pressure for all compositions.

To correct this discrepancy, we first updated the equation of state parameters for FeO 351 using the values from [42]. Next, we adjusted the reference chemical potential (F_0 in the 352 [46] thermodynamic database) to fit the mean value of K_D at 40 GPa. These changes pro-353 duce a reasonable pressure-dependence for KD (Figure 4b) self-consistently using equation 354 6, whether mixing in the Mws system is assumed to be ideal or non-ideal. The interaction 355 parameter W is then inverted from the experimental data presented in this study (Figure 356 4) using the forward model for K_D in equation 6. The resulting thermodynamic parameters 357 for FeO are shown in Table 2. This model successfully reproduces the partitioning obser-358 vations in ferrous-iron-bearing systems in this and previous studies [8,10,12], as well as 359 previous observations of the stability of iron-rich bridgmanite [19] (Figure 5). 360

Table 2. Thermodynamic parameters for FeO employed in partitioning and phase equilibrium361modeling: 1 bar volume V_0 , 1 bar bulk modulus K_0 , pressure-derivative of the bulk modulus K_0' ,362Grüneisen parameter γ_0 , second-order Grüneisen parameter q, Debye temperature θ_0 , reference363chemical potential F_0 , and Margules parameter for ferropericlase W. Values for these parameters364from [46] were updated per measurements by [42] and further modified to produce partitioning365behavior in agreement with experimental data.366

	Previous parameters	Revised parameters
V_0 (cm ³ /mol)	12.26ª	12.26
K ₀ (GPa)	179 ª	149 ^b

(4)

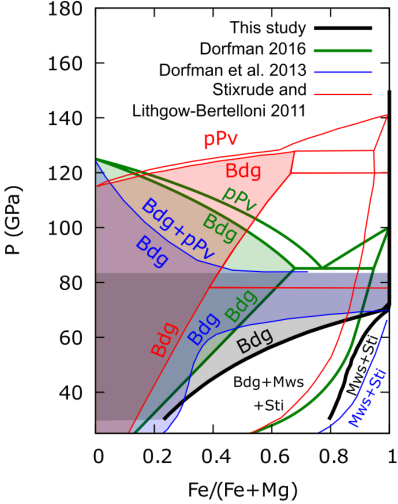
(5)

(6)

(7)

Ko'	4.9 ^a	3.6 ^b
γ_0	1.53ª	1.53
9	1.7ª	1.7
θ ₀ (K)	454ª	454
F ₀ (x10 ³ J/mol)	-242ª	-235°
W (x10 ³ J/mol)	13 ^a	14.746°

^aStixrude and Lithgow-Bertelloni (2011). ^bFischer et al. (2011). ^cThis study.



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Figure 5. Phase diagram of MgSiO₃-FeSiO₃ system at ~2000 K based on experimental observations369(blue: [19]) and thermodynamic calculations in previous work (red: [46], green: [47]) and this370study (bold black lines). Thermodynamic calculations using excess mixing volume term in this371study successfully reproduce experimental observations of very high solubility of FeSiO₃ in bridg-372manite at deep lower mantle pressures >80 GPa. Regions in pressure-composition space where373single phase bridgmanite are stable are highlighted for each phase diagram.374

4. Discussion

4.1. Multivariable effects on partitioning

The experiments presented here examine substantially more Fe²⁺-rich compositions 377 than those explored in previous studies of partitioning in Earth's deep lower mantle, and 378 thus provide systematic constraints on effects of compositional differences on partitioning 379 in iron-rich heterogeneities. Previous studies have observed two major trends of partitioning with increasing pressure/depth for different compositions [18]: in San Carlos olivine, 381 with no Fe³⁺, silicate-oxide K_D is flat or decreases between 50-100 GPa and increases across 382 the post-perovskite transition [8,11–13,16], while pyrolite, with high Fe³⁺/ Σ Fe, K_D exhibits 383 a more complex behavior in the Bdg stability field, and decreases across the pPv transition 384 [9,15,48]. Below 50 GPa in other olivines, presumably with low Fe³⁺/ Σ Fe, K_D decreases with 385 total Fe# [10,36,49], and this work demonstrates that the trend for iron-rich compositions 386 at higher pressures continues to be shifted to lower K_D values relative to San Carlos olivine. 388

Differences in partitioning behavior between Fe-rich and Fe-poor compositions may 389 be due to differences in the conditions at which high-to-low spin transitions occur in Fe-390 bearing mantle phases. The decrease in K_D in low Fe³⁺/ Σ Fe olivine was linked to the spin 391 transition of Fe²⁺ in magnesiowüstite [23]. In more Fe-rich olivine, any decrease in K_D can 392 be expected to occur at higher pressures due to the increase in high-to-low spin transition 393 pressure with Fe# in magnesiowüstite [40,41]. The spin transition in Fe#~70 magnesio-394 wüstite in Fa45 can be estimated to begin at ~80 GPa based on observations of similar 395 compositions [40,41,50]. The spin transition in FeO does not occur until ~120 GPa, partic-396 ularly at high temperatures corresponding to conditions during partitioning [e.g. 43] (Fig-397 ure 3a). We can assume that in Fa72 samples, with magnesiowüstite compositions close to 398 pure FeO, the spin transition does not occur in magnesiowüstite in the bridgmanite sta-399 bility field. However, a gradual decrease is observed in K_D in Fa72 in this study (Figure 4). 400This suggests that at high Fe-contents, a decrease in K_D may arise from pressure effects 401 alone on thermoelastic properties of Bdg and Mws. 402

Spin transitions in bridgmanite may increase the partition coefficient, but this has 403 been less clear due to multiple possible crystallographic sites and valence states for Fe 404 [26]. Increases in K_D due to spin transitions in Bdg have been proposed due to a high-to-405 low spin transition in Fe³⁺ at <30 GPa [25]. Since olivine starting materials for these exper-406 iments are nominally Fe³⁺-free, a spin transition in Fe³⁺ will only have a significant effect 407 on K_D if during Bdg/Mws synthesis a substantial amount of Fe²⁺ is oxidized to Fe³⁺. Oxidi-408 zation of Fe²⁺ was limited in Bdg and Bdg-Mws mixtures synthesized from ~100% ferrous 409 starting materials under identical loading and heating protocols to experiments in this 410 study, resulting in Fe³⁺/2Fe of less than 13-25% [51]. In addition, substantial dispropor-411 tionation of Fe²⁺ to Fe³⁺ and Fe⁰ metal is not supported by TEM observations detailed 412 above. A high-to-low spin transition in Fe^{3+} is thus unlikely to significantly affect parti-413 tioning in these experiments. Although a high-to-low spin transition does not occur in 414 Fe²⁺ at mantle-relevant pressures in bridgmanite with low Fe# [20], higher Fe-content may 415 promote this transition in bridgmanite at lower pressures [22,27,52,53]. For Fe#75, a frac-416 tion of Fe²⁺ in Bdg may adopt the low spin state by 80 GPa [22]. Although Bdg in Fa72 417 samples reaches Fe#>40, this is not sufficiently Fe-rich to promote a spin transition in Fe²⁺. 418 The lack of effects of any spin transition in bridgmanite on partitioning is consistent with 419 the strictly decreasing trend of K_D in Fa72 in the bridgmanite stability field (Figure 4). 420

Observations of the partitioning of Fe between bridgmanite and magnesiowüstite 421 versus post-perovskite and magnesiowüstite provide important constraints on the post-422 perovskite transition in Fe-bearing compositions in the mantle and the D" discontinuity. 423 Most measurements of phase equilibria by X-ray diffraction support a lower-pressure, 424 broader pPv transition in Fe-rich compositions [2,19,28,54], which implies that Fe parti-425 tions from Bdg into the pPv phase. Partitioning of Fe from Bdg to pPv is also supported 426 by observations of higher silicate-oxide K_D for pPv than for Bdg [8,11,12]. However, a few 427 studies have suggested that Fe partitions in the opposite direction, based on a higher pPv 428 transition pressure in the presence of Fe [29] and direct partitioning measurements be-429 tween coexisting silicates [14,30,55]. The conflict has been attributed to differences in the 430 $Fe^{3+}/\Sigma Fe$ content of starting materials and synthesized silicates [14,18,26], which is not 431 well-known for all experiments and challenging to determine both in situ and ex situ. The 432 experiments in this study support a higher KD for post-perovskite-magnesiowüstite rela-433 tive to bridgmanite-magnesiowüstite, suggesting a shallower, broader depth for the post-434 perovskite transition in iron-rich lower mantle heterogeneities. 435

Dense mantle heterogeneities may have formed through the crystallization of a basal 437 magma ocean, subduction of dense material, or a combination of multiple mechanisms. 438 Although some models have proposed that such regions could remain relatively chemi-439 cally isolated over Earth's history, preserving primordial material rich in incompatible 440 elements [e.g. 56], depending on the amount of mixing the compositions of heterogeneous 441 regions must be a time-integrated result of chemical behavior during their segregation 442 and since. For example, the basal magma ocean hypothesis may imply that the final dense 443 residue is depleted in Si and enriched in Fe relative to a dominantly bridgmanite lower 444 mantle [5]. The density of large low shear velocity provinces could be consistent with iron-445 enrichment (Deschamps et al., 2012), but whether subducting slabs mix into these regions 446 is not known. To test all models of modern compositions of dense heterogeneities using 447 modern observations of seismic properties, we need to be able to model chemical parti-448tioning between arbitrary mixtures of bridgmanite and oxide phases. 449

Based on our modeled composition- and pressure dependence of iron partitioning, 450we obtain compositions and physical properties of the major phases in the deep lower 451 mantle for a heterogeneous lower mantle containing iron-rich regions. We examine com-452 positions with total Fe# 10, 30, 45 and 72 at selected pressures 40, 60, 80, 100 and 125 GPa 453 (corresponding to depths of roughly 1050, 1470, 1870, 2250 and 2700 km, respectively). 454 Note again that the partitioning values from this work represent low ferric iron and fixed 455 ~peridotitic SiO₂-content. We assume that K_D is independent of mol% SiO₂, expressed as 456 mol% bridgmanite in a bridgmanite – magnesiowüstite phase assemblage. 457

Model compositions for variable Fe#, mol% SiO₂ and pressure are combined with 458 measurements of physical properties of bridgmanite and magnesiowüstite to determine 459 the observable seismic properties of mantle heterogeneities. Recent experimental equation 460 of state measurements over a wide range of compositions have confirmed that the densi-461 ties of bridgmanite and post-perovskite vary linearly with Fe²⁺-content (Supplementary 462 Figure S11) [4]. The Fe#- and pressure-dependence of density of ferropericlase/magnesio-463 wüstite requires a more complex model due to the composition-dependent spin transition. 464 Below the onset of the spin transition, oxide density varies linearly with Fe-content, but 465 at higher pressures the spin transition results in systematically higher densities for fer-466 ropericlase relative to FeO. Densities of mantle phase assemblages at lower mantle pres-467 sures and room temperature (Figures 6-7) were modeled as weighted means between the 468 silicate and oxide densities (see Appendix A). A room-temperature density calculation 469 ignores any difference in thermoelastic properties between Fe-poor and Fe-rich systems, 470 but requires minimal extrapolation from existing experimental data. This model also does 471 not consider Fe^{3+} -rich mantle compositions, in which K_D is substantially different both 472 above and below the post-perovskite transition [26] and a spin transition in bridgmanite 473 affects density [e.g. 57]. Another source of uncertainty in the model is limited experi-474 mental constraints on temperature-dependence of partitioning at deep mantle conditions, 475 with 100s K differences in temperature between the experimental data (Figure 4). How-476 ever, modeled densities are not highly sensitive to uncertainty in partitioning: 50% error 477 bars on K_D result in uncertainty in modeled density of the resulting assemblage of ~0.2-4781% in the deep mantle. 479

While it is well-known that increasing Fe# of the mantle phase assemblage increases 480 its density, the effect of variable SiO₂-content on mantle dynamics is perhaps less intuitive. 481 In average lower mantle rock with Fe#10, the change in density due to difference in mol% 482 SiO_2 , $\partial \rho / \partial SiO_2$, is nearly zero (Figure 6). Increasing the fraction of bridgmanite in the 483 mantle slightly increases net density in the shallow lower mantle (40 GPa and lower pres-484 sures). With increasing pressure, differences in compressibility between bridgmanite and 485 ferropericlase result in a slightly negative change in density with increasing bridgmanite 486 fraction. The post-perovskite transition increases the density of the silicate and $\partial \rho / \partial SiO_2$ 487 becomes slightly positive again at 125 GPa. Adding iron to the mantle not only increases 488 the density, it also modifies $\partial \rho / \partial SiO_2$ (Figures 6-7). An Fe#30 dense mantle heterogeneity 489 exhibits negative $\partial \rho / \partial \operatorname{SiO}_2$ at all modeled pressures (Figure 6). For very Fe-rich 490 compositions at deep mantle pressures, $\partial \rho / \partial SiO_2$ becomes comparable in magnitude to 491 $\partial \rho / \partial Fe\#$ (Figure 7). The depth- and composition-dependence of $\partial \rho / \partial SiO_2$ may provide a 492 mechanism for compositional stratification in the mantle, leading to Fe-bearing, Si-depleted LLSVPs and ULVZs. 494

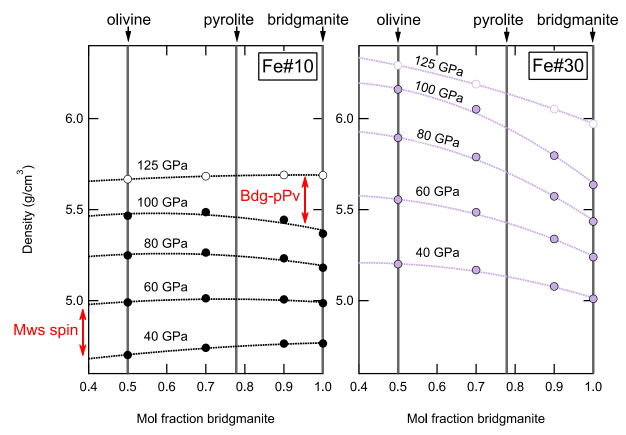


Figure 6. Density of a lower mantle phase assemblage comprising ferrous-iron-bearing bridgmanite, ferropericlase, and post-perovskite as a function of mol% silicate and pressure. The % bridgmanite marked "pyrolite" corresponds to the pyrolite Mg/Si ratio, but does not address other elemental abundances in pyrolite (e.g. Al, Ca). Densities are calculated based on partitioning data in this study and equation of state data for bridgmanite, ferropericlase, and post-perovskite as detailed in Supporting Text. Filled symbols = bridgmanite and ferropericlase phase assemblage, open symbols = post-perovskite and ferropericlase. Compositions are modeled at pressures 40, 60, 80, 100 and 125 GPa (dotted lines = isobars) for fixed bulk Fe# a) Fe#10, representing average mantle, and b) Fe#30, representing dense mantle heterogeneity.

Heterogeneous mantle SiO₂-content may also explain enigmatic differences in seismic velocities V_s and V_P in LLSVPs [56]. The deep lower mantle has been noted for anticorrelation of V_s and V_P , which neither thermal differences nor iron content alone can explain [4]. However, differences in Fe-content in combination with SiO₂-content produce complex effects on density and compressibility. Further studies on composition, pressure, and temperature dependence of thermoelastic properties will test whether dense heterogeneities such as LLSVPs are enriched or depleted in SiO₂.

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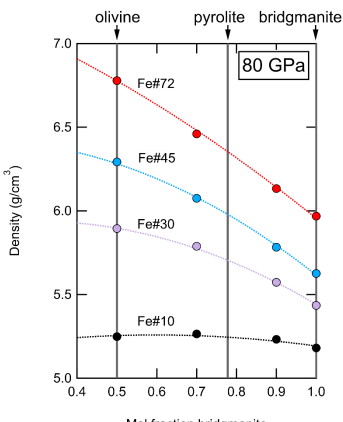
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Mol fraction bridgmanite

Figure 7. At 80 GPa, modeled densities of lower mantle phase assemblages comprising ferrousiron-bearing bridgmanite and ferropericlase. Model compositions (symbols) vary in mol% bridgmanite and bulk iron content. The % bridgmanite marked "pyrolite" corresponds to the pyrolite Mg/Si ratio, but does not address other elemental abundances in pyrolite (e.g. Al, Ca). Dotted lines for constant bulk Fe# are provided as guides to the eye.

5. Conclusions

In order to systematically investigate the effects of iron enrichment on the properties 517 of dense LLSVPs and ULVZs, partitioning of iron between magnesiowüstite, bridgmanite, 518 and post-perovskite was examined in Fe2+-rich olivine compositions, Fa72 and Fa45, at 519 pressure and temperature conditions spanning the entire range for Earth's lower mantle, 520 33-128 GPa and 1900-3000 K. Both in situ X-ray diffraction and ex situ transmission electron 521 microscopy were used to determine synthesized phase assemblages and compositions of 522 each phase. Experimental equations of state for bridgmanite can be used to determine 523 composition within ~20 Fe# of the composition measured by EDX over a compositional 524 range up to at least Fe#50. In all samples, magnesiowüstite composition was enriched in 525 Fe relative to the starting material, demonstrating that Fe favors the oxide over the silicate 526 phase throughout the lower mantle. In Fa72 samples, both EDX and XRD indicate that 527 magnesiowüstite was within 5% Fe/(Mg+Fe) of pure FeO (i.e. Fe#95-100) between 50-100 528 GPa. Fe-content in bridgmanite increases with pressure up to ~80 GPa, reaching a maxi-529 mum Fe# of 51 in Fa72 at 82 GPa. At 128 GPa, the Fe-content of the silicate phase is higher 530 than that measured at lower pressures due to the higher silicate-oxide partitioning coeffi-531 cient of Fe into post-perovskite relative to bridgmanite. The observations show no evi-532 dence of increase in partition coefficient due to a spin transition in bridgmanite. Decrease 533 of the partition coefficient from 60-100 GPa in Fa72 is unlikely to be related to a spin tran-534 sition in magnesiowüstite as the spin transition pressure in Fe#95-100 magnesiowüstite is 535 greater than 100 GPa. Observed partitioning behaviors are successfully reproduced by 536 self-consistent thermodynamic modeling with updated equation of state parameters for 537

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iron-rich bridgmanite and FeO. These results provide important constraints on the effect
 of Fe-content on partitioning of iron and resulting physical properties of iron-rich lower
 mantle heterogeneities. In iron-rich low shear velocity provinces, different effects of iron
 on densities of magnesiowüstite and bridgmanite result in increased effect of SiO₂-content
 to reduce the density of the bulk phase assemblage. Dynamics of iron-rich regions over
 Earth's history may have resulted in Si-depletion at the base of the mantle in LLSVPs and
 ULVZs.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Supplementary Figures S1-S11, Supplementary Tables S1-S2. 546

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Appendix A

Physical properties of iron-rich mantle phases

Interpretation of *in situ* X-ray diffraction data and inferred physical properties of 572 mantle phase assemblages require pressure-volume-composition equation of state data 573 for bridgmanite (Bdg), ferropericlase-magnesiowüstite (Mws), and post-perovskite (pPv). 574

For quantitative analysis of Bdg composition at high pressures, recent compression 575 measurements of single-phase Bdg with a wide range of Fe# synthesized in the laser-576 heated diamond anvil cell [19] were used to fit a pressure-lattice spacing-composition 577 equation of state. The equation of state calibrates composition as a function of pressure 578 and the *a* lattice parameter of bridgmanite, the direction most sensitive to Fe-content. For 579 lattice parameter data from three studies of Fe-bearing Bdg by powder diffraction in 580 quasi-hydrostatic media [19,39,58], compositions indicated by this calibration scatter 581 around known compositions of starting materials with a standard deviation of ~5% (Sup-582 plementary Figure S7b). Note that data used for equation of state fits in these studies in-583 clude only data obtained for Bdg observed without accessory SiO2 or (Mg,Fe)O, and thus 584 the compositions of the starting materials are assumed to be identical to the synthesized 585 Bdg. Precision of composition estimates by XRD for this study and other typical partition-586 ing studies will be lower due to the need to avoid internal pressure calibrants, which may 587 complicate chemistry and/or result in Soret diffusion [35]. 588

The composition of Mws is more difficult to model based on to unit cell volume than 589 ferrous-iron-bearing Bdg due to the composition-dependent spin transition. For cubic 590 Mws, unit cell volumes of B1-structure (Mg,Fe)O and B8 FeO were used to determine the 591 pressure-volume-composition-spin state equation of state [41,42,50,59-68]. Magnesio-592 wüstite with Fe#≥95 may adopt the B8 structure at the pressure conditions of these exper-593 iments [69], but the diffraction peaks observed from this phase are not resolved well 594 enough to determine its structure (Figure 1). Qualitatively, Mws volumes in Fa45 samples 595 fall between previous measurements of Fe#58 and Fe#100 (FeO) compositions (Supple-596 mentary Figure S8). The measured volume of magnesiowüstite formed from Fa72 is con-597 sistent with nearly pure FeO. 598

These equation of state data for Bdg and Mws and composition-dependent equation 599 of state of post-perovskite [44,70,71] were used to compute densities of mantle phases 600 based on compositions set by partitioning constraints at model pressures of 40, 60, 80, 100, 601 and 125 GPa. Mws pressure-volume-composition data were interpolated at each model 602 pressure using quadratic trends (Supplementary Figure S9). 603

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