

Phase relations of Fe-Si alloy in Earth's core

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[1] Phase relations of an $Fe_{0.85}Si_{0.15}$ alloy were investigated up to 240 GPa and 3000 K using in situ X-ray diffraction in a laser-heated diamond anvil cell. An alloy of this composition as starting material is found to result in a stabilized mixture of Si-rich bcc and Si-poor hcp Fe-Si phases up to at least 150 GPa and 3000 K, whereas only hcp-Fe_{0.85}Si_{0.15} is found to be stable between approximately 170 GPa and 240 GPa at high temperatures. Our extended results indicate that $Fe_{0.85}Si_{0.15}$ alloy is likely to have the *hcp* structure in the inner core, instead of the previously proposed mixture of *hcp* and *bcc* phases. Due to the volumetric dominance of the hcp phase in the hcp + bcc coexistence region close to the outer-core conditions, the dense closest-packed Fe-Si liquid is more relevant to understanding the properties of the outer core. Citation: Lin, J.-F., H. P. Scott, R. A. Fischer, Y.-Y. Chang, I. Kantor, and V. B. Prakapenka (2009), Phase relations of Fe-Si alloy in Earth's core, Geophys. Res. Lett., 36, L06306, doi:10.1029/2008GL036990.

1. Introduction

[2] Geophysical and cosmochemical evidence indicates that iron is the most abundant component in the Earth's core [Birch, 1952]. However, there is a density deficit between seismic observations of Earth's core and high pressuretemperature iron, implying the presence of a certain amount of element(s) lighter than iron at an abundance of 6-10 wt%for the outer core and 2-5 wt% for the inner core [e.g., Poirier, 1994; Li and Fei, 2003]. Silicon has been proposed to be a major alloving light element in the Earth's core [e.g., Poirier, 1994; Li and Fei, 2003] based on its cosmochemical abundance, solubility in liquid Fe at high pressures and temperatures [e.g., Takafuji et al., 2005; Sakai et al., 2006], and thermoelastic properties in Fe-Si alloys [e.g., Lin et al., 2003a, 2003b; Hirao et al., 2004]. These recent results indicate that an outer core containing about 8-10 wt% Si and inner core containing about 4 wt% Si in Fe would satisfy seismological constraints, although other light elements such as oxygen and sulfur could also exist in the core [e.g., Takafuji et al., 2005; Sakai et al., 2006]. Studying the phase relations and thermoelastic properties of the Fe-rich Fe-Si alloy is thus essential for our understanding of Earth's core.

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[3] Pure Fe is found to be stable in the hexagonal closestpacked (*hcp*) structure over a wide range of pressures and temperatures approaching suspected Earth's core conditions, whereas reports on the occurrence of a doubled *hcp*-Fe and an orthorhombic Fe at high pressures and temperatures are now mostly believed to be a result of experimental problems [e.g., *Shen et al.*, 1998; *Ma et al.*, 2004; *Kuwayama et al.*, 2008]. Theoretical calculations further suggest that the *hcp*-Fe transforms into a body-centered cubic (*bcc*) Fe at conditions close to its melting line at core pressures [e.g., *Vocadlo et al.*, 2003; *Belonoshko et al.*, 2008], but this has not been confirmed experimentally.

[4] Under ambient conditions, Fe-rich Fe-Si alloys with a Si content relevant to Earth's core (<10 wt%) are stable in the *bcc* structure. Theoretical studies have shown that addition of a few weight percent of Si into Fe can stabilize the bcc Fe-Si alloy at the pressure-temperature conditions of the core [Cote et al., 2008a, 2008b; Vocadlo et al., 2003]. Whereas earlier high pressure-temperature experiments have shown that Si alloyed with Fe can stabilize the bcc phase up to at least 84 GPa (compared to ~ 10 GPa for pure Fe) and 2400 K [Lin et al., 2002a; Lin, 2002], recent studies, however, have suggested that Fe-rich Fe-Si alloys dissociate into almost pure Fe and a b2-structured FeSi compound [Dubrovinsky et al., 2003], or a mixture of face-centered cubic (fcc) and hcp phases [Asanuma et al., 2008]. Since these phases may differ in physical properties such as sound velocities, knowing the stable crystal structure of the Fe-Si alloy under relevant conditions is thus essential for our understanding of Earth's core. To decipher the phase relations of the Fe-Si alloy at extended pressure-temperature conditions relevant to Earth's core, here we have studied the phase relations of an Fe0.85Si0.15 alloy (containing 8 wt% Si) up to 240 GPa and 3000 K using in situ X-ray diffraction in a laser-heated diamond anvil cell (DAC).

2. Experiments

[5] The starting Fe_{0.85}Si_{0.15} alloy was obtained from Goodfellow Corporation. X-ray diffraction and electron microprobe analyses showed that the sample was in the *bcc* structure and chemically homogenous. Samples measuring approximately $5-15 \mu$ m thick and 25μ m in diameter were loaded into symmetric DACs with 75–300 μ m beveled diamonds or $60-180-300 \mu$ m double-beveled diamonds using Re gaskets. We also used cubic boron nitride (BN) gasket inserts in some of the experiments to prepare samples with sufficient thickness for laser heating and diffraction measurements and to avoid diffraction peaks from Re gaskets. Samples were sandwiched between dried NaCl layers in the DACs, and the NaCl was used as the thermal insulator, pressure medium, and pressure calibrant. Pressures were determined from the lattice parameter and equation of state

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Figure 1. Representative X-ray diffraction patterns of $Fe_{0.85}Si_{0.15}$ at high pressures and temperatures. (a) Detailed spectra at 143 GPa, (b) detailed spectra at 193 GPa, and (c) full diffraction spectra after quenching from highest temperatures at high pressures (see Figure 2 for the highest temperatures). Vertical ticks in Figure 1c represent modeled diffraction peaks of *bcc*, *hcp*, *fcc*, and *b2* phases. *NaCl*, sodium chloride in the *CsCl* structure; *Re*, rhenium gasket. Note that the *b2* and *fcc* peaks were not observed in these spectra. The wavelength of the monochromatic X-ray source was 0.3344 Å.

of NaCl before heating [Fei et al., 2007], and were also measured after heating. In situ high pressure-temperature X-ray diffraction experiments were conducted at the Geo-SoilEnviroCARS sector of the Advanced Photon Source (APS), Argonne National Laboratory (ANL) [Prakapenka et al., 2008]. We used two diode-pumped ytterbium infrared fiber lasers operating in single-mode continuous wave, with a wavelength of 1064 nm, and focused flat-top shapes approximately 15 μ m in diameter at the sample position to laser-heat the sample from both sides of the DAC. A monochromatic beam 0.3344 Å in wavelength and 5 μ m in diameter (at Full Width at Half Maximum (FWHM)) was used as the incident X-ray source and was well aligned with the laser-heating spots using the X-ray fluorescence of the sample, and the diffracted X-rays were collected by a CCD (MAR165). Graybody temperatures were determined by fitting the thermal radiation spectra between 670 and 830 nm to the Planck radiation function. The temperature uncertainty (1σ) , averaged from multiple temperature measurements, ranges from 100 K to 250 K in most of the experiments.

3. Experimental Results

[6] The phase relations of the Fe_{0.85}Si_{0.15} alloy have been investigated up to 240 GPa and 3000 K (Figures 1 and 2). Prior to laser heating, we compressed the *bcc*-Fe_{0.85}Si_{0.15} starting material to pressures between 100 and 240 GPa at room temperature and found that all transformed to the *hcp* structure. Each sample was then laser heated, and diffraction patterns were collected at a series of temperature steps up to maximum heating temperatures as high as 3000 K (see auxiliary material for details); these data were used to constrain the phase relations as indicated in Figure 2.¹ Upon heating, the *hcp*-Fe_{0.85}Si_{0.15} transitioned to a mixture of the Si-poor *hcp* and Si-rich *bcc* phases at pressures up to at least 150 GPa (Figure 2), whereas the *hcp*-Fe_{0.85}Si_{0.15} phase is observed to be stable between approximately 170 GPa and 240 GPa at high temperatures. Following previous experiments on the identification of the Fe-Si polymorphs [*Lin* et al., 2002a; *Lin*, 2002], the occurrence of the *bcc* phase can be identified by the *bcc* (110) peak in between the *hcp* (002) and (101) peaks at high temperatures, although the intensity of the *bcc* (110) peak is relatively weaker compared with



Figure 2. Observed phases from an Fe_{0.85}Si_{0.15} starting material superimposed on a phase diagram of pure Fe (solid lines). The phase diagram for Fe is modified from Nguyen and Holmes [2004] and includes data from static DAC experiments [Shen et al., 1998; Ma et al., 2004]. Typical error bars for our pressure and temperature measurements on $Fe_{0.85}Si_{0.15}$ are shown in the bottom right corner. Open circles, hcp-Fe_{0.85}Si_{0.15} only; solid triangles, coexistence of *hcp* and *bcc* phases; dashed lines, phase boundary between hcp-Fe_{0.85}Si_{0.15} and *hcp/bcc* Fe-Si alloys from this study, *Lin* et al. [2002a], and Lin [2002]; grey hexagon, bcc-Fe_{0.9}Ni_{0.1} at 225 GPa and 3400 K [Dubrovinsky et al., 2007]; dashdotted and dotted lines, theoretically predicted melting curve of Fe by Alfe et al. [2002] and Laio et al. [2000], respectively; open triangle, square, and diamond, melting data on pure Fe from shock-wave experiments by Ahrens et al. [2002], Brown and McQueen [1986], and Nguyen and Holmes [2004], respectively.

¹Auxiliary materials are available in the HTML. doi:10.1029/2008GL036990.



Figure 3. Unit cell parameters of the Si-rich *bcc* (red open triangles) and Si-poor *hcp* phases (blue open circles) at high pressures and room temperature. Lattice parameters of the *hcp*-Fe_{0.85}Si_{0.15} phase alone (without coexistence with the bcc phase; black open squares) are also plotted for comparison. The coexisting *bcc* and *hcp* phases were produced at 148 GPa and high temperatures (Figure 2) and decompressed at room temperature. Insert, axial ratio (*c/a*) of the *hcp* phase.

lower-pressure results (Figure 1). These results show that the volume ratio of the *bcc* phase to the *hcp* phase increases with increasing temperature and decreases with increasing pressure (Figure 1) (see also *Lin* [2002] for detail of the diffraction patterns in the *bcc* + *hcp* coexistence region), indicating that the *hcp* phase is the stable phase at higher pressures.

[7] Since most intense diffraction peaks of the suspected hcp, bcc, fcc, and b2 phases of Fe-Si alloys are very close to each other according to previous reports [Lin et al., 2002a; Dubrovinsky et al., 2003; Asanuma et al., 2008] (Figure 2), we moved the CCD to various distances from the sample to increase the spatial resolution of the diffraction peaks for phase identification in particular regions of interest. We observed no X-ray diffraction evidence of the previously reported fcc or b2 phases of the Fe-Si alloy. The molar volume of the *bcc* phase is about 1% smaller than that of the coexisting hcp at 143 GPa and 300 K, whereas its incompressibility based on experimental data between 60 GPa and 143 GPa is similar to those of the *hcp*-Fe_{0.85}Si_{0.15} [Lin et al., 2003a] and hcp-Fe phases [Mao et al., 1990] (Figure 3). Our measured c/a axial ratio of the hcp-Fe_{0.85}Si_{0.15} phase remains almost constant at 1.60 (± 0.01) up to 240 GPa. Using unit cell parameters determined from this study and the compositional difference of 4 atomic % in Si between the lower-density Sirich bcc and higher-density Si-poor hcp phases reported by Lin et al. [2002a; Lin, 2002], the density difference between the bcc and hcp phases is approximately 2% at 100–150 GPa.

4. Discussion and Geophysical Applications

[8] Compared to the phase diagram of pure Fe (Figure 2), it is evident that hcp-Fe_{0.85}Si_{0.15} is stable as a single phase between 170 GPa and 240 GPa at high temperatures, whereas it separates into a mixture of the *bcc* and *hcp* Fe-Si phases at lower pressures up to at least 150 GPa at high temperatures. To explain X-ray diffraction patterns of Fe-Si alloys, here we further discuss the stacking sequences of the Fe polymorphs. The layered dense-packed structure of Fe and Fe-rich alloys

can result in various stacking polymorphs—hcp phase with ABAB stacking sequence along [001], fcc with ABCABC stacking along [111], and doubled *hcp* phase with *ABAC* stacking along [001]. Because of the similarities in the stacking layers, the difference in the Gibbs free energy between these phases at high pressures and temperatures is very small, and variations in experimental conditions such as temperature gradient have been shown to result in the occurrence of the *fcc* or doubled *hcp* phase at high pressures and temperatures. The *hcp* and *fcc* phases share the same lattice plane in the stacking layers along [001] and [111], respectively, which are manifested in their overlapping diffraction peaks in hcp (002) and fcc (111) in Fe and Fe-Si alloys over a wide pressure-temperature range (Figure 1b) [e.g., Shen et al., 1998; Lin et al., 2002a; Lin, 2002; Mikhaylushkin et al., 2007]. The large splitting between the assigned hcp (002) and fcc (111) peaks for $Fe_{0.93}Si_{0.07}$ by Asanuma et al. [2008], however, contradicts the aforementioned statement and cannot be explained by the occurrence of the metastable *fcc* phase alone, indicating that this assignment is most likely incorrect.

[9] Dubrovinsky et al. [2003] reported that Fe-rich Fe-Si alloys dissociate into almost pure *hcp*-Fe and *b2*-structured FeSi with the 1:1 atomic ratio at high pressures and temperatures, indicating the occurrence of an immiscibility gap between the phases. Their transition boundary between the *hcp* and *hcp* + b2 regions had a negative slope, and the transition from the low-pressure bcc Fe-Si alloy to the high-pressure b2-structured FeSi phase is discontinuous [Dubrovinsky et al., 2003], inconsistent with our results and previous experimental and theoretical studies [Lin et al., 2002a; Cote et al., 2008a, 2008b]. The b2-structured FeSi with the 1:1 atomic ratio reported by *Dubrovinsky et al.* [2003] is the fully ordered form of the *bcc* Fe-Si alloy, requiring a significant amount of Si partitioned into the phase. We note that beside different origins of the samples Dubrovinsky et al. [2003] mainly used electrically-heated DAC experiments with hours of heating before the observation of the b2-structured FeSi phase, while their laser-heated DAC experiments were conducted offline. It is thus conceivable that a small amount of carbon diffusion from the diamond anvils to the sample in the long time heating experiments [Prakapenka et al., 2003] could stabilize this phase, which in turn may affect the slope of the phase boundary. In our on-line high pressure-temperature experiments we have detected in-situ effect of temperature cycling on appearance-disappearance of the *bcc* phase that is necessary to constrain phase boundary slope. Based on our current and previous studies [Lin et al., 2002a], we observed the bcc phase coexisted with the *hcp* phase but found no evidence for the b2 (100) reflection. We note that the compositional difference in Si between the bcc and hcp phases is approximately 3-4 atomic % [Lin et al., 2002a], which is clearly insufficient to form the fully ordered b2-structured FeSi phase.

[10] Previous studies estimate that an inner core containing about 4 wt% Si in Fe would satisfy seismological constraints. Our results on $Fe_{0.85}Si_{0.15}$ alloy (with 8 wt% Si) up to 240 GPa and 3000 K show that it is stable in the *hcp* phase above 170 GPa at high temperatures (Figure 3), suggesting Fe-Si alloy is likely in the *hcp* structure in the inner core, instead of a previously proposed mixture of *hcp* and *bcc* phases [*Lin et al.*, 2002a]. However, the additional effect of Ni in stabilizing the *fcc* or *bcc* phases and the predicted occurrence of *bcc*-Fe in the core all remain to be further investigated [e.g., *Lin et al.*, 2002b; *Vocadlo et al.*, 2003; *Mao et al.*, 2006; *Dubrovinsky et al.*, 2007; *Belonoshko et al.*, 2008; *Kuwayama et al.*, 2008].

[11] The observed mixture of the *bcc* and *hcp* alloys up to at least 150 GPa and 3000 K may have implications for our understanding of the properties of the outer core (Figure 2). The Si-rich *bcc* and Si-poor *hcp* phases should have different melting points at their coexisting pressures due to their compositional and structural differences, which could provide a mechanism for Si partitioning between liquid and solid Fe in the core. Based on the concept that the local structure and properties of a liquid mimics that of a relevant solid [*Sanloup et al.*, 2000; *Shen et al.*, 2004], the volumetric dominance of the *hcp* phase in the *hcp* + *bcc* coexistence region indicates that the dense closest-packed Fe-Si liquid is more relevant to understanding the properties of the liquid outer core.

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