Revision 1

The axial ratio of hcp Fe and Fe–Ni–Si alloys to the conditions of Earth’s inner core

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Abstract

The Earth’s iron-rich inner core is seismically anisotropic, which may be due to the preferred orientation of Fe-rich hexagonal close packed (hcp) alloy crystals. Elastic anisotropy in a hexagonal crystal is related to its $c/a$ axial ratio; therefore, it is important to know how this ratio depends on volume (or pressure), temperature, and composition. Experimental data on the axial ratio of iron and alloys in the Fe–Ni–Si system from 15 previous studies are combined here to parameterize the effects of these variables. The axial ratio increases with increasing volume, temperature, silicon content, and nickel content. When an hcp phase coexists with another structure, sample recovery and chemical analysis from each pressure-temperature point is one method for determining the phase’s composition and thus the position of the phase boundary. An alternate method is demonstrated here, using this parameterization to calculate the composition of an hcp phase whose volume, temperature, and axial ratio are measured. The hcp to hcp+B2
phase boundary in the Fe–FeSi system is parameterized as a function of pressure, temperature,
and composition, showing that a silicon-rich inner core may be an hcp+B2 mixture. These
findings could help explain observations of a layered seismic anisotropy structure in the Earth’s
inner core.

Introduction

In the upper part of the Earth’s iron-rich inner core, seismic waves traveling parallel to
the planet’s rotational axis propagate ~3% faster than waves traveling in the equatorial plane
(Morelli et al. 1986; Poupin et al. 1983). Seismic data suggest the existence of layered
structures and hemispherical variations of this anisotropy (e.g., Irving and Deuss 2011a, 2011b;
mechanisms for explaining these properties have been considered (e.g., Alboussière et al. 2010;
Bergman 1997; Buffett and Wenk 2001; Jeanloz and Wenk 1988; Karato 1999; Reaman et al.
2011; Yoshida et al. 1996). This anisotropy is frequently attributed to preferred orientation of Fe-
rich alloy crystals in the inner core, due to iron’s strong single-crystal elastic anisotropy (Jeanloz
and Wenk 1988; Morelli et al. 1986; Stixrude and Cohen 1995). The c/a axial ratio of a
hexagonal crystal directly influences its elastic anisotropy (e.g., Steinle-Neumann et al. 2001;
Vočadlo et al. 2009; Wenk et al. 1988); therefore, the c/a ratio of preferentially aligned crystals
of a candidate core material can be related to the anisotropy of Earth’s inner core. For example,
c/a ratios of iron alloys can serve as input in calculations of elastic moduli or in models of core
anisotropy, as has previously been done in the case of pure hexagonal close packed (hcp) iron
(e.g., Steinle-Neumann et al. 2001; Vočadlo et al. 2009).
The subsolidus phase relations and equation of state of pure iron at high pressures have been studied numerous times (e.g., Anzellini et al. 2013; Boehler et al. 2008; Dewaele et al. 2006; Jephcoat et al. 1986; Komabayashi et al. 2009; Ma et al. 2004; Mao et al. 1990; Ono et al. 2010; Sakai et al. 2011; Tateno et al. 2010; Uchida et al. 2001; Yamazaki et al. 2012) due to its key geophysical applications. Some studies report a trend of the c/a ratio of hcp iron decreasing with increasing pressure (Boehler et al. 2008; Dewaele et al. 2006; Jephcoat et al. 1986; Yamazaki et al. 2012), while others report seemingly no trend, a very weak trend, or a trend that changes with pressure (Jephcoat et al. 1986; Ma et al. 2004; Mao et al. 1990; Ono et al. 2010). Similarly, some studies find that the c/a ratio of iron increases with increasing temperature (Boehler et al. 2008; Sakai et al. 2011; Tateno et al. 2010; Yamazaki et al. 2012), in agreement with most theoretical calculations (Belonoshko et al. 2003; Gannarelli et al. 2005; Modak et al. 2007; Sha and Cohen 2006; Steinle-Neumann et al. 2001; Wasserman et al. 1996), while others do not (Ma et al. 2004). A few studies on iron (Boehler et al. 2008; Jephcoat et al. 1986) and Fe–Ni–S alloy (Sakai et al. 2012) fit their room temperature data to a function describing how the axial ratio changes with pressure, but it would be more useful to compile the extensive literature on the c/a ratios of iron alloys into a single parameterization of their volume, temperature, and compositional dependence; this is a goal of the present study.

Earth’s inner core is known to contain several weight percent of nickel and of one or more elements lighter than iron (Birch 1952; Jephcoat and Olson 1987). Silicon is one of the leading candidates for comprising this light element component, based on its abundance, presence in the metal of some chondritic meteorites, partitioning behavior, and non-chondritic isotope ratio in the mantle (e.g., Allegre et al. 1995; McDonough 2003; Shahar et al. 2009). Density functional calculations have indicated that Fe–Si hcp alloys can have significantly higher
elastic anisotropy than pure iron (Tsuchiya and Fujibuchi 2009), so it is important to consider the
composition of the alloy in addition to its structure when interpreting seismic anisotropy of the
inner core. Silicon-bearing alloys in the Earth’s inner core may be an hcp+B2 mixture (Fischer et
al. 2013), depending on composition and temperature, which could offer an alternate explanation
for anisotropy. However, the exact phase stability of the hcp+B2 mixture of Fe–Si alloys remains
elusive, due to the difficulty of obtaining compositional measurements of coexisting phases in
experiments at the extreme pressures and temperatures of the Earth’s core.

One of the goals of the present study is to combine some of the many available datasets
on the c/a ratio of pure iron and alloys in the Fe–Ni–Si system, to develop a single expression for
the variation in c/a ratio as a function of volume, temperature, and composition. This
parameterization can be used to calculate the compositions of the hcp component of hcp+B2
mixtures of Fe–9wt%Si (Fe–9Si) and Fe–16wt%Si (Fe–16Si) at high pressures using literature
data (Fischer et al. 2012, 2014). This will allow us to put better constraints on the Fe–Si phase
diagram at high pressures and temperatures, in particular the crystal structure of an Fe–Si alloy at
inner core conditions, and to enhance our understanding of anisotropy in Fe–Ni–Si alloys at core
conditions.

Methods

For our study of the c/a ratio of hcp iron, we selected numerous X-ray diffraction datasets
from among the studies on the equation of state and phase relations of iron. These datasets were
chosen based on their relatively smaller degree of scatter (see Discussion), the inclusion of
published c/a ratios, and general compatibility with other modern results on the trends of the c/a
ratio of iron with pressure and temperature. The studies used here consist of both diamond anvil
cell (Anzellini et al. 2013; Boehler et al. 2008; Dewaele et al. 2006; Fischer et al. 2011; Ono et al. 2010; Sakai et al. 2011; Tateno et al. 2010) and multi-anvil press (Uchida et al. 2001; Yamazaki et al. 2012) studies. We generally selected more recent datasets, which exhibit more precise data due to improvements in experimental techniques.

These data were all collected using in situ synchrotron X-ray diffraction, using a variety of different pressure standards and calibrations. Consideration of the pressure calibration is critical when comparing results of different studies; here we largely circumvent this difficulty by parameterizing $c/a$ as a function of volume instead of pressure. In a few studies, only the pressure, temperature, and $c/a$ ratio was reported, and an equation of state was necessary to determine the measured volume. Boehler et al. (2008) do not report unit cell volumes, and used the ruby fluorescence pressure scale of Mao et al. (1986). We corrected their pressure measurements to the ruby scale of Dorogokupets and Oganov (2007), which was calibrated against the pressure scale of Dewaele et al. (2006). We then calculated the volume of iron in their experiments from the corrected pressure and the reported temperature using the equation of state of Dewaele et al. (2006). Likewise, Tateno et al. (2010) used the hcp iron equation of state of Dubrovinsky et al. (2000) for their pressure calibration, so we used the Dubrovinsky et al. (2000) equation of state to extract unit cell volumes from the pressures and temperatures reported by Tateno et al. (2010).

To investigate the effects of nickel and silicon on the $c/a$ ratio of hcp Fe alloys, we used data from Fischer et al. (2014) on Fe–9Si, Lin et al. (2002a) on Fe–10Ni, Lin et al. (2002b) on Fe–8Si, Sakai et al. (2011) on Fe–10Ni and Fe–5Ni–4Si, Komabayashi et al. (2012) on Fe–10Ni, Tateno et al. (2012) on Fe–10Ni, and Tateno et al. (2015) on Fe–9Si and Fe–7Si. Lin et al. (2002a, 2002b) do not report volumes, so the volumes were calculated using their reported
pressures and temperatures and the equations of state of pure iron (Dewaele et al. 2006) and Fe–$9\text{Si}$ (Fischer et al. 2014). The equation of state of Fischer et al. (2014) was calibrated against an equation of state of B2 KBr (Fischer et al. 2012), which in turn was calibrated against the equation of state of Dewaele et al. (2006), so these scales should be consistent.

**Results**

The data we have compiled on the $c/a$ ratio in hcp Fe–Ni–Si alloys are listed in Supplemental Table S1. It includes 928 measurements taken from 15 different studies, listing the $c/a$ ratio, its uncertainty where available, volume, temperature, lattice parameter $a$, and mole fraction of silicon and nickel for each measurement. There are 632 measurements of pure Fe, 100 of approximately Fe$_{0.8}$Ni$_{0.2}$ (Fe–10Ni), 6 of Fe$_{0.88}$Ni$_{0.04}$Si$_{0.08}$ (Fe–5Ni–4Si), 10 of Fe$_{0.88}$Si$_{0.12}$ (Fe–6.5Si) and 180 of approximately Fe$_{0.84}$Si$_{0.16}$ (Fe–9Si). Data on pure Fe span ~6–340 GPa (Dewaele et al. 2006) and 300–4890 K, data on Fe–Ni alloys span ~25–340 GPa and 300–4700 K, and data on Fe–Si alloys span ~13–407 GPa and 300–5910 K.

Figure 1a shows the $c/a$ ratio of pure iron from numerous studies as a function of volume and temperature (Anzellini et al. 2013; Boehler et al. 2008; Dewaele et al. 2006; Fischer et al. 2011; Ono et al. 2010; Sakai et al. 2011; Tateno et al. 2010; Uchida et al. 2001; Yamazaki et al. 2012). This figure illustrates that there is still considerable scatter within and between the datasets shown here; however, there is even greater scatter in the older datasets not chosen for inclusion in this study (e.g., Jephcoat et al. 1986; Ma et al. 2004; Mao et al. 1990). Regardless, Figure 1a demonstrates a trend of increasing $c/a$ ratio with increasing volume and with increasing temperature. Similarly, the $c/a$ ratios of hcp Fe–Ni alloys containing ~10 wt% Ni are shown in Figure 1b (Komabayashi et al. 2012; Lin et al. 2002a; Sakai et al. 2011; Tateno et al. 2010; Uchida et al. 2001).
Figure 1c shows the \( c/a \) ratio of hcp Fe–9Si where it does not coexist with any other phases (Fischer et al. 2014; Lin et al. 2002b; Tateno et al. 2015). The \( c/a \) ratio in these alloys increases with both volume and temperature, similar to that of pure iron (Figure 1a). If the axial ratio of hcp iron approaches its ideal value of 1.633 at core conditions, this could prevent anisotropy in the inner core by preferential alignment of hcp crystals (e.g., Steinle-Neumann et al. 2001). However, the data in Figure 1 indicate that the temperature dependence at inner core conditions is not strong enough for the \( c/a \) ratio to approach this value for pure Fe or Fe–Ni–Si alloys.

Figure 2 illustrates the \( c/a \) ratio of Fe–9Si changing as a function of temperature during a single heating cycle at ~145 GPa (Fischer et al. 2014). For reference, data on the \( c/a \) ratio of pure iron from several studies at ~130–155 GPa are also shown (Anzellini et al. 2013; Boehler et al. 2008; Fischer et al. 2011; Tateno et al. 2010). Within experimental precision, the temperature dependence of the measured axial ratio at this pressure appears to be the same for Fe–9Si and for pure Fe, since the trends in Figure 2 are parallel. The \( c/a \) ratio of the alloy is higher than that of iron by ~0.009 at these conditions.

Discussion

Axial ratios as a function of volume, temperature, and composition

The \( c/a \) ratio of an Fe–rich hcp alloy is a function of volume \( (V) \), temperature \( (T) \), and composition \( (X_{\text{Si}}, X_{\text{Ni}}) \). Using the data shown in Figure 1 and listed in Table S1, we have parameterized its dependence on these variables in the Fe–Ni–Si system as an unweighted linear fit to the data. We found the best fit to the compiled data with the following relationship:

\[
c/a = 1.551 + (-2.6 \times 10^{-5})T + 0.0094V + (1.5 \times 10^{-5})T^2 + V + \\
\]
(4.4 \times 10^{-4})X_{Si} + (2.8 \times 10^{-4})X_{Ni} \tag{1}

with \( T \) in Kelvin, \( V \) in \( \text{cm}^3/\text{mol} \), and \( X_{Si} \) and \( X_{Ni} \) in mole fraction (e.g., \( X_{Si} = X_{Ni} = 0 \) for pure iron and \( X_{Si} = 16 \) for Fe–9Si). The variance-covariance matrix describing this fit is shown in Supplemental Table S2. Based on the amount of scatter in the data, it was not justifiable to fit any additional terms to Equation 1; for example, allowing the temperature dependence of the \( c/a \) ratio to vary with composition did not significantly improve the quality of the fit.

Figures 1 and 2 contain curves calculated from this parameterization compared to the data. As volume increases, the \( c/a \) ratio increases. The \( c/a \) ratio always increases with increasing temperature, but this effect is strongest at higher volumes, and becomes weaker with decreasing volume. Based on the available data, we could not justify fitting any more than linear compositional terms. This implies that the axial ratios of alloys in the Fe–Ni–Si system have similar volume and temperature dependences, regardless of composition, and are simply shifted from each other based on their silicon and nickel contents.

Residuals to this fit from various studies are illustrated as a function of temperature and volume in Figure 3, which demonstrates the adequacy of Equation 1 to describe the data. The root mean squared (rms) average misfit between measured \( c/a \) ratios and those calculated from Equation 1 is 0.004 for the studies included in the fit, with the rms misfits for individual datasets ranging from 0.002 to 0.007. While these misfits are small in absolute value, they equate to \(-10\%\) of the range seen in the data. In comparison, previous studies not included in the fit due to a higher degree of scatter exhibit an estimated 2\( \sigma \) variation of \( >0.01 \). The \( c/a \) ratio is likely affected by deviatoric stresses as well, which may explain the greater misfit in the 300 K data, but in this analysis we estimate that high experimental temperatures are sufficient to relax the hcp alloy and minimize deviatoric stress effects on the axial ratio.
Sakai et al. (2012) studied the effects of sulfur on the \(c/a\) ratio of hcp iron–nickel alloys at 300 K and high pressures. They found an approximately linear relationship between \(c/a\) and pressure, which is different from the trend we observe in this more extensive cross-study analysis. Their data indicate that adding 2.8 mol% S to Fe–9Ni alloy lowers the \(c/a\) ratio by ~0.005, though this varies with pressure due to the different pressure trend reported in their study.

Using the axial ratio of an Fe–Si alloy to determine its composition

Our fit to the available literature data relates the \(c/a\) ratio, temperature, volume, and composition of an alloy in the Fe–Ni–Si system. Therefore, in circumstances where independent measurements of the \(c/a\) ratio, temperature, and volume of an Fe–Si alloy are available from X-ray diffraction and spectroradiometry (or thermocouple sensor), this parameterization may be used to calculate the alloy’s silicon content. In Fe–9Si and Fe–16Si, coexisting hcp+B2 structures are observed in some regions of phase space (Fischer et al. 2012, 2013). When two Fe–Si phases are present, silicon partitions between them as a function of pressure and temperature such that the compositions of the phases are unknown without sample recovery and analysis from each \(P-T\) point of interest. The method presented here provides an alternative to this restrictive process.

Figure 4 illustrates the results of this method, applied to synchrotron X-ray diffraction measurements of Fe–Si alloys at 125 and 145 GPa. At each \(P-T\) point, we have used the measured \(c/a\) ratio and Equation 1 to calculate the composition of the hcp phase that coexists with the B2 structure, thus providing the composition along the hcp to hcp+B2 phase boundary. We used data on Fe–9Si (Fischer et al. 2014) and Fe–16Si (Fischer et al. 2012), which are shown
in Figure 4 to be mutually consistent in their definitions of the phase boundary. The in situ
crossing of this phase boundary determined in Fe–9Si (Fischer et al. 2013) agrees with the
calculated compositions along the phase boundary within uncertainty at both pressures, verifying
our results. As is evident in Figure 4, this method is most useful when applied to data at
temperatures far from any observed phase boundary crossings, to best constrain the slope of the
boundary in T-X space.

Under experimental P-T conditions where an hcp Fe–9Si alloy does not coexist with any
other phase, its composition is the same as the bulk starting composition. We have compared its
known composition under these conditions to its composition calculated from its c/a ratio to
determine the precision of our method, finding a root mean square (rms) misfit of 3.9 wt%
silicon, based on all 147 high temperature measurements spanning 45–407 GPa (Fischer et al.
2014; Lin et al. 2002b; Tateno et al. 2015). We take this as an estimate of the uncertainty in the
calculations shown in Figure 4. The misfit might be greater at different compositions; since the
fit is based only on alloys with up to 9 wt% Si, we caution against its extrapolation to alloys with
considerably higher silicon contents.

Figure 4 illustrates that at temperatures approaching the eutectic temperature in the Fe–
FeSi system (3700(300) K at 125 GPa, 3900(300) K at 145 GPa) (Fischer et al. 2012, 2013), the
composition of hcp alloy that coexists with B2 alloy nearly approaches pure iron. This result is
consistent with ab initio studies, which find that hcp is the stable phase of iron at inner core
conditions but that there is a small energy difference between hcp and bcc structures, with bcc
stabilized by the addition of silicon (Vočadlo et al. 2003). At these pressures, eutectic melting
from an hcp+B2 mixture is likely over a wide range of silicon contents in Fe–Si alloys,
extending down to low silicon contents. The slopes we find for this phase boundary (Figure 4)
are slightly shallower than the boundaries reported in Fischer et al. (2013), whose slopes were
less well constrained. This result extends the stability of the B2 structure to lower silicon
contents at high temperatures, which implies that partial melting of Fe–Si alloys can produce a
substantial density contrast between the melt and coexisting Si-poor metal. This concept is
illustrated schematically in Figure S1. A 4.5–7% density contrast is observed at Earth’s inner
core boundary (Dziewonski and Anderson 1981; Kennett et al. 1995; Masters and Gubbins
2003), which represents the point at which solid Fe-rich alloy crystallizes from a metallic melt.
At one bar the melting loop in the Fe–Si system is too narrow to explain such a large density
contrast, but the much larger compositional loop at high pressures, shown in this study and in
Fischer et al. (2013), supports the possibility that silicon could be the major light element in the
core, consistent with the seismological constraints.

Calculation of the Fe–Si phase diagram in the inner core

Since our parameterization of the relationship between volume, temperature, c/a ratio,
and silicon content spans a large P-T range up to >400 GPa and >5900 K, it can be used to
calculate phase boundaries at inner core conditions. The inner core is at pressures of 329–364
GPa (Dziewonski and Anderson 1981) and is thought to be nearly isothermal (e.g., Pozzo et al.
2014). The temperature of the inner core is anchored at the inner core–outer core boundary
(ICB), which is at the liquidus of the core’s Fe-rich alloy. An extrapolation of recent results on
the melting of pure Fe suggest that it melts at ~6200 K at 329 GPa (Anzellini et al. 2013). At
pressures of 50–140 GPa, we observe a ~200 K melting point depression in Fe–Si alloys relative
to pure Fe (Fischer et al. 2012, 2013). Consequently, we use here an estimated ICB temperature
of ~6000 K for an Fe–Si core. This is an approximate lower bound on the ICB temperature of a
postulated Fe–Si core, because the melting data in the Fe–Si system are for the solidus and the
inner core is crystallizing along the liquidus.

To determine the phase diagram of an Fe–Si core, we first calculated the hcp to hcp+B2
phase boundary based on experimental data from thirteen heating cycles ranging from 45 to 200
GPa. We used $\alpha$ $\beta$ ratios from Fischer et al. (2012, 2014), following the method described above
and illustrated in Figure 4 at two different pressures. We then performed a weighted linear fit to
these results, describing the relationship between pressure, temperature, and silicon content along
the phase boundary. The resultant hcp to hcp+B2 boundary in the Fe–Si system is described by
the equation:

$$\text{wt\% Si} = 16.15 - 0.00555 \times T + 0.0520 \times P$$

where temperature $T$ is in Kelvin, pressure $P$ is in GPa, and silicon content is in weight percent.

A term describing the pressure dependence of the slope (constant $\times P \times T$) was not used, as it was
found to be statistically insignificant at the 90% confidence level. The variance-covariance
matrix describing this fit is shown in Supplemental Table S3. The rms misfit between the
calculated silicon content (from Equation 1) and the fit described by Equation 2 is 2.8 wt\% Si,
comparable to our estimated (rms) uncertainty on the silicon content calculation. Phase
boundaries calculated from Equation 2 are shown in Figure 4, illustrating compatibility with the
observed phase boundary crossings in Fe–9Si. The hcp+B2 mixture is stabilized by increasing
temperature, increasing silicon content, or decreasing pressure. This equation indicates that near
inner core boundary conditions (329 GPa, 6000 K), an Fe–Si alloy containing greater than
0.0(21) wt\% Si will be stable as a two-phase hcp+B2 mixture. For a temperature of 5500 K, an
Fe–Si alloy containing greater than 2.7(19) wt\% Si will be an hcp+B2 mixture. The effects of
nickel on this phase boundary remain uncertain.
Figure 5 illustrates a projection of this result to inner core conditions, for an inner core temperature of 5500 K or 6000 K. At 6000 K, an Fe–Ni–Si alloy containing 6.0(7) wt% Si would match the inner core’s density, based on extrapolating the equation of state of Fischer et al. (2014) and correcting for a Ni/Fe atomic ratio of 0.058 (McDonough 2003), consistent with the findings of Tateno et al. (2015). Figure 5 shows that near modern ICB conditions, the stable structure of an Fe–Si alloy with 6 wt% Si should be an hcp+B2 mixture. This phase boundary shifts to increasing silicon content with decreasing temperature, but an hcp+B2 mixture should be stable for Fe–6Si at 329 GPa for temperatures above ~4900 K.

This fit predicts a phase transition in Fe–9Si at 329 GPa and 4350(300) K. This temperature falls intermediate between extrapolations of phase boundaries from Fischer et al. (2013) and earlier studies (Lin et al. 2009; Kuwayama et al. 2009). A recent study (Tateno et al. 2015) measured the hcp/hcp+B2 phase boundary in Fe–9Si to over 400 GPa. Their phase boundary implies a transition temperature of ~4600 K at 329 GPa, in agreement with the findings of this study within uncertainty.

Implications

The axial \( c/a \) ratio in iron and Fe–Ni–Si alloys is sensitive to volume, temperature, and composition, and it has been parameterized here as a function of these variables based on a meta-analysis of experimental studies spanning a large range of pressures and temperatures. The axial ratio increases with volume, temperature, silicon content, and nickel content. The parameterization of the axial ratio as a function of these variables can be used to calculate the composition of an hcp Fe–Si alloy if its \( c/a \) ratio, temperature, and volume (or pressure) are known. Though not a substitute for direct compositional measurements, this method offers a new
application of equation of state data, a means to estimate the composition of the high P-T Fe–Si phase in situ, and a mechanism for filling in details of phase diagrams, allowing for more robust extrapolations of phase relations in pressure and temperature.

This parameterization allows predictions of the c/a ratio of an Fe–Ni–Si alloy of specified composition at inner core conditions, which are necessary to understand the elastic anisotropy of the hcp alloy (Gannarelli et al. 2005). The experiment-based parameterization presented here can inform future ab initio work relating elastic constants to the c/a ratio, for a better understanding of how seismic anisotropy varies with composition in hcp Fe–Ni–Si alloys.

The stability field of an hcp+B2 mixture in Fe–Si alloys at 125 and 145 GPa extends almost to pure iron at high temperatures. This large compositional loop is consistent with silicon being the light element in the Earth’s core, based on seismological observations of a large density contrast between the inner and outer core (Dziewonski and Anderson 1981; Kennett et al. 1995; Masters and Gubbins 2003). It also implies eutectic melting in the Fe–Si system over a large compositional range.

Our calculations of the Fe–Si phase diagram suggest that if silicon is an important part of the core’s light element component, then a two-phase hcp+B2 mixture may be stable at inner core conditions. In a two component system, there can only be two stable phases at the ICB (hcp+melt), in which case the hcp structure would be stable in the inner core, but in a ternary or higher order system with additional light element(s) present, hcp and B2 phases could co-crystallizing in the Earth’s inner core along a cotectic. (In this case, the density contrast at the ICB would be attributable mostly to the additional light element(s).) As the inner core grew, it may have crystallized different compositions containing different relative amounts of hcp and B2 Fe–Si-rich alloy. The hcp and B2 phases have different anisotropies, so this may lead to
variations of anisotropy with depth. The ab initio simulations of Belonoshko et al. (2008) indicate a much stronger anisotropy in bcc Fe than in hcp Fe, though results of other studies (e.g., Tsuchiya and Fujibuchi 2009) are in conflict with this low anisotropy of hcp Fe. Future studies on the anisotropy of bcc-like Fe-alloys at inner core conditions are needed to clarify this issue. Similarly, it is possible that the proposed inner core translation (Alboussière et al. 2010) could cause hemispherical variations in anisotropy due to variations in phase proportion. In addition to hemispherical and radial variations in phase proportions, the hcp phase will have a different \( c/a \) ratio based on its composition and volume, with its anisotropy decreasing as the \( c/a \) ratio approaches its ideal value of 1.633. The combination of a two-phase mixture in the inner core and variations in the \( c/a \) ratio of hcp phases may help explain the observed seismic anisotropy patterns in Earth’s core.

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**Figure Captions:**


Figure 2: Measured c/a axial ratio of Fe–9Si compared to that of pure iron as a function of temperature. Data on Fe–9Si (grey symbols) are from a heating cycle at ~145 GPa (Fischer et al. 2014). Data on pure iron (black symbols) span ~130–155 GPa. Iron data come from a variety of studies (Anzellini et al. 2013; Boehler et al. 2008; Fischer et al. 2011; Tateno et al. 2010), with a single heating cycle shown from each study. Lines are calculated for 145 GPa (based on equations of state of Dewaele et al. (2006) and Fischer et al. (2014)) from Equation 1 (solid line: Fe–9Si; dashed line: pure iron). The trends of the data are parallel, indicating that the c/a ratios of Fe–9Si and pure iron have similar temperature dependences at this pressure.

Figure 3: Residuals to Equation 1. Symbols are as in Figure 1. A: Pure hcp iron. B: Fe–10Ni alloy. C: Fe–9Si alloy. All data are color-coded by temperature according to the legend.
Figure 4: Phase boundaries in the Fe–FeSi system calculated using the c/a ratio of intermediate alloys at A: ~125 GPa and B: ~145 GPa. Each data point represents an observation of coexisting hcp and B2 structures in either Fe–9Si (blue diamonds) or Fe–16Si (orange open circles). Data are from Fischer et al. (2012, 2014). The c/a ratio of the hcp phase was used to calculate its composition along the phase boundary using Equation 1. Black crosses indicate upper and lower bounds on the transition for Fe–9Si based on in situ X-ray diffraction measurements (Fischer et al. 2013). Solid black lines are phase boundaries calculated from Equation 2; dashed lines are 95% confidence intervals. Error bars in composition are a root mean square misfit.

Figure 5: Phase diagram of Fe–Si alloys in the inner core, calculated using Equation 2 and shown in pressure-composition space at a fixed temperature of 6000 K (black lines) or 5500 K (grey lines). Solid lines: hcp to hcp+B2 phase boundary. Dashed lines: amount of silicon needed to match the inner core’s density at these conditions (Fischer et al. 2014). Yellow bands indicate uncertainties at 6000 K. If silicon is the core’s dominant light element, the inner core may be a mixture of hcp and B2 phases.

Supplemental Table Captions:

Table S1: Compilation of c/a ratios from 15 studies, as a function of temperature, volume, lattice parameter a, and mole fractions of Ni and Si. Uncertainties on the c/a ratio are listed where available.
Table S2: Variance-covariance matrix describing the fit of $c/a$ as a function of $T$, $V$, and composition (Equation 1), with $T$ in Kelvin, $V$ in cm$^3$/mol, and $X_{Si}$ and $X_{M}$ in mole fraction. Diagonal terms describe the variance in each coefficient, while off-diagonal terms describe the covariance between terms. Matrix is symmetric by definition.

Table S3: Variance-covariance matrix describing the silicon content along the hcp to hcp$+\text{B2}$ phase boundary in the Fe–Si system as a function of $T$ and $P$ (Equation 2), with silicon content in weight percent, $T$ in Kelvin, and $P$ in GPa. Diagonal terms describe the variance in each coefficient, while off-diagonal terms describe the covariance between terms. Matrix is symmetric by definition.

Supplemental Figure Caption:

Figure S1: Schematic temperature-composition phase diagrams in the Fe–Si system. A: A wide hcp$+\text{B2}$ two-phase field (red arrow), as shown in this study, allows for a larger compositional contrast (blue arrow) between coexisting solid and melt at inner core boundary pressures. This makes it possible for a Si-rich core to be compatible with seismic observations of a large density contrast between the inner and outer core (e.g., Masters and Gubbins 2003). B: In contrast, a narrow two-phase field would preclude the possibility of a large compositional contrast between coexisting solid and melt in the Fe–Si system, making silicon a less viable candidate for the core’s dominant light element.
Figure 1

A:

B:

C:

Fe

Fe–10Ni

Fe–9Si

Volume (cm$^3$/mol)

c/a ratio

300–800 K
800–1300 K
1300–1800 K
1800–2300 K
2300–2800 K
2800–3300 K
3300–3800 K
3800–4300 K
4300–4800 K
>4800 K
Figure 2
Figure 3

A:

Fe

B:

Fe–10Ni

C:

Fe–9Si

Volume (cm³/mol)
Figure 4

A:

B:

Temperature (K)

125 GPa

145 GPa

wt% silicon

hcp + B2

Fe-9Si

Fe-16Si

hcp