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Research Paper

The effect of nitrogen on the compressibility and conductivity of iron at high pressure



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ARTICLE INFO

Keywords: High pressure Iron nitrides Synchrotron X-ray diffraction Electrical conductivity

ABSTRACT

Although nitrogen in the Earth's interior has attracted significant attention recently, it remains the most enigmatic of the light elements in the Earth's core. In this work, synchrotron X-ray diffraction (XRD) and electrical conductivity experiments were conducted on iron nitrides (Fe_2N and Fe_4N) in diamond anvil cells (DACs) up to about 70 GPa at ambient temperature. These results show that iron nitrides are stable up to at least 70 GPa. From the equation of state (EOS) parameters, iron nitrides are more compressible than iron carbides. Moreover, using the van der Pauw method and Wiedemann-Franz law, the electrical and thermal conductivity of samples were determined to be much lower than that of iron carbides. The conductivities of Fe_2N and Fe_4N were similar at 20–70 GPa, suggesting no evident effects by varying the N stoichiometries in iron nitrides. Iron nitrides are less dense and conductive but more compressible than carbides at 0–70 GPa. This study indicates that less nitrogen than carbon can explain geophysical phenomena in the deep Earth, such as the density deficit.

1. Introduction

Nitrogen, the fifth most abundant element in the Solar System, is highly significant in life as well as to our planet (Bebout et al., 2013). Compared to other terrestrial planets like Venus and Mars, the anomalously high nitrogen content in Earth's atmosphere might be linked to the distinctive nitrogen progresses on Earth, such as life evolution and plate tectonics (Owen et al., 1977; Lécuyer et al., 2000). While N_2 is the primary constituent in our planet's atmosphere, iron nitrides are likely stored throughout all geological reservoirs in the Earth's interior. However, the storage and circulation of deep nitrogen remain hotly debated (Halliday, 2013). To date, nitrogen is one of the least studied elements under conditions relevant to the Earth's deep interior (Litasov et al., 2014).

Based on its high volatility and low concentration in some meteorites, it is largely presumed that nitrogen is only abundant on the Earth's surface but scantly in the deep Earth (Allègre et al., 2001). It is evident that negligible nitrogen concentrations (several ppm) are in some stony meteorites (Sugiura et al., 1998). On the other hand, the primary

building blocks of Earth (i.e., enstatite and carbonaceous chondrites) and many other meteorites can contain much more nitrogen ranging from a few hundred ppm to even more than 1000 ppm (Moore and Keil, 1969; Javoy, 1997). For instance, Sugiura et al. (1998) found that the amount of nitrogen in the taenite portion of iron meteorites can be up to 1 wt.%. The nitrogen content of mantle-derived diamonds from kimberlites and metamorphic rocks could be up to 1500 ppm and 1.1 wt.%, respectively (Cartigny, 2005). Notably, Fe₄N (roaldite) has been observed in some iron meteorites (Rubin, 1997). Thus, iron meteorites can hold a substantial amount of nitrogen up to at least 1 wt.%, especially under reducing conditions (Adler and Williams, 2005).

The nitrogen content in the deep Earth may be significantly underestimated. Based on a chondritic comparison and terrestrial compilation, "missing N" reservoirs exist in the deep Earth (Johnson and Goldblatt, 2015). Johnson and Goldblatt (2015) proposed that the total mass of nitrogen in the Earth's mantle is likely about 2–6 times that of the atmosphere's N $_2$ (4 × 10^{18} kg) through subduction, while the core holds $\sim 10^{20}$ kg N. In particular, iron meteorites represent core remnants from differentiated and subsequently disintegrated planetesimals and they are

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Peer-review under responsibility of China University of Geosciences (Beijing).

https://doi.org/10.1016/j.gsf.2020.04.012

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mostly analogous to the central part of our planet. Thus, a small amount of the Earth's total N may remain on the Earth's surface, and the predominant portion could be hidden deep inside the Earth (Bebout and Penniston-Dorland, 2016; Yoshioka et al., 2018). Deep nitrogen is often considered to be N2, NH3, and its protonated form NH4, which can substitute K⁺, Na⁺, and Ca²⁺ in silicate minerals (Eugster and Munoz, 1966; Watenphul et al., 2010; Mikhail and Sverjensky, 2014; Sokol et al., 2017b). In subduction zones, those forms of nitrogen would be mostly recycled back to the surface by slab-derived fluids, while a small portion could be delivered further into the deep mantle (Chen et al., 2019). However, these previous studies barely took into account the reaction of nitrogen with iron or the existence of iron nitrides in diamond inclusions. Due to their siderophile nature at high-pressure and high-temperature environments, iron nitrides are plausibly one of nitrogen's most common forms in the deep Earth (Johnson and Goldblatt, 2015; Bebout and Penniston-Dorland, 2016).

Thus far, Fe7N3 and Fe3N have been extensively investigated. Hexagonal close-packed Fe7N3 could be stable up to at least 3000 K at 135 GPa (Adler and Williams, 2005; Kusakabe et al., 2019). Moreover, ab initio calculations predicted that Fe7N3 is dynamically and thermodynamically stable under the Earth's core conditions (Sagatov et al., 2019). In particular, β -Fe₇N₃ is structurally identical or very close to Fe₇C₃, including their elastic property (Minobe et al., 2015; Li et al., 2016). It is also suggested that carbon and nitrogen can mutually coexist and replace each other in the Earth's and other planetary cores (Sagatov et al., 2019). Meanwhile, another important type of iron nitride, Fe₃N, is stable to at least 30 GPa and 1473 K (Litasov et al., 2017). This phase could contain up to 2.5 wt.% C in equilibrium with C-rich melt, and the C-rich ε -Fe₃N nitride may be stable in local N-rich mantle domains, formed by redox freezing of N-rich carbonatite melts in the reduced mantle (Sokol et al., 2017a). Other iron nitrides such as Fe₂N and Fe₄N were primarily studied for their phase equilibria at only ambient pressure (Jacobs et al., 1995; Tessier et al., 2000). However, little is known about the high-pressure behavior of meteorite nitrogen components Fe₂N and Fe₄N (Nielsen and Buchwald, 1981; Sugiura, 1998). Therefore, it is indispensable to study the compressibility and conductivity properties of Fe₂N and Fe₄N at high pressure, allowing us a better understanding of the nature and role of these potential deep-nitrogen hosts in the Earth's deep interior (Adler and Williams, 2005; Kaminsky and Wirth, 2017).

In this study, the high-pressure stability and P-V EOS of Fe₂N and Fe₄N were determined using DACs coupled with synchrotron XRD. Further, conductivity measurements were carried on Fe₂N and Fe₄N samples up to 70 GPa at room temperature using the van der Pauw four-probe method. We found that both Fe₂N and Fe₄N significantly affect the geophysical properties of iron. Our results on Fe₂N and Fe₄N offer new insights into the evolution of iron nitrides with pressure and have been further applied to decipher the role of iron nitrides in the storage and circulation of deep nitrogen.

2. Materials and methods

Iron nitride powder samples were purchased from Beijing Antechnology Co., Ltd (Fe₂N Lot#: RA191101 and Fe₄N Lot#: RA191301) with lattice parameters of a=4.4181 Å, b=5.4207 Å, and c=4.7809 Å for orthorhombic Fe₂N, and a=3.8023 Å for cubic Fe₄N, respectively, based on in-house XRD measurements (Fig. S1). Polycrystalline samples were gently grounded and subsequently pressed between two opposing diamond anvils to form a disk of ~10 μ m thick. The sample disk was then loaded into a DAC for further high-pressure measurements.

Synchrotron angular dispersive XRD experiments were carried out in a DAC with 200 μm culets at beamline 13BM-C of the GeoSoilEnviroCARS (GSECARS) at the Advanced Phonon Source (APS), Argonne National Laboratory (ANL). A monochromatic X-ray beam with a wavelength of 0.4340 Å was used to carry out the measurement. The incident X-ray beam was focused to a spot size of 15 $\mu m \times$ 15 μm , measured at the full width at half maximum. LaB₆ powder was used to calibrate the sample-to-detector

distance, the tilt angle, and the rotation angle of the image plate relative to the incident X-ray beam. A MAR165 CCD detector was used to record diffracted X-rays. XRD images were collected at an exposure time of 120 s at intervals of $\sim 1-3$ GPa. A tungsten gasket was pre-indented to 40 μm thick with a hole of 100 µm in diameter drilled at the center to serve as the sample chamber. Neon was loaded into the sample chamber as the pressure-transmitting medium to provide quasi-hydrostatic conditions using the high-pressure gas loading system at the Center for High Pressure Science and Technology Advanced Research (HPSTAR). The pressure was calibrated using a ruby ball and a piece of Au placed next to the sample. The ruby pressure scale was crosschecked with the equation of state of Au (Mao et al., 1986; Ye et al., 2018). The pressure uncertainty was about 1-2 GPa throughout experiments. Diffraction patterns were integrated using the Dioptas program (Prescher and Prakapenka, 2015). The lattice parameters of the samples were derived using the UnitCell program. The EOS parameters obtained by fitting the P-V data to a third-order Birch-Mur-

$$P = \frac{3}{2} K_{T0} \left[\left(\frac{V_0}{V} \right)^{\frac{7}{3}} - \left(\frac{V_0}{V} \right)^{\frac{5}{3}} \right] \times \left\{ 1 + \frac{3}{4} \left(K_0 - 4 \right) \left[\left(\frac{V_0}{V} \right)^{\frac{2}{3}} - 1 \right] \right\}$$
 (1)

where P is pressure, K_{T0} is the isothermal bulk modulus at ambient pressure, V_0 and V are the volume at ambient conditions and high pressures, respectively, and K_0 is the pressure derivative of K_{T0} at 1 bar.

High-pressure four-probe electrical resistance experiments were carried out in a symmetrical DAC with 260 μm culet, coupled with a source meter (Keithley 2400) and a nanovoltage meter (Keithley 2182A) with an output current of $\sim\!1-5$ mA at the HPSTAR. A tungsten gasket was preindented to 50 μm thick. A hole of 200 μm in diameter was drilled at the center of the indentation using laser ablation. Subsequently, the mixture of powder cubic boron nitride (cBN) and epoxy was packed into the hole and compressed to $\sim\!20$ GPa, serving as a gasket insert. Afterwards, a hole of 70 μm in diameter was drilled at the center of the cBN gasket insert to act as an insulating sample chamber. Four Pt foils were used to serve as electrodes with 4 μm thick or less. The four-probe method can determine the resistivity of an arbitrary-shaped sample with an even thickness. It can also minimize the resistivity effect from the contact resistance, and the detailed process to obtain resistance was reported by van der Pauw (1958).

3. Results and discussion

3.1. Equation of state of Fe₂N and Fe₄N

XRD patterns of Fe₂N and Fe₄N were collected up to 65.4 and 71.4 GPa, respectively, at 300 K with 1-3 GPa intervals (Fig. 1). At ambient conditions, Fe₂N has an orthorhombic structure with space group Pbcn (Laniel et al., 2018), which persisted up to the highest pressure in our studies, as can be seen in Fig. 1a. Fe₄N crystallizes in a cubic structure with space group Pm3m, consistent with previous work (Ishimatsu et al., 2003). Under pressure, Fe₄N remained stable in the pressure range of this experiment up to 71.4 GPa while we noted that the Fe₄N sample for high-pressure XRD measurements contained Fe₃N and Fe impurities (Fig. 1b), probably formed during the synthesis process of Fe₄N (Fig. S1). Adler and Williams (2005) had reported that the (200) diffraction peak of Fe₄N disappeared at ~30 GPa, which they claimed as a structural phase transition. However, we did not observe such a transition for Fe₄N in our experiments. Particularly, all diffraction peaks of Fe₄N could be clearly recognized up to the highest pressure studied here (Fig. 1b). Note that Adler and Williams (2005) used the mixture of 17% Fe₄N and 83% Fe₇N₃ as starting materials and the methanol-ethanol-water (MEW, 16 : 4 : 1) mixture as a pressure-transmitting medium. So, it can be speculated that the diffraction peaks of Fe₄N could have partly overlapped with Fe₇N₃. Another reason could be the non-hydrostatic nature of MEW at 30 GPa, as it is known to solidify by 15.2 GPa (Klotz et al., 2009).

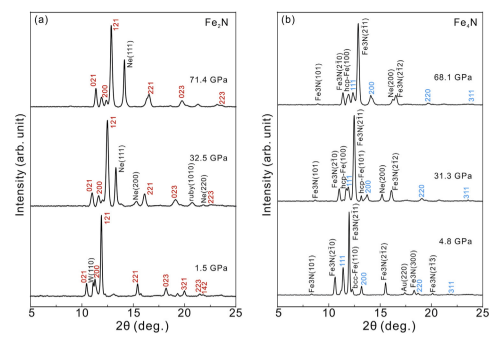


Fig. 1. Representative XRD patterns of iron nitrides Fe₂N and Fe₄N with pressure. Red labels in Fig. 1a are the peaks of Fe₂N while the blue ones in Fig. 1b are the peaks of Fe₄N. Both Au and ruby were used as pressure calibrants. The pressure medium was neon (Ne) and the gasket was tungsten (W) ($\lambda = 0.4340 \text{ Å}$).

The volume of iron nitrides (Fe₂N and Fe₄N) are plotted as a function of pressure in Fig. 2a and b, respectively. Interestingly, the volume of Fe₂N showed a change at 10–20 GPa (Fig. 2a). However, the crystal structure of Fe₂N remained the same up to 71 GPa (Fig. 1a). This could be associated with an isostructural phase transition in Fe₂N at 10–20 GPa. In Fig. 3, the c/a and b/a values of Fe₂N were plotted against pressure. Fig. 3 gives more apparent evidence of discontinuity at about 10 GPa in both c/a and b/a ratios, which can be correlated with an isostructural transition. The Raman spectra (Figs. S2 and S3) also showed that Raman shifts of modes 1 and 2 displayed subtle changes at approximately 9 GPa, while the discontinuity in the Raman mode 3 is more obvious. These results indicated an isostructural phase transition in Fe₂N around 10–20 GPa, which is likely associated with a magnetic moment collapse or electron spin transition of iron in Fe₂N.

Lattice parameters of Fe₂N and Fe₄N are plotted as a function of pressure in Fig. 4. In orthorhombic Fe₂N, the lattice parameter a showed the largest reduction of 9.2% at ~50 GPa, whereas the b and c parameters reduced by 6.4% and 5.8%, respectively. It indicates the strong anisotropic effect of pressure on the a-axis of Fe₂N. The reduction of a-parameter in cubic Fe₄N was 6.2%, suggesting a lower reduction than the

 $\mbox{Fe}_2\mbox{N},$ indicating that $\mbox{Fe}_4\mbox{N}$ may be more stable in the high-pressure area.

The EOS parameters of Fe₂N and Fe₄N were derived by fitting the P–V data to the third-order Birch-Murnaghan EOS, see Fig. 2. Fitting parameters for Fe₂N are as follows; the bulk modulus is $K_0 = 131.1(1)$ GPa with $V_0/Z = 29.6(1)$ Å³ with a fixed K' at 4 at 0–10 GPa before the phase transition, and at 20–65 GPa the bulk modulus is $K_0 = 149.5(1)$ GPa with $V_0/Z = 28.8(1)$ Å³ with a fixed K' at 4. Fitting parameters for Fe₄N are as follows; the bulk modulus is $K_0 = 144.4(1)$ GPa with $V_0/Z = 55.3(1)$ Å³ with a fixed K' at 4.

In Fig. 5, the compressibility of Fe_2N and Fe_4N was plotted and compared with reported compressibility of other iron nitrides such as Fe_7N_3 and $Fe_3N_{1.26}$ (Adler and Williams, 2005; Litasov et al., 2017). In Fig. 5, it can be inferred that the compressibility of those iron nitrides is comparable. Furthermore, in Fig. 5, the compressibility of Fe_2N and Fe_4N is also compared with the compressibility of hcp-Fe and iron carbides such as Fe_7C_3 , Fe_2C , and Fe_3C (Litasov et al., 2015). Iron nitrides are generally more compressible than hcp-Fe and the iron carbides, indicating the relatively strong effect of nitrogen on the enhancement of the compressibility of iron compared to carbon.

The pressure-dependent densities of the $\mathrm{Fe_2N}$ and $\mathrm{Fe_4N}$ samples were

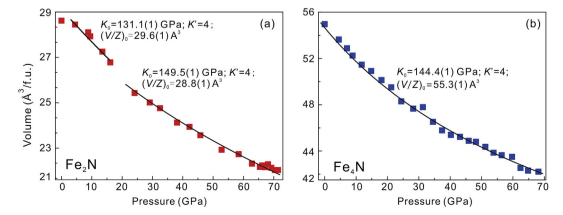


Fig. 2. The volume changes of Fe_2N and Fe_4N with pressure. The error bars are smaller than the symbols. The third-order Birch-Murnaghan fits to the data are shown by the solid curves.

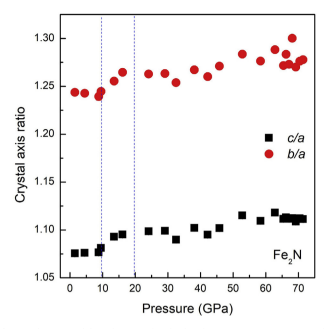


Fig. 3. The c/a and b/a changes of orthorhombic Fe₂N versus pressure. The error bars are smaller than the symbols. The dotted lines are the auxiliary lines.

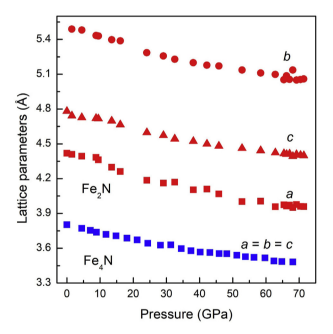


Fig. 4. The lattice parameters of Fe_2N and Fe_4N as a function of pressure. The error bars are smaller than the symbols.

plotted in Fig. 6. The densities of hcp-Fe, Fe $_3$ C, Fe $_7$ C $_3$, and Fe $_2$ C are also plotted in Fig. 6. The densities of both iron nitrides are comparable. The effect of the nitrogen content on the density of iron-based nitrides seems to be constant at this stoichiometric range. The densities of iron nitrides were lower than those iron carbides (Fe $_3$ C, Fe $_7$ C $_3$, and Fe $_2$ C) (Gao et al., 2008; Mookherjee, 2011; Nakajima et al., 2011; Chen et al., 2014; Litasov et al., 2015; Prescher et al., 2015). Thus, like compressibility, nitrogen can more significantly affect the density of iron compared to carbon (Mao et al., 1990). The nitrogen element itself plays an important role in affecting the properties of iron rather than its content per iron atom. We also speculate that compared to carbon, lower nitrogen content can explain the \sim 10% density deficit of the outer liquid core and the 3%—

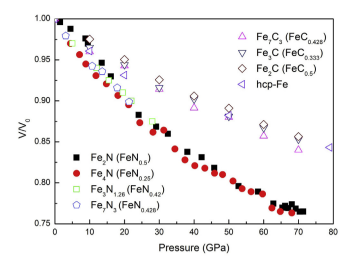


Fig. 5. The compression behaviour of Fe $_2$ N and Fe $_4$ N in comparison with other iron nitrides and carbide. The error bars are smaller than the symbols. Solid squares: Fe $_2$ N (this study); solid circles: Fe $_4$ N (this study); open squares: Fe $_3$ N_{1.26} (Litasov et al., 2017); open pentagon: Fe $_7$ N₃ (Adler et al., 2005); open triangles, inverted triangles, diamonds and oblique triangles: Fe $_7$ C₃, Fe $_3$ C, Fe $_2$ C and hcp-Fe, respectively (Litasov et al., 2015).

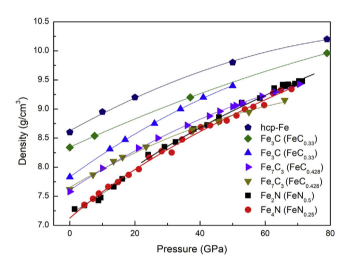


Fig. 6. The density of iron nitrides and carbides as a function of pressure. The solid curves represent fits to the data and extrapolations on the basis of the Birch-Murnaghan EOS. The error bars are smaller than the symbols. Squares: Fe_2N (this study); circles: Fe_4N (this study); diamonds: Fe_3C (Mookherjee et al., 2011); triangles: Fe_3C (Gao et al., 2008); oblique triangles: Fe_7C_3 (Prescher et al., 2015); inverted triangles: Fe_7C_3 (Chen et al., 2014); and pentagon: Fe_7C_3 (Mao et al., 1990).

5% density deficit of the solid inner core in comparison to the experimentally-derived density of iron with seismological data.

3.2. Electrical and thermal conductivity of Fe₂N at high pressure

The electrical conductivity of Fe₂N was plotted against pressure in Fig. 7. The conductivities of iron nitrides were measured to be as low as $2-3 \times 10^3$ S m⁻¹ at ambient pressure, which is consistent with some previous reports (Jiang and Jiang, 2019; Tao et al., 2019). Further, it is observed that the electrical conductivity of Fe₂N decreased at 0–10 GPa and increased after 10 GPa. This observation is consistent with volume discontinuity observed for Fe₂N in high-pressure XRD studies (Fig. 2a).

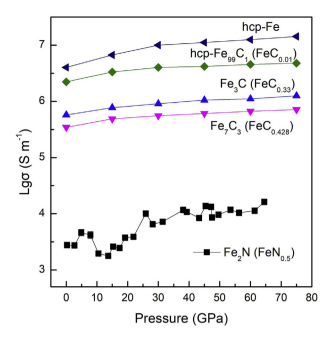


Fig. 7. The electrical conductivity of Fe_2N with pressure. The error bars are smaller than the symbols. Squares: Fe_2N (this study); inverted triangles, triangles, diamonds, and oblique triangles: Fe_7C_3 , Fe_3C , $hcp-Fe_{99}C_1$, and hcp-Fe, respectively (Zhang et al., 2018).

In Fig. 7, we also compare the conductivity of Fe₂N with the reported electrical conductivity of hcp-Fe, Fe₇C₃, Fe₃C, and Fe₉₉C₁ (Zhang et al., 2018). These results indicate that the nitrogen can have a more significant effect on iron than carbon. Nitrogen shows a much stronger alloying effect on the electrical conductivity of iron than carbon. Further, the conductivity of Fe₂N increased gradually with pressure. With extrapolation to higher pressure, the conductivity of Fe₂N would be close to other iron-carbon alloys, which suggest the strong effect of pressure on the conductivity of iron nitrides compared to other iron alloys (Zhang et al., 2018). Pressure also has a significant effect on the solubility of nitrogen in iron (Mysen, 2019).

Electronic contribution plays a key role in the thermal and electrical conductivity of metals. Hence, the thermal conductivity of metals can be estimated from its electrical conductivity, using the Wiedemann-Franz law (Anzellini et al., 2013): $k_e = L\sigma T$. Where k_e is the thermal conductivity contributed by electrons, L is the Lorenz number, σ is the electrical conductivity, T is the temperature, and L is usually adopted by the ideal value ($L_0 = 2.44 \times 10^{-8}$ W Ω K⁻²) at high pressure and temperature (Gomi and Hirose, 2015; Ohta et al., 2016). The thermal conductivity of Fe₂N is calculated using the same approach and is plotted in Fig. 8. Similar to electrical conductivity (Fig. 7), nitrogen can significantly decrease the thermal conductivity of iron, and like other physical properties, the nitrogen element also has a significant influence on the thermodynamic properties of iron compared to carbon.

3.3. Iron nitrides and carbides in the deep earth

As substantial studies have been carried out on nitrogen in the deep Earth, the decision to rule out the nitrogen element as a light element in the deep mantle and core seems an arbitrary choice. Our XRD results show that both the mainstream iron nitrides Fe_2N and Fe_4N can be thermodynamically stable at high pressure. Compared to carbon in iron phases under high pressure, nitrogen showed a stronger effect on the geophysical properties of iron. Nitrogen can effectively decrease the density and the electrical and thermal conductivity of iron with relatively lower content. Thus, nitrogen could coexist with other light elements in the deep Earth.

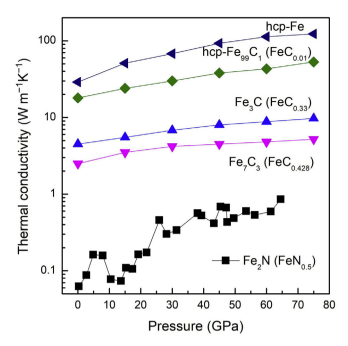


Fig. 8. The pressure-dependent thermal conductivity of Fe_2N . The data were converted from relative electrical conductivity using Wiedemann-Franz law at 300 K. The symbols are the same as Fig. 7. The error bars are smaller than the symbols.

Otherwise, it is plausible that carbon plays an important role in the deep Earth with iron. We compared Fe₂N and Fe₄N with different iron carbides and found they are interestingly similar to each other, further proving that iron nitrides can be solid solutions in iron carbides at high pressure (Schnepp et al., 2015; Litasov et al., 2016). This fact may have potentially significant implications that the carbon content is lower than previous predictions in the deep Earth with nitrogen to achieve the goal of seismological features observed in the Earth's lower mantle and core. Iron-nitride exhibited higher stability at high pressure. Thus, iron-carbide can be replaced by iron-nitride through the reaction of N-bearing fluid with iron-carbide due to their very close structural similarity, and the N, itself, is siderophile under reducing conditions (Johnson and Goldblatt, 2015; Litasov et al., 2017). Actually, previous studies about inclusions in diamonds have revealed that up to 20 mol.% carbon can be replaced by nitrogen in iron-carbides (Kaminsky et al., 2015; Smith et al., 2016).

Many studies have proved that nitrogen content is greatly underestimated in the deep Earth (Johnson and Goldblatt, 2015; Yoshioka et al., 2018). Through the Earth's history, nitrogen stored in the mantle and core may have increased based on the deduction for the difference in nitrogen input and output (Mysen, 2019). Subduction zones can slide into the mantle, even the core (Mao et al., 2017), and have proven to be major venues for nitrogen flux into and out of the Earth (Busigny et al., 2011). Moreover, lots of previous studies indicate a variety of minerals can be effective carriers to bringing nitrogen into the deep Earth through subduction zones such as clay minerals (Cedeño et al., 2019), mica (Sokol et al., 2018) and feldspar (Watenphul et al., 2009). Our experimental results clearly show the significant effect of nitrogen on the compressibility, density, electrical, and thermal conductivity of iron. The effect of carbon content on iron was noticeable, while the effect of nitrogen was subtle. However, the nitrogen element has a more remarkable influence on the thermodynamic properties of iron compared to the carbon element at high pressure. Nitrogen could exist in the deep Earth as many forms of iron nitrides or solid solutions of iron carbides. Therefore, it is desirable to study the behaviors of nitrogen and its derivatives in the Earth's deep interior further.

4. Conclusions

In summary, synchrotron XRD and electrical conductivity studies were carried out for Fe₂N and Fe₄N in DACs up to \sim 70 GPa at ambient temperature. XRD and electrical conductivity experiments confirmed the isostructural phase transition in Fe₂N at 10–20 GPa. Fe₄N has no structural transition in the experimentally investigated pressure range. For Fe₂N EOS fitting parameters are as, $K_0 = 131.1(1)$ GPa, $V_0/Z = 29.6(1)$ Å³, K' fixed at 4 for 0–10 GPa range and $K_0 = 149.5(1)$ GPa, $V_0/Z = 28.8(1)$ Å³, K' fixed at 4 for 20–65 GPa range. For Fe₄N the EOS fitting parameters are as, $K_0 = 144.4(1)$ GPa, $V_0/Z = 55.3(1)$ Å³, K' at 4.

The density, compressibility, and electrical and thermal conductivity of iron nitrides at high pressures were compared with iron carbides. Iron nitrides displayed lower density, higher compressibility, and much lower electrical and thermal conductivity than iron carbides. Thus, nitrogen would have a more remarkable influence on the high-pressure properties of iron than carbon. Our work also shows nitrogen itself plays an important role in affecting the properties of iron rather than its content per iron atom. The present results indicate that the role that nitrogen plays in the deep Earth has been previously underestimated and should be reevaluated.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

We acknowledge the use of synchrotron X-ray diffraction at the 13BM-C of GSECARS, Advanced Photon Sciences, Argonne National Laboratory. Yukai Zhuang is supported by the China Postdoctoral Science Foundation (18NZ021-0213-216308). GeoSoilEnviroCARS is supported by the National Science Foundation- Earth Sciences (EAR – 1634415) and the Department of Energy-GeoSciences (DE-FG02-94ER14466). 13BM-C is partially supported by COMPRES under NSF Cooperative Agreement EAR -1606856. This research used resources of the Advanced Photon Source, a U.S. Department of Energy (DOE) Office of Science User Facility operated for the DOE Office of Science by Argonne National Laboratory under Contract No. DE-AC02-06CH11357.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi. org/10.1016/j.gsf.2020.04.012.

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