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Pressure-induced structural and OPENspin transitions of Fe3S⁴

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Greigite (Fe3S4), isostructural with Fe3O4 has recently attracted great scientific interests from material science to geology due to its complicated structure and electronic and magnetic configurations. Here, an investigation into the structural, magnetic and electronic properties of Fe3S4 under high pressure has been conducted by first-principle calculations based on density functional theory. The results show that a first-order phase transition of Fe3S4 would occur from the inverse spinel (SP) structure to the Cr3S4 type (CS) structure at 3.4GPa, accompanied by a collapse of 9.7% in the volume, a redistribution of iron cations, and a half-metal to metal transition. In the CS-Fe3S4, Fe²⁺ located at octahedral environment firstly undergoes a transition from high-spin (HS) state to low-spin (LS) state at 8.5GPa and Fe3⁺ subsequently does at 17GPa. The Equation of State for different phases of Fe3S4 are also determined. Our results not only give some clues to explore novel materials by utilizing Fe3S4 but also shed light on the fundamental information of Fe3O4, as well as those of other SP-AB2X4 compounds.

Spinel-structured (SP) AB_2X_4 (X = O, S, Se) compounds with space group $Fd\overline{3}m$ (Z = 8) [see Supplementary Fig. S1(a)], such as $MgA1_2O_4$, FeCr₂O₄, Fe₃O₄ and FeCr₂S₄, have attracted great scientific interests from geophysics to material science. For example, the phase transition of $(Mg,Fe)_{2}SiO_{4}$ from olivine to wadsleyite, and subsequently to ringwoodite explains the seismic discontinuity in 410 km and 520 km in Earth's interior, respectively^{[1,](#page-6-0)2}. In addition, the discovery of a new class of multiferroic materials, in which ferroelectricity and ferromagnetism, as well as other unusual physical properties are compatible with a cubic spinel symmetry holds promise for new generations of functional material[s3](#page-6-2)[,4](#page-6-3). These significant physical phenomena are possibly attrib-uted to intimate coupling between structural, electronic and magnetic properties in the material^{[5,](#page-6-4)6}.

High pressure, as one important thermodynamic parameter, can strongly affect a material's structures and electronic and magnetic configurations, which inspires a renewed interest in the fundamentally physical and chemical properties of spinel-structured AB_2X_4 compounds. Besides dissociating into the assemblage of simpler compounds, three orthorhombic phases of CaTi₂O₄-type (space group: *Bbmm*, Z=4, CT), CaMn₂O₄-type (space group: *Pbcm*, $Z = 4$, CM) and CaFe₂O₄-type (space group: *Pnma*, $Z = 4$, CF) structures have been identified by both experiments and theoretical calculations as high-pressure polymorphs of oxide spinels with the initial chemical formula[7–12](#page-6-6). In addition, some of them can experience the *I*41/*amd* structure, a subgroup of the SP structure, before transforming into the denser orthorhombic structures^{13,[14](#page-6-8)}. While for thiospinels and selenospinels, the phase transition to a Cr_3S_4 -type (space group: $I2/m$, $Z = 2$, CS) structure upon compression has been confirmed as well $15,16$.

In particular, the high-pressure behavior of magnetite (Fe₃O₄), an inverse spinel, is fundamentally important for understanding the oxidation state of Earth's interior. A phase transition of $Fe₃O₄$ to a monoclinic structure was observed at ~25GPa, but more details about the high-pressure structure, such as space group and atomic positions were not provided¹⁷. Later, both CM-type and CT-type structures were proposed to be candidates for the high-pressure post-spinel phase of Fe₃O₄^{[8,](#page-6-12)[9,](#page-6-13)[18](#page-6-14),[19](#page-6-15)}. However, the accurate high-pressure polymorph of Fe₃O₄ remains unclear. Additionally, the electronic and magnetic behaviors of $Fe₃O₄$ under extreme conditions are under debate. Transformation of the electron charge density from the octahedral to the tetrahedral site of $Fe₃O₄$ results in a phase transition to a normal spinel structure that was proposed at \sim 7 GPa^{[20](#page-6-16)}, but data obtained by high-pressure single crystal X-ray diffraction experiments did not support such a conclusio[n21](#page-6-17). The amplitude of the Fe *K*-edge X-ray magnetic circular dichroism (XMCD) signal of $Fe₃O₄$ was observed to decrease discontinuously by 50% between ~12 and ~16 GPa, which was interpreted by a high-spin (HS) to intermediate-spin (IS) transition of Fe²⁺ on the octahedral sites^{[22](#page-6-18)}. However, Baudelet *et al.*^{[23](#page-6-19)} found neither an inverse-to-normal phase transition nor a

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spin transition in Fe₃O₄ up to ~41 GPa by XMCD experiments. On the other hand, using first-principle calculation, Ju *et al.*[24](#page-6-20) predicted a very complicated spin transition sequence from HS state to IS state and further to low-spin (LS) state of iron above 30GPa. Another calculation found that the HS state of iron was stable at least up to 45 GPa instead²⁵. Therefore, there exist significant controversies on high-pressure polymorphs and electronic and magnetic properties of compressed Fe₃O₄, which restrain our understanding of oxide buffer and magnetic field involving $Fe₃O₄$ in the lower mantle.

Since various structural, electronic and magnetic transitions of $Fe₃O₄$ can only be observed at very high pressure that are challenging for accurate detections, the high-pressure studies of compounds analogous to $Fe₃O₄$, that may experience similar phase transitions at relatively low pressure conditions, are urgent. Greigite (Fe₃S₄) is isostructural with Fe₃O₄ and can be considered as the sulfide counterpart of Fe₃O₄. Hence, Fe₃S₄ is a good candidate for investigations of the high-pressure behavior of Fe₃O₄. In addition, Fe₃S₄ has been reported to play an important role not only on hydrogen storage^{26,27}, lithium-ion batteries²⁸ and water dissociation^{29,30} but also on paleo-magnetism^{[31](#page-7-5),[32](#page-7-6)}. The saturation magnetization of Fe₃S₄ with ferrimagnetic (FIM) property was reported in a wide range of values from 1.06 to 3.35 *μB*/f.u., which was attributed to different samples used in the experiment[s33–37.](#page-7-7) However, any result on the magnetic moment is much lower than that of Fe₃O₄ (~4.0*µ_B*/f.u.), attributed to an increased degree of covalence between iron and sulfur compared to oxygen ligands or by greater delocalization of the 3*d* electrons in Fe₃S₄^{[38](#page-7-8)}. An interesting phenomenon, the Verwey transition was observed in Fe₃O₄, where the resistivity abruptly increased implying a metal-to-insulator transition by charge ordering when the temperature decreased to \sim 120 K³⁹. However, neither low-temperature magnetic nor structural transition was evident for Fe₃S₄ by high-resolution neutron powder diffraction (NPD) and polarized neutron diffraction (PND)³⁸. Later, two research groups reported opposite results about whether or not the Verwey transition exists in Fe₃S₄ by first-principle calculations^{40,41}. Generally speaking, the structural, electronic and magnetic properties of Fe₃S₄ are also controversial and more complex than we have expected. However, to the best of our knowledge, no work has been published about the high-pressure behavior of Fe₃S₄. Hence, our study aims to explore the structural stability and electronic and magnetic configurations of $Fe₃S₄$ under high pressure by first-principle calculations based on density functional theory (DFT), which could provide more fundamental details about the physical and chemical properties of Fe₃S₄ and aid to our understanding of the high-pressure behavior of Fe₃O₄.

Results

The corresponding Equation of State (EoS) parameters of different Fe₃S₄ phases are listed in [Table 1](#page-1-0). Based on those parameters, we calculate static enthalpy differences among various candidate phases as a function of pressure [\(Fig. 1](#page-2-0)). The FIM-SP phase with the lowest enthalpy is the ground state at ambient pressure in agreement with previous studies^{[38](#page-7-8),[40](#page-7-10)}. Upon compression, the CT-, CM-, CF-type Fe₃S₄ are unstable under all calculated pressure range no matter what types of magnetic structures they are, which indicates that the phase transition from the spinel structure to orthorhombic structures is energetically unfavorable up to at least 30 GPa. A structural phase transition from the SP structure to the CS structure keeping ferrimagnetic is predicted to occur at 3.4GPa and 0K. The transition point is relatively low compared with ~20 GPa in Fe₃O₄, and more similar to ~7 GPa in FeCr₂S₄^{[15](#page-6-9)}. Another two phase transitions are predicted to occur at 8.5 GPa and 17 GPa, respectively due to spin state transitions, which will be discussed later.

Figure 1. Static enthalpy differences among various candidate phases as a function of pressure. FIM, FM, AFM and NM represent ferrimagnetic, ferromagnetic, antiferromagnetic and nonmagnetic structures, respectively.

Figure 2. Calculated net magnetic moments (**a**) and sub-lattice magnetic moments per Fe of A-site and B-site (**b**) as a function of pressure. The magnetic moments of Fe on the B-site are defined as positive and therefore those of Fe on the A-site in the FIM-SP and FIM-CS phases are negative. The grey, pink, blue and yellow regions represent FIM-SP, FIM-CS, FM1-CS and FM2-CS phases, respectively. The previous data marked with red (experimental results, Exp.), blue (Exp.), green (theoretical calculations, The.) and pink (The.) points are extracted from refs [38,](#page-7-8)[40](#page-7-10) and [42](#page-7-12), respectively for comparison.

[Figure 2](#page-2-1) displays the net and individual sub-lattice magnetic moments as a function of pressure, respectively, where the available experimental and theoretical data at ambient pressure are included for comparison^{[38](#page-7-8),[40](#page-7-10),42}. The calculated sub-lattice magnetic moments of SP-type Fe₃S₄ at ambient pressure are $-3.12 \mu_B$ (m_A) and 3.30 μ_B (m_B) , indicating a ferrimagentic structure in agreement with the data obtained by NPD^{[38](#page-7-8)}. This result suggests that the simulation is well improved by the DFT+*U* method (see Supplementary text and Figure S2). The calculation also presents HS electronic configurations for Fe sites, where the effective spin in the tetrahedral site is lower than that in the octahedral one. The sub-lattice magnetic moments on both sites decrease with the increasing pressure. However, the net magnetic moment of FIM-CS $Fe₃S₄$ increases upon compression from 3.4 GPa to 8.5 GPa because the magnetic moment of Fe_A decreases more rapidly than Fe_B does. At 8.5 GPa, the ferromagnetic (FM) CS phase with LS Fe_A and HS Fe_B (named as FM1-CS) becomes energetically favorable compared with the FIM-CS phase, accounting for a significant increase in the net magnetic moment which is even higher than that at ambient pressure. Subsequently, a pressure-induced HS-LS transition of Fe_B in the FM-CS phase (named as FM2-CS) results in a large drop in the net magnetic moment above 17GPa. Under further compression, the net magnetic moment and sub-lattice magnetic moments decrease slowly without any abrupt changes up to 30GPa.

The electronic density of states (DOS) of Fe at each equivalent site in four phases (FIM-SP, FIM-CS, FM1-CS and FM2-CS) are shown in [Fig. 3](#page-3-0). In the FIM-SP phase [[Fig. 3\(a\)](#page-3-0)], both Fe_A and Fe_B are identified in HS state. The DOS in the spin-down state of Fe_B crosses the Fermi level while there is a gap of 0.2 eV in the spin-up state of Fe_A,

Figure 3. Calculated Fe DOS of each equivalent atom site of different Fe₃S₄ phases by GGA + *U* methods. (**a**) DOS of Fe in FIM-SP phase at 0GPa, (**b**) DOS of Fe in FIM-CS phase at 3.7GPa, (**c**) DOS of Fe in FM1-CS phase at 9.2GPa and (**d**) DOS of Fe in FM2-CS phase at 27.2GPa. The Fermi level is indicated by the vertical line in each DOS.

which is in agreement with similar theoretical studies^{[40](#page-7-10)}. Besides, it is commonly accepted that the DFT calculation systematically underestimates the band gaps of materials. Therefore, the half-metallic character for FIM-SP Fe₃S₄ at ambient pressure could be confirmed by our calculation. At 3.7 GPa, the DOS of the spin-up state of Fe_A on the bottom of conduction bands changes significantly, indicating a variation of local environment of Fe_A due to the SP-CS phase transition. It is obvious that the contribution of electrons of Fe to the top of valance bands in the high-pressure phase [[Fig. 3\(b\)](#page-3-0)] is more pronounced than that in the low-pressure phase [[Fig. 3\(a\)\]](#page-3-0), contributing to the metallic property of the FIM-CS phase. However, whether or not FIM-CS Fe₃S₄ becomes metallic upon compression needs further experimental verification due to the underestimation of band gaps of materials by DFT calculations as mentioned above. At 9.2 GPa [\[Fig. 3\(c\)\]](#page-3-0), the orbitals in the spin-up state of Fe_A in the FM1-CS phase mainly form the top of valance bands, which could be interpreted as the HS-LS transition of Fe_A . While the DOS in the spin-down state of Fe_B does not change significantly, suggesting that Fe_B remains the HS state under such conditions. Upon compression, a LS state of Fe_B is also observed [\[Fig. 3\(d\)](#page-3-0)]. In the meanwhile, Fe₃S₄ keeps metallic in the high-pressure CS phase throughout the calculated pressure range.

Discussion

In order to identify the valence state of iron cations in the CS structure, we have calculated the charge density on the FIM-CS (020) section at 3.7 GPa (see Supplementary Figure S3). The calculation illustrates the strong covalence between iron and sulfur due to the existence of high charge density in-between Fe-S bonds. It can be also observed that the distribution of charge around S near to Fe_A is more regular than that of S located near Fe_B (top-left corner or bottom-right corner), implying that Fe_B possesses a stronger polarization than Fe_A does. This implies that Fe_B possesses a smaller radius or higher chemical valence compared with Fe_A, either of which indicates that Fe³⁺ occupies the B-site while Fe²⁺ occupies the A-site in the CS structure. Therefore, we suggest a redistribution of iron cations between A-sites and B-sites at the SP-CS transition. In such a case, Fe2⁺ and Fe3⁺ occupy 2*a* (A-site) and 4*i* (B-site) positions, respectively, in the CS structure contrary to the low-pressure inverse spinel structure, where Fe2⁺ and half of Fe3⁺ occupy 16*d* (B-site) positions while the rest of Fe3⁺ occupy 8*a* (A-site) positions. Moreover, considering the magnetic properties of Fe₃S₄ displayed in [Fig. 2](#page-2-1), we propose that the HS-LS transitions of Fe2⁺, occupying A-site, and subsequently Fe3⁺, occupying B-site, in the high-pressure CS phase occur at 8.5 GPa and 17 GPa, respectively. The above-mentioned abrupt changes (local environment of Fe³⁺ at the SP-CS transition and spin state of iron cations) are significant and therefore could be detected by Mössbauer

Figure 4. Calculated results of bond distances and lattice parameters of different Fe₃S₄ phases as a function **of pressure.** (a) Fe_A-S bond distances and (b) Fe_B-S bond distances in FIM-SP, FIM-CS, FM1-CS and FM2-CS phases as a function of pressure. (**c**) axis and (**d**) *β* angle of FIM-CS, FM1-CS and FM2-CS phases as a function of pressure.

spectroscopy. For example, an increase of coordination number of Fe³⁺ at the SP-CS transition would lead to an abrupt increase of isomer shift. In addition, the spin transitions of iron cations would result in a discontinuous decrease of magnetic hyperfine field and isomer shift^{[43,](#page-7-13)44}. At the same time, the quadrupole splitting of Fe²⁺ might disappear while that of Fe³⁺ would increase discontinuously at the HS-LS spin transition^{44,45}. However, the accurate description of changes of hyperfine interaction parameters for $Fe₃S₄$ under high pressure requires further investigation due to the different nature of Fe-S and Fe-O bonds.

The compressions of bond distances and lattice parameters of different phases as a function of pressure are shown in [Fig. 4.](#page-4-0) The redistribution of iron cations between A-sites and B-sites leads to abrupt changes of Fe-S bonds at the SP-CS transition at 3.4 GPa [[Fig. 4\(a\) and \(b\)\]](#page-4-0). Upon compression, both Fe²⁺-S1 and Fe²⁺-S2 bonds decrease abruptly at 8.5 GPa while all Fe³⁺-S bonds decrease discontinuously at 17 GPa, corresponding to the above-mentioned HS-LS transition sequence of iron cations. It should be noted that although the Fe³⁺-S2 and $Fe³⁺-S3$ bonds change abruptly at 8.5 GPa, we do not consider these to be due to the spin transition of $Fe³⁺$. Instead, the HS-LS transition of Fe²⁺ with abrupt decrease in the radius of Fe²⁺ results in a distortion of CS struc-ture, causing some of Fe³⁺-S bonds to change discontinuously. [Figure 4\(c\)](#page-4-0) presents the fact that the compression of the CS phase is highly anisotropic, i.e., the compressibility of *c*-axis is larger than that of *a*-axis or *b*-axis. The spin transitions of Fe²⁺ and Fe³⁺ cause an abrupt reduction of *c*-axis at 8.5 GPa and 17 GPa, respectively while these transitions affect *a*-axis and *b*-axis only slightly. As shown in Figure S1(b), layers of Fe²⁺ and Fe³⁺ alternate with a cation ordering of Fe^{2+} - Fe^{3+} - Fe^{2+} - Fe^{3+} - Fe^{2+} along the *c*-axis in the CS structure. Therefore, the decrease in radius of either Fe²⁺ or Fe³⁺ could exert an influence on *c*-axis. The compression coefficients of the *c*-axis are -6.92×10^{-3} GPa⁻¹ for FIM-CS phase, -2.18×10^{-3} GPa⁻¹ for FM1-CS phase and -1.70×10^{-3} GPa⁻¹ for FM2-CS phase, indicating that the spin transitions of iron make the *c*-axis less compressible. Additionally, the *β* angle increases rapidly from 3.4GPa to 8.5GPa, illustrating a higher degree of distortion [\[Fig. 4 \(d\)\]](#page-4-0). This is in agreement with our analysis above.

[Figure 5](#page-5-0) displays the compressions of volumes of different phases as a function of pressure, and the corre-sponding parameters have been listed in [Table 1.](#page-1-0) The K_0 of FIM-SP phase is in agreement with previous calculation of 62.8 GPa⁴². A smaller *K₀* of FIM-CS phase, 46.6 GPa, compared to that of FIM-SP phase is obtained in our calculations. This relation between spinel structure and high-pressure post-spinel structure is quite typical and the intrinsic mechanism needs further exploration^{[10,](#page-6-22)15}. The K_0 of FM1-CS and FM2-CS phases increase significantly. Moreover, these three phase transitions, FIM-SP to FIM-CS, FIM-CS to FM1-CS and FM1-CS to FM2-CS, are all first-order with volume reductions of 9.7% at 3.4GPa, 5.1% at 8.5GPa and 3.1% at 17GPa, respectively.

Figure 5. Calculated results of volume per formula unit of different Fe₃S₄ phases as a function of pressure. The solid lines segments are fitted by the third-order Birch-Murnaghan Equation of State. The volume collapse of each phase transition is marked. HS and LS represent high-spin and low-spin states, respectively. The previous data marked with red (Exp.), blue (Exp.), green (The.) and pink (The.) points are extracted from refs [38](#page-7-8)[,40](#page-7-10) and [42,](#page-7-12) respectively for comparison.

An increase of coordination number of Fe³⁺ from four to six mainly contributes to the 9.1% volume collapse. Although the number of Fe³⁺ is twice as that of Fe²⁺ in the chemical formula Fe₃S₄, the volume collapse at 8.5 GPa is larger than that at 17 GPa. Both the decrease in radius of $Fe²⁺$ due to the spin transition and consequently a structural distortion most likely account for the seemingly anomalous volume reduction.

Since the radius of S^{2-} is larger compared to that of O^{2-} , the transformation from SP structure to CM-, CTor CF-type structure for many $\overline{AB_2S_4}$ compounds is energetically unfavorable. Indeed, it has been reported that many of SP-type transition-metal AB₂S₄ compounds transform into CS structure, such as $MnCr_2S_4$, FeCr₂S₄ and $CoCr_2S_4^{15,46}$ $CoCr_2S_4^{15,46}$ $CoCr_2S_4^{15,46}$. The SP-CS phase transition is reconstructive, where the sulfur sub-lattice changes from an fcc to an hcp arrangement for lower energy. It is worthwhile to mention that previous investigations on $F\in C_rS_4$ have shown that the sample is composed solely of the CS phase upon decompression to ambient pressure, indicating an irreversible phase transition¹⁵. Recent studies on $ZnCr_2Se_4$ under high pressure have confirmed a reversible SP-CS phase transition at 17 GPa by XRD, however^{[16](#page-6-10)}. Whether or not the CS phase of Fe₃S₄ can be pressure-quenched needs confirmation by experimental methods.

For high-pressure polymorph of Fe₃O₄, an occupation of Fe²⁺ in A-sites in either CM phase or CT phase is presumed. However, recent theoretical studies have shown that the total electron numbers of A-sites and B-sites remain unchanged in $Fe₃O₄$ with increasing pressure. The study therefore concluded that the high-pressure phases of Fe₃O₄ still possess the chemical formula as Fe₃O₄ at ambient conditions²⁴. As for Fe₃S₄ in our studies, the redistribution of iron cations between A-sites and B-sites occurs at the SP-CS transition. The redistributed occupations of iron cations compared to low- and high-pressure phases of Fe₃O₄ discussed above account for different magnetic properties under high pressure in terms of the spin state of iron cations. In addition to the spin transition of Fe²⁺, Ju *et al.*²⁴ observed that Fe³⁺ on both A-sites and B-sites underwent a HS-IS transition in the CM-type $Fe₃O₄$ between 50 and 60 GPa in Fe₃O₄. However, this claim is not necessarily supported by their calculated sub-lattice magnetic moments or bond distances. In particular, the calculations predicted that the net magnetic moment of Fe₃O₄ is lost at 65 GPa and recovered to a nonzero value after the CM-CT transition above 65 GPa, implying Fe₃O₄ can be magnetized in the deep lower mantle. This was caused by the fact that Fe³⁺ on the A-site and B-site were not identical. Specifically, Fe^{3+} on the B-site possesses a lower IS-LS transition pressure than Fe^{3+} on the A-site does. In Fe3S4, both HS and LS state of iron occur under different conditions while the IS state of $Fe²⁺$ or $Fe³⁺$ is not observed in our calculations. Although the net magnetic moment changes differently as a function of pressure with respect to the results obtained by Ju *et al.*[24](#page-6-20) due to different occupations of iron cations, the intrinsic spin transition sequence, where Fe^{2+} possesses a lower spin transition pressure and Fe^{3+} possesses a higher spin transition pressure, is basically in agreement with the exception for the absence of an IS state of iron cations in our calculations. The present studies on Fe₃S₄, a sulphide counterpart of Fe₃O₄, may contribute to provide details towards a more comprehensive view towards Fe₃O₄. Additionally, compared to previous studies on Fe₃O₄, our calculations predict a FM1-CS phase (CS-typeFe^{LS}(Fe^{HS})₂S₄) with a significantly positive jump in the net magnetic moment at the transition, which is even higher than that of SP-type Fe_3S_4 at ambient pressure. If such a prediction could be confirmed by experiments, it would be also of interest for exploring new magnetic materials.

Conclusions

In conclusion, the high-pressure behavior of $Fe₃S₄$ has been studied based on first-principle density functional calculations. A first-order phase transition from the SP-type structure to a monoclinic CS-type structure, rather than to an orthorhombic structure, at 3.4 GPa has been established. This reconstructive transition leads to an increase in the coordination number of Fe^{3+} from four to six with a 9.7% volume reduction and a metallic nature of the high-pressure phase of Fe_{3S4}. According to the calculated charge density of CS-Fe_{3S4} as well as bond distances of Fe with respect to S as a function of pressure, a redistribution of iron cations with occupation of $Fe²⁺$ on A-sites and all of Fe³⁺ on B-sites in the CS-type structure at 3.4 GPa is observed, which has a significant effect on the magnetic properties of high-pressure $Fe₃S₄$ upon further compression. The calculations predict HS-LS transitions of Fe^{2+} and subsequently Fe^{3+} , where an abrupt decrease in magnetic moments at 8.5 GPa and 17 GPa result in a volume collapse of 5.1% and 3.1%, respectively. Finally, the Equation of State for different phases of $Fe₃S₄$ are also determined.

Methods

Five candidate structures (SP, CT, CM, CF and CS) described in the literature were considered for Fe₃S₄ in our simulation. First-principle calculations based on density functional theory were performed using the projected augmented wave (PAW) method implemented in Vienna *ab-initio* simulation package (VASP[\)47–49.](#page-7-17) The Perdew-Burke-Ernzerhof (PBE) version of the generalized gradient approximations (GGA) was selected to treat the exchange correlation potential⁵⁰. The kinetic energy cut-off was set to 600 eV. The energy convergence criterion for the electronic self-consistent calculation was 10[−]⁴ eV. A DFT+ *U* method was introduced in the simulation to correctly describe the strong electronic correlation⁵¹. $U=2.5 \text{ eV}$ and $J=1 \text{ eV}$ were applied into all iron in all GGA+ *U* calculations (see Supplementary text and Figure S2). The spin-polarization of iron without spin-orbit coupling was included in calculations to obtain the accurate cell parameters and energies. For SP, CT, CM and CF phases, ferrimagnetic (FIM), ferromagnetic (FM) and anti-ferromagnetic (AFM) structures were included. For CS phase, FIM, FM, non-magnetic (NM), and three AFM (AFM1, AFM2 and AFM3) structures were considered in the calculation. The magnetic arrangement of AFM3 structure was used from ref. [52,](#page-7-20) thus a $2 \times 1 \times 2$ supercell was constructed. Computations were performed at various volumes for each crystalline phase. The atomic positions, unit-cell parameters and individual magnetic moments were allowed to relax at each given volume to obtain the minimum total energy. The Monkhorst-Pack scheme was used for Brillouin zone sampling. The k-points grids were set as $4 \times 4 \times 4$ for SP phase, $2 \times 2 \times 8$ for CT phase, $8 \times 2 \times 2$ for CM phase, $2 \times 8 \times 2$ for CF phase and $4 \times 8 \times 2$ for CS phase ($2 \times 8 \times 1$ for AFM3 CS phase). The DOS were obtained by the static calculation, utilizing the tetrahedral smearing method with Blöchl corrections. The k-points grids were set as $8 \times 8 \times 8$ for SP phase and $12 \times 16 \times 8$ for CS phase in DOS calculation.

Once the minimum total energies of each phase were obtained at different volumes, they were fitted to the third-order Birch-Murnaghan EoS to compute the volume per formula unit (V_0) , bulk modulus (K_0) , its pressure derivative (K_0) and energy (E_0) at zero pressure^{[53](#page-7-21),[54](#page-7-22)}. In addition, the enthalpy $(H = E + PV)$ of each phase was compared with each other to identify the most stable structure under the given pressure.

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Author Contributions

X.W. conceived the research. S.H. and D.K performed the theoretical calculations. S.H., D.K., J.N. and S.Q. analyzed the data. S.H., X.W. and S.Q. wrote the manuscript. All authors reviewed the manuscript.

Additional Information

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