Pressure-induced high-spin/low-spin disproportionated state in the Mott insulator FeBO₃

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The pressure-induced Mott insulator-to-metal transitions are often accompanied by a collapse of magnetic interactions associated with delocalization of 3d electrons and high-spin to low-spin (HS-LS) state transition. Here, we address a long-standing controversy regarding the high-pressure behavior of an archetypal Mott insulator FeBO₃ and show the insufficiency of a standard theoretical approach assuming a conventional HS-LS transition for the description of the electronic properties of the Mott insulators at high pressures. Using high-resolution x-ray diffraction measurements supplemented by Mössbauer spectroscopy up to pressures ~ 150 GPa, we document an unusual electronic state characterized by a “mixed” HS/LS state with a stable abundance ratio realized in the R3c crystal structure with a single Fe site within a wide pressure range of ~ 50–106 GPa. Our results imply an unconventional cooperative (and probably dynamical) nature of the ordering of the HS/LS Fe sites randomly distributed over the lattice, resulting in frustration of magnetic moments.

Iron borate, FeBO₃, is one of a few materials that are transparent in a broad range of visible light and have a spontaneous magnetization at room temperature, which makes it attractive in applications for visible or x-ray light magneto-optical devices⁹. It is a canted antiferromagnet with the Néel temperature TN ~ 348 K and weak...
ferromagnetism. Optical spectroscopy shows that FeBO₃ is a Mott insulator with a large energy gap of ~2.9 eV (18 and ref. therein). Formally, FeBO₃ can be considered as a part of a more general FeOX family (e.g., FeFeO₃, FeGaO₃, etc.), with ferric Fe³⁺ ion playing a major role in determining the electronic and magnetic properties of FeXO₃. Recent extensive high-pressure studies of FeBO₃ reveal that, similar to many other ferrites, in the vicinity of ~50 GPa it undergoes an isostress phase transition corroborating with a significant reduction of the lattice volume and coinciding with an abrupt magnetic collapse and a Mott insulator-to-semiconductor transition. However, we notice that FeBO₃ shows some very specific features following an affirmed HS-LS transition at ~50 GPa. Most notably, it exhibits apparently resilient non-metallic behavior above 100 GPa. This behavior is different, e.g., to FeGaO₃ and FeO₃, which exhibit the classical band-width controlled Mott transition at ~50–106 GPa. Our observations are unexpected given the presence of only a single Fe³⁺ structural position in the crystal structure of FeBO₃. This behavior clearly distinguishes the behavior in FeBO₃ from a “conventional” spin-state crossover observed in other ferric systems. We propose a model explaining this unusual electronic state based on a complex interplay between the spin and lattice degrees of freedom.

Results

X-ray diffraction. We perform four independent SC-XRD experiments, which are mutually consistent for their measured pressure range. The exceptional quality of the data is reflected in the parameters of fit (see Supplementary information). In particular, we found that at ambient pressure FeBO₃ adopts the rhombohedral R3c crystal structure (see inset Fig. 2b and Supplementary Fig. S2), in agreement with previous PWD and SC XRD [20 and ref. therein]. The R3c structure is conserved at least up to ~105 GPa (Figs. 1, 2). At about 50 GPa we observe a doubling of the reflections within the diffraction patterns attributed to the onset of the high-pressure (HP) phase characterized by the same space group (R3c) but with significantly reduced unit-cell volume and lattice parameters (Supplementary Fig. S1). Our SC-XRD data confirm that multiple domains of the HP phase grow on top of the HS low-pressure (LP) state grains preserving the same orientation. Here we extend the previous PWD-XRD studies and shed new light on the process of phase transformations in FeBO₃. We observe a finite range (~50–55 GPa) of the HP-LP phase coexistence on the same grains. Although, this isostress phase transition is related to an increased strain and broadening of the peaks with a slight increase of mosaicity, as well as of dislocation density, we were able to solve the crystal structure as a function of pressure, extracting structural parameters with a high precision.

Our results for the compressional behavior of the LP R3c phase V(P) are fitted with a second-order Birch-Murnaghan (BM2) equation of state (EOS) (see Fig. 2a). At 1 bar and 298 K the calculated bulk modulus and equilibrium unit-cell volume are K₀ = 200.14(7) GPa and V₀ = 268.2(2) Å³, respectively. For the HP R3c phase we obtain Kₗ = 153.8(7) GPa and Vₗ = 257.5(8) Å³. The volume and bulk modulus at 50 GPa (V₅₀ and K₅₀) for the LP phase are 224.98 Å³ and 384.9 GPa and 230.33 Å³ and 417.2 GPa for SC and PWD, respectively. For the HP phase the values are 208.22 Å³ and 335.8 GPa and 211.67 Å³ and 343.5 GPa for SC and PWD, respectively. We observe a large unit-cell volume reduction at the isostress transition above ~50 GPa, ΔV/V ≈ 7.5%. It is attributed to the shrinkage of the Fe–O interatomic distances and the corresponding decrease of the FeO₆ octahedral volume (Fig. 2b,c).

At about 106 GPa we observe a splitting of the (100) reflection (in hexagonal notation) in the SC-XRD pattern (see Fig. 1a,b, Supplementary Fig. S3), implying a structural phase transition with a lowering of the unit-cell symmetry. Based on the SC-XRD we solve the new structure (designated as HP2) to be the monoclinic with the space group (see Supplementary information). In particular, we found that at ambient pressure FeBO₃ adopts the rhombohedral R3c crystal structure (see inset Fig. 2b and Supplementary Fig. S2), in agreement with previous PWD and SC XRD [20 and ref. therein]. The R3c structure is conserved at least up to ~105 GPa (Figs. 1, 2). At about 50 GPa we observe a doubling of the reflections within the diffraction patterns attributed to the onset of the high-pressure (HP) phase characterized by the same space group (R3c) but with significantly reduced unit-cell volume and lattice parameters (Supplementary Fig. S1). Our SC-XRD data confirm that multiple domains of the HP phase grow on top of the HS low-pressure (LP) state grains preserving the same orientation. Here we extend the previous PWD-XRD studies and shed new light on the process of phase transformations in FeBO₃. We observe a finite range (~50–55 GPa) of the HP-LP phase coexistence on the same grains. Although, this isostress phase transition is related to an increased strain and broadening of the peaks with a slight increase of mosaicity, as well as of dislocation density, we were able to solve the crystal structure as a function of pressure, extracting structural parameters with a high precision.

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In comparison to SC-XRD, for the PWD data (Fig. 2a) we observe a larger unit-cell volume at a given pressure, larger bulk modulus values for the LP phase, and a broader pressure range for the phase transition around 50 GPa; all these are indicative of deviatoric stress and enhanced strain contributions, i.e., grain-grain interactions (for more details see Supplementary and Refs. 27–32). However, comparing the SC and PWD data we notice overall GPa; all these are indicative of deviatoric stress and enhanced strain contributions, i.e., grain-grain interactions larger bulk modulus values for the LP phase, and a broader pressure range for the phase transition around 50 GPa. Most notably, it exhibits apparently resilient non-metallic behavior above 100 GPa. This behavior is different, e.g., to FeGaO₃ and FeO₃, which exhibit the classical band-width controlled Mott transition at ~50–106 GPa. Our observations are unexpected given the presence of only a single Fe³⁺ structural position in the crystal structure of FeBO₃. This behavior clearly distinguishes the behavior in FeBO₃ from a "conventional" spin-state crossover observed in other ferric systems. We propose a model explaining this unusual electronic state based on a complex interplay between the spin and lattice degrees of freedom.

In our work, we present a detailed study of the electronic structure, local magnetic state of Fe³⁺ ions, and phase stability of FeBO₃ up to pressures ~150 GPa, combining room and low-temperature ⁵⁷Fe Mössbauer spectroscopy with single crystal (SC) and powder (PWD) X-ray diffraction. Our results reveal that the simultaneous magnetic and isostress phase transition in FeBO₃ at ~50 GPa, which was previously considered as a HS-LS transition, is in fact the transition to a "mixed" HS/LS state. It is characterized by a stable (although a weakly temperature-dependent) abundance ratio of the HS/LS states of ~1:3 in a wide pressure range of ~50–106 GPa. Our observations are unexpected given the presence of only a single Fe³⁺ structural position in the crystal structure of FeBO₃. This behavior clearly distinguishes the behavior in FeBO₃ from a "conventional" spin-state crossover observed in other ferric systems. We propose a model explaining this unusual electronic state based on a complex interplay between the spin and lattice degrees of freedom.
was limited to ~115 GPa since its extension to higher pressures leads to degradation of the EOS fit quality and results in a significant deviation of the obtained parameters from those calculated for SC EOS. For 60–115 GPa

Figure 1. X-ray single crystal (a) and powder (b) diffraction patterns of FeBO$_3$ at RT at various pressures ($\lambda = 0.2898 \, \text{Å}$ and $0.3738 \, \text{Å}$, respectively). Note a splitting of the (10\overline{2}) reflection in the SC and powder XRD pattern at 129 and 145.8 GPa, respectively, signifying lowering of the original symmetry. * marks an unidentified peak, which disappears at higher pressures.
range the performed fit using a BM2 EOS results in $K_0 = 149(12)$ GPa, $V_0 = 262(2)$ Å$^3$ suggesting an appreciable “pressure overestimation” in PWD measurements (see27–30).)

Mössbauer spectroscopy. In Fig. 3 we display Mössbauer spectra of polycrystalline FeBO$_3$ for different pressures recorded at room temperature (RT). In agreement with previous publications$^{22}$, the only observed spectral component upon compression up to ~ 45 GPa within the LP phase is that of the HS state (LP-HS, $S_z = 5/2$, $^6A_1g$) characterized by a magnetically-split sextet and a small quadrupole splitting ($QS \approx 0$ mm/s). At $P \geq 46$ GPa two new doublets emerge: (1) a more intense component with $QS \approx 2$ mm/s and small isomer shift ($IS \approx 0.03$ mm/s); and (2) a less intense component with $QS \approx 1.7$ mm/s and $IS \approx 0.34$ mm/s. Correspondingly, within the coexistence range (~ 46–65 GPa) the spectra are the superposition of three components: the magnetically ordered LP phase with $H_{hf} \approx 48$ T and $IS \approx 0.27$ mm/s, and two high-pressure doublets. We note that a similar observation was previously reported by Sarkisyan et al.$^{22}$ for PWD FeBO$_3$ at a limited pressure range of ~ 48–58 GPa. Considering the Ref.$^{22}$, the authors suggested different behavior of powder and single crystal data. According to their MS results, a two-doublet-structure is not evident in the SC sample up to ~ 55 GPa, in contrast to the powder. We note the rather poor statistics of the SC data at high pressures in Ref.$^{22}$, which potentially prevented detecting small features. Indeed, there is a clear evidence of a small shoulder at 46.6 GPa in the vicinity of 0 mm/s in their Fig. 1 corresponding to single crystal measurements. In order to resolve a potential controversy, we performed our own experiment with large grain material and Ne pressure medium. In this experiment, we tried to avoid possible preferred orientation effects, and, instead of one big piece of a single crystal$^{22}$, we measured tens of pieces of about 10 μm size, obtained by crushing of a SC sample. Noteworthy, in this experiment we did not detect any appreciable difference compared to the case of a powder sample grinded to a size of ~ 1 μm. This MS result in conjunction with the XRD results suggests that the differences between the powder and SC cases are rather small to claim as different scenarios in these cases.

**Figure 2.** Pressure dependencies of the unit-cell volume divided by $Z$ unit formulas ($Z = 6$ and 4 for the $R\bar{3}c$ and $C2/c$ phases, respectively) (a), determined in the powder and single crystal XRD studies; FeO$_6$ octahedral volume (b) and average Fe–O, B–O distances (c) for FeBO$_3$. The solid, dashed, dash-dot and short-dash lines in (a) are fits with the Birch–Murnaghan equation of state (see text). The panel (b) insets show the $R\bar{3}c$ and $C2/c$ crystal structures, respectively. The red and green spheres correspond to the oxygen and boron atoms.
Upon compression above ~ 65 GPa the magnetically-split LP component disappears and only quadrupole-split components (1) and (2) are observed at RT (Fig. 3) with almost the same abundances ratio of ~ 3:1. The parameters QS and IS of the more intense doublet are typical for that of the LS Fe$^{3+}$ state (S = 1/2, $I_2^g$) and coincide with the values obtained for the high-pressure Fe$^{3+}$ state in SC FeBO$_3$.$^{22}$ At the same time, the parameters of the less intense doublet are more typical for that in the Fe$^{3+}$ HS state.$^{33}$ The pressure-dependencies of the isomer shift of the various components and their relative abundances are summarized in Fig. 3b. The relative abundance of site i was determined from the respective areas $A_i$ under the absorption peaks for each component using the relation $A_i = K n_i f_i$ where $K$ is a constant, $n_i$ is the abundance of component i, and $f_i$ is its recoil-free fraction. We assumed as a first approximation that at each pressure the recoil-free fraction values $f_i$ for the three components are the same.

To further clarify the nature of the HP MS components we perform low-temperature MS measurements at temperatures down to 3 K. In Fig. 4 we show the MS spectra for ~ 85 and 140 GPa collected at various temperatures using in-house and synchrotron Mössbauer spectroscopy. Our results reveal a magnetic splitting for both sites at low temperatures although characterized by a very different magnetic splitting: 0.57(5) and 14.75(5) mm/s (at 85 GPa); the corresponding hyperfine field values are $H_{hf} = 1.9(2)$ and $49.5(2)$ T, respectively. Based on the obtained IS and $H_{hf}$ values we finally attribute these components to the HP LS and HP HS states, respectively.

We note that the relative abundances of these components slightly change upon cooling. In fact, the relative abundances of the HS state seem to slightly increase upon temperature decrease (see Fig. 3b). Our detailed low-temperature measurements performed at ~ 115 GPa (Fig. 4c) allow us to estimate the Néel temperature of the HP HS state. The obtained value of $T_N \sim 60(10)$ K is significantly lower than the $T_N \sim 600$ K of LP HS state at ~ 50 GPa.$^{21}$ We note that for the HP phase $T_N$ is almost the same for both components, HS and LS, and that our estimate for $T_N$ is in good qualitative agreement with that obtained from the NFS data for the HP phase (~ 50 K at the range 50–55 GPa)$^{21}$, suggesting that the antiferromagnetic state with $T_N \sim 60$ K persists in a broad pressure range of ~ 50–106 GPa. Our results therefore suggest that around 50 GPa the major portion of the Fe$^{3+}$ ions in FeBO$_3$ undergoes a transition into the LS state, while the rest of the Fe$^{3+}$ ions remain in the HS state (paramagnetic at room temperature).

Interestingly, the phase transition in the mixed HS-LS state above ~ 50 GPa does not lead to a change of the unit-cell symmetry or superstructure formation$^{34,35}$ in FeBO$_3$. In fact, our high-resolution SC and PWD XRD with high accuracy show the R3c unit-cell symmetry characterized by a single Fe$^{3+}$ site up to ~ 106 GPa, implying a cooperative dynamical ordering of the HS/LS Fe$^{3+}$ sites.
Discussion

Our data indicate that under ambient conditions and up to high compression ~ 106 GPa, the crystal lattice of FeBO₃ has the rhombohedral symmetry with space group \( R\bar{3}c \) with a single crystallographic position of the Fe³⁺ ions. In agreement with previous studies, at ~ 50 GPa FeBO₃ undergoes an isostructural phase transition corroborating with a lattice volume collapse of ~ 7.5%. We note differences in the onset pressures and the pressure range of spin crossover as discerned by XRD and MS, which may be attributed to the different pressure transmitting media used and how the degree of nonhydrostaticity affects the electronic transition (see Refs. 31,32). In addition, we note that in the synchrotron XRD measurements the signal derives from a small central part of the sample, whereas in Mössbauer pressure studies the signal is collected from a much larger ~ 2/3 inner region of the sample diameter. In the latter case this results in a potential importance of pressure gradient effects which, on top of the deviatoric stress effect, could be impactful in determining phase transition pressures and the transition pressure range. It was previously shown that at the transition the charge drop gaps from ~ 2.9 to 0.5 eV and then gradually decreases within the HP phase. Previously this transition was considered as a conventional (complete) HS–LS transition of all Fe³⁺ ions in FeBO₃. However, our high-resolution single crystal study gives previously inaccessible structural information which shows the inconsistency of this assumption. We report that the Fe–O distances are shortened by ~ 3.1% and the FeO₆ octahedral volume drops by \( \Delta V/V_{\text{oct}} \approx 9.2\% \). The octahedral volume reduction is remarkably smaller (by ~ 3%) compared to what is observed in confirmed complete Fe³⁺ HS–LS transitions, e.g., in CaFe₂O₄ and FeOOH, where \( \Delta V/V_{\text{oct}} \) ~ 12–12.4%. In Fe₂O₃ the octahedral volume change is even larger, of ~ 14%. However, in this case spin transition coincides with a structure change. In contrast to FeBO₃, the latter values are in good agreement with the theoretical values tabulated by Shannon.

Our ambient temperature MS shows that despite the appearance of the LS state at ~ 50 GPa, a significant part of the Fe³⁺ ions remains in the HS state up to the highest pressures studied here. Most notably, above ~ 65 GPa the abundance of the HS state is almost unaffected by compression and remains rather high, even at pressures above ~ 100 GPa (see Fig. 4). The obtained MS results correlate with a change of the Fe³⁺ site volume. The ~ 3% deficiency in the octahedral volume change for the ~ 50 GPa transition is consistent with the idea of a partial spin transition. Our SC XRD and MS data reveal that for a wide range of pressures and at RT only 75(3)% of Fe³⁺ ions are in the LS state, while the rest remain in the HS state, i.e., the abundance ratio of the HS-to-LS states is of ~ 1:3.

Furthermore, based on the low-temperature MS we verify that the HS/LS abundance ratio shows a weak temperature dependence. To further clarify this point we performed isobaric PWD–XRD measurements, testing the assumption that a change in this ratio may correlate with the unit-cell volume changes. In fact, our low-temperature measurements conducted at 78 GPa show an appreciable negative thermal expansion of the \( R\bar{3}c \) unit-cell at ~ 180–295 K (see Fig. 5, Supplementary Fig. S8) confirming a possible change of the HS/LS abundances ratio with temperature. Below ~ 180 K a conventional \( V(T) \) behavior is observed indicating stabilization of the HS/LS ratio. The observed negative thermal expansion is associated with a crystal volume change by ~ 0.9% upon cooling (Fig. 5), which suggests a rise by ~ 8% of the abundance of the HS state, in agreement with the MS data.

Our results therefore suggest a formation of a stable “mixed” spin state within a very broad pressure range of ~ 50–106 GPa in FeBO₃. Realization of such steady spin-disproportionated state assumes a cooperative ordering of the Fe³⁺ sites with the HS and LS moments. The unforeseen coincidence of the Néel temperatures for the HS and LS states supports a cooperative nature of the mixed spin state. Moreover, since in the \( R\bar{3}c \) structure all the Fe³⁺ occupy identical crystallographic sites, the site selection for HS and LS is random, and not fixed, which

Figure 4. Mössbauer spectra of FeBO₃ at different temperatures at 85 GPa (a), 140 GPa (b) and 115 GPa (c). Spectra at 115 and 140 GPa were collected using synchrotron MS. Spectra collected at 115 GPa allow us to define the Néel temperature of ~ 60(10) K.
results in frustration of magnetic moments. This cooperative phenomenon could potentially be dynamic with dynamical correlations playing a major role within a wide pressure range preceding delocalization of the 3d electrons. While it seems to be plausible that such a dynamical effect may arise from the instantaneous HS-LS interaction of electronic origin, this topic needs further detailed theoretical and experimental considerations. Our numerous attempts to detect a possible formation of a superlattice (reduction of the unit-cell symmetry) in FeBO$_3$ at ~50–106 GPa using SC-XRD have not been successful, which is consistent with the proposed dynamical features (critical nature) of spin correlations. We note that recent theoretical model calculations propose the formation of a (static) spin-disproportionated state in the case of the thermally driven HS-LS transition in LaCoO$_3$.

We observed significant increase of the compressibility and hence decrease of bulk modulus from $K_0$ ~ 200 (LP) to ~154 GPa (HP) for the fixed bulk modulus pressure derivative $K'' = 4$. These results reveal an unusual softening of the lattice that follows the spin transition above ~50 GPa, which is in contrast to the anticipated hardening of the lattice at the HS-LS and/or Mott transition. Furthermore, this agrees with the behavior of the octahedral FeO$_6$ volume obtained above 50 GPa, which points to an averaged spin-state not consistent with the pure HS or LS one. While there is a possibility that quantum spin fluctuations are occurring on a time scale faster than our experimental measurements, we note that the pure HS and LS states are distinguished on the MS time-scales (~10$^{-7}$ s) with no significant broadening observed.

Above ~108 GPa SC FeBO$_3$ undergoes a distortion of the rhombohedral unit-cell resulting in a structural transition to the monoclinic C2/c phase, a subgroup of the original R3c. The phase transition is accompanied by an additional reduction of the Fe–O distances by ~1.3% and octahedral volume by ~3%. This suggests an additional electron density deformation and consequently redistribution of 3d electrons on the Fe$^{3+}$ site. Since the total octahedral volume drop accumulated throughout the different phase transitions up to ~108 GPa in FeBO$_3$ is about 12.2% (i.e., in agreement with Shannon), we can deduce that the R3c to C2/c phase transition is associated with a completion of the HS/LS spin crossover for all Fe$^{3+}$ in FeBO$_3$. The C2/c phase is characterized by a reduced compressibility, as that expected for a complete LS state (Fig. 2a). Considering that the pressure transformation at ~50 GPa is accompanied by a Mott insulator-to-semiconductor transition with a collapse of the charge gap from ~2.9 to 0.5 eV, we propose that the R3c to C2/c phase transformation above ~106 GPa, associated with a completion of the Fe$^{3+}$ transition into the LS state, may result in metallization of FeBO$_3$.

In contrast to SC, in PWD FeBO$_3$ characterized by strong deviatoric stress and grain-grain strain the structural transition is very sluggish, far from completion even at ~145 GPa. In agreement with this, PWD MS at ~140 GPa shows only some small decrease in the abundance of the HS state and of $H_{hf}$. We believe that the above consideration is also applicable in the case of recent electrical transport measurements performed without any pressure-transmitting medium.

**Summary**

We have shown that the interplay between electronic correlations, lattice, and spin states results in the formation of a complex electronic and magnetic behavior of FeBO$_3$ under pressure. In particular, we observe a remarkable coexistence of the HS and LS states in the original R3c structure, characterized by a single Fe$^{3+}$ site, stable within a broad pressure range of ~50–106 GPa. We propose that the spin-disproportionated phase is driven by a cooperative ordering of the HS/LS states, randomly distributed over the R3c lattice, suggesting a potential dynamical nature of the HS-LS correlations. This results in frustration of magnetic moments which is manifested by a large suppression of the Néel temperature to ~60 K above ~50 GPa, compare to $T_N$ ~ 600 K at ~50 GPa before Figure 5. Temperature dependence of the relative unit-cell volume for FeBO$_3$ at ~78 GPa. For comparison we show also the temperature dependence of the relative unit-cell volume for HS Fe$_2$O$_3$, which belong to the same FeXO$_3$ family, at ambient pressure (solid line) calculated from Ref. 40. The dashed line is to guide the eye for a conventional thermal expansion behavior of FeBO$_3$ at the range 10–180 K.
the transition\textsuperscript{21}. Only above ~ 106 GPa for SC FeBO\textsubscript{3}, we found the transition to the lower symmetry C2/c phase which is associated with a further spin-state alteration and possible metallization of FeBO\textsubscript{3}. Our observations emphasize a remarkable importance of spin fluctuations and correlation effects for understanding the electronic structure and magnetic behavior of strongly correlated systems preceding the Mott transition. Overall, our results significantly improve understanding of the pressure-induced evolution of the electronic and magnetic properties of the Mott insulators. Our proposed novel scenario of the spin-state transformation may have important implications not only for the theoretical picture of compounds undergoing a spin-state transition, but also for understanding of quantum criticality of the Mott transitions. We believe that this topic deserves further detailed theoretical and experimental considerations.

Methods
The experiments were performed with high-quality single crystals of FeBO\textsubscript{3} (enriched to 96\% with \textsuperscript{57}Fe isotope when needed for MS experiments)\textsuperscript{43}. Polycrystalline samples were obtained by grinding the FeBO\textsubscript{3} single crystal. Custom diamond anvil cells (DACs) and DACs of symmetric design were used to induce high pressure, with Ne, He or N\textsubscript{2} serving as a pressure-transmitting medium. Pressure was determined using the ruby R\textsubscript{1} fluorescence line as a pressure marker\textsuperscript{44}, as well as the Ne, Au or Pt unit-cell volume in the case of various x-ray diffraction studies. \textsuperscript{57}Fe Mössbauer studies were performed using a 10 mCi \textsuperscript{57}Co (Rh) point source in a variable temperature (5–300 K) cryostat. The spectra at high pressures, 115 and 140 GPa, were collected using energy-domain synchrotron Mössbauer spectroscopy (SMS) carried out at the beamline ID18 at ESRF (Grenoble). SC XRD experiments were performed at the Extreme Conditions Beamline P02.2 at PETRA III (Hamburg, Germany), ID15B beamlines of ESRF (Grenoble) and the 13ID-D GSECARS beamline of APS (Argonne); PWD experiments at the ID27 and ID09A beamlines of ESRF and the 12.2.2 beamline of ALS (Berkeley). Further technical details about the methods used can be found in the Supplementary information S1.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Author contributions
Experiments were conceived by M.P., G.Kh.R., W.X., and K.G. High pressure Mössbauer spectroscopy measurements were performed by W.X., S.L., M.S., and E.G. High pressure SC X-ray diffraction measurements were performed by W.D., K.G., E.B., M.B., E.G., S.L., M.S., G.Kh.R. and M.H. High pressure PWD X-ray diffraction measurements were performed by E.G., S.L., M.S., G.Kh.R. and M.H. The data analysis was carried out by G.Kh.R., K.G., W.X., E.G., E.B. and M.B. S.L. Manuscript was prepared by G.Kh.R., K.G., I.L. and E.G. with contributions from all co-authors.

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