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Citation for published version:
Monserrat, B, Martinez-Canales, M, Needs, RJ & Pickard, CJ 2018, 'Helium-Iron Compounds at Terapascal Pressures', Physical Review Letters, vol. 121, no. 01, pp. 015301.
https://doi.org/10.1103/PhysRevLett.121.015301

Digital Object Identifier (DOI):
10.1103/PhysRevLett.121.015301

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
Physical Review Letters

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Helium-iron compounds at terapascal pressures

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(Dated: June 11, 2018)

We investigate the binary phase diagram of helium and iron using first-principles calculations. We find that helium, which is a noble gas and inert at ambient conditions, forms stable crystalline compounds with iron at terapascal pressures. A FeHe compound becomes stable above 4 TPa, and a FeHe2 compound above 12 TPa. Melting is investigated using molecular dynamics simulations, and a superionic phase with sublattice melting of the helium atoms is predicted. We discuss the implications of our predicted helium-iron phase diagram for interiors of giant (exo)planets and white dwarf stars.

Matter under extreme compression exhibits rich and unexpected behaviour, such as unconventional chemistry [1, 2], structure [3], and phases [4]. Inside planets and stars, electrons and nuclei are subject to extreme conditions of pressure and temperature, and the exploration of new physics and chemistry under these conditions is necessary for the study of astrophysical processes. In this context, we use quantum mechanical calculations to explore the phase diagram of helium and iron, two of the most abundant elements in the Universe.

Helium nuclei formed in the early Universe during Big Bang nucleosynthesis, and the primordial 25% mass fraction of helium makes it the second most abundant element after hydrogen. In addition, thermonuclear reactions within the interiors of stars fuse hydrogen to form helium. Therefore, helium is found inside many astrophysical objects, from planets, to stars, to white dwarf stars, and it plays a central role in their behaviour. For example, recent experimental and theoretical work has shown that helium metallises at TPa pressures [10–12], which is higher than previously anticipated. As a consequence it has been suggested that the cooling rate of white dwarf stars is slowed by their helium-rich atmospheres, and therefore current estimates of their ages need to be revised.

Helium has two electrons in the closed-shell 1s state, and is chemically inert under ambient conditions. The only known helium compounds are either metastable, involving ionised species such as HeH+ [17]; or are formed by weak van der Waals interactions, such as helium inside C60 [18]. Recently, a helium-sodium compound has been reported above pressures of about 0.1 TPa [19].

Iron has one of the highest binding energies per nucleon (the highest is 62Ni) and is therefore also very abundant [20]. It accounts for about 80% of the Earth’s core mass [7], where it is found at pressures up to 0.35 TPa, and it is responsible for the magnetic field surrounding the planet [21]. Iron is not expected to exhibit magnetic order at TPa pressures, and it is predicted to occur in a series of closed-packed non-magnetic crystal structures [22]. Iron compounds with hydrogen, carbon, oxygen, silicon, and sulfur have been investigated at pressures of about 0.35 TPa due to their importance for the composition of the Earth’s core [23].

In this work we investigate the possibility that, under extreme compression, helium might form stable compounds with iron. The high abundances of helium and iron make it crucial to understand the helium-iron phase diagram for astrophysical modelling of the interiors of giant planets, including the increasing number of exoplanets being discovered [24], and iron-core white dwarf stars [25].

Our strategy is to search for high-pressure compounds of helium and iron using first-principles quantum mechanical density functional theory (DFT) methods as implemented in the CASTEP code [26], and the ab initio random structure searching (AIRSS) method [27]. The
stability of a compound \( s \) with respect to the constituent elements can be evaluated by calculating the Gibbs free energy of formation per atom \( \Delta G_s = (G_s - (G_{He}N_{He} + G_{Fe}N_{Fe}))/N_{He} + N_{Fe} \) as described in the text, is indicated by the gradients and approaches \(-4\) eV/atom for both stoichiometries at 100 TPa.

We show the static lattice phase diagram of the helium-iron system in the pressure range 1–100 TPa in Fig. 1. Helium is predicted to adopt the hexagonal closed-packed (hcp) crystal structure at TPa pressures [11]. Iron exhibits a sequence of phase transitions at TPa pressures, starting with the hcp structure which transforms to the face-centered cubic (fcc) structure in the range 7–22 TPa, then it transforms back to the hcp structure up to pressures of 35 TPa, above which it transforms into the body-centered tetragonal (bct) structure, which approaches the body-centered cubic (bcc) structure with increasing pressure [22, 30].

The structure searches find several compounds of helium and iron that are energetically competitive in the TPa pressure range, and the most stable have stoichiometries \( \text{FeHe} \) and \( \text{FeHe}_2 \) (see Fig. 1). The \( \text{FeHe} \) stoichiometry first forms at 4 TPa in a structure of orthorhombic space group \( \text{Cmcm} \) containing 8 atoms in the primitive cell, and at 50 TPa it transforms to a \( \text{Fm3m} \) structure (rock-salt structure). The \( \text{FeHe}_2 \) stoichiometry appears in three distinct structures which have similar energies. The first is an orthorhombic structure of space group \( \text{Cmcm} \) with nine atoms in the primitive cell, which forms around 12 TPa. The second has a space group of \( \text{I41/amd} \) symmetry with six atoms in the primitive cell, and becomes the most stable \( \text{FeHe}_2 \) structure above 47 TPa. The third has \( \text{P6/mmm} \) space group and three atoms in the primitive cell, but is not thermodynamically stable. Structure files for all of the helium-iron compounds are provided as Supplemental Material [29].

The helium-iron compounds that form at the lowest pressures have the \( \text{FeHe} \) \( \text{Cmcm} \) and the \( \text{FeHe}_2 \) \( \text{Cmmm} \) structures shown in Fig. S4. The iron atoms form open channels containing helium chains in the \( \text{FeHe} \) \( \text{Cmcm} \) structure (Fig. S4a). At 10 TPa, the minimum He-He distance is 0.98 Å, the He-Fe distance is 1.16 Å, and the Fe-Fe distance is 1.47 Å. The volume per formula unit in \( \text{FeHe} \) is 2.76 Å\(^3\), compared to 0.50 Å\(^3\) in hcp helium and 2.27 Å\(^3\) in both hcp and fcc iron, which add to a combined volume of 2.77 Å\(^3\) per formula unit. In the \( \text{FeHe}_2 \) \( \text{Cmmm} \) structure (Fig. S4c), the helium atoms form hexagonal layers incorporated inside iron channels that are wider than those present in \( \text{FeHe} \). The minimum He-He distance is 0.89 Å at 10 TPa, the He-Fe distance is 1.19 Å, and the Fe-Fe distance is larger at 1.54 Å. The volume per formula unit in \( \text{FeHe}_2 \) is 3.24 Å\(^3\), which is smaller than that of the elements (total of 3.27 Å\(^3\)). The smaller volumes of the compounds favour their formation under pressure via the enthalpy term in the Gibbs free energy.

We next investigate the effects of temperature on the formation of helium-iron compounds upon increasing pressure. If the effects of nuclear motion are neglected, \( \text{FeHe} \) forms at pressures above 4.1 TPa, and the inclusion of quantum and thermal nuclear motion lowers this pressure to 2.7 TPa at 10,000 K. \( \text{FeHe}_2 \) only forms at a higher pressure of about 12 TPa, and therefore we focus on the \( \text{FeHe} \) compound to study the formation of helium-iron compounds under pressure.

We use \textit{ab initio} molecular dynamics simulations in conjunction with the Z-method [31] to estimate the melting temperature of \( \text{FeHe} \). These calculations are
performed using the `Quantum Espresso` package [32], and the details are provided in the Supplemental Material [29]. The melting temperatures of helium and iron differ by thousands of degrees, suggesting that FeHe might exhibit superionicity, that is, sublattice melting of the helium component while the iron atoms oscillate around their crystallographic positions. Superionicity has been discussed before [33], for example in a lithium-based conductor at ambient pressure [34], and in the melting of ice and ammonia at extreme pressures [8]. Indeed, our molecular dynamics simulations demonstrate that, upon increasing temperature, the helium chains melt within the iron channels in FeHe before the iron channels themselves melt. Interestingly, metallic superionic compounds are uncommon [35], and FeHe provides a nice platform to further investigate their properties.

In Fig. 3 we show the proposed phase diagram for the formation of helium-iron compounds under pressures up to 10 TPa. At low pressures, helium and iron do not mix. Below about 4,000 K at 1 TPa and 6,000 K at 3 TPa, both materials are found in the solid state, but helium melts above this temperature. Iron only melts at much higher temperatures, of the order of 15,000 K [36]. Upon increasing pressure, helium and iron form a FeHe compound between 2 and 4 TPa, depending on the temperature. FeHe undergoes sublattice melting of the helium atoms at temperatures between 13,000 and 18,000 K, depending on the pressure. The superionic phase is stable in a wide temperature and pressure range, and melting is completed at around 17,000 K at 4 TPa, and above 19,000 K at 10 TPa.

Our results suggest that the FeHe compound should form at the pressures accessible to dynamic compression experiments. Furthermore, the formation pressure of FeHe is predicted to be within the pressure range found at the core of Jupiter, with a core-mantle boundary pressure of 4.2 TPa and temperature of 20,000 K, and at the highest pressures found at the centre of Saturn, with a core-mantle boundary pressure of 1 TPa [14, 37]. The interiors of exoplanets with masses similar or larger than that of Jupiter will also be subject to pressures higher than those required to form FeHe. This raises the possibility that helium is captured by iron within the interior of these planets, and potentially bound to other elements. The atmosphere of Saturn is indeed depleted of helium [38], and the capture of helium in compounds in its interior could contribute to this phenomenon. This could also affect the helium composition of the atmospheres of giant exoplanets. White dwarf stars are subject to more extreme conditions, with helium-rich atmospheres subject to tens of terapascals, and the interiors to even higher pressures. Due to cooling, white dwarf stars exhibit temperatures in the range from only a few thousand Kelvin to hundreds of thousands Kelvin [39], raising the possibility that even the solid FeHe phases appear in these stars. The formation of helium compounds with other elements could alter the cooling rates of white dwarf stars, which are largely determined by the atmospheric composition, and as a consequence affect current estimates of their ages. Our results indicate that, in contrast to the inertness of helium at ambient pressure, accurate models of the composition of planets and stars should treat helium as a compound-forming element.

In conclusion, we have used first-principles methods to study the binary phase diagram of helium and iron. We have found that compounds can form at pressures of several TPa, suggesting that they might be found inside giant (exo)planets and white dwarf stars. We have also predicted that the most stable FeHe compound exhibits a superionic phase with sublattice melting of the helium atoms within a wide range of temperatures and pressures. Overall, our results show that helium can form compounds at terapascal pressures.

B.M. acknowledges support from the Winton Programme for the Physics of Sustainability, and from Robinson College, Cambridge, and the Cambridge Philosophical Society for a Henslow Research Fellowship. R.J.N. and C.J.P. acknowledge financial support from the Engineering and Physical Sciences Research Council (EPSRC) of the UK [EP/J017639/1 and EP/G007489/2]. C.J.P. is also supported by the Royal Society through a Royal Society Wolfson Research Merit award. The calculations were performed on the Cambridge High Performance Computing Service facility and the Archer facility of the UK’s national high-
performance computing service (for which access was obtained via the UKCP consortium [EP/K014560/1 and EP/K013688/1]).

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Supplemental Material for “Helium-iron compounds at terapascal pressures”

COMPUTATIONAL DETAILS

Density functional theory

Density functional theory calculations have been performed using the plane-wave pseudopotential CASTEP code [S26] with the PBE functional [S40]. The extreme pressures we study force us to use tailored pseudopotentials with core radii of 0.6 bohr for helium and 1.0 bohr for iron, and our calculations explicitly include the 1s states of helium and the 3s3p3d4s states of iron. The pseudopotential strings used are:

He 1|0.6|27|31|36|10(qc=11)
Fe 3|1.0|33|38|44|30U:40:31:32(qc=11)

These hard pseudopotentials require plane-wave cut-off energies of 1500 eV. For the searches we have used Brillouin zone k-point grids of density $2\pi \times 0.07$ Å$^{-1}$, and finer grids of density $2\pi \times 0.025$ Å$^{-1}$ for the final results and the vibrational calculations. Structures were relaxed to achieve uncertainties in the pressure below 0.1 TPa.

AIRSS

The structure searches have been performed using the AIRSS method [S27, S41] at pressures of 5 TPa, 10 TPa, and 50 TPa, and with stoichiometries $\text{Fe}_x\text{He}_y$ for $x, y = 1, \ldots, 6$. The total number of structures generated in the searches is 7,371.

Vibrational calculations

Harmonic vibrational free energies have been calculated using finite-displacements [S42] with the nondiagonal supercells method [S28]. The system sizes required to obtain converged results are shown in Table I, and the accuracy required could not have been achieved unless nondiagonal supercells had been used, as the computational cost of using small radii pseudopotentials is significant. Results accurate to 1 meV/atom have been obtained by diagonalising the corresponding dynamical matrices over a fine vibrational Brillouin-zone grid. Anharmonic vibrations have been previously found to be negligible in helium at TPa pressures [S11].

| TABLE I. Size of largest supercell required for the corresponding vibrational BZ grid. |
|--------------------------------|-----------------|-----------------|
|                                | BZ grid size    | Diagonal supercells | Nondiagonal supercells |
| Fe-hpc  | 4 × 4 × 4        | 64 cells (128 atoms) | 4 cells (8 atoms) |
| Fe-hcp  | 4 × 4 × 4        | 64 cells (128 atoms) | 4 cells (8 atoms) |
| Fe-fcc  | 4 × 4 × 4        | 64 cells (64 atoms)  | 4 cells (4 atoms)  |
| FeHe-Cmcm  | 3 × 3 × 3        | 27 cells (216 atoms) | 3 cells (24 atoms) |
| FeHe$_2$-P6/mmm  | 5 × 5 × 5        | 125 cells (375 atoms) | 5 cells (15 atoms) |
| FeHe$_2$-I$_4$/amd  | 3 × 3 × 3        | 27 cells (162 atoms) | 3 cells (18 atoms) |
| FeHe$_2$-Cmmm  | 4 × 4 × 4        | 64 cells (576 atoms) | 4 cells (36 atoms) |

THERMODYNAMIC STABILITY

In this section we show the relative stability of the various crystal structures for iron, helium, FeHe, and FeHe$_2$. For any pressure-temperature conditions, the thermodynamically stable structure is used in the main text.

The relevant structures of iron are the face-centered cubic (fcc), body-centered cubic (bcc), body-centered tetragonal (bct), and hexagonal closed-packed (hcp). Their relative static lattice enthalpies are shown in Fig. S1a. In agreement
with previous reports [S22], we find that fcc iron becomes stable compared to hcp iron at pressures in the range 7.4 TPa to 21.7 TPa, and a body-centered tetragonal (bct) iron is the most stable structure above 35 TPa. The bct structure approaches the bcc structure with increasing pressure. The effects of temperature are shown in Fig. S1b up to 20 TPa. Including quantum and thermal nuclear motion destabilises fcc iron, so that at 10,000 K it only becomes stable at 9.4 TPa.

Helium is only found in the hcp structure in the entire pressure range from 1 TPa to 100 TPa. A comparison with the bcc and fcc structures is provided in Fig. S2.

For the helium-iron FeHe stoichiometry, there are two different structures that are energetically competitive: (i) a structure of space group Cmcm with 8 atoms in the primitive cell, and (ii) a structure of space group Fm-3m with 2 atoms in the primitive cell (rock-salt structure). The relative enthalpies of these structures are shown in Fig. S3.

For the helium-iron FeHe stoichiometry, there are three different structures that are energetically competitive: (i)
FIG. S5. Phase diagram for the helium-iron compounds of FeHe$_2$ stoichiometry.

FIG. S6. Convex hull diagram for the helium-iron system at the static lattice level and for a range of temperatures. The solid circles indicate structures on the convex hull, and the solid lines connect them. The open circles indicate structures that are not on the convex hull, and these are connected by dotted lines as a guide to the eye.

a structure of space group $P6/mmm$ with 3 atoms in the primitive cell, (ii) a structure of space group $I4_1/amd$ with 6 atoms in the primitive cell, and (iii) a structure of space group $Cmmm$ with 9 atoms in the primitive cell. These structures are depicted in Fig. S4, and their relative enthalpies and low pressure phase diagram are shown in Fig. S5. Combining the He, Fe, FeHe, and FeHe$_2$ results, we construct the convex hull diagram for the helium-iron mixtures, shown in Fig. S6.

SUPERIONICITY AND MELTING

We investigate superionicity and melting in the FeHe $Cmcm$ structure using molecular dynamics simulations performed with QUANTUM ESPRESSO [S32]. We used pseudopotentials converted from CASTEP with the following strings:

He 1|0.7|27|31|36|10(qc=9.2)
Fe 3|1.2|30|35|40|30U:40:31:32(qc=10)

We note that the computational expense of the molecular dynamics calculations forced us to use pseudopotentials with larger core radii than those used for the lattice dynamics calculations. Nonetheless, we have calculated the phonon density of states at 10 TPa for the FeHe $Cmcm$ structure and confirmed that the hard and soft pseudopotentials lead to consistent results (see Fig. S7). We used an energy cutoff of 90 Ry (1225 eV), and a charge density cutoff of 720 Ry. The electronic Brillouin zone was sampled using a $2 \times 2 \times 2$ k-point grid for a $4 \times 2 \times 4$ supercell of the conventional cell of the $Cmcm$ structure, containing a total of 256 atoms.
The molecular dynamics simulations were used to investigate the melting curve of the FeHe \( Cmcm \) compound, for which the Z-method [S31] was used. In this method, the temperature is increased to the desired value using an NVT ensemble, and then relaxed using an NVE ensemble. This procedure is iterated while checking whether a component (He or Fe) is diffusing or the whole lattice has melted by monitoring both the temperature and the mean square displacement (MSD) of the relevant species. Representative runs at 5 and 10 TPa are shown in Fig. S8.

The results in Fig. S9 are used to prove that the iron sublattice can withstand helium diffusion and therefore sustain the superionic phase. We first equilibrated FeHe at 10 TPa and 17,500 K. Next, we equilibrated at 19,000 K, a temperature at which helium melts and the relaxed temperature then decreases to 18,500 K. The iron sublattice survives for more than 1.5 ps, at which point we stop the simulation. We have performed additional tests in which the iron sublattice remains crystalline up to 10 ps. These molecular dynamics runs provide strong evidence for the existence of the superionic phase in FeHe.

It is interesting to note that the Lindemann criterion correctly predicts the existence of a superionic phase. This criterion associates nuclear vibrations with structural changes and states that a solid melts when the ratio of the root-mean-square nuclear vibrational amplitude and the interatomic separation exceeds some critical value, \( \sqrt{\langle u^2 \rangle}/a \gtrsim \lambda_c \), which is usually taken to be \( \lambda_c = 0.1 \). The Lindemann criterion using the smallest He–He distance as the interatomic separation \( a \) predicts that the helium sublattice melts at temperatures of a few thousand Kelvin, but at those temperatures the Fe–Fe Lindemann ratio is significantly smaller than \( \lambda_c \). These results suggest that, upon increasing temperature, the helium chains melt within the iron channels in FeHe before the iron channels themselves melt. We note that quantitatively, the Lindemann criterion results significantly deviate from the molecular dynamics results.
FIG. S9. Molecular dynamics runs of \textit{Cmcm} FeHe in which the helium sublattice is melted but the iron sublattice is not, demonstrating the existence of a superionic phase in FeHe.

**HELIUM AND IRON MELTING**

We have calculated the helium melting curve using MD in a similar manner to that described above for FeHe \textit{Cmcm}. We will report full details of this calculation in a separate publication.

The iron melting curve is taken from Ref. [S36], in which density functional theory in conjunction with a two-phase coexistence approach was used.