Helium-hydrogen immiscibility at high pressures

Yu Wang1,2, Xiao Zhang1,2, Shuqing Jiang1, Zachary M. Geballe3, Teerachote Pakornchote3,4, Maddury Somayazulu3, Vitali B. Prakapenka5, Eran Greenberg5, Alexander F. Goncharov1-3

1Key Laboratory of Materials Physics and Center for Energy Matter in Extreme Environments, Institute of Solid State Physics, Chinese Academy of Sciences, Hefei, Anhui 230031, China
2University of Science and Technology of China, Hefei 230026, Anhui, People’s Republic of China
3Geophysical Laboratory, Carnegie Institution of Washington, 5251 Broad Branch Road, Washington, DC 20015, USA
4Department of Physics, Chulalongkorn University, Bangkok, 10330, Thailand
5Center for Advanced Radiations Sources, University of Chicago, Chicago, Illinois 60637, USA

Abstract

Hydrogen and helium are the most abundant elements in the universe and they constitute the interiors of gas giant planets. Thus, their equations of states, phase, chemical state, and chemical reactivity at extreme conditions are of great interest. Applying Raman spectroscopy, visual observation, and synchrotron X-ray diffraction in diamond anvil cells (DAC), we performed experiments on H2-He 1:1 and D2-He 1:10 compressed gas mixtures up to 100 GPa at 300 K. By comparing with the available data on pure bulk materials, we find no sign of miscibility, chemical reactivity, and new compound formation. This result establishes a new baseline for future investigations of miscibility in the He-H2 system at extreme P-T conditions.

Introduction

As the two lightest elements, hydrogen and helium are ubiquitous in universe. Moreover, they are the essential elements in the composition of the gas giant planets Jupiter and Saturn in our solar system 1-7. The existence of helium in the molecular hydrogen ocean and the sedimentation of the helium due to helium-hydrogen separation at high pressures has a significant effect on the evolution of the interiors of gas giants 8-12.

Recently, great progress has been achieved both theoretically and experimentally in understanding the transformation of chemical bonding and formation of novel materials under extreme conditions of high pressure 13-18. Under pressure, the electronic states of atoms and molecules change, giving elements new chemical character, which can create polymerized states 19,20, compounds with unusual stoichiometry 13,21,22, electrides, and
multicenter bonding \(^{23,24}\). It has been found that even chemically inert materials such as rare gas solids made of noble gases become chemically reactive at high pressures \(^{16,17,25}\). Thus, the chemical inertness of He (which is the second most inert element after Ne) at high pressures \(^{16}\) cannot be taken for granted. In fact, He forms binary mixed van der Waals compounds with molecular nitrogen - He(N\(_2\))\(_{11}\) \(^{26}\) and neon - NeHe\(_2\) \(^{27}\) and it was reported to form an electride compound with Na above 113 GPa \(^{16}\). In addition, molecular hydrogen is known to react easily with many elements, forming hydrides with variable composition, including polyhydrides at high pressures \(^{15,28-31}\). Moreover, hydrogen forms binary van der Waals compounds with other diatomic molecules, such as N\(_2\) \(^{32,33}\). The latter compounds were found to react to form N-H chemical bonds above 47 GPa \(^{32,34,35}\). Previous experimental investigations in He-H\(_2\) mixtures showed very small miscibility (if any) at room temperature up to 14 GPa \(^{36,37}\), which enabled single crystal X-ray diffraction studies of hydrogen to 120 GPa \(^{38}\). Theoretical investigations of fluid H\(_2\)-He mixtures show immiscibility at 100s of GPa, over a temperature range up to at least 3000 K \(^{39-42}\). However, to the best of our knowledge there are no predictions about the formation of any compound at low temperatures. In contrast, a strong chemical association, miscibility, and structural changes in He-H\(_2\) mixtures were reported recently at pressures as low as 12.6 GPa \(^{43}\). This report has been critiqued in a very recent report \(^{44}\), where the Raman peaks of new phases observed by Lim and Yoo \(^{43}\) have been attributed to N\(_2\) impurity. Here we present Raman spectroscopy and X-ray diffraction (XRD) results in 1:1 H\(_2\)-He and 1:10 D\(_2\)-He mixtures up to 100 GPa at 300 K. In contrast to Lim and Yoo \(^{42}\), we show immiscibility of H\(_2\) and D\(_2\) with He, as well as the absence of the He-H\(_2\) and He-D\(_2\) compounds. The complex Raman spectra of He-H\(_2\) and N\(_2\) doped He reported by Lim and Yoo \(^{43}\) are due to contamination by N\(_2\) and O\(_2\), respectively. Our results support the phase diagram reported in Ref. 37 and extend it to higher pressure, assuming that miscibility is not hindered by kinetics.

We performed experiments with the research grade (99.999\% purity) 1:1 H\(_2\)–He and self-produced approximately 1:10 D\(_2\)–He mixtures made of high purity research grade components. The choice of the compositions was arbitrary as Lim and Yoo \(^{43}\) reported their effects in a wide range of He-H\(_2\) concentrations. The gas mixture was loaded into DACs by compressing up to 0.17 GPa at room temperature. To trap a 1:10 ratio of D\(_2\) to He in our high-pressure sample chamber, we pumped D\(_2\) to a pressure of 17 MPa and then pumped He to a total pressure of 170 MPa, waited for about 1 hour for the gases to mix, and then sealed the DAC. The connecting gas lines and the high-pressure loading chamber were carefully purged before gas mixing and loading. Raman experiments were performed at ISSP (Hefei, China), GSECARS (APS, Argonne National Lab), and the Geophysical Laboratory (Carnegie Institution of Science) using similar setups which include solid state lasers (473, 488, 532, and 660 nm excitation wavelengths), ultralow low-frequency notch filters (down to 10 cm\(^{-1}\) Raman coverage), and wavelength dispersive single grating spectrographs with CCD detectors \(^{16,34,45}\). X-ray diffraction measurements were performed at the undulator XRD beamline at GSECARS. Typical X-ray beam size in all the experiments was 2-3 \(\mu\)m. Pressure was
determined via ruby fluorescence scale\textsuperscript{46}, Raman of the stressed diamond anvil \textsuperscript{47}, gold and ruby XRD sensors \textsuperscript{48, 49} (the latter one below 50 GPa). In the control XRD experiments on He the sample was laser annealed up to 2000 K at 70 GPa using nanocarbon immersed in He; this did not affect the results substantially indicating that He remains in quasihydrostatic conditions. All the measurements have been performed at room temperature. The figures presenting the proposed phase diagram and the additional XRD and Raman data are in Supplemental Material \textsuperscript{50}.

After gas loading the H\textsubscript{2}–He gas mixture to approximately 0.2 GPa, the system is a single miscible fluid (Figs. 1 and S1 \textsuperscript{50}). The Raman spectra show the bands characterizing H\textsubscript{2} molecules: rotons and vibron corresponding to change in rotational and vibrational molecular states. We find that the spectra are independent of the sample position and the sample looks uniform, indicating miscibility (Fig. 1). The Raman frequency of the vibron mode in a disordered state strongly depends on the H\textsubscript{2} concentration and the matrix material \textsuperscript{37, 51}. It is shifted to higher frequency compared to pure bulk H\textsubscript{2} because of the difference in environment. Upon the transition to a state where H\textsubscript{2} solidifies (Fig. S1 \textsuperscript{50}), the vibron mode shifts down in frequency abruptly due to a cooperative effect of intermolecular coupling in the ordered state. The sample shows three distinct areas in this regime: H\textsubscript{2}-rich solid S\textsubscript{1}, He-rich fluid F\textsubscript{2}, and a mesoscopic mixed region (Figs. 1, 2); the Raman spectra of hydrogen clearly characterizes all these three areas (Fig. 1).

**Figure 1 (left)**. Microphotograph and corresponding Raman spectra of H\textsubscript{2}-He mixtures at different phases. He-rich areas are marked in blue “He”. H\textsubscript{2}–rich areas are marked in red “H\textsubscript{2}”. A mixture He and H\textsubscript{2} is marked in green “He+H\textsubscript{2}”. a) Pressure below 5 GPa. Both hydrogen and helium are in a fluid state. b) Pressure above the solidification line of H\textsubscript{2}–rich fluid in Fig. S1 \textsuperscript{50}. Hydrogen is solid and helium is in a fluid state. Corresponding spectra are in same color. c) Pressure above the solidification line of He–rich fluid. Both hydrogen and helium are in solid state. Pressure is inferred from the position of the hydrogen vibron of F\textsubscript{1}+F\textsubscript{2} and S\textsubscript{1} as ruby is absent in this set of experiments to avoid the fluorescence, which can interfere with the Raman detection of.
weak hydrogen vibron modes.

**Figure 2 (right).** Temporal changes in morphology of H$_2$-He mixtures in the stability range of S$_1$-F$_2$ conglomerate when also exposed to a low-power laser beam (at ~7 GPa). Smaller hydrogen-rich crystals surrounded by fluid helium migrate with time to merge with the bigger hydrogen crystals.

No measurable H$_2$-vibron frequency irregularity is observed through the solidification of He, and the vibron frequency in both S$_1$+F$_1$ and S$_1$+S$_2$ states follows that of a pure bulk hydrogen, indicating the formation of a H$_2$-rich solid in our experiments (Fig. 3). We find no sign of the S$_1’$ and S$_3$ solids that seemed to be created in the experiments of Ref. 43. In the previous work, phases S$_1’$, S$_2$, and S$_3$ were characterized by “novel” Raman bands at 140, 2400, and 3200 cm$^{-1}$, and additional higher frequency H$_2$ vibron modes. The present measurements are sensitive to these Raman signatures, and yet no signal is detected (Fig. 4). Raman experiments in the D$_2$-He system up to 54 GPa reveal a similar behavior and also no extra lines have been recorded (Figs. S2 and S3 50). These latter experiments were designed to investigate the isotope effect on the H-He vibron frequency at 2400 cm$^{-1}$ proposed by Lim and Yoo 43, but the experiments yielded no measurable Raman peak at 2400 cm$^{-1}$ or at an isotopically shifted frequency.

Our Raman measurements aimed at the He-rich solid S$_2$ show a small peak at exactly the same frequency as that of the vibron mode of H$_2$-rich solid S$_1$, strongly suggesting that it is originated in the S$_1$ region of the sample chamber where it is the dominant Raman signal (Fig. 4). In both S$_1$ and S$_2$ solids, there are no other Raman bands at 140, 2400, and 3200 cm$^{-1}$, and additional higher frequency H$_2$ vibron modes as reported in Ref. 43 (Figs. 1, 4 and Fig. S3 50). Based on the absence of any extra Raman peaks in S$_2$ and the signal to noise ratio exceeding 1000:1 for the H$_2$(D$_2$)-vibron, we conclude that miscibility of H$_2$(D$_2$) in He is less than 0.1% and there is no detectable formation of any chemical bonds between He and H(D)43. We note that if a compound of He and H$_2$ formed, it would be an inclusion compound stabilized via an enthalpy gain due to long-range Coulomb interactions 52. However, we find no sign of an extra lattice mode (Figs. 1, 3, 4), which would likely to appear in this case (cf. Ref. 43).
Figure 3. The pressure dependence of the Raman frequencies of the H₂-rich fluid and solid in He medium as a function of pressure. Black solid and gray lines and light blue crosses are the literature data (see Ref. 54 and References therein) for H₂. An inset shows the details of low-pressure behavior, the results of previous measurements are from Ref. 36. Our spectra indicate the presence of HD molecules (<2%) as the major impurity in H₂. Crosses in the right panel show the previously measured pressure dependence of the vibron mode of HD molecules which formed in 70:30 H₂-D₂ gas mixture.

The low-frequency Raman spectra of H₂-rich solid S₁ probe the quantum rotational transitions (rotons), which are weakly pressure dependent, and a translational mode (phonon), which is strongly pressure dependent (Fig. 3). The phonon mode is relatively weak and it can be observed via the rotational-vibrational coupling with the roton modes, which can be clearly viewed as a dip that first appears at the left side of an S₀(1) roton mode at nearly 600 cm⁻¹ and then moves with pressure to the right side of this band (Fig. S4). The pressure dependencies of the Raman modes show an avoided crossing of the phonon roton modes (Fig. 3). These measurements demonstrate that S₁ solid is well ordered as otherwise one would expect to see a relaxation of the Raman selection rules. Furthermore, the comparison of the low-frequency spectra show that they are indistinguishable from those of the pure bulk H₂ (Fig. S4). Our experiments in He-D₂ system demonstrated qualitatively similar results (Fig. S3). On the contrary, measurements of these excitations in mixed disordered H₂-D₂ crystals show a broader spectrum and no coupling of the phonon and roton modes.
Figure 4. Raman spectra of H$_2$-He mixtures at different pressures in the range of rotational-translational (a), second-order diamond (b), and vibrational (c) modes. The spectra of hydrogen- (S$_1$) and helium- (S$_2$) rich parts are shown in red and blue respectively. The spectra of S$_2$ solid have a small contribution from the S$_1$ solid, which surrounds it.

Figure 5. X-ray diffraction patterns of H$_2$-He mixtures at 37.5 GPa measured using a wide angle scan to make detectable single crystal reflections (see, for example, Ref. 56). The x-ray wavelength is 0.3344 Å. Blue and red vertical ticks with the peak indexing are presented for the He and H$_2$ respectively. Yellow vertical ticks correspond to the positions of the ruby XRD peaks, which were used to determine the pressure $^{49}$. The right panel is the 2D XRD image (cake) of the He rich solid.
Synchrotron XRD measurements have been performed on the same H₂-He sample at 37.5 GPa and 102 GPa (Fig. 5). The DAC has been rotated along the vertical (Ω) axis to match the Bragg conditions in single crystals of He- and H₂-rich solids. We have been able to observe a few low-hkI diffraction peaks, which identify the lattice symmetry of these crystals (Fig. 5; Table 1). The results show that both solids form an hcp lattice, in agreement with previous reports on He up to 53 GPa 38, 47, 57, 58 and H₂ (e.g. Ref. 38). These data have been compared to the results of three other experimental XRD runs (without hydrogen) up to 74 GPa, where He was the sample or served as a transmitting medium, and the lattice parameters were determined similarly to the described above (Fig. 6). The (101) XRD peak of the H₂ rich solid is documented in our H₂-He experiment at 37.5 GPa, and its position is consistent with hcp H₂ reported previously (e.g., Ref. 38).

The lattice parameters and the unit cell volume of He measured here are in a good agreement with those measured previously (Fig. 6), in which pure He was used as a sample 57, 58. A small disagreement may be related to systematic errors due to the use of the energy dispersive method in these works. Our results for He-H₂ system and for pure bulk He are in a good agreement with each other (Fig. 6). This indicates that He-rich solid S₂ prepared in mixtures of H₂ and He can accommodate very little H₂ impurity (if any). The upper limit of the H₂ doping content can be estimated using Vegard’s law based on the difference in the lattice parameters of H₂ and He which both form hcp lattices in the explored pressure range. For example, at 100 GPa the difference in the lattice parameters is about 10%. The lattice parameters of He solid determined in this work are accurate to about 0.1%. Thus, the maximum amount of H₂ admixture in He rich solid determined from our XRD experiments is about 1%.
Our Raman and XRD experiments in He-H$_2$ mixtures up to 102 GPa show that H$_2$ and He form the corresponding solid phases. Each phase does not reveal any measurable difference in structural, thermodynamic, or vibrational properties to those of pure materials. Providing that hydrogen and helium are sufficiently mobile at high pressures \( (e.g. \text{Refs. 60-62}) \) and the observed immiscibility is not the kinetic effect, this finding extends the main result of Ref. 36 to higher pressure, strongly suggesting that there is no measurable miscibility of H$_2$ and He in solid phases. How can these results be reconciled with the report about the additional Raman bands at 140, 2400, 3200 cm$^{-1}$, and above 4280 cm$^{-1}$ in the He-rich solid \( 43 ? \)

First of all, we note that these new bands closely correlate with those reported in the Raman spectra of the H$_2$-N$_2$ mixed system reported previously\(^{32,34,35} \). These bands can be naturally assigned to the excitations of the H$_2$-N$_2$ van der Waals crystals. In this regard, the disappearance or substantial diminishing of the 2400 cm$^{-1}$ (N$_2$ vibrons) and 4200 cm$^{-1}$ (H$_2$ vibrons) bands and appearance of the 3200 cm$^{-1}$ band (N-H stretch) reported in Ref. 43 at 50 GPa is closely correlated with the polymerization of the H$_2$-N$_2$ mixed crystal at these conditions \(^{34,35} \). In accord with this, our XRD results indicate that hcp He-rich solid remains stable up to at least 102 GPa similarly to the behavior of pure He suggesting that N$_2$ impurity goes to the H$_2$ rich solid in experiments of Lim and Yoo \(^{43} \). As such, we rule out the recently reported chemical association, miscibility and structural change of He and H$_2$ \(^{43} \). Similar conclusions have been reached in the independent investigations reported recently \(^{44} \). Lim and Yoo \(^{43} \) point out that the Raman bands in the 2400 cm$^{-1}$ range are somewhat dissimilar in N$_2$ doped He (and also pure bulk N$_2$) and their He-H$_2$ system. However, they compare different systems so these differences can be easily reconciled. Indeed, N$_2$ and He at high pressures form a \( (N_2)_1 \) He crystal \(^{63} \), the N$_2$-vibron spectra of which are somewhat different from pure bulk N$_2$. Furthermore, we argue that N$_2$ impurities in Lim and Yoo’s He-H$_2$ experiments go into the H$_2$ crystal, not the He crystal, so their control experiment in the N$_2$-He system is not representative. In contrast to the N$_2$-He system, the N$_2$ vibron spectra in the H$_2$-N$_2$ system \(^{34,44} \) are very similar to those reported by Lim and Yoo’s in terms of a number of the bands and their frequencies. Please note that the N$_2$ vibron spectra of the H$_2$-N$_2$ system depend on the composition, so a small disagreement is possible as the H$_2$ content is expected to be larger in the experiments of Lim and Yoo \(^{43} \) compared to studies in which N$_2$ was loaded intentionally\(^{32,34,35,44} \). Moreover, one should take into account the difference in the spectra of bulk solids \( (e.g., \text{bulk pure N}_2) \) and N$_2$ impurities in H$_2$. In the pure bulk materials there are collective effects, which results in splitting of the vibrons and their shifts due to intermolecular coupling. On the contrary, the impurity molecules, which are uncoupled, will have different vibron frequencies. This
has been carefully documented for H₂ vibron impurity modes by Loubeyre et al. ⁵¹.

Second, Lim and Yoo⁴³ performed a control experiment in the N₂-He experiment and found yet another set of extra Raman lines, which cannot be attributed to nitrogen molecules. Therefore, they argued about the N₂ contamination in their He-H₂ system. We have compared their frequencies in the N₂-He system at different pressures with the literature data on the pressure dependence of the Raman frequencies in O₂ ⁶⁴, ⁶⁵. The agreement is almost perfect (Fig. S5 ⁵⁰) making clear that Lim and Yoo reported the behavior of O₂ doped N₂-He system, not a pure N₂-He system. Our data and the data of Turnbull et al. ⁴⁴ do not show the 1600 cm⁻¹ peak and others at lower frequencies because these works studied pure systems that do not contain N₂ and O₂.

In conclusion, our combined Raman and XRD experiments on H₂(D₂)-He system up to 102 GPa at 300 K show the occurrence of only two solids, H₂ and He. We constrain the impurity concentration in these solids to be below 1 %, thus revealing a very large miscibility gap at the explored conditions of pressure and temperature. Our experiments explain the extra Raman bands reported recently by Lim and Yoo ⁴³ as due to the presence of N₂ and O₂ impurities in their samples. As such, we rule out the recently reported chemical association, miscibility and structural change of He and H₂ ⁴³.

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|      | hkl | d   | 2θ  | a     | c     |
|------|-----|-----|-----|-------|-------|
| He   | 101 | 1.449 | 13.250 | 1.894(2) | 3.093(3) |
| He   | 100 | 1.640 | 11.700 |       |       |
| H₂   | 101 | 1.656 | 11.587 | 2.177(4) | 3.495(6) |
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Supplemental Materials to

**Helium-hydrogen immiscibility at high pressures**

Yu Wang\(^1,2\), Xiao Zhang\(^1,2\), Shuqing Jiang\(^1\), Zachary M. Geballe\(^3\), Teerachote Pakornchote\(^3,4\), Maddury Somayazulu\(^3\), Vitali Prakapenka\(^5\), Eran Greenberg\(^5\),

Alexander F. Goncharov\(^1,3\)

\(^1\)Key Laboratory of Materials Physics and Center for Energy Matter in Extreme Environments, Institute of Solid State Physics, Chinese Academy of Sciences, Hefei, Anhui 230031, China

\(^2\)University of Science and Technology of China, Hefei 230026, Anhui, People’s Republic of China

\(^3\)Geophysical Laboratory, Carnegie Institution of Washington, 5251 Broad Branch Road, Washington, DC 20015, USA

\(^4\)Department of Physics, Chulalongkorn University, Bangkok, 10330, Thailand

\(^5\)Center for Advanced Radiations Sources, University of Chicago, Chicago, Illinois 60637, USA

Alexander Goncharov
E-mail: agoncharov@carnegiescience.edu

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Figs. S1 to S5
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Figure S1. Binary phase diagram of H₂-He mixtures adapted from Loubeyre et al. ¹. Dashed line with arrow shows our experiment path. F₁, S₁, F₂ and S₂ signify fluid H₂, solid H₂, fluid He, and solid He, respectively. The areas colored green correspond to miscibility regions, while those color gray- are immiscible. Thick red and blue lines correspond to H₂ and He solids, observed in this study via Raman and XRD techniques, respectively.
Figure S2. The pressure dependence of the vibron Raman frequency of the D$_2$-rich fluid and solid in He medium as a function of pressure. Red squares are the data of this study and gray triangles are the results of previous experiments in pure bulk D$_2$. ²
**Figure S3.** Raman spectra of D₂-He mixtures at different pressures in the range of rotational-translational (a), second-order diamond (b), and vibrational (c) modes.
Figure S4. Raman spectra of H₂-rich solid formed in the mixed 50:50 H₂-He sample at 96 GPa compared to those of pure bulk H₂ at 86 and 120 GPa.
Figure S5. Pressure dependencies of the Raman frequencies of O$_2$ taken from the literature$^{3,4}$ compared to the reported by Lim and Yoo Raman frequencies of the N$_2$-He system$^5$. The data suggest that Lim and Yoo investigated the contaminated by O$_2$ system.
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