Novel Strongly Correlated Europium Superhydrides

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ABSTRACT: We conducted a joint experimental— theoretical investigation of the high-pressure chemistry of europium polyhydrides at pressures of 86–130 GPa. We discovered several novel magnetic Eu superhydrides stabilized by anharmonic effects: cubic EuH₉, hexagonal EuH₁₀, and an unexpected cubic (Pm₃n) clathrate phase, Eu₆H₄₆. Monte Carlo simulations indicate that cubic EuH₉ has antiferromagnetic ordering with Tₘ of up to 24 K, whereas hexagonal EuH₁₀ and Pm₃n-EuH₁₀ possess ferromagnetic ordering with Tₘ = 137 and 336 K, respectively. The electron–phonon interaction is weak in all studied europium hydrides, and their magnetic ordering excludes s-wave superconductivity, except, perhaps, for distorted pseudohexagonal EuH₉. The equations of state predicted within the DFT+U approach (U = J were found within linear response theory) are in close agreement with the experimental data. This work shows the great influence of the atomic radius on symmetry-breaking distortions of the crystal structures of superhydrides and on their thermodynamic stability.

The high-pressure synthesis of new compounds of metal hydrides is a rapidly developing field in view of the unusual chemistry and record-high Tₘ superconductivity of some hydrides. Some extraordinary compounds, such as CSH₉, LaH₁₀, YH₆ and YH₉, ThH₆ and ThH₁₀, UH₇, CeH₉, PrH₁₀, and NdH₁₂ have already been synthesized. Some of these hydrides, for example, superhydrides of yttrium, lanthanum, and thorium, demonstrate remarkable superconductivity. However, magnetic ordering emerging in the hydrides of many f metals at low temperatures partially or completely destroys their superconductivity (polyhydrides of praseodymium PrH₁₀ and neodymium NdH₁₂). As we have recently shown for NdH₁₂, taking into account the “spin splitting” of the electron bands near the Fermi level rules classical superconductivity out because of the high-energy gap that electrons need to overcome for s pairing. Nevertheless, the possibility of d pairing remains in such materials. On the other hand, studies of lanthanoid superhydrides are motivated by the need for further development in the description of strongly correlated systems. In the studies of the Pr–H¹¹ and Nd–H₁₁² systems, we have found that the convex hulls obtained with and without the inclusion of spin–orbit coupling (SOC) as well as with or without magnetism and the Hubbard-like correction (U = J)¹⁴ differ significantly. In this work, we continue the study of high-pressure chemistry of lanthanoid hydrides focusing on the Eu–H system.

Under normal conditions, europium has the electronic configuration [Xe]⁴f⁷⁶s² and an unusually low density and melting point, and a bcc lattice with a relatively large cell parameter of a = 4.581 Å.¹⁵ Europium is an anomalous lanthanoid: it is usually divalent and has an abnormally high atomic radius of ~200 pm, about 10% larger than those of Y, Sm, and Gd. Under pressure, it undergoes a series of phase transitions (bcc → hcp → C2/c → Pnma¹⁶) accompanied by the emergence of superconductivity above 80 GPa at 1.8–2.75 K due to the conversion of the Eu atoms from the divalent to the trivalent state with zero magnetic moment. Europium readily reacts with hydrogen, forming Pnma-EuH₂. With excess hydrogen under pressure, the reaction proceeds further, and trivalent tetragonal EuH₃–x may be obtained at about 10 GPa.¹⁶,¹⁷ In this work, we continue the study of the Eu–H system at pressures of up to 130 GPa.

Experiment. To investigate the formation of new chemical compounds in the Eu–H system at high pressures, we loaded three high-pressure diamond anvil cells (DAC nos. E1–E3) with metallic europium and sublimated ammonia borane NH₃BH₃ (AB), used as both a source of hydrogen and a pressure-transmitting medium. A tungsten gasket was pressed to 20 μm. Additional parameters of the high-pressure diamond anvil cells are listed in Supporting Information Table S1.

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Laser heating of the metal Eu sample in the AB medium at 1600−1800 K for 0.1 s yielded a complex mixture of various europium hydrides observed in DAC nos. E1 and E3. The experimental X-ray diffraction (XRD) patterns are shown in Figure 1. The analysis of the original diffraction images demonstrated the existence of a fine-grained phase with a set of reflections (inset of Figure 1b) that can be indexed in space group \( Pm\bar{3}n \) with \( a = 5.858 \) Å and \( V = 25.13 \) Å\(^3\)/f.u. at 130 GPa, corresponding to composition \( EuH_{5+x} \), \( 0 < x < 1 \). Similar diffraction patterns have been previously observed in the U−H system8 (and attributed to \( \beta\)-UH\(_3\)) and in the investigation of the Eu−H system by Ma et al.,20 where the authors attributed it to \( Pm\bar{3}n-EuH_5 \).

Two other sets of reflections may be indexed in space group \( Fm\bar{3}m \) or \( F\bar{4}3m \) (\( a = 4.947 \) Å, \( V = 30.27 \) Å\(^3\)/f.u. at 130 GPa) and \( P6_3/mmc \) (\( a = 3.591 \) Å, \( c = 5.509 \) Å, \( V = 30.76 \) Å\(^3\)/f.u. at 130 GPa), which may be proposed by analogy with the chemistry of the Pr−H11 and Nd−H12 systems (Figure 1a,b). Both compounds have a cell volume close to that of the Eu/H 1:9 composition. At high pressure (120−130 GPa, DAC no. E1), hexagonal polyhydride EuH\(_4\) is dominant, whereas a small amount of the cubic modification was found, but below 90 GPa the situation is opposite (DAC no. E3, Figure 1c,d). Changing the pressure in cell nos. E1 and E3 allowed us to get the pressure dependence of the cell volume and compare it with the theoretically calculated equations of state (EoS’s) for the obtained phases (Figures 2 and 3). As demonstrated below, this comparison makes it possible to determine the hydrogen content in the discovered compounds and confirm our guesses about EuH\(_9\). Thus, we found that \( F\bar{4}3m-EuH_9 \) is present in both diamond anvil cells in the pressure range from 86 to 130 GPa, while the formation of \( P6_3/mmc-EuH_9 \) requires higher pressure (see also Figure S2).

Additional laser heating of DAC nos. E1 at 130 GPa and E2 at 74 GPa under the same conditions (∼1600 K, 0.1 s) with consequent registration of the X-ray diffraction at synchrotron radiation facility SPring-8 (Japan, \( \lambda = 0.413 \) Å) left the diffraction pattern almost unchanged. The analysis showed that the samples still contained a mixture of cubic and hexagonal EuH\(_9\) and \( Pm\bar{3}n \)-phases (Supporting Information Figures S2 and S3 and Figures S12−S15, respectively). The reaction products were unevenly distributed over the volume of the sample: \( P6_3/mmc-EuH_9 \) (Supporting Information Figures S2 and S4, high granularity) or \( Pm\bar{3}n-EuH_{9+} \) (Supporting Information Figures S12 and S15, low granularity) may dominate other compounds. In DAC no. E2, after compressing to 89 GPa, we mainly found \( Pm\bar{3}n-EuH_{9+} \) (Supporting Information Figure S15). Thus, additional laser heating neither improved the purity of the studied mixture nor yielded any

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**Figure 1.** Experimental XRD patterns and Le Bail refinements of Eu hydrides. Le Bail refinements of \( F\bar{4}3m-EuH_9 \), \( P6_3/mmc-EuH_9 \), and \( Pm\bar{3}n-EuH_{16} \) at (a) 121 GPa and (b) 130 GPa. (Inset) Experimental XRD pattern (cell no. E1). The experimental data, model fit for the structure, and residues are shown in red, black, and green, respectively. Unidentified reflections (possibly \( I4/mmm-EuH_{16} \)) are indicated by asterisks. The reliable parameters for \( F\bar{4}3m-EuH_9 \), \( P6_3/mmc-EuH_9 \), and \( Pm\bar{3}n-EuH_{16} \) at 121 GPa are \( Rp = 21.4\% \) and \( Rwp = 35.6\% \); at 130 GPa, \( Rp = 25.3\% \) and \( Rwp = 34.6\% \). Le Bail refinements of \( F\bar{4}3m-EuH_9 \) and \( Pm\bar{3}n-EuH_5 \) at (c) 86 GPa and (d) 94 GPa for cell no. E3. The insets of panels a and c of Figure 1 show the sample chamber before and after laser heating of the cell nos. E1 and E3, respectively.

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new europium hydrides. The formation and stability of $Pm\bar{3}n\text{-}EuH_{5+}x$ was confirmed to at least 89 GPa.

Crystal Structure Search. In order to more accurately establish the composition and compute various physical properties of the synthesized phases, we performed a variable-composition search for stable compounds in the Eu–H system at pressures of 50, 100, 130, and 150 GPa using the Universal Structure Predictor: Evolutionary Xtallography (USPEX) algorithm.21−23 The spin−orbit coupling (SOC) and Hubbard-like correction term ($U−J$) were not taken into account in the evolutionary search, but were included when plotting the convex hulls (Figure 2). The $U−J$ values were found for all novel europium hydrides using the first-principles linear response approach.24 It was possible to compute $U−J$ using VASP25−27 by three single-point calculations. The obtained results (Supporting Information Table S7 and Figures S4−S7) are close to $U−J=4.5−5$ eV for all studied compounds.

To construct the convex hulls, we used the experimental XRD data from the previous studies of the phase transition of metallic europium. Bi et al.16 have shown that at 75−92 GPa europium has a $Pnma$ crystal lattice, with $a=4.977$ Å, $b=4.264$ Å, $c=2.944$ Å, and $V=15.62$ Å³/atom at 75 GPa.

Figure 2. Calculated convex hulls of the Eu–H system at (a) 50, (b) 100, and (d) 150 GPa without SOC and $U−J=0$ and (c) at 130 GPa with SOC and specific $U−J$ (see Supporting Information, Table S7).

Figure 3. Equations of state and crystal structures of synthesized Eu−H phases. (a) Pressure dependence of volumes of europium hydrides. The theoretical results taking into account SOC are shown by solid ($U−J=0$ eV) and dashed−dotted ($U−J$ values from Supporting Table S7) lines. The experimental data for EuH$_x$ from ref 20 are shown by hollow squares. Crystal structures of ideal $F\bar{4}3m$-EuH$_9$ (b) and $P6_3/mmc$-EuH$_9$ (c) at 121 GPa and $Pm\bar{3}n$-Eu$_8$H$_{46}$ (d) at 130 GPa. Purple and blue spheres represent Eu and H atoms, respectively.
Numerical simulations show that the unit cell volume \( V_{\text{Pm3m-Eu}} \) shows almost no dependence on the value of \( U - J \) in the range of 0 to 5 eV or on SOC, but the pseudopotential used in calculations strongly affects this volume: \( V_{\text{Pm3m-Eu}} = 15.82 \) and 13.49 Å\(^3\)/atom for 17-electron and 8-electron pseudopotentials, respectively. Because of much better agreement with experiment, the Eu pseudopotential with 17 valence electrons was used in all further calculations.

\( I4/mmm-EuH_\nu \), a compound from the previously found hydrides \( I4/mmm-XH_\nu \) (X = Ce, Pr, Nd), and \( \text{Im3m-EuH}_6 \) are placed on the convex hull of the Eu–H system in the experimental range of pressure from 100 to 130 GPa (Figure 2). At low pressures of \( \sim 50 \) GPa, only \( I4/mmm-EuH_\nu \) is expected, whereas higher polyhydrides are unstable. The inclusion of SOC and \( U - J \) and the analysis of the entropy factor (Supporting Information, Table S9 and Figure S10) do not change the set of stable Eu–H phases, although at 130 GPa and 0 K \( \text{EuH}_3 \) is slightly (20 meV/atom) above the convex hull. However, the analysis of the equation of state (Figure 3) points to several deviations of \( V(P) \) of the predicted europium hydrides from the experimental volumes.

First, the predicted unit cell volume of \( \text{Pm3n-EuH}_6 \) is 22.8–23.9 Å\(^3\)/f.u. at 130 GPa (without SOC, \( U - J = 0 \) eV and with SOC and \( U - J = 5 \) eV, respectively), whereas the experimental value is 25.13 Å\(^3\)/f.u., about 1 to 2 Å\(^3\) higher. The situation holds for other \( U - J = 0 \)–7 eV. To solve the problem, get closer to the experimental volume, and find a better structural solution, we increased the hydrogen content \( x \) in \( \text{Pm3n-EuH}_{1+x} \) performing two USPEX searches at 130 GPa with fixed ratios of \( \text{Eu}/\text{H} \) 1:6 and 8:46 (1:5.75, Supporting Information Table S10). The latter composition was proposed by analogy with the well-known \( \text{Pm3n} \) phases of Ge, Si, and Sn clathrates of alkali and alkaline earth metals, such as \( \text{K}_3\text{Ge}_{23} \), \( \text{Rb}_2\text{Ge}_{25} \), \( \text{Ba}_3\text{Si}_{46} \), and so forth (so-called Zintl clathrates\(^{25} \)). For the former composition, the best solution corresponds to the recently synthesized \( \text{Im3n-EuH}_6 \),\(^{26} \) however, we did not detect this compound in our experiments.

As a result of the second USPEX search, \( \text{Pm3n-EuH}_{1.46} \) was found to be the most stable phase with a cell volume of 25.21 Å\(^3\)/Eu. This structure features \( \text{H}_8 \) cages around each Eu atom (among 24 hydrogens, 8 are closest, with Eu–H distances below 1.87 Å). Among the 46 hydrogen atoms in the unit cell, 16 form pairs that can be described as stretched \( \text{H}_2 \) molecules (\( d(H-H) = 0.99 \) Å vs 0.74 in the free \( \text{H}_2 \) molecule), and the remaining 30 hydrogen atoms can be described as single atoms (their shortest H–H distance is 1.33 Å at 130 GPa). This phase corresponds to the experimental XRD and EoS, lies close to the convex hull with and without SOC and \( U - J \) (Figure 2), and can explain the experimental data for the “\( \text{Pm3n-EuH}_5 \)” phase (Figure 3) obtained by Ma et al.\(^{20} \)

Second, ideal \( \text{P6}_3/mmc-EuH_6 \) is out of the thermodynamic stability area (e.g., Figure 2c). It has been shown recently that similar behavior is observed in the Y–H system,\(^{23} \) where ideal \( \text{P6}_3/mmc-YH_6 \) is also unstable thermodynamically and distorts to stable \( \text{P1-Y}_3\text{H}_{36} \) which has almost the same XRD pattern (Figure S1). Using this structure as a prototype, we calculated the enthalpy of similar \( \text{P1-Eu}_3\text{H}_{36} \) (which may be relaxed and symmetrized to a pseudo-hexagonal \( \text{Cmcm-EuH}_9 \)) and found that this distorted structure (Supporting Information Figure S1) is more stable than the ideal \( \text{P6}_3/mmc-EuH_6 \). Similarly, \( \text{I43m-EuH}_6 \) distorts to form more stable pseudocubic \( \text{P1-EuH}_6 \).

The calculated unit cell volume of \( \text{Pm3n-EuH}_{1.46} \) is 31.8–32.4 Å\(^3\)/f.u. with \( U - J = 0 \) and 5 eV, respectively (Supporting Information Figure S8), at 130 GPa, which is 1 to 2 Å\(^3\) higher than what we found in the experiment for the cubic \( \text{EuH}_4 \) phase (Figures 1 and 3a). Therefore, the existence of \( \text{Pm3n-EuH}_{1.46} \), which is isostрукtrual to previously found \( \text{Fm3m-LaH}_{10} \), is not confirmed by the current experimental data.

It is curious to track how a series of stable europium hydrides change with pressure. At 50 GPa, all europium polyhydrides except \( \text{EuH}_4 \) are metastable (Figure 2a). An increase in pressure to 100 GPa stabilizes a wide range of polyhydrides, including the above-described \( \text{Pm3n-EuH}_{1.46} \), \( \text{I3m-EuH}_6 \), pseudocubic \( \text{P1-EuH}_{10} \), and pseudohexagonal \( \text{P1-EuH}_6 \) which is close to \( \text{Cmcm} \) (Figure 2b). Further increasing the pressure to 150 GPa leads to the stabilization of higher-symmetry structures, and distortions of ideal \( \text{I43m-EuH}_6 \) and \( \text{P6}_3/mmc-EuH_6 \) cease to play a significant role (Figure 2d). An increase in temperature to 2000 K leads to the destabilization of \( \text{EuH}_4 \) in favor of \( \text{EuH}_5 \) which may explain the unsuccessful experimental attempts to change the phase composition by
additional laser heating (Supporting Information Figure S10). Anharmonic calculations based on molecular dynamics and machine learning potentials of interatomic interactions (details in the Supporting Information) show that ideal P6$_3$/mmc-EuH$_9$, F43m-EuH$_9$, and Pm3n-EuH$_{46}$ are dynamically stable, although in the harmonic approximation they should undergo distortions (Supporting Information Figures S26 and S29–S31). Having predicted the stable compositions and structures of europium hydrides, we now want to gain insight into their nature.

**Superatomic Orbitals and Chemical Bonding.** The description of electronic structure in terms of delocalized canonical orbitals is almost completely devoid of an explicit chemical bonding picture of a given system. To understand the chemical bonding in europium polyhydrides, we first analyzed model clusters (blocks separated from the infinite crystal lattice) via the adaptive natural density partitioning (AdNDP) algorithm. The AdNDP is a localization technique that follows general ideas of the natural bond orbital (NBO) analysis proposed by Weinhold and Landis. The main advantage of this method is the possibility to represent a chemical bonding pattern in terms of both Lewis bonding elements (lone pairs, two-center two-electron (2c–2e) bonds) and delocalized bonding elements (nc–2e bonds) by partial diagonalization of the one-body reduced density matrix. This technique is widely used in materials science and in describing various Zintl clusters.

The chemical bonding analysis can offer insights into the reasons for the stability, electronic properties, magnetism, and chemical activity of investigated systems. It has been shown that the chemical bonding analysis of cluster models qualitatively agrees with the analysis of solid-state structures. In this work, we used EuH$_{26}$ as a model of P6$_3$/mmc-EuH$_9$, EuH$_{26}$ as a model of Pm3n-EuH$_{46}$, and EuH$_{26}$ as a model of F43m-EuH$_9$ (Figure 4). Charges were chosen so as to adjust the total number of electrons per Eu atom in a studied fragment and consider those electrons shared between the cluster and the neighboring atoms.

Starting our search from localized bonding elements such as 2c–2e, 3c–3e, and so forth, we found no highly occupied localized bonds in the present structures. Because of the absence of highly occupied localized bonding elements, the model clusters could be completely described in terms of fully delocalized canonical molecular orbitals (MOs). As was expected, for such sphere-like clusters, the canonical MOs mimic the spherical harmonics forming so-called superatomic orbitals. Further analysis of the superatomic orbitals of corresponding clusters reveals that unpaired electrons are located mostly on the F-type orbitals (Figure 4a–f), whereas the S-, P-, and D-type orbitals are doubly occupied.

Following the analysis of the model clusters, we performed the solid-state AdNDP (SSAdNDP) calculations for ideal cubic and hexagonal EuH$_9$ and Zintl-like clathrate Pm3n-EuH$_{46}$. Such an analysis was first implemented for polyhydrides in the current work. SSAdNDP revealed the same bonding picture as was found in the model clusters. A cognate pattern could also be seen in the comparison of the electron localization function (ELF) plots of the solid-state structures and model clusters (Supporting Information Figure S19). The corresponding occupancies of valence superatomic orbitals are presented in Supporting Information Table S6. For all europium polyhydride structures, there are seven almost ideally (1.00e, where e is the elementary charge) singly occupied superatomic F orbitals, which are responsible for the magnetic properties.

The corresponding nc–2e occupancy numbers (where n is the number of hydrogen atoms in the cage) of superatomic D and P orbitals are (1.72–1.39)e for P6$_3$/mmc-EuH$_9$, (1.17–1.09)e for Pm3n-EuH$_{46}$, and (1.57–1.38)e for F43m-EuH$_9$. Because of the delocalization of conducting electrons, which is common for metals, these values are about (0.3–0.9)e lower than those that we found for clusters.

To understand the charge distribution in the studied structures, Bader charge analysis was performed. In all structures, the Eu atom bears a positive charge of +1.15e, whereas a partial negative charge of −0.31e to −0.08e was found for all hydrogen atoms. This behavior is common for metal hydrides, and similar results have been found in previous works. The average Bader charges are presented in Supporting Information Table S18.

Both cubic and hexagonal EuH$_9$ as well as Eu$_8$H$_{46}$ demonstrate a large magnetic moment (absolute value) in the unit cell (Figure S7), slightly decreasing as the pressure increases. At 100 GPa, the magnetic moment of EuH$_9$ is 3.96μ$_B$ per Eu atom for the P6$_3$/mmc modification, 6.32μ$_B$ per Eu atom for the Pm3n-EuH$_{46}$ structure, and about 5.88μ$_B$ per Eu atom for cubic EuH$_9$. The magnetization arises mostly from the f electrons of the Eu atoms. We observed the same behavior for the corresponding clusters (Supporting Information Table S5): the ground spin state of EuH$_{27}^{17+}$ and EuH$_{24}^{17+}$ is nonet (S$_{\text{total}}$ = 4), whereas for EuH$_{26}^{19+}$ the ground spin state is septet (S$_{\text{total}}$ = 3).

We performed a comparative analysis of superatomic orbitals of europium hydrides, recently synthesized cubic and hexagonal PrH$_9$ and hexagonal NdH$_9$ (Supporting Information Table S17). For PrH$_9$, the shapes of five obtained nc–2e bonds do not completely correspond to the D type: two orbitals have a D-type nature, and the three other nc–2e bonds have the F-type nature. This probably happens because of the ambiguous behavior of f and d orbitals of lanthanoids. Similar behavior—the mixing of F and D orbitals—was also observed for the molecular clusters of these hydrides and for the NdH$_{16}^{14+}$ cluster, which has D-type and F-type orbitals with very similar energies.

The analysis of MOs revealed that all clusters could be described within the superatomic concept, according to which the most stable clusters emerge from completely filled superatomic shells. However, CeH$_{32}$ (Supporting Information Figure S18) has an additional 2P electron that contradicts this concept. We believe that this P electron plays an important role in the metallic and superconducting (T$_C$ ≈ 110 K (unpublished)) properties of cerium polyhydrides as was previously shown for LaH$_{10}$. The overall approach of the localization of superatomic orbitals in hydrogen cages could also be extended to other superconducting polyhydrides. For example, strong localization of the multicenter bond was found for H$_{32}$ cages present in LaH$_{10}$ and YH$_{10}$ superconductors. As in the case of Ce polyhydride, 2P electrons in these structures were found to have relatively high occupation numbers. The 2S superatomic orbital, which is present in clusters, is absent in solid-state superatoms. We think that these two electrons, localized on the H cage of clusters, may participate in metallic interactions and cannot be localized via the SSAdNDP technique. The detailed comparison of occupancies of the superatomic orbitals in different polyhydrides is presented in Supporting Information Table S17.

**Magnetic Structure.** To analyze the magnetic structure of europium hydrides, we studied a series of ferromagnetic (FM)
and antiferromagnetic (AFM) configurations. The enthalpies of the optimized structures are presented in Supporting Information Table S11. During the calculation, the ideal cubic and hexagonal structures of hydrides are distorted into more stable low-symmetry pseudocubic and pseudohexagonal modifications. Most of these distorted structures have XRD pattern close to the ideal ones. To simplify the magnetic calculations, in several cases we used high-symmetry EuH₉ prototypes (F4₃m and P6₃mmc) instead of thermodynamically more stable distorted P1-Eu₄H₃₆. The magnetic structure of the pseudohexagonal Cmcm-EuH₉ has nevertheless been studied in detail (Supporting Information).

The most stable collinear magnetic state of F4₃m-EuH₉ is AFM 1, which distorts to P1 during relaxation. The second most stable configuration is AFM 3, which distorts to Immm2. Finally, among the magnetic states that preserve the original ideal F4₃m symmetry, the most stable is AFM 2. The same is observed for P6₃mmc-EuH₉. The most stable collinear magnetic state is AFM 3, but during relaxation the hexagonal structure is distorted and transformed to pseudohexagonal Cmcm, which lies on the convex hull (Figures 2 and 4). The second most stable configuration is AFM 4, which distorts to C2. Among the magnetic states that preserve the ideal P6₃/mmc symmetry, the most stable is FM. The most stable collinear configuration for Eu₈H₄₆ is FM. Unlike the EuH₉ phases, the magnetic configurations of Eu₈H₄₆ do not show significant distortions from the ideal Pm3n geometry after relaxation.

For each of our three phases, we used the relaxed geometries of the most stable magnetic states to study magnetic anisotropy by running calculations with SOC. We computed the single-point enthalpy of these structures in both the FM and in one AFM state, with the magnetic moments aligned along different directions (x, y, z, xy, xz, and yz). The results are summarized in Supporting Information Table S12. We expect a small amount of magnetic anisotropy for all phases with EuH₉ stoichiometry, for which we identified the most stable orientation of the magnetic moments. We do not expect any magnetic anisotropy in Pm3n-Eu₈H₄₆ because different orientations of the magnetic moments have almost the same enthalpy up to the numerical errors.

To study the Néel and Curie temperatures of F4₃m-EuH₉, P6₃/mmc-EuH₉, and Pm3n-Eu₈H₄₆ we modeled the magnetic interaction within the Ising Hamiltonian and obtained the critical temperatures for both EuH₉ and Eu₈H₄₆ in our Heisenberg model from a Monte Carlo simulation as implemented in the VAMPIRE code (Supporting Information Table S14). We obtained the critical temperature Tₙ = 24 K for cubic EuH₉, while hexagonal EuH₉ and Pm3n-Eu₈H₄₆

Figure 5. Electron density of states (DOS) of europium hydrides at 130 GPa. (a, c, e) Contributions of different spin orientations to the total DOS. (b, d, f) Contributions of the Eu and H atoms to the total DOS. The DOS of the Eu sublattice separated from hydrogen at 130 GPa is shown in red; the DOS of the Hsub lattice separated from the Eu atoms at 130 GPa is shown in blue.
possess ferromagnetic ordering with $T_C = 137$ and 336 K, respectively.

**Electron–Phonon Interaction.** Strong electron–phonon coupling (EPC) is expected for lanthanides and their polyhydrides under pressure. It has been shown for superhydrides of Ce, Pr, Nd, Y, and Th. The superconducting (SC) properties and the strength of the electron–phonon interactions decrease as the number of $f$ electrons increases and manifestations of magnetism become more pronounced.

The numerical analysis using the UppSC code12 performed for hexagonal NdH$_6$ (120 GPa)12 showed that if the effective spin splitting in the band structure $\Delta(k) = (T - T_i) / 2$, where $T$ is the spin-resolved electron dispersion, is about 0.5 eV or higher then superconductivity will be completely suppressed at all $\mu^* \geq 0$. We found that a similar spin splitting gap reaches 4–4.5 eV in both EuH$_6$ and Eu$_3$H$_{46}$ (Figure S5a, c, e), which is 2 orders of magnitude larger than for all known SC gaps. This fact excludes singlet superconductivity and $s$ pairing in europium hydrides.

Despite this pessimistic conclusion, we estimated the parameters of the electron–phonon interaction in europium hydrides as if magnetism were absent. The results show that the distorted H sublattice in both modifications of EuH$_6$ could be a superconductor with a $T_C$ of up to 21 K ($\mu^* = 0.1$), an EPC coefficient $\lambda = 0.61$, and $\omega_{\text{ph}} = 386$ K for hexagonal EuH$_6$ and $T_C = 27$ K and $\lambda = 0.51$ for cubic EuH$_6$. Compared with LaH$_{10}$ ($T_C \geq 250$ K$^{3,4}$), this indicates the enormous role played by the metal atom (even in the absence of magnetism) and surprisingly different electron–phonon coupling properties of the otherwise very similar lanthanoids. The crucial role of the hydride-forming element and its electronic structure in the $T_C$ of the hydride has been discussed before.84

The projection of the total density of states of both EuH$_6$ on the H atoms, $N_H(E_F)$, is 0.41–0.48 states/eV/Eu (Figure Sb, df) at 130 GPa. An estimate of the logarithmic frequency made on the basis of the experimental equation of state yields $\omega_{\text{ph}} = 975–1060$ K for EuH$_6$ at 100 GPa, agreeing with calculations. Both of these factors would be favorable to superconductivity if electron–phonon coupling were stronger and if there were no magnetic ordering. An exception may exist for pseudohexagonal Cmcm-EuH$_6$, which does not exhibit magnetism at finite temperatures (Supporting Information).

Compared to PrH$_6$ and NdH$_6$, $f$ electrons in europium hydrides are localized much deeper (2 eV) below the Fermi level, and $N(E_F)$ is ~0.95–1.17 states/eV/Eu in EuH$_6$, similar to the values for LaH$_{10}$ and YH$_6$ with a high contribution of the H sublattice (~50% of $N(E_F)$). If we consider the metal and hydrogen sublattices in EuH$_6$ separately, disregarding the interaction between them (Figure Sb, df), then we will see that the influence of this interaction is completely negative for both $N_M(E_F)$ and $N_H(E_F)$, which decrease by 1.8–2-fold. The calculation of the superconducting properties of the formally separated Eu and H sublattices of $F^{-43}m$-EuH$_6$ at 130 GPa shows that the critical temperature of hydrogen (“$H_2$”) in the cubic sublattice reaches 150–200 K, while interaction with Eu leads to about a 5 times lower $T_C$ of ~27 K. Thus, the metal lattice plays the role of a stabilizer of metallic hydrogen but reduces the density of states $N_M(E_F)$ at the Fermi level and the critical temperature of superconductivity.

In conclusion, continuing the studies of the stabilization of metallic hydrogen in compressed metal–hydrogen systems, we synthesized three novel europium superhydrides with probably slightly distorted structures due to the abnormally large radius of the Eu atom: $P6_3/mmc$-EuH$_{10}$, $F^{-43}m$-EuH$_{10}$, and an unexpected clathrate phase, $Pm3n$-EuH$_{46}$. The discovered cubic and hexagonal modifications of EuH$_9$ are the new members of the nonahydrde family, which currently includes the polyhydrides of Th, U, Y, Ce, Pr, and Nd. Similar to the Nd–H system, all europium hydrides are magnetic and strongly correlated systems with a significant contribution of the SOC interaction. They possess antiferromagnetic (cubic EuH$_6$) or ferromagnetic (hexagonal EuH$_6$, Eu$_3$H$_{46}$) ordering with $T_N \approx 24–336$ K. Although the projection of the DOS on the H sublattice in both phases of EuH$_6$ is similar to that of high-$T_C$ superconducting hydrides, such as LaH$_{10}$ and YH$_6$, the electron–phonon coupling is weak. Magnetic order and a large effective spin splitting (>4 eV) in the electron band structure of EuH$_6$ make classical $s$-wave superconductivity impossible. An exception may take place for pseudohexagonal Cmcm-EuH$_6$, which does not exhibit magnetism at finite temperatures.

The chemical bonding analysis was implemented for polyhydrides for the first time. The analysis points to the contribution of 2P superatomic electrons as an important distinguishing feature in determining high-temperature superconductivity. It was found that the superatomic description of polyhydrides could be extended to other important superconductors, such as LaH$_{10}$ and YH$_6$.

By analyzing the whole series of La–Ce–Pr–Nd–Eu superhydrides, we conclude that the role of magnetism is gradually increasing in this row, which leads to the suppression of $s$-wave superconductivity. Most of the higher superhydrides of these metals have antiferromagnetic ordering, whereas the lower hydrides have ferromagnetic ordering. Europium hydrides, in addition to YH$_{36}$ and BaH$_{12}$, give us yet another indication of the great role of distortions of the ideal hexagonal and cubic structures in their stabilization. Without considering these distortions, a theoretical explanation of the thermodynamic stability of europium superhydrides would be impossible.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcl.0c03331.

Detailed descriptions of the used experimental and computational methods; structural information on the predicted and synthesized structures; additional data for the Le Bail refinements; information on the magnetic structure, natural population analysis, electronic structure, phonons, anharmonic calculations, and elastic properties (PDF)

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D.V.S., D.Z., and A.G.K. contributed equally to this work. X.H. and A.R.O. conceived this project. D.Z., D.V.S., I.A.T., A.G.I., A.G.G., and X.H. performed the experiment. D.V.S., D.Z., A.R.O., and T.C. prepared the theoretical calculations and analysis. M.G. studied the magnetic properties. A.R.O. and T.C. wrote and revised the paper. All authors discussed the results and offered useful ideas.

Notes
The authors declare no competing financial interest.

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