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High-Pressure Synthesis of Magnetic Neodymium Polyhydrides

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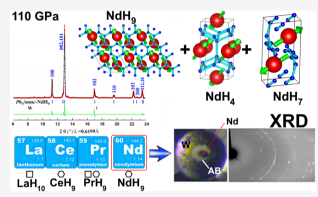
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ABSTRACT: Ongoing search for room-temperature superconductivity is inspired by the unique properties of the electron—phonon interaction in metal superhydrides. Encouraged by the recently found highest- $T_{\rm C}$ superconductor fcc-LaH₁₀, here we discover several superhydrides of another lanthanoid, neodymium. We identify three novel metallic Nd—H phases at pressures ranging from 85 to 135 GPa: I4/mmm-NdH₄, C2/c-NdH₇, and $P6_3/mmc$ -NdH₉, synthesized by laser-heating metal samples in NH₃BH₃ media for in situ generation of hydrogen. A lower trihydride Fm3m-NdH₃ is found at pressures from 2 to 52 GPa. I4/mmm-NdH₄ and C2/c-NdH₇ are stable from 135 to 85 GPa, and $P6_3/mmc$ -NdH₉ is stable from 110 to 130 GPa. Measurements of the electrical resistance of NdH₉ demonstrate a possible superconducting transition at \sim 4.5 K in $P6_3/mmc$ -NdH₉. Our theoretical calculations predict that all of the



neodymium hydrides have antiferromagnetic order at pressures below 150 GPa and represent one of the first discovered examples of strongly correlated superhydrides with large exchange spin-splitting in the electronic band structure (>450 meV). The critical Néel temperatures for new neodymium hydrides are estimated using the mean-field approximation to be about 4 K (NdH_4), 251 K (NdH_7), and 136 K (NdH_9).

■ INTRODUCTION

Since 1968, it is widely discussed that dense metallic hydrogen, if ever produced, could be a high-temperature superconductor. The main reason is its very high Debye temperature (due to low atomic mass) and very strong electron-phonon coupling.^{2,3} However, as creating metallic hydrogen requires immense pressures of ~500 GPa, 4-6 a confirmation of high- $T_{\rm C}$ superconductivity in pure hydrogen is still pending. Instead, in the search for hydrogen-induced hightemperature superconductivity, most researchers have turned to hydrogen-rich hydrides.⁷ Hydrides are promising for hightemperature superconductivity⁸⁻¹⁰ under lower pressure as compared to pure metallic hydrogen. Particularly, hydrides are convenient for experimental investigations because of a very high diffusion coefficient of atomic hydrogen, which facilitates the formation of new chemical compounds and makes it possible to perform laser-assisted chemical synthesis in diamond anvil cells in milliseconds.¹¹

Just a few years ago, a series of theoretical and experimental studies of various metal hydrides demonstrated that heavy elements relatively easily form metallic superhydrides (i.e., hydrides containing more hydrogen than expected on the basis of atomic valences) at pressures below 200 GPa, such as fcc-LaH₁₀, 12,13 P4/nmm-LaD₁₁₋₁₂, 10 P6₃/mmc-ThH₉ and $Fm\overline{3}m$ -ThH₁₀, 14 P6₃/mmc-UH₇, $Fm\overline{3}m$ -UH₈, 15 P6₃/mmc-CeH₉, 16,17 and P6₃/mmc- and $F\overline{4}3m$ -PrH₉, 18 whereas lighter atoms tend

to form molecular semiconducting hydrides with low symmetry, for instance, ${\rm LiH_6}^{19}$ and ${\rm NaH_7}^{20}$ Extremely high values of $T_{\rm C}$ have been predicted for MgH₆, 21 CaH₆, 22 and YH₁₀, 23 but these compounds have not been synthesized yet.

One of the most important properties of high-symmetry polyhydrides is a very strong electron-phonon interaction. The critical temperature (>250 $K^{9,10}$) and upper critical magnetic field (H_C up to 140 T) achieved in fcc-LaH₁₀ have already surpassed by far the parameters of other known compounds, for example, cuprates.²⁴ Analysis of theoretical results in the field shows that the increasing number of d- and f-electrons in the electron shells of a hydride-forming atom leads to the enhancement of magnetism, which can suppress superconductivity, and increasing relative contribution of dand f-orbitals to the total electron density of states at the Fermi level $N(E_{\rm F})$ is associated with weakening of electron-phonon coupling.²⁵ Thus, the main driver of the study of lanthanoid hydrides is their unusual crystal structures and possible interplay between the classical phonon-mediated superconductivity and magnetic ordering.

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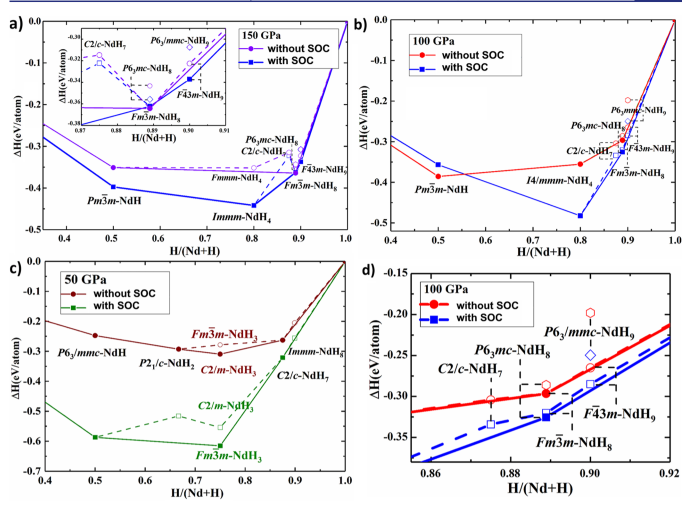


Figure 1. Thermodynamic convex hulls of the Nd-H system calculated with and without spin-orbit coupling (SOC) and magnetism at (a) 150 GPa, (b and d) 100 GPa, and (c) 50 GPa. The enlarged part of (b) at 100 GPa is shown in (d).

Currently, high-pressure experimental studies of lanthanoid hydrides are not sufficiently developed, and further investigations of the synthetic pathways and crystal structure of these compounds will contribute to building a deeper understanding of the chemistry and physical properties of metal hydrides. Previously, Chesnut and Vohra²⁶ studied the crystal structure of metallic Nd at pressures up to 150 GPa and determined the phase sequence occurring as the pressure increases: $dhcp \rightarrow fcc \rightarrow dfcc \ (hR24) \rightarrow hP3 \rightarrow monoclinic \rightarrow$ α -U. Like other lanthanides, Nd can readily absorb hydrogen at high temperatures and form hydrides: NdH2 with cubic closepacked and NdH3 with hexagonal close-packed Nd sublattice were found at ambient pressure.²⁷ Continuing the investigations of lanthanide-hydrogen systems, we study here the crystal structures and properties of compounds in the Nd-H system in the pressure range of 0-140 GPa. Three novel superhydrides, NdH₄, NdH₇, and NdH₉, are synthesized, which display a fascinating combination of magnetism and superconductivity.

RESULTS AND DISCUSSION

Stable Phases Predicted by Theoretical Calculations. Before the experiment, we carried out several independent variable-composition searches for stable compounds in the Nd–H system at 50, 100, and 150 GPa using the USPEX^{28–30} code. The results of the structure search (Figure 1) exhibit

large differences depending on the inclusion or exclusion of spin-orbit coupling (SOC) and magnetism. Results with SOC and magnetism indicate that Pm3m-NdH, tetrahydride Immm-NdH₄, cubic Fm3m-NdH₈, and F43m-NdH₉ superhydrides are stable compounds at 150 GPa, while C2/c-NdH₇ lies ~0.05 eV/atom above the convex hull. P63/mmc-NdH9, which is similar to recently discovered CeH₉ and PrH₉, lies 0.035 eV/ atom above the convex hull (Figure 1a). At 100 GPa (Figure 1b,d), we see stabilization of the I4/mmm modification of NdH₄ and appearance of P6₃mc-NdH₈ on the convex hull. At the same time, C2/c-NdH₇ becomes closer (~ 0.015 eV/atom) to the convex hull and becomes stable at about 80 GPa. The calculations at 50 GPa without SOC and magnetism show that P6₃/mmc-NdH, P2₁/c-NdH₂, C2/m-NdH₃, and C2/c-NdH₇ are stable (Figure 1c). However, inclusion of spin-orbit coupling leads to dramatic changes in the set of stable phases: P2₁/c-NdH₂ disappears, while Fm3m-NdH₃ becomes thermodynamically stable. Molecular polyhydride C2/c-NdH₇ maintains its stability at 50 GPa.

Experimental Synthesis of Atomic I4/mmm-NdH₄ and **Molecular** C2/c-NdH₇. The synthesis of neodymium hydrides was performed in a diamond anvil cell (DAC) that we denoted as Z1, containing a piece of Nd compressed in the NH₃BH₃ medium to 94 GPa and heated to 1700 K, according to the reaction: Nd + NH₃BH₃ \rightarrow NdH_x + c-BN. The Raman signal of H₂ was detected at 4180.9 cm⁻¹, and the

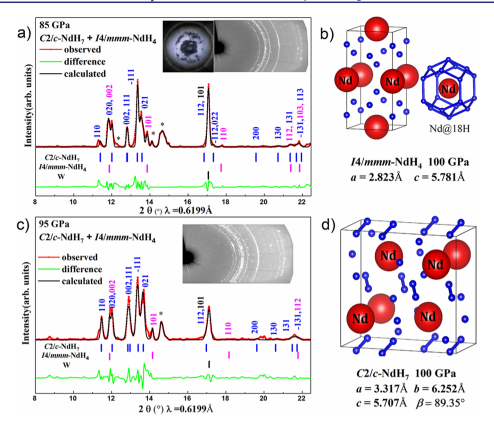


Figure 2. Experimental X-ray diffraction (XRD) patterns and Le Bail refinement of I4/mmm-NdH₄ and C2/c-NdH₇ at (a) 85 GPa and (c) 95 GPa. The experimental data and model fit for the structure are shown in red and black, respectively; the residues are indicated in green, while the unexplained peaks are marked by asterisks. The crystal structures of I4/mmm-NdH₄ and C2/c-NdH₇ at 100 GPa are shown in (b) and (d), respectively.

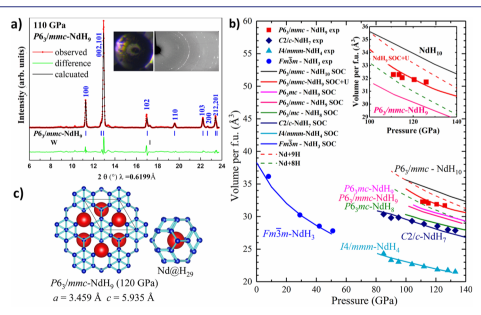


Figure 3. (a) Experimental XRD pattern and Le Bail refinement of $P6_3/mmc$ -NdH₉. The experimental data and model fit for the structure are shown in red and black, respectively; the residues are indicated in green. (b) The equation of state of the synthesized Nd–H phases; theoretical results include spin—orbit coupling and magnetism. Inset: The distinction between $P6_3mc$ -NdH₈, $P6_3/mmc$ - and $P6_3/mc$ -NdH₉, and $P6_3/mmc$ -NdH₁₀ phases. (c) The crystal structure of $P6_3/mmc$ -NdH₉ and parameters of the unit cell.

corresponding pressure is 96 GPa. The rather complex diffraction pattern observed after laser heating shows that the reaction products are dominated by C2/c-NdH₇ with a small impurity of tetragonal I4/mmm-NdH₄ (Figures 2 and S1). Exploring this sample, we determined the experimental

equation of state (EoS) of C2/c-NdH $_7$ and I4/mmm-NdH $_4$ in the pressure range from 85 to 135 GPa (Figure 3b) and found close agreement with the EoS obtained by DFT calculations.

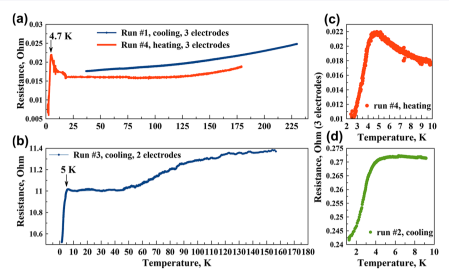


Figure 4. Dependence of electrical resistance of the synthesized sample on the temperature: (a, c, and d) in three-electrodes geometry; (b) in two-electrodes geometry. Runs 1–4 correspond to sequential steps of the laser heating.

In I4/mmm-NdH₄, which is isostructural to the recently found I4/mmm-ThH₄, 14,34 CeH₄, 17 and CaH₄, 35 the shortest H–H distance is $d_{\min}(H-H)=1.55$ Å at 100 GPa. Each Nd atom is bounded to 10 H atoms with d(Nd-H)=2.02-2.08 Å. The hydrogen sublattice in NdH₄ is represented by the atomic hydrogen with a relatively high contribution of 1s-electrons to the electronic density of states $N(E_F)$. The experimental cell parameters of the discovered compounds are shown in Table S4. Fitting the experimental pressure—volume data in the pressure range from 85 to 135 GPa by the third-order Birch—Murnaghan equation of state gives $V_{100}=45.7(2)$ ų, $K_{100}=525(\pm15)$ GPa, and $K_{100}'=2.7(\pm1.2)$.

The experimentally discovered C2/c-NdH $_7$ structure, which is close to the recently predicted C2/m-AcH $_8$ ³⁶ and $P2_1/m$ -ThH $_7$,³⁴ at 100 GPa contains quasimolecular hydrogen H $_2$ units with d(H-H)=0.92 Å. The remaining hydrogen (\sim 30%) is in the atomic form. Each Nd atom is bounded to 17 H atoms with d(Nd-H)=1.98-2.06 Å. The experimental cell parameters of this phase are shown in Table S5. Fitting the experimental pressure–volume data in the pressure range from 85 to 135 GPa by the third-order Birch–Murnaghan equation of state gives $V_{100}=118.3(1)$ Å 3 , $K_{100}=522(\pm23)$ GPa and $K_{100}'=2.4(\pm1.5)$.

Synthesis of Superhydride $P6_3/mmc$ -NdH₉. The synthesis of higher neodymium hydrides was performed in a Z2 diamond anvil cell with Nd particle compressed in the NH₃BH₃ medium to 113 GPa and heated to 1800 K. The Raman signal of H₂ was detected at 4142.9 cm⁻¹, and the corresponding pressure is 115 GPa. The obtained diffraction pattern (Figure 3a, Figure S2b, and Table S6) corresponds to hexagonal $P6_3/mmc$ -NdH₉, the structure of which is very close to the previously described hexagonal $P6_3/mmc$ -ThH₉. (Figure 3c). Both known theoretical investigations of the high-pressure chemistry of the Nd–H system 13,25 claim the existence of cubic NdH₈ and NdH₉, while the experiment shows the presence of only one unexpected hexagonal $P6_3/mmc$ -NdH₉ phase.

The calculated volumes of the ideal $P6_3/mmc$ -NdH₉ structure are close to the experimental values, although the theoretical cell parameters have some deviations from the observed ones, for instance: $a(\exp) = 3.639$ Å, a(theory) = 3.459 Å, $c(\exp) = 5.560$ Å, and c(theory) = 5.935 Å at 120

GPa. These deviations, as well as the stability of hexagonal NdH₉, can be almost completely explained by DFT+U approach with U-J=5 eV, chosen by using linear-response calculations (inset in Figure 3b; see the Supporting Information for details) and introduced for describing correlation effects. Note that such values of U-J are commonly used for modeling f-electrons in Nd. $^{37-41}$ Additional ab initio studies show that the $P6_3/mmc$ -NdH₉ structure is dynamically stable (Figure S5).

Because of the visible difference in cell parameters, we also considered the possible presence of additional hydrogen in the structure: NdH_{9+x} (x=0-0.5), where the degree of nonstoichiometry (x) was determined by a linear interpolation of the dependence of the cell volume on the H content between 9 and 10 H atoms per Nd atom. In addition, we investigated $P6_3mc$ -NdH₈ and $P6_3/mmc$ -NdH₁₀ structures as possible candidates. NdH₈ is similar to predicted $P6_3mc$ -PrH₈ and dynamically stable, lying on the convex hull at 100 GPa (Figure 3b). However, its predicted volume $V_{120}(NdH_8) = 29.04 \, \text{Å}^3$ is much smaller than the experimental value of 31.88 Å³. The proposed $P6_3/mmc$ -NdH₁₀ is thermodynamically and dynamically unstable, and its predicted cell volume differs significantly from the measured one (Figure 3b).

NdH₉ has the same hexagonal structure as the whole family of reported hexagonal superhydrides: YH₉, CeH₉, PrH₉, and ThH₉. In $P6_3/mmc$ -NdH₉, H₂₉ cages have the nearest H–H distance of 1.272 Å, the longest among all previously studied lanthanoid polyhydrides at 120 GPa, while the nearest Nd–H distance at this pressure is 1.973 Å. The experimental cell parameters of the compound are listed in Table S6. Fitting the experimental pressure—volume data in the range from 110 to 126 GPa using the third Birch—Murnaghan equation of state with fixed $K_{100}' = 4$ gives $V_{100} = 32.8(1)$ Å³ and $K_{100} = 719(\pm 41)$ GPa.

After the destruction of diamonds in the Z2 DAC, followed by decompression from 106 to 51 GPa, the recorded XRD patterns demonstrated the presence of only one hydride phase: the metallic and magnetic $Fm\overline{3}m$ -NdH₃ (Figure S2c) with the experimental cell parameter a=4.814 Å at 50 GPa, in agreement with the earlier predictions. Fitting the experimental pressure—volume data for this phase in the pressure range from 7 to 50 GPa using the third-order Birch—

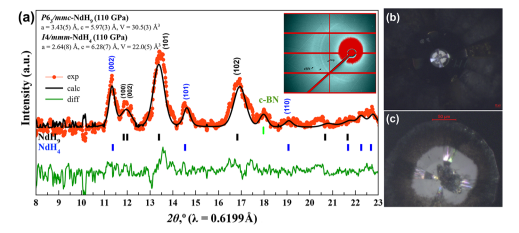


Figure 5. (a) X-ray diffraction pattern and Le Bail refinement of the NdH₉ and NdH₄ structures found in the electrical cell at 110 GPa. Reflection from c-BN (a = 3.441 Å) corresponds to \sim 75 GPa; (b,c) optical microscopy of the culet of four-electrodes electrical cell and sample after laser heating.

Murnaghan equation of state gives $V_0 = 42.3(3) \text{ Å}^3$, $K_0 = 48.4(9)$ GPa, and $K_0' = 4.5(0)$.

Measurements of the Electrical Resistance of P6₃/ mmc-NdH₉. The study of possible superconductivity in neodymium polyhydrides was performed in a Z3 cell with four 500 nm thick molybdenum electrodes prepared by magnetron sputtering (E = 200 V) and UV lithography on a diamond culet of 80 μ m in diameter. The cell was loaded with a 30 μ m particle of Nd and ammonium borane layer (thickness \sim 15–20 μ m) and then compressed to a pressure of 110 GPa using c-BN/epoxy insulating gasket. After pulsed laser heating of the sample during 3 s (total of 4 runs) over 1600 K and subsequent cooling in a cryostat, the electrode system was partially damaged, and several electrodes sequentially lost contact with the sample. In these cases, three-electrode (pseudo van der Pauw) and two-electrode schemes were used (Figure 4). As a result of the resistance (R) measurements in the range of 1.6-230 K, we have detected a sharp reproducible drop of R(T) at 4.5 \pm 0.5 K, possibly caused by the superconducting transition in NdH₉. At the same time, no clear and reproducible transitions or resistance drops were detected above 5 K.

To study the phase composition of the synthesized sample in the electrical cell, X-ray diffraction was performed using Beijing Synchrotron Research Facilities (BSRF, China). The wavelength of the synchrotron radiation was $\lambda=0.6199$ Å, with an exposure time of 10 min and a beam diameter of ~50 μ m. Because of the presence of the electrodes, beam widening, and its weak intensity, the X-ray diffraction pattern cannot be quantitatively interpreted; however, the obtained diffractogram (Figure 5) qualitatively corresponds to a distorted hexagonal $P6_3/mmc$ -NdH_{9-x} mixed with tetragonal I4/mmm-NdH₄, the volume of which is close to the predicted value.

Magnetic Properties of Neodymium Hydrides. Ab initio calculations show that $P6_3/mmc$ -NdH₉ exhibits metallic properties and magnetism (Figure S6). The contribution of hydrogen atoms to $N(E_{\rm F})$ is very low: 7.44 eV⁻¹ f.u.⁻¹ (97%) comes from Nd, while hydrogen gives only 0.22 eV⁻¹ f.u.⁻¹ (3%) at 120 GPa. It is expected that such a high electron density at the Fermi level drives an instability against spontaneous magnetization (see Stoner criterion⁴²). Despite a clear analogy between Nd–H and Pr–H¹⁸ systems (hexagonal XH₉ and tetragonal XH₄ polyhydrides), there is a

significant difference in magnetic properties associated with the number of f-electrons. All Nd—H phases demonstrate strong magnetic properties (Figure 6a and b), more pronounced than

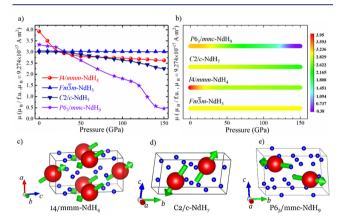


Figure 6. Magnetic properties of Nd–H compounds. Magnetism of Nd hydrides at pressures up to 150 GPa. (a) Magnetic moments (per Nd atom) at various pressures and (b) magnetic map of the Nd–H system as a function of pressure. Computed spin configurations for the $2\times2\times1$ supercell of NdH₄ (c) and unit cells of NdH₇ (d) and NdH₉ (e). Arrows indicate magnetic moment directions on the Nd-

those of the corresponding praseodymium hydrides, except recently discovered I4/mmm-PrH₄, 43 which also possesses AFM ordering at 100 GPa (see Tables S8 and S9). This, probably, excludes the possibility of a classical s-wave superconductivity due to the large exchange spin-splitting in electron band structure (>450 meV). On the other hand, the coexistence of magnetism and a relatively strong electron—phonon interaction leaves open the possibility for more complex mechanisms of superconductivity as found, for example, in cuprates and Fe-containing pnictides. 24,44

Simple spin-polarized calculations show that NdH₇ and NdH₄ maintain the magnetic moment of ~2.5–3.5 $\mu_{\rm B}$ per Nd atom at low pressures, then start to slowly lose magnetism as pressure increases, while cubic NdH₃ keeps an almost constant magnetic moment, around 3 $\mu_{\rm B}$ per Nd atom in the pressure range from 0 to 150 GPa. The magnetization of NdH₉ rapidly

decreases with rising pressure and vanishes at \sim 230 GPa (Figure 6a,b).

To understand the exact spin configurations at finite pressure, we have employed 10 different spin configurations using single-unit cells for NdH₇ and NdH₉, and 13 spin configurations and a 2 × 2 × 1 supercell for NdH₄. Using noncollinear calculations, the magnetic anisotropy energy (MAE) for Nd–H phases along different directions has been computed. Each magnetic configuration has been calculated at fixed lattice parameters and corresponding pressure. The results summarized in Table S8 clearly indicate the antiferromagnetic (AFM) character of NdH₄, NdH₇, and NdH₉ compounds (Figure 4c–e and Tables S8 and S9).

We have identified that NdH₄, NdH₇, and NdH₉ compounds possess AFM collinear [112], [144], and [23 $\overline{1}$] orders, respectively. The Néel temperature can be estimated using the mean-field approximation $T_{\rm N}^{\rm MF} = \sum_{i,j} \frac{I_{i,j}S^2}{3k_{\rm B}} \approx \min \frac{|E_{\rm FM} - E_{\rm AFM}|}{6k_{\rm B}}$ as ~5 K (NdH₄), ~251 K (NdH₇), and ~136 K (NdH₉). As it will be demonstrated below, the Néel temperatures are significantly higher than the expected superconducting $T_{\rm C}$.

Electron–Phonon Interaction in Neodymium Hydrides. The study of the electron–phonon (el–ph) interaction and superconducting properties of neodymium hydrides is complicated because of two reasons: (1) possible interplay between magnetic ordering and el–ph interaction (see below) for which there is no relevant theory at this moment, ^{24,44} and (2) complex structure of the electronic density of states N(E) (or DOS) with multiple van Hove singularities (vHs) near the Fermi level (Figures S6 and S11) caused by f-electrons of Nd. The latter makes us go beyond the commonly accepted "constant DOS approximation" in calculations of the critical temperature and take into account the exact structure of the electronic density of states.

One of the consequences of the described effect is a strong dependence of the parameters of el–ph interaction on the Gaussian broadening (σ) parameter used in Quantum ESPRESSO (QE) (see the Supporting Information). A similar situation was previously observed for praseodymium superhydrides $P6_3/mmc$ -PrH $_9$ and $F\overline{4}3m$ -PrH $_9$, where the results of QE calculations are strongly dependent on σ and on the chosen Pr pseudopotential.

To solve this problem, we applied a method developed by Lie and Carbotte⁴⁷ and analyzed the solution of the linearized Eliashberg equations,⁴⁸ taking into account the detailed structure of N(E). As it was shown,⁴⁷ the N(E) affects the Eliashberg equations via the following integral1:

$$\overline{N}(\overline{|\omega_i|}) = \frac{1}{\pi} \int_{\infty}^{-\infty} \frac{\overline{|\omega_i|}}{E^2 + \overline{|\omega_i|^2}} \frac{N(E)}{N(0)} dE$$
 (1)

the contribution of which is determined by the $E\approx 0$ (= $E_{\rm F}$) region in N(E). To illustrate this, we calculated this expression for the studied compounds: NdH₄, NdH₇, and NdH₉ (Figure S11). Our results show that the maximum positive influence of N(E) on $T_{\rm C}$ is for NdH₇ and NdH₉, while for NdH₄ this effect is negative.

Harmonic calculations (for now neglecting magnetism) with using numerical solution of the isotropic set of Eliashberg equations and $\sigma = 0.025$ Ry (which gives good convergence) show that NdH₉ at 120 GPa has quite a small electron—phonon coupling constant $\lambda = 0.43$ and logarithmic average

phonon frequency ω_{\log} = 602 K. Contribution of the hydrogen sublattice to the DOS is very small (1–3%); that is why the matrix elements of the el-ph interaction are ~0. As it is shown in Figure S11, accounting for the exact N(E) structure leads to the increase of critical temperature. Calculations within the "constant DOS approximation" led to a 10% lower $T_{\rm C}$ = 3.7 K (μ * = 0.1).

Combining the results of the VASP calculations with those of QE, we have obtained a more reliable Eliashberg function (Figure S13) to describe the electron—phonon interaction in NdH₉ (see the Supporting Information). This $\alpha^2 F(\omega)$ leads to much higher $\lambda=2.82$ and lower $\omega_{\log}=272$ K. The resulting critical temperature is 63 K ($\mu^*=0.1$) according to the Allen—Dynes formula. However, consideration of antiferromagnetic ordering and spin-splitting leads to an almost complete suppression of superconductivity (see calculations with UppSC code).

The electron–phonon interaction in the molecular C2/c-NdH₇ at 100 GPa is even weaker than that in NdH₉: $\lambda = 0.23$, $\omega_{\log} = 842$ K, and formal $T_{\rm C}$ ($\mu^* = 0.1$) is ~ 0.01 K. This may be associated with the almost zero ($\sim 1\%$) contribution to $N(E_{\rm F})$ from hydrogen (Figure S6b). Numerical solution of Eliashberg equations with accounting for the exact DOS structure at formal $\mu^* = 0$ leads to a maximum possible $T_{\rm C}$ of 2.7 K, while within the "constant DOS approximation" we obtained a much lower value of 1.2 K.

Unexpectedly, a pronounced electron–phonon coupling is predicted for I4/mmm-NdH₄ (Figures S10 and S11). At 100 GPa, this compound has $\lambda=0.54$, $\omega_{\rm log}=857$ K that corresponds to $T_{\rm C}=13.3$ K ($\mu^*=0.1$). Because of the presence of a clear maximum of N(E) close to $E_{\rm F}$, the determination of superconducting $T_{\rm C}$ strongly depends on the chosen σ -smoothing (Figures S10 and S11). This has a negative effect on $T_{\rm C}$, and within the "constant DOS approximation" the critical temperature is 15.2 K.

Eliashberg Calculations with UppSC Code. In conventional superconductors, Cooper pairs are formed from electrons in time-reversed states, and the relevant order parameter is proportional to the anomalous average $\langle c_{k\uparrow}c_{-k\downarrow}\rangle$. Because of the opposite spin of the paired electrons, an applied magnetic field tends to break the Cooper pair apart, and eventually at some critical field the superconducting state will be destroyed. In magnetic superconductors due to the underlying magnetic state, electrons feel such an effective magnetic field, which has a detrimental influence on the superconducting state. This effective magnetic field lifts the spin degeneracy, therefore inducing a spin-splitting "gap" in the electron energy dispersion. Electrons at the Fermi level now need to overcome this energy to form spin singlet Cooper pairs.

To consider the possible effect of the magnetic structure on the electron–phonon interaction in $P6_3/mmc$ -NdH₉, we analyzed the spin-resolved electronic band structure (Figure S6) and computed solutions of the Eliashberg equations using the UppSC code (see the Supporting Information). We found that the value of the spin-splitting at the Fermi level is too large (450 meV with U-J=0, and 890 meV at U-J=5 eV) for any finite superconducting solution, as confirmed by our calculations, which yield $T_{\rm C}=0$ K even for $\mu^*=0$. Given that experiments find $T_{\rm C}\approx 4.5$ K, it is worthwhile to discuss possible scenarios. Aside from the potential presence of superconducting impurities (Mo,Nd)C_xH_y or magnetic transitions, it is possible that, within our current calculations, we

overestimated the spin-splitting (h) around the Fermi level. To investigate this scenario, we solved the Eliashberg equations for several values of h, T, and μ^* . We found that for $\mu^* = 0.1$ and h = 20 meV, $T_C \approx 5$ K in agreement with experiment. Therefore, it may be that high-pressure NdH_o could in principle be the first example material of the hydride family where superconductivity and AFM coexist, but at low temperature. Such a coexistence is not a priori to be excluded as has been shown, for example, in the case of unconventional AF super-Another possible scenario is that the lowtemperature superconducting state may involve some form of a spin triplet superconducting order parameter. A possible candidate could be an equal spin triplet state.⁵² Further investigations toward such an exciting direction would require solving the fully anisotropic Eliashberg equations, 53 which goes beyond the scope of the present work.

CONCLUSIONS

Following the remarkable discovery of LaH₁₀, here we synthesized three novel neodymium polyhydrides, P6₃/mmc-NdH₉, C2/c-NdH₇, and I4/mmm-NdH₄, through the in situ laser-assisted decomposition of NH3BH3 with the simultaneous absorption of the released hydrogen by metallic Nd. Hexagonal NdH₉ is the next member of the P6₃/mmc family of La, Ce, Pr, Th, and U nonahydrides. For all of the synthesized phases, the equations of state and unit cell parameters are in satisfactory agreement with our DFT or DFT+U(U-I=5 eV)calculations. Preliminary measurements of the electrical resistance of NdH₉/NdH₄ sample point to a possible SC transition at 4.5 ± 0.5 K at 110 GPa and the absence of superconductivity above 5 K. Although P63/mmc-NdH9 has the highest H-content, the large spin-splitting in the electron band structure (>450 meV) and antiferromagnetic ordering almost rule out classical s-wave superconductivity. Thus, the intensity of superconducting properties declines in the La-Ce-Pr-Nd series of superhydrides, while magnetic properties become more pronounced.

Theoretical calculations predict that all of the neodymium hydrides exhibit strong magnetism at pressures below 150 GPa and possess collinear antiferromagnetic order, similar to *I*4/*mmm*-PrH₄, examined as a reference. The critical Néel temperatures for the newly synthesized neodymium hydrides were estimated using the mean-field approximation to be about 5 K (NdH₄), 251 K (NdH₇), and 136 K (NdH₉).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.9b10439.

Experimental methods, computational details, structural information, electron and phonon properties of neodymium hydrides, magnetic properties of neodymium and praseodymium hydrides, superconductivity in neodymium hydrides and additional references (PDF)

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