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Synthesis of Superconducting Cobalt Trihydride

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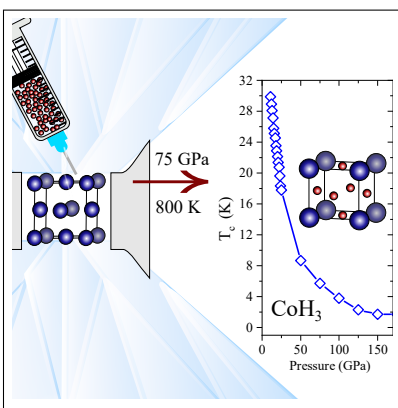
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Abstract

The Co-H system has been investigated through high-pressure high-temperature x-ray diffraction experiments combined with first principles calculations. On compression of elemental cobalt in a hydrogen medium, we observe *fcc* cobalt hydride (CoH) and cobalt dihydride (CoH₂) at 33 GPa. Laser heating CoH₂ in a hydrogen matrix at 75 GPa to temperatures in excess of ~ 800 K, produces cobalt trihydride (CoH₃) which adopts a primitive structure. Density functional theory calculations support the stability of CoH₃. This phase is predicted to be thermodynamically stable at pressures above 18 GPa and to be a superconductor below 23 K as critical temperature, T_c . Theory predicts that this phase remains dynamically stable upon decompression above 11 GPa where it has a maximum T_c of 30 K.

Graphical TOC Entry



The physical properties of a host metal can be profoundly altered by the presence of hydrogen. Metallic structures have been predicted that when hosting high density atomic hydrogen they could exhibit novel properties such as high-temperature superconductivity.¹ The potentials of these materials are exemplified by the recent high-temperature superconductivity experimentally observed in the rare-earth hydrides.^{2,3} Prior to these findings, superconductivity was also proposed to emerge in the transition metal hydrides (TMH).⁴⁻⁶

The transition metals are good electron donors, as they tend to donate their two external s^2 electrons thus stabilizing the H^- anion over molecular hydrogen. Under moderate pressures, almost all transition metals form monohydride compounds.⁷⁻⁹ Platinum, for example, is observed to form two hexagonal monohydride variants, PtH-I and PtH-II, at pressures above 27 GPa.¹⁰ *Ab initio* calculations initially predicted PtH-II to be a superconductor, with a critical temperature, T_c , below 17 K at 90 GPa, however with the inclusion of anharmonic effects, the T_c was substantially reduced to < 1 K.^{5,10,11} Recent electrical measurements found that at 30 GPa, PtH-II has a T_c of 7 K, which reduced on further compression.¹² Through a combination of high pressure and/or high temperature, TMH with more unusual stoichiometries become energetically competitive.¹³⁻²² Iron exhibits several polyhydride species, of which FeH_5 ($I4/mmm$), synthesised at 120 GPa and temperatures >1500 K, is the most hydrogen-rich and possesses a unique layered structure.²³ Potential superconductivity in FeH_5 has been a subject of debate, with two theoretical works predicting remarkably high T_c values ranging between 46 - 51 K, and conversely, another computational study finding no superconductivity.²⁴⁻²⁶ However, the latter study did suggest that trihydrides heavier than FeH_3 , with a cubic $Pm\bar{3}m$ structural type, could facilitate superconductivity through the addition d electrons to the hybrid $3d$ electron shell of the metal-hydrogen band.²⁶

Cobalt is a prime transition metal candidate to also exhibit a trihydride form, being positioned in the same group as iridium and period neighbour iron, both of which form cubic trihydrides.^{14,23} Indeed, a recent theoretical study predicted the formation of CoH_3 above 30 GPa sharing the same $Pm\bar{3}m$ structure as both FeH_3 and IrH_3 .²⁷ However, both

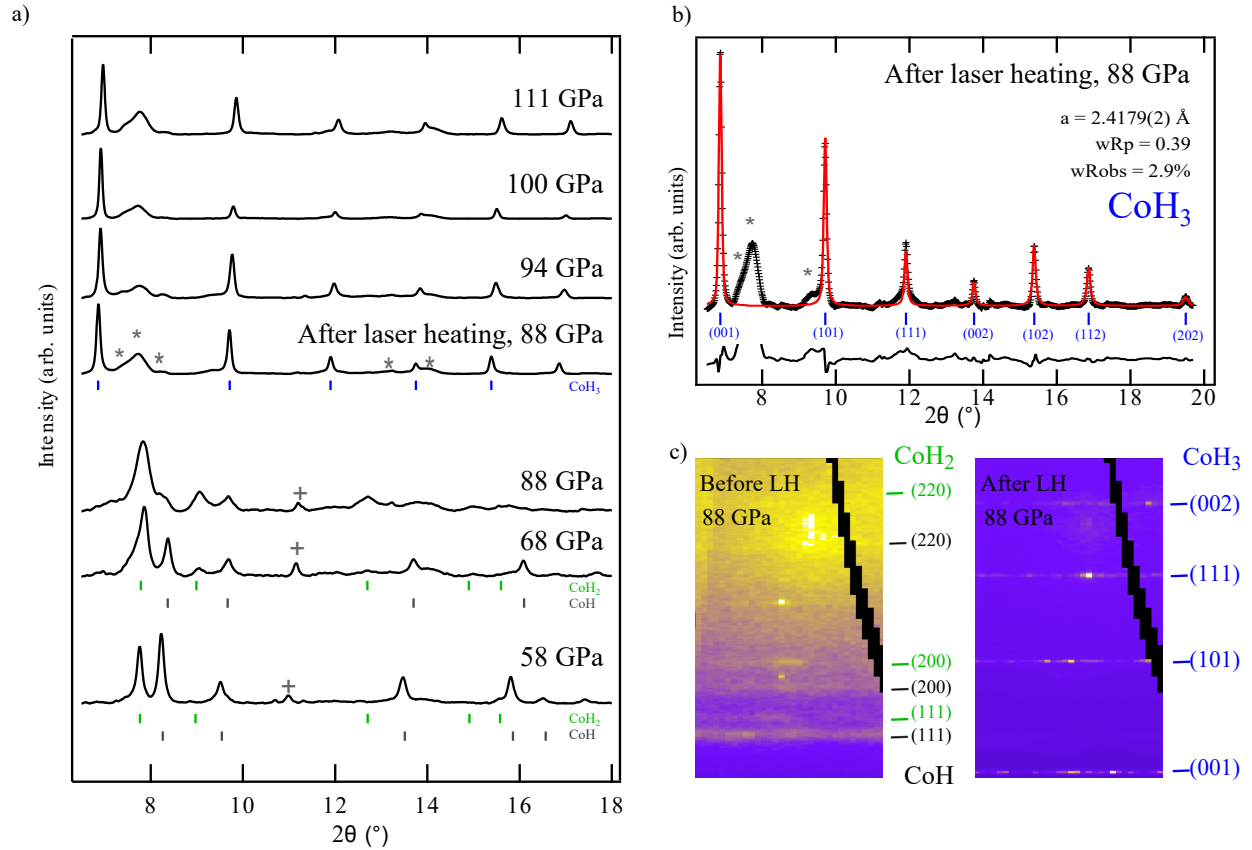


Figure 1: (a) High pressure x-ray diffraction patterns ($\lambda = 0.2895 \text{ \AA}$) taken on compression to 111 GPa. Laser heating at 88 GPa yields the synthesis of $Pm\bar{3}m$ cobalt trihydride. Asterisks correspond to Re and ReH, whilst crosses correspond to W. (b) Rietveld refinement fit of CoH₃ structures, unfitted peaks marked by * correspond to ReH and Re. Refinement parameter are $wRp=0.39$ and $wR_{obs}=2.9$. (c) Diffraction plates of the sample at 88 GPa before (left) and after (right) laser heating. In all panels, tick marks indicate the reflections of CoH (grey), CoH₂ (green), CoH₃ (blue).

these studies and a further computational work, predict CoH_3 to have a T_c below 1 K at pressures above 150 GPa.^{26,27} Experimentally, the synthesis of CoH_3 has remained elusive, with only the formation of face-centred cubic CoH above 4 GPa and the stepwise transition to CoH_2 above 45 GPa.²⁰

Here, we demonstrate the synthesis of cobalt trihydride in a laser-heated diamond-anvil cell through a series of synchrotron X-ray diffraction experiments. On compression, we observe the known formation of both CoH and CoH_2 , which we find to co-exist to at least 75 GPa. Laser heating CoH and CoH_2 in a hydrogen matrix at pressures and temperatures above 75 GPa and 700 K leads to the complete transformation of the sample, identified by new diffraction peaks. We identify the new compound as CoH_3 and find it to be isostructural to FeH_3 , with Co atoms occupying the vertices of the simple cubic unit cell and H atoms occupying the vacant face centers.¹⁵ Total energy and electron-phonon calculations reveal that CoH_3 remains energetically stable down to 18 GPa, at which pressure it exhibits superconductivity with a T_c of 23 K.

After gas loading at 0.2 GPa, mixtures of Co and H_2 were compressed to 33 GPa. [After one week](#) at this pressure, x-ray diffraction patterns reveal the synthesis of both CoH and CoH_2 compounds, in agreement with previous work.²⁰ CoH has space group symmetry $Fm\bar{3}m$, with unit-cell length $a = 3.4849(7)$ Å (at 60 GPa), while CoH_2 has unit cell length $a = 3.7533(8)$ Å (at 60 GPa) within the same group. Interestingly, CoH_2 is not isostructural to FeH_2 which adopts a tetragonal $I4/mmm$ structure,¹⁵ but instead adopts the same structure as RhH_2 at 14 GPa.¹³ Remarkably, we find that both CoH and CoH_2 co-exist up to pressures of 88 GPa (see Figure 1(a)), with no indication of further hydrogenation. We also do not observe a transition on compression to tetragonal ($I4/mmm$) CoH_2 , which was predicted to become energetically favourable above 42 GPa.²⁷

The laser heating of metals in a high-pressure hydrogen environment has been a successful synthetic tool in yielding hydrogen-rich metal hydrides,^{3,17,30,31} and was utilised here to explore the synthesis of cobalt polyhydride species. Samples of CoH/CoH_2 and H_2 were

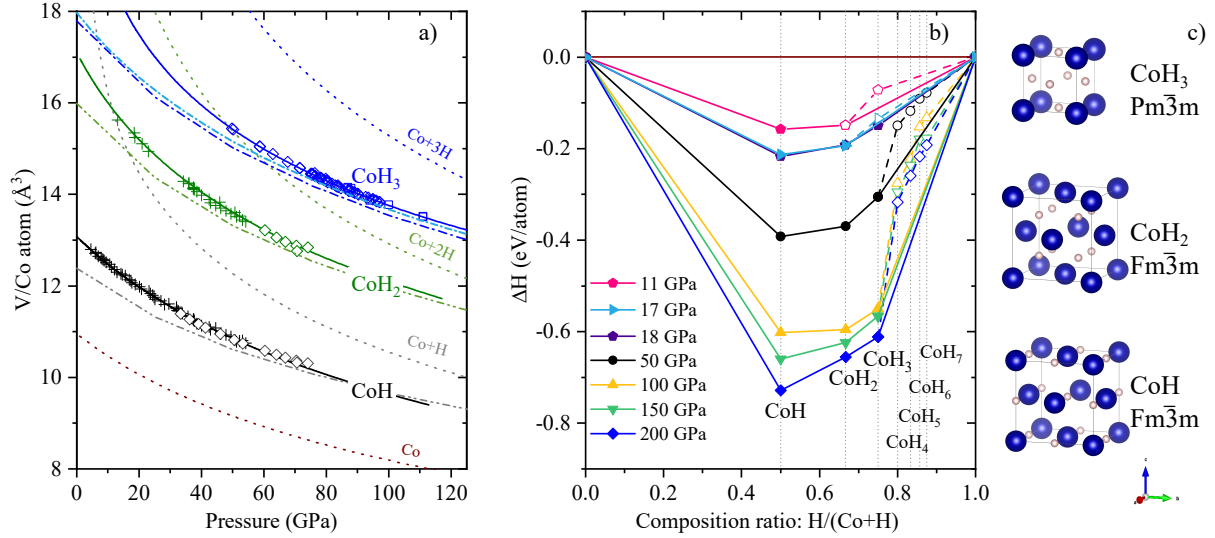


Figure 2: (a) Volume per Co atom for Co-H compounds. Diamond and square symbols correspond to measurements from two experimental runs, whilst crosses represent data taken from Wang *et al.*²⁰ Solid lines correspond to the experimental equation of states (EOS) calculated using data from this work and Wang *et al.*²⁰ Dashed dotted lines were calculated through DFT using ultrasoft pseudopotentials (QuantumESPRESSO code). For CoH₃ the light blue dashed-dotted line was calculated with VASP, using ‘hard’ PAW pseudopotentials and the optB88-vdW dispersion-corrected functional. Dotted lines represent the EOS derived from the atomic volumes of Co and H.^{28,29} (b) Convex hull construction for CoH_x compounds at a sequence of pressures. Empty (filled) symbols denote metastable (stable) phases. (c) Crystal structures of cobalt polyhydrides, CoH ($Fm\bar{3}m$), CoH₂ ($Fm\bar{3}m$) and CoH₃ ($Pm\bar{3}m$). Co atoms are represented by blue spheres and H atoms are represented by pink spheres.

compressed to pressures of 75 GPa and 88 GPa, and held here for 24 and 12 hours respectively, observing no time induced transformation. These samples were then laser heated to about <800 K (Figure S1 and see methods for further details). Upon quenching of the sample, we see the disappearance of the diffraction lines attributed to CoH/CoH₂, which are supplanted by new diffraction peaks (see Figure 1(a,c) and S1³²). All of the new diffraction peaks can be indexed with a simple cubic unit cell ($Pm\bar{3}m$) with lattice parameter $a = 2.4358(2)$ Å at 75 GPa. Rietveld refinements of this structure (Figure 1(b)) show Co atoms to be on the vertexes of the unit cell, with final agreement factors of wRp 0.39 and a wR_{obs} 2.9. Samples were compressed up to 111 GPa, the highest pressure reached in this study, and subsequently decompressed down to 52 GPa (below which the sample was lost due to anvil failure) to determine the equation of state (EoS) and evaluate the stability of the compound. The volume per Co atom as a function of pressure of CoH, CoH₂, and the synthesised product were fitted with third-order Birch-Murnaghan P - V EoS's (see Figure 2 (a)) with the fit parameters shown in table S1. The experimentally determined EoS gives volumes greater than Co+2H, and comparison between the experimental EOS and that determined theoretically for CoH₃ present a good agreement. The determined structure also matches that of which was previously predicted for CoH₃,²⁷ and is isostructural to FeH₃,¹⁵ synthesised through similar methods.

We have performed our own variable-composition structure search to find energetically stable structures in the Co-H system up to 400 GPa using the evolutionary crystal structure prediction method USPEX package and the particle swarm optimisation method as implemented in the CALYPSO code (Figures S2-S4).³³⁻³⁷ Through enthalpy calculations for the most stable structures, we find CoH₃ to adopt a $Pm\bar{3}m$ unit cell identical to FeH₃^{15,38} and IrH₃,¹⁴ with the H atoms occupying the 3c site $(0, \frac{1}{2}, \frac{1}{2})$. This, together with the computationally determined EOS, is in good agreement with experiments. CoH₃ is on the convex hull above 18 GPa, , however experimentally we only see CoH and CoH₂ which we attribute to kinetics barriers. CoH and CoH₂ have the highest formation enthalpies, 0.602 and 0.596 eV

per atom respectively at 100 GPa (see Figure 2) and also share the same *fcc*-Co sub lattice. Pure Co adopts a *hcp* phase at pressures below 105 GPa, above which it transforms to *fcc*-Co.²⁹ So although *fcc*-Co is not the ground state at 33-55 GPa, it is likely there is a low energy barrier³⁹ to form metastable *fcc*-Co, which can then be filled with hydrogen to form CoH and/or CoH₂. Experimentally, both compounds can be formed on compression alone, with laser heating accelerating the process.²⁰

CoH₃, however, is very different, as it has a *simple cubic* Co sub lattice. Since it is not close packed a major rearrangement of the Co lattice is required to allow the formation of CoH₃. Neither low pressure and high temperature or long times at pressures below 50 GPa are sufficient to initiate the CoH₃ formation,²⁰ requiring pressures above 75 GPa and laser heating to trigger the CoH₃ synthesis. In agreement with the convex hull calculations, and Gibbs' phase rule, only CoH₃ is formed after the laser heating in an excess H₂ environment. Our DFT calculations also find an unreported structural transition in CoH₂ above 275 GPa to a tetragonal *P4/nmm* phase (Figure S4).

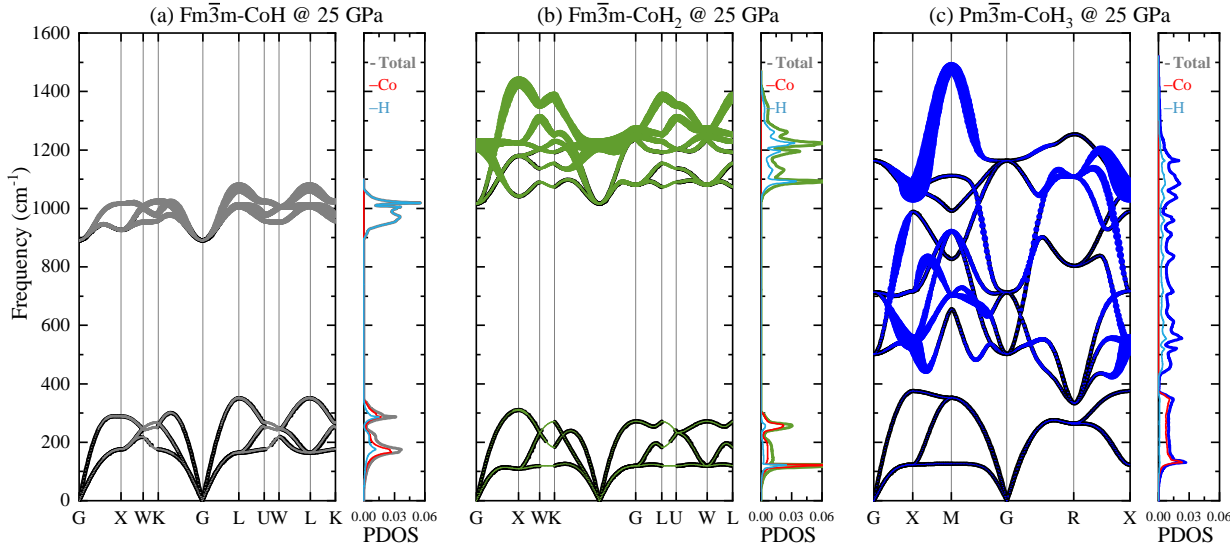


Figure 3: Phonon dispersion curves and Projected phonon density of states at at 25 GPa for; (a) CoH, (b) CoH₂, (c) CoH₃. The phonon dispersions $\omega_\nu(\mathbf{q})$ for each phonon mode ν at momentum space point \mathbf{q} are drawn as circles with radii proportional to the magnitude of the electron-phonon linewidths $\gamma_{q\nu}$.

Recent theoretical work anticipated that transition metal trihydrides isostructural to

FeH₃ but with heavier elements, like Co, would be candidates to undergo superconducting transitions.²⁶ Our DFT calculations of the electronic structure and electronic bands demonstrate metallicity of the three hydrides at all pressures throughout their stability ranges (see Figure S5 - S8). At the lowest pressure, the electronic density of states (DOS) of CoH₃ (see Fig. S5) has important contributions from hydrogen orbitals around the Fermi level, indicative that it could exhibit superconductivity.

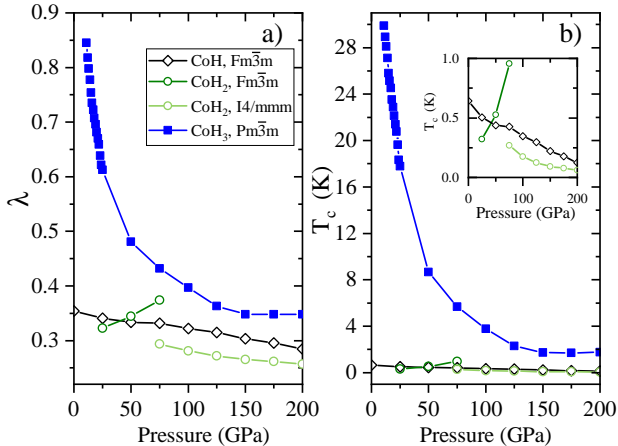


Figure 4: (a) Electron-phonon coupling constant λ for the calculated favoured cobalt hydrides phases as a function of pressure. (b) Critical temperature, T_c , for CoH, CoH₂ and CoH₃ as a function of pressure. Insert shows T_c as a function of pressure between 0 and 1 K. In all panels, CoH is represented as black diamonds, CoH₂ as green circles and CoH₃ as blue squares.

In Figure 3 we show the calculated phonon dispersion curves and projected phonon density of states (PDOS) for the low pressure phases of CoH, CoH₂ and CoH₃ at 25 GPa (see Figures S9-S12 for higher and lower pressure results). Phonon structures of all the hydrides depict no imaginary frequencies along the high symmetry k path, which demonstrates their dynamical stability. For $Fm\bar{3}m$ -CoH, as shown in Figure 3(a), a phonon gap between 400 cm^{-1} and 900 cm^{-1} divides the spectrum into two regions: the low-frequency acoustic branches and high-frequency optical branches. The optical modes are associated with almost exclusive motion of H atoms, whereas the acoustic modes are associated with the concerted motions of both Co and H atoms. From Figure 3(b), CoH₂ has a similar gapped phonon

spectrum compared to CoH, but with higher optical frequency branches up to 1400 cm^{-1} in the hydrogenic sublattice. In CoH₃, there is a big change in the phonon dispersion and PDOS. The phonon gap has shrunk and the larger number of optical branches occupies a much wider frequency region. From CoH to CoH₃, the magnitude of the electron-phonon linewidths increases with the higher hydrogen content, which may lead to a higher zone-averaged electron-phonon coupling parameter (EPC) and superconducting critical temperature, T_c .

We estimate the EPC and T_c using the McMillan-Allen-Dynes formula^{40–42}

$$T_c = \frac{\omega_{ln}}{1.2} \exp \left[-\frac{1.04(1 + \lambda)}{\lambda - \mu^*(1 + 0.62\lambda)} \right], \quad (1)$$

where ω_{ln} is the logarithmically averaged phonon frequency, and taking the Coulomb parameter as $\mu^* = 0.1$. The EPC λ and T_c are plotted in Figure 4 as a function of pressure. For CoH and CoH₂, the calculated λ are below ~ 0.45 in the studied pressure range. Such weak EPC result in T_c below 2 K. However, as seen in Figure 4, CoH₃ shows a different and remarkable trend with λ significantly increased with decreasing pressure, which in turn leads to a rapid T_c increase upon decompression, reaching a predicted maximum of 30 K at 11 GPa.

There are some quantitative differences between our results and those obtained in previous calculations.²⁷ For CoH₃ at 200 GPa we obtain $\lambda = 0.35$ and $T_c = 2$ K, while Wang *et al.*²⁷ obtain $\lambda = 0.19$ and $T_c \sim 0$ K. We have repeated our calculations with parameters [very similar to those](#) used by Wang *et al.* (see figs. S13-S23) and reproduce the EPC and T_c data (Figure S23, [\$\lambda = 0.21\$ and \$T_c = 4\$ mK](#)). We postulate that the choice of pseudopotential – norm-conserving (NC) in Wang *et al.*²⁷ and ultrasoft (US) here – is the source of the quantitative differences we see. The US pseudopotentials are better suited than the NC to describe the EOS of pure Co and H⁴³ [but give very similar EOS results for CoH₃; both also agree with VASP calculations using the PAW method, see Figure S13 in the SM.](#) The choice of pseudopotential has a larger effect on phonon properties than on electronic properties;

Nonetheless, using either approach, the EPC and T_c share the same trend upon compression, but on different absolute scales (see Figure S23) - with the NC pseudopotentials, a T_c of CoH₃ is predicted to reach 15 K at 13 GPa the US pseudopotentials give the same value if the screened Coulomb interaction μ^* would be increased to 0.18). These quantitative uncertainties might be best resolved by using more sophisticated theoretical approaches such as directly solving the Eliashberg equation or the superconducting DFT method.^{44,45}

The strong negative pressure dependence of T_c in CoH₃ at these low pressures can be explained by the hardening of the phonon frequencies leading to a weakening of EPC λ . As shown in Figures 3 and S9-S12, when the pressure increases from 10 GPa to 300 GPa, the maximal frequencies of optical phonon branches for CoH₃ hardens from 1500 cm⁻¹ to 2400 cm⁻¹, while DOS $N(0)$ (see figures S6 to S8) and magnitude of electron-phonon linewidths $\gamma_{\nu q}$ remain almost unchanged. A similar trend in T_c was experimentally observed in PtH-II, from 7 K at 30 GPa to 5 K at 40 GPa.¹²

In our experimental runs, diamond failure on decompression prevented us to study the stability of CoH₃ at pressures lower than 52 GPa. At 52 GPa, we predict CoH₃ to have T_c of 8 K, which could increase to 30 K at 11 GPa. We speculate that due to the large stability range predicted for CoH₃ (Figure S1), that low temperature could maintain the stability of CoH₃ down to its maximal T_c conditions, and such behaviour should be explored in future studies.

The predicted superconducting character of CoH₃ proves that TMH₃ heavier than FeH₃ do facilitate more efficient electron-phonon coupling. We have demonstrated the formation of CoH₃ synthesised at 75 GPa and stable in decompression to at least 52 GPa. CoH₃ adopts a cubic $Pm\bar{3}m$ structure which is very stable both in compression up to 111 GPa and in decompression down to 52 GPa. Our DFT results predict that this structure should be superconducting with a T_c which increased with decreasing pressure, reporting a maximum of 30 K at 11 GPa. These results give an indication that superconductivity could also be observed in metal hydrides at low pressure conditions.

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Supporting Information Available

The Supporting Information is available free of charge at <https://>

- Experimental and theoretical methods
- Equation of state parameters, electronic and phonon band structures, experimental and calculated unit-cell parameters.

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