Supplementary Material for: Chemically-Assisted Pre-Compression of Hydrogen Molecules in Alkaline-Earth Tetrahydrides

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METHODS

Experimental details

Elemental Ca (99.9%, Alfa Aesar), Sr (99.99%, Sigma Aldrich), Ba (99 %,Sigma Aldrich) (or BaH$_2$ (99.5 %, ABCR)) were loaded into diamond anvil cells in an argon atmosphere glove box and and subsequently gas loaded with research grade hydrogen (99.9995 %) at 0.2 GPa. In one of our experiments Sr was loaded with ammonia borane (NH$_3$BH$_3$) as a hydrogen source. A small chip of ruby or gold was included as a pressure calibrant and cross-checked against the vibron frequency of pure H$_2$, the Raman edge of stressed diamond and the known equation of state of the dihydride precursor. Ultra low-fluorescent diamonds with culet diameters ranging from 50-200 µm were used with rhenium gaskets. Samples were heated uniaxially in situ by directly coupling them to an yttrium-aluminum-garnet (YAG) continuous wave laser with wavelength $\lambda =$1064 nm. Collected emission spectra were fitted using a two-parameter grey-body Planck distribution.

Angle-dispersive X-ray diffraction patterns were recorded at 16-ID-B, Advanced Photon Soure, United States, ID15B at the European Synchrotron Radiation Facility, France and P02.2 Petra-III, Germany. Incident beam energies in the range of 25–37 keV were used. The calibration of detector position, primary processing, azimuthal integration and background subtraction were performed using the DIOPTAS v0.5.5 software. Le Bail refinements were carried out in JANA2006 and for Rietveld refinements the POWDERCELL 2.4 program was used.

Raman spectroscopy measurements were conducted using 514.5 nm excitation wavelengths via a custom-built micro-focused Raman system provided with 1800 gr/mm and 600 gr/mm gratings and a Pylon 1340x100BRX liquid nitrogen cooled camera.

Computational details

Density functional theory calculations were performed with the CASTEP code using the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional. Geometry optimization calculations of
the tetrahydride structures were carried out using ultrasoft pseudopotentials generated by CASTEP on-the-fly. A cutoff energy of 700 eV and a k-point grid spacing better than 0.04 Å⁻¹ gave well converged energies and forces for all the structures.

Other exchange-correlation functionals were tested: BLYP¹⁵ and the semi-empirical dispersion correction (SEDC) method of Tkatchenko and Scheffler¹⁶ to account for the van der Waals interactions. Table S1 contains the structural parameters for the three alkaline-earth metal hydrides at 25 GPa obtained with the different functionals. BLYP volumes are bigger than those obtained with the PBE functional by 3-5-4.6 %, whereas the c/a radii and intramolecular distances are lower by 1-9-2.6 % and 3.2%. The inclusion of the van der Waals contributions leads to a considerable decrease of the volume by up to 8.2 % from PBE. Experimental volumes lie between PBE and BLYP values. The inclusion of van der Waals contributions gives too low volumes and the comparison with the experiment actually become worse.

|        | CaH₄ |         | SrH₄ |         | BaH₄ |         |
|--------|------|---------|------|---------|------|---------|
|        | Volume | c/a | d₉–H | Volume | c/a | d₉–H | Volume | c/a | d₉–H |
| PBE    | 30.67 | 1.653 | 0.782 | 36.15 | 1.653 | 0.782 | 42.95 | 1.6903 | 0.784 |
| BLYP   | 31.73 | 1.622 | 0.757 | 37.53 | 1.617 | 0.757 | 44.92 | 1.6459 | 0.759 |
| PBE+TS | 28.29 | 1.6476 | 0.782 | 33.17 | 1.7199 | 0.791 | 39.48 | 1.7919 | 0.796 |

Table S1: Volumes per alkaline-earth metal, c/a radii and intramolecular distances for the I₄/mmm structures of the hydrides at 25 GPa.

Random structure searches were performed with AIRSS to explore the BaH₄ landscape. Ab initio molecular dynamics (AIMD) calculation in CASTEP were run with the NPT ensemble and with Barium assigned a fictitious mass number of 2 to more effectively sample the configuration space. A topological analysis of the electron density and the electron localization function was done with the CRITIC2 code¹⁷ with the energy cutoff increased to 1000 eV to assure convergence. In Table S2 we present the structural parameters for I₄/mmm BaH₄ at 40 GPa estimated with PBE, BLYP and PBE+TS, and we included the experimental data. It can be seen that the choice of a different functional does not solve the discrepancy for c/a between theory and experiment, with an even lower c/a using BLYP.

There have been different approaches to determine the atomic charges in crystals. Probably,
Table S2: Volumes per formula unit, $c/a$ radii and intramolecular H-H covalent bond length for the $I4/mmm$ BaH$_4$ at 40 GPa.

| Compound | Lattice parameters ($\AA$) | Atoms | $x$ | $y$ | $z$ |
|----------|---------------------------|-------|-----|-----|-----|
| CaH$_4$ ($I4/mmm$) | $a=b=3.129$ $c=5.331$ $\alpha=\beta=\gamma=90^\circ$ | Ca(2a) | 0.0000 | 0.0000 | 0.0000 |
| 50 GPa | H(4d) | 0.0000 | 0.5000 | 0.2500 |
| | H(4e) | 0.0000 | 0.0000 | 0.4258 |
| SrH$_4$ ($I4/mmm$) | $a=b=3.298$ $c=5.641$ $\alpha=\beta=\gamma=90^\circ$ | Sr(2a) | 0.0000 | 0.0000 | 0.0000 |
| 50 GPa | H(4d) | 0.0000 | 0.5000 | 0.2500 |
| | H(4e) | 0.0000 | 0.0000 | 0.4298 |
| BaH$_4$ ($I4/mmm$) | $a=b=3.445$ $c=6.074$ $\alpha=\beta=\gamma=90^\circ$ | Ba(2a) | 0.0000 | 0.0000 | 0.0000 |
| 50 GPa | H(4d) | 0.0000 | 0.5000 | 0.2500 |
| | H(4e) | 0.0000 | 0.0000 | 0.4345 |
| BaH$_4$ ($I4/mmm$) | $a=b=2.637$ $c=6.492$ $\alpha=\beta=\gamma=90^\circ$ | Ba(2a) | 0.0000 | 0.0000 | 0.0000 |
| 200 GPa | H(4d) | 0.0000 | 0.5000 | 0.2500 |
| | H(4e) | 0.0000 | 0.0000 | 0.3156 |
| BaH$_4$ ($R3m$) | $a=b=4.289$ $c=9.708$ $\alpha=\beta=90^\circ$ | Ba(3b) | 0.0000 | 0.0000 | 0.5000 |
| 10 GPa | H(6c) | 0.0000 | 0.0000 | 0.7646 |
| | H(6c) | 0.0000 | 0.0000 | 0.0403 |

Table S3: Calculated structural parameters of the predicted $I4/mmm$ phases of CaH$_4$, SrH$_4$ and BaH$_4$ at 50 GPa, BaH$_4$ at 200 GPa and the $R3m$ phase of BaH$_4$ at 10 GPa using PBE. We have calculated the ELF of the metallic sublattices (without hydrogens).

one of the most straightforward and mathematically rigorous is the Quantum Theory of Atoms in Molecules developed by Bader.\textsuperscript{18} It is based on the topology of the electron density. It shows maxima (cusps) on the atoms and decays exponentially when moving away from them. This allows the partition of the crystalline space in topological atoms defined by the union of the electron density maxima with their attraction maxima and delimited by surfaces obeying the zero-flux condition for the electron density. Their charge can be calculated from the integration of the electron density within these regions. These regions are non-overlapping and additive, recovering the total volume of the crystal.

The electron localization function was introduced by Becke and Edgecombe in 1990 for the
Table S4: PBE calculated volume per metallic atom in the $I4/mmm$ structure for the alkaline-earth tetrahydrides versus pressure

| P(GPa) | CaH$_4$ | SrH$_4$ | BaH$_4$ |
|--------|---------|---------|---------|
| 0      | 42.25   | 51.24   | 64.26   |
| 5      | 38.42   | 46.00   | 56.29   |
| 10     | 35.73   | 42.50   | 51.33   |
| 15     | 33.68   | 39.89   | 47.82   |
| 20     | 32.04   | 37.84   | 45.12   |
| 25     | 30.67   | 36.15   | 42.95   |
| 30     | 29.50   | 34.73   | 41.15   |
| 35     | 28.49   | 33.50   | 39.61   |
| 40     | 27.60   | 32.42   | 38.27   |
| 45     | 26.81   | 31.47   | 37.17   |
| 50     | 26.10   | 30.61   | 36.05   |

analysis in real space of the electron localization, and later reinterpreted by Savin in terms of the Pauli kinetic energy density ($t_p$). By definition, ELF is a relative measure of the electron localization with respect to the homogenous electron gas (HEG). In general, the ELF value approaches 1 in regions of the space where electron pairing occurs (e.g., atomic shells, bonds and lone pairs).

In analogy with QTAIM, a partition of the space based on the ELF can be performed. It consists of non-overlapping basins with well-defined chemical interpretation (cores, bonds, lone pairs). Moreover, the basin charges come from integration of the electron density within these regions.

We have also calculated the radius of the H$_2$ ELF basin along the line joining its middle point and the alkaline-earth metal. It corresponds to the distance from the middle point to the minimum of the ELF profile along the line connecting it to the alkaline-earth metal.

Regardless of the Ca/Sr/Ba host and structure, the positions of the hydrogens are dictated by the electronic structure of the host lattices, as we find that hydrogens in the tetrahydrides are located on the ELF maxima of the empty metallic lattices (see Fig. 2).

Because $I4/mmm$ is a subgroup of $fcc$ and $bcc$, $fcc$ Ba can be described within the $I4/mmm$ space group with cell parameters $a=a_{fcc}/\sqrt{2}$, $c=a_{fcc}$. Thus, the $fcc$ Ba ELF topology is equivalent to that of $I4/mmm$ Ba: octahedral ELF maxima with bigger ELF values are the preferred site for the H$_2^{8-}$ units, whereas the H$^-$ atoms are located on the tetrahedral sites, Fig. 2b of the main text.
|                | $\text{ELF}_{\text{max}}\text{(oct)}$ |                | $\text{ELF}_{\text{max}}\text{(tet)}$ |
|----------------|-----------------|-----------------|-----------------|
| **Ca (I4/mmm)**| 0.794 14.4282   | 0.472 1.4581    |
| **Sr (I4/mmm)**| 0.705 13.1247   | 0.406 2.2979    |
| **Ba (I4/mmm)**| 0.477 9.6122    | 0.318 2.6687    |
| **Ba (Fm\text{3}m)**| 0.653 19.1620  | 0.384 5.0335    |

Table S5: Calculated ELF values at the octahedral and tetrahedral ELF maxima of the pure metallic sublattices (without hydrogens) of $\text{I4/mmm}$ CaH$_4$, SrH$_4$ and BaH$_4$ at 50 GPa and the $\text{Fm\text{3}m}$ Ba sublattice at 10 GPa. The volumes of their corresponding basins are included in $\text{Å}^3$. We have also calculated the ELF of the metallic sublattice of the experimental BaH$_4$ at 50 GPa with $c/a=2.17$. In this case the elongated basin centered on the octahedral sites ($2b$ sites) breaks into 2 separate maxima on $4e$ sites with ELF value=0.480, with the ELF value 0.336 at the maxima on tetrahedral sites ($4d$ sites). Calculated $\text{I4/mmm}$ BaH$_4$ structure at 200 GPa has $c/a > 2.2$ and becomes more stable than the low $c/a$ $\text{I4/mmm}$ BaH$_4$ structure. This structure at 200 GPa does not have any H$_2$ molecules as all H-H distances are above 1.5 Å, and all the H$_2$, H-) bonds are broken. In the presence of excess hydrogen, it is calculated to be unstable with respect to Ba$_8$H$_{46}$.

Raman calculations and vibrational modes symmetry assignment are computed by CASTEP as part of the raman calculation, as implemented by Keith Refson. The visualization of the vibrations within the lattice related to the Raman modes was done using Jmol.
Table S6: Raman shift of the $I4/mmm$ tetrahydrides theoretically calculated$^{23}$ and experimentally observed. Experimental $fcc$ BaH$_4$ is theoretically approached taking the $R3m$ as template frozen H$_2$ molecules. DFTP was used to calculate vibrational frequencies.$^{23}$ Experimental bands labelled with —— overlap with diamond first order Raman mode and therefore are not observable. Differences between BaH$_4 I4/mmm$ theoretical and experimental are due to the different lattice parameters. Differences between $R3m$ and $fcc$ BaH$_4$ are due to structural differences and calculations not accounting for the H$_2$ rotation. Differences between theory and experiment are very significant in BaH$_4 I4/mmm$ because as said in the main text although the space groups are the same, the lattice parameters are different and the experimental structure is not stable. Difference between the theoretical $R3m$ and experimental $fcc$ are expected as the $R3m$ is taken just as a way to mimic the cubic structure, and $R3m$ distorts in compression.

|          | DFPT $E_{2g}$ | Exp. $E_{2g}$ | DFPT $E_{2g}$ | Exp. $E_{2g}$ | DFPT $A_{1g}$ | Exp. $A_{1g}$ | DFPT $A_{1g}$ | Exp. $A_{1g}$ |
|----------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|
|          | $H_2^+$ lib.  | $H_2^+$ lib.  | $H_2^+H^-$    | $H_2^+H^-$    | $H^-$ lib.    | $H^-$ lib.    | $H^-$ lib.    | $H^-$ lib.    |
| CaH$_4$  | 747           | 675           | 1355          | ——            | 1425          | ——            | 3606          | 3670          |
| SrH$_4$  | 852           | 790           | 1306          | ——            | 1436          | ——            | 3597          | 3600          |
| BaH$_4$  | 988           | 970           | 1212          | 1282          | 1445          | ——            | 3509          | 3330          |
| BaH$_4$  | 1159          | 940           | 1265          | 1125          | 1311          | ——            | 2900          | 3300          |

Figure S1: Volume per Ba/Sr/Ca atom as function of pressure for their corresponding hydrides. The volumes of BaH$_2$; BaH$_4$; SrH$_2$ and SrH$_4$ are from this work. BaH$_{12}$ is taken from Chen et al.$^{24}$ and Ba$_8$H$_{46}$ from Pena-Alvarez, et al.$^{25}$ Sr$_2$H$_3$, Sr$_8$H$_{48}$, SrH$_6$ and SrH$_9$ are from Semenok, et al.$^{26}$ compared with the empty turquoise triangle representing SrH$_9$ from this study. CaH$_2$ and CaH$_4$ are from Mishra, et al.$^{27}$ and from Wu, et al.$^{28}$ whilst CaH$_6$ is from Ma, et al.$^{29}$
Figure S2: Comparison between experimentally derived and calculated lattice parameters for the tetrahydrides. a) Volume per Ba/Sr/Ca atom as a function of pressure. b) Lattice parameters for the I4/mmm structure as a function of pressure. c) c/a ratios for the I4/mmm structure as a function of pressure.
Figure S3: X-ray diffraction patterns of Ca, Sr and Ba tetrahydrides, with their Rietveld refinement shown as solid lines and the raw data as symbols. Tick marks indicate the Bragg peaks of identified phases.
Figure S4: X-ray diffraction patterns of Calcium and hydrogen compressed to 52 GPa, before (bottom) and after laser heating (top). CaH₂ is refined to hexagonal \( P6_3/mmc \) and CaH₄ is refined to tetragonal \( I4/mmm \) (\( \lambda = 0.2904\,\text{Å} \)). Tick marks indicate Bragg peaks from the labelled phases.
Figure S5: a) Representative X-ray diffraction patterns of strontium hydride before laser heating (43 GPa), after laser heating (44 GPa) and upon decompression. SrH$_4$ is refined to tetragonal $I4/mmm$ ($\lambda = 0.2922$ Å). Tick marks indicate Bragg peaks from the labelled phases. b) Representative X-ray diffraction patterns of barium hydrides. BaH$_4$ is refined to $I4/mmm$ upon compression to 45 GPa and on decompression to 35 GPa. At 17 GPa BaH$_4$ is refined to the $fcc$ phase ($\lambda = 0.2922$ Å). c) Representative X-ray diffraction patterns of barium hydrides upon decompression from the Ba$_8$H$_{46}$ phase.$^{25}$ At 18 GPa and below, the patterns are refined to $fcc$ BaH$_4$($\lambda = 0.4115$ Å). Asterisks correspond to BaH$_2$. 
Figure S6: Rietveld refinement of the XRD patterns collected from an Sr + NH\textsubscript{3}BH\textsubscript{3} experiment, collected before (top) and after (middle and bottom) laser heating at a pressure of 74 GPa. After the first laser heating SrH\textsubscript{2} is formed and after a subsequent laser heating the patterns can be indexed to an fcc structure of which the lattice parameters and volumes agree with the previously reported SrH\textsubscript{9}.\textsuperscript{26}
Figure S7: DFT enthalpy differences as a function of pressure for candidate structures of BaH$_4$ from AIRSS with different symmetries vs $C2/c$ – II. The group of stable structures at lower pressures are small distortions from fcc, and are predicted to become fcc at 300K.
Figure S8: Band structure of $I4/mmm$ BaH$_4$ at 0 GPa
Figure S9: Band structure of $I4/mmm$ BaH$_4$ at 50 GPa
Figure S10: a) The $I4/mmm$ pseudo-octahedral metallic coordination shell of the $H_2^5^{-}$ units (with $D_{4h}$ symmetry). In CaH$_4$ and SrH$_4$, this can be viewed as a small distortion of the cation positions from $Fm\bar{3}m$ with the $H_2^5^{-}$ units parallel to the $c$ axis. The larger distortion seen for $I4/mmm$ BaH$_4$ ($c/a > 2$) leads to a more elongated octahedron consistent with the larger H-H distance. b) $R\bar{3}m$ BaH$_4$, a theoretical stable structure found in AIRSS with the molecule oriented on ($\frac{1}{2}, \frac{1}{2}, \frac{1}{2}$); this has a more regular octahedron, similar to the experimentally observed $Fm\bar{3}m$ with a higher ELF-value at the octahedral maximum than $I4/mmm$. Other stable structures found by AIRSS have similar Ba sublattices with various other molecular orientations and associated distortions of the octahedral site. With enforced $I4/mmm$ symmetry a local minimum energy structure exists with $c/a = 2.3$, no molecular hydrogens, and each Ba in a cage of 18 hydrogens. This BaH$_4$ $I4/mmm$ structure becomes more stable than the molecule-containing version at higher pressures around 160 GPa (Ba$_8$H$_{46}$ is still the ground state).
Figure S11: Schematic representation of the librational mode (left) and the H-H vibrational stretching mode (right) within $I4/mmm$ tetrahydrides.
Figure S12: Representative Raman spectra for \textit{fcc} BaH$_4$ + H$_2$ upon compression and decompression. Grey spectra correspond to selected pressures from a different experimental run.
Figure S13: a) Charges of the metals (M), the hydrogens located in tetrahedral sites ($H^\text{y−}$) and of molecular hydrogen located in the octahedral sites ($H_2^\text{δ−}$) based on the Bader topology. b,c) Radius of the $H_2^\text{δ−}$ Bader basin within the $M\text{H}_4$ along the line joining its middle point and the alkaline-earth metals: b) across the $I4/mmm$ ab plane, and octahedral diagonal on the $R3m$; c) across the $I4/mmm$ c axis. Inserted schemes show: b) the $I4/mmm$ ab plane with 4 metallic atoms on the corners hosting one $H_2^\text{δ−}$ in the center perpendicular to the plane; c) $I4/mmm$ c axis with 2 metallic atoms on the corners hosting one $H_2^\text{δ−}$ in the center along the axis. The black arrows represent the $H_2^\text{δ−}$ Bader basin radius.

Figure S14: a) Radius of the ELF basin of the $H^−$ along the line joining it with the $H_2^\text{δ−}$ middle point.
Figure S15: ELF results for the tetrahydride structures as a function of pressure, per columns from left to right is $H_2$, $H^-$ and $M^{2+}$. Per rows, from top to bottom: Basin charge (e), Basin charge density (e/Bohr$^3$), % Volume, Volume.
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