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Configuration and Dynamics of Hydride lons in the Nitride-Hydride Catalyst Ca₃CrN₃H

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sample have an occupancy of at least 95%. We argue that this high H^- occupancy may be related to the material's high efficiency as a catalyst for ammonia synthesis.

1. INTRODUCTION

Mixed-anion materials, that is, solid-state materials containing more than one anionic species in the same phase, e.g., oxidehydrides and nitride-hydrides, can exhibit unique properties that are unattainable with their single-anion counterparts.^{1,2} Such properties include photocatalytic activity,^{3,4} ferroelectricity,⁵ superconductivity,^{6,7} and photoluminescence,⁸ to name a few. Much research in this field has been focused on materials with the antiperovskite structure, i.e., X₃BA, where X is a cation and A and B are anions, due to their functionality as ionic conductors and/or as catalysts.^{9–11} Of specific concern in this study is the antipervoskite hexagonal phase of the nitridehydride Ca₃CrN₃H. This material is obtained by a novel topochemical reaction of the orthorhombic structured Ca₃CrN₃.¹² Importantly, it shows promise as a catalyst for ammonia synthesis,¹³ and its efficient catalytic activity is partly attributed to the diffusion of H⁻ within the crystal structure.¹

Powder X-ray diffraction (PXRD) measurements of Ca_3CrN_3H have shown¹² that it is an X_3BA -type antiperovskite with $X = Ca^{2+}$, $B = H^-$, and $A = [CrN_3]^{5-}$, with the Ca^{2+} ions organized in face-sharing HCa₆ octahedra stacked along the crystallographic *c* axis and bridging $[CrN_3]^{5-}$ planar moieties (Figure 1). It belongs to the space group $P6_3/m$. Interestingly, this channel-like structure accommodating hydride ions is unique within the antiperovskite class, to the best of our knowledge. However, because of the insensitivity of X-rays to hydrogen species, the PXRD patterns of Ca_3CrN_3H and its

hydrogen-free hexagonal counterpart Ca₃CrN₃ barely differ and are hardly distinguishable.¹² Therefore, the actual H⁻ stoichiometry and local coordination environment of the H⁻ anions in Ca₃CrN₃H were not determined in the previous PXRD study.¹² Indeed, Rietveld refinement of the PXRD data together with thermogravimetric and mass spectrometric analyses indicate that Ca₃CrN₃H is nonstoichiometric and rather corresponds to the composition Ca₃CrN₃H_x, with x between 0.7 and 1.¹² In particular, the local coordination environment (configuration) of H⁻, which may be inferred from the vibrational dynamics of the material,¹⁴ is unclear.

A powerful technique to investigate the configuration and vibrational dynamics of hydrogen species in solid-state materials is given by inelastic neutron scattering (INS) coupled to density functional theory (DFT) calculations.^{14–16} This is because neutrons are very sensitive to hydrogen species, and the frequencies and intensities of INS spectra can be generally computed on an absolute scale.¹⁷ Therefore, the experimentally determined vibrational frequencies of H⁻ in Ca₃CrN₃H may, when compared to the DFT-determined

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Figure 1. Schematic crystal structure of Ca_3CrN_3H , adapted with permission from ref 12. Copyright [2024] [John Wiley and Sons]. The dashed lines indicate the unit cell used in the computer calculations (see Section 3). The red lines indicate a HCa_6 octahedron.

spectra, serve as a fingerprint of both the configuration and dynamics of the H⁻ present. Sometimes, machine-learning molecular dynamics (MLMD) calculations can be chosen instead of traditional DFT approaches for the benefit of a reduced computational cost, allowing calculations in larger cells and at several temperatures, and because it can probe the anharmonic behavior of the vibrations.^{18,19} Using a combination of INS and MLMD calculations, we here investigate the configuration and vibrational dynamics of H⁻ in Ca₃CrN₃H. The results unravel significant interactions between neighboring H⁻ in a unique, chain-like, structural arrangement of H⁻, from which we can also determine the actual H⁻ stoichiometry in the sample.

2. EXPERIMENTAL DETAILS

2.1. Sample Synthesis. Approximately 4 g of Ca₃CrN₃H powder was prepared following the procedure as reported in ref 12. A starting material of Ca₃CrN₃ was first treated with H₂ gas (>99.999999%) at 450 °C for 2 h and then treated at 400 °C for 24 h (heating and cooling rates of 100 °C/h) with a mixed gas of N₂/H₂/Ar in the respective proportions of 22.5:67.5:10. The purity and crystal structure of the obtained powder sample was verified by PXRD at the beamline BL02B2 at the SPring-8 synchrotron X-ray facility in Japan.²⁰ BL02B2 was used with an incident photon wavelength of $\lambda = 0.41947(1)$ Å and a Debye–Scherrer camera, and the powder sample was contained in a Lindemann glass capillary of 0.3 mm inner diameter. All manipulations of the sample were carried out in a glovebox filled with He gas. The crystal structure was found to belong to the *P*6₃/*m* space group with unit-cell parameters *a* = 7.2280(3) Å and *c* = 5.0642(2) Å at 300 K.

2.2. INS. The INS measurements were performed on 4SEA-SONS^{21,22} at the Japan Proton Accelerator Research Complex and on IN1-Lagrange^{23,24} at the Institut Laue Langevin. Approximately 1.3 g of powder sample was evenly distributed inside an aluminum foil that was subsequently rolled into a 14 mm diameter and 44 mm long

cylinder. The sample was inserted into a cylindrical aluminum cell and sealed with a stainless steel metal ring.

The measured INS intensity contains information about vibrational dynamics in Ca₃CrN₃H and is proportional to the dynamical structure factor $S(Q, \omega)$, where Q and ω are the momentum and energy transfer, respectively. With regard to the measurements on 4SEASONS, we used incident energies of $E_i = 299$ (16.6 meV), 156 (6.8 meV), 95.2 (3.5 meV), and 64.2 meV (2.1 meV), where the values in parentheses refer to the energy resolutions at zero energy transfer. The frequency of the Fermi chopper was set to 600 Hz, and the analyzed Q and ω ranges were 0.5–22.5 Å⁻¹ and 5–260 meV, respectively. The measurements were performed at T = 7 K and lasted for 11 h. The data reduction was done with the Utsusemi software²⁵ and included normalization to monitor counts and efficiency correction of the detectors. With regard to the measurements on IN1-Lagrange, we used a Cu(220) monochromator and a pyrolytic graphite crystals analyzer, with which we analyzed the ω range of 40– 140 meV, with an energy resolution of \sim 2% of the incident energy. As an indirect-geometry spectrometer, IN1-Lagrange has a well-defined trajectory in (Q,ω) space, in which $\omega \propto Q^{2,26} S(Q,\omega)$ was obtained from 2 h long measurements upon heating from 10 to 300 K. The data reduction was done with the software Mantid²⁷ and included normalization to monitor counts. Mantid was also used for the data analysis of the reduced 4SEASONS and IN1-Lagrange data.

2.2.1. Theoretical Considerations. We note that the total cross section, σ , for Ca, Cr, N, and H is 2.8, 3.5, 12, and 82 barns,²⁸ respectively, which suggests that the predominant part of the scattering on Ca₃CrN₃H comes from H⁻ and N⁻. Moreover, we note that H⁻ scatters mainly incoherently (the incoherent cross section of H is 80 barns²⁸), whereas N⁻ scatters mainly coherently (the coherent cross section of N is 11 barns²⁸), which implies that the measured INS spectra will contain both incoherent and coherent contributions. However, at relatively large Q (i.e., when $Q \gg 2\pi/d \approx 2.5$ Å⁻¹, with $d \sim 2.5$ Å being the nearest-neighbor distance in Ca₃CrN₃H), the coherent intensity is smoothed out due to the finite Q resolution of the instrument.²⁹ As a result, for one-phonon scattering events, $S(Q, \omega)$ can be expressed in the incoherent approximation²⁹ as

$$S_{1\text{ph}}(Q, \omega) = \sum_{a} \frac{\sigma_a}{4\pi} \frac{Q^2}{3m_a} e^{-2W_a} \frac{\text{PDOS}_a(\omega)}{2\omega} (n(\omega) + 1)$$
(1)

where *a* indexes the atoms, m_a is the atomic mass, $2W_a = Q^2 u_a^2$ is the Debye–Waller exponent, ¹⁷ u_a^2 is the mean square displacement (msd) of atom *a*, PDOS_{*a*}(ω) denotes the partial vibrational density of state associated with atom *a*, and $n(\omega)$ is the phonon population factor. Inverting eq 1, the generalized DOS can be calculated as

$$DOS(\omega) = \sum_{a} \frac{\sigma_{a}}{3m_{a}} PDOS_{a}(\omega)$$
$$= \int dQS_{1ph}(Q, \omega) e^{2W_{a}} \frac{2\omega}{Q^{2}} \frac{4\pi}{n(\omega) + 1}$$
(2)

where $u_{\rm H}^2$ was obtained by fitting S(Q) profiles (see the Supporting Information), and takes on a value of approximately 0.02 Å².

3. COMPUTATIONAL DETAILS

The MLMD calculations were conducted using the package Graphics Processing Units Molecular Dynamics (GPUMD v3.9.5)^{30,31} based on a neural network-based machine-learning potential called neuroevolution potential (NEP v4),³¹ and they were used to compute vibrational properties of Ca₃CrN₃H_x with $x \leq 1$. The training procedure of the NEP was conducted following an approach successfully used before on similar systems,^{32–34} as described below.

3.1. NEP Training Procedure. First, an ab initio MD (AIMD) calculation was conducted in the NVT ensemble using the Vienna Ab-initio Simulation Package $(VASP)^{35-37}$ with a supercell containing 185 atoms, including 17 randomly

distributed H⁻ and 7 randomly distributed H⁻ vacancies $(V_{H^{-}})$, corresponding to the stoichiometry Ca₃CrN₃H_{0.71}, which is close to the H⁻ content previously estimated (0.7-1).¹² The thermostat was set at a temperature decreasing from 400 to 200 K, and the calculation was run during 8 ps by steps of 2 fs. VASP was set with the projector augmented wave (PAW)³⁸ method and the Perdew–Burke–Ernzerhof (PBE)³⁹ exchange-correlation functional. The calculation was conducted at the Γ -point, and a cutoff energy of 500 eV was applied to the plane wave basis. Electronic structures were converged to energies within 10^{-5} eV. In the AIMD trajectory, snapshots were extracted every 100 fs (a total of 72 snapshots), and the energy, forces, and stresses in each snapshot were used to train a NEP (further referred to as the first-generation NEP). The NEP training used radial and angular descriptor cutoffs at 8 and 4 Å, respectively, 30 neurons, and converged within the 1000000 maximal number of generations. Second, the first-generation NEP was used to run MD simulations of 20 ps on 10 supercells including 0-18 randomly distributed V_{H} , corresponding to the stoichiometries $Ca_3CrN_3H_x$ with x in the range of 0.25-1, at 100, 200, 400, and 600 K using the NPT ensemble. In each trajectory, snapshots every 100 fs were taken after the temperature and volume stabilization (a total of 400 snapshots). The energy, forces, and stresses in each snapshot were subsequently computed with VASP using the same settings as above, which constituted the second-generation training data set. Finally, a new NEP was trained using the firstand second-generation data sets (a total of 472 snapshots), with the same descriptors, neurons, and maximal number of generations as above. This NEP accurately reproduces the forces and energies calculated with DFT (see Supporting Information for details) and is the one used in further calculations.

3.2. Harmonic Phonons and Dispersion Curves. The vibrational modes and energies were first calculated in the harmonic approximation using the NEP with a finite displacements approach as implemented in the software packages Phonopy^{40,41} and Calorine.³¹ A supercell of $2 \times 2 \times 2$ unit cells (a unit cell is depicted in Figure 1), containing 128 atoms, was used to model Ca₃CrN₃H, and a similar supercell containing 120 atoms was used to model Ca₃CrN₃H_{0.5}. These supercells were relaxed to forces below 10^{-4} eV/Å using the NEP. Then, the force constants were calculated using the NEP and used to compute the phonon eigenvectors and eigenfrequencies. The atom-projected (partial) vibrational density of states (PDOS) was calculated on a *q*-mesh of $31 \times 31 \times 31$ points and can be expressed as⁴⁰

$$PDOS_{a}(\omega, \ \hat{\mathbf{n}}) = \frac{1}{N} \sum_{(\mathbf{q}, \nu)} \delta(\omega - \omega_{\nu}(\mathbf{q})) |\hat{\mathbf{n}} \cdot \mathbf{e}_{\nu}(a, \ \mathbf{q})|^{2}$$
(3)

Here, N = 8 is the number of unit cells in the calculation, *a* is the atom index, $\hat{\mathbf{n}}$ is the unit projection direction vector, and $\omega_{\nu}(\mathbf{q})$ and $\mathbf{e}_{\nu}(a, \mathbf{q})$ are the eigenfrequency and eigenvector of mode (\mathbf{q}, ν) . The phonon dispersion curves were calculated along a hexagonal path of 459 *q*-points within the hexagonal Brillouin zone.

The validity of the harmonic phonon calculations using the NEP was confirmed by repeating the calculation using DFT to compute the force constants (details in the Supporting Information).

The experimental $S(Q, \omega)$ was simulated with the software package OCLIMAX,⁴² using the calculated PDOS and the geometry of 4SEASONS. The calculation included multiphonon contributions up to the 10th order.

3.3. MLMD Vibrational Density of State. The NEP was finally used to run MD simulations of $Ca_3CrN_3H_x$ with *x* in the range of 0.64–1, modeled by large supercells containing 1466–1536 atoms, and at temperatures in the range of 10–300 K. The simulations were first conducted during 200 ps in the NPT ensemble for temperature and volume stabilization and then conducted during 100 ps in the NVE ensemble for the vibrational DOS analysis. Additional calculations were conducted by using the NVT ensemble for temperature and pressure stabilization, followed by a run with the NVE ensemble for analysis. The PDOS was computed from the Fourier transform of the velocity autocorrelation function, as implemented in GPUMD.

Due to the presence of H⁻ vacancies, 3 inequivalent H⁻ configurations can be found (considering only nearest neighboring H⁻ sites) in the supercells, here labeled as H⁻– H⁻–H⁻, H⁻– H⁻– V_{H⁻}, and V_{H⁻}– H⁻– V_{H⁻} (see Figure 2). H⁻–H⁻–H⁻ corresponds to H⁻ which are flanked by two



Figure 2. Schematics of the 3 inequivalent H^- configurations existing in the AIMD calculation.

 H^- in first neighbor positions, $H^- - H^- - V_{H^-}$ corresponds to H^- having a unique H^- first neighbor, and $V_{H^-} - H^- - V_{H^-}$ corresponds to H^- which are isolated with no first neighbor H^- .

4. RESULTS AND DISCUSSION

4.1. INS Spectra. Figure 3a shows the $S(Q, \omega)$ map of Ca₃CrN₃H, as measured on 4SEASONS at 7 K for the four different incident energies. Because the intensity below 40 meV contains contributions from the aluminum container, which was not subjected to a separate measurement in our experiment, this part was ignored in our detailed analysis. Above 40 meV, the INS data are characterized by a series of bands; however, for energies higher than 140 meV, the bands are related to second-order transitions, i.e., to scattering events involving two-phonon modes,¹⁷ which will not be analyzed explicitly here. The assignment to second-order transitions is supported both by our computational results and the experimental band energies, which are found to be multiples of the energies of the respective fundamental modes in the interval 90-130 meV (see Supporting Information). For our more detailed analysis, we hence focused on the energy range between 40 and 140 meV. This analysis showed that the INS spectrum can be adequately reproduced by a sum of 8, wellseparated Gaussian components ($\omega_1 - \omega_8$), plus a flat background (Figure 3b). For comparison, Figure 3b,c show



Figure 3. (a) Combined presentation of overlaying $S(Q, \omega)$ maps (A, B, C, D) of Ca₃CrN₃H, as measured on 4SEASONS at 7 K with four incident neutron energies: 64.2 meV (A), 95.2 meV (B), 156 meV (C), and 299 meV (D). In order to be comparable, spectra A, B and C were normalized such that the intensity between 20 and 40 meV is continuous along Q. Spectrum D was normalized such that the maximum intensity at 95 meV matches the maximum intensity of spectrum C at 95 meV. (b) $S(\omega)$ obtained by Q-integration of spectrum C, together with a fit of the peaks with Gaussians (shaded areas) and a theoretical INS spectrum of Ca₃CrN₃H obtained from the phonon calculation and OCLIMAX (dashed line). (c) PDOS of Ca₃CrN₃H from the phonon calculation using the NEP, weighted by the total neutron cross section and the atomic mass of each element. Ha⁻_a, Hb⁻_b, and Hc⁻_c denote the PDOS of H⁻ projected along the a, b, and c axis, respectively.

the theoretical spectrum, as obtained from the phonon calculation using NEP. As can be observed, the theoretical spectrum shows the same main features and is overall similar to the experimental spectrum, which suggests that the modeled structure provides a good description of the structure of the real material. Therefore, by comparing the experimental and calculated vibrational energies, we can assign the bands to specific vibrational motions. The suggested band assignment is summarized in Table 1. As can be seen from the table, the calculation successfully models the vibrational energies of the Cr–N bending mode (ω_1), with a deviation of less than 3 meV from the experimental data. However, the contributions from hydrogen of the theoretical spectrum are globally downshifted with 3–9 meV compared to the experimental data. This can be attributed to a general overestimation of the equilibrium

Table 1. Experimental Vibrational Energies (in meV) from Fits with Gaussian Components (See Figure 3b), Calculated Frequencies (in meV), and Suggested Band Assignment for Ca₃CrN₃H

band	exp.	calc.	proposed assignment
ω_1	50.7	48.5	Cr–N bending motions
ω_2	86.9	81.2	in-phase H^- motions along c
ω_3^{a}	91.3		
ω_4	94.9	88.2	H ⁻ motions in <i>ab</i> -plane
ω_5	97.8	91.1	H ⁻ motions in <i>ab</i> -plane
ω_6	106	104	H^- motions along <i>a</i> , <i>b</i> , and <i>c</i>
ω_7	113	110	H ⁻ motions along <i>a</i> , <i>b</i> , and <i>c</i>
ω_8	125	116	out-of-phase H^- motions along c
			-file

 ${}^a\omega_3$ accounts for the asymmetry of the surrounding ω_2 and ω_4 peaks and was not calculated.

volume by the PBE functional, as it was already observed in $BaTiO_3$ and the oxide-hydride $BaTiO_{2.75}H_{0.125}^{\ 14,43}$

4.2. Hydrogen Local Coordination Environment. We observe that the in-phase and out-of-phase vibrations (ω_2 and ω_8 , respectively) of H along the *c* axis (H_c⁻) are located at different energies, indicating that the H_c⁻ vibrations are dispersive, which is confirmed by the phonon dispersion curve of Ca₃CrN₃H, as shown in Figure 4a. Such a dispersive



Figure 4. Dispersion curve and PDOS of (a) Ca_3CrN_3H and (b) $Ca_3CrN_3H_{0.5}$, as obtained from the NEP in harmonic calculations. The purple line in the dispersion curves highlights the dispersive and nondispersive character of the H_c^- vibrations in Ca_3CrN_3H and $Ca_3CrN_3H_{0.5}$, respectively.

character reflects a significant interaction between neighboring H⁻ anions, due to their relative close proximity (2.5 Å along the *c* axis). Conversely, no dispersion is expected in Ca₃CrN₃H_{0.5} (see Figure 4b), as the lack of H⁻ in every other H⁻ site extends their mutual distance up to 5 Å along the *c* axis, thereby reducing their interaction. As a result, in Ca₃CrN₃H_{0.5}, the PDOS of H⁻_c has a higher intensity and a lower vibrational energy than in Ca₃CrN₃H. These observations imply that the vibrational dynamics of H⁻ are correlated to the occupation of their surrounding H⁻ sites, and, as a

result, that the average H^- occupation in Ca₃CrN₃H can be probed *via* its INS spectrum.

We turn now to the MLMD calculations of the compositions $Ca_3CrN_3H_x$ with x in the range of 0.64–1. Figure 5 shows the



Figure 5. Comparison of the experimental and computed DOS: (blue lines) DOS for $Ca_3CrN_3H_x$ with *x* in the 0.64–1 range extracted from the MLMD trajectories at 100 K, weighted by the total neutron cross section and atomic masses. (orange lines) DOS for Ca_3CrN_3H and $Ca_3CrN_3H_{0.5}$ from the phonon calculation, weighted by the total neutron cross section and the atomic masses. (black lines) experimental DOS extracted from the INS data using eq 2 with (continuous black line) the data recorded at $E_i = 156$ meV and (dashed black line) the data recorded at $E_i = 95.2$ meV. The computed DOS lines are shifted vertically for clarity and scaled horizontally by a factor of 1.055 in order to match the experimental data.

computed DOS for the various H⁻ contents, together with the experimental DOS. A main observation is the excellent agreement between the computed and experimental DOS for x = 1. With decreasing x, we observe a gradual decrease in the intensity of the high-energy peak ω_{8} , as related to out-of-phase vibrations, and a down-shift and broadening in the energy of the peak ω_2 , related to in-phase vibrations, already at x = 0.95. In the experiment, no such down-shift is observed, and the peak ω_8 is well visible, which indicates that for our sample x > 0.95. We note that this high H⁻ content may be at the origin of the efficient ammonia synthesis activity of Ca₃CrN₃H_x, as the surface H⁻ is suspected to participate in the hydrogenation of the N₂ molecules.¹³

The down-shift of the ω_2 peak and the decrease in intensity of the ω_8 peak with decreasing x can be understood by separating the contributions of each H⁻ configuration in the PDOS, as shown in Figure 6 for x = 0.64. The PDOS for the 3 configurations are overall quite similar but show some marked differences. In particular, we observe that the ω_8 peak is only present for the configuration $H^--H^--H^-$, whereas the ω_2 peak down-shifts in energy and increases in intensity for V_H-- $H^--V_{H^-}$. As a result, it is the transformation of configurations of the type $H^--H^--H^-$ into configurations of the type $H^- H^--V_{H^-}$ and $V_{H^-}-H^--V_{H^-}$ with decreasing *x* which induces a decrease in intensity of the ω_8 peak and a down-shift and broadening of the ω_2 peak. Another observation is a weak down-shift of the peaks ω_3 , ω_4 , and ω_5 (referred to as the ensemble $\omega_{3,4,5}$ with decreasing *x*. $\omega_{3,4,5}$ is related to vibrations in the *ab*-plane, and the observed down-shift is due to an



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Figure 6. PDOS from each H⁻ configuration in Ca₃CrN₃H_{0.64} extracted from the MLMD trajectory at 100 K. The supercell includes 43 configurations of the type H⁻-H⁻-H⁻, 62 configurations of the type H⁻-H⁻ - H⁻, and 17 configurations of the type V_{H⁻} - H⁻ - V_{H⁻}. The lines are scaled horizontally by a factor of 1.055 in order to match the experimental data.

increase of the lattice constants a and b with decreasing x (see Supporting Information for details).

4.3. Temperature Dependence. Next, we turn to investigate how the H^- vibrational dynamics are affected by temperature. Figure 7 shows variable temperature (10–300 K) INS spectra, as measured on IN1-Lagrange, together with the



Figure 7. (a) $S(\omega)$ of Ca₃CrN₃H as measured on IN1-Lagrange at various temperatures. Insets show close-up views for the regions around 80 and 120 meV, respectively. (b) Computed PDOS of H⁻ for Ca₃CrN₃H as extracted from the MLMD trajectories stabilized in the NPT ensemble at the same temperatures. The PDOS lines are scaled horizontally by a factor of 1.055 in order to match the experimental data.

computed PDOS of H⁻ for Ca₃CrN₃H, in the same temperature range. In the experiment, the most pronounced effects of increasing the temperature are a gradual decrease in intensity by approximately 30% of the ensemble of peaks $\omega_{3,4,5}$, a broadening of the ω_2 peak, and a broad increase in intensity around 80 meV. The gradual decrease in intensity can be partially attributed to the Debye–Waller factor, as the msd of H⁻ increases with increasing temperature. However, the comparison with the MLMD calculations shows that most of it is to be attributed to a decrease in intensity and a broadening of the ensemble of peaks $\omega_{3,4,5}$ in the PDOS of H⁻. Similarly, the broadening of the ω_2 peak and the increase in intensity around 80 meV is well reproduced in the MLMD calculations, which indicates that these features originate from a change in the PDOS of H⁻.

Overall, the temperature-dependent features observed experimentally can be attributed to a global broadening of PDOS with increasing temperature. Such a broadening is due to phonon-phonon interactions resulting in an anharmonic behavior of the related modes.⁴⁴ Specifically, an analysis of the NEP shows that one of the modes associated with the ensemble of peaks $\omega_{3,4,5}$ deviates significantly from a harmonic behavior (see Supporting Information for details). Moreover, it is found that the ensemble of peaks $\omega_{3,4,5}$ and the peak ω_2 are upshifted in the PDOS when the MLMD calculation is conducted in the NVT ensemble, that is, without volume stabilization, which also indicates anharmonicity. In the NPT ensemble, as the calculation is conducted here, the upshift due to anharmonicity is compensated by a global down-shift due to thermal expansion (see Supporting Information for details). Anharmonic H⁻ vibrations in Ca₃CrN₃H could suggest additional motions of H⁻, such as rattling motions within the HCa₆ octahedra,^{44,45} or may be due to the interaction of the *s* electron pairs of H^- anions with the nearby *p* electrons of Ca²⁺ cations, known as the lone pair electron mechanism.⁴ Interestingly, we note that Ca₃CrN₃H is electrically conducting,¹² and anharmonicity generally leads to a reduction of the thermal conductivity,⁴⁶ which suggests that this material may have good thermoelectric properties.

Furthermore, we note that the ω_8 peak does not show a significant temperature dependence in the experiment, while it broadens and up-shifts in the calculated PDOS. This apparent mismatch may indicate that the anharmonic features of the out-of-phase H⁻ vibrations are not captured in detail by the calculation. The weak temperature dependence of the peaks ω_6 and ω_7 observed in the experiment and in the PDOS may be due to a significant coupling between the H⁻ motions probed here and the N³⁻ vibrations, which are weakly temperature dependent. In fact, the msd of N³⁻ is weakly temperature dependent in the range of 10-300 K due to stiff Cr-N bonds (according to the DFT calculations, the Cr-N bonds are approximately 50 times stronger than the Ca-H bonds), and the coupling of H⁻ vibrations with this mode is indicated by a strong participation ratio of N at the frequencies ω_6 and ω_7 (see Supporting Information for details).

5. CONCLUSIONS

To conclude, our combined INS and MLMD simulation study of the technologically relevant novel nitride-hydride Ca_3CrN_3H has unravelled the vibrational character of H⁻, as well as the local configuration of H⁻ in this material's unique, chain-like structural arrangement of H⁻. In particular, the results reveal a generally anharmonic dynamics of H⁻ with significant interactions between neighboring H⁻ in the material, and it is found that most H⁻ are surrounded by two another H⁻ and that the H⁻ sites have an occupancy of at least 95%. This high H⁻ occupancy may be related to the material's high efficiency as a catalyst for ammonia synthesis. In effect, our results indicate that the vibrational properties of Ca₃CrN₃H may be used as indicators for catalytic activity. Such an approach may as well be applicable to other mixed-anion materials, such as Ca₂N: e⁻, a two-dimensional electride turning into the nitride-hydride Ca₂NH during catalytic activity,⁴⁷ or the oxide-hydride BaTiO_{2.5}H_{0.5} in which the lattice hydride also plays a crucial role in the catalytic activity.⁴⁸

ASSOCIATED CONTENT

3 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.4c02897.

Additional INS data and additional calculation data (PDF)

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Notes

The authors declare no competing financial interest.

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