

Neutron Characterization in support of the DOE Hydrogen Storage Sub-Program

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ST067

Timeline

- Project start date: 10/2010
- Project end date: 10/2014*

*Project continuation and direction determined annually by DOE

Budget

- FY13 DOE Funding: \$200K
- Planned FY14 DOE Funding: \$200K
- Total DOE Project Value: \$600K

[NIST continues to provide access to neutron facilities and FTEs for the DOE Hydrogen Storage Projects.](#)

NIST Associates

Nina Verdal	Hui Wu
John J. Rush	Wei Zhou
Matt Hudson	Wan Si Tang
Zeric Hulvey	

Barriers addressed

- A. System Weight and Volume
- O. Lack of Understanding of Hydrogen Physisorption and Chemisorption

Partners

Caltech, Delaware, GM, HRL, Maryland, Michigan, Missouri-St. Louis, Molecular Foundry, NREL, Ohio State, Penn, Penn State, RCB Hydrides, Sandia, UC Berkeley, Utah, Washington U.-St. Louis

Overall: Support the DOE-EERE-funded hydrogen-storage projects by providing timely, comprehensive characterization of materials and storage systems using state-of-the-art neutron methods.

- Direct partner synthesis efforts based on the understanding gained through the use of these methods.
- Demonstrate the fundamental characteristics of useful hydrogen-storage materials.

Neutron methods

- determine elemental compositions of materials (prompt- γ activation analysis and neutron reflectometry of H stoichiometries and profiles)
- determine location of H and crystal structures of materials (neutron diffraction superior to XRD for “seeing” light H and D)
- determine bonding of absorbed H (unlike IR and Raman, neutron vibrational spectroscopy “sees” all H vibrations for straightforward comparison with first-principles calculations)
- elucidate H diffusion mechanisms (faster dynamics timescale of neutron quasielastic scattering complements NMR; transport mechanisms gleaned from momentum transfer dependence)

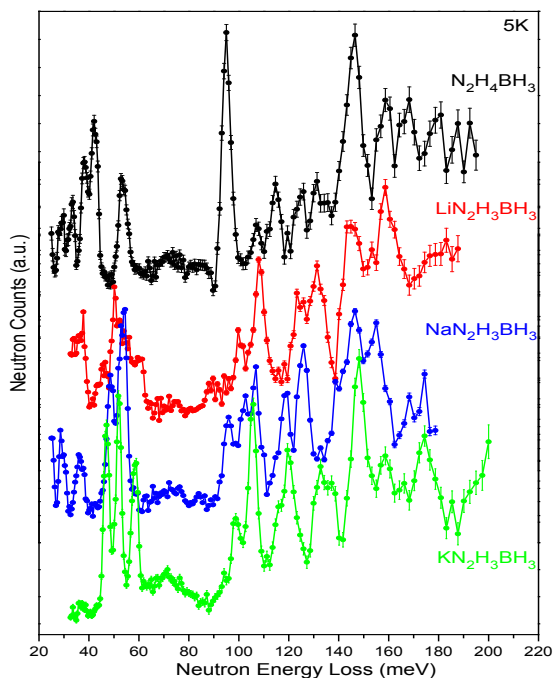
Technical Accomplishments

Neutron methods, synthesis expertise, and DFT computations were used to further the goals of the DOE Hydrogen-Storage Sub-Program.

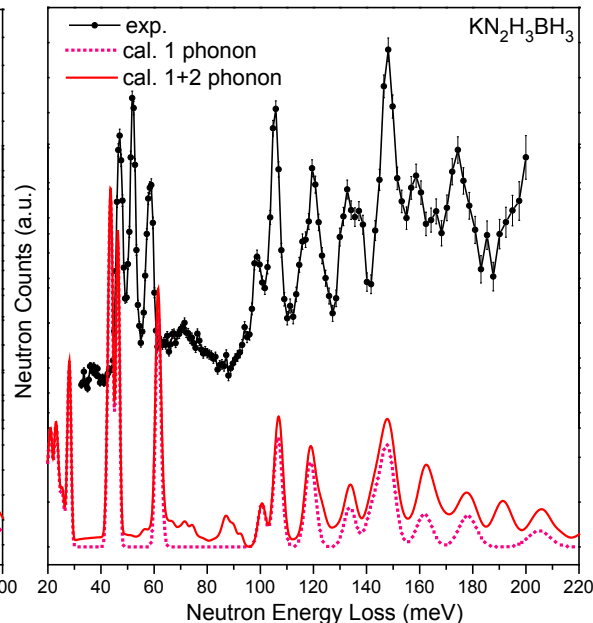
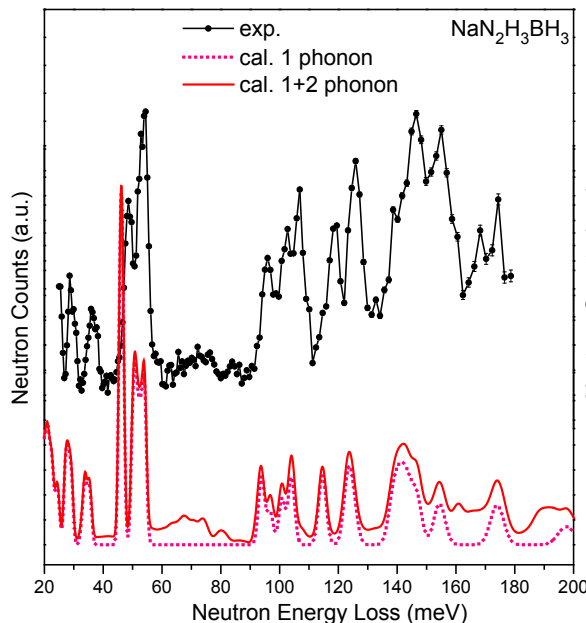
Synthesis and Characterization of Novel Compounds **NIST**

Collaboration with Maryland, GM, Dalian Institute of Chemical Physics, Sichuan University

Motivation: synthesizing new compounds potentially promising for hydrogen storage



Comparison of neutron vibrational spectra for $N_2H_4BH_3$ and $AN_2H_3BH_3$ (A = Li, Na, K)



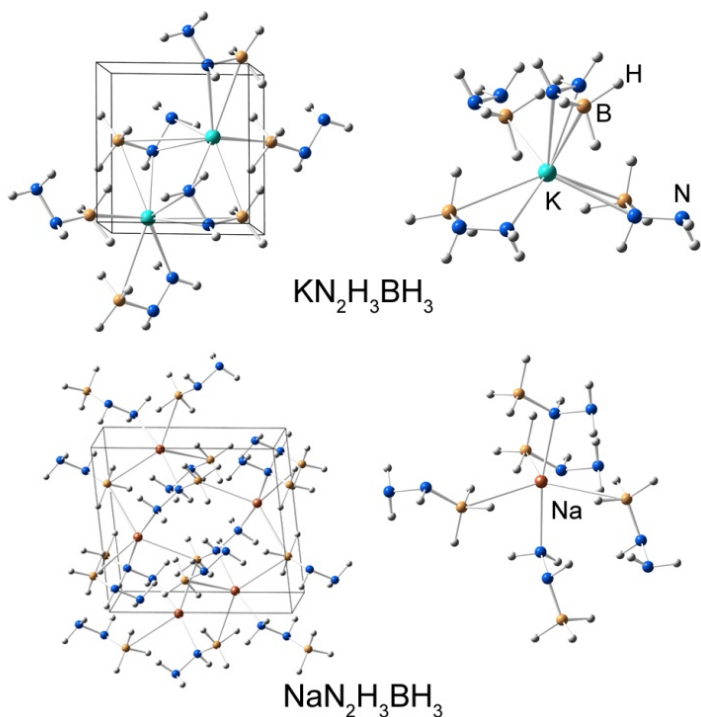
DFT phonon simulations are in agreement with neutron vibrational spectra for the recently synthesized $NaN_2H_3BH_3$ and $KN_2H_3BH_3$.

Agreement between neutron vibrational spectroscopy and density functional theory corroborates structures determined by diffraction methods.

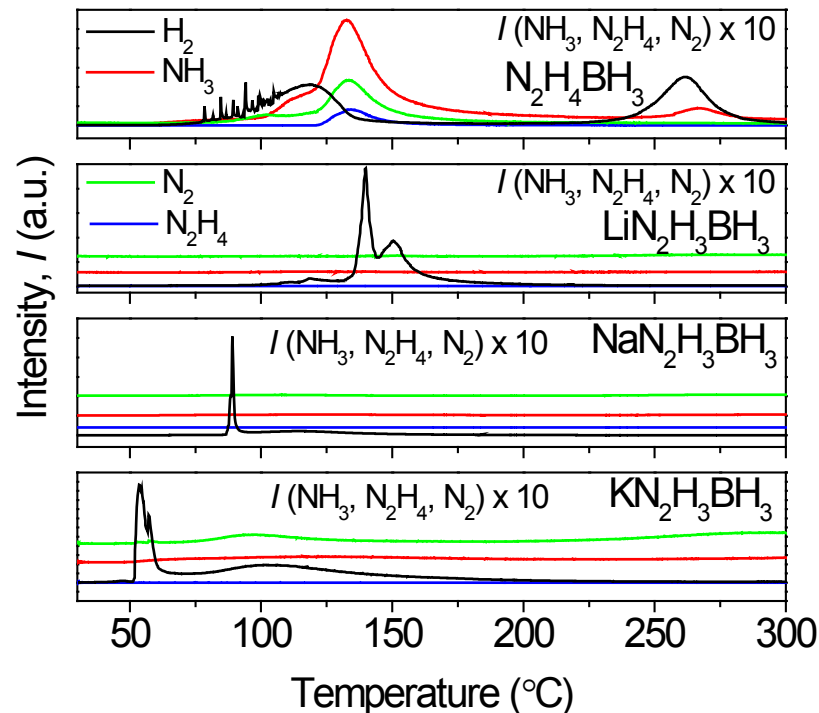
Synthesis and Characterization of Novel Compounds **NLST**

Collaboration with Maryland, GM, Dalian Institute of Chemical Physics, Sichuan University

Motivation: synthesizing new compounds potentially promising for hydrogen storage



New $\text{NaN}_2\text{H}_3\text{BH}_3$ and $\text{KN}_2\text{H}_3\text{BH}_3$ structures determined from diffraction data



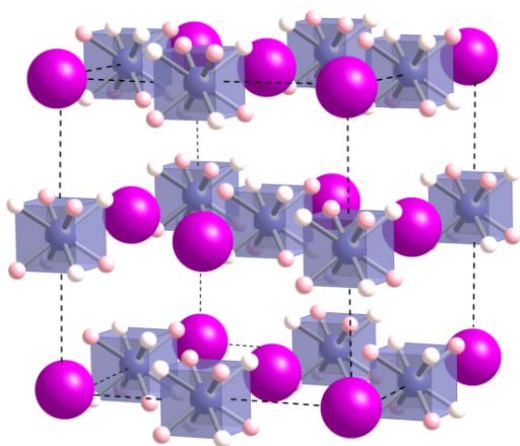
TPD-MS measurements of $\text{N}_2\text{H}_4\text{BH}_3$, $\text{LiN}_2\text{H}_3\text{BH}_3$, $\text{NaN}_2\text{H}_3\text{BH}_3$ and $\text{KN}_2\text{H}_3\text{BH}_3$

Compared to $\text{N}_2\text{H}_4\text{BH}_3$, the alkali-metal substituted hydrazidotrihydridoborates display improved dehydrogenation behavior with no N_2H_4 emission and greatly suppressed NH_3 release.

Subtle Borohydride Phase Transitions

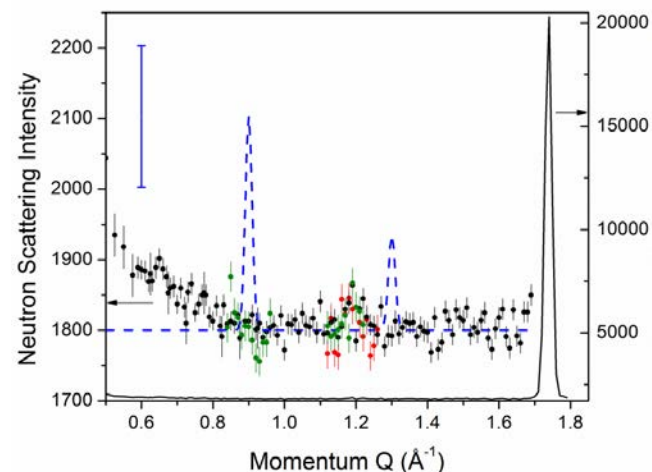
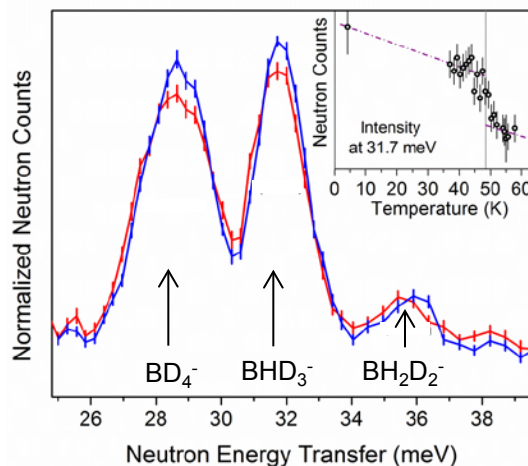
Collaboration with Maryland, Michigan, Delft University

Motivation: characterizing the bonding in borohydride materials



A disordered RbBD₄ structure was determined from NPD data at both 5 K and 60 K.

Neutron vibrational spectra of torsions in RbBD₄ (with 4 % H) at 38 K (blue) and 55 K (red). Inset: neutron scattering intensity at 31.7 meV vs. temperature, denoting the phase transition at 48.5 K.

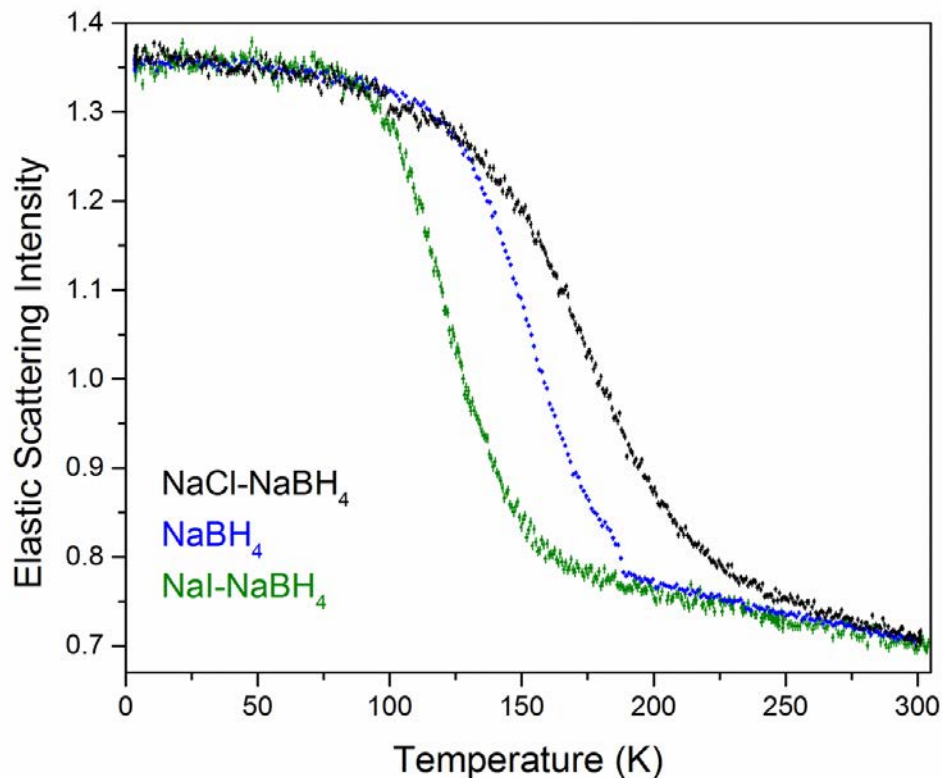


Neutron powder diffraction data for RbBD₄ at low Q below (4 K, green; 38 K, black) and above (58 K, red) the phase transition. Dashed lines represent previously suggested superlattice peaks at 4 K.

Neutron diffraction/spectroscopy suggest that only short-range BD₄⁻ order exists in RbBD₄. 8

Motivation: understanding BH₄⁻ rotational dynamics in modified NaBH₄

Neutron-Scattering Fixed-Window Scans for Bulk and Halide-Anion-Substituted NaBH₄



Anion Sizes
I⁻ > BH₄⁻ > Cl⁻

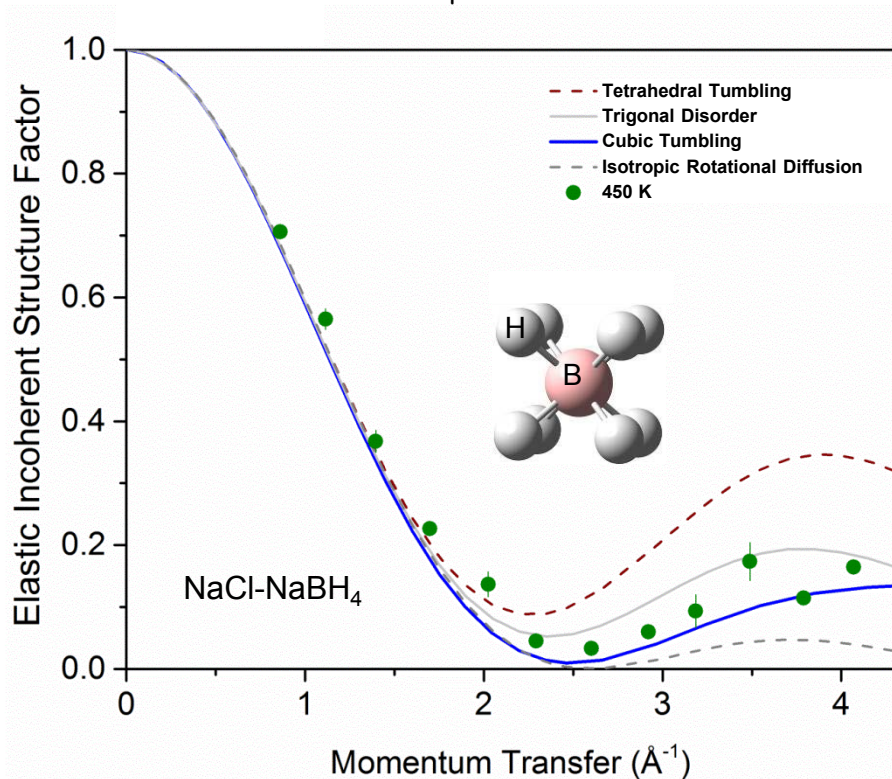
As the average anion size increases:

- The BH₄⁻ rotational jump rate increases at a given temperature.
- the temperature corresponding to the onset of rapid rotations decreases.

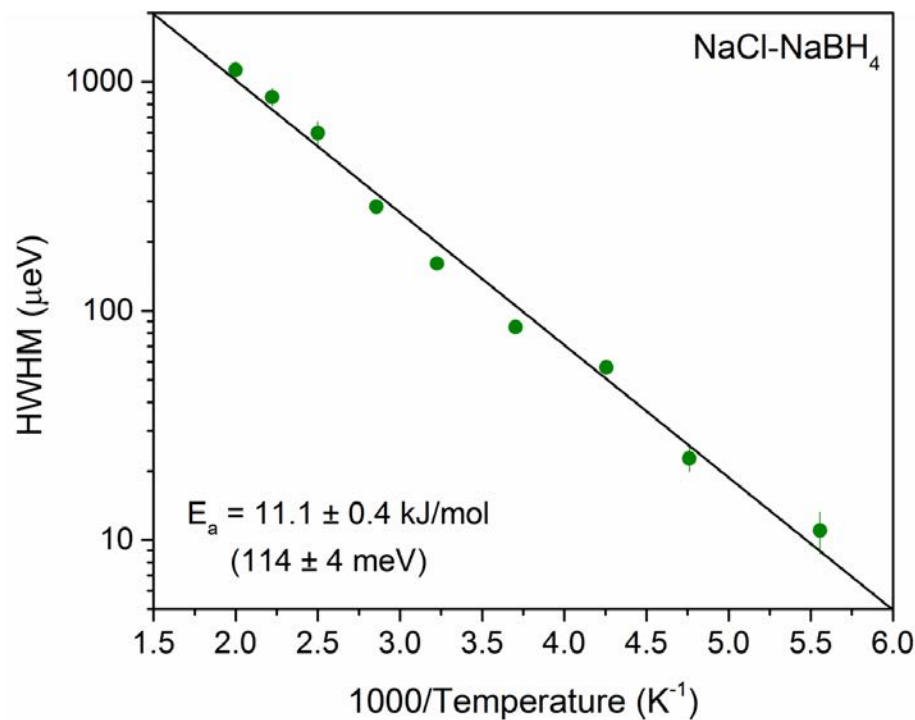
Fixed-window scans allow for a relative comparison of BH₄⁻ anion rotational mobilities.

Motivation: understanding BH₄⁻ rotational dynamics in modified NaBH₄

Elastic Incoherent Structure Factor (EISF) data for NaCl-NaBH₄



The activation energy for BH₄⁻ reorientation is similar to that for bulk NaBH₄.



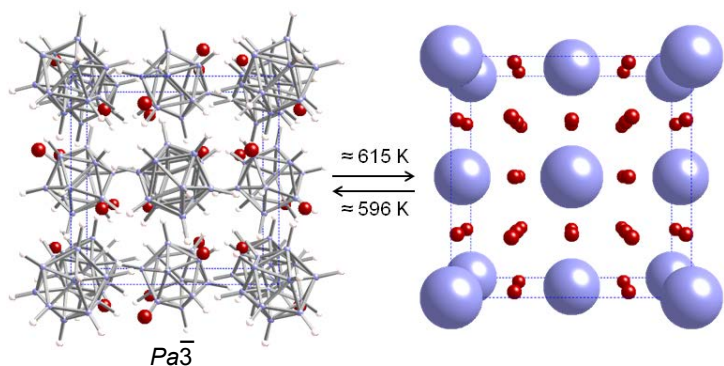
The cubic tumbling mechanism, where the H atoms jump to the eight corners of a cube, is in best agreement with the EISF data. 10

Li₂B₁₂H₁₂ and Na₂B₁₂H₁₂ Phase Transitions

Collaboration with Maryland, GE, Sandia, IMP Ekaterinburg

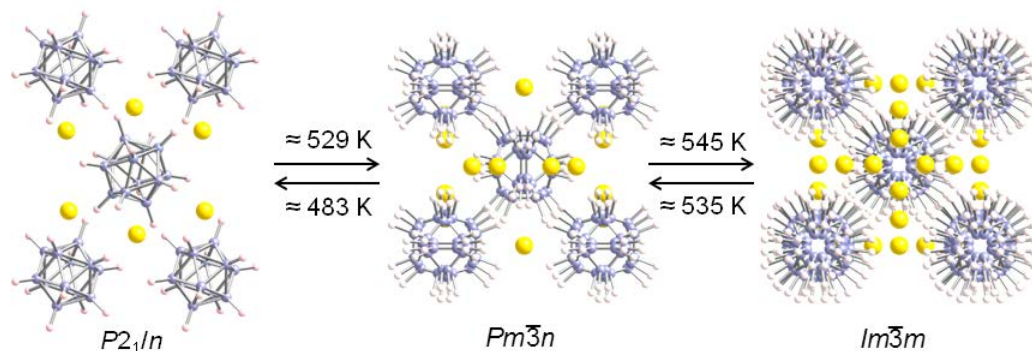
Motivation: characterizing borohydride dehydrogenation products

X-ray and neutron powder diffraction were used to elucidate the structures of Li₂B₁₂H₁₂ and Na₂B₁₂H₁₂ below and above their phase transitions near 615 K and 529 K, respectively.



Proposed Li₂B₁₂H₁₂ transition from an ordered cubic structure to an orientationally disordered anion sublattice harboring disordered cation sites

Na₂B₁₂H₁₂ transition from an ordered monoclinic structure to a sequence of (anion and cation) disordered quasi-body-centered-cubic structures



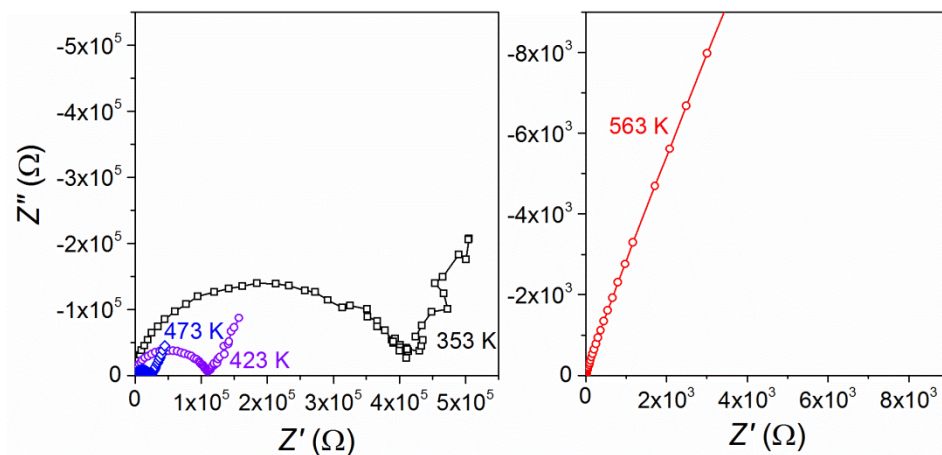
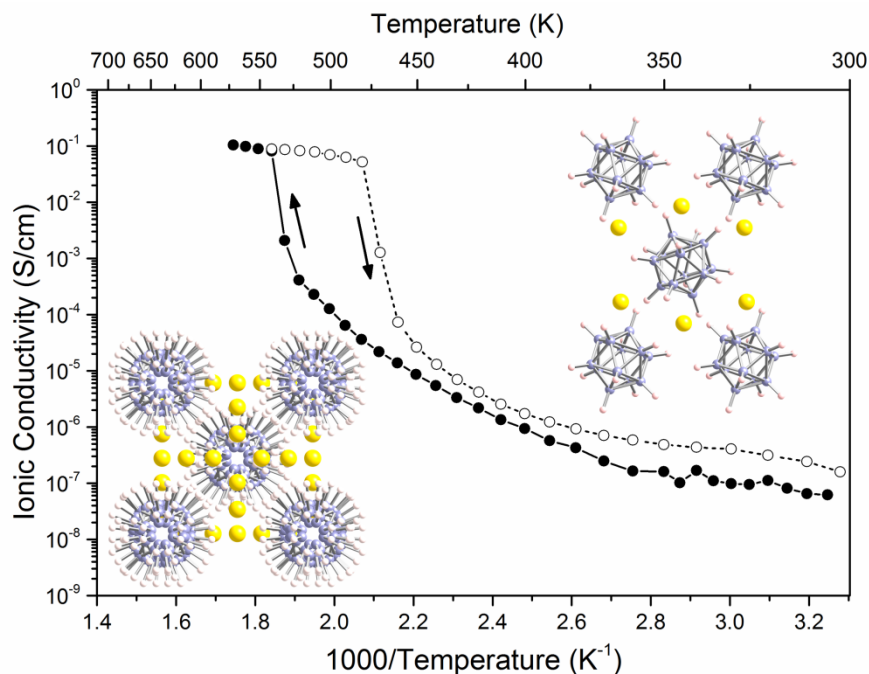
Both Li₂B₁₂H₁₂ and Na₂B₁₂H₁₂ undergo order-disorder phase transitions resulting in orientationally mobile anions and vacancy-rich cation sublattices.

Sodium Superionic Conduction in $\text{Na}_2\text{B}_{12}\text{H}_{12}$

Collaboration with Maryland, GE, Sandia, IMP Ekaterinburg, Tohoku University

Motivation: characterizing conductivity properties of cubic $\text{Na}_2\text{B}_{12}\text{H}_{12}$

AC impedance measurements of $\text{Na}_2\text{B}_{12}\text{H}_{12}$
ionic conductivity vs. temperature

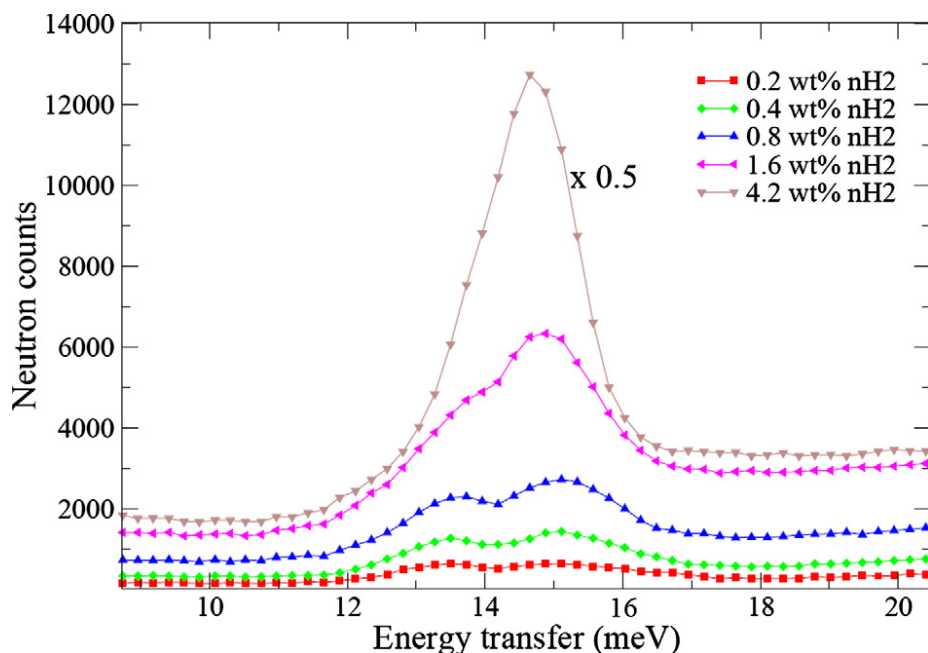


Complex impedance plots of $\text{Na}_2\text{B}_{12}\text{H}_{12}$ at various temperatures during heating run

$\text{Na}_2\text{B}_{12}\text{H}_{12}$ exhibits dramatic superionicity above its order-disorder phase transition, aided by the large mobile anions and the appearance of cation vacancies.

Motivation: investigating surface compression effects in C nanopores

Neutron vibrational spectra at 4 K of normal H₂ adsorbed in polyfurfuryl-alcohol-derived nanoporous carbon (PFA-NPC40) at different loadings



The rotational tunneling behavior of adsorbed H₂ in this nanoporous carbon confirmed the hindrance to rotational motion.

Measurements using heavier gases showed similar changes in the nature of the lateral forces during adsorption. Lateral forces were attractive at low uptakes and pressures but became repulsive at higher uptakes and pressures.

Adsorbates at high uptakes can be in a highly compressed state with the appearance of repulsive lateral forces that significantly affect the energetics of adsorption.

Collaborations

U.S. Partners (Type of Institution): What we provide to them

CalTech (Univ.): neutron measurements of various porous materials and borohydrides.

Delaware (Univ.): neutron measurements of various zeolites.

GE (Industry): neutron characterization of complex hydrides.

GM (Industry): neutron characterization of complex hydrides.

HRL (Industry) neutron measurements of BH_4^- dynamics in nanoconfined metal borohydrides.

Maryland (Univ.): neutron and x-ray measurements of alanates, borohydrides, and related materials.

Michigan (Univ.): neutron measurements of BH_4^- dynamics in neat and nanoconfined metal borohydrides.

Missouri-St. Louis (Univ.): neutron measurements of various alanates, borohydrides, and related materials.

NREL (Gov.): neutron measurements / isotherm measurements of various spillover related materials.

Ohio State (Univ.): neutron and x-ray measurements of borated compounds.

Penn (Univ.): neutron measurements of amidoboranes and other storage-related materials.

Penn State (Univ.): neutron measurements of various boron doped carbons.

RCB Hydrides (Industry): neutron and x-ray measurements and DFT calculations of various alanates, borohydrides, and related materials; neutron imaging of hydrogen-storage beds.

Sandia (Gov.): neutron measurements and DFT calculations of various alanates, borohydrides, and related materials.

Molecular Foundry (Gov.): neutron measurements of various metal-organic framework materials

U.C. Berkeley (Univ.): neutron measurements of various alanates, borohydrides, and related materials.

Remainder of FY2014:

- Continue to perform neutron-based structural and spectroscopic characterizations of new materials in conjunction with the needs of the DOE-funded projects, including novel bulk and nanoconfined complex hydride materials.
- For example, start characterization work on new hydrogenated metal silicides

Neutron methods continue to provide crucial, non-destructive characterization tools for the DOE Hydrogen-Storage Sub-Program.

- Agreement between NVS and DFT corroborates the recently synthesized $\text{NaN}_2\text{H}_3\text{BH}_3$ and $\text{KN}_2\text{H}_3\text{BH}_3$ structures that were determined by diffraction methods.
- Compared to $\text{N}_2\text{H}_4\text{BH}_3$, the alkali-metal substituted hydrazidotrihydridoborates display improved dehydrogenation behavior with no N_2H_4 emission and greatly suppressed NH_3 release.
- Neutron diffraction and spectroscopy indicate that BD_4^- anion ordering in RbBD_4 does exist but is short-ranged and does not involve any superlattice ordering.
- Neutron-scattering fixed-window scans are shown to be valuable for making relative comparisons of BH_4^- anion rotational mobilities in bulk and halide-substituted NaBH_4 .
- The cubic tumbling mechanism, where the H atoms jump to the eight corners of a cube, is in best agreement with the quasielastic neutron scattering data for NaCl-NaBH_4 solid solution above room temperature.
- $\text{Li}_2\text{B}_{12}\text{H}_{12}$ and $\text{Na}_2\text{B}_{12}\text{H}_{12}$ undergo order-disorder phase transitions at ~ 615 K and 529 K, respectively, resulting in orientationally mobile anions and vacancy-rich cation sublattices.
- $\text{Na}_2\text{B}_{12}\text{H}_{12}$ exhibits dramatic superionicity above its order-disorder phase transition, aided by the large mobile anions and the appearance of cation vacancies.
- Molecular hydrogen and other gaseous adsorbates at high uptakes in nanoporous carbon can be in a highly compressed state with the appearance of repulsive lateral forces that significantly affect the energetics of adsorption.