

HyMARC: Technical Activities at NIST

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National Renewable Energy Laboratory & National Institute of Standards and Technology 2019 DOE Annual Merit Review

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Overview



Timeline and Budget

• Timeline:

Phase 1: 10/1/2015 to 9/30/2018 Phase 2: 10/1/2018 to 9/30/2022

• Budget:

2 post-docs are provided for this effort through HyMARC/NREL

• Barriers Addressed:

General:

A. Cost; B. Weight and Volume; C. Efficiency;

E. Refueling Time;

Reversible Solid-State Material:

- M. Hydrogen Capacity and Reversibility;
- N. Understanding of Hydrogen Physi- and Chemisorption;

O. Test Protocols and Evaluation Facilities;

HyMARC Collaborators

- LBNL Jeff Long, Martin Head-Gordon
- PNNL Tom Autrey, Mark Bowden
- SNL Vitalie Stavila,
- LLNL Brandon Wood
- HyMARC seedling Eric Majzoub (Univ. Missouri-St. Louis)













An synergistic collaboration and research effort among HyMARC participants to:

- <u>develop</u> and <u>enhance</u> hydrogen-storage core capabilities, i.e. characterization techniques
- <u>validate</u> claims, concepts, and theories of hydrogen-storage materials
- <u>double</u> hydrogen storage energy density (increase from 25g/L to 50 g/L)







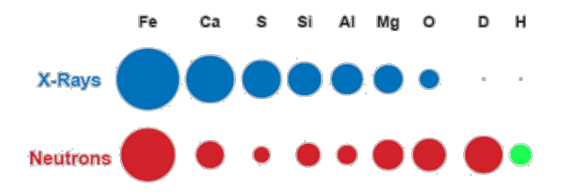




Relevance: Impact of Neutron Analyses



- Neutrons provide unique specificity towards determination of hydrogen properties
 - Enables identification of isotopically-labelled hydrogen location within complex structures
 - Enables identification of hydrogen dynamics within complex structures



http://www.ne.ncsu.edu/nrp/npdf.html













HyMARC

LBNL/NREL)

NIST

Pacific Northwest

Utilize neutrons to characterize and validate hydrogen storage media

NIST provides neutron-scattering-based characterization of materials of interest within HyMARC and other DOE-funded projects

- Solvent addition as a solution for enhancing hydrogen storage properties of magnesium borohydride (Mg(BH₄)₂)
- Quantum rotational tunneling of BH₄⁻ anions in lithium benzimidazolateborohydride Li₂(bIm)BH₄
- H₂ adsorption in a Cu(I) MOF

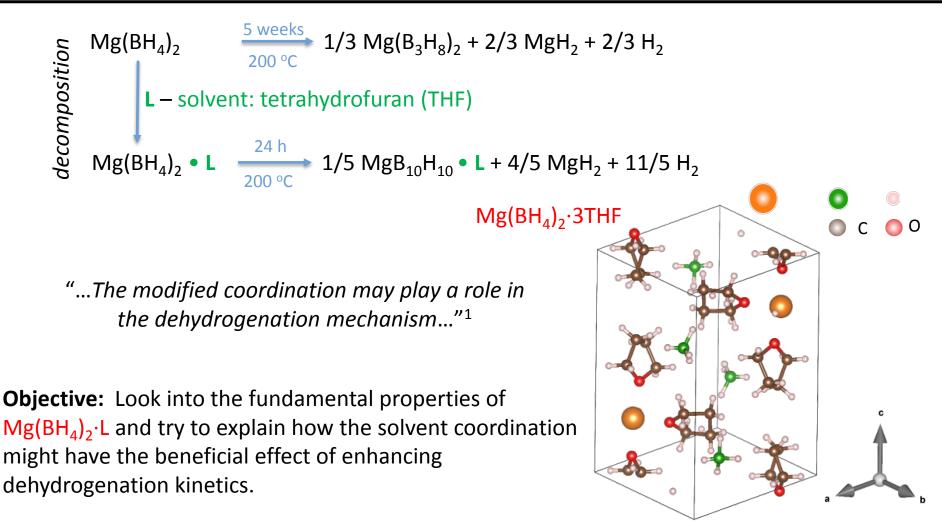
U.C. Berkeley/



Accomplishments and Progress:

Enhancing hydrogen storage properties of $Mg(BH_4)_2$





[1] M. Chong et al. *Inorganics* **2017**, *5*, 89.





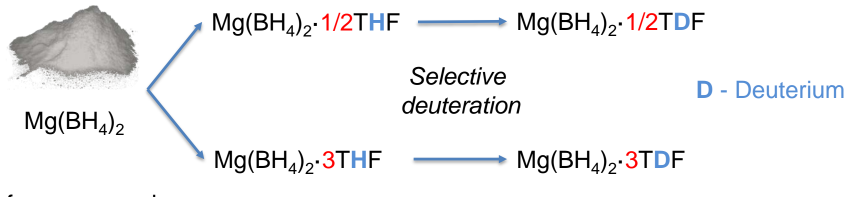




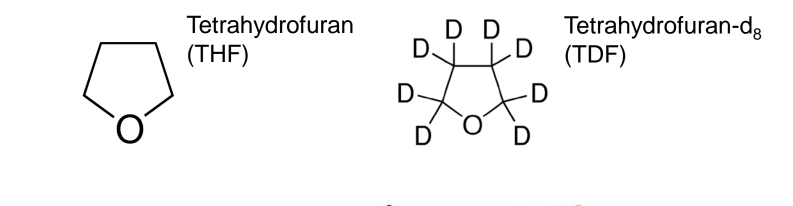
Accomplishments and Progress: Enhancing hydrogen storage properties of Mg(BH₄)₂



Samples: $Mg(BH_4)_2 \cdot L$ were prepared by adding an excess of solvent to $Mg(BH_4)_2$ at room temperature.



Reference samples:



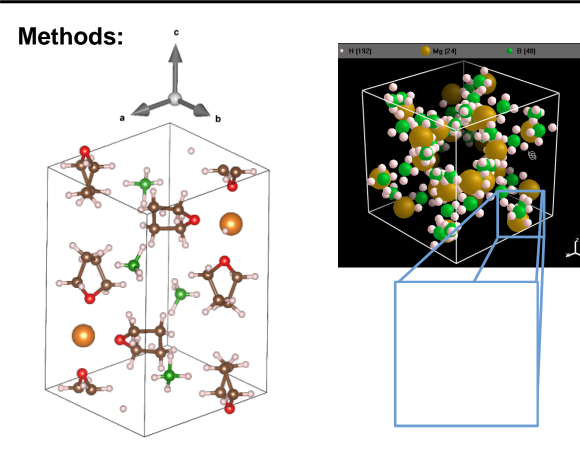
Pacific Northwest

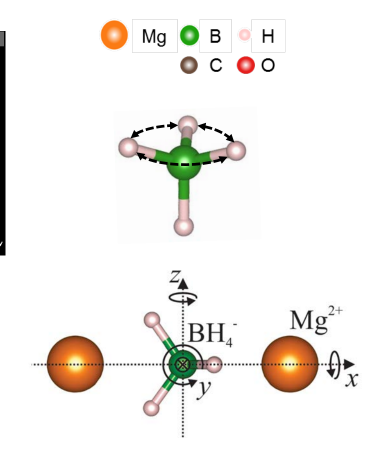




Accomplishments and Progress: Enhancing hydrogen storage properties of Mg(BH₄)₂







Neutron powder diffraction (NPD): structure and phases Neutron vibrational spectroscopy (NVS): vibrational modes Quasielastic neutron scattering (QENS): reorientational dynamics





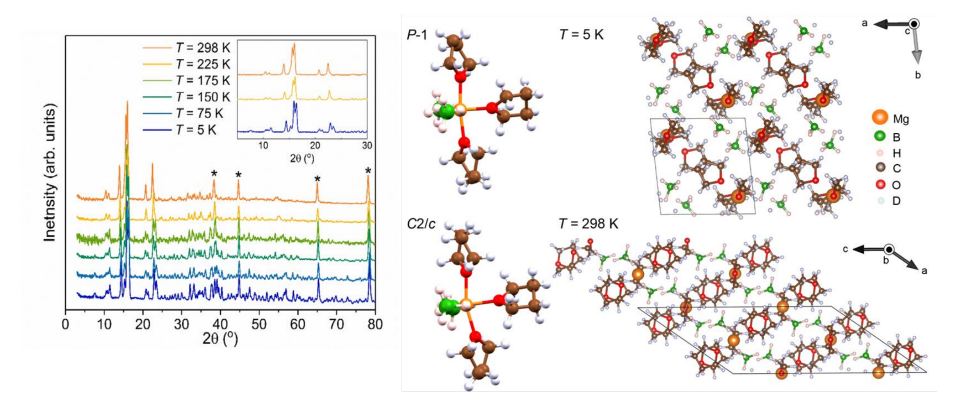


Accomplishments and Progress: Enhancing hydrogen storage properties of $Mg(BH_4)_2$

Sandia National

Laboratories

Neutron powder diffraction measurements of $Mg(BH_4)_2$ ·3TDF upon heating from 5 K to 298 K indicate a phase transition in the temperature range between 175 and 225 K, from triclinic (P-1) to monoclinic (C2/c) structure.



National Laborator



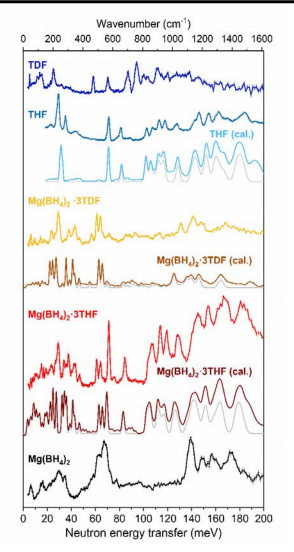
WMARC

Accomplishments and Progress: Enhancing hydrogen storage properties of Mg(BH₄)₂

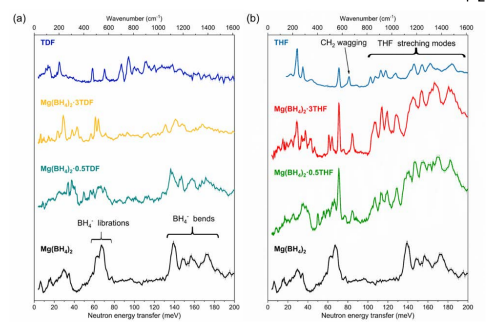
Lawrence Livermore

National Laboratory





- Vibrational properties are strongly influenced by the THF environment.
- In particular, there is a shift towards lower energies of the BH₄⁻ librational and bending modes with THF present as a result of changes in the bond lengths and force constants. Splitting in spectral features is also observed and explained by the lowering of the overall symmetry of Mg(BH₄)₂·3THF.

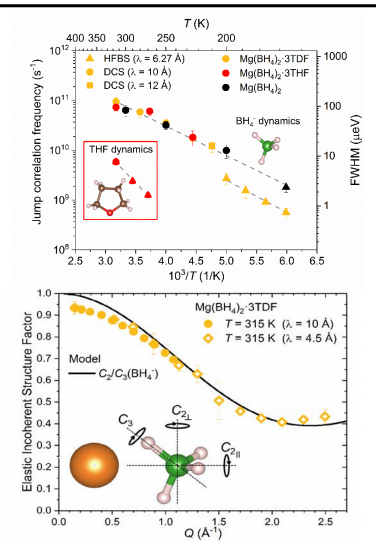


Pacific Northwest



Accomplishments and Progress: Enhancing hydrogen storage properties of $Mg(BH_4)_2$





Sandia

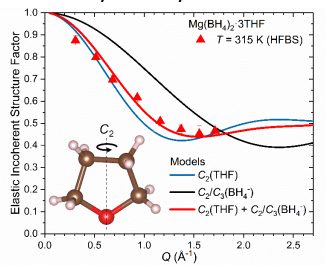
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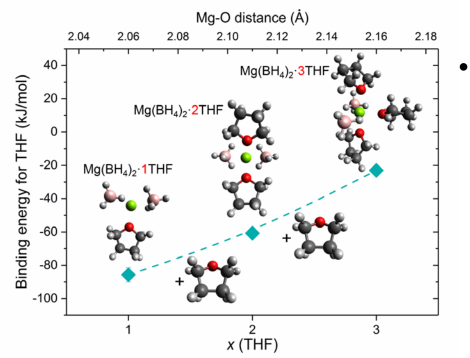
National Laboratory

- Orientational mobilities of the BH₄⁻ anions obtained from QENS are not particularly sensitive to the presence of THF or TDF and compare well with the mobilities of BH₄⁻ anions in unsolvated Mg(BH₄)₂
- the THF molecules in Mg(BH₄)₂·3THF are also found to be orientationally mobile, undergoing 180° reorientational jumps around their C₂ molecular symmetry axis.





Accomplishments and Progress: Enhancing hydrogen storage properties of Mg(BH₄)₂



Computational results for the solid state using the crystal structures suggest the static binding energy of each THF molecule averages to -27 kJ/mol THF. On the other hand, DFT calculations for the hypothetical $Mg(BH_4)_2 \cdot 1THF$ complex suggests the binding energy is greater for the first THF, then the subsequent $\cdot 2THF$ and $\cdot 3THF$ adducts.

From this combined experimental and computational study of THF adducts of Mg(BH₄)₂, we find little direct interaction between the THF and the BH₄⁻ anion. We propose that using fractions of THF to Mg(BH₄)₂ is beneficial in (i) preventing weakly bound THF from coming free from the Mg²⁺ cation and (ii) disrupts the stability of crystalline phase leading to a lower melting point and enhanced kinetics.



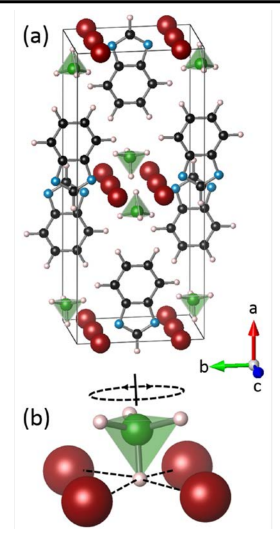






Accomplishments and Progress: Quantum BH₄⁻ Rotational Tunneling in Li₂(blm)BH₄





- Reorientational BH₄⁻ motion is known to contribute strongly to the balance of energies determining the thermodynamic stability of borohydrides.
- Therefore, information on the reorientational dynamics is important for understanding the fundamental properties of these compounds.
- The hydrogen dynamics in lithium benzimidazolateborohydride, Li₂(bIm)(BH₄) (bIm = C₇H₅N₂) was studied using various neutron scattering techniques and DFT calculations, in order to get better insights into the possible application of this material in hydrogen storage.







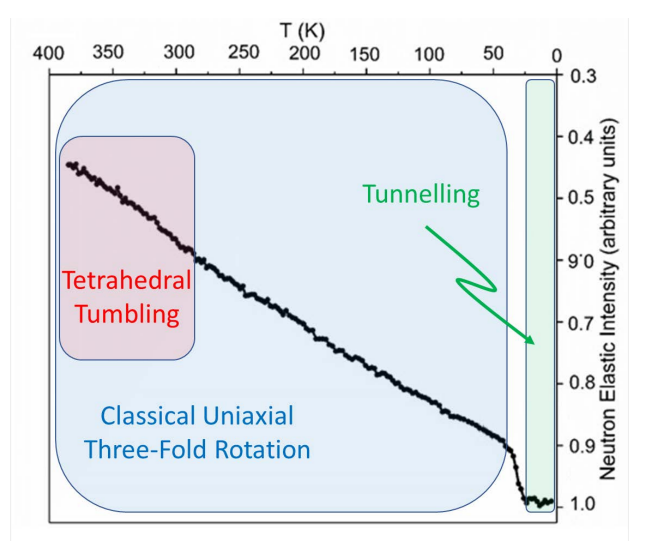




Accomplishments and Progress:

Quantum BH₄⁻ Rotational Tunneling in Li₂(blm)BH₄

Neutron-elastic-scattering fixed-window scan on the High-Flux Backscattering Spectrometer (HFBS) upon heating at 1 K min⁻¹ from 4 K to 385 K at 1.2 Å⁻¹ neutron momentum transfer.











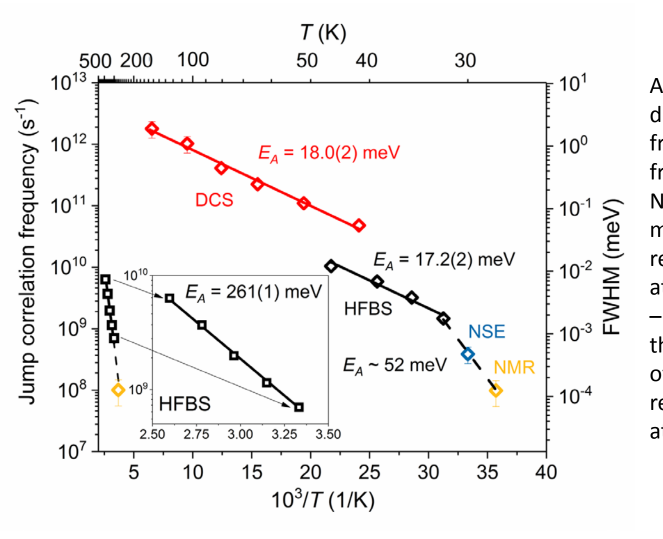




Accomplishments and Progress:

Quantum BH₄⁻ Rotational Tunneling in Li₂(blm)BH₄





Arrhenius-type plots of the derived jump correlation frequencies τ_1^{-1} versus 1/Tfrom the various QENS and NMR results for the much more rapid three-fold reorientations of the three H atoms associated with the -BH₃ fragment, as well as for the much slower exchange of these H atoms with the remaining, Li₄-anchored H atom.

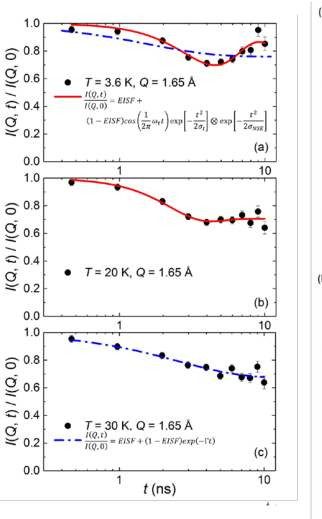


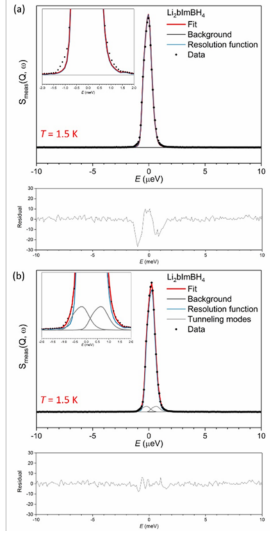






Accomplishments and Progress: Quantum BH_4^- Rotational Tunneling in Li₂(blm) BH_4





It is a rare observation of rotational tunneling of the BH₄anion via neutron scattering, and possibly the second neutron-spinecho (NSE) observation of tunnelling phenomena after an early work on dimethylacetylene.

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Both NMR and neutron scattering results for $Li_2(bIm)(BH_4)$ are described in terms of a gradual transition from the regime of lowtemperature quantum dynamics (rotational tunneling of BH₄⁻ anions) to the regime of classical three-fold uniaxial jump reorientations at higher temperatures.





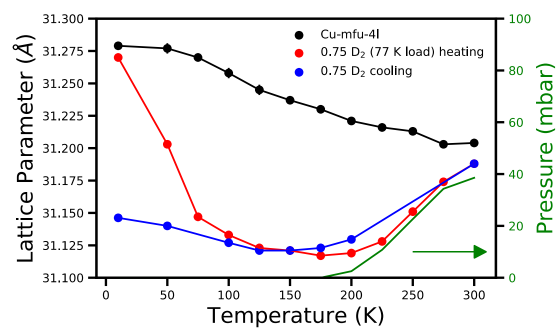






Accomplishments and Progress: H₂ adsorption in a Cu(I) MOF





Gas adsorption isotherms and Infrared spectra indicate a strong temperature dependence of H₂ adsorption characteristics in a Cu(I) MOF.

We performed variable-temperature diffraction to monitor how the H_2 is interacting with the framework.

• The bare framework shows a moderate negative thermal expansion (NTE).

encel

- 0.75 D₂:Cu dosed at 77 K and then cooled to base (prior to heating) shows a strong NTE.
- Recooling to base shows a large hysteresis away form the physisorbed system at low temperature. Rietveld refinement is consistent with two types of H₂, one being much closer to the Cu(I) site.





Summary



- Orientational mobilities of the BH₄⁻ anions in Mg(BH₄)₂·xTHF adducts are found not to be particularly sensitive to the presence of THF and compare well with the mobilities of BH₄⁻ anions in unsolvated Mg(BH₄)₂. The THF molecules in Mg(BH₄)₂·3THF are also found to be orientationally mobile, undergoing 180° reorientational jumps around their C₂ molecular symmetry axis.
- Both NMR and neutron scattering results for Li₂(bIm)(BH₄) are described in terms of a gradual transition from the regime of low-temperature quantum dynamics (rotational tunneling of BH₄⁻ anions) to the regime of classical three-fold uniaxial jump reorientations at higher temperatures. This uncommon behavior is due to the unique BH₄⁻ coordination in this compound.
- Gas adsorption isotherms and infra-red spectra of H₂ in a Cu(I) MOF indicate a strong temperature dependence of the H₂ adsorption behavior. Rietveld refinement analysis of complementary diffraction data is consistent with two types of H₂, one being much closer to the Cu(I) site.













- NREL/NIST collaboration
 - Characterizing ultra-microporous materials using neutron diffraction and neutron spectroscopy
- NREL/NIST collaboration with LBNL and PNNL
 - Characterizing hydrogen adsorption in metal organic framework materials using neutron diffraction and neutron spectroscopy
 - Characterizing various hydrogen storage materials at the Advanced Photon Source
 - Various neutron scattering characterization of $Mg(BH_4)_2 \cdot xTHF$











Project was not reviewed last year.











Proposed Future Work



- Perform QENS measurements on recently synthesized Mg(¹¹BH₄)₂·xTHF and Mg(¹¹BH₄)₂·xTDF materials to gain insights into the reorientation dynamics and mobilities of BH₄⁻ anions and provide more information on the interaction between THF and Mg(BH₄)₂.
- Continue neutron diffraction/NVS characterizations of new MOF materials

Any proposed future work will depend on the available funding.









Publications



- M. Dimitrievska, V. Stavila, A. V. Soloninin, R. V. Skoryunov, O. A. Babanova, H. Wu, W. Zhou, W. S. Tang, A. Faraone, J. D. Tarver, B. A. Trump, A. V. Skripov, and T. J. Udovic, The Nature of Decahydro-*Closo*-Decaborate Anion Reorientations in an Ordered Alkali-Metal Salt: Rb₂B₁₀H₁₀, J. Phys. Chem. C, 122, 15198-15207 (2018).
- M. Dimitrievska, J.-N. Chotard, R. Janot, A. Faraone, W. S. Tang, A. V. Skripov, and T. J. Udovic, Tracking the Progression of Anion Reorientational Behavior between α-phase and β-phase Alkali-Metal Silanides by Quasielastic Neutron Scattering, J. Phys. Chem. C, 122, 23985-23997 (2018).
- 3. M. Dimitrievska, P. Shea, K. E. Kweon, M. Bercx, J. B. Varley, W. S. Tang, A. V. Skripov, V. Stavila, T. J. Udovic, B. C. Wood, Carbon Incorporation and Anion Dynamics as Synergistic Drivers for Ultrafast Diffusion in Superionic LiCB₁₁H₁₂ and NaCB₁₁H₁₂, **Adv. Energy Mater.**, 1703422 (2018).
- 4. W. S. Tang, M. Dimitrievska, V. Stavila, W. Zhou, H. Wu, A. A. Talin, T. J. Udovic, Order–Disorder Transitions and Superionic Conductivity in the Sodium nido-Undeca(carba)borates, **Chem. Mater.**, 29, 10496–10509 (2017).
- 5. Asgari, M., Semino, R., Schouwink, P., Kochetygov, I., Trukhina, O., Tarver, J.D., Bulut, S., Yang, S., Brown, C.M., Ceriotti, M., Queen, W.L., An In-Situ Neutron Diffraction and DFT Study of Hydrogen Adsorption in a Sodalite-Type Metal–Organic Framework, Cu-BTTri, **Eur. J. Inorganic Chem.** 2019, 1147-1154 (2019).
- Kapelewski, M.T., Runčevski, T., Tarver, J.D., Jiang, H.Z.H., Hurst, K.E., Parilla, P.A., Ayala, A., Gennett, T., FitzGerald, S.A., Brown, C.M., Long, J.R., Record High Hydrogen Storage Capacity in the Metal–Organic Framework Ni₂(m-dobdc) at Near-Ambient Temperatures, Chem. Mater. 30, 8179-8189 (2018).
- 7. Strobel, T.A., Ramirez-Cuesta, A.J., Daemen, L.L., Bhadram, V.S., Jenkins, T.A., Brown, C.M., and Cheng, Y., Quantum Dynamics of H₂ Trapped within Organic Clathrate Cages, **Phys. Rev. Lett.** 120, 120402 (2018).









Presentations



- 1. M. Dimitrievska: "Neutron Scattering Studies of Hydrogenous Materials for Next-Generation Energy Storage", ACS National Meeting & Exposition, New Orleans, LA, Mar. 2018.
- 2. M. Dimitrievska: "Role of Solvent Adducts in Hydrogen Dynamics of Metal Borohydrides: Neutron-Scattering Characterization", ACS National Meeting & Exposition, New Orleans, LA, Mar. 2018.
- 3. M. Dimitrievska: "Carbon Incorporation and Anion Dynamics as Synergistic Drivers for Ultrafast Diffusion in Superionic LiCB₁₁H₁₂ and NaCB₁₁H₁₂", MRS Spring meeting, Phoenix, AR, Apr. 2018.
- 4. M. Dimitrievska: "HySCORE: Technical Activities at NIST", DOE-EERE-FCTO Annual Merit Review, Washington DC, Jun. 2018.
- 5. M. Dimitrievska: "Complex Borohydrides as Superionic Electrolytes", Review of the NIST Center for Neutron Research at the National Institute of Standards and Technology (NIST), Gaithersburg, MD, Jul. 2018.
- 6. M. Dimitrievska: "Neutron Backscattering Studies of Hydrogenous Materials for Next-Generation Energy Storage", National Science Foundation Site Visit Review of the Center for High Resolution Neutron Scattering (CHRNS), Gaithersburg, MD, Apr. 2018.
- 7. M. Dimitrievska: "Role of Solvent Adducts in Hydrogen Dynamics of Metal Borohydrides—Neutron-Scattering Characterization", American Conference on Neutron Scattering, Collage Park, MD, Jun. 2018.
- 8. C.M. Brown: "Shedding Light on Molecular Separations in Metal-Organic Frameworks through Neutron Scattering", International MOF Conference 2018, Auckland, New Zealand, Dec. 2018.
- 9. C.M. Brown: "Shedding Light on Industrial Separations in Metal-Organic Frameworks through Neutron Scattering", ACS Regional Meeting, Atlanta, GA, Nov. 2018.
- 10. C.M. Brown: "Neutron Scattering Studies of Small Molecules Adsorbed in Metal-Organic Frameworks", Materials Science & Technology Conference, Columbus, OH, Oct. 2018.
- 11. C.M. Brown: "Neutron Measurements of Hydrogen Storage Materials", ACS National Meeting, New Orleans, LA, Mar. 2018.
- A. Faraone: "Tunnelling H Motion in Lithium Benzimidazolate-Borohydride Li₂(blm)(BH₄): Neutron Spin Echo, Quasielastic Neutron Scattering and Nuclear Magnetic Resonance Results", 'Neutron Spin-Echo Spectroscopy 2018 – 40 Years of User Operation of NSE' Workshop, Grenoble, France, Oct. 2018.
- A. V. Skripov: "Low-Temperature Rotational tunneling of BH₄⁻ Groups in Lithium Benzimidazolate-Borohydride Li₂(bIm)(BH₄): Nuclear Magnetic Resonance and Neutron Scattering Studies", 16th International Symposium on Metal-Hydrogen Systems (MH2018), Guangzhou, China, Nov. 2018.





