





Synthesis of Nanophase Materials for Thermodynamically Tuned Reversible Hydrogen Storage

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Overview

Timeline

- Project start date: October 1, 2004
- Project end date: January 31, 2010
 Percent complete: 60%

Budget

- Total project funding DOE share \$1M (5 yrs) Contractor share \$250k (5 yrs)
- Funding for FY07 DOE share \$200k Contractor share \$50k
- Funding for FY08 DOE share \$200k Contractor share \$50k

Barriers

- (B) Weight and volume of on board hydrogen storage systems
- (M) Reversibility of high capacity solid state storage materials
- (N) Kinetics (fueling/refueling times) associated with current solid state storage materials

Partners

Interactions/ collaborations:

John Vajo, HRL Laboratories Ewa Rönnebro, Sandia Natl. Lab. David Sholl, Georgia Tech. Jason Graetz, Brookhaven Natl. Lab. J.C. Zhao, GE/Ohio State Univ. Terry Udovic, NIST R. C. Bowman, Jr. JPL



Objectives

- To understand if thermodynamically tractable reactions based on hydride destabilization, that should be reversible but appear not to be, are kinetically limited.
- To enable short hydrogenation times associated with refueling, that will require short solid-state and gas-solid diffusion path lengths.
- To address the problems associated with large, light-metal-hydride enthalpies (hydrogen fueling/refueling temperatures) and develop strategies to address thermodynamic issues surrounding the use of these materials through hydride destabilization. Systems of interest determined through "theoretical screening" by MHCoE partner members.
- To understand issues related to grain growth and surface/interface energies, which are vital in order to understand the kinetics of hydrogenation/dehydrogenation reactions.
- To follow up on previously studied reactions with phase identification via x-ray diffraction, NMR and transmission electron microscopy.

Technical accomplishments: The ScH₂ + 2LiBH₄ → ScB₂ + 2LiH + 5/2H₂ System: *Experimental Assessment of Chemical Destabilization*^a

- Destabilization reaction with ideal thermodynamic properties (DFT)
 - $-\Delta H_{300K} = 34.1 \text{kJ/mol}^b$
 - 8.91 wt% capacity
- Isothermal kinetic desorption measurments (450°C)
 - 4.5 wt% desorption
- Powder XRD detects only LiH and ScH₂ crystalline phases in desorption product, i.e. no reaction under these conditions between ScH₂ and LiBH₄.
- Stability of ScH₂, ScB₂ plays critical role in determining overall kinetics.

^aHydrogen Sorption Behavior of the ScH₂-LiBH₄ System: Experimental Assessment of Chemical Destabilization Effects, *Justin Purewal, Son-Jong Hwang, Robert C. Bowman, Jr., Ewa Rönnebro, Brent Fultz and Channing Ahn, accepted for publication in J. Phys. Chem. C (2008).*



^bCalculated by Center partner <u>David Sholl</u> and published as Alapati, S. V.; Johnson, J. K.; Sholl, D. S., J. Alloys Compd. 2007, 446–447, 23.

Technical accomplishments: The ScH₂+2LiBH₄ \rightarrow ScB₂ + 2LiH + 5/2H₂ System (continued): *MAS-NMR Spectroscopy*



(a) Bloch decay ¹¹B MAS NMR spectra with neat LiBH₄ (Sigma-Aldrich) added as a reference. No change of LiBH, peak observed after ball milling. ¹¹B MAS and CPMAS spectra (a and b) show formation of elemental boron in the amorphous phase (broad shoulder at ~ 5 ppm) and formation of intermediate phase (with peak at -12 ppm), recently identified^{*} as $[B_{12}H_{12}]^{2-}$ species. Note that a ¹¹B MAS NMR spectrum of ScB₂+2LiH system after absorption treatment at high H₂ pressure (896 bar, and 460°C for 48 hr) also included where the broader spinning sidebands in this spectrum are due to unreacted ScB₂. The small peak at -41 ppm, indicates very limited LiBH₄ formation (~ 3%) in the reaction product. So reverse reaction is seen but under technologically challenging conditions.

(b) ¹¹B MAS and CPMAS NMR spectra (contact time=0.1 ms) of desorbed sample.

*NMR Confirmation for Formation of $[B_{12}H_{12}]^{2-}$ Complexes during Hydrogen Desorption from Metal Borohydrides, Son-Jong Hwang, Robert C. Bowman, Jr., Joseph W. Reiter, Job Rijssenbeek, Grigorii L. Soloveichik, Ji-Cheng Zhao, Houria Kabbour, and Channing C. Ahn, J. Phys Chem. C, 112, 3164-3169, 2008.

Technical accomplishments: The ScH₂+2LiBH₄ \rightarrow ScB₂ + 2LiH + 5/2H₂ System (continued): *MAS-NMR Spectroscopy*



) Un-reacted ScH₂ still present in desorption product, ScB₂/other Sc compounds not detectable in ⁴⁵Sc MAS NMR spectra presented in c). Note that ⁴⁵Sc MAS NMR signal significantly reduced after the desorption reaction, tentatively attributed to inhomogeneous separation of ScH₂ and LiBH₄ phases during the desorption reaction and correspondingly nonuniform distribution of ScH₂ particles in resulting NMR sample.

d) ⁶Li MAS NMR spectra also show distinctive signature of phase changes from LiBH₄ (-1.1 ppm) to Li₂B₁₂H₁₂ (-0.5 ppm). Presence of LiH confirmed by independent ⁶Li CPMAS NMR measurements (not shown) because extremely long spin lattice relaxation behavior inhibits observation during recording of MAS spectra. As with XRD data, NMR results identify decomposition products of LiBH₄, but show no evidence of ScH₂ reacting to form the ScB₂ product destabilization phase.

Technical accomplishments: The ScH₂+2LiBH₄ \rightarrow ScB₂ + 2LiH + 5/2H₂ System (continued): *Raman Spectroscopy*





Top: Optical micrograph of a $ScH_2+2LiBH_4$ mixture after desorption at 400°C for 20 hours. Desorption product segregates into two phases that are distinguished by their white and dark colors, respectively. The dark phase and the as-milled mixture not Raman active as determined by our spatially resolved analysis.

Bottom: White phase (trace a) shows similarities to the spectrum of dehydrogenated LiBH_a (trace b). The B-H bending and stretching modes of LiBH₄ around 1300 and 2300 cm⁻¹ present in both samples, and not affected by decomposition at temperatures below 400°C. The broad peak at 1100 cm⁻¹ corresponds to Raman modes from B-B bonding in amorphous boron. The additional B-H bending and stretching modes around 500-1000 cm⁻¹ and 2500 cm⁻¹ consistent with calculated and measured B-H vibrational modes in $Li_2B_{12}H_{12}$. These results suggest that the ball-milled ScH₂+2LiBH₄ mixture segregates back into its initial components upon melting and solidifying of LiBH₄. The dark phase contains ScH_2 while the white phase contains LiBH₄ and its thermal decomposition products. The solid triangles, open circles, and open triangles identify modes from Li₂B₁₂H₁₂, LiBH₄, and amorphous boron respectively. 7

LiBH₄ reversibility in Ca(AIH₄)₂ - 2LiBH₄



Proton, Li, B, Al NMR analysis of as milled, desorbed and rehydrogenated samples.

Initial work found partial reversibility in LiBH₄ when alanate made via mechano-synthesis* (~70% yield). ~88% LiBH₄ reversibility in Ca(AIH₄)₂-2LiBH₄ system although no evidence of CaB₂ formation. H, Li, and B NMR show that LiBH₄ can be rehydrogenated although the data for Al indicate that the Ca(AIH₄)₂ compound has undergone dissociation and the Al in this phase has transformed to the elemental state.

LiBH₄ reversibility in Ca(AIH₄)₂(pure)- 2LiBH₄ cont'd

AlH₃ (reference)

Solution synthesis $Ca(AIH_4)_2$



With ¹¹B MAS NMR, reversible formation of LiBH₄ is verified for 3 destabilization reactions containing aluminum hydrides.

Formation of $Li_2B_{12}H_{12}$ was also observed for all three systems. This is a competing reaction which limits reversibility of $LiBH_4$ formation.

LiBH₄ reversibility in AIH₃ - 2LiBH₄ cont'd



(**above**) Kinetic H₂ desorption data for a final temperature of 400°C. (**right**) ¹H MAS NMR revealing the H content in the direct synthesis of $AIH_3 + 2LiBH_4$ system.



- LiBH₄ reversibly desorbs/reabsorbs H_2 when mixed with
 - Ca(AIH₄)₂ direct ball-milling synthesis
 - $Ca(AIH_4)_2$ synthesized in solution (pure)
 - AIH₃ (Dow Chemical)
- Catalytic role of elemental AI is key in reversible formation of LiBH₄

LiBH₄ reversibility in Ca(AIH₄)₂ - 2LiBH₄: XRD Qualitative Phase Analysis



Formation of elemental aluminum is observed in all cases. Appears to have catalytic effect in reversible formation of LiBH₄. (**Right**) Formation of CaH₂ occurs as a side-reaction with Ca(AlH₄)₂. (**Left**) Peaks marked ??? have previously been identified as AlB₂ in the literature, but they don't match our reference pattern for AlB₂.

No data available yet for direct ball-mill synthesized Ca(AIH₄)₂

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Solution synthesis

LiBH₄ reversibility in Ca(AIH₄)₂ - 6LiBH₄



Proton, Li, B, Al NMR analysis of as milled, desorbed and rehydrogenated samples.

H desorbed and adsorbed in LiBH₄ in this system. ~40% LiBH₄ reversibility measured, less than in 2LiBH₄ case. No NMR evidence of CaB₆ formation. Ca(AIH₄)₂ decomposes to AI metal. Formation of 981 and 782 ppm peaks in ²⁷AI NMR spectra of desorbed material observed for first time, but identities unknown.

Technical accomplishments: Reaction pathways in borohydride systems

Direct thermal decomposition or destabilization system reaction pathways not easy to predetermine.

An important class of metal borohydride systems unexpectedly observed to produce a $(B_{12}H_{12})^{2-}$ intermediate phase as determined by NMR.

In collaboration with JPL and GE, LiBH₄, Mg(BH₄)₂, LiSc(BH₄)₄, Ca(AIH₄)₂-LiBH₄, studied by ¹¹B magic angle spinning and cross-polarization magic angle spinning as described in table below.

TABLE 1: Reactions Examined^a

reaction	reactants	pretreatment	T _{desorb} /P _{final} [in total vessel volume]	H ₂ desorbed (wt % of total sample)
1	LiBH ₄ (Sigma Aldrich)	as received	400, 500 °C, 620 Torr [1.1 liters] or vacuum	varied
2	$Mg(BH_4)_2$	after second desorption	450 °C, ramp heating in vacuum	11.5
3	SeCl ₃ +3LiBH ₄	ball milled	450 °C, ramp heating, 10351 Torr [24.4 cm ³]	7.7
4	Ca(A1H ₄) ₂ +6LiBH ₄	ball milled	350 C, 280 Torr [1.1 liters]	8.5

^a Unless noted, pressures are final values, after desorbing into initially evacuated volumes. Quantity of desorbed H₂ is given as weight percent of original sample weight.

ppm

-20

0

20

40



¹¹B NMR spectra at left, of powders collected after hydrogen desorption reactions of LiBH₄ (see Table 1). (a) LiBH₄ (as received, Sigma-Aldrich, note the scale adjusted by 1/2), (b) desorbed at 400° C, (c) desorbed at 500°C, (d) desorbed at 500°C under vacuum, (e) elemental boron in amorphous phase (Sigma-Aldrich), (g) ¹¹B MQMAS spectrum of sample d. Spinning side bands are marked with a *. The dashed line in 2D MQMAS spectra is the chemical shift axis.

Technical accomplishments: (cont'd) *Reaction pathways in borohydride systems*



At left, (a) ¹¹B CPMAS variable contact experiment of powders collected after hydrogen desorption reactions of LiBH₄ and MgBH₄, and pure K₂B₁₂H₁₂. The broken line shows ¹H relaxation during the spin-lock period. (b) ¹¹B CPMAS NMR spectra of desorbed materials (see Table 1). The contact time of 0.05 ms was used. ¹¹B CPMAS spectra of elemental amorphous boron (a-B, note the scale change (1/100)), K₂B₁₀H₁₀, and K₂B₁₂H₁₂ reference compounds are added for comparison.

At left ¹¹B NMR spectra in solution phase after powders from desorption reactions were dissolved in H₂O. (c) ¹H decoupled, (d) ¹H coupled. Spectra from aqueous solutions of the K₂B₁₂H₁₂ compounds are compared to confirm the presence of $[B_{12}H_{12}]^{-2}$ unit in the dissolved LiBH₄ and Mg(BH₄)₂ decomposition products. J(B-H) was measured to be 125 Hz.

Technical accomplishments: Transmission electron microscopy and analysis of Mg in carbon aerogel

As part of an effort in looking into the possible effects of size on the thermodynamic properties of hydrogenation/ dehydrogenation, and in conjunction with HRL Laboratories, we have been looking at elemental distributions of Mg in carbon aerogels. Because of the lack of "wettability" of Mg on typical carbon surfaces (seen at right), schemes that promote the incorporation of Mg, especially into the small pore dimensions of aerogels needs to be verified.



Sample 50559-60-1-c above showing aerogel clusters in bright field image at left. Mg image on right formed by using Mg L23 edge. The "white specular" regions are results of artifacts from difficulties with background subtraction. Region in upper right of these images shown in images at right (and rotated 90° anticlockwise). Mg L23 image at 1/2 scale in upper right. Carbon K image at 1/2 scale in lower right.



Image at left shows attempt to melt Mg in carbon crucible and shows no evidence of wetting. Original ingot morphology was cylindrical and turned spheroidal with poor contact angle. Image at right shows similar attempt when Ni film of several hundred Å is deposited into crucible before melt attempt. Some Mg coating of surface evident, presumably from Mg vapor phase.



Technical accomplishments: Transmission electron microscopy and analysis of Mg in carbon aerogel cont'd

The typical microstructure can be dominated by regions of Mg metal or hydride that are decoupled from the aerogel regions as shown in the bright field image below,



Characteristic of Mg particles in the aerogel microstructure are particles with smooth surfaces. To the upper right is high angle annular dark field image of the same region (where the bright regions are Rutherford scattered electrons with intensity that goes as Z^2) as above but where the image is inverted horizontally. While Mg₂Ni is commonly observed within the aerogel, the overall composition of typical aerogel regions is greater than this stoichiometry, indicating that excess Mg can be incorporated into the aerogel microstructure. While most of the Mg is found to go into larger pores, evidence for high Mg concentrations on the aerogel structure in nano phase form can be discerned.



Spectrum Spectrum 1 Spectrum 2 Spectrum 3 Spectrum 4 Spectrum 5 Spectrum 6 Spectrum 7 Spectrum 8	In stats. Yes Yes Yes Yes Yes Yes Yes Yes	Mg 70.89 104.73 99.94 100.30 48.33 118.87 52.81 94.51	Ni 29.11 -4.73 0.06 -0.30 51.67 -18.87 47.19 5.49
Spectrum 9	Yes	94.51 88.85	5.49 11.15

FY 2008-09 plans

- Reaction path investigation in AIH₃: LiBH₄ system to understand reversibility mechanism.
- Wetting and aerogel pore size enthalpy dependent studies.
- Continued NMR studies to identify reaction species that lack definitive crystallographic features.
- Enthalpy studies in micropore sized aerogels.
 - Pressure infusion approach to force metal/hydride into non-wetting pores.
- Coordination with T. Baumann (LLNL) in synthesis of aerogels with pore size range down to 1 to 2 nm. Surface/interface effects will play a dominant role in reaction enthalpy behavior at these size dimensions.
- Use of surface treatments/additives to enhance wetting of hydrides in aerogel pores.
- Confirmation of lowered enthalpies and diffusion paths in hydrides encapsulated within aerogel pores.