### ENVIRONMENTAL REACTIVITY OF SOLID STATE HYDRIDE MATERIALS

### Joshua R. Gray and Donald L. Anton<sup>\*</sup> Savannah River National Laboratory, Aiken, SC 29803, USA

#### ABSTRACT

In searching for high gravimetric and volumetric density hydrogen storage systems, it is inevitable that higher energy density materials will be used. In order to make safe and commercially acceptable condensed phase hydrogen storage systems, it is important to understand quantitatively the hazards involved in using and handling these materials and to develop appropriate mitigation strategies to handle potential material exposure events. A crucial aspect of the development of risk identification and mitigation strategies is the development of rigorous environmental reactivity testing standards and procedures. This will allow for the identification of potential hazards and implementation of risk mitigation strategies. Modified testing procedures for shipping air and/or water sensitive materials, as codified by the United Nations, have been used to evaluate two potential hydrogen storage materials, 2LiBH<sub>4</sub>·MgH<sub>2</sub> and NH<sub>3</sub>BH<sub>3</sub>. The modified U.N. procedures include identification of self-reactive substances, pyrophoric substances, and gas-emitting substances with water contact. The results of these tests for air and water contact sensitivity will be compared to the pure material components where appropriate (e.g.  $LiBH_4$  and  $MgH_2$ ). The water contact tests are divided into two scenarios dependent on the hydride to water mole ratio and heat transport characteristics. Air contact tests were run to determine whether a substance will spontaneously react with air in a packed or dispersed form. Relative to  $2LiBH_4 \cdot MgH_2$ , the chemical hydride NH<sub>3</sub>BH<sub>3</sub> was observed to be less environmentally reactive.

### **1.0 INTRODUCTION**

Hydrogen is a viable candidate as an energy storage medium due to its inherent cleanliness upon oxidation and its ready utilization in fuel cell applications. In searching for ever higher gravimetric and volumetric density hydrogen storage materials and systems, it is inevitable that higher energy materials will be studied and used. To make safe and commercially acceptable systems, it is important to understand, quantitatively, the hazards involved in using and handling these materials and to develop appropriate safety systems to handle unforeseen accidental events.

The majority of research programs currently underway deal with the development of new solidstate hydrogen storage materials and their reversibility. There have been few programs concentrating on development of systems to utilize these materials and fewer programs still comprehensively dealing with the inherent safety considerations inherent to any system built using these materials. Adherence to international standards for safety will ultimately be required of any energy storage system which commercially exploits these materials. Many of the materials currently being considered as solid state hydrogen storage media are sensitive to air, water and water vapor [1,2]. In the spirit of system engineering, it is particularly enlightening to determine the relative environmental reactivity of the various candidate compounds under consideration. By understanding the nature of the environmental reactivity of these materials, amelioration and risk mitigation methods can be devised which will reduce the hazard associated with any inadvertent exposure to acceptable levels. Through development of global solid-state hydrogen storage safety programs, it is anticipated that international standards for testing and risk mitigation can be developed and implemented.

The current work is part of the IPHE collaboration (International Partnership for a Hydrogen Economy) that seeks to understand the physical hazards involved in synthesis, handling and utilization of these materials as hydrogen storage media as well as development of methods to mitigate these hazards which would result in commercially acceptable high density hydrogen storage system designs. A team of laboratories expert in hydrogen storage: Savannah River National Laboratory, Sandia National Laboratory & United Technologies Corp. (USA), Forschungszentrum Karlsruhe (Germany), National Institute for Advanced Industrial Science and Technology (Japan) and Université du Québec à Trois-Rivières (Canada), brings a broad spectrum of expertise in solid sate hydrogen storage focused to fundamentally understand these phenomena. This international collaboration will hasten the development and acceptance of these systems.

The objective of the presented study is to fundamentally understand the safety issues associated with solid state hydrogen storage materials through development of standard testing techniques to quantitatively evaluate both materials and systems. The results of these tests will be used to understand the environmental reactivity of hydrides and physisorption and to develop amelioration methods and systems to mitigate the hazards and handling and use to acceptable levels.

### 2.0 EXPERIMENTAL

### 2.1 Standardized Materials Testing

A set of materials testing procedures was developed based on internationally accepted standards drawn from United Nations [3] testing procedures. These tests include exposure to air, humidity, and water. These tests are described in the following sections. Generally, the

U.N. test procedures were modified by the addition of thermocouples to monitor the temperature within the hydrogen storage material, or in near proximity to the material, as appropriate. All tests were video recorded.

## 2.2 Water Reactivity Tests

The purpose of the water reactivity test is to verify if the substance, when in contact with water, burns or emits flammable gases. The experimental details followed UN-RTDG part-3, test-N5 in which three separate tests were conducted. (i) The test substance was formed into a small pile approximately 20 mm high and 30 mm diameter with a hollow in the top. A few drops of water are added to the hollow. If spontaneous ignition occurred at any stage, the substance is classified as a water reactive substance emitting flammable gases. No further testing is necessary. (*Water Drop Test*) (ii) A small quantity (approximately 2 mm diameter) of the test substance was dropped in a trough of distilled water at 20°C. (*Water Immersion Test*) (iii) A small quantity (approximately 2 mm diameter) of the test substance was placed on the center of a filter paper which is floated flat on the surface of distilled water at 20°C in a suitable vessel (i.e. a 250 ml beaker). The filter paper is to keep the substance in one place, under which condition the likelihood of spontaneous ignition of any gas is greatest. (*Surface Contact Test*)

## 2.3 Burning Rate Test

The purposes of the burn rate test are the classification of rapidly combustible solids, differentiation of ignitable, rapidly burning and dangerous burning substances and assessment of the relative hazard of rapidly combustible solids. The test procedure details followed UN-RTDG part-3, test-N1 [3]. The powdery hydrogen storage material was deposited as a strip on a platform to measure the burning rate. The powder strip with 250 mm of length and a 100 mm<sup>2</sup> cross-section was ignited from one end and burning propagation time measured for 100 mm after an initial stabilization period. A series of 6 thermocouples was fitted along the axis of burn rate propagation at regular intervals, so that the temperature at a point ~5 mm into the pile could be monitored as a function of time. Additionally, these tests were video recorded to provide qualitative data acquisition, as well as a second manner to reproducibly calculate the burn rate. The substance is classified as Packing Group II if burning time less than 45 seconds.

# 2.4 Spontaneous Combustion Test

The purpose of the spontaneous combustion test is to classify spontaneously combustible materials into two types: (i) Pyrophoric Substances, solid or liquid mixtures which, even in small quantities, ignite within 5 minutes of coming in contact with air, and (ii) Self-heating substances, substances which, in contact with air and without an energy supply, are liable to self-heating. These substances will ignite only when in large quantities (i.e. kilogram), and after a long period of time (i.e. hours or days).

The details of the test procedure followed UN-RTDG div. 4.2 [3]. The powder samples were loaded in 25x25x25 mm cubic baskets made of stainless steel screen with 0.05 mm openings and an uncovered top surface. 3 chromel-alumel thermocouples, with 0.3 mm diameter were inserted into the cubic sample container at positions in the center, face-center, and corner of the powdery sample to monitor temperature. The basket was housed in a cubic container cover made from a stainless steel net with a mesh opening of 0.60 mm, and slightly larger than the sample container. The cube is set in a hot air-circulating oven nominally at 150°C for at least 24 hours or until spontaneous ignition or hazardous self-heating was observed. It was experimentally observed that there was a 10°C overshoot of the measured internal oven temperature at setpoint and the nominal

setpoint temperature. The changes of the temperature at the chosen locations of the cube were recorded for the duration of the test.

# 2.5 Pyrophoricity Test

The purpose of the pyrophoricity test is to verify the ability of a solid to ignite on contact with air and determine the time to ignition. The test procedure followed UN-RTDG part-3, test-N2 [3]. An aliquot of  $1\sim2$  ml of the sample was poured from approximately a one meter height onto a non-combustible surface. Observation was made as to whether the substance ignited during dropping or within 5 minutes of settling. This procedure was performed six times or until a positive result was obtained. The substance was classified as pyrophoric if ignition occurred during one of the free-dropping tests.

# 2.6 Materials

Two hydrogen storage materials were tested for the work reported.  $2\text{LiBH}_4$ ·MgH<sub>2</sub> was selected as a material representative of the metal hydride class of storage materials [4], and NH<sub>3</sub>BH<sub>3</sub> was selected as a representative of the chemical hydride class of storage materials [5]. The raw materials were purchased from commercial vendors and ballmilled to give a uniform starting condition for the tests. Materials were typically SPEX milled in 3g amounts. All materials were handled in an inert atmosphere Ar environment glove box, and exposure to air when in transfer to the testing equipment was minimized as much as possible. Powder X-Ray Diffraction was performed on the materials after ballmilling, which confirmed only the presence of the starting materials. In order to discharge the materials, 3-10g amounts of material were loaded into a pressure manifold and heated to a temperature higher than the literature reported discharge temperature for the material of interest [4,5].

# **3.0 RESULTS AND DISCUSSION**

The results of the air and water reactivity tests are presented below with the water reactivity tests presented first followed by the results of the air reactivity tests. The materials are also subject to rigorous chemical thermodynamic and kinetic characterizations using sophisticated calorimetric techniques. The results of the standardized tests and the chemical characterization are being used to support a numerical simulation effort to increase understanding of the chemistry and mechanisms involved in the material reactivity.

# 3.1 2LiBH<sub>4</sub>·MgH<sub>2</sub>

The results of the water reactivity tests for  $2\text{LiBH}_4$ ·MgH<sub>2</sub> in the fully charged state are shown below in Fig. 1. Fig. 1(a) depicts the results from the Water Drop Test, where it can be seen that the material is highly reactive with the addition of a few drops of H<sub>2</sub>O to a relatively large amount of the hydride material. In Fig. 1(b), the results of the Water Immersion Test are shown. The material tends to form a film on the surface of the water, with very small spark-like events noted in some cases. At longer times, gas is evolved from the materials in the film. In Fig. 1(c), the results of the Surface Contact Test are shown, where it can be seen that the material is immediately reactive with the H<sub>2</sub>O soaked filter paper, which is in constant contact with the water bath in the beaker.

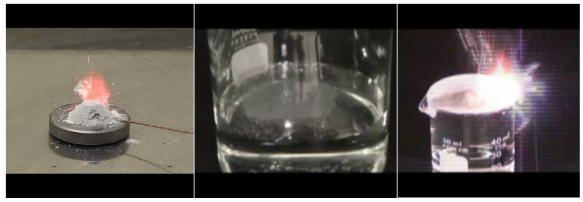


Figure 1. Results from the Water Contact Test for 2LiBH<sub>4</sub>·MgH<sub>2</sub>. a) Water Drop Test. b) Water Immersion Test, and c) Surface Contact Test.

The difference in the observed water reactivity of  $2\text{LiBH}_4 \cdot \text{MgH}_2$  is hypothesized to be the result of the different heat dissipation rates in the three scenarios and the rate of hydrogen generation. In the Water Drop Test, hydrogen gas is rapidly evolved via the hydrolysis of the constituent LiBH<sub>4</sub> and MgH<sub>2</sub> materials; additionally, elemental Mg and Li may also be present in these samples after the ball milling process. These hydrolysis reactions are well known to be highly exothermic [2], and therefore the possibility of reaching the ignition temperature of the H<sub>2</sub> is possible. In the Water Drop Test, there is no appreciable heat sink to reduce the temperature of the sample and surrounding air. Comparing this to the Water Immersion Test, hydrogen gas is clearly evolved via hydrolysis, as in the Water Drop Test. However, with the presence of the relatively large amount of water in intimate contact with the material, the heat is more quickly wicked away thus preventing a reactive event. The Surface Contact Test is a hybrid between the two scenarios, where the mass transport of the water to the hydride for reaction is impeded by the presence of the filter paper, as is the transfer of the heat to the water which is restricted to only the bottom surface of the pile through the filter. It is obvious that this material is sufficiently reactive to build up enough heat energy and H<sub>2</sub> to give rise to an ignition event.

The results of the Self-Heating Test for  $2\text{LiBH}_4$ ·MgH<sub>2</sub> are shown below in Fig. 2. Data were collected at 5 minute intervals during this experiment, therefore the number of data points is somewhat sparse over the time interval which the experiment occurred. On the right of Fig. 2 (Fig. 2.b), is a background control experiment that was performed on a SiO<sub>2</sub> sample which was loaded into the sample container and placed into the oven. As can be seen, the temperature measured at a point in the interior of the oven, but exterior to the sample, shows the characteristic 10°C overshoot that is typical of the current setup. The other thermocouples (center, face-center, and corner) all follow the same temperature trajectory. They heat up initially until they reach a steady-state of approximately 150°C after which the readings remain constant until the heat source from the oven is turned off. The thermocouple reading gradually fall back to room temperature as the oven cools. In contrast, inspection of Fig. 2.a shows all 4 thermocouples at room temperature at t=0 s. After 5 min, the highest reading is experienced at the face-center location (T = ~400°C), with the 2<sup>nd</sup> highest temperature at the corner (T = ~320°C), and the temperature at the center of the sample relatively low at T = ~240°C).

It should be noted that the criterion for failure of the self-heating test is for the sample to heat above the ambient temperature by 60°C or more; thus the sample has already experienced dangerous self-heating at this point in the experiment. After the passage of another 5 minutes, the temperature of the center is now the highest measurement ( $T = \sim 410^{\circ}C$ ), followed by the

temperature of the corner (T =  $\sim$ 340°C), followed by the temperature of the face-center (T =  $\sim$ 290°C). As time proceeds, the center increases until a maximum reading of  $\sim$ 450°C, after which the sample cools until it reaches a steady state of approximately 150°C, consistent with the background temperature from the SiO<sub>2</sub> sample. These results are interpreted as the procession of a reaction front, which starts at the face-center couple first, then proceeding towards the slightly more interior corner couple, and finally to the central couple. The relative insulation surrounding the central couple, provided by the reacted material, causes the couple to retain more heat and reach higher temperatures. These results are consistent with temperature enhanced hydrolysis and oxidation of the hydride materials resulting from the contact with the ambient air in the oven at the elevated temperature of the test.

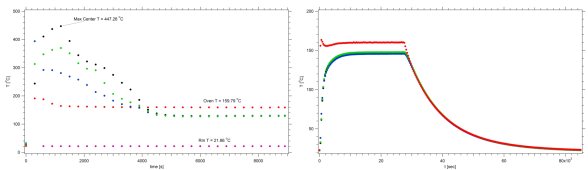


Figure 2. Thermocouple readings for the Self-Heating Test performed on  $2LiBH_4 \cdot MgH_2$ . a) Active  $2LiBH_4 \cdot MgH_2$  material. b) SiO<sub>2</sub> material used for background reference. (Red = Oven Temperature, Black = Sample Center Temperature, Blue = Sample Side Temperature, Green = Sample Corner Temperature)

Shown below in Fig. 3 are the results of the Burn Rate Test for the  $2\text{LiBH}_4$ ·MgH<sub>2</sub> material. In Fig. 3.a, a photograph is shown which was taken ~ 5 s into the burn rate test, after which time the flame had propagated the entire length of the sample. It is obvious that the material is combustible by the metrics employed in this test. On the right in Fig. 3.b., the results of the thermocouple measurements are shown. In the upper right of the figure is an inset showing a blown-up depiction of the initiation period from 750s – 800s. The burn rate that is reported uses an average of the time optically measured using the video recording data, and the time measured using the thermocouple results. The calculated burn rate for this experiment is 52 mm/s. This value is very similar to the value previously measured for NaAlH<sub>4</sub> by Mosher *et al* [6] of 51 mm/s. The lack of symmetry in the thermocouple measurements between the couples which are reflectively displaced from the midpoint of the sample length axis is attributed to spatial non-uniformity of the sample during the test.

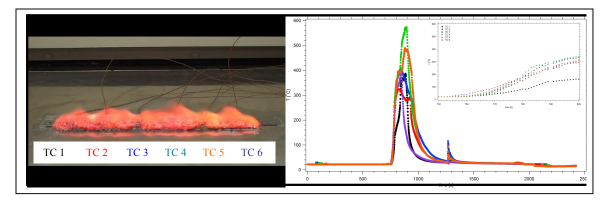


Figure 3. Results of the Burn Rate Test for the 2LiBH<sub>4</sub>·MgH<sub>2</sub> material. a) Photograph of Burning Material after 5 seconds, b) Thermocouple Readings

The  $2\text{LiBH}_4$ ·MgH<sub>2</sub> was also tested using the procedure for the U.N. Pyrophoricity classification. Approximately 1g of the material was dropped into a box with glass sides with the bottom made from industrial grade aluminum sheeting. The test nominally stipulates a 5 min waiting period. No reaction of the material in the air during the material drop, or within 15 minutes after the material was allowed to remain on the ground, was observed. Therefore, the material is deemed to non-pyrophoric by the metrics used in this test.

# **3.2 NH<sub>3</sub>BH<sub>3</sub>**

The results of the Water Reactivity tests for  $NH_3BH_3$  are shown below in Fig. 5. The results are given in the same order as the previous case; the Water Drop (a), Water Immersion (b), and the Surface Contact (c), respectively. As can be seen from the representative photographs, no discernable reactivity was observed using the metrics associated with the modified U.N. protocols. Whereas the  $2LiBH_4$ ·MgH<sub>2</sub> material was observed to be reactive in all three tests, with significant reactivity observed in the Water Drop and Surface Contact tests, in the case of ammonia borane there is virtually no reactivity. This is consistent with the results of the calorimetric tests that have been performed on this material, as well as other literature on  $NH_3BH_3$  [7,8]. The results would indicate that water contact may not be a significant risk mitigation factor for the  $NH_3BH_3$  materials in the same manner that is might be for a metal hydride based on hydride and borohydride chemical structures.



Figure 5. Results from the Water Reactivity Tests for NH<sub>3</sub>BH<sub>3</sub>. a) Water Drop Test. b) Water Immersion Test, and c) Surface Contact Test.

The results of the Self-Heating Test for the NH<sub>3</sub>BH<sub>3</sub> material are shown below in Fig. 6. In this case, data was collected with greater frequency than that for the 2LiBH<sub>4</sub>·MgH<sub>2</sub> material, so that the temperature ramp of the oven is captured by the ambient thermocouple readings. As can be seen in Fig. 6.a, the set point was hit within about 5 or 6 minutes after the experiment began. After soaking at the set point for approximately 5 minutes (for a total of 11 minutes exposure), the NH<sub>3</sub>BH<sub>3</sub> undergoes a strong exothermic event and green flame was observed within the oven (photographic data not available). The ambient thermocouple, which is located just exterior to the sample, rapidly spiked to ~500°C, after which it fell to around 200°C - 220°C for the remainder of the test. The thermocouples at the face-center and corner locations virtually track one another through the test, reaching a maximum temperature of ~440°C. In Fig. 6.b., after the experiment it is shown that the NH<sub>3</sub>BH<sub>3</sub> material has been extruded through the metal screen of the sample holder. When the material is removed from the exterior of the sample holder, it was found that there was no damage to the sample holder. This is consistent with experimental observations that place the melting point of NH<sub>3</sub>BH<sub>3</sub> at 104°C, and the first H<sub>2</sub> desorption temperature at 120°C [5]. The NH<sub>3</sub>BH<sub>3</sub> material undergoes some degree of melting, followed by hydrogen discharge at

120°C. These temperatures are well below the autoignition temperature of H<sub>2</sub> (~571°C); therefore, there must be some degree of temperature-assisted hydrolysis or similar reaction to provide the initial energy for the ignition event.

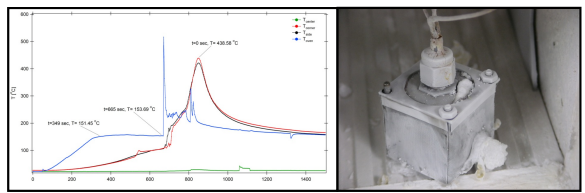


Figure 6. Results of the Self-Heating Test performed on NH<sub>3</sub>BH<sub>3</sub>. a) Thermocouple Readings. b) Photograph of sample container showing the volume expansion of NH<sub>3</sub>BH<sub>3</sub> upon heating.

The results of the Burn Rate test for  $NH_3BH_3$  are shown below in Fig. 7. As can be seen, a strong green flame was observed, characteristic of the flame color for boron. The thermocouple data is shown in Fig 7.b., where it can be seen that the temperature within the burning  $NH_3BH_3$  pile approached nearly 500°C, approximately the same temperature as the maximum reading of the ambient thermocouple from the Self-Heating test. The calculated burn rate for the  $NH_3BH_3$  material is 33 mm/s, 37% slower than the burn rate observed for the  $2LiBH_4$ ·MgH<sub>2</sub> and  $NaAlH_4$  materials [6].

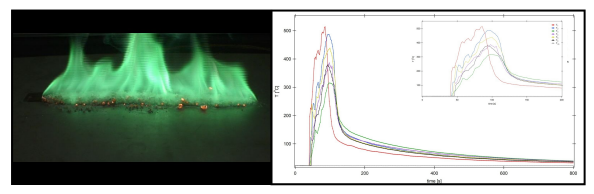


Figure 7. Results of the Burn Rate Test for the NH<sub>3</sub>BH<sub>3</sub> material. a) Photograph of Burning Material after 5 seconds, b) Thermocouple Readings

The findings of the current U.N. testing procedures for  $2\text{LiBH}_4 \cdot \text{MgH}_2$  and  $\text{NH}_3\text{BH}_3$  are shown below in Table 1. It can be seen that the more reactive material is the metal hydride  $2\text{LiBH}_4 \cdot \text{MgH}_2$  as compared to the less environmentally reactive  $\text{NH}_3\text{BH}_3$ . The differences in reactivity may be an important factor in the development of risk mitigation and engineering amelioration strategies developed for systems which employ these types of materials for hydrogen storage applications.

Material / UN Test	State	Pyrophoricity	Self-Heat	Burn Rate	Water Drop	Surface Contact	Water Immersion
2LiBH4'MgH2	С	70 I	Self-heated ~300 oC within 5 min at as Toven = 150 o is approached.	Flame propagated in 5 sec with burn rate of 52 mm/sec.	2 H2O drops required for near-instant combustion.	Material combusted	No ignition event recorded. Gas evolved at longer times. (5 min)
	D	Not tested	Not tested	Not tested	1 H2O drop required for near-instant ignition	Reaction observed with no flame	Reaction observed with no flame
NH3BH3	С	No ignition event. Hygroscopic material absorbed H2O from air.	Self-heated ~300 oC within 10 min, 5 min at Tover=150 oC	Flame propagated in 6 sec with burn rate of 33 mm/sec	No reactivity	No ignition event recorded. Gas evolved at longer times. (5 min)	No reactivity detected
	D	Not tested	Not tested	Not tested	No reaction	No reaction	No reaction

Table 1. Results of the U.N. Standardized Tests for  $2\text{LiBH}_4$ ·MgH<sub>2</sub> and NH<sub>3</sub>BH<sub>3</sub>. (C = Charged State, D = Fully Discharged State)

### 4.0 CONCLUSIONS AND RECOMMENDATIONS

The materials  $2\text{LiBH}_4$ ·MgH<sub>2</sub> and NH<sub>3</sub>BH<sub>3</sub> have been evaluated using modified versions of United Nations test methodologies for shipping and packing classification. It was found that  $2\text{LiBH}_4$ ·MgH<sub>2</sub> is reactive with both air and water under the conditions used in the current tests. In contrast, it was found that NH<sub>3</sub>BH<sub>3</sub> is relatively inert to water reactivity, but is reactive with air under the test conditions. These results, in combination with the thermodynamic and kinetic analysis, are being used to develop risk mitigation strategies for the hydride materials that have been considered within the program thus far. These results and proposed risk mitigation strategies will be disseminated so that users of these materials may take proper safeguards to minimize risk associated with their use and potential air and water exposure. Future work will consider additional materials such as AlH<sub>3</sub> and the use of numerical simulations to evaluate critical environmental exposure scenarios, with a strong focus on the implementation of the results from these standardized tests for system optimization.

### **5.0 ACKNOWLEDGEMENTS**

This work was funded by the United States Department of Energy. The authors would like to thank their IPHE partners for many discussions, including Dan Mosher of United Technologies Research Center, Dan Dedrick of Sandia National Laboratory, Nobuhiro Kuriyama of the National Institute of Advanced Industrial Science and Technology, Kyle Brinkman of Savannah River National Laboratory and Will James of Savannah River National Laboratory. Thanks also to Tom Autrey of Pacific Northwest National Laboratory, and Ned Stetson of the United States Department of Energy for their insight and discussions pertaining to this work.

#### **6.0 REFERENCES**

- 1. Finholt, A.E.; Bond, A.C. Jr.; Schlesigner, H.I. "Lithium Aluminum Hydride, Aluminum Hydride, and Lithium Gallium Hydride, and Some of their Applications in Organic and Inorganic Chemistry." *J. Amer. Chem. Soc.*, **69**, pp. 1199. (1947)
- Zalosh, "Metal hydride fires and fire suppression agents." J. Loss. Prevention, 21, pp. 214. (2008)
  DOT/UN Doc., Recommendations on the Transport of Dangerous Goods, Manual of Tests and Criteria, 3rd Revised Ed., ISBN 92-1-139068-0, (1999).
- 4. Vajo, J.; Skeith, S.; and Mertens, F. "Reversible Storage of Hydrogen in Destabilized LiBH<sub>4</sub>." *J. Phys. Chem. B.*, **109**, pp. 3719. (2005)

5. Karkamkar, A.; Aardahl, C.; and Autrey, T. "Recent Developments on Hydrogen Release from Ammonia Borane." *Material Matters*, **2**, pp. 6 (2007)

6. Mosher et al, J. Alloys and Compounds, 446, pp. 707 (2007)

7. Xu, Q. and Chandra, M., "A portable hydrogen generation system: Catalytic hydrolysis of ammoniaborane." *J. Alloys and Compounds*, **446-447**, pp. 729. (2007)

8. Chandra, M. and Xu, Q. "Dissociation and hydrolysis of ammonia-borane with solid acids and carbon dioxide: An efficient hydrogen generation system." *J. Power Sources*, **159**, pp. 855. (2006)