Main Group Element Chemistry for Hydrogen Storage and Activation

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> Project ID # STP5

This presentation does not contain any proprietary or confidential information

# Overview

#### Timeline

- Project start date: FY05
- Project end date: FY09
- Percent complete: New Start

#### Budget

- Total project funding: \$1,780,690
  - DOE share: \$1,273,711(requested)
    UA share: \$506,979
- \$0K in FY04
- Funding for FY05: \$432,077K
   \$226,510(DOE), \$205,567 (UA)

#### Barriers

- Barriers addressed
  - Cost
  - Weight and Volume
  - Efficiency
  - Regeneration

#### Partners

DOE Center of Excellence for Chemical Hydrogen Storage: with LANL, PNNL UW, UC-Davis, Penn State, UA, UPenn, Northern Arizona, UCLA a Millenium Cell, Rohm and Haas, and US Borax.

## Objectives

- New Project: Start CY2005
- Focus is the development of new chemistries to enable DOE to meet the technical objective: "By 2010, develop and verify onboard hydrogen storage systems achieving 2 kWh/kg (6 wt%), 1.5 kWh/L, and \$4/kWh.; by 2015, 3 kWh/kg (9 wt%), 2.7 kWh/L, and \$2/kWh" by using chemical hydrogen storage systems
- Develop and implement imidazolium-based H<sub>2</sub> activation chemistry
- Develop and implement systems based on polyhydrides of main group elements, *e.g.*, phosphorous
- Develop and implement cyanocarbon systems for H<sub>2</sub> storage
- Provide computational chemistry support (thermodynamics, kinetics, properties prediction) to the experimental efforts of the DOE Center of Excellence for Chemical Hydrogen Storage to reduce the time to design new materials and develop materials that meet the 2010 and 2015 DOE objectives.

## Approach

- Use novel chemistry approaches to synthesize compounds for easily reversible addition/elimination of H<sub>2</sub> based on our novel, stable carbene chemistry for use in H<sub>2</sub> storage systems.
- Use first principles computational chemistry approaches on advanced computer architectures to predict the electronic structure of molecules to obtain thermodynamic and kinetic information in support of the design of hydrogen storage materials and of catalysts to effect easy release and addition of H<sub>2</sub>.
- Develop a thermodynamic approach for chemical H<sub>2</sub> storage based on exploiting ΔH and ΔG coupled with Le Chatelier's principle to manage H<sub>2</sub> addition and release in chemical compounds. Potential approach for cold-start issues.
- Experimental close interactions with partners: LANL, UC-Davis
- Computational interactions with LANL, PNNL, UW, UPenn, UC-Davis, UCLA

## Technical Accomplishments/ Progress/Results

- Structural characterication of an imidazolium borohydride.
- Formation of carbene  $H_2$  adducts.
- Model compound available for cyanocarbon·H<sub>2</sub> adducts.
- Reliable computational chemical prediction of  $H_2$  release energetics from borane amines and carbenes –demonstration of methodology and further development of  $\Delta G/\Delta H$  concepts.

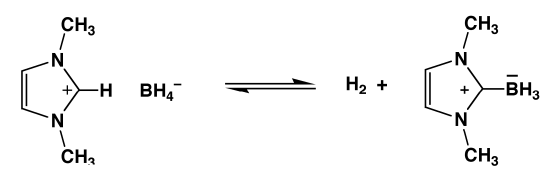
# Examples of % by weight of $\rm H_2$ of compounds under study

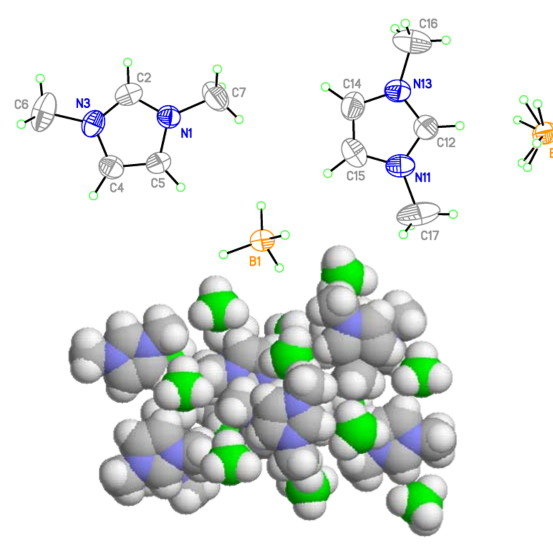
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Reaction or compound	<u>% H</u> 2
$NH_4BH_4(s) \rightarrow NH_3BH_3 + H_2$ $NH_3BH_3 \rightarrow NH_2BH_2 + H_2$	6.1% 6.5%
$[PH_4][BH_4]$ (release all H <sub>2</sub> ) PH <sub>3</sub> BH <sub>3</sub> (release all H <sub>2</sub> ) [BH <sub>4</sub> ][NH <sub>4</sub> ] (release all H <sub>2</sub> )	16% 13% 24%
carbene-derived compounds	7-14%
$R^2$ $N$ $C^2$ $N$ $+$ $C-H$ $N$ $+$ $N$ $+$ $N$ $+$ $N$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$	$BH_4^-$

 $\mathbf{R}^1$ 

#### Imidazolium Borohydride Solid State Structure Determined



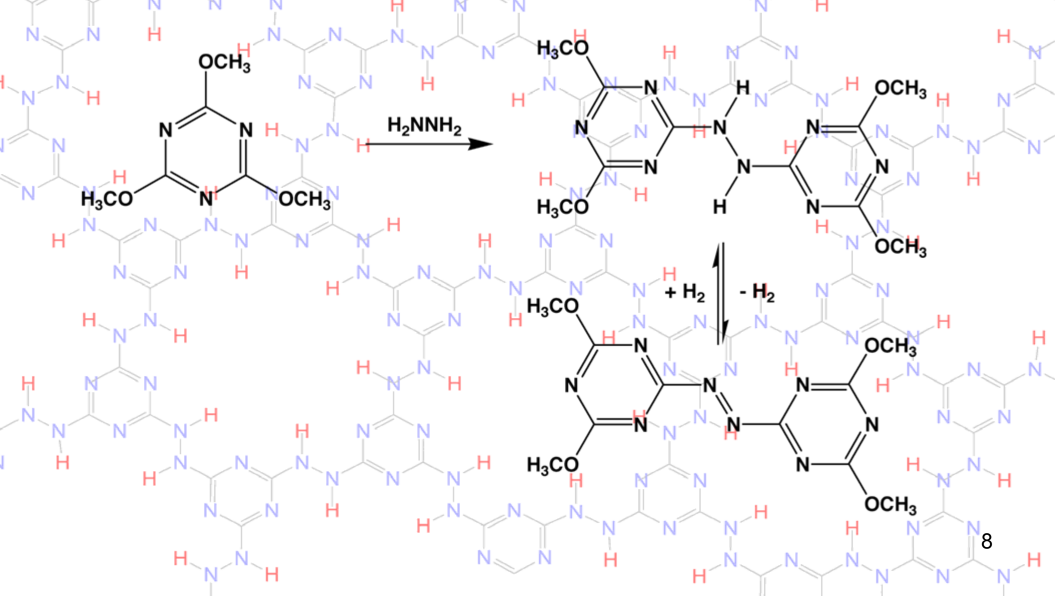


The first X-ray structure of an imidazolium borohydride was determined.

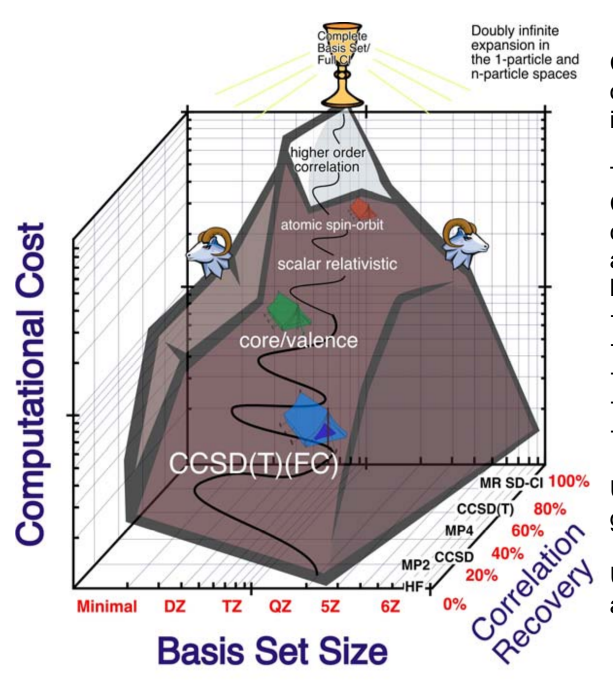
Structure shows interactions between hydrogen at C-2, C-4 and the borohydride and evidence for an H-H "hydrogen bond"

#### **Cyanocarbon**·H<sub>2</sub> Adducts Synthesis

Work initiated on a 2-dimensional cyanocarbon polymer (illustrated in the background) as a candidate for hydrogen uptake and release. The first unimolecular models have been synthesized. Characterization of these materials is in progress.



#### High Level Computational Chemistry for H<sub>2</sub> Storage



Calculate accurate molecular heats of formation (±1 kcal/mol) by ab initio molecular orbital theory:

Total atomization energy at the CCSD(T) level extrapolated to the complete basis set limit using the augmented-correlation consistent basis sets

- + core-valence corrections
- + scalar relativistic
- + spin orbit
- + zero point energy
- + thermal corrections (0  $\rightarrow$  298K)

Use atomic heats of formation to get molecular heats of formation

Use MP2 or CCSD(T) geometries and frequencies

## Reaction Energies in borane amines in kcal/mol @ 298K for chemical hydride storage

Reaction energies for the sequential release of  $H_2$  from borane amines (gas phase) based on the most accurate heats of formation now available (calculated by us).

$BH_3NH_3 \rightarrow BH_2NH_2 + H_2$	∆H = -5.1
$BH_2NH_2 \rightarrow HBNH + H_2$	∆H = 31.4
$HBNH \rightarrow BN + H_2$	∆H = 134.3

 $BH_3NH_3$  will be a good source of  $H_2$  as the release of  $H_2$  from this species is not far from thermoneutral. Cannot make T too high as entropy also favors release of  $H_2$ .

C2 gaseous hydrocarbons cannot release  $H_2$  (except at high T due to T $\Delta$ S).

The reaction energies for forming BN/C2 and BHNH/CHCH are ~ equal. Manifestation of similarity of the isoelectronic  $C_2H_{2m}$  and  $BNH_2m$  systems except for m = 3.

## Bond Energies in borane amines in kcal/mol @ 0K in kcal/mol for chemical hydride storage

Reaction	B.E.	Bond type
$\begin{array}{l} BH_3-NH_3 \to BH_3 + NH_3 \\ CH_3-CH_3 \to 2 \ CH_3 \\ BH_2=NH_2 \to BH_2 + NH_2 \\ CH_2=CH_2 \to 2 \ CH_2 \ ({}^3B_1) \\ BHNH \to BH + NH \\ HCCH \to 2 \ CH \end{array}$	25.9 87.9 139.6 171.9 178.5 228.1	dative C-C σ B-N π + B-N σ C-C π + C-C σ

For C<sub>2</sub>H<sub>4</sub>, the C-C  $\pi$  bond is ~65 kcal/mol giving a C-C  $\sigma$  bond energy of 107 kcal/mol. Due to sp<sup>2</sup> –sp<sup>2</sup> bonding, not sp<sup>3</sup>-sp<sup>3</sup> bonding.

Estimate the B=N  $\pi$  bond energy by calculating the rotation barrier in BH<sub>2</sub>NH<sub>2</sub> Rotation Barrier at 0K = 30.0 kcal/mol =  $\pi$  bond energy

Low  $\pi$  bond energy yields a very strong B-N single bond of 109.6 kcal/mol, comparable to the C-C  $\sigma$  bond in C<sub>2</sub>H<sub>4</sub>. The high B-N  $\sigma$  +  $\pi$  bond energy in BH<sub>2</sub>NH<sub>2</sub> is consistent with the fact that it does not release H<sub>2</sub> readily.

# Can the salt [BH<sub>4</sub><sup>-</sup>][NH<sub>4</sub><sup>+</sup>] serve as an H2 storage system?

Estimate the lattice energy  $U_{L}$  of the salt from the empirical expression:  $U_{L} = 2 I [\alpha V_{m}^{-1/3} + \beta]$  where *I* is the ionic strength (I = 1),  $V_{m}$  is the molecular (formula unit) volume  $\alpha$  and  $\beta$  are empirical parameters. Equation good to  $\pm 5$ kcal/mol

Use experimental volumes for  $BH_4^-$  and  $NH_4^+$  to obtain a lattice energy of 151.3 kcal/mol and  $\Delta H(0K)$  for the salt of -9.3 kcal/mol.

The reaction from the ionic solid to the gas phase products  $[BH_4^{-}]NH_4^{+}](s) \rightarrow BH_3NH_3(g) + H_2 \qquad \Delta H (0K) = 0.2 \text{ kcal/mol}$ is essentially thermoneutral and the salt would be a good source of H<sub>2</sub>.

Another possibility for the formation of  $H_2(g)$  is formation of  $NH_3BH_3(s)$   $[BH_4^{-}]NH_4^{+}](s) \rightarrow BH_3NH_3(s) + H_2 \qquad \Delta H(0K) = -16.8 \text{ kcal/mol}$ The calculated cohesive energy for  $BH_3NH_3$  is 17 kcal/mol and the enthalpy change for the hydrogen release reaction is substantially more exothermic than if  $BH_3NH_3(g)$  is produced.

# Stability of a salt requires that electron transfer cannot occur between the cation and anion

-Electron affinity (EA) of  $NH_4^+$  is very low as  $NH_4$  is a Rydberg molecule with only a weak binding of H to  $NH_3$  if at all. Estimate EA( $NH_4^+$ ) as the energy of the following reaction giving -4.84 eV

 $NH_4^+ + e^- \rightarrow NH3 + H$ 

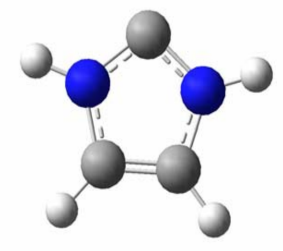
-Ionization potential (IP) of  $BH_4^{-1}$  is given by the reaction

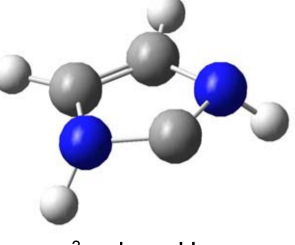
 $BH_4^- \rightarrow BH_3 + H + e^-$ 

as BH<sub>4</sub> is also a very weakly bonded system and EA(BH<sub>3</sub>) is very small (0.038 ± 0.015 eV, *J. Chem. Phys.*, **1989**, *90*, 795). The ionization potential of BH<sub>4</sub><sup>-</sup> is low, 3.89 eV.

The fact that  $EA(NH_4^+)$  and  $IP(BH_4^-)$  are comparable within 1 eV of each other is consistent with the fact that this salt can be synthesized.

#### Carbene Structures for H<sub>2</sub> Storage Systems

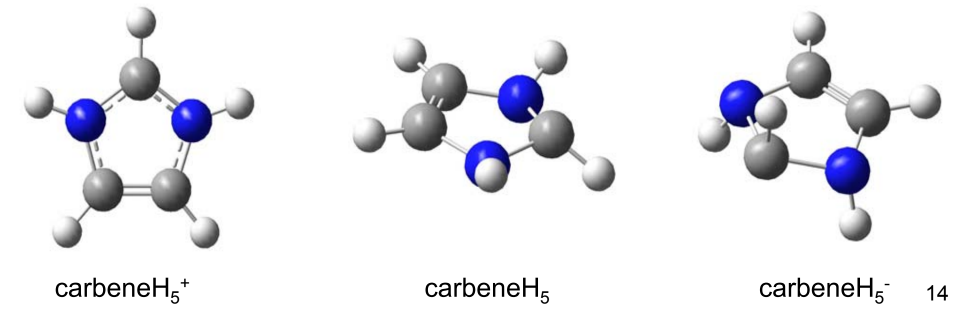




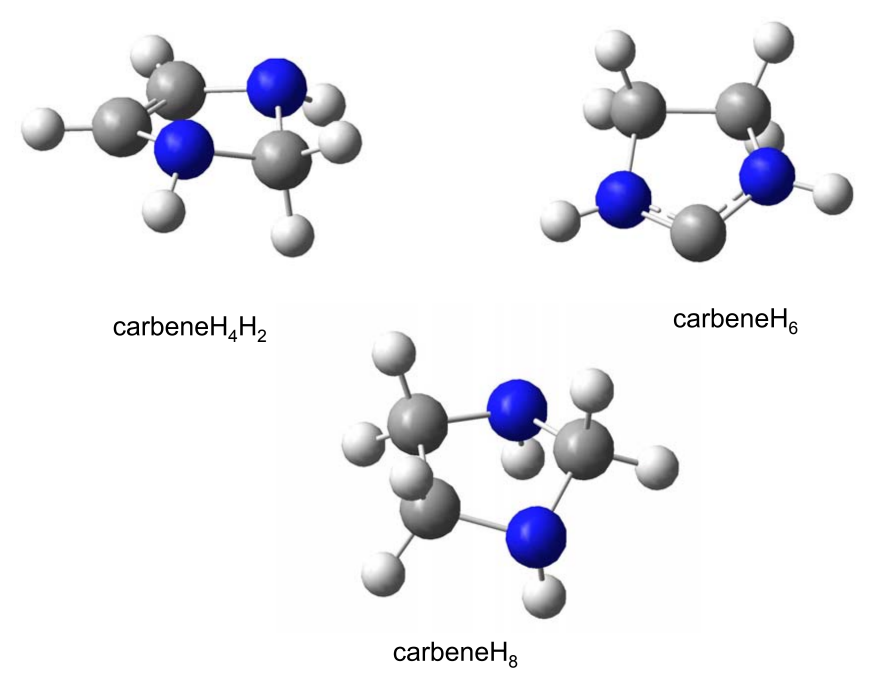
 $carbeneH_4$ 

<sup>3</sup>carbeneH<sub>4</sub>

Based on Arduengo's stable carbene



#### Carbene structures for H<sub>2</sub> Storage Systems



#### Carbene Reaction Energetics (kcal/mol) for H<sub>2</sub> storage at 298K

Carbene +  $H_2 \rightarrow \text{carbeneH}_2$   $\Delta H = -13.9$ Adding  $H_2$  to the carbene is exothermic by 13.3 kcal/mol -- very nice in managing release because we can use  $\Delta G$  to pull it off using Le Chatlier's Principle. T $\Delta S(298K) = +8.2$  kcal/mol

 $\label{eq:249.5} Carbene + H^+ \rightarrow carbene H^+ \qquad \Delta H = -249.5 \\ The PA of the simplest carbene is 249.1 kcal/mol. Very basic!$ 

Carbene  $\rightarrow$  <sup>3</sup>carbene $\Delta H = 90.5$ The singlet triplet splitting of the carbene is very large, 89.8 kcal/mol!

Carbene + H  $\rightarrow$  carbeneH $\Delta H = -31.8$ The C-H bond energy for adding an H to the carbene is only 32 kcal/mol.

CarbeneH +  $e^- \rightarrow$  carbeneH<sup>-</sup>  $\Delta$ H = 5.0 Addition of H- to the simplest carbene leads to autodetachment of the  $e^-$ .

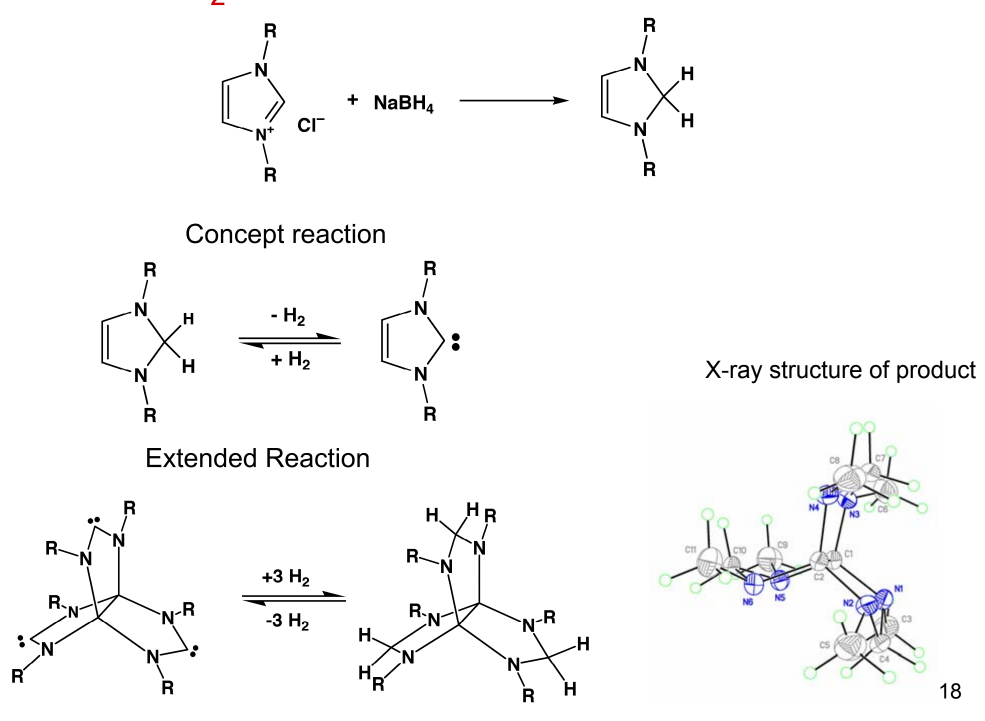
 $\begin{array}{ll} \mbox{Carbene} + \mbox{H}_2 \rightarrow \mbox{carbene}\mbox{H}_6 & \Delta \mbox{H} = -8.6 \\ \mbox{Adding}\mbox{ H}_2 \mbox{ to hydrogenate the double bond is exothermic by only 8.6 kcal/mol as} \\ \mbox{compared to -31 kcal/mol for hydrogenation of $C_2$\mbox{H}_4$.} \end{array}$ 

CarbeneH<sub>6</sub> + H<sub>2</sub>  $\rightarrow$  carbeneH<sub>8</sub>  $\Delta$ H = -32.5 Adding H<sub>2</sub> to the hydrogenated carbene is exothermic by 32.5 kcal/mol.

## Future Work

- Experimental and computational characterization of cyanocarbons for H<sub>2</sub> storage
- Experimental and computational studies on Carbene-H<sub>2</sub> adducts.
- Experimental and computational studies on imidazol(in)ium borohydrides and carbeneborane adducts
- Continue computational chemistry studies in overall support of DOE Center of Excellence for Chemical Hydrogen Storage

Carbene-H<sub>2</sub> Adducts: Proposed Experimental Work



## Computational studies of other potential molecular systems for H<sub>2</sub> chemical hydrogen storage

Look at isoelectronic molecules in the Periodic Table.

Use our advanced electronic structure methods on parallel computers to predict reaction energies for the following

```
\begin{split} & [\mathsf{A}\mathsf{IH}_4^{-}]\mathsf{NH}_4^{+}](\mathsf{s}) \to \mathsf{A}\mathsf{IH}_3\mathsf{NH}_3 + \mathsf{H}_2 \\ & \mathsf{A}\mathsf{IH}_3\mathsf{NH}_3 \to \mathsf{A}\mathsf{IH}_2\mathsf{NH}_2 + \mathsf{H}_2 \\ & \mathsf{A}\mathsf{IH}_2\mathsf{NH}_2 \to \mathsf{H}\mathsf{A}\mathsf{INH} + \mathsf{H}_2 \end{split}
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```
\begin{array}{l} [\mathsf{BH}_4^{-}]\mathsf{PH}_4^{+}](\mathsf{s}) \to \mathsf{BH}_3\mathsf{PH}_3 + \mathsf{H}_2\\ \mathsf{BH}_3\mathsf{PH}_3 \to \mathsf{BH}_2\mathsf{PH}_2 + \mathsf{H}_2\\ \mathsf{BH}_2\mathsf{PH}_2 \to \mathsf{HBPH} + \mathsf{H}_2 \end{array}
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 \begin{array}{l} [\mathsf{A}\mathsf{IH}_4^{-}]\mathsf{P}\mathsf{H}_4^{+}] (s) \to \mathsf{A}\mathsf{IH}_3\mathsf{P}\mathsf{H}_3 + \mathsf{H}_2 \\ \mathsf{A}\mathsf{IH}_3\mathsf{P}\mathsf{H}_3 \to \mathsf{A}\mathsf{I}\mathsf{H}_2\mathsf{P}\mathsf{H}_2 + \mathsf{H}_2 \\ \mathsf{A}\mathsf{I}\mathsf{H}_2\mathsf{P}\mathsf{H}_2 \to \mathsf{H}\mathsf{A}\mathsf{I}\mathsf{P}\mathsf{H} + \mathsf{H}_2 \end{array}
```

# Predict the thermodynamics and rate constants for reactions of alkoxyboron compounds with metal hydrides

Examples of disproportionation/conproportionation equilibria:

$$\begin{array}{rcrcrcrcrcrc} 2 \ HB(OCH_3)_3^{-} &\to H_2B(OCH_3)_2^{-} \ + \ B(OCH_3)_4^{-} \\ HB(OCH_3)_3^{-} \ + \ H_2B(OCH_3)_2^{-} \ \to \ B(OCH_3)_4^{-} \ + \ H_3B(OCH_3)^{-} \\ HB(OCH_3)_3^{-} \ + \ H_3B(OCH_3)^{-} \ \to \ B(OCH_3)_4^{-} \ + \ BH_4^{-} \end{array}$$

Reaction profile for hydride exchange:

 $HB(OCH_3)_3^- + B^*(OCH_3)_3 \rightarrow B(OCH_3)_3 + HB^*(OCH_3)_3^-$ 

Thermodynamics of hydride exchange with other hydrides  $HSiMe_3 + B(OCH_3)_{4-} \rightarrow Me_3Si(OCH_3) + HB(OCH_3)_3^ B(OCH_3)_3 + HOCH_3 + base \rightarrow H-base + B(OCH_3)_4^ HSi(OCH_3)_3 + B(OCH_3)_4^- \rightarrow Si(OCH_3)_4 + HB(OCH_3)_3^ HAI(OCH_3)_2 + B(OCH_3)_4^- \rightarrow AI(OCH_3)_3 + HB(OCH_3)_3^ HAIMe_2 + B(OCH_3)_4^- \rightarrow Me_2AI(OCH_3) + HB(OCH_3)_3^-$ 

Similar reactions involving other main group elements including:  $HGeMe_3$ ,  $HGe(OCH_3)_3$ ,  $HSnMe_3$ 

# Other Reaction modeling and Cyber-Infrastructure support

• Borane amine reactions: Predictions of reaction mechanisms including dehydro-oligomerization reactions, Lewis acid base addition reactions, hydride extraction, proton loss

• Predict nmr chemical shifts, infrared spectra, and UV-vis spectra for use in analyzing experimental data

- Boron hydride cluster modeling
- Electronic infrastructure support
  - Develop website for Center partners to share data, publications, presentations

- Implement electronic communication approaches to enable Center partners to communicate better

- Data management: Develop and construct database for thermodynamic, kinetic, and spectroscopic data for the Center

#### University of Alabama: Experimental Effort Timeline: Go/No GoDecision point is end of year 3

Task	Year 1	Year 2	Year 3	Year 4	Year 5
Synthesis of pendant functionalized imidazoles and imidazolines as candidates for $H_2$ activation					
Find conditions for $H_2$ activation by imidazole based systems					
H <sub>2</sub> elimination from imidazolium and/or imidazolinium borohydrides					
H <sub>2</sub> uptake by imidazolylidene boranes and/or imidazolinylidene boranes					
Synthesize and characterize LiH-carbene complexes					
Synthesize hindered imidazole-2-thione for red/ox studies					
Oxidize imidazole-2-thione with chemical reagents					
"Hydride" reduction of Imidazolethione oxid state +2					
Synthesis of oxidation state +4 sulfur derivatives					
Reduction of high oxidation sulfur species					
Synthesize carbene-SiH <sub>4</sub> complexes					
MW/struct. control of 2D cyanocarbon polymer					
Reduction of Cyanocarbon polymer					
Reversible Cyanocarbon H <sub>2</sub> uptake					
Optimize carbene MW					
Optimize cyanocarbon MW					
Develop catalysts for H2 release from carbene-based compounds					
Develop catalysts for H2 release from cyanocarbons					
Develop production chemistry					

#### University of Alabama: Computational Effort Timeline: Go/No Go Decision Points in collaboration with Center Experimental Efforts

Task	Year 1	Year 2	Year 2	Year 4	Year 5
Benchmark of density functional theory (DFT) method against accurate molecular orbital methods and experiment					
Accurate predictions of heats of formation of main group compounds for $H_2$ storage/release					
Computational results for B-O $\rightarrow$ B-H conversions – birate complexation/reduction					
Imidazolium borohydrides and carbene-borane adducts					
Cyanocarbon chemistry					
Carbene-main group compound adducts					
Polyhedral borane chemistry					
Initial database constructed for thermodynamic and kinetic data					
Continuous evaluation of go-no go decisions for different compounds and approaches by combining computational and experimental results in combination with partners					
Web site construction and maintenance					
Computational chemistry calculations on proposed Tier 3 compounds to predict thermochemistry and kinetics for novel H <sub>2</sub>					
storage compounds and materials based on C, N, and main group elements					
Computational chemistry calculations on proposed catalysts for Center including main group, organic, and boron-based					
compounds					

### **Publications and Presentations**

#### Publication:

"Thermodynamic Properties of Molecular Borane Amines and the [BH4-][NH4+] Salt for Chemical Hydrogen Storage Systems from Ab Initio Electronic Structure Theory," D. A. Dixon and M. Gutowski, *J. Phys. Chem. A*, 2005, accepted, April

#### **Presentations:**

Invited Presentation at the 229<sup>th</sup> National American Chemical Society Meeting, San Diego CA, March, 2005, Division of Fuel Chemistry, Computational Methods and Modeling in Fuel Chemistry Symposium (+ 2 page extended abstract): "High Level Computational Approaches to the Prediction of the Thermodynamics of Chemical Hydrogen Storage Systems and Hydrocarbon Fuels" David A. Dixon, Maciej Gutowski, Lisa Pollack, Theresa L. Windus, Wibe de Jong, University of Alabama, and Pacific Northwest National Laboratory

2<sup>nd</sup> Undergraduate Research Symposium, Arts and Sciences, University of Alabama, April 8, 2005

(1) "High level computational approaches to the prediction of the thermodynamics of chemical hydrogen storage systems," Jacob Batson and David A. Dixon, 2<sup>nd</sup> place in sciences

(2) "Cyanocarbons and Their Role in Chemical Hydrogen Storage," Michael Phillips and Anthony J. Arduengo

## Hydrogen Safety

#### The most significant hazards associated with this project are: Compound Flammability and Toxicity

- § At present no unique hazards have arisen as a result of these studies. Our approach avoids the use of free  $H_2$  which minimizes safety issues
- § All new reduced compounds are handled under inert atmospheres (N<sub>2</sub>) until combustion hazards are determined.
- § All cyanocarbon compounds are treated as toxic and worker exposure is strictly controlled.

#### Laboratory Safety

#### Our approach to deal with this hazard is:

- § All new reduced compounds are handled under inert atmospheres (N<sub>2</sub>) until combustion hazards are determined.
- § All cyanocarbon compounds are treated as toxic and worker exposure is strictly controlled
- § All procedures are reviewed by other laboratory personnel before execution.
- § Working alone in a chemical laboratory is not permitted.
- § A strict "Take-Two" policy is followed by all experimentalists.