



Reversible Hydrogen Storage Materials – Structure, Chemistry, and Electronic Structure

Ian Robertson and Duane Johnson (Angus Rockett, Stephen House and Pam Martin, Lin-Lin Wang, Jason Reich) University of Illinois 3 April 2010

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Overview



Timeline

- Project start date
 March 2005
- Project end date
 - February 2010
 - Application to extend project has been approved.
- Percent complete 90%

Budget

- Total project funding
 - \$1 600 000 committed
 - \$1 231 250 funded
- FY09: \$100,000
- Funds Remaining: \$132,000

Barriers

- Issues addressed
 - A: Understanding chemical response to hydrogen
 - B: Understanding system weight and particle packing
 - C: Theory support to address critical issues blocking experimental progress.

Partners

- Project lead: Sandia National Laboratory & MHCoE
- Brookhaven National Laboratory
- Savannah River National Laboratory
- University of Pittsburg, Georgia Tech and University of Hawaii ²





• Illinois' main purpose within the MHCoE is

- Advance the understanding of the microstructural and modeling characteristics of complex hydrides, helping with selection and down-selection.
- Provide more reliable theoretical methods to assess H-storage materials, including key issues affecting materials under study.
- Help identify down-selects, via experimental and theoretical characterization.
- Provide characterization expertise to other MHCoE and other critical DoE programs, as suggested by Program Officers.

• Some Illinois recent impacts

- Improved 3-D reconstructions and visualization to address the question of the dispersion of particles in scaffold.
- In MgH_{2} , AIH_{3} and MOFs, complete understanding dispersion of materials.
- In MgH₂ showed that there is no nanoparticle size effect for dehydrogenation enthalpies, although there is an effect on kinetic barriers with TM catalyst.
- Determined universal behavior for core-shell preference in binary nanoparticles.
- Demonstration that the amount of Ti needed to activate the system is less than introduced initially.
- In $Ca(BH_4)_2$ established the existence of numerous $CaB_{12}H_{12}$ polymorphs in intermediate phase, and the origin for observed amorphous-like behavior.





- <u>Task 1: Experimental investigation of</u> <u>MHCoE partner materials</u>
 - Determining microstructural and compositional changes in hydrogen storage materials due to hydrogenation and dehydrogenation cycles.

<u>Task 2: Incorporation and development</u> of new models

 Predicting and understanding structural, thermodynamic, and kinetic properties of hydrogen-storage materials, including size effects, with direct connection to ongoing experimental measurements, including in MgH₂, Alanes, and Borohydrides





• Electron tomography capability for compositional imaging has been improved significantly, enabling better resolution in the through thickness direction.

Confirming silver particles are distributed inside a MOF. (MOF is removed to show the particles.)



expanded

expanded and rotated.



Electron tomography shows the particles distributed throughout the scaffold. The new capability allows full three-dimensional rotation and exploded view capabilities to better visualize and characterize the particles.
 Current effort is exploring the location of MgH₂ in a scaffold – collaboration with the University of Hawaii.



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Role of Ti on particle size during cycling

Jensen (U. Hawaii) Robertson (U. Illinois)







Dedicated surface study using UHV **Scanning Tunneling Microscopy** to determine atomic-scale effects of Ti on AIH₃ storage system



- 20 ML of AI deposited on Si (111) substrate
- Annealed to form atomically flat Al (111) islands, tens of nm wide

Al islands (a) prior to and (b) following exposure to $5x10^{-8}$ mbar of UHP H₂ for 5 hours. Scan parameters: scan size = 100x100 nm², V = -1.136 V, (a) I = 0.360 nA, (b) I = 0.456 nA.

No change in morphology observed during exposure to H₂ gas
 This is not surprising, as H₂ has negligible sticking to pure Al



H₂ interaction changes when Ti is present on the surface



Following deposition of 0.02 ML of Ti at 300K and exposure to H₂:



 Ti apparent heights (0.2-0.35 Å) and diameters (6-7.5 Å) correspond with previously observed and calculated values* for first-layer substitutional positions.



Time-resolved study of interaction of H₂ with AI(Ti) surface





- No evidence of Ti motion or agglomeration during exposure.
- Formation of vacancy clusters (1.2-1.5 Å deep)
 - Grew in number and width over the course of the exposure.
- Surface AIH_x monomers formed at edges of vacancy clusters.
- Surface AIH_x apparent heights (2-4 Å) and apparent diameters (4-6 Å) are consistent with values observed for surface alane monomers (2.8 Å and ≈ 4 Å, respectively)**.



Changes in the I-V curves on the addition of hydrogen





The I-V curves suggest the formation of a small band-gap on hydrogen charging. This has required use of a different STM with superior spectroscopic capabilities. Experiments and *ab initio* calculations are in progress to verify this result. DFT results are being directly connected to STM results for AI, Ti-doped AI, and AIH_x cases.





- Characterization tools are available to determine the three-dimensional distribution as well as the size of particles either on surfaces or embedded in scaffolds.
- The role of Ti on clean AI surfaces appears to be to enhance the dissociation of molecular hydrogen to atomic hydrogen.
- In NaAlH₄ ball-milling decreases particle size, thereby enhancing kinetics; Al₃Ti has a possible role as a grain refiner; and sintering occurs during dehydrogenation and de-sintering during hydrogenation.



Characterization of Ca(BH₄)₂: Reactions and Products







Both reactions show CaB₁₂H₁₂ is stable, but with many polymorphs (denoted by +).
Same ground-state as *Ozolins et al.* [*JACS* 2009, 131, 230] but with many polymorphs.
Results show that CaB₁₂H₁₂ is destabilized at higher T (but the kinetic barrier is high).
Due to CaB₁₂H₁₂ stability, H-release will decline each cycle at lower T (observed) as it arises from remnant CaH₂. Full release only possible at high T.



Are the CaB₁₂H₁₂ polymorphs observed in cycled material?



Experimental (TEM and X-ray Diffraction) scattering indicates amorphous state. Debye-Waller, finite-size (Sherrer), or volume-fraction effects related to the coexisting phase alter the location of peaks, so we compare locations in 2θ.



- Multiple CaB₁₂H₁₂ polymorphs are indicated by matching structure in diffraction.
- Enthalpy and structure characterization suggest dehydrogenation in repeated cycling arises from CaH₂, rather than the very stable CaB₁₂H₁₂, limiting low-T Ca(BH₄)₂ as reversible H-storage media. Higher T can be reversible (previous slide).



Hydrogen Desorption from MgH₂: Nanoparticles and Bulk-Terminated Surface



Two Relevant Energies

- **1. Thermodynamics** enthalpy difference between two states.
- 2. Kinetics

– energy barriers between these states.

Two Critical Main Questions

Is there a *size effect* for H-desorption for each case? For a finite cluster, is the MgH₂ cluster at a global energy minimum? Both full and single-site calculated desorption are affected by this.









- $(MgH_2)_n$ clusters of < 5 F.U. (\geq 5 F.U.) have a smaller (larger) desorption energy than bulk.
- Similar to recent, higher accuracy, QMC results [Wu et al., JACS 2009, 131, 13918].
- 31 MgH₂ is an amorphous cluster, similar to de Jongh et al. [JACS 2005, 127, 16675] but unlike that of Larsson et al. [PNAS 2008, 105, 8227], which are metastable.



Hydrogen Desorption From Low-Energy Mg₃₁H₆₂ Amorphous Cluster





- ✓ We reproduce *spurious exothermic* results of Larsson et al. [PNAS 2008] that used bulk-terminated (locally relaxed) rutile-based clusters.
- ✓ In contrast, there is no exothermic H-desorption, due to large relaxation of bulkterminated cluster to the amorphous state (found via *simulated annealing*).
- ✓ $\Delta E_{H-desorp}$: "surface site" 246 kJ/mol-H₂ vs. bulk-surface site 256 kJ/mol-H₂.
- ✓ For fully desorbed $\Delta E_{H-desorp}$ is 67 kJ/mol-H₂ in cluster vs. 62 kJ/mol-H₂ in bulk.
- ✓ Similar results for Ti and Fe catalyst site (to be published).



Desorption From MgH_2 (110) Surface: Is there a size effect or catalyst effect?



Compare bulk-terminated surface with nanocluster results.





- \checkmark ΔE_{H-desorp}: 256 kJ/mol-H agrees with cluster "surface site" 246 kJ/mol-H₂.
- ✓ Step-edge site should be comparable to "edge site" in cluster (ongoing).
- ✓ Essentially NO thermodynamic size effect for desorption energy.
- \checkmark The barrier reduction for H removal kinetic size effect is under study.



Two-H Desorption from MgH₂(110) Surface: VASP Energetics and Kinetic Barriers



- For 2-H desorption, the lowest-energy structures involve in-plane H, albeit the BH1-PH3 and BH1-SBH1 pairs are close in energy.
- However, see next slide, BH1-PH3 has the lowest-kinetic barrier.
- Large difference between 1-H and 2-H cases, same as Du et al. [J Phys Chem C, 2007].
- Similar to nanoclusters, we need to consider the effect of large structural distortions for initial and final configurations, especially with catalyst (see next).
- Bulk diffusion barrier for H is ~0.2 eV, so rates controlled by desorption energy.



Two-H Desorption from MgH₂(110) with Catalyst: **VASP Energetics and Kinetic Barriers**





- For 2-H desorption, the lowest-energy structures (like NPs) involve dramatic local reconstructions around the catalyst site, with Ti (V) preferring 8 H (7 H) Coord. No.
- **Kinetic barrier results:** For 2-H desorption, there is a ~20% change of barrier due • to V (catalyst) site. A "concerted" bulk-to-surface H motion is found.
- Initial Ti site results show no barrier. However, a careful search for true "final" state configuration (involving concerted motions) is underway via ab initio MD, as barriers are sensitive to initial-to-final state choices for NEB.



Universal Core-Shell Preference in Binary, Transition-Metal Nanoclusters for Catalysis





Segregation Energy in Nanocluster

•Supported principally by DOE/Catalysis and partially by DOE/MHCoE

Energy Difference between Surface Segregated and Internal "red" atom



A negative (positive) Segregation Energy indicates that the surface is the (un)favorable site.

- In contrast to most suggestions, core-shell preference is determined by only two quantities: cohesive (or surface) energy and size.
- Molecular adsorption can change the preference (changes surface energy).
- All known experimental results are reproduced with this correlation map.
- Correlation also reproduces surface-alloy segregation preference.
- Universal behavior shown via "tight-binding/Debye-Hückl" model.

Wang and Johnson, J. Amer. Chem. Soc. 131, 14023-14029 (2009).





- The core-shell behavior in transition-metal nanoparticles is governed by cohesive energy (or surface energy) and size differences, which predicts all observed structure.
- No nanoparticle size effect for dehydrogenation enthalpy exists; there is a reduction in kinetic barrier with catalyst.
- We are able to reproduce quantitatively the experimental observations. Notably errors exist in QC literature that use small clusters to approximate surfaces: So, there are no exothermic reactions in MgH₂ system.





- Sandia National Laboratory –Determining if nanoparticles coat the surface or fill the internal structure in a MOF. (Mark Allendorf non MHCoE collaboration)
- University of Hawaii Determine if MgH_2 particles are distributed inside the scaffold. Also, theory activity with to determining properties of MgH_2 clusters.
- Brookhaven National Laboratory determining the structure and composition of AI-H system. Surface study of Hydrogen interactions on surfaces and the effect of Ti.
- Savannah River National Laboratory characterization of aluminum- based hydrides
- U. of Pittsburgh theoretical characterization of metal- $B_{12}H_{12}$ (de)hydrogenated intermediate phases, including amorphous.
- Georgia Tech theoretical characterization of metal- $B_{12}H_{12}$ (de)hydrogenated intermediate phases, including amorphous.





- Determine if the preliminary results from the surface study are valid, especially the spectroscopy results.
- Determine if atomic hydrogen on clean AI surfaces interacts in the same manner as molecular hydrogen when Ti is present.
- Repeat the study but with the addition of contaminants carbon and oxygen to determine how they will impact the hydrogen surface interaction.
- Complete *ab initio* calculations of projected density-of-states and Fermi's golden rule to interpret the above STM experimental results.
- These studies will provide insight as to how to manipulate the surface composition to assist in the regeneration of aluminum hydrogen systems as well as enhancing our understanding of the role of Ti in general.
- Determine if the MgH₂ nanoparticles are distributed throughout the scaffold.
- Complete MgH₂ Kinetic barrier calculations for nanoparticle and bulkterminated surface (initial results show ~20% reduction vs. size).





Supplemental Slides



How does hydrogen interact with clean surfaces in the presence and absence of a catalyst ?

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- Existing work strongly suggests that how hydrogen interacts with pure and contaminated surfaces is crucial to understanding the fundamental processes of hydrogen uptake and release as well as impact of catalyst species. To investigate we have probed the interaction of molecular hydrogen with clean AI surfaces using Scanning Tunneling Microscopy
 - Thin film of AI has been grown epitaxially on Si(111) substrate





Si(111) with AI epitaxial layer chosen due to know growth conditions and similarity to other hydrogen-storage materials. (See, e.g., Hasan, et al. Vacuum 41 (1990) 1121.)



Contaminants change response of surface to hydrogen exposure !





Two types of Ti atoms:

- 0.3-0.4 Å high, 4 Å wide → surface Ti?
- 0.2 Å high, 6 Å wide \rightarrow subsurface Ti?
- 0.3-0.4 Å high Ti disappear, leaving almost entirely 0.2 Å high Ti.
- Instead of many small vacancy clusters, a few very large ones.
- Formation of large secondary surface islands.
- Fewer, but larger, highly mobile surface features (AIH $_x$?).



Transition-Metal Surface Alloys for Catalysis: "Design Maps" for Surface Enrichment



We apply same segregation energy arguments to surfaces.

fcc(111) Surface Segregation Energy





- Again, we observe "universal" behavior for Core-shell (bulk/surface) preference from NP to bulk surfaces – driven by cohesive/surface energy and size only!
- "Maps" must be amended to include ligand-modified (functionalized) cases, which can alter (even reverse) surface enrichment.
- Correlation reproduces known surface-alloy segregation.
- Observed surface preference can depend on orientation, e.g., Ni-Pt has Ni on(110).