# **Optimization of Nano-Carbon Materials for Hydrogen Sorption**



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## Hydrogen Sorption Center of Excellence

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The presentation contains no proprietary, confidential, or otherwise restricted information

# Overview

# Timeline

- Start: 01 February 2005
- End: 31 January 2010
- 60% complete

# Budget

- Total project funding
  - DOE: \$1,065,209
  - Contractor: \$269,181
- Funding in FY07 \$235,000
- Funding for FY08 \$235,000

Barriers to be addressed

 Estimating hydrogen storage capacity of 3D-foams near ambient conditions

**Barriers** 

- Conversion of VANTA into solid superacids with high surface area and charge by introduction of electronegative tetrafluoroborate
- Identifying the catalytic role of metal on support for hydrogen adsorption and its spillover
- Understanding thermodynamics and kinetic mechanisms of spillover

# **Partners**

• NREL, Air Products Corp, regular teleconferences, face-toface meetings (Houston, Golden, CO, Washington, Gaithersburg, ORNL TT, MRS San Francisco, MRS Boston).



OVERALL: Model materials structures' interaction with hydrogen, optimize their makeup for storage and assess the volumetric and gravimetric capacity. Provide recommendations for the synthetic goals (e.g. pore/channel size, metal enhancement routes).

### 2007:

- Identify the obstacles (thermodynamics and kinetic) for the spillover for suggesting the materials design to overcome them
- Implications of metal aggregation on the hydrogen storage
- Estimator of volumetric and gravimetric capacities of foam, comparison with NTbundles
- Enhance binding of H<sub>2</sub> by introducing charge into the carbon lattice by adding a highly stable superacid anion that also acts as a spacer

### 2008:

- continue the above and
- Synthesis of metal- and electronegative-group- (F, BF<sub>3</sub>) enhanced VANTA (vertically aligned nanotube arrays, contrast to fibers) for H<sub>2</sub> adsorption
- Explore doping as an anchor to metal/metal cluster, role of bridges, and dopants on the threshold of spillover
- Hydrogen sorption on novel structure of B (fullerenes, sheets, nanotubes)

# **APPROACH**



+/- 1 meV

# **1** Rapid computation of capacity

 via Grand Canonical Monte Carlo simulations for the structures of interest based on judicious choice of potential (of paramount importance)

depths of potential obtained from experiment the (weaker) and ab initio calculations (stronger) differ by 2 meV, greatly affecting the capacity estimates

> • perform GCMC with both potentials to determine the range where the storage capacity is.



For technical accomplishment 2

# **2** Experimental

• VANTA arrays of 100 μm long carbon nanotubes, grown in large amounts with **CVD** methods. The alignment enhances gas transport into the array and thermal conduction.

• The aligned nature also lends itself to insertion of bulky anions, e.g. BF<sub>3</sub>-Δ

# APPROACH

Me-enhanced sorption on carbon receptors



 $\mu(H_2-gas) > \mu(H@metal) > \mu(H@support)$ 

Identify thermodynamically favorable and kinetically accessible paths Reconcile strong H binding to Me with possibility of spillover, by comparing the chemical potential of H on metal to that on receptor Role of bridges, defects and dopants on the overall process of spillover

**4** Stability of Me-atom anchors against aggregation, analysis of binding energies and diffusion barriers through DFT calculations

Energy states and barriers calculated with DFT and using NEB method

### **Capacity** of 3D-foams for H<sub>2</sub> storage

### technical accomplishments 1



Ding et al JCP 127, 164703 (2007)

 Nanometer pores and channels
Geometrical surface ~2600 m<sup>2</sup>/g fully accessible

Capacity

- Sensitive to choice of potential
- At 77 K is comparable to MOF's

Foams could meet both volumetric and gravimetric goals

$$u(r) = 4\varepsilon [(\sigma/r)^{12} - (\sigma/r)^6]$$
  $u(r) = Ae^{-\alpha r} + C_6 r^{-6}$ 

- ε = 3.7 eV/molecule
- σ **= 3.0 Å**

Wang et al J Low Temp Phys 41, 611 A = 1100 eV/molecule C<sub>6</sub> = -17 eV Å<sup>6</sup>/molecule

Patchkovski et al PNAS 102 10439



# Comparison of foam with (10,0)-NT bundle

technical accomplishments 1



Storage capacity as function of wall-wall distance D Volumetric % peaks at wall-wall distance of D  $\sim$  7 Å agrees with previous studies

Storage capacity of foams is superior to bestseparated, similar diameter nanotubes bundle (easier to synthesize)

# Excess H<sub>2</sub>



## **Experimental analogues produced**

### technical accomplishments 1

### **Zeolite-templated nanostructured carbon materials**



Z. Yang, Y. Xia, and R. Mokaya, J Am Chem Soc 129, 1673 2007; T. Roussel, et al J Phys Chem C 2007, 111, 15863



technical accomplishments 1`

Which is "ideal pore" ? C<sub>60</sub> interior *attracts* one H<sub>2</sub>

What if we could utilize the *repulsion* for storage



Existing giant fullerenes as ~C<sub>1000</sub> sustain great internal pressure, permitting H storage.

How it can be placed there remains yet unexplored



Fullerene Nanocage Capacity for Hydrogen Storage

System for synthesis of metal- and electronegative-group- (*F*-, *BF*<sub>3</sub>-) enhanced VANTA



Reaction system for generation of fullerene superacid boron tetrafluoride salts, to enhance H2 polarization for storage

technical accomplishments 2

Parameters for best growth of VANTA scaffolds have been defined and VANTA are now available for addition of electronegative groups for enhanced  $H_2$  adsorption.

A variable temperature reactor has been built and tested for controlled exposure of samples to fluorine and boron trifluoride.

One gram samples have been converted to a carbon nanotube salt and will be tested for hydrogen uptake in collaboration with NREL and Air Products. **Spillover kinetics** 

### technical accomplishments 3



# Vacancy formation and front stability





calculations at T = 0

### technical accomplishments 3

- Near-front kinetics and stability of the hydrogenated graphene island have been tested
- The new CH phase can absorb H around it
- Formation energy of vacancy (or DV) is high, rendering the front propagation via vacancy diffusion unlikely at storage temperatures

## Metal mobility role to be investigated

**Overcoming nucleation barrier of spillover** 

technical accomplishments 3



Nucleation barrier decreases with lower T and higher P Can be overcome by possibly "bridges" [R Yang] and the precursor H-patches [NREL work]

### technical accomplishments 4

## Metal clustering and effect on H<sub>2</sub> capacity



- Metal clustering is energetically and kinetically feasible at room temperature
- The diffusion frequency is ~10<sup>12</sup> exp(-E<sub>b</sub>/k<sub>b</sub>T)~10<sup>5</sup>-10<sup>10</sup> s<sup>-1</sup> at room temperature Krasnov, Ding, Singh, and Yakobson, JPCC, 2007, 111, 17977; method: B98/6-31G, Gaussian03
- Can be prevented by pinning on defects or via doping [NREL: Zhao et al PRL 94, 2005]
- Transition from Kubas sorption to H on large Me-clusters can lead to spillover 14

# **H** on catalyst: spillover-saturation window

### technical accomplishments 3

Determine the energy levels for H on free Pd<sub>4</sub> cluster and compare it with the onset of spillover onto C-receptor



\* calculations at 0 K, free cluster



• Several isomers were studied, only lowest energy ones are shown

 First H<sub>2</sub> gets adsorbed via dissociation, subsequent H<sub>2</sub> via Kubas type of interaction

Cluster saturates with  $9H_2$  (n = 18), spillover is favored after  $6H_2$  (n = 12)

## **H** desorption from catalyst

Here we answer the following questions:

What would be the energy cost to remove H atom from saturated cluster? Does it depend on the type of H dissociative/Kubas?



- Cost to remove dissociated (H) and non-dissociated (Kubas H-H) H is almost same
- Removing odd number of H costs more energy, we believe that the difference will go down for the larger size clusters, this could be a bottle neck for the sustainable spillover through smaller clusters

# H on catalyst on receptor/support: spillover-saturation window Effect of <u>support</u> on the H energy on metal



technical accomplishments 3



Spillover Threshold:

• On graphene metal saturates sooner than in the free cluster

- Onset of spillover is very close to metal full saturation
- First H<sub>2</sub> again gets adsorbed via dissociation, subsequent via Kubas
- The cluster saturates with 5H<sub>2</sub>

Kinetics: activation barriers for spillover Can metal hop from saturated cluster onto pristine graphene?



• Final state is not stable. H atom returns to metal spontaneously (implies possibility of reverse spillover)

• H further away from metal is stable, but high in energy (2.54 eV). Spillover on a pristine graphitic surfaces would be difficult. Curvature, defects, or modifications can facilitate

### technical accomplishments 3



• Dramatic decrease in energy, H on the phase 0.15 eV more stable—first sign of possibility of spillover

• The combination of saturated metal cluster with the hydrogenated phase formation would enhance the spillover process

# future work FY08-FY09

### spillover dynamics

- 1 Further energy and thermodynamics calculations: effects of size of metal cluster, dopants, bridges, on the H migration barriers from activator to receptor (along with v-signatures for experimental detection). Extend to other receptor geometry/materials (MOF, Met-Car).
- Quantitative analysis of the <u>front propagation model</u> in contrast to conventional <r<sup>2</sup>> ~ D×t diffusion. Compute the mobility barriers for the metal cluster to study the "scooter" effect, in which the chemically adsorbed H does not move, rather the metal moves.

### **3D-foams**

2 Generate several foams and optimize the storage capacity, using GCMC code for evaluating experimentally accessible surface areas. Ability to generate isotherms "on the fly" for any C-based structure. Consider doping the 3D-foams for better capacity.

### VANTA synthesis

3 Optimization of VANTA for area and  $H_2$  storage. Conversion of  $\pi$ -delocalized carbon systems (e.g. giant fullerenes, graphene sheets) and nitrogen doped carbon systems, to superacid salts. Test for hydrogen storage.

### **Me-centers stability**

4 Generate tailor made yet experimentally feasible C-based materials to achieve stronger metal binding, on defects or dopant-sites of the carbon lattice.

## summary

**Relevance:** Path-finding to reach DOE goals by modeling of major options: *spillover,* sorption on *3D-foams, metal-enhancement, doping* 

Approach: Quantum *ab initio* and empirical representations of H-carriers interaction, to assess their retaining ability. Selectively test synthesis options.

#### **Technical Accomplishments and Progress:**

Storage capacity of 3D-foam assessed, found to be comparable with the best MOFs.
Catalytic activity of metal clusters, relevant to spillover was determined and onset of spillover both in air and at receptor was computed

• Hydrogenated phase and saturated clusters are identified as the key element for spillover.

• Reactor constructed for conversion of carbon systems to superacid salts, VANTA's have been converted to superacid salts

• Addressed aggregation of metal atom at elevated temperatures, detrimental for storage

**Tech Transfer/Collaborations:** Partnership with NREL, Air Products, NIST (spectroscopy). Partnership with U Michigan and U Nevada Las Vegas (external).

Future Research: ● Emphasis on theory spillover dynamics, including other material-receptors (in collaboration with Yang's project) ● Optimized 3D-foam generation for storage and large surface and screening of Me-centers via doping and defect for stability against aggregation. ● Nitrogen doped carbon materials, large fullerenes and graphene sheets as high area superacid materials for hydrogen storage.

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