

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**

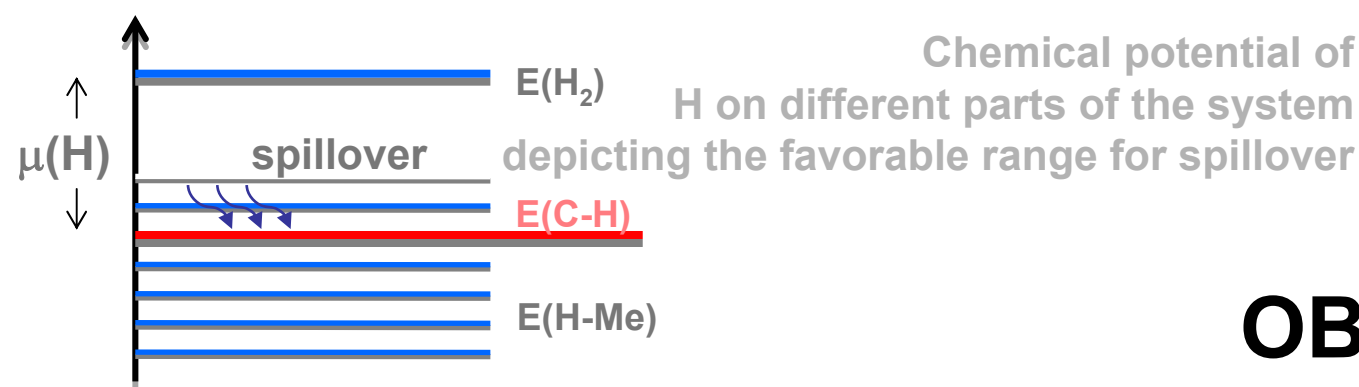
Barriers

Barriers to be addressed

- **Estimating hydrogen storage capacity of 3D-foams near ambient conditions**
- **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-to-face meetings (Houston, Golden, CO, Washington, Gaithersburg, ORNL TT, MRS San Francisco, MRS Boston).



OBJECTIVES

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).

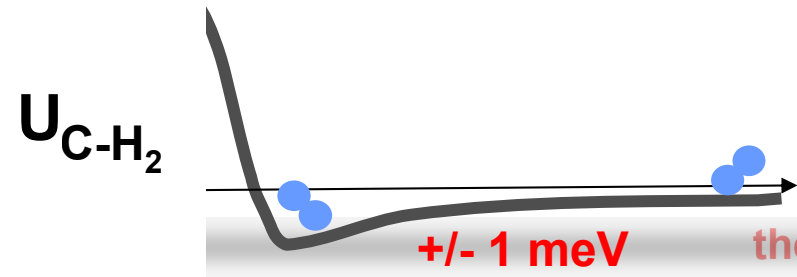
2007:

- 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 NT-bundles
- Enhance binding of H_2 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_3^-) enhanced VANTA (vertically aligned nanotube arrays, *contrast to fibers*) for H_2 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



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

1 Rapid computation of capacity

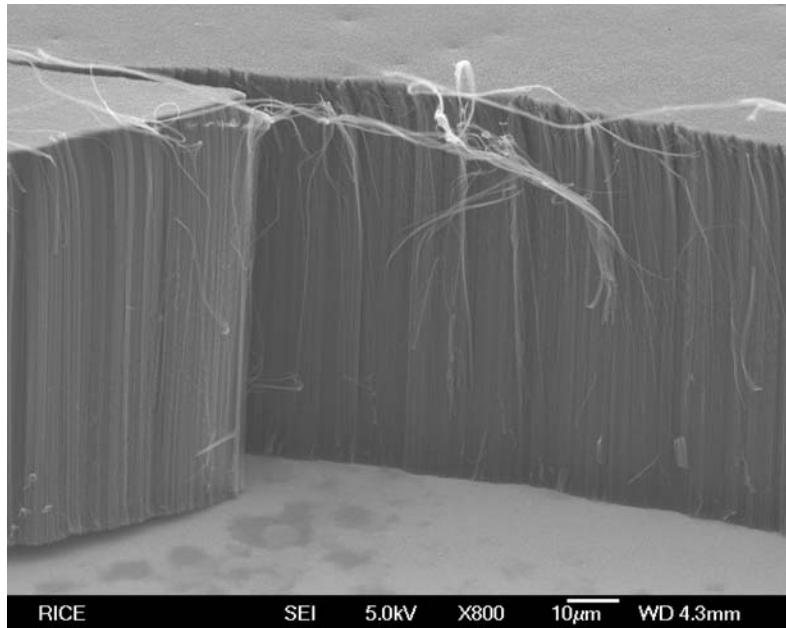
- via Grand Canonical Monte Carlo simulations for the structures of interest
- based on judicious choice of potential (of paramount importance)

- perform GCMC with **both** potentials to determine the range where the storage capacity is.

For technical accomplishment 2

2 Experimental

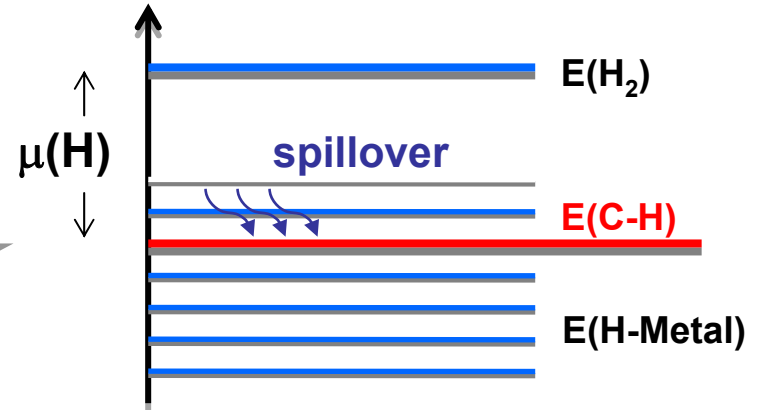
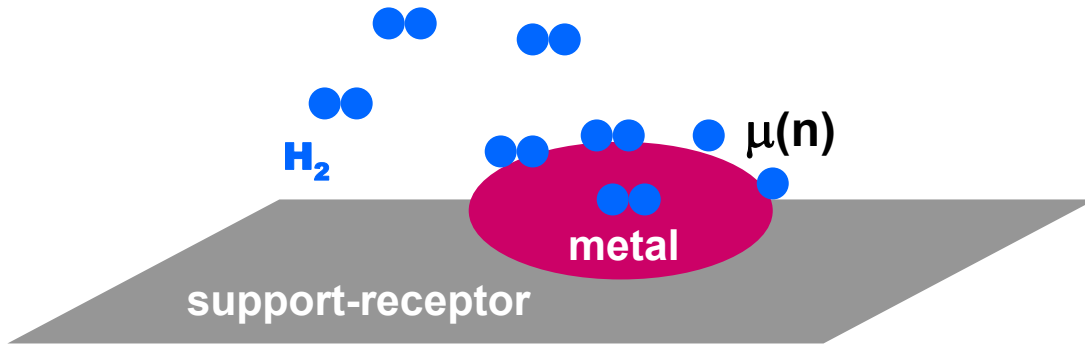
- VANTA arrays of 100 μm long carbon nanotubes, grown in large amounts with CVD methods. The alignment enhances gas transport into the array and thermal conduction.
- The aligned nature also lends itself to insertion of bulky anions, e.g. BF_3^-



APPROACH

Me-enhanced sorption on carbon receptors

3 Spillover thermodynamics, kinetic paths



$$\mu(\text{H}_2\text{-gas}) > \mu(\text{H@metal}) > \mu(\text{H@support})$$

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

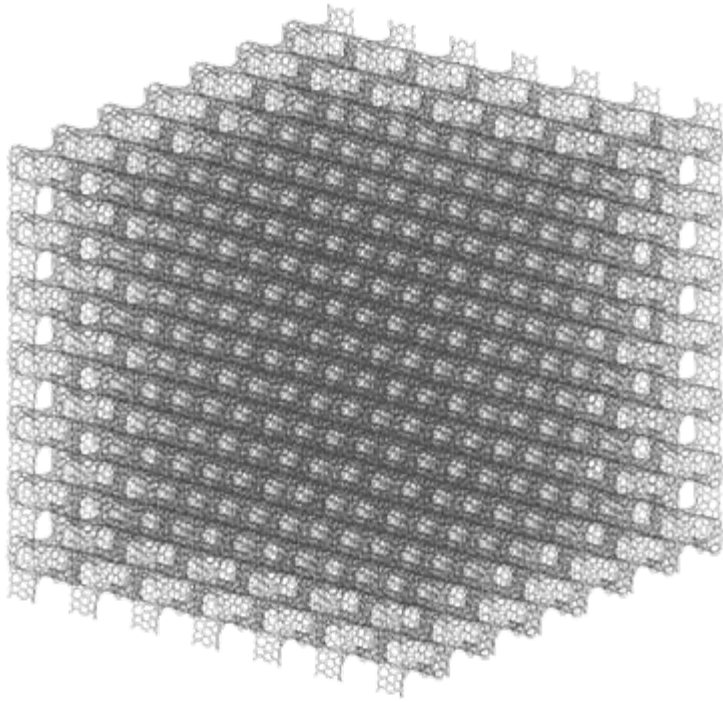
Role of bridges, defects and dopants on the overall process of spillover

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 H₂ storage

technical accomplishments 1



Ding et al JCP 127, 164703 (2007)

- Nanometer pores and channels
- Geometrical surface ~2600 m²/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

$$u(r) = 4\varepsilon[(\sigma/r)^{12} - (\sigma/r)^6] \quad u(r) = Ae^{-\alpha r} + C_6r^{-6}$$

$$\varepsilon = 3.7 \text{ eV/molecule}$$

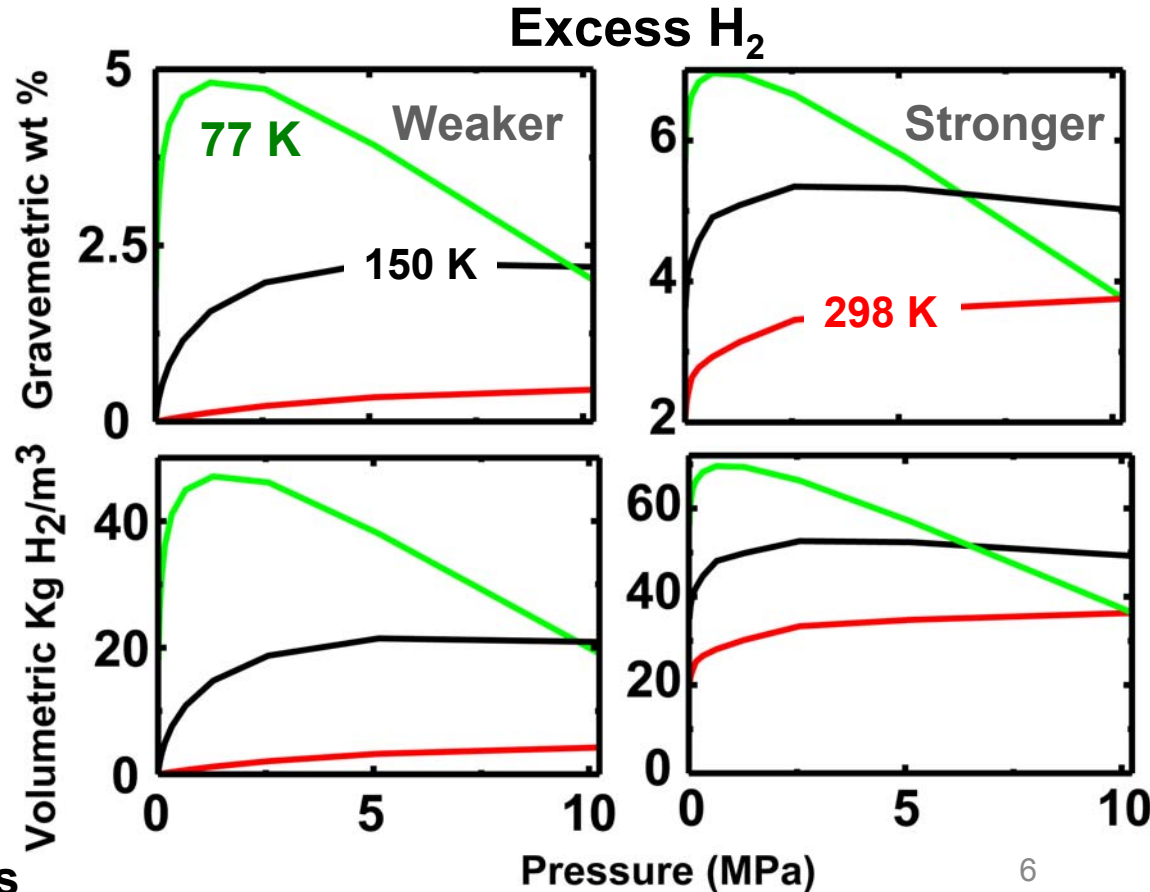
$$A = 1100 \text{ eV/molecule}$$

$$\sigma = 3.0 \text{ \AA}$$

$$C_6 = -17 \text{ eV \AA}^6/\text{molecule}$$

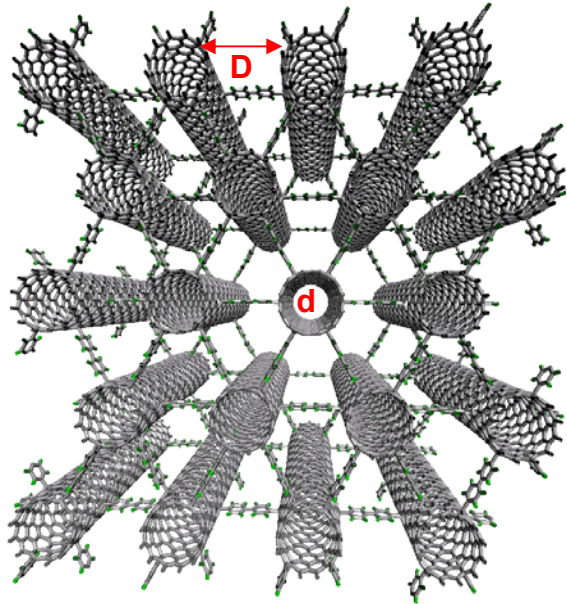
Wang et al J Low Temp Phys 41, 611

Patchkovski et al PNAS 102 10439



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

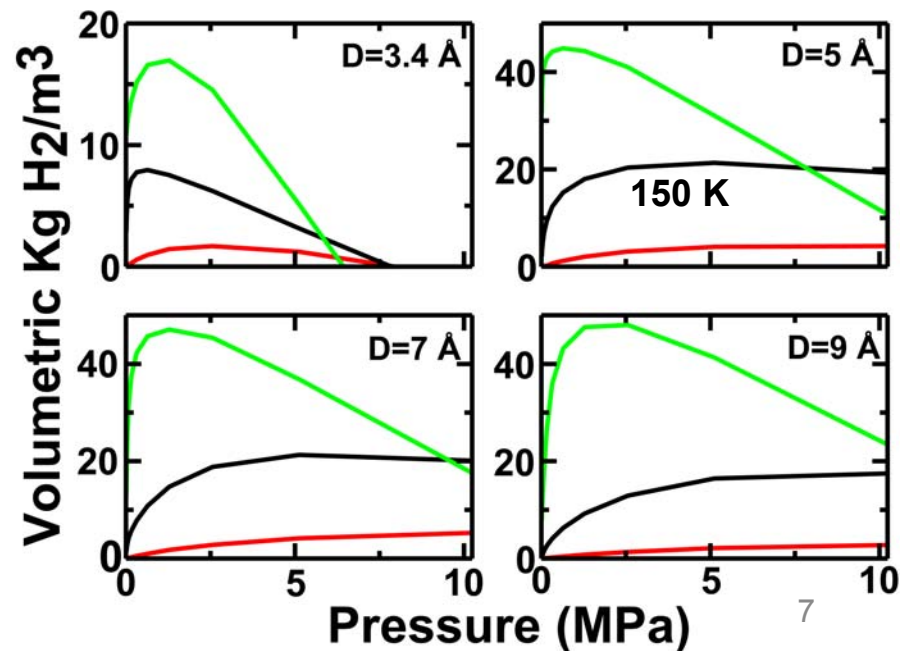
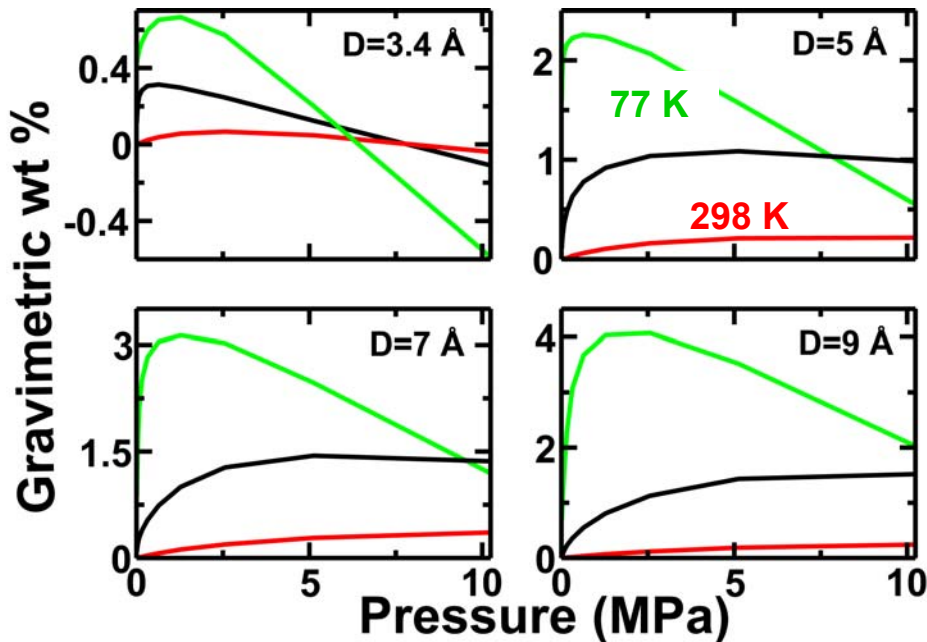
technical accomplishments 1



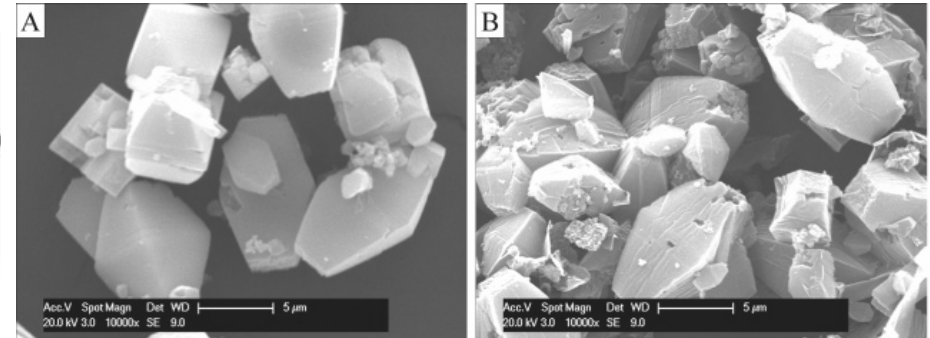
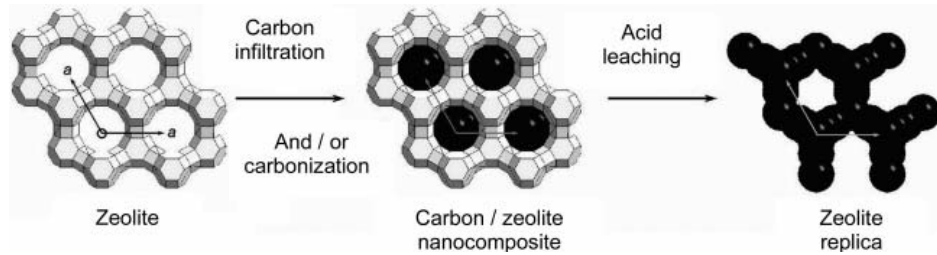
Storage capacity as function of wall-wall distance **D**
Volumetric % peaks at wall-wall distance of $D \sim 7 \text{ \AA}$
agrees with previous studies

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

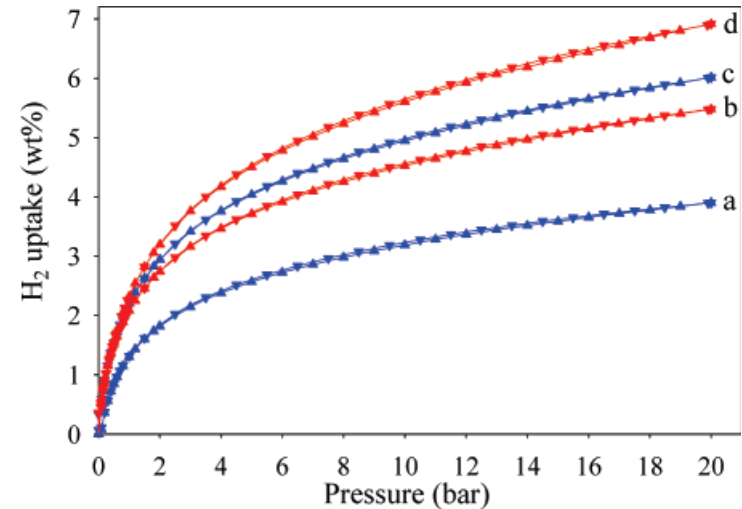
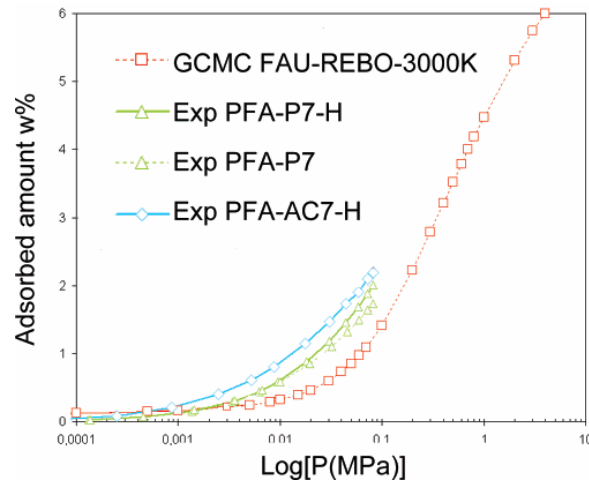
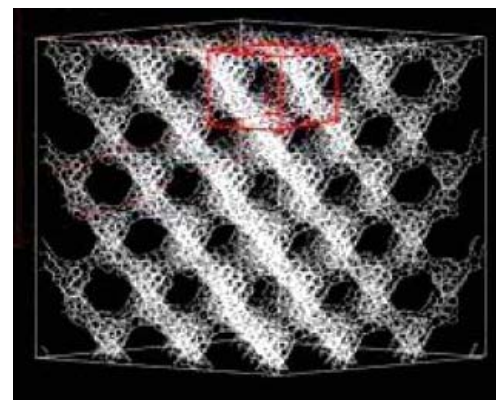
Excess H_2



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

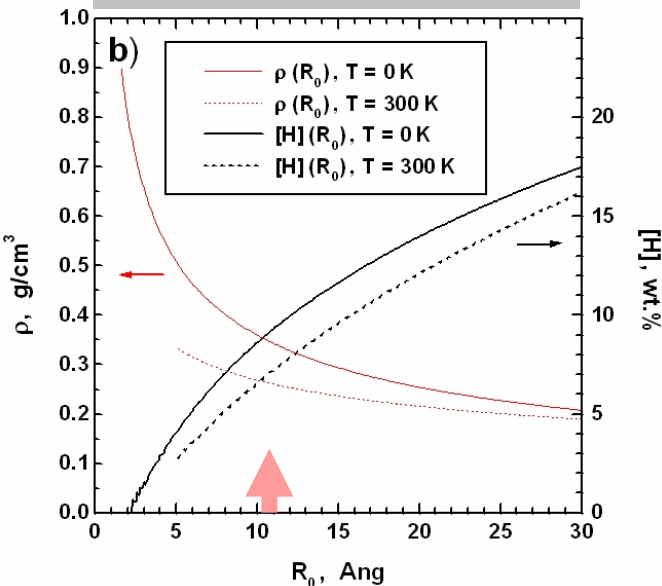
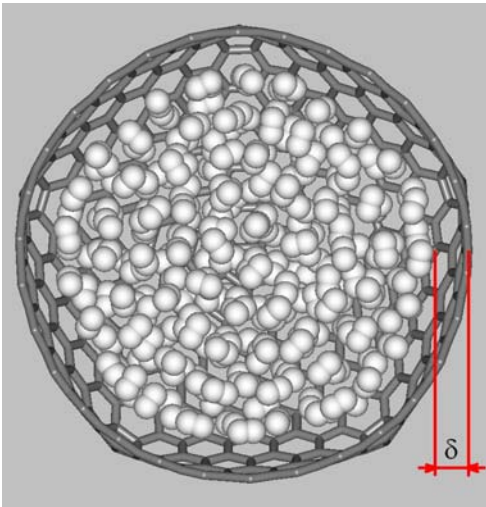
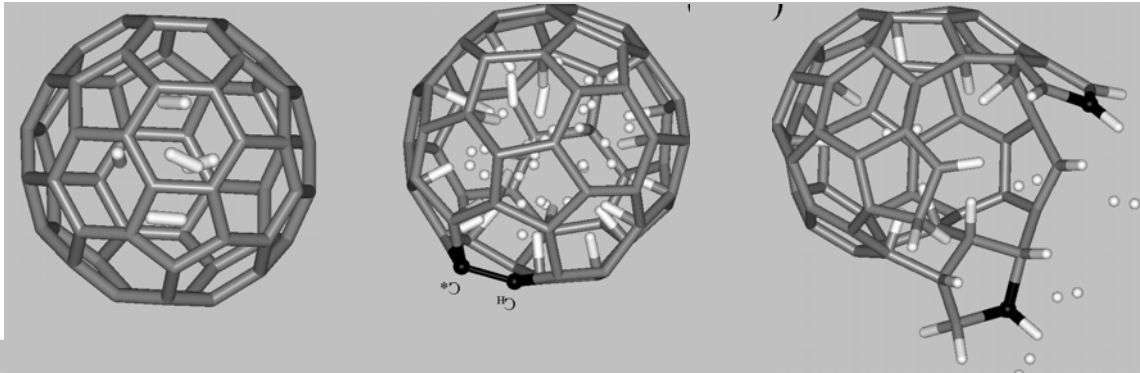


Hydrogen uptake at 77 K

Which is "ideal pore" ?

C_{60} interior attracts one H_2

What if we could utilize the *repulsion* for storage



Existing giant fullerenes as $\sim C_{1000}$ sustain great internal pressure, permitting H storage.

How it can be placed there remains yet unexplored

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NANO

LETTERS

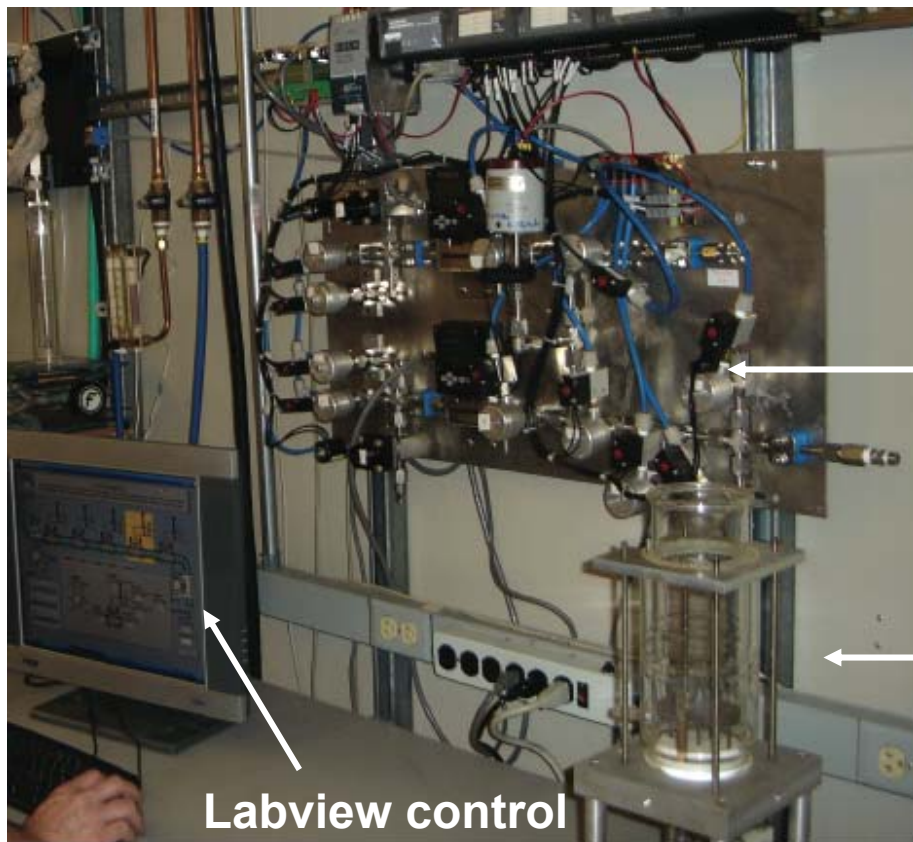
http://pubs.acs.org/NanoLett

VOLUME 8 NUMBER 3 MARCH 2008

Fullerene Nanocage Capacity for Hydrogen Storage

System for synthesis of metal- and electro-negative-group- (F^- , BF_3^-) enhanced VANTA

technical accomplishments 2



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

F₂ and BF₃
metering
controls

Reactor

Labview control

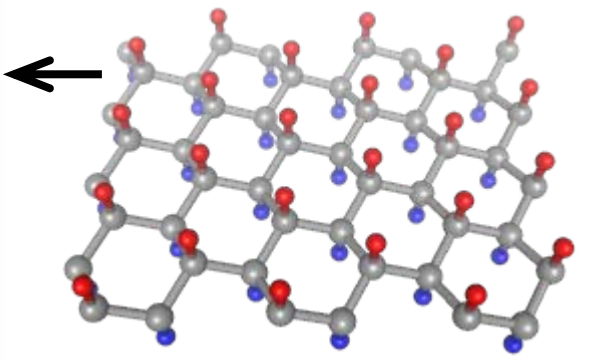
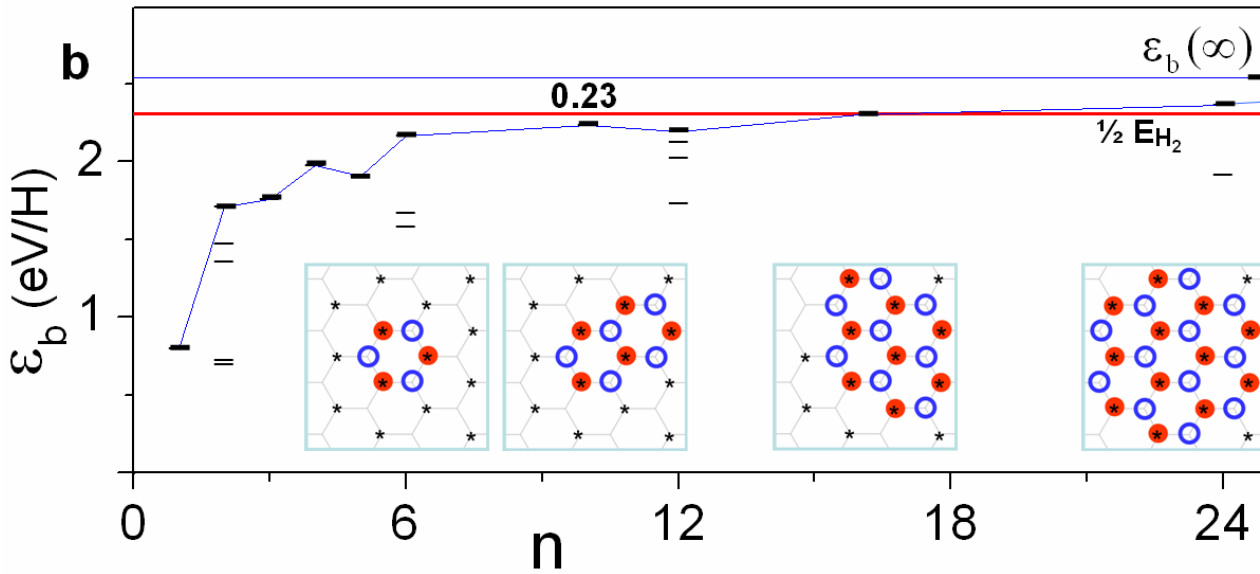
Parameters for best growth of VANTA scaffolds have been defined and VANTA are now available for addition of electronegative groups for enhanced H₂ adsorption.

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

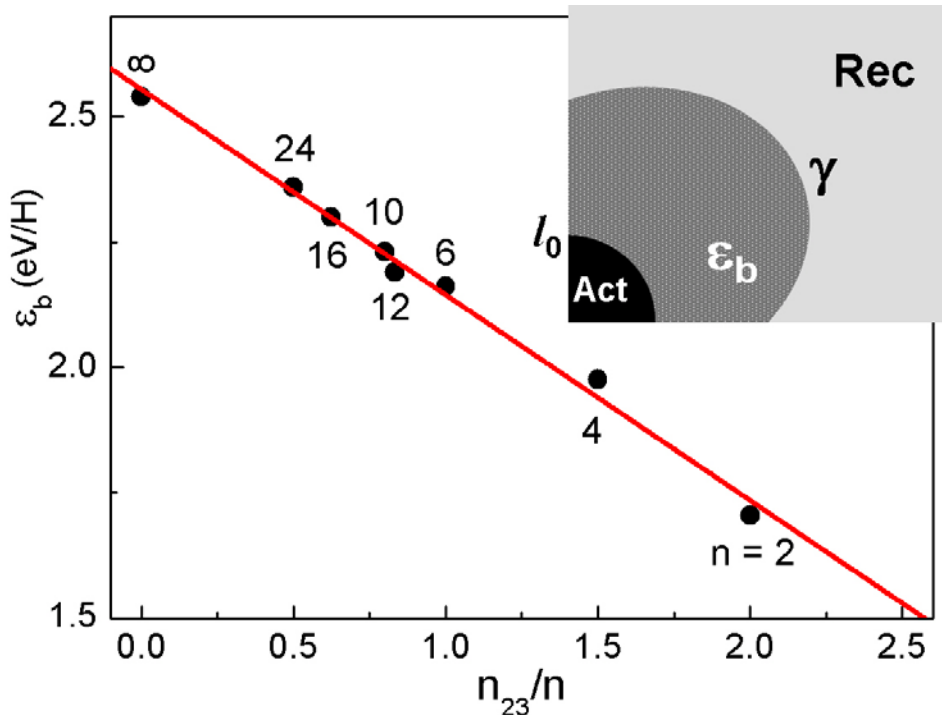
One gram samples have been converted to a carbon nanotube salt and will be tested for hydrogen uptake in collaboration with NREL and Air Products.

Spillover kinetics

technical accomplishments 3

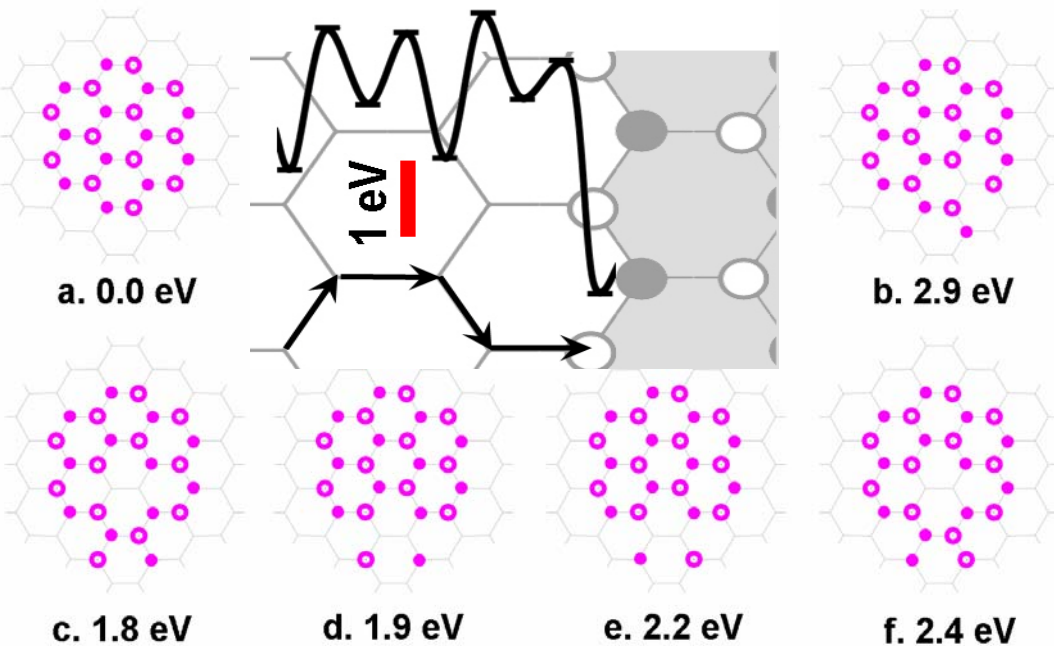
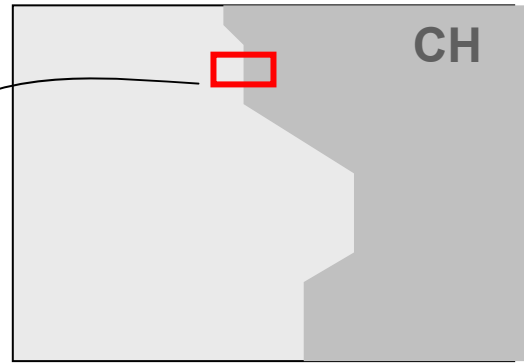


H tends to form even-numbered, compact, counter-layered clusters with H_2 energy crossover near $n = 20$



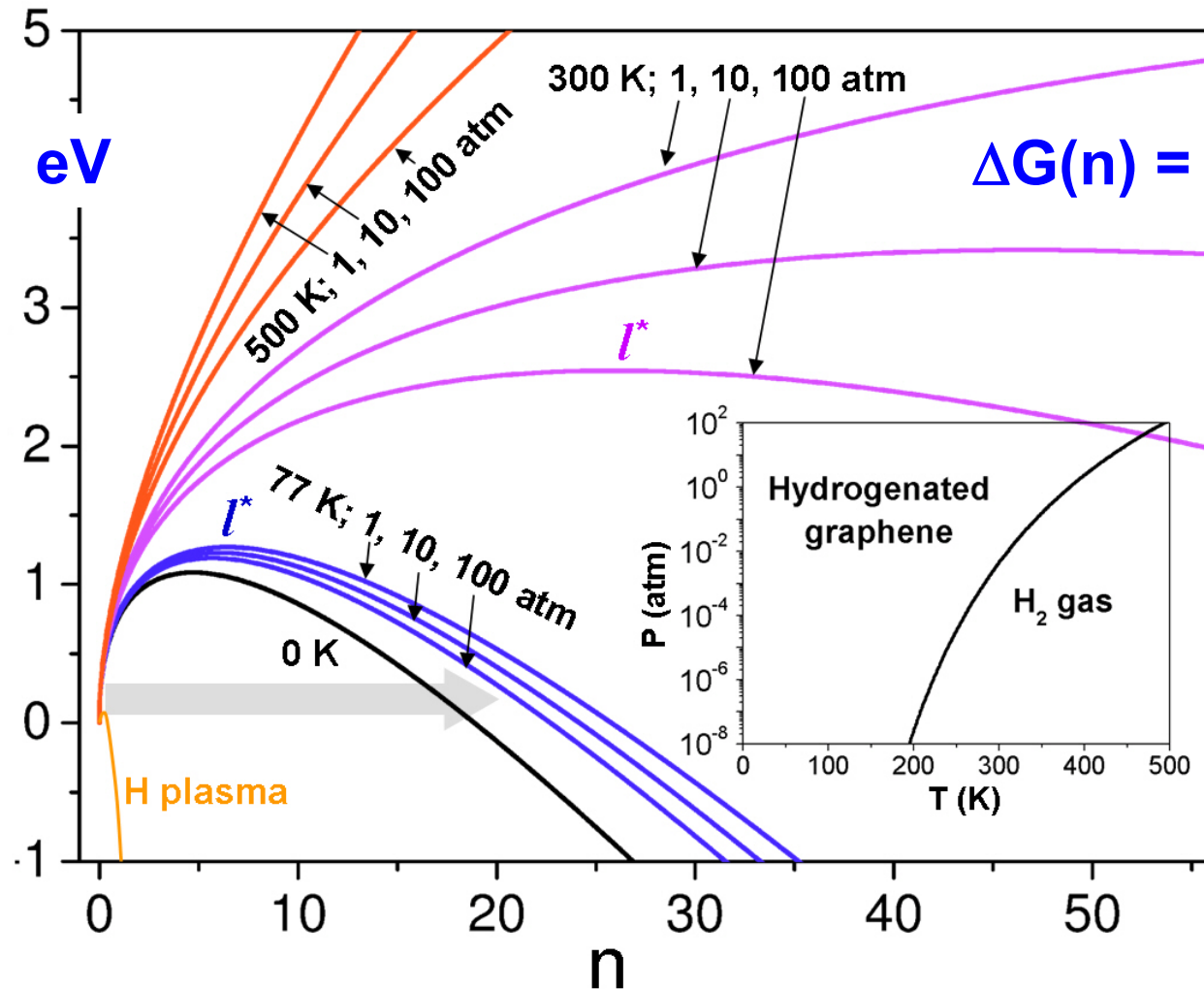
remarkably linear dependence
 $\epsilon_b(n) = \epsilon_b(\infty) - \gamma \cdot (n_{23}/n)$
supports the notion of hydrogenated phase nucleation, with effective 0.23 eV binding, proper for H_2 adsorption/desorption

Vacancy formation and front stability



- 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**

* calculations at T = 0

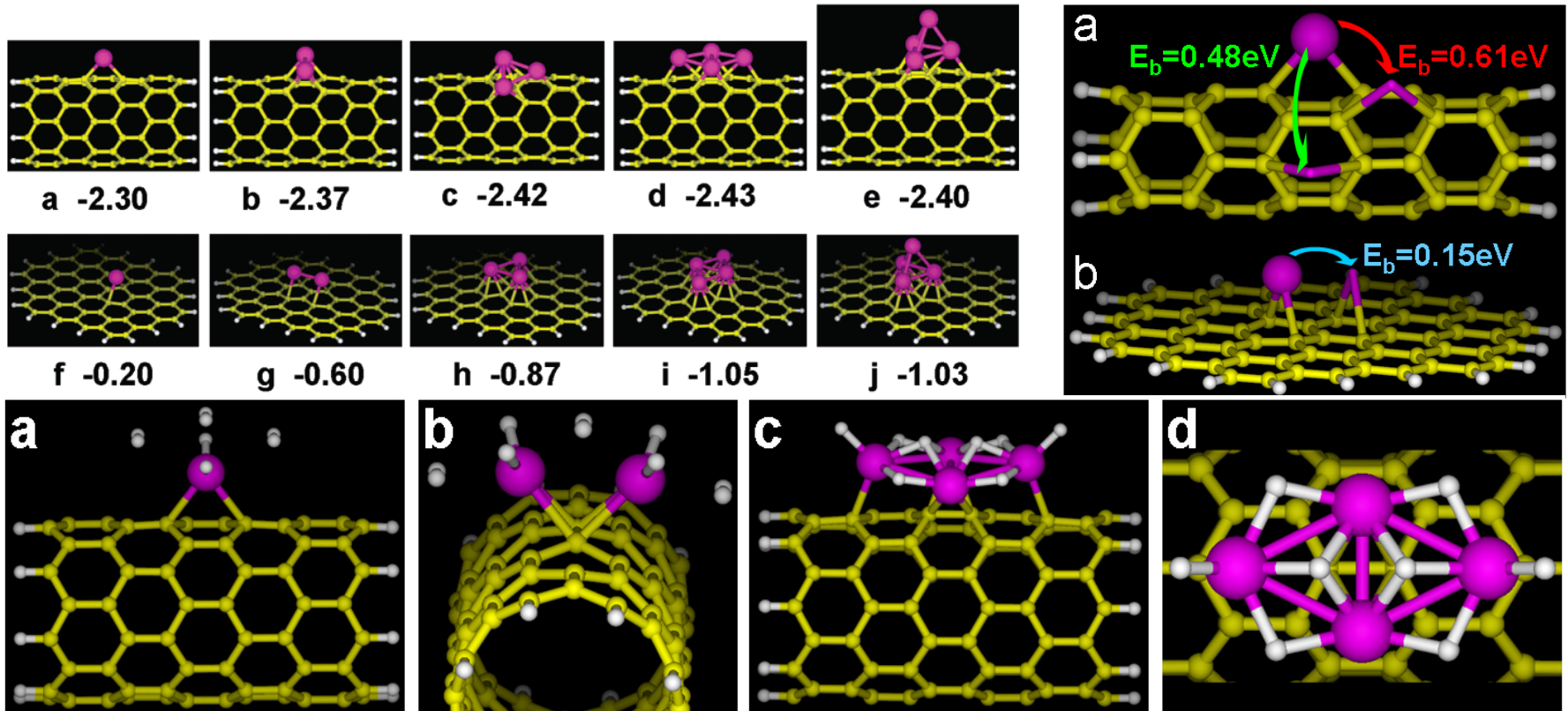


$$\Delta G(n) = -[\epsilon_b(\infty) + \mu_H] n + \gamma\sqrt{n}$$

Phase diagram of the chemisorbed H, may occur at normal conditions, e.g. 380 K, 1 atm

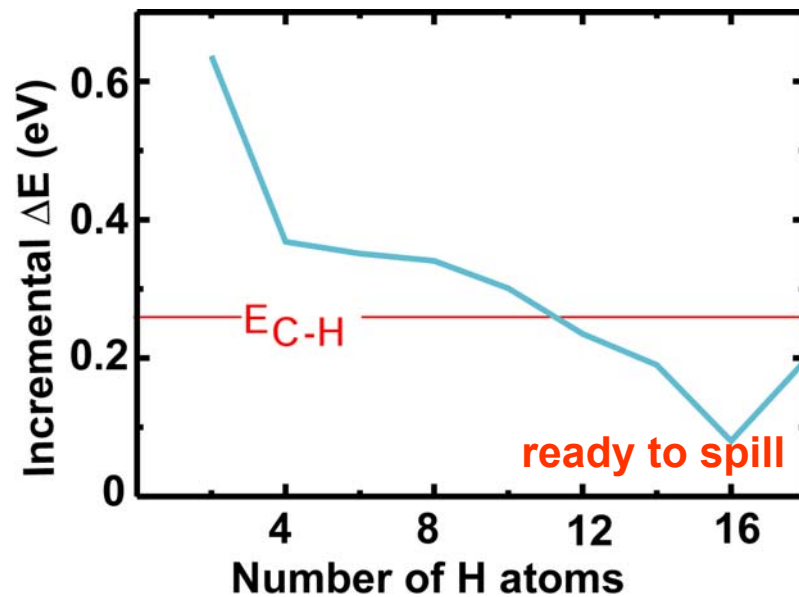
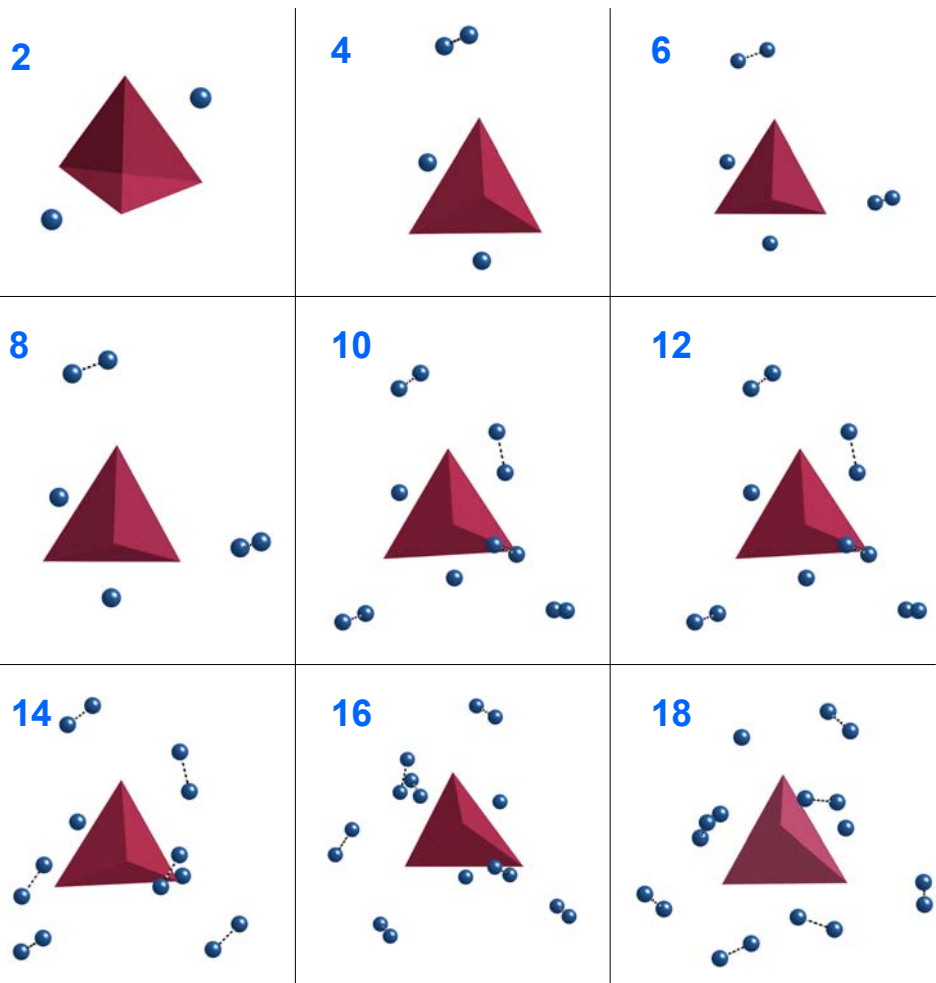
Nucleation barrier decreases with lower T and higher P
 Can be overcome by possibly “bridges” [R Yang] and the precursor H-patches [NREL work]

Metal clustering and effect on H₂ capacity



- Metal clustering is energetically and kinetically feasible at room temperature
- The diffusion frequency is $\sim 10^{12} \exp(-E_b/k_b T) \sim 10^5 - 10^{10} \text{ s}^{-1}$ 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

Determine the energy levels for H on free Pd₄ cluster and compare it with the onset of spillover onto C-receptor



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

- First H₂ gets adsorbed via dissociation, subsequent H₂ via Kubas type of interaction

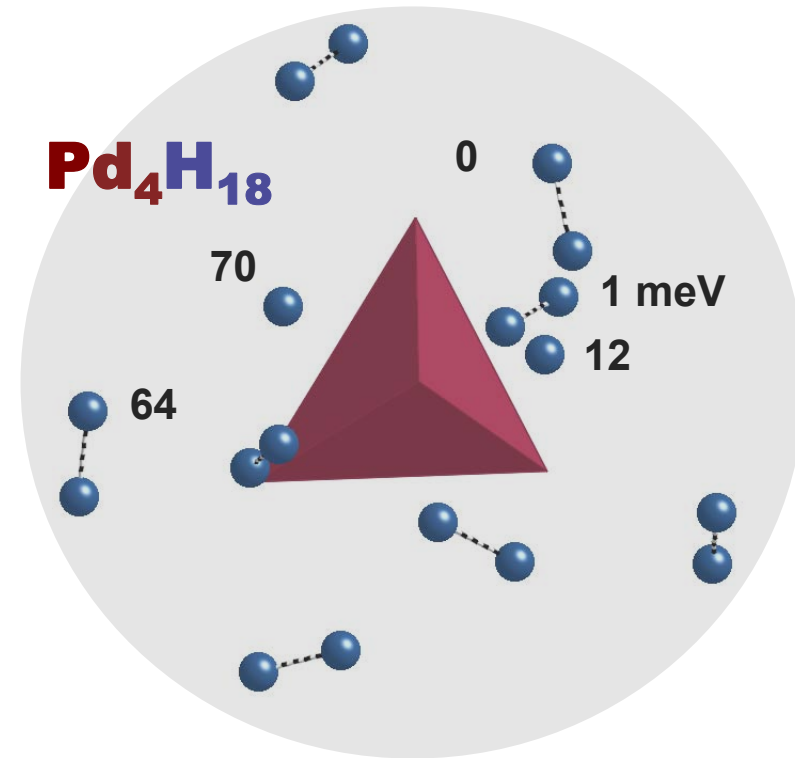
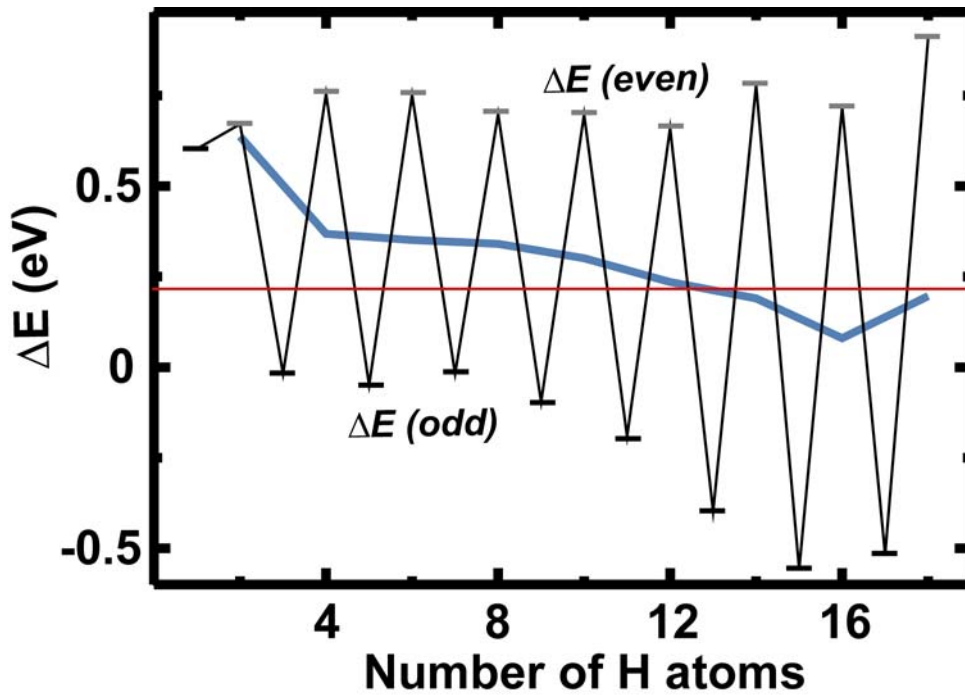
* calculations at 0 K, free cluster

Cluster saturates with 9H₂ (n = 18), spillover is favored after 6H₂ (n = 12)

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?

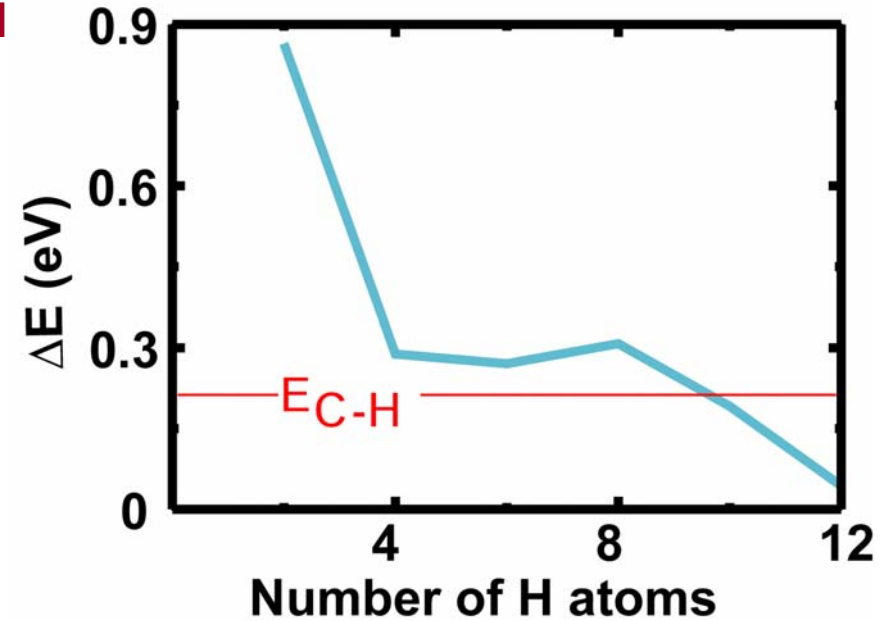
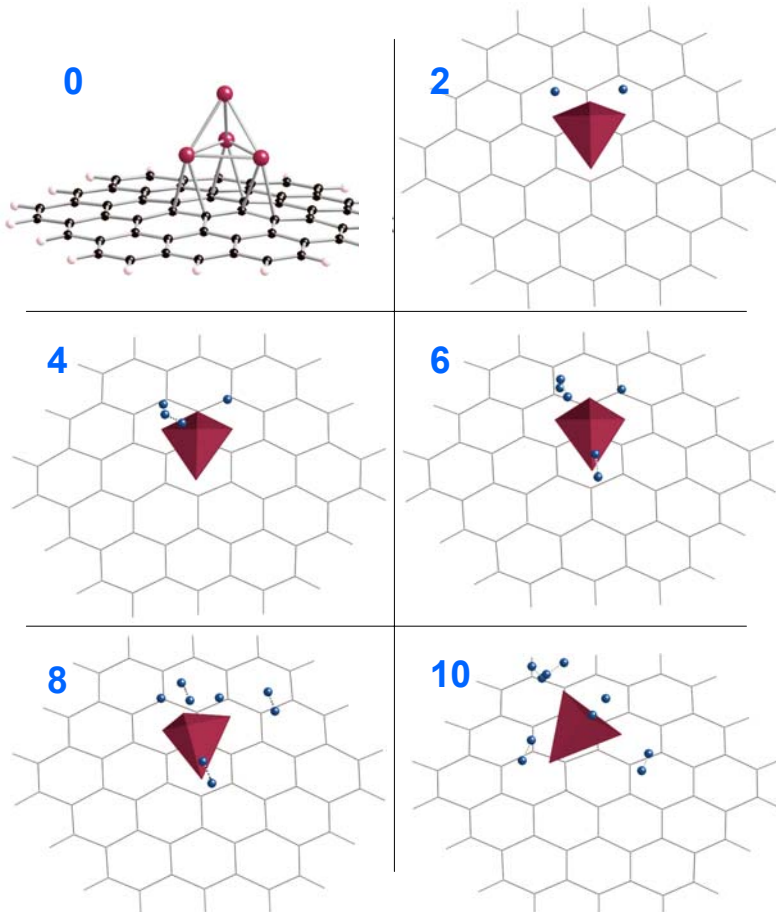


- Cost to remove dissociated (H) and non-dissociated (Kubas H-H) H is almost same

- 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 support on the **H** energy on **metal**

technical accomplishments 3



Spillover Threshold:

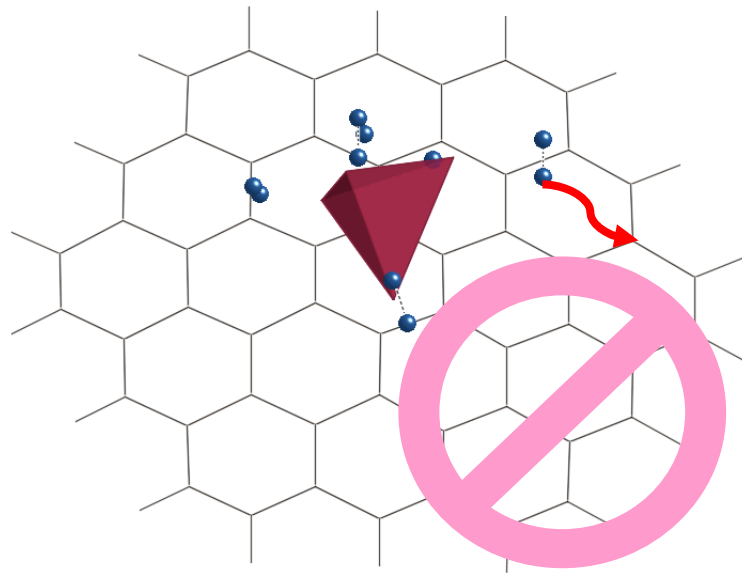
- On graphene metal saturates sooner than in the free cluster
- Onset of spillover is very close to metal full saturation

* calculations at 0 K, free cluster

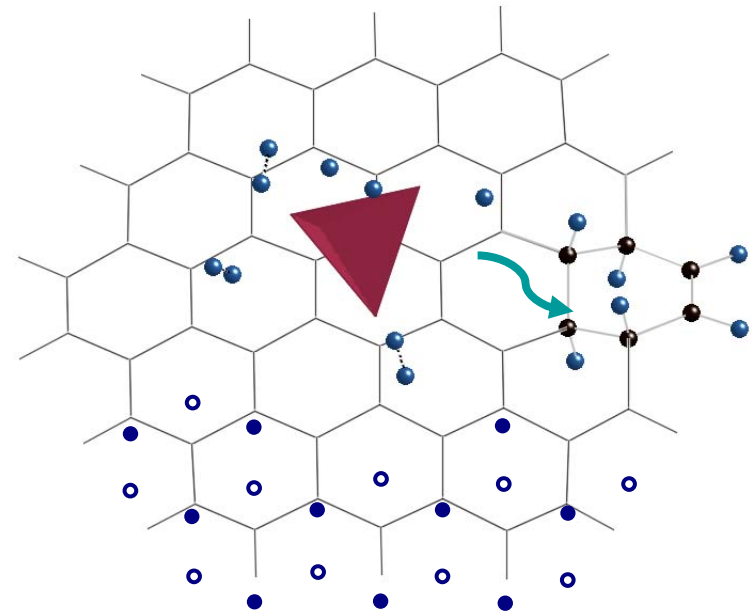
- First H₂ again gets adsorbed via dissociation, subsequent via Kubas
- The cluster saturates with 5H₂

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



- 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

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 front propagation model in contrast to conventional $\langle r^2 \rangle \sim D \times t$ 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 H₂ storage. Conversion of π -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.

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.