



A Synergistic Approach to the Development of New Hydrogen Storage Materials, Part I

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Overview

Timeline

- Project start: 12/1/04
- Project end: 11/30/09
- Percent complete: 70%

Budget

- Total funding expected: \$2.9M
 - \$1.8M from DOE to UC Berkeley
 - \$600k from DOE to LBNL
 - \$500k in cost-sharing
- Funding FY08: \$600k
- Funding FY09: \$567k

Barriers

- Identify new materials enabling a hydrogen storage system achieving:
 - 2 kWh/kg (6 wt %)
 - 1.5 kWh/L (0.045 kg/L)
 - 4 \$/kWh

Partners

- ChevronTexaco
- General Motors Corporation
- Electric Power Research
 Institute

Overall Program

Synthesis of porous polymers (Fréchet)

Synthesis of porous coordination solids (Long)

Calculations of H₂ binding energies (Head-Gordon)

Synthesis of destabilized hydrides (Richardson) Part I*

Part II

(BES)

H₂ storage characterization instrumentation (Mao)

Metal/metal hydride nanocrystals (Alivisatos)

Synthesis of nanostructured boron nitrides (Zettl)

Theory for boron nitride materials (Cohen and Louie)

H₂ Adsorption in a Hypercrosslinked Polymer



poly(chloromethylstyrene-codivinylbenzene)

Hypercrosslinked Polyaniline



Size Exclusion of Gases in Ultrananopores



Nanoporous Nitrogen-Containing Polymers



Crosslinked aromatic rings

Reagent	Surf. area, m ² /g	Pore volume, mL/g		
	Langmuir	Total	Nano	Ultra
Diiodobenzene	192	0.01	0.01	0.09
Tribromobenzene	384	0.13	0.11	0.06

Stronger H₂ Adsorption



Hypercrosslinked Polypyrrole

 Cs_2CO_3

DMSO

Crosslinking with alkyl groups:





Crosslinking with boron:





Size Exclusion of Gases in Ultrananopores



Hydrogen Storage in Completely-Activated MOF-5



- Currently best known material for cryogenic hydrogen storage at 77 K
- Performance at 298 K is poor owing to weak interaction of H₂ with surface

A Beryllium-Based Metal-Organic Framework



- Unprecedented structure with Be₁₂(OH)₁₂ rings and 12 and 15 Å channels
- Nitrogen adsorption isotherm affords BET surface area of 4020 m²/g

Low-Pressure H₂ Uptake in Be₁₂(OH)₁₂(BTB)₄



- Weak interaction of H₂ with surface, as desired for cryogenic storage
- At pressures up to 100 bar, expect gravimetric storage above MOF-5

Strong H₂ Binding in MOF-5 Functionalized with Cr⁰



- Orbital interactions lead to strongly-bound H₂ complex that is too stable
- Need to generate charge-induced dipole interaction of 15-20 kJ/mol

Room Temperature H₂ Uptake in Mn-BTT*



- Exposed Mn²⁺ sites lead to isosteric heat of adsorption of up to 10.1 kJ/mol
- Need to increase strength of binding and concentration of open metal sites

Paddlewheel Frameworks



 $M_3(BTC)_2$ (M = Cr, Cu, Zn, Mo)

	surface area (m²/g)		
	BET	Langmuir	
Cr ₃ (BTC) ₂	2340	2720	
Cu ₃ (BTC) ₂ ^a	1944	2260	
Zn ₃ (BTC) ₂	collapsed		
Mo ₃ (BTC) ₂ ^b	1280	2010	
Mo ₃ (BTC) ₂	1800	2100	

^a J. Am. Chem. Soc. **2006**, 128, 3494 ^b J. Mater. Chem. **2006**, 16, 2245

- Preparation of Cr₃(BTC)₂ is new and activation of Mo₃(BTC)₂ is improved
- Enables comparison of H₂ binding at the open M^{II} coordination sites

H_2 Uptake in $M_3(BTC)_2$ (M = Cr, Mo)



- First assessment of strength of H₂ binding to a Cr²⁺ center
- Expect better results for Co²⁺ and Ni²⁺ owing to a smaller ionic radius
- Attempts to synthesize analogues with other metal ions are underway

H₂ Uptake in Mg₂(DOBDC)



Matzger et al. J. Am. Chem. Soc. 2008, 130, 10870

- Open Mg²⁺ sites lead to an isosteric heat of adsorption as high as 12.8 kJ/mol
- Neutron diffraction (Craig Brown, NIST) shows Mg···D₂ distance of 2.5 Å

Calculation of Substituent Effects

Metal chosen as Cr⁰



Cr

R

Effect of Substituent (R)



 Electron-donating groups enhance binding, while electron-withdrawing groups reduce binding

-Tunability is 7% of binding

–Energies are for three bound H₂ molecules

- Correlates with back-donation, electrostatics
- Quantitative information; qualitative insight

-BDC²⁻ substituents can fine-tune binding

-Coarse-tuning must come from different metals

Effect of Metal Substitution



• Heavier isoelectronic elements:

 $(C_6H_6)Cr(H_2)_3$ binding per H2 of 68 kJ/mol $(C_6H_6)Mo(H_2)_3$ binding per H2 of 84 kJ/mol

• Lighter transition elements:

 $(C_6H_6)Cr(H_2)_3$ binding per H2 of 68 kJ/mol $(C_6H_6)Ti(H_2)_4$ binding per H2 of 32 kJ/mol

• Shows coarse tuning is possible

Still need to examine synergy of these effects

Computational Study of H₂ Binding in Cu-BTT



[Cu₄X(N₄CH)₈]⁻ fragment

ωB97X-D/6-31G* calculations:

Х	E/kJ/mol	
F	-10.9	
CI	-10.9	
Br	-13.0	
Ι		

- Measurements of H₂ binding energy within HCu[(Cu₄Cl)₃(BTT)₈] underway
- We will attempt to synthesize HCu[(Cu₄Br)₃(BTT)₈]

Computational Study of H₂ Binding in "Zn-BTT"



[Zn₄X(N₄CH)₈]⁻ fragment

ωB97X-D/6-31G* calculations:

Х	E _{Cu} /kJ/mol	E _{zn} /kJ/mol
F	-10.9	-13.8
CI	-10.9	-15.9
Br	-13.0	-16.3
I		

- Suggests significant improvement in binding energy for Zn-BTT frameworks
- We will therefore attempt to synthesize $Zn_3[(Zn_4Cl)_3(BTT)_8]_2$

Destabilization of Metal Hydrides



- Attempts at alloying of Mg in order to reduce ΔH
- Success in partial substitution to form $Mg_{1-x}A_x$ (A = Mn, Fe, Ni)
- Some increases in plateau pressures, but poor kinetics
- Attempts to substitute Na and Li for Mg are underway

Addition of MgF₂ Enhances Utilization of MgH₂



- MgF₂ slows desorption, but increases amount desorbed despite added weight
- Fluoride is distributed over particle surface; no evidence for bulk substitution (XRD)

Fluoride Effect Persists through Repeated Cycling





MgH₂ + 3 mol% MgF₂ after 2nd desorption @ 300° C

Without fluoride, sintering and coarsening reduce surface area and contribute to Mg isolation High-resolution TEM shows sharp faceting and marked inhibition of Mg grain growth in fluoridecontaining samples.



Best results are for 3 mol% MgF₂ added

 $\rm MgH_2$ after 2nd desorption @ 300° C

Metal Catalyst Activity not Inhibited by Fluoride



Activity of added vanadium not inhibited by the presence of fluoride

Good utilization at 250 °C

Still below 1 wt % at 200 °C

5wt .% V+ MgH₂ milled for 2hrs(300 rpm) 1 mol % MgF₂ added (600 rpm for 1hr)

- Future work: higher energy milling and alternative fluoride sources
- In addition, the effect of fluoride addition on ternary hydrides will be studied