## Sub-Nanostructured Non-Transition Metal Complex Grids for Hydrogen Storage

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Project ID # ST7

This presentation does not contain any proprietary or confidential information

# Overview

#### Timeline

- Project start date: 1/20/04 (contractual) April 2004 (actual)
- Project end date: 1/19/07
- Percent Completed: 30 %

#### Budget

- Project funding: \$1,201,862
  - DOE share: \$ 959,317
  - Contractor share: \$242,545
- Funding received in FY04: \$310,407
- Funding for FY05: \$ 190,000

# Overview (con.)

- DOE On-Board Hydrogen Storage Barriers
  - -C. Efficiency
  - -D. Durability
  - -E. Refueling Time
  - -M. Hydrogen Capacity and Reversibility
  - -N. Lack of Understanding of Hydrogen Physisorption and Chemisorption

#### Partners

- Steven M. Kuznicki, U. of Alberta; zeolite chemistry
- Suha Yazici, Graftec; electrochemistry
- David Sholl, Carnegie Mellon U.; computational (pending)

## OBJECTIVE

- To develop the techniques to make nanostructured continuous threedimensional metal alloy grid for hydrogen storage
- To make and characterize the grids for state-of-art alloys

# **TECHNICAL APPROACH**

- 1. Use nanoporous (e.g. zeolite) templates as mold on cathode
- 2. Electrodeposit metals in the pores to grow nanostructured grids
- 3. Dissolve the mold and recover the metal grid

## WHY NANOSTRUCTURE ?

► Large surface area <u>increase H<sub>2</sub> dissociation rate</u>

Smaller diffusion distance increase H-transport rate in metal

≻Flexible/open structure <u>decrease decrepitation by cycling</u>

Energy release/absorption & mass/energy transfer throughout matrix <u>improve heat transfer characteristics</u>

≻Large surface area increase storage capacity by physical adsorption

➢ Possible Increase in capacity by quantum effects ? (30-50 atom clusters)

#### Increase dynamic hydrogen uptake !

# Example: Mg grown in faujisite



Cubic superlattice; a=2.4 nm

Characteristic length (pore or solid); L = 1nm

Porosity;  $\epsilon = 0.5$ 

Surface area;  $\tau$  = 500 m<sup>2</sup>/gm

## **Expected improvements**

Base case: 1 mm solid metal particle

	Comment		Increase	
H <sub>2</sub> dissociation rate	scales with 1/L		10 <sup>6</sup>	
H diffusion rate	scales v	s with ε/L (		).5x10 <sup>6</sup>
Energy flux in metal	k(1-ε)δ <sup>2</sup> T/δx <sup>2</sup> conduction <b>lower</b> ε	(εDδC/δx)*C (mass flux)*e additional,	₀δ∆T/δx enthalpy small	r <sup>s</sup> (∆H+C <sub>p</sub> ∆T) <i>reaction</i> additional, large
Physical adsorption	Scales with 1/L		10 <sup>6</sup>	

## **ISSUES/QUESTIONS**

• Volumetric capacity reduced by a factor of  $\epsilon$  compared to bulk metal

• Will the metal sinter during cycling?

# TASKS

#### YEAR: 1 2 3

- 1. Mold/Cathode preparation
- 2. Pure metal electrodeposition *characterizations*
- 3. Mold removal, grid recovery
- 4. Pure metal hydrogen storage
- 5. Mixed metal electrodeposition *characterizations*
- 6. Mixed metal hydrogen storage



## TASK 1: Mold/Cathode Fabrication

#### Least Attractive

- I. <u>Multicrystal</u> particles <u>compacted</u> on a metallic substrate.
- 2. <u>Multicrystal</u> film of <u>inter-</u> <u>grown</u> crystals in-situ synthesized on a metallic substrate.



- 3. <u>Multicrystal</u> film containing large <u>oriented</u> crystals <u>grown</u> on a metallic substrate.
- 4. Very large single nanoporous crystal grown on a metallic substrate ?



Most Attractive

### Electrodeposition in compacted powder



		IN CREVICE		
С	31 at%	C	45 at%	
0	43	0	30.4	
Cu	4	Cu	2.3	
Al	9.9	Al	9.5	
Si	12.6	Si	12.8	



## **TASK 1.1 Compaction Summary**

•Faujisite powder (Si/Al=1.4, 1  $\mu$ m particle) exchanged to Cu compacted on Ni mesh used as cathode

•Cu electrodeposited in the pores

•EDX shows Cu enrichment in zeolite crystals

•Cu/Al ratios: 0.51 left (closest to anode), 0.4 top, 0.35 right
•excessive Cu growth in between zeolite particles (like tree roots in stony soil)

# electrodeposition in zeolite particles works hard to prevent growth in between particles

## Anodized Ti-cathode for in-situ growth



- NaOH treatment at 60 C, cure at 500 C
- $TiO_2$  is formed (anatase)
- Ti metal is also exposed
- <u>Texture 0.1 to 10 μm, controlled by</u> treatment conditions (0.1 μm shown)



- Purple =  $TiO_2$  (for adhesion of zeolite)
- Blue = Ti metal (<u>for electrical</u> <u>conductivity</u>)

## Faujisite growths on anodized-Ti



Same chemistry, different magnification

#### Two other chemistry's



picked the best adhesion chemistry for electrodeposition

#### **Electrodeposition in synthesized zeolite film**



## Task 1.2 Anodized Ti + Faujisite + Cu-deposition

- Metal (Ti) is microstructured to TiO<sub>2</sub> to promote zeolite film growth with good adhesion
- Cu is electrodeposited in zeolite pores

- > No Cu growth between particles
- "mushroom" overgrowth will be prevented
- > Need to grow thicker films
- Try also anodized aluminum

## **TASK 2: Electrodeposition**

•Vast electroplating/electrodeposition literature is primarily geared to optimize the <u>uniformity<sup>1</sup></u> of <u>thin</u>  $\underline{\text{films}^2}$  on <u>large smooth<sup>3</sup></u> cathode surfaces of a <u>different<sup>4</sup></u> metal.

•We are trying to deposit <u>non-uniform<sup>1</sup></u>, "<u>thick</u>" <u>bulk<sup>2</sup></u> metal in <u>small nanopores<sup>3</sup> of a non-conducting</u> material. (The cathode/anode conductor can be the <u>same<sup>4</sup></u> metal.)

# Need to fine-tune electrodeposition at nano-scale.

# Electrodeposition in track-etched polycarbonate films (PCF)

Commercially available membranes used as cathode mold
Non-conducting polycarbonate "mold" (like zeolites)

- •Pore sizes from 1  $\mu$ m, down to 0.015  $\mu$ m = 15 nm (150x larger than zeolite pores, <u>hard to scale</u>)
- •Film thickness up to 10  $\mu$ m (zeolite crystal size, <u>close</u> scale) •Porosity up to 10% (less than zeolites, <u>easy</u> to scale)

Analyze effects of electrolyte chemistry, deposition conditions, and mold characteristics for electrodeposition at nanoscale

## Cu deposit in 10 $\mu\text{m}$ holes of PCF



Solution side at 1,000x

Caps coalesce, diameter closer to 10  $\mu m$ 



Cathode conductor side at 1,000x

No growth on Au coating

Holes in center of deposited Cu suggest inwards growth from walls

## Task 2: Electrodeposition fine tuning

PCF as "proxy" to fine tune electrodeposition parameters
Identified controlling radial versus axial growth
Significant effect of pore size

#### Future

Extrapolate to nanoscale electro deposition (below 15 nm)
 Bring-in H<sub>2</sub> hydride electrochemistry



#### HRTEM images of faujisite after Cu electrodeposition





- •1 nm lines spaced at 1 nm
- •No such lines in normal faujisite images

(Courtesy of Hitachi High Technologies America)

### RESULTS

HRTEM of Cu "mushrooms" grown from nanometer pores



MCM\_005.tif MCM Print Mag: 126000x @7.0 in 10:47 03/31/05

100 nm HV=100kV Direct Mag: 50000x AMT Camera System

## Next Year Activities

#### Remainder FY05:

- Multicrystal thicker zeolite grown on metals (Ti and Al)
- Oriented large crystal growth of nanoporous materials on cathodes
- Extrapolation of size effect on electrodeposition
- Switch from Cu to Pd (and Mg?) electrochemistry
- Characterization by HRTEM, STEM, AFM/STM, EDX

# Next Year Activities (Con.)

#### <u>FY06:</u>

- Mold removal, grid recovery and surface area measurements
- Pure palladium (and magnesium?) for hydrogen storage testing
- Mixed metal electrodeposition chemistry

# **Auxiliary Slides**

- Publications and Presentations
- 2 papers under review, 1 presentation scheduled for AIChE Annual meeting in Cincinnati, OH (Nov. 2005)
- The most significant hydrogen hazard associated with this project is:
- The experiments to date did not require hydrogen. Limited uptake rate and equilibrium measurements will be performed in FY06 in our Rubotherm © magnetic suspension balance using small/lab quantities of H2 vented into fume-hood.
- Our approach to deal with this hazard is: (Not applicable)