

# Hydrogen Storage through Nanostructured Porous Organic Polymers (POPs)

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## Overview

## Timeline

- Project start: July 2007
- Project end: October 2011
- % complete: 75%

## **Budget**

- Total project funding: \$2 Million
  - DOE share: \$1.88 Million
  - Contractor share: \$120 K
- Funding received in FY2010
  - \$500 K
- Funding for FY2011
  - \$400 K

## **Barriers**

- Barriers addressed
  - A. System Weight and Volume
  - B. System cost
  - C. Efficiency
  - D. Durability/Operability

## **Partners**

- Interactions/collaborations
  - Argonne National Laboratory (Lead)
  - U of Chicago (Subcontractor)
  - HSCoE Members
    - NREL
    - UNC
  - Non HSCoE Members
    - U of Hawaii (Sample exchange)
    - GM (Sample exchange)

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## **Objective - Relevance**

- To design, synthesize, and evaluate nanostructured porous organic polymers (POPs) as new hydrogen storage adsorbents for transportation applications
- To support polymer materials development with modeling/simulation and advanced structural characterizations

# Potential Advantages of POP H<sub>2</sub> Adsorbent & Their Impact on Technology Barriers

- System Weight and Volume POPs are light weight and can be converted to high volumetric density by engineering process such as compression, pelletizing, etc.
- System Cost POPs can be scaled-up for industrial production with the existing infrastructure at competitive cost.
- Efficiency POP-H<sub>2</sub> interaction is based on physi-adsorption/desorption principle with minimum parasitic energy consumption.
- Durability/Operability POPs are stable under the temperature and humidity conditions required for hydrogen storage application.

## **Approach - Milestones**

Month/ Year	Milestones	Status Update			
11/10	Complete the design, synthesis, and characterization of B-doped POPs	<b>80% Completed</b> . Surface properties, $H_2$ storage capacities and $\Delta H_{ads}$ were Investigated for three new three B-doped POPs. Finding is accepted for publication.			
02/11	Complete the synthesis of an ultra-high surface area aromatic POPs	<b>Completed</b> . Duplicated a literature report on porous aromatic framework as benchmark comparison for POP capacity and heat of adsorption investigation.			
05/11	Complete adsorption kinetics and charging time investigation for selected POPs	<b>Completed</b> . Adsorption/desorption kinetics of two representative POPs were studied.			
08/11	Complete H <sub>2</sub> storage capacity and heat of adsorption optimization through design/synthesis of new metal-doped POPs	<b>60% completed</b> . Seven new transition metal doped POPs were prepared. Measurement of surface property, $H_2$ adsorption uptake and isosteric heat of adsorption 50% finished.			

The focus of FY2011 is to improve and understand the key factor for isosteric heat of adsorption targeting for ambient temperature storage application

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## Approach - Development Strategy

	New Polymer Exploration (UofC/ANL)		Characterization & Optimization (ANL)		Modeling & Simulation (ANL)
•	New POP synthesis through rational design at molecular level	•	H <sub>2</sub> storage capacity & heat of adsorption measurement	•	H <sub>2</sub> -POP interaction study via <i>ab</i> <i>initio</i> , DFTB & MD methods
ŀ	Structure characterization	•	Surface property characterization	•	Advanced characterization
•	Post synthesis modification	•	Engineering process development		

- Prepare high surface area & narrow/adjustable pore size polymers through rational design and synthesis
- Incorporate "metallic" feature into polymer through conductive backbone or metal doping
- Improve POP-H<sub>2</sub> interaction by incorporating heteroaromatic functional groups
- Develop fundamental understanding through modeling and advanced characterization
- Over 100 POPs in three categories, aromatic, heteroaromatic and metal doped systems with high surface areas and narrow pore size distribution were designed and prepared
- Hydrogen uptake capacities of 0.055 (kg<sub>H2</sub>/kg<sub>ads</sub>) and 0.022 (kg<sub>H2</sub>/L<sub>ads</sub>), and the isosteric heat of adsorption of ~10 kJ/mol were achieved
- The correlations between surface property, H<sub>2</sub> storage capacity and adsorption enthalpy were found through combined experiment/simulation effort for better understanding on H<sub>2</sub>-POP interaction

### Technical Approach - Summary on Hydrogen Storage via Aromatic POPs



- Over 50 aromatic POPs were prepared, high BET surface (> 3200 m<sup>2</sup>/g) and tunable pore sizes (7Å to 10Å) achieved
- H<sub>2</sub> uptakes up to 5.5% at 77K and 0.5% at RT were achieved, heat of adsorptions are usually limited at ~6 kJ/mol
- High SSA leads to higher gravimetric hydrogen uptake at 77 K, but not necessarily higher volumetric uptake

### Technical Approach - Summary on Hydrogen Storage via Heteroaromatic POPs



- Over 30 heteroaromatic POPs were prepared containing B, N, S, etc., high BET surface (> 1000 m<sup>2</sup>/g) and narrow pore sizes (~8Å) achieved
- H<sub>2</sub> uptakes ~ 3% at 77K and the heat of adsorptions > 9 kJ/mol were achieved
- Improvement of  $\Delta H_{ads}$  is element-dependent, for example, S and N  $\rightarrow \Delta H_{ads} \downarrow$ , B  $\rightarrow \Delta H_{ads} \uparrow$

## Technical Approach - Summary on Hydrogen Storage via Metal Doped POPs

POPs with different TM-ligand coordinations were prepared ...



- Over 25 transition metal (Fe, Co, Ni...) doped POPs were prepared with BET surface (~2000 m<sup>2</sup>/g) and narrow pore sizes (~8Å) achieved
- $H_2$  uptakes of ~ 4% at 77K and the heat of adsorptions as high as ~ 10 kJ/mol were achieved
- Incorporating TMs clearly improves the isosteric heat of adsorption. New metals (Ti, Mg, V, etc.) and possibly new coordination chemistry need to be explored to enhance  $\Delta H_{ads}$  in the 15 ~20 kJ/mol range

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... for direct hydrogen storage evaluation ...

## FY2011 Technical Accomplishment 1 - Design & Synthesis of High Surface Area Carborane-containing POPs

#### **Rationale**

- Computational modeling studies from HSCoE suggest non-dissociative binding energy of 19.2 kJ/mol between H<sub>2</sub> and boron doped carbon cluster (Kim, et. al. *Phys. Rev. Lett.* 2006)
- Isotherm and spectroscopic studies from HSCoE found enhanced △H<sub>ads</sub> over B-doped graphitic carbon (T. Chung, et. al. JACS 2008, A. Kleinhammes, et. al. JPCC 2010)
- High surface area B-doped polymer can serve as precursor of high SSA adsorbent via further activation

#### Synthetic Scheme - An Example



#### HSCoE Studies on H<sub>2</sub> in BC<sub>3</sub> system



Simulation by Cooper/Cheng Airproducts, HSCoE

 $\Delta H_{ads}$  = 12~13 kJ/mol observed by Chung's group



Desired B-C framework Chung – Penn State, HSCoE

# FY2011 Technical Accomplishment 1 - Surface Properties & H<sub>2</sub> Storage Capacity of Carborane-containing POPs







BPOP-1, BET SSA =  $422 \text{ m}^2/\text{g}$ 

BPOP-2, BET SSA =  $864 \text{ m}^2/\text{g}$ 

BPOP-3, BET SSA =  $1037 \text{ m}^2/\text{g}$ 



# FY2011 Technical Accomplishment 1 - Improving $\Delta H_{ads}$ through B-doped POPs

H<sub>2</sub> adsorption isotherms at different T

<u><u>AHads</u> as function of hydrogen uptakes</u>



	BET SSA (m²/g)	Langr. SSA (m²/g)	Tot Pore Vol (cm <sup>3</sup> /g)	µ-pore Volume (cm³/g)	Pore Diameter (nm)	H <sub>2</sub> Gr. Uptake @ 77K (kg <sub>H2</sub> /kg <sub>Ads+H2</sub> )	B/C ratio	ΔH <sub>ads</sub> (kJ/mol)
BPOP-1	422	592	0.14	0.04	0.68	0.014	1/1.8	10.2
BPOP-2	864	1164	0.57	0.30	0.76	0.021	1/1.6	9.0
BPOP-3	1037	1497	1.12	0.33	0.77	0.028	1/3.0	8.2

•  $\Delta H_{ads}$  improves with higher B content in POPs, but it decays quickly with increase of hydrogen loading

■ Carborane POPs will be evaluated as precursors of adsorbents with improved ∠H<sub>ads</sub> through further activation

## FY 2011 Technical Accomplishment 2 - Design & Synthesis of **TM/Polyporphyrin POPs**

#### Rationale

- Unsaturated transition metal/hydrogen interaction could form H<sub>2</sub>. TM bond, leading to improved heat of adsorption ( $\Delta H_{ads}$ ) without dissociation (Kubas interaction)
- Computational modelings suggest enhanced  $\Delta H_{ads}$  can be achieved by decorating TM in graphene plane (M. Yoon et. al. ORNL) or macrocyclic plane (Kim & Zhang NREL/RPI)
- ANL/UC team successfully prepared and characterized several TM doped POPs with high surface area and narrow pore size distribution



#### Synthetic Scheme: Fe-Co/Por

#### HSCoE theoretical studies on H<sub>2</sub> over TM doped surfaces



V7 x V7 TM-Decorated Graphene Model M. Yoon, et.al., HSCoE - ORNL

Suitable ranges of  $\Delta H_{ads}$  for storage are suggested through non-dissociative binding of H<sub>2</sub>/square-planar TM coordination site



H<sub>2</sub>/HEME Model Kim & Zhang, HSCoE - NREL/RPI



# FY 2011 Technical Accomplishment 2 - Design & Synthesis of TM/Polyporphyrin POPs



## FY 2011 Technical Accomplishment 2 - Improving H<sub>2</sub> Storage Capacity & Heat of Adsorption over TM/Polyporphyrin POPs



■ Transition metal (Ni, Co, and Fe) addition improves *△H<sub>ads</sub>*, mechanism needs to be studied

Other promising metals (Ti, V, Mg, etc. ) need to be explored with different doping method

# FY 2011 Technical Accomplishment 3 - Design & Synthesis of TM/POPs Coordinated through Hydroxyquinoline

#### <u>Rationale</u>

- POPs crosslinked through TM-hydroxyquinoline bonds are very stable towards humidity and other contaminants
- Coordination geometry between TMs and ligand (square-planar vs. tetrahedral) may shed light on hydrogen-metal interaction

#### Synthetic Scheme: Metal Coordinated Hydroxyquinoline



Co/HQ-1, BET SSA: 580 m<sup>2</sup>/g; Ni/HQ-1, BET SSA: 596 m<sup>2</sup>/g

Other TM/HQ POPs was also prepared with high SSAs

The tetrahedral (Co) metal-ligand coordination led to POP with a slightly larger pore size than that with square-planar (Ni) ligation

## FY 2011 Technical Accomplishment 3 - Surface Properties, $H_2$ Storage Capacity & $\Delta H_{ads}$ of TM/HQ-1s



	BET SSA (m²/g)	Langr. SSA (m²/g)	Tot. Pore Vol. (cm <sup>3</sup> /g)	µ-pore Volume (cm³/g)	Pore Diameter (nm)	H <sub>2</sub> Gr. Uptake @ 77K (kg <sub>H2</sub> /kg <sub>Ads+H2</sub> )	∆H <sub>ads</sub> (kJ/mol)
Ni/HQ-1	596	807	0.59	0.22	0.82	0.017	8.0
Co/HQ-1	580	786	0.58	0.22	0.87	0.017	9.9

Co-hydroxyquinoline POP produced a higher isosteric heat of adsorption (9.9 kJ/mol) than its nickel counterpart, suggesting a possible metal dependent H2-TM/POP interaction.

## FY 2011 Technical Accomplishment 4 -Adsorption/Desorption Kinetics Study



Adsorption/desorption time dependence at liquid nitrogen temperature



- The temporal profiles between adsorption and desorption are nearly identical
- The system reaches equilibrium in a faster pace at ambient temperature than 77 K, all within 20 seconds

## Summary Table

#### H<sub>2</sub> storage capacities for selected POPs by Argonne – UofC Team in FY2011

Sample	BET SSA (m²/g)	<b>Gr. Uptake</b> (77K, 40 bars) (kg H <sub>2</sub> /kg adsorbent+H <sub>2ads</sub> )	Vol. Uptake <sup>a</sup> (77K, 40 bars) (kg H <sub>2</sub> /L adsorbent)	<b>Gr. Uptake</b> (RT, 70 bars) (kg H <sub>2</sub> /kg adsorbent+H <sub>2ads</sub> )	<b>Vol. Uptake</b> (RT, 70 bars) (kg H <sub>2</sub> /L adsorbent)	μ-Pore /Total Pore (cm <sup>3</sup> g <sup>-1</sup> / cm <sup>3</sup> g <sup>-1</sup> )	Skeleton Vol. <sup>b</sup> (cm <sup>3</sup> g <sup>-1</sup> )	<b>⊿H<sub>ads</sub></b> (kJmol⁻¹)
PAF-1 (A)	3143	0.055	0.020	0.0053	0.0018	1.08/2.08	0.82	6.5
РТА-З (Н)	870	0.029	0.021	0.0024	0.0017	0.40/0.66	0.80	6.6
BPOP-1 (H)	422	0.013	0.014	0.0016	0.0017	0.04/0.14	0.81	10.2
BPOP-2 (H)	864	0.021	0.014	0.0018	0.0012	0.30/0.57	0.94	9.0
BPOP-3 (H)	1037	0.027	0.016	0.0030	0.0015	0.33/1.12	0.87	8.2
Ni/Por-1 (M)	1704	0.035	0.021	0.0038	0.0022	0.66/0.91	0.85	8.0
Fe-Co/Por (M)	1571	0.034	0.022	0.0041	0.0026	0.63/0.97	0.65	7.4
Ni/HQ-1 (M)	596	0.017	0.015	0.0019	0.0017	0.22/0.59	0.56	8.0
Co/HQ-1 (M)	580	0.017	0.016	0.0019	0.0017	0.22/0.58	0.53	9.9

A – Aromatic POP; H – Heteroaromatic POP; M – Metal doped POP

- a. Volumetric capacity is calculated based on the measured skeleton density plus total pore volume density.
- b. Skeleton volume is measured using helium as calibration gas
- c. Rate of  $\Delta H_{ads}$  (kJ/mol) change as function of gravimetric uptake  $C_g$  (%) at ambient temperature near the zero coverage point.
  - System with the highest gravimetric capacity does not necessarily have the highest volumetric capacity
  - Transition metal and boron doping can improve adsorption enthalpy

## Collaboration

#### Partnership within and outside of Hydrogen Sorption Center of Excellence

- Teaming between Argonne National Laboratory (prime) and The University of Chicago (subcontractor)
- Members of DOE HSCoE under the clusters of "Engineered Nanospace" (RC1) and "Substituted Materials" (RC2)
- Collaboration with UNC (HSCoE member) on <sup>1</sup>H NMR experiment
- Collaboration with NREL (HSCoE member) on measurement validation
- Information exchange with ORNL (HSCoE member) and RPI on computational modeling and simulation

### **Technology Transfer through HSCoE**

- Valuable inputs on our adsorption apparatus test validation
- New ideas and direction, examples include B and metal doped polymers
- Collaboration opportunities in polymer characterization, examples include NMR study
- Up-to-date information on new developments in sorption based materials

## **Future Work**

- Complete the investigation on TM exchanged polyporphyrin POPs
- Complete the investigation on improving heat of adsorption for B-doped POP via activation
- Evaluate potential application of other emerging technologies to sorption based hydrogen storage and recommend further research direction to DOE
- Prepare final project report

#### **FUTURE DIRECTION**

- POP is becoming a great platform as hydrogen adsorbent due to its high surface area and narrow pore (vs. carbon) and excellent chemical stability (vs. MOF)
- Unmodified aromatic POPs will unlikely reach desired H<sub>2</sub> adsorption enthalpy for room temperature application even with increased surface area or adjustable porosity
- To achieve near-ambient temperature application, the surface of POP needs to be modified by incorporating metal or other elements promoting H<sub>2</sub> binding
- New surface modification techniques, departing from the conventional synthetic approaches, have the potential to produce high binding energy sites predicted by theory
- POPs with tailored surface property and chemical composition can also serve as precursor of fabricating new adsorbent after additional chemical/physical processing