

# Optimization of SWNT Production and Theoretical Models of H<sub>2</sub>-SWNT Systems For Hydrogen Storage

Carried out in the DOE Center of Excellence  
on Carbon-based Hydrogen Storage

Boris Yakobson and Robert Hauge

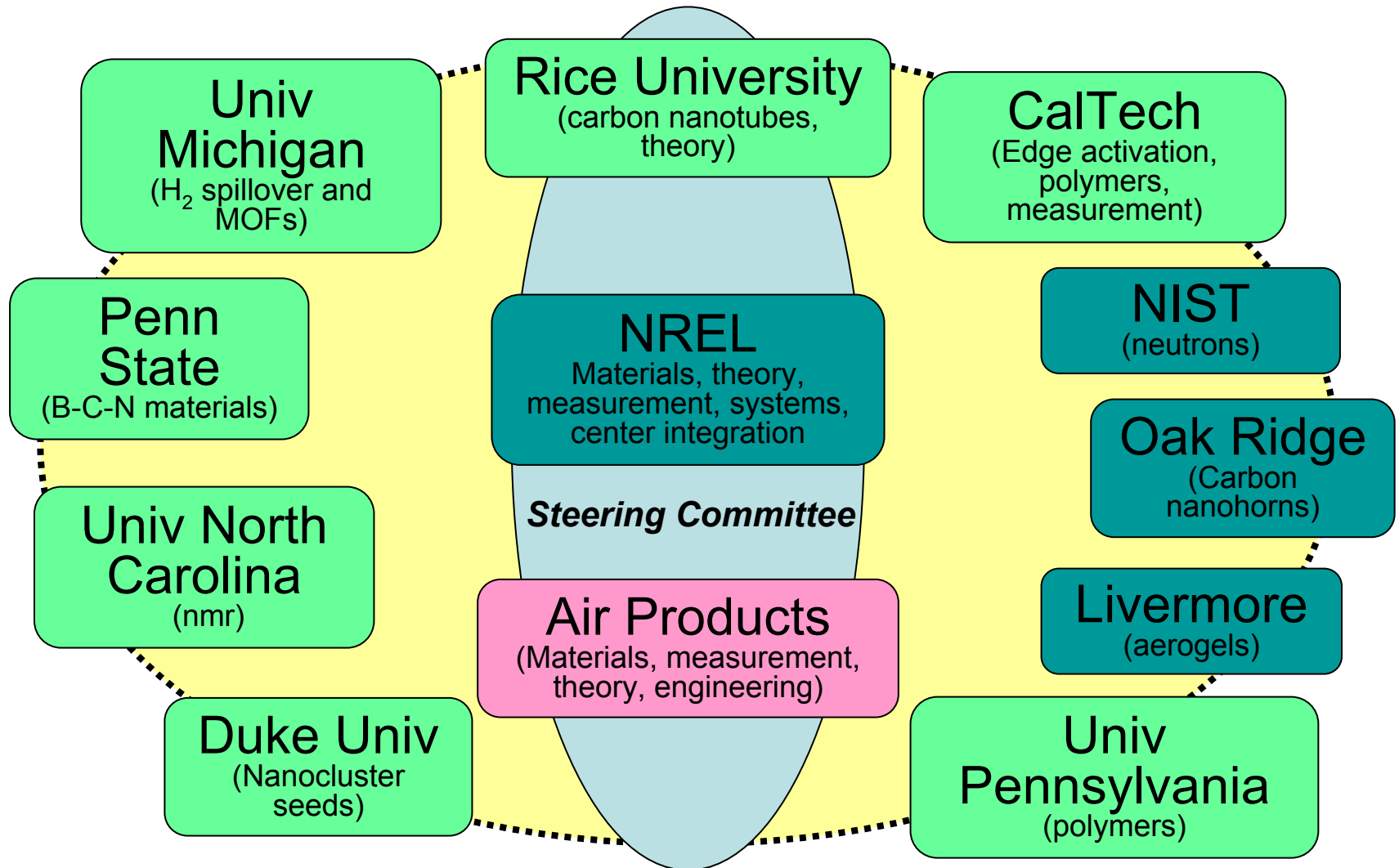
Rice University

May 25, 2005

Project ID  
#STP38

# CbHS Center of Excellence Partners

9 university projects (at 7 universities), 4 government labs, 1 industrial partner



# **Optimization of SWNT Production and Theoretical Models of H<sub>2</sub>-SWNT Systems for Hydrogen Storage**

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## **General Objectives:**

**To produce single wall carbon nanotubes by the HiPco process that are optimal for molecular hydrogen adsorption**

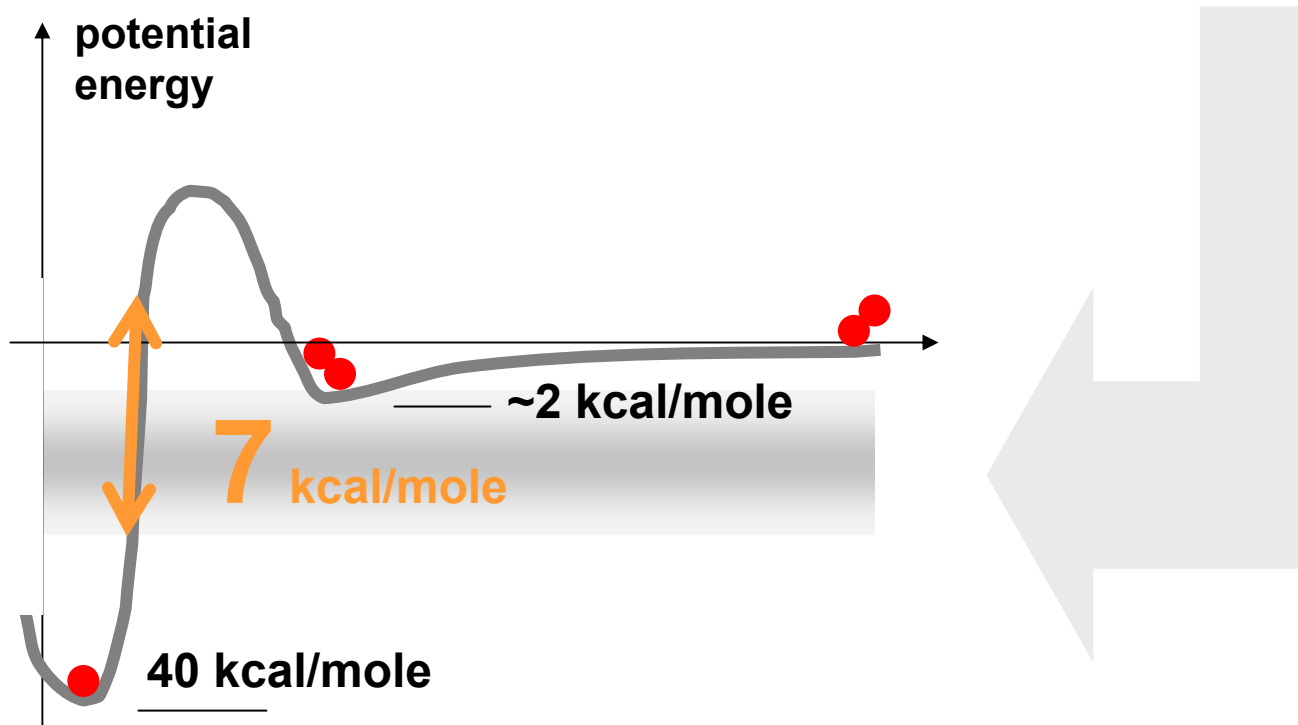
**To expand theoretical analysis and computations of hydrogen interaction with carbon nanotubes produced at Rice by HiPco technology.**

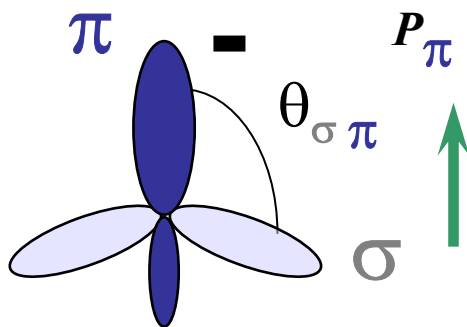
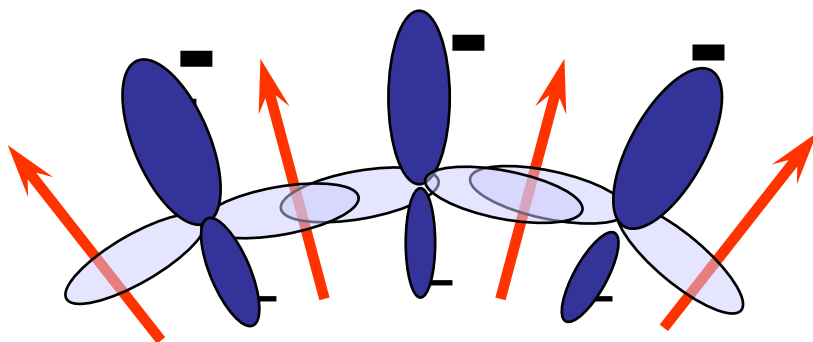
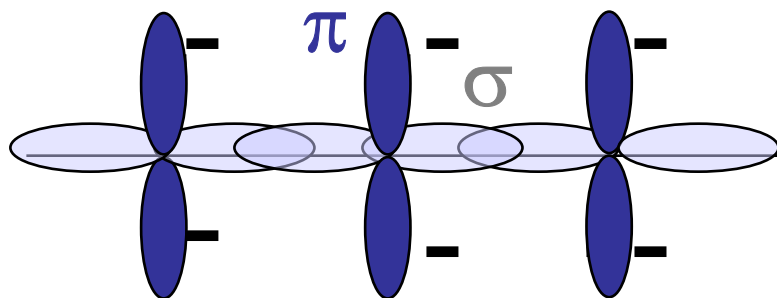
**Obtain quantitative maps of binding energies for various chiral and diameter types of SWNT.**

**To derive recommendations, wherever possible, upon the best diameter and/or chirality of the HiPco SWNT**

## Challenge:

Can the desirable range of binding energies be achieved for some types of nanotubes, due to their static or dynamic curvature, local electrical fields, and/or possible role of metal-dopant?

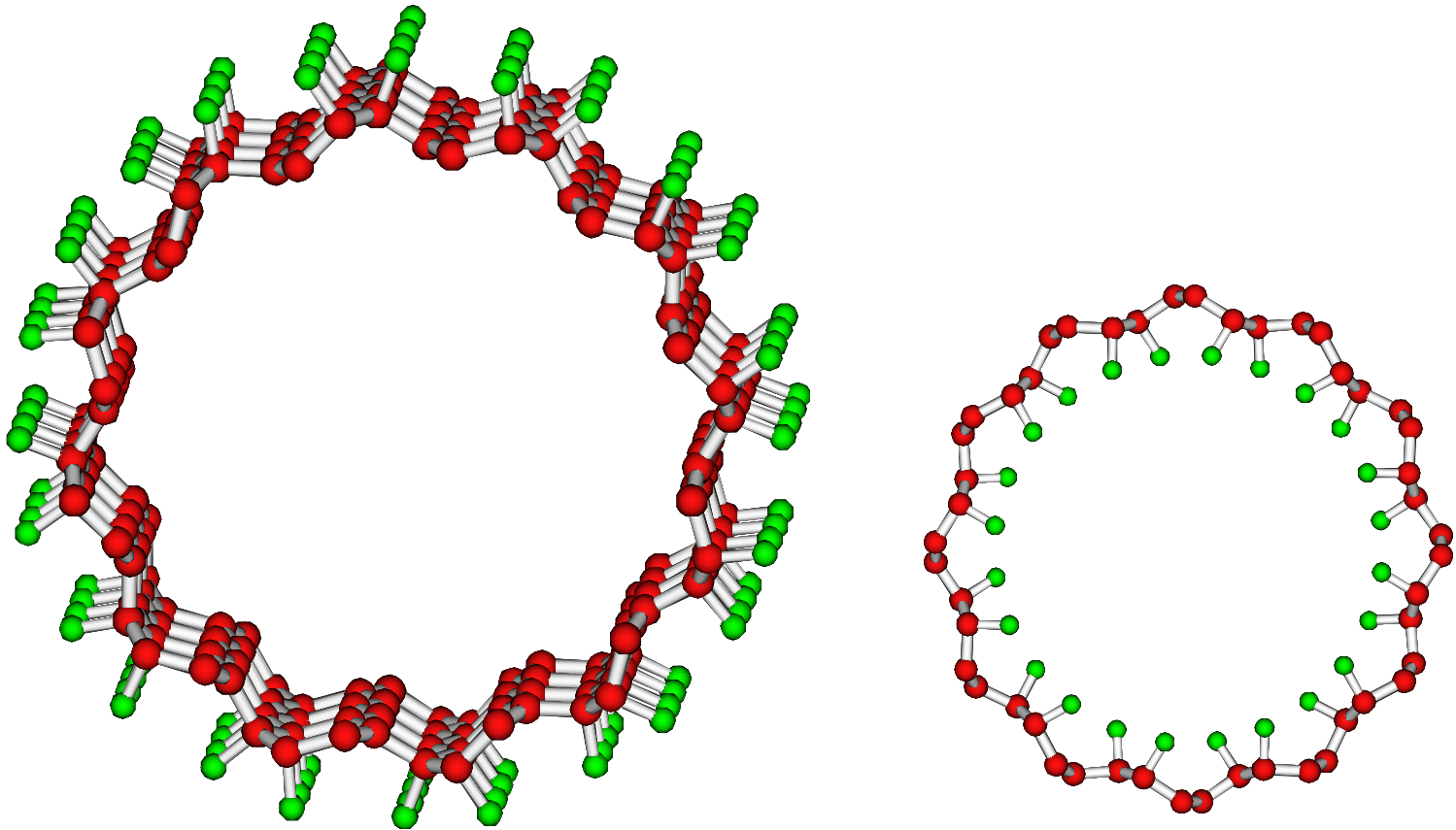




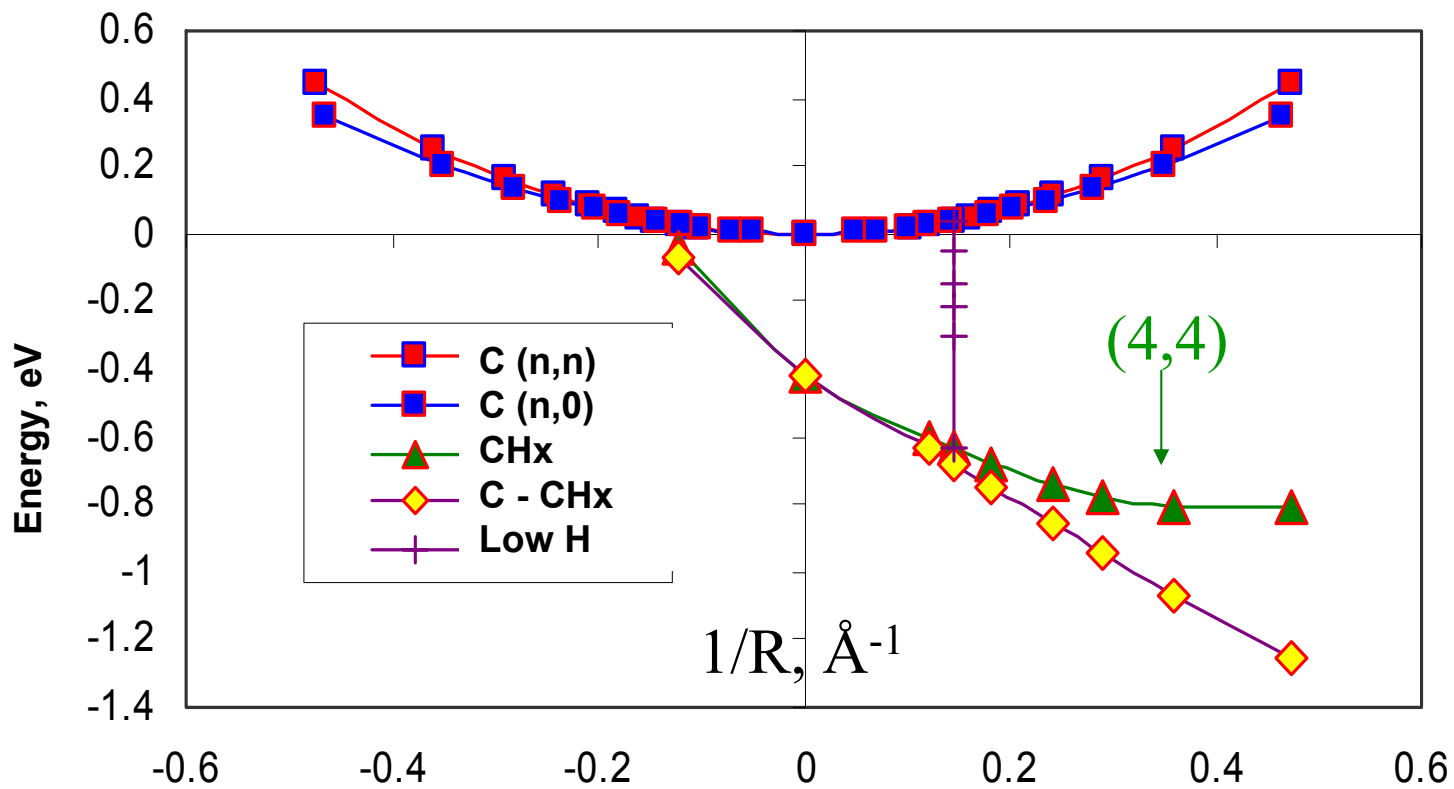
**Will further explore curvature-induced polarization and evaluate local field and their ability to change the energy of physisorption**

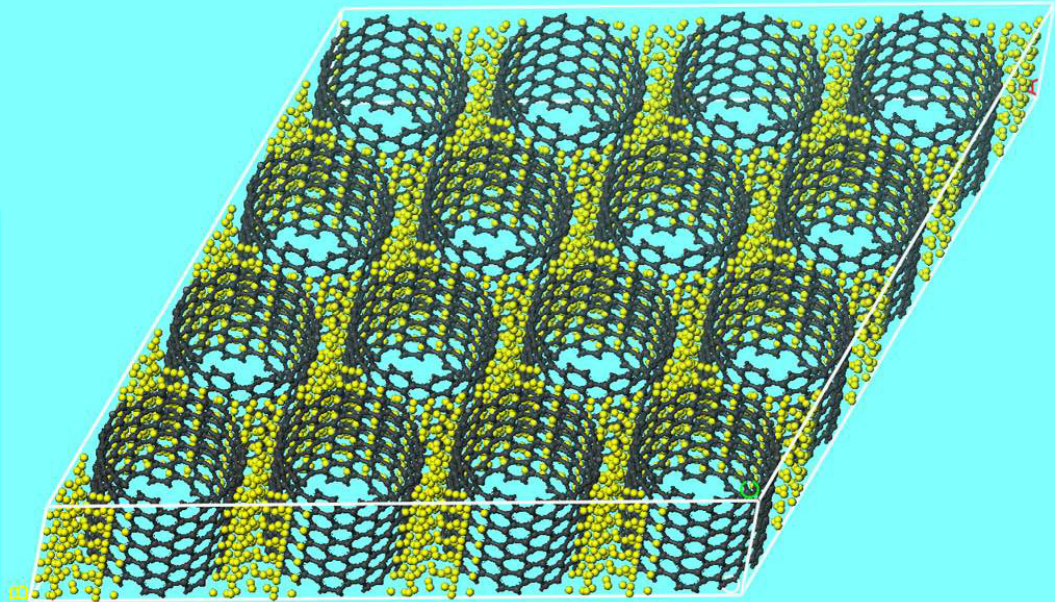
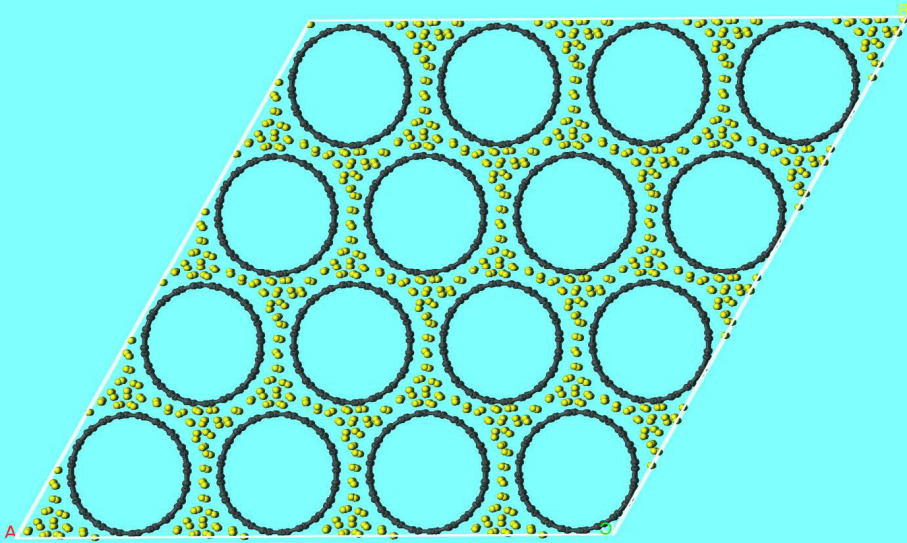
Preliminary/past work  
 Yakobson *et al.*,  
*Chem Phys Lett* v **360**, p 182

**Compute binding energies and the “saturation limit” for nanotubes of different diameters and chiral types, for both exterior and interior attachment (shown here is armchair type)**



Generate quantitative maps of binding energy as a function of radius (as shown here) and chirality, as a difference of total energies of **pristine** and **H-saturated** tubular structures

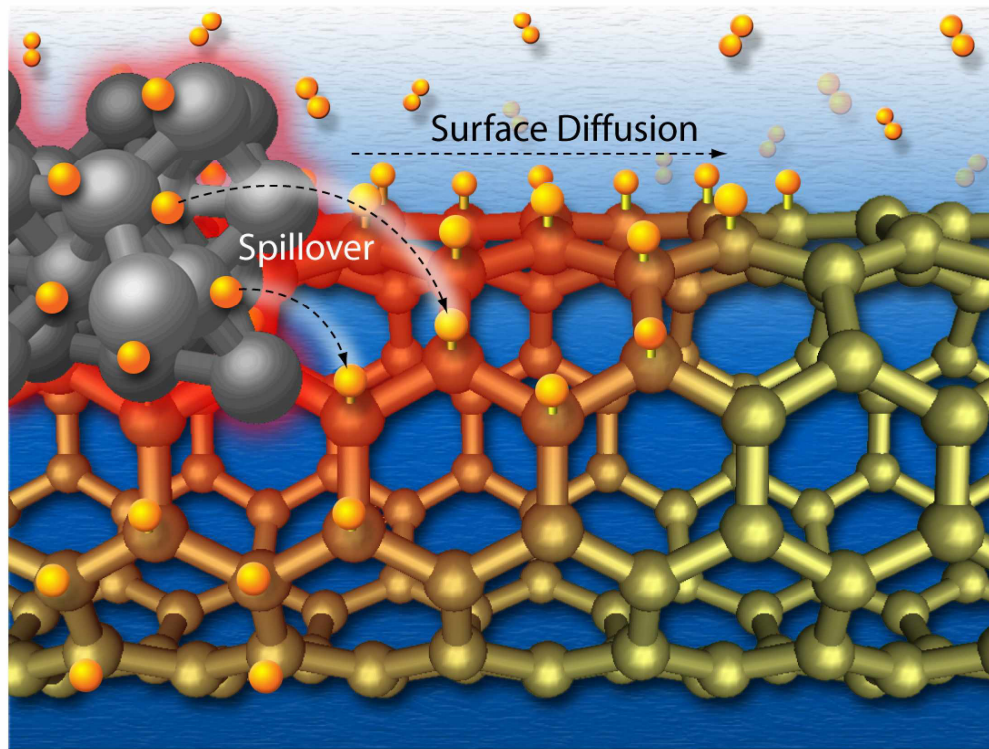




From analysis of H-SWNT interaction, move to a large scale **molecular dynamics** simulations, as shown, including **thermodynamic-integration** for computing thermodynamic potentials of the joint SWNT-array-hydrogen systems, as storage battery prototypes.

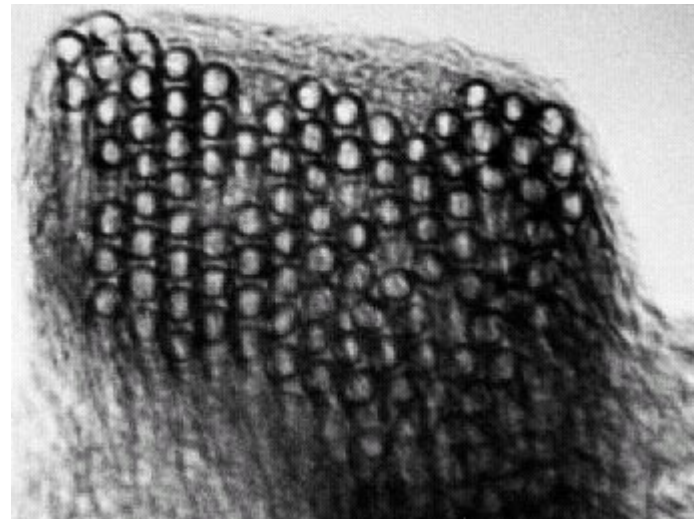
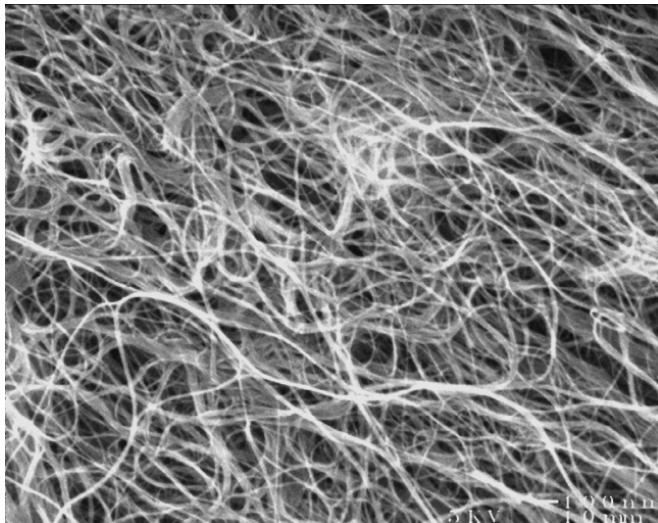


Besides thermodynamics of H-bonding, we will investigate **kinetics** of hydrogen redistribution along the SWNT surface. Quantum computations of the **activation barriers** for H-hopping will be input into and **transition state theory** in order to evaluate time-scale of storage cycle and the necessity of metal catalyst spillover.



# Why SWNT?

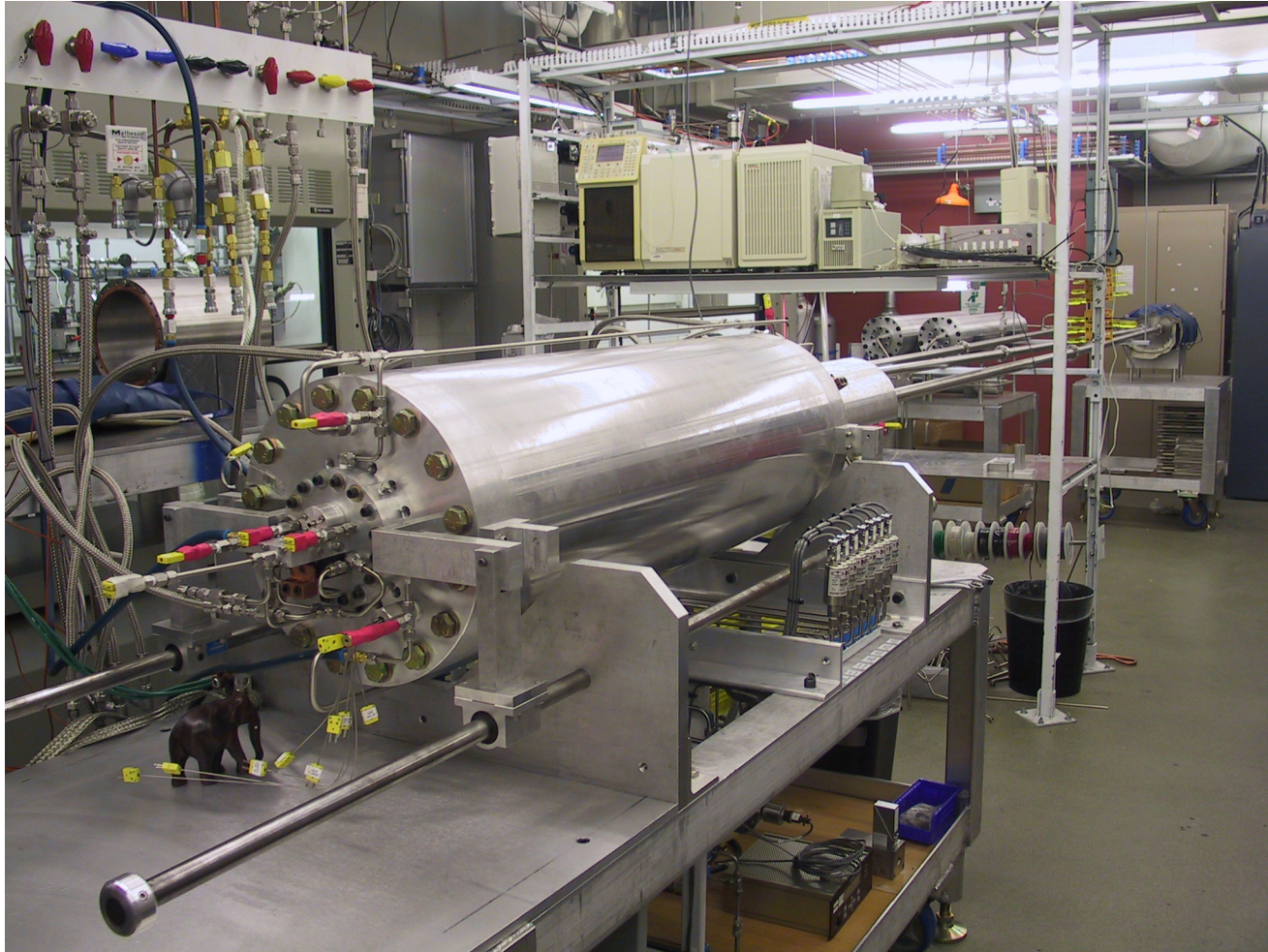
- The Scale and Perfection of DNA
- The Strongest Possible Fiber
- Thermal Conductivity of Diamond
- The Unique Chemistry of Carbon
- Maximum Possible Surface Area
- Selectable Electrical Properties
  - Metallics Better Than Copper
  - Semiconductors Better Than InSb or GaAs
- *The Ultimate Engineering Material*



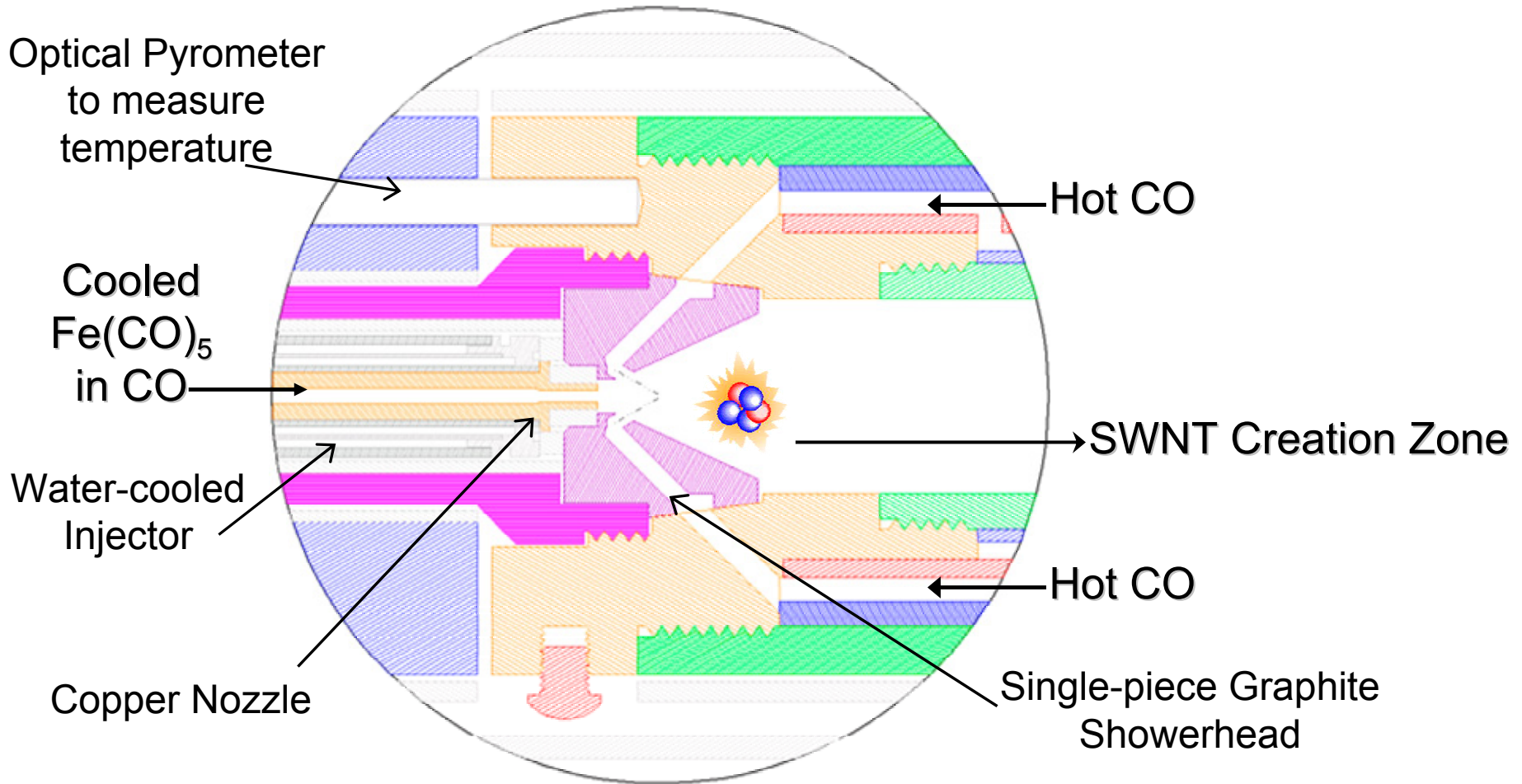
# The HiPco Process

- High Pressure CO (HiPco) at ~1000 °C is mixed with room temperature gas containing gas phase iron pentacarbonyl
- Standard operating conditions are iron carbonyl at 52 mtorr in the reactor with an overall reactor pressure of 30 atm
- The iron nucleates forming small iron clusters a few atoms in size
- These iron clusters then catalyze the Boudouard reaction
- $2 \text{CO} \rightarrow \text{CO}_2 (\text{g}) + \text{C} (\text{s})$
- Nucleated catalyst particles grow SWNTs
- The SWNTs are filtered out and the CO is recycled to create a continuous process

# HiPco Reactor



# HiPco Mixing Zone



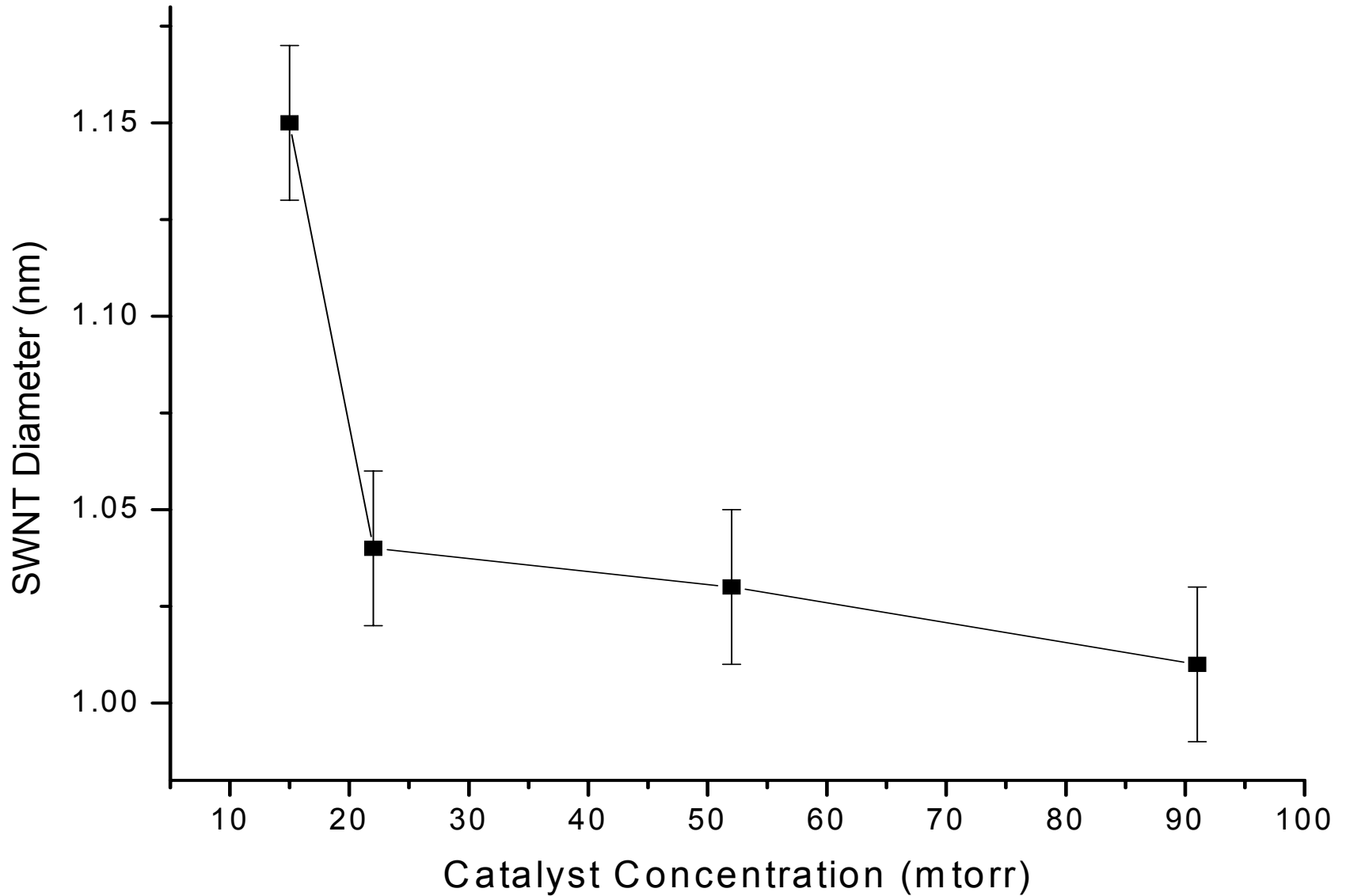
# Single Wall Carbon Nanotube Production

This project aims to advance HiPco™ production in order to enable a strong material-supply base for hydrogen storage.

We will take advantage of **tunability** of HiPco™ and guide the production conditions based on detailed theoretical models of **H<sub>2</sub>**-swnt interactions and their possible enhancement

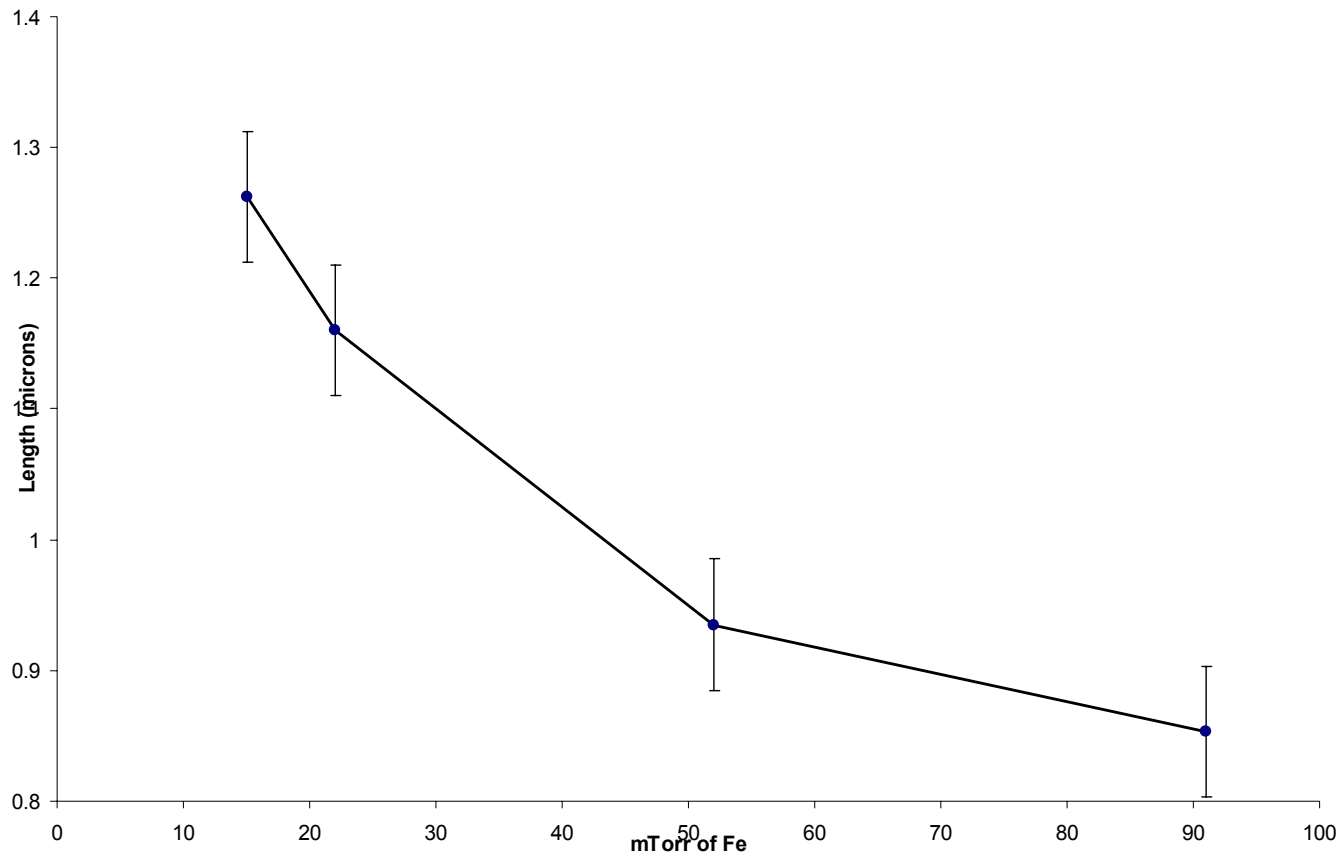
# **Effects of Metal Catalyst Concentration Variation On Single Wall Carbon Nanotube Properties In the HiPco Reactor**

# Average SWNT Diameters





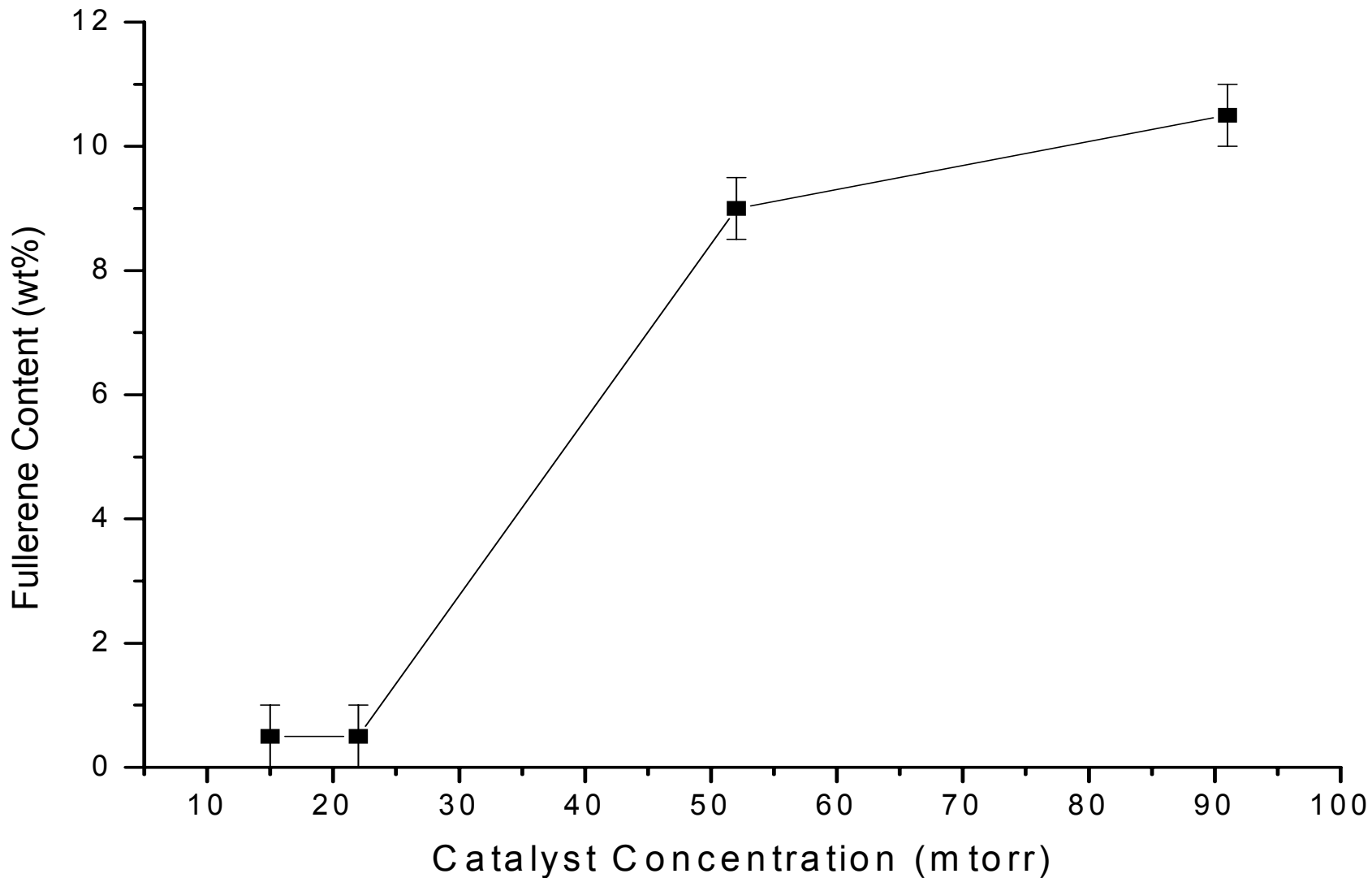
# Average SWNT Lengths



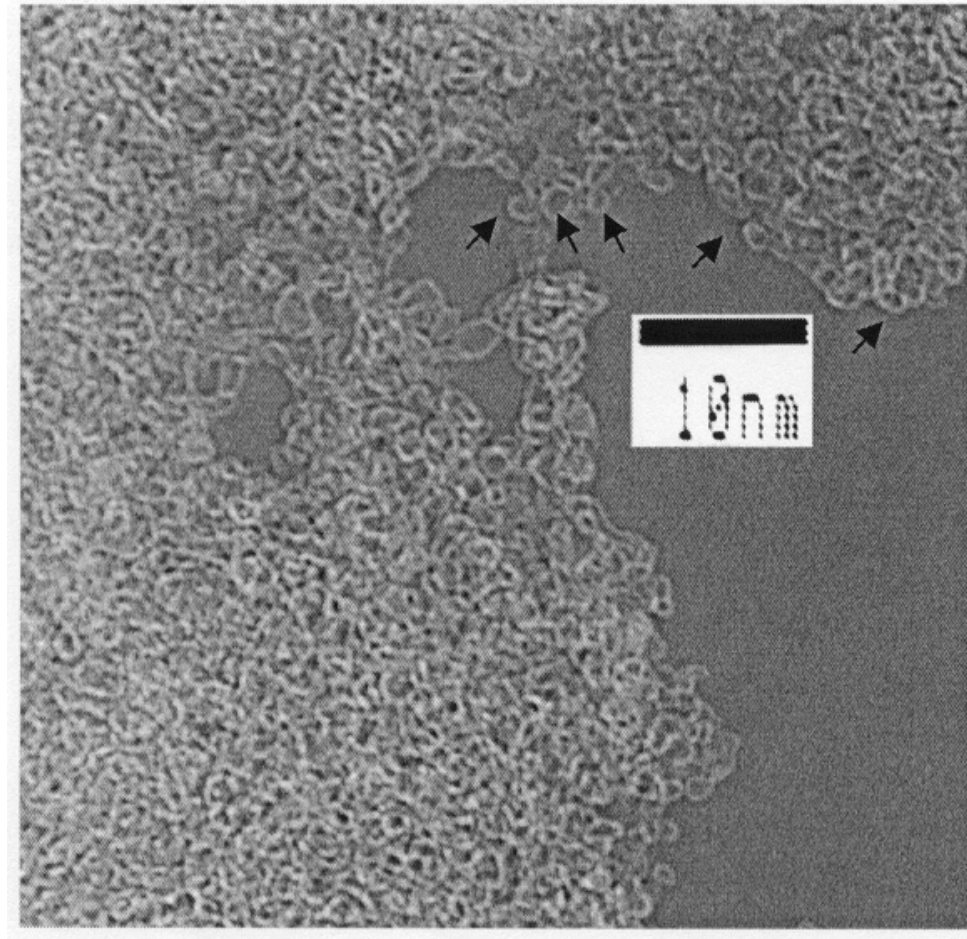
# Superfullerenes

**Superfullerenes, the end caps of single wall carbon nanotubes joined together, will be studied as a subset of swnt and in combination with swnt as high surface area supports for molecular hydrogen adsorption.**

# Large Fullerene Production



# Large Fullerenes



- Large fullerenes extracted from raw HiPco Sample by fluorination

# Milestones, Interactions & Safety

- **Milestones (fy05)**
  - Compute binding energies & saturation limit for various SWNT types
  - Compute quantitative maps of H binding energies afa tube radius
  - Estimate H hopping barriers along SWNT
  - Production of 50 grams of HiPco material optimized for H<sub>2</sub> adsorption
- **Milestones (fy06)**
  - Compute binding energies & saturation limit for various SWNT types
  - Compute quantitative maps of H binding energies afa tube radius
  - Estimate H hopping barriers along SWNT
  - Production of 50 grams of HiPco material optimized for H<sub>2</sub> adsorption
- **Interactions**
  - Smalley & Heben on SWNT Growth
  - Modelers on Hydrogen Storage Team (
- **Safety**
  - No lab safety or materials issues expected
- **Go/no-go decision**
  - Go/no-go decision of the use of swnt as hydrogen storage material at the end of fy06

# Overview

## Timeline

- Project start date: FY05
- Project end date: FY09
- New Start

## Budget

- **Expected Total Funding**
- **Total \$1,072,515**
  - DOE share \$857,996
  - Contractor share \$214,520
- **Funding for FY05 \$200,004**

## Barriers

Reversible Solid-State Material Storage Systems:  
Hydrogen Capacity and Reversibility  
Lack of Understanding of Physisorption and Chemisorption *variability*  
Test Protocols and Evaluation Facilities

## Partners

- Interactions - NREL, Air Products
- Collaborations - Duke Univ., ORNL