# Gas Adsorption in Metal Organic Frameworks : an experiment using the NCNR Disk Chopper Spectrometer

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## Overview

- Issues with Energy Storage:
  - The how and the why.
- Applications of Neutron Scattering
  - Locations of molecules
  - Dynamics/binding strengths
- Outlook for this experiment
- Conclusion



#### Why alternative fuels?

- Reduce dependence on foreign oil
- Harness renewable energy sources
- New opportunities for agriculture
- Clean air in cities
- Reduce transportation costs
- Reduce greenhouse gas emissions

#### What are alternative fuels?

- Ethanol (from corn, wood, ...)
- Natural gas; 85% of NG used in U.S. is domestic
   (NG; from domestic gas/oil fields, deep-sea methane hydrate fields, landfills, biomass)
- Biodiesel (from soybeans, vegetable oils, ...)
- Hydrogen (from NG, water & electricity, coal, ...)
- Electricity (from nuclear/hydroelectric/solar/wind power plants)



#### **Current natural-gas vehicles**

- Low emission of
  - hydrocarbons (ozone, smog)
  - $-NO_{x}$
  - particulate matter
  - Up to 40% reduction of CO<sub>2</sub>
- Clean Cities Coalitions:
  - Los Angeles: 1500 CNG buses
  - Kansas City: 200 CNG public utility vehicles
  - U.S.: 130,000 CNG vehicles
  - worldwide: over 5 million CNG vehicles

Alternative fuel systems (BAF Tech.)

In 2006, Gasoline was \$2.84 per gallon, diesel was \$2.98 per gallon, and CNG was \$1.90 per gasoline gallon equivalent!

http://www.eere.energy.gov/afdc/resources/pricereport/price\_report.html















#### **Fuel Storage**

Compressed natural gas (CNG) is stored on board vehicles at high-pressure (3,000 psi)

Liquefied natural gas (LNG) must be cooled to -162 °C.

LNG requires only 30 percent of the space of CNG to store the same amount of energy.

# Why?



\$1.2 Billion to develop the technology needed for commercially viable hydrogen-powered fuel cells (2003)



#### However:

H<sub>2</sub> has 3x energy content by **MASS** c.f. gasoline

Gasoline has 4x energy content by **VOLUME** c.f H<sub>2</sub>



Schlapbach and Zuttel (2001) Nature 414: 353-358

# **Targets**

#### **Methane**

#### Hydrogen

180 (cc CH<sub>4</sub>)/cc 35bar (500 psi)/25K

Parameter	'07	'10
Energy <sub>(system)</sub> (wt%) Volumetric (g/L)	4.5 36	6 45
Fuel cost (\$ per gge)	3	1.5
Reversible, safe		

gge: gallon gasoline equivalent

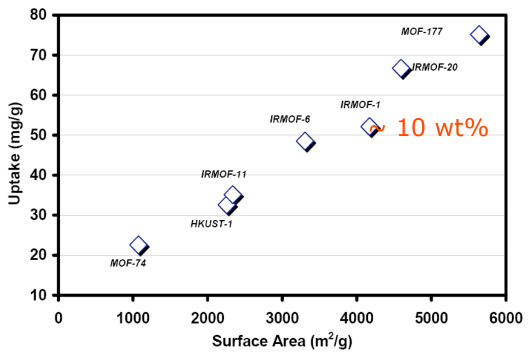
-achieved using carbonized corncobs (*Pfeifer, University of Missouri, 2007*)

-IRMOF-6 155 cc/cc (Eddaoudi, Science 2002)

-IRMOF-1 ~115 cc/cc (*Zhou, in prep.*)

Gravimetric and volumetric of best MOFs @77K ~7 wt%, ~36g/L (e.g. *Dinca, JACS, 2006*) – NOT *SYSTEM* 

# Hydrogen Storage in MOFs



(Wong-Foy et al. JACS 128, 3494 (2006))

MOF-5 (IRMOF-1) can adsorb  $\sim$ 10 wt% H<sub>2</sub> (<10 K)

(Yildirim et al. PRL 95, 215504 (2005))

# Hydrogen Adsorption Enthalpy

```
HKUST-1
                                                 ~6.6 kJ/mol <sup>1</sup>
                                                 ~7.4 kJ/mol<sup>2</sup>
Prussian blue analogus
MOF-74
                                                              ~8.3 kJ/mol <sup>1</sup>
                                                 ~8.7 kJ/mol <sup>3</sup>
Zn<sub>3</sub>(1,4-benzeneditetrazolate)<sub>3</sub>
IRMOF-11
                                                 ~9.1 kJ/mol <sup>1</sup>
                                                 ~9.4 kJ/mol 4
Cu_{1.5}[(Cu4Cl)_3BTT_8]
PCN-9
                                                 ~10.1 kJ/mol <sup>5</sup>
                                                 ~10.1 kJ/mol 6
Mn_{1.5}[(Mn_4CI)_3BTT_8]
```

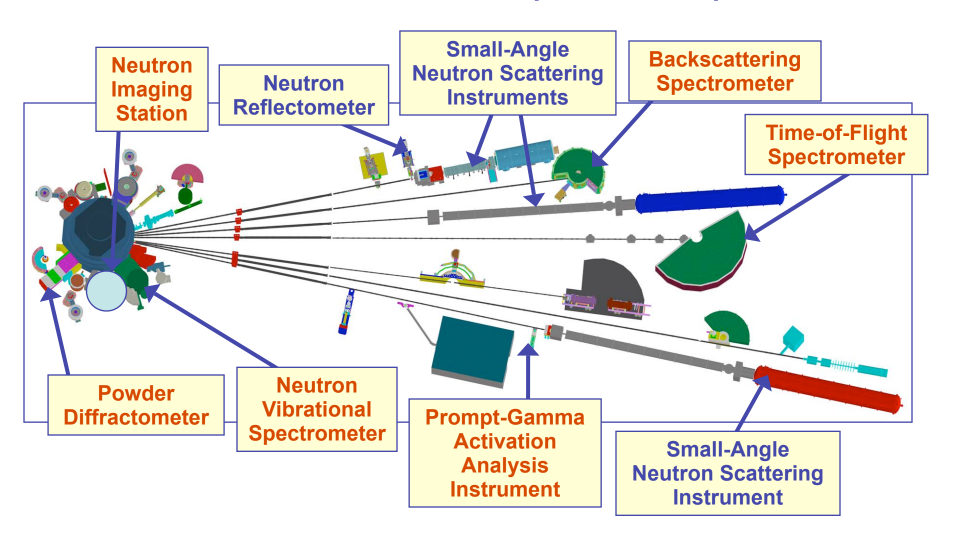
# ~15 kJ/mol would be ideal for hydrogen storage material working at room temperature.

(S. K. Bhatia, A. L. Myers, *Langmuir* 22, 1688 (2006))

#### Reference:

- 1. Rowsell et al., J. Am. Chem. Soc. 128, 1304 (2006)
- 2. S. S., Kaye et al., J. Am. Chem. Soc. 127, 6506 (2005)
- 3. M. Dinca et al., J. Am. Chem. Soc. 128, 8904 (2006)
- 4. M. Dinca et al., Angew. Chem. Int. Ed., in press (2007)
- 5. S. Ma et al., J. Am. Chem. Soc. 128, 11734 (2006)
- 6. M. Dinca et al., J. Am. Chem. Soc. 128, 16876 (2006)

# NIST Center for Neutron Research (NCNR)



#### HKUST-1

Cu<sub>3</sub>(1,3,5 benzenetricarboxylate)<sub>2</sub>

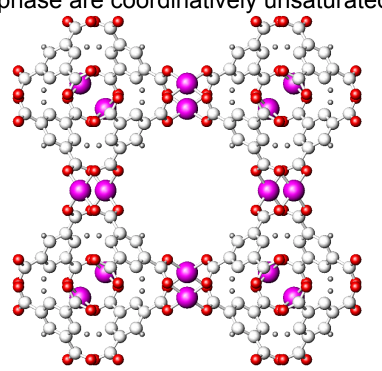
The Cu atoms in the fully dehydrated phase are coordinatively unsaturated

Desolvated crystals exhibit :

Total H<sub>2</sub> uptake of ~3 wt % at 77 K and 90 bar

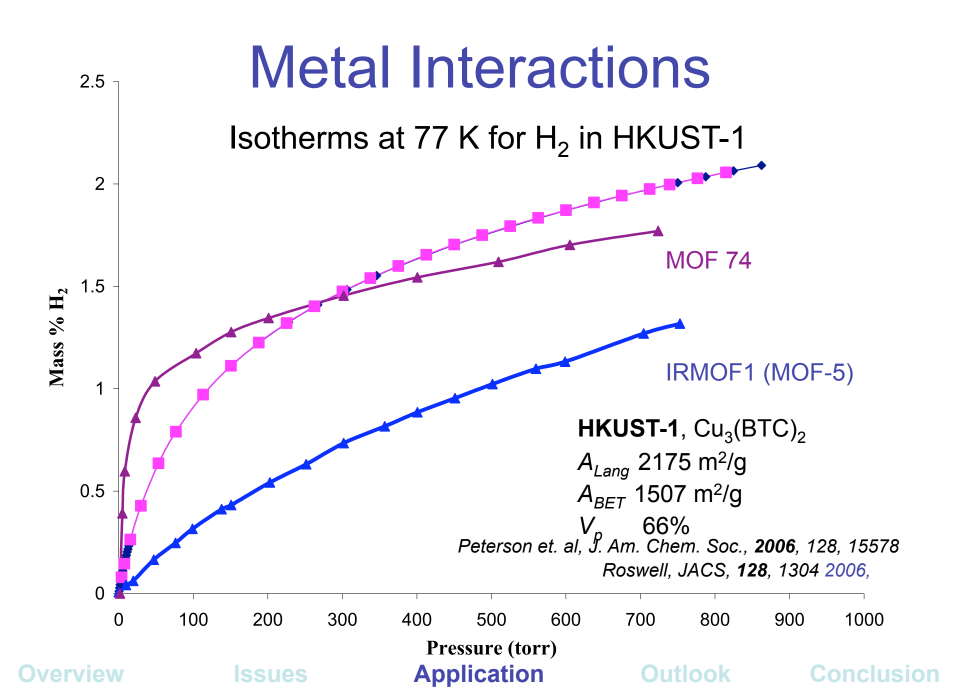
At 27 g H<sub>2</sub>/L provides a storage density <40% of that of liquid H<sub>2</sub>

A maximum isosteric heat of adsorption of 6.6 kJ/mol



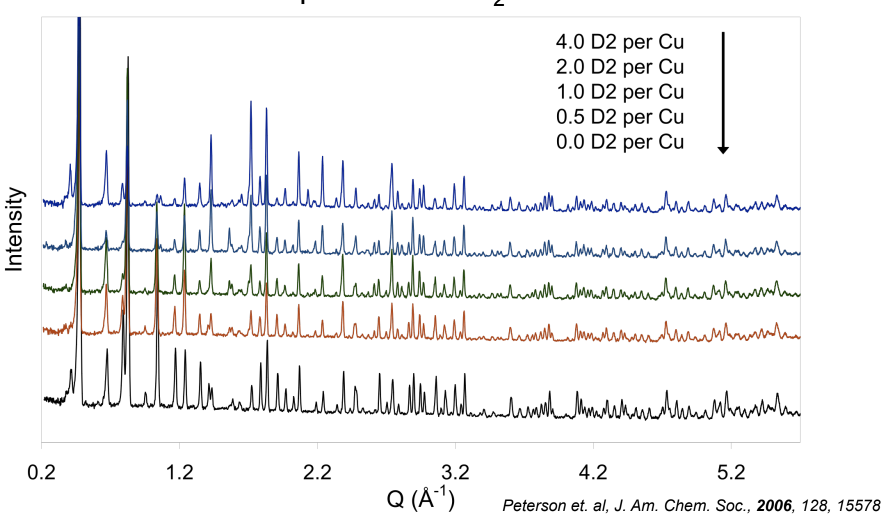
Chui, Science, **283**, 1148 1999 Roswell, JACS, **128**, 1304 2006 Wong-Foy, JACS., **128**, 3494 2006

Prestipino, Chem. Mater., 18 (5), 1337 2006



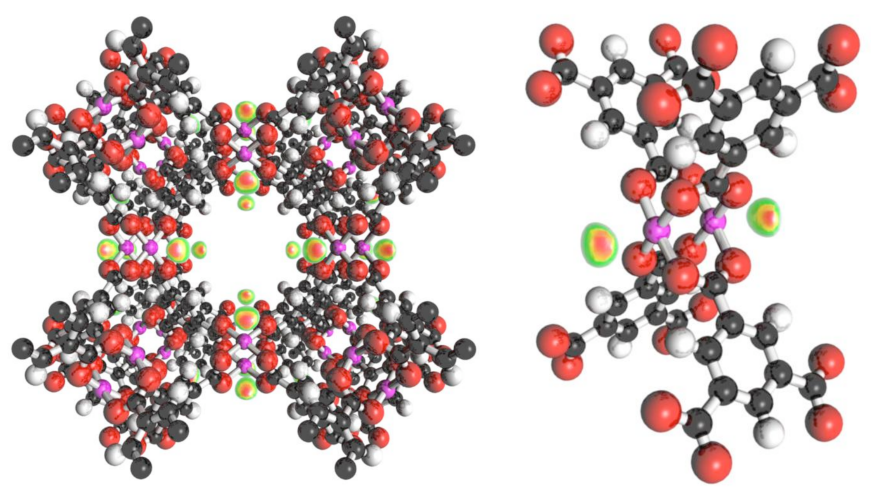
#### **Metal Interactions**

Diffraction patterns for D<sub>2</sub> in HKUST-1



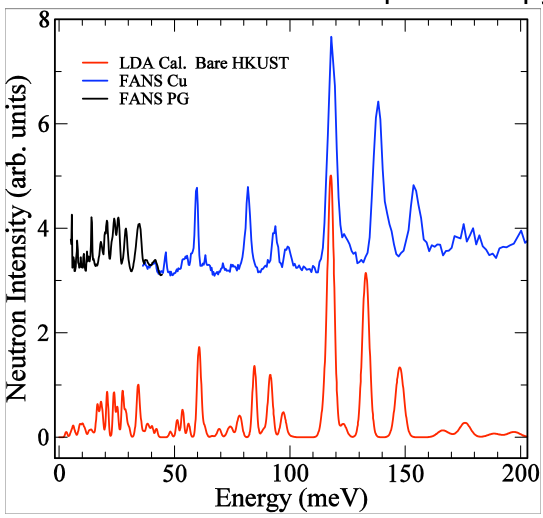
#### **Metal Interactions**

Fourier Difference to locate D<sub>2</sub> in HKUST-1



## Framework and DFT

Bare HKUST-1 Spectroscopy



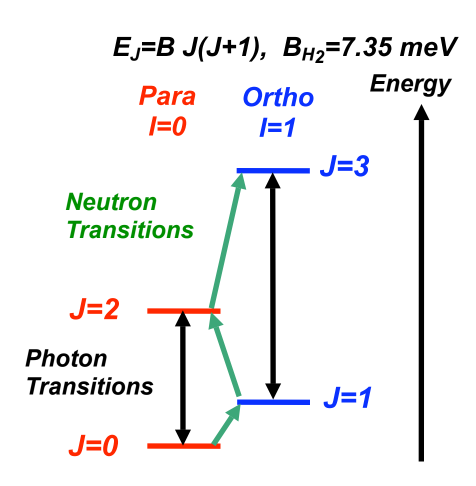
unpublished

## Hydrogen Transitions

Para has a nuclear spin I=0. This constrains J to be even.

Ortho has a nuclear spin I=1. This constrains J to be odd.

Transition between ortho and para species can occur through flipping the nuclear spin.

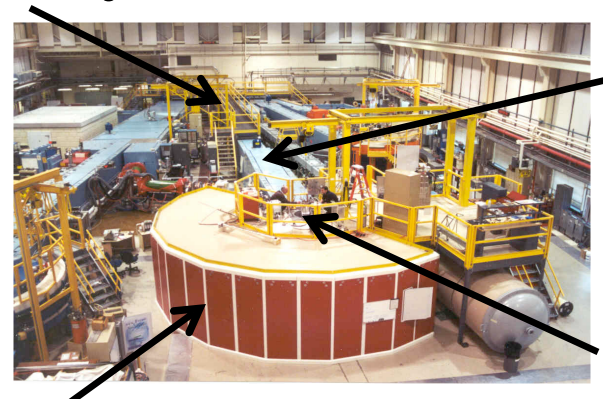


(Neutron energy loss)

## TOF spectroscopy

Disc Chopper Spectrometer

(1) The neutron guide

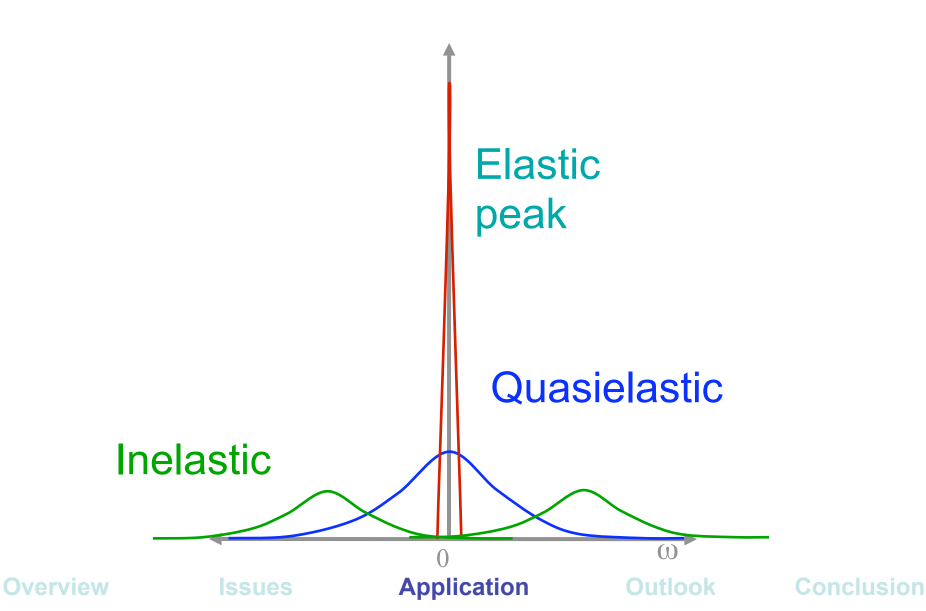


(2) The choppers

(4) The flight chamber and the detectors

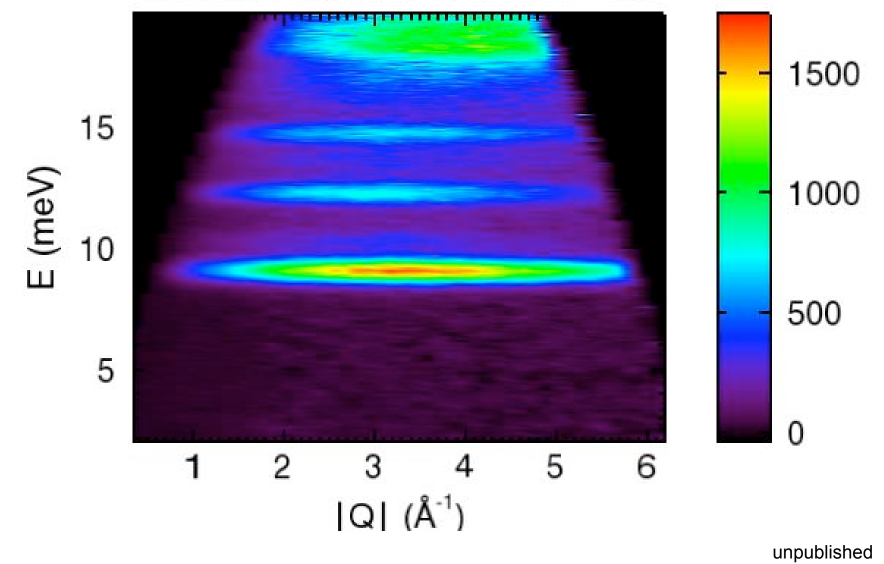
(3) The sample area

# **Total Scattering**

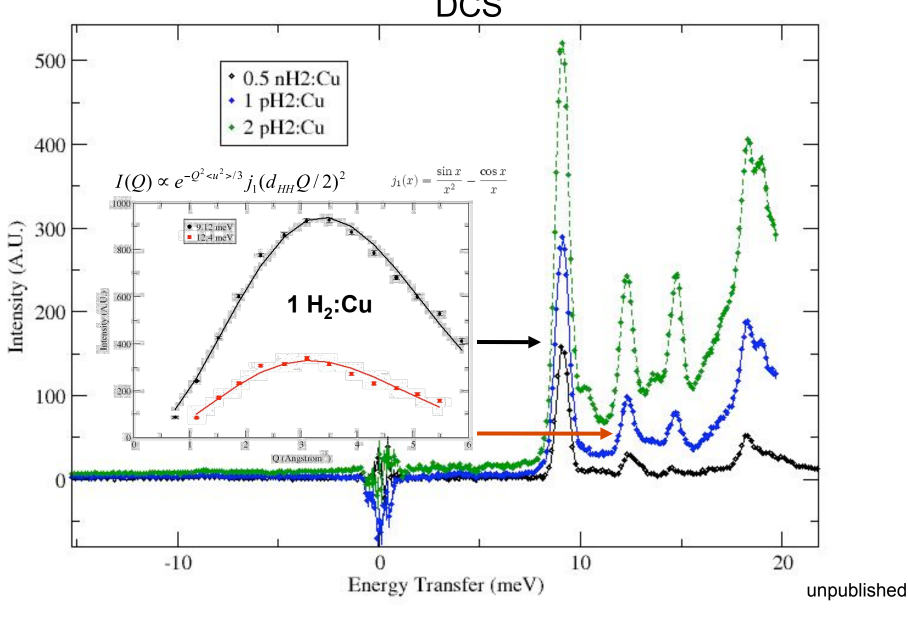


# Metal Interactions

HKUST-1 2 p-H2:Cu 1.81A T=4K

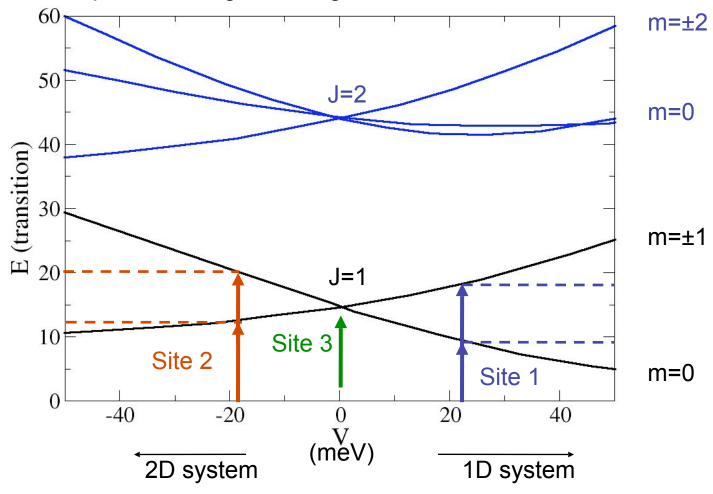






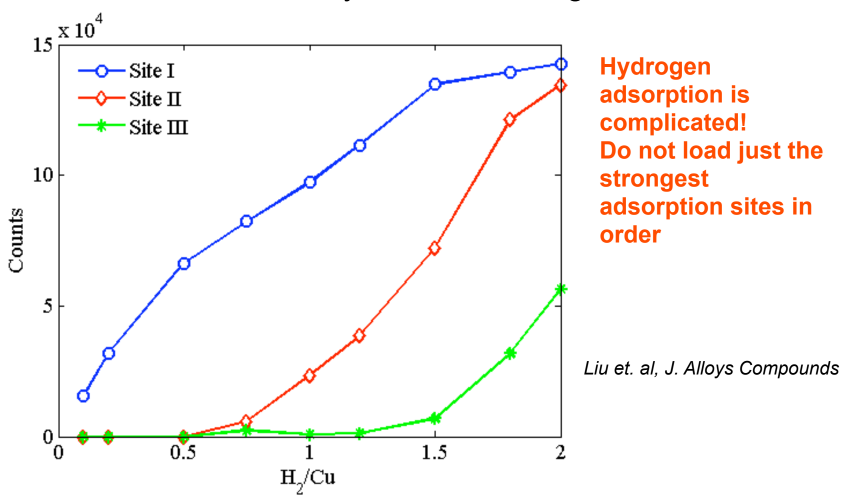
# Metal Interactions Spectroscopy

The transition tells us about the symmetry and strength of the local potential. A larger rotational barrier *implies* a stronger binding.



## **Metal Interactions**

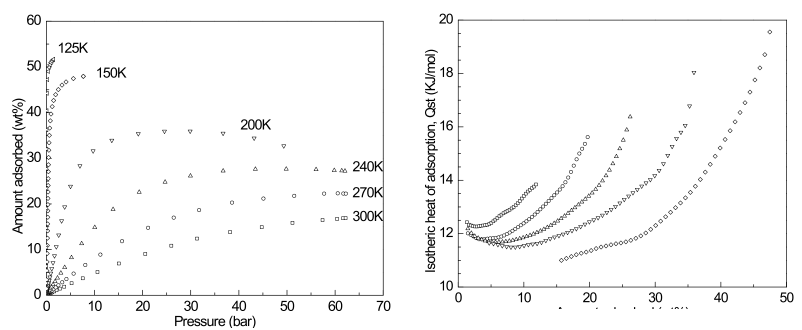
Extract Intensity as fn of loading...



## Outlook

- Experience Practical TOF spectroscopy
  - sample choice
  - geometry consideration
- Learn something about the instrument
  - Wavelength / Resolution / Intensity
- Data Reduction
- Data Analysis and Interpretation
  - Tunneling spectroscopy
  - Quasi-elastic spectroscopy
    - spatial and temporal information

# Adsorption isotherms



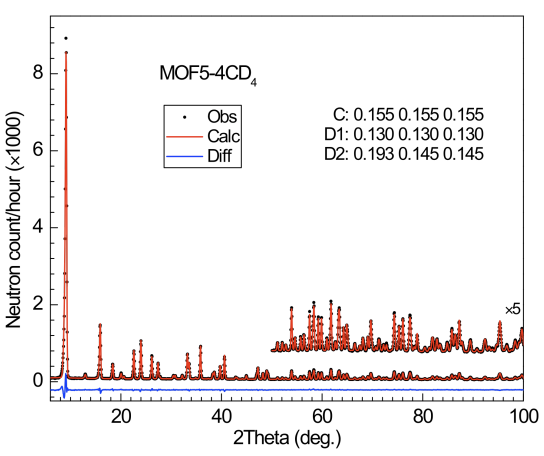
The maximal excess adsorption capacity of CH<sub>4</sub> in MOF-5 51.7 wt%, or 24 CH<sub>4</sub> per MOF-5 formula (i.e., 4Zn). This is reduced to ~15 wt% (115 cc/cc) at room temperature, 35bar.

The excess isosteric heat of adsorption (calculated using the Clausius-Clapeyron equation) for the initial CH₄ adsorption in MOF-5 is ~12.2 KJ/mol. At high concentration, Qst increases with increasing amount adsorbed, indicating the importance of the interactions between adsorbed CH<sub>4</sub> molecules. **Outlook** Application

(Zhou, in prep.)

#### Where are the methane molecules?

The adsorption sites were directly determined from the difference-Fourier analysis of neutron powder diffraction data. Initial adsorption occurs at the MOF5 "cup site" with a well defined CH<sub>4</sub> orientation. We did not see any well-defined sites for additional adsorption.



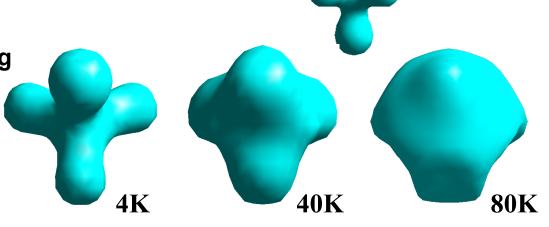
The neutron powder diffraction pattern of MOF-5:4CD₄ at 4 K with the Rietveld refinement.

(Zhou, in prep.) Conclusion

#### Where are the methane molecules?

The isosurface of the difference-Fourier (DF) neutron scattering-length density superimposed with the ZnO<sub>4</sub> clusters of the MOF-5 host structure, indicating the location of the first methane adsorption sites. This is a "direct measurement" (like taking a picture) of the methane molecules packed in the solid with a well defined orientation.

T-dependent neutron scattering is further used to visualize the methane orientational dynamics with increasing temperature.



Outlook Conclusion

# Types of Experiments

- Translational and rotational diffusion processes, where scattering experiments provide information about time scales, length scales and geometrical constraints; the ability to access a wide range of wave vector transfers, with good energy resolution, is key to the success of such investigations
- Low energy vibrational and magnetic excitations and densities of states
- Tunneling phenomena
- Chemistry --- e.g. clathrates, molecular crystals, fullerenes
- Polymers --- bound polymers, glass phenomenon, confinement effects
- Biological systems --- protein folding, protein preservation, water dynamics in membranes
- Physics --- adsorbate dynamics in mesoporous systems (zeolites and clays) and in confined geometries, metal-hydrogen systems, glasses, magnetic systems
- Materials --- negative thermal expansion materials, low conductivity materials, thermo-electrics, hydration of cement, carbon nanotubes, proton conductors, metal hydrides

## Conclusions

- Neutrons can tell us where atoms are located.
- Neutrons can tell us how a lattice vibrates->

   very sensitive to the local potential
- Neutrons can tell us adsorbate-framework interaction strengths