



University of
Massachusetts
Amherst

Spectroscopic Signatures of Nitrogen-Substituted Zeolites

Item Type	article;article
Authors	Hammond, Karl D;Dogan, Fulya;Tompsett, Geoffrey A;Conner, Wm. Curtis;Grey, Clare P;Auerbach, Scott M
Download date	2025-03-23 11:23:18
Link to Item	https://hdl.handle.net/20.500.14394/47956



Spectroscopic Signatures of Nitrogen-Substituted Zeolites

KARL D. HAMMOND, GEOFFREY A. TOMPSETT, W. CURTIS CONNER, JR., SCOTT M. AUERBACH

E-Mail: khammond@ecs.umass.edu.

Address: 159 Goessmann Lab / University of Massachusetts / Amherst, MA 01003.

FAX: (413) 545-1647

Abstract

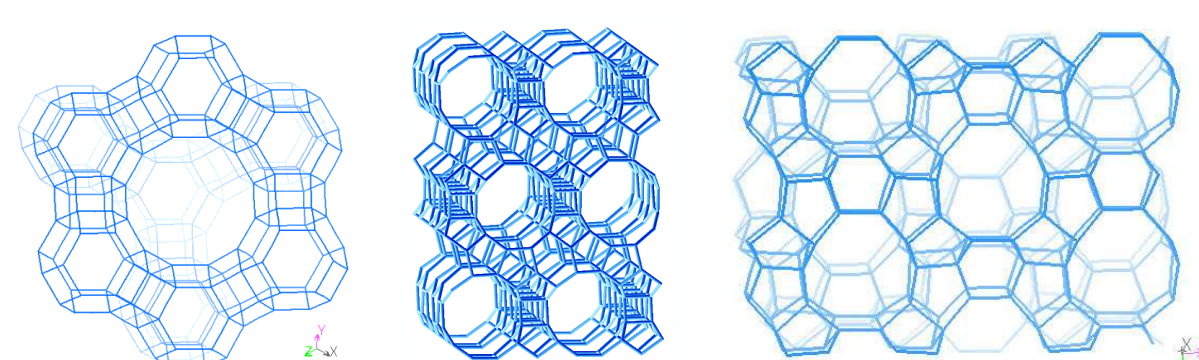
- Zeolites are crystalline microporous materials
 - Zeolites are typically acids
 - Treating zeolites with amines at high temperatures has been shown to produce basic (alkaline) catalysts in which nitrogen is substituted for oxygen
 - Goal: Find a way to characterize nitrogen substituted zeolites

Methods

- Experiment: nuclear magnetic resonance spectroscopy, infrared spectroscopy, Raman spectroscopy, X-ray diffraction, physical adsorption
- Calculations: cluster models, Reaction energies, chemical shielding/chemical shifts (NMR), quadrupolar NMR, vibrational spectra (infrared/Raman).

Materials of interest

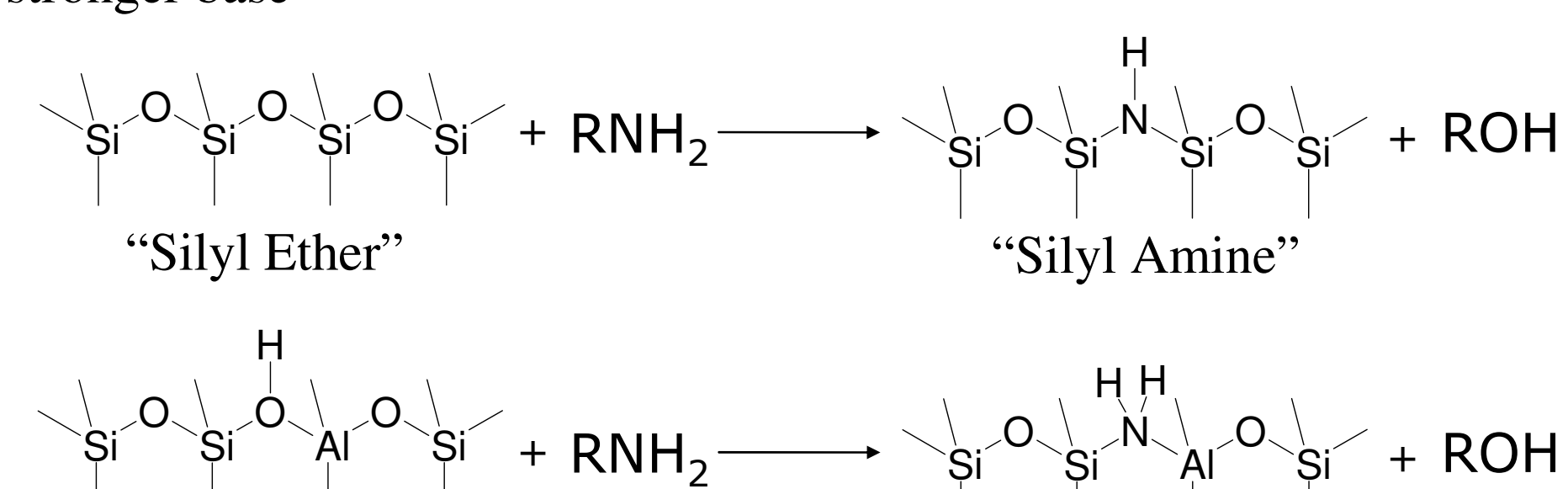
- Y zeolite (FAU structure)
- Beta zeolite (BEA structure)
- ZSM-5 (MFI structure)



We find compelling evidence from experiments and simulations that nitrogen incorporates into zeolite frameworks. Questions remain of the stability of these materials.

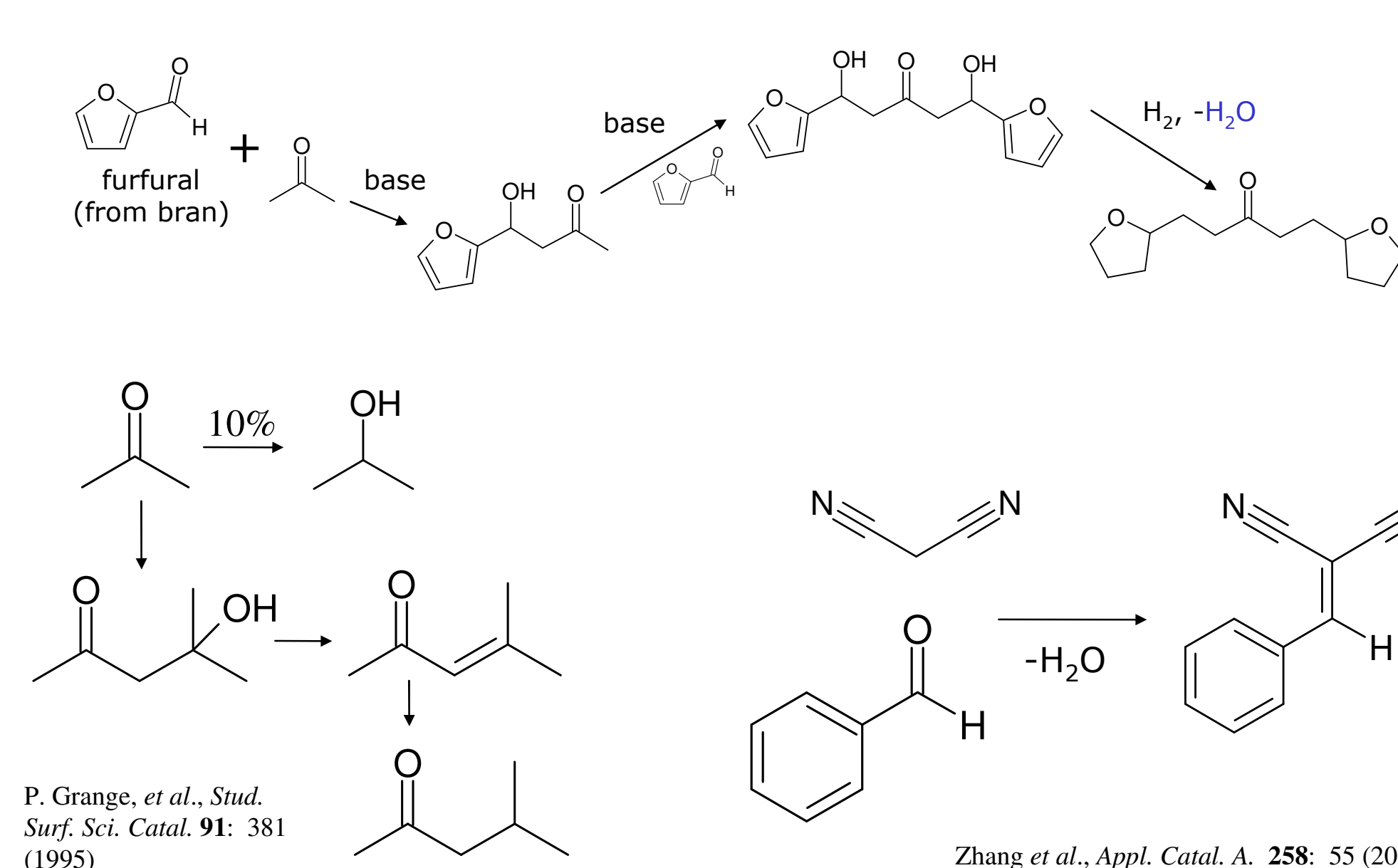
Zeolites as Acid/Base Catalysts

- Zeolites are usually strongly acidic
 - Low-aluminum zeolites are stronger acids
 - High-aluminum zeolites are weak(er) acids
- The alkaline or alkaline-earth forms are weakly basic: NaY, CsX, K-ZSM-5
 - Low-aluminum zeolites (e.g. K-ZSM-5) are very weak bases
 - High-aluminum zeolites (e.g. CsX) are merely weak bases
- IDEA: Replace some oxygen in the zeolite with nitrogen to produce a stronger base



Examples of Uses for Zeolitic Base Catalysts

Condensation reactions

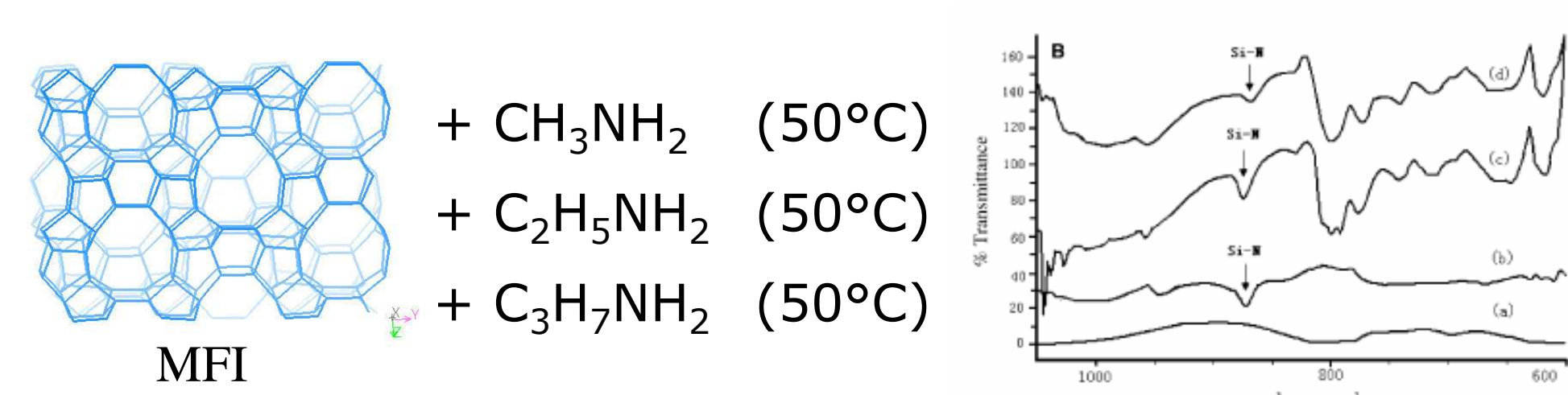
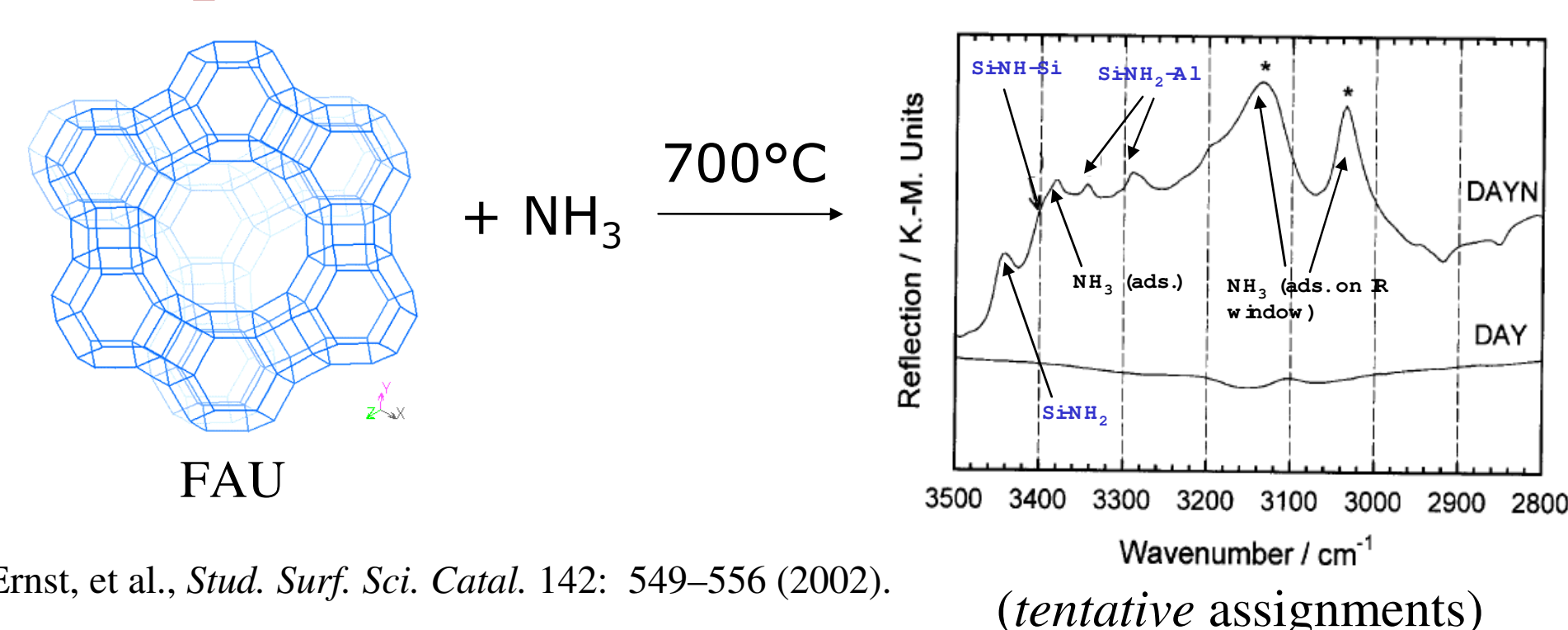


How to Make Zeolites into Bases

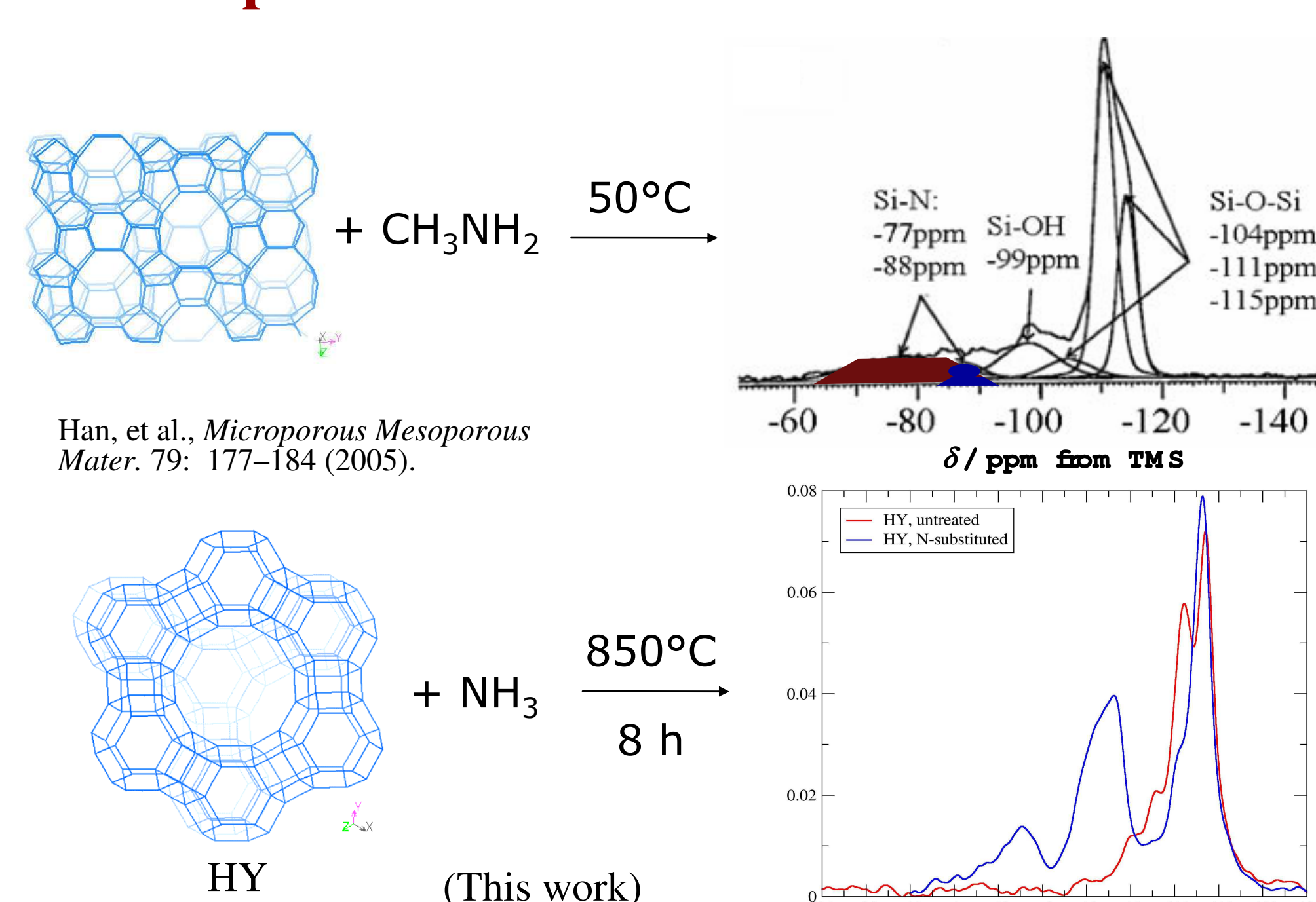
- Step 1: Heat
 - Removes water
 - Removes some hydroxides on surface
- Step 2: Expose to a nitrogen source:
 - Ammonia (NH₃)
 - Alkylamines (CH₃NH₂, C₂H₅NH₂, etc.)
 - Others: Polysilazane (SiCl₄+NH₃ deposited and pyrolyzed), Silicon nitride (Si₃N₄), Azides (NaN₃, etc.), Nitrogen gas (N₂)



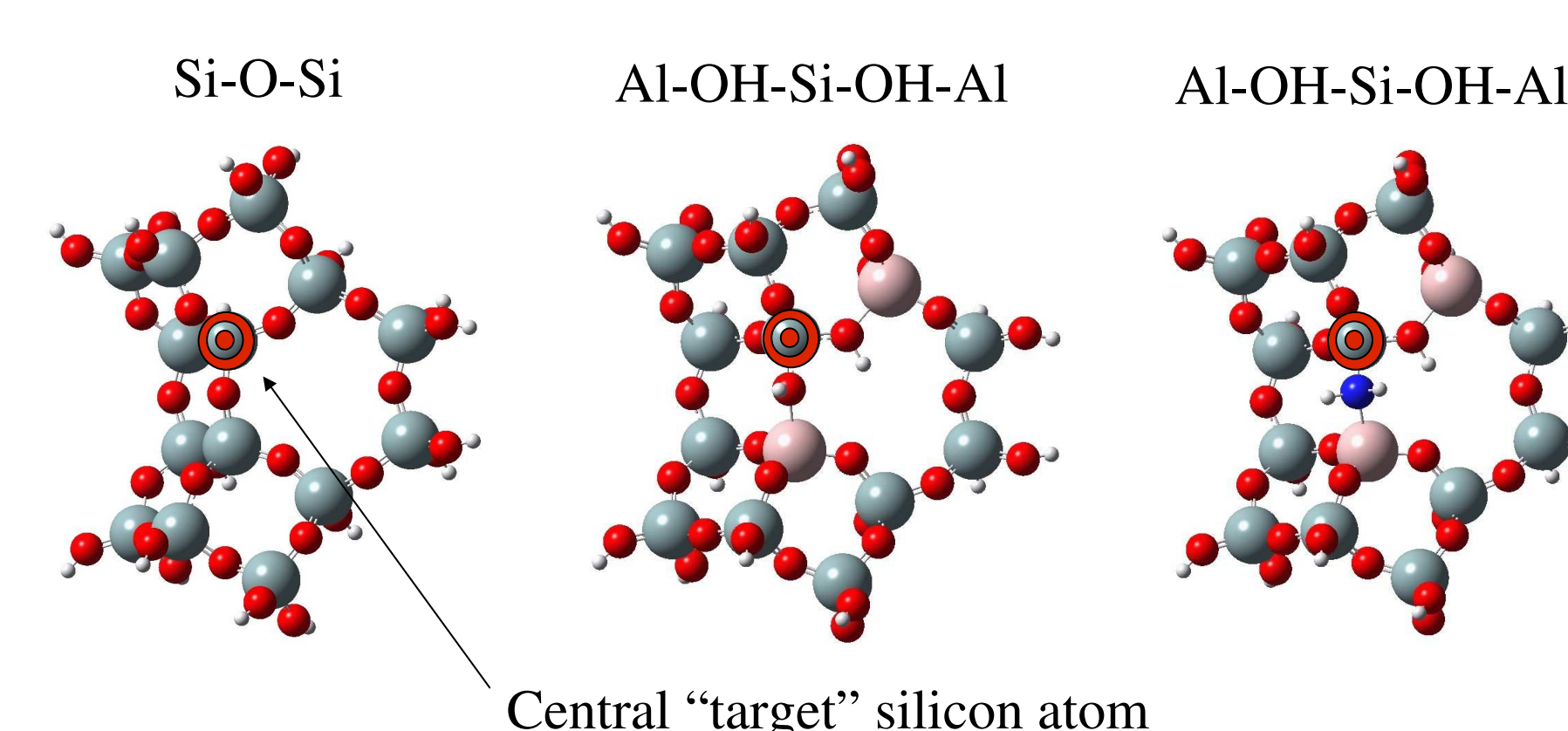
Experimental Characterization: Infrared



Experimental Characterization: NMR



Cluster Calculations



- Things to vary from cluster to cluster:
- Aluminum atoms nearby
 - Placement of hydrogen
 - Number of nearby nitrogen vs. oxygen atoms

Chemical Shift Calculations

$$\Delta E = \hbar \gamma \Delta I (1 - \sigma) B$$

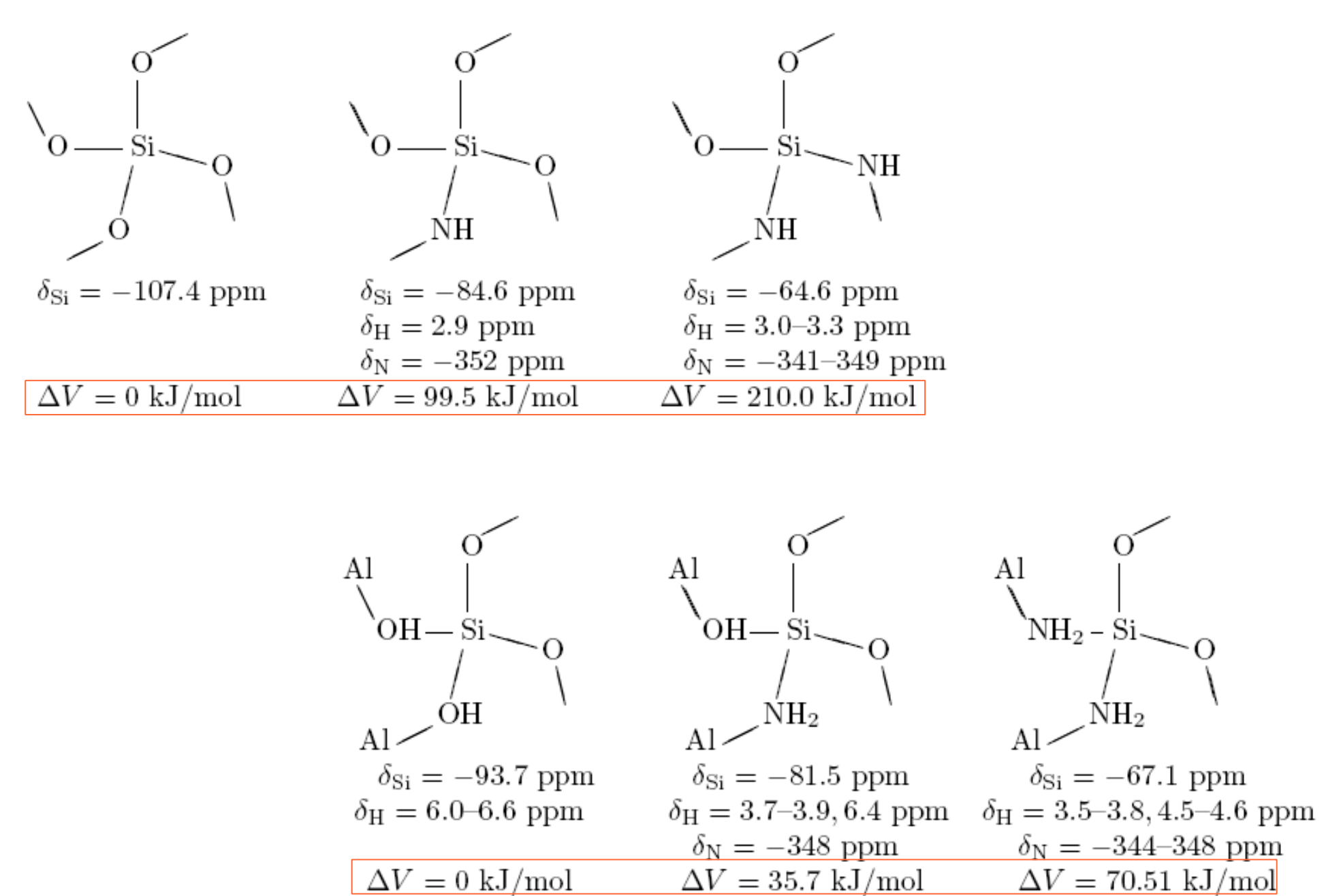
Change in magnetic moment (constant for each nucleus)

External magnetic field (constant)

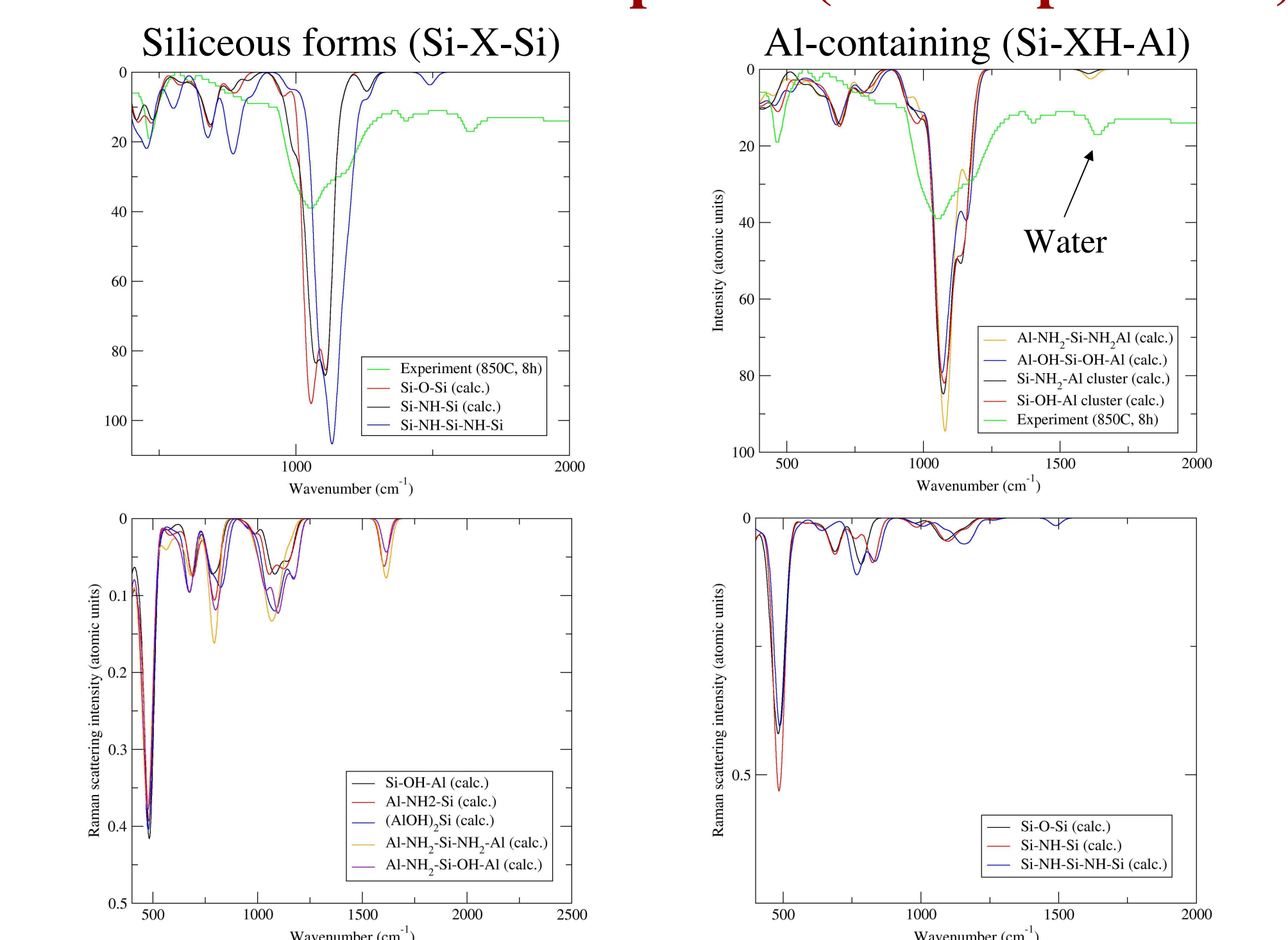
Shielding constant (function of electron density)

$$\delta = \frac{\Delta E - \Delta E_{\text{ref}}}{\Delta E_{\text{ref}}} = \frac{V - V_{\text{ref}}}{V_{\text{ref}}} = \frac{\sigma_{\text{ref}} - \sigma}{1 - \sigma_{\text{ref}}} \approx \sigma_{\text{ref}} - \sigma$$

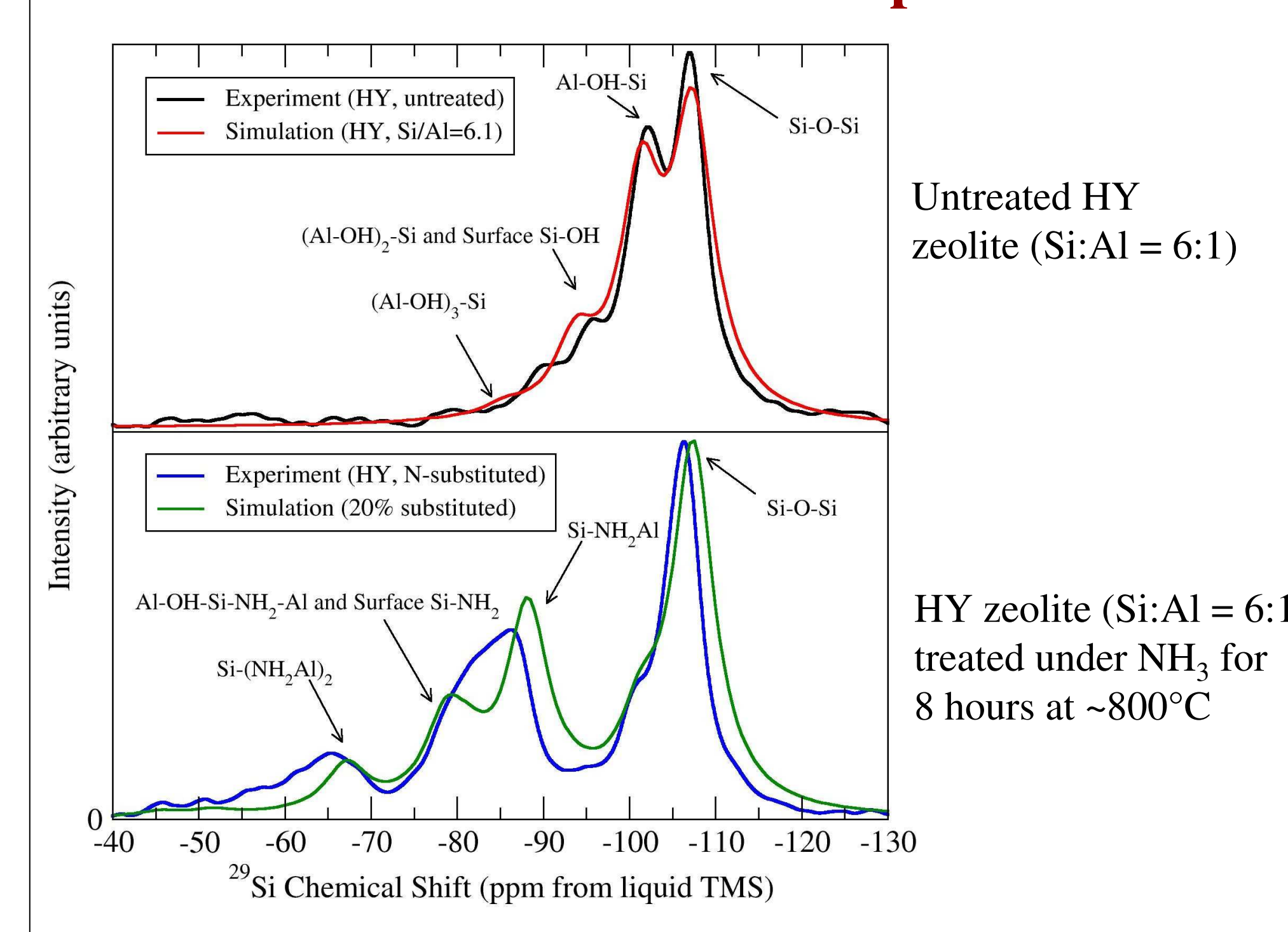
Calculated Energies of Amine Substitutions



Calculated Infrared Spectra (with Experiment)



Calculated ²⁹Si NMR Spectrum



Conclusions and Future Work

- Nitrogen prefers to substitute between Si and Al, not between Si and Si
- Approximately 20% of total oxygens are substituted for nitrogen after 8 hours of treatment. Of those, 87% take place inside the framework
- Current/Future Work:
 - High-resolution physical adsorption
 - Stability of treated zeolites
 - Extraframework aluminum simulations
 - Non-silicon NMR spectroscopy