

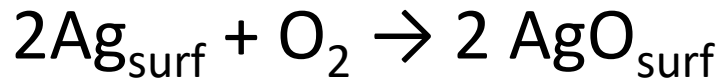
Chemisorption

- Pulse titration

- **Homework**

- Silver (2.2 wt% on silica)

- Surface reaction:



@T = 170 °C (no bulk oxidation!)

- $m_{\text{sample}} = 0.5021 \text{ g}$

- $V_{\text{pulse}} = 0.00925 \text{ cm}^3 \text{ O}_2 @ \text{RT} @ 1 \text{ atm}$

- Max TCD signal = 0.022

- D = ?

| Pulse | TCD signal |
|-------|------------|
| #1 | 0 |
| #2 | 0 |
| #3 | 0 |
| #4 | 0 |
| #5 | 0 |
| ... | ... |
| #28 | 0 |
| #29 | 0 |
| #30 | 0.002 |
| #31 | 0.005 |
| #32 | 0.011 |
| #33 | 0.016 |
| #34 | 0.019 |
| #35 | 0.021 |
| #36 | 0.022 |

Heterogeneous catalysis (C9981)

Lecture 6

Catalysts characterization - continuation

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Catalyst characterization

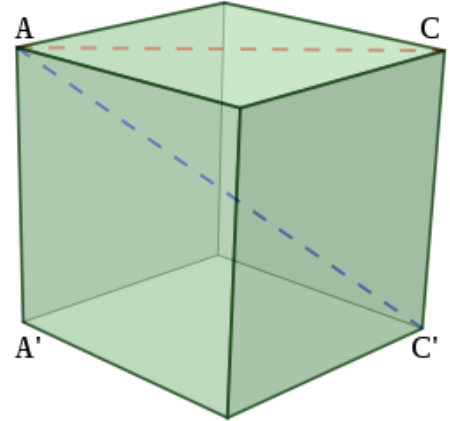
- Outline
 - Solid state NMR techniques
 - **Hydrophilicity vs. Hydrophobicity meas**
 - Water sorption
 - Dynamic water sorption
 - Inverse gas chromatography
 - Microcalorimetry
 - Quartz crystal microbalance
 - **In situ and operando techniques**

Solid state NMR techniques

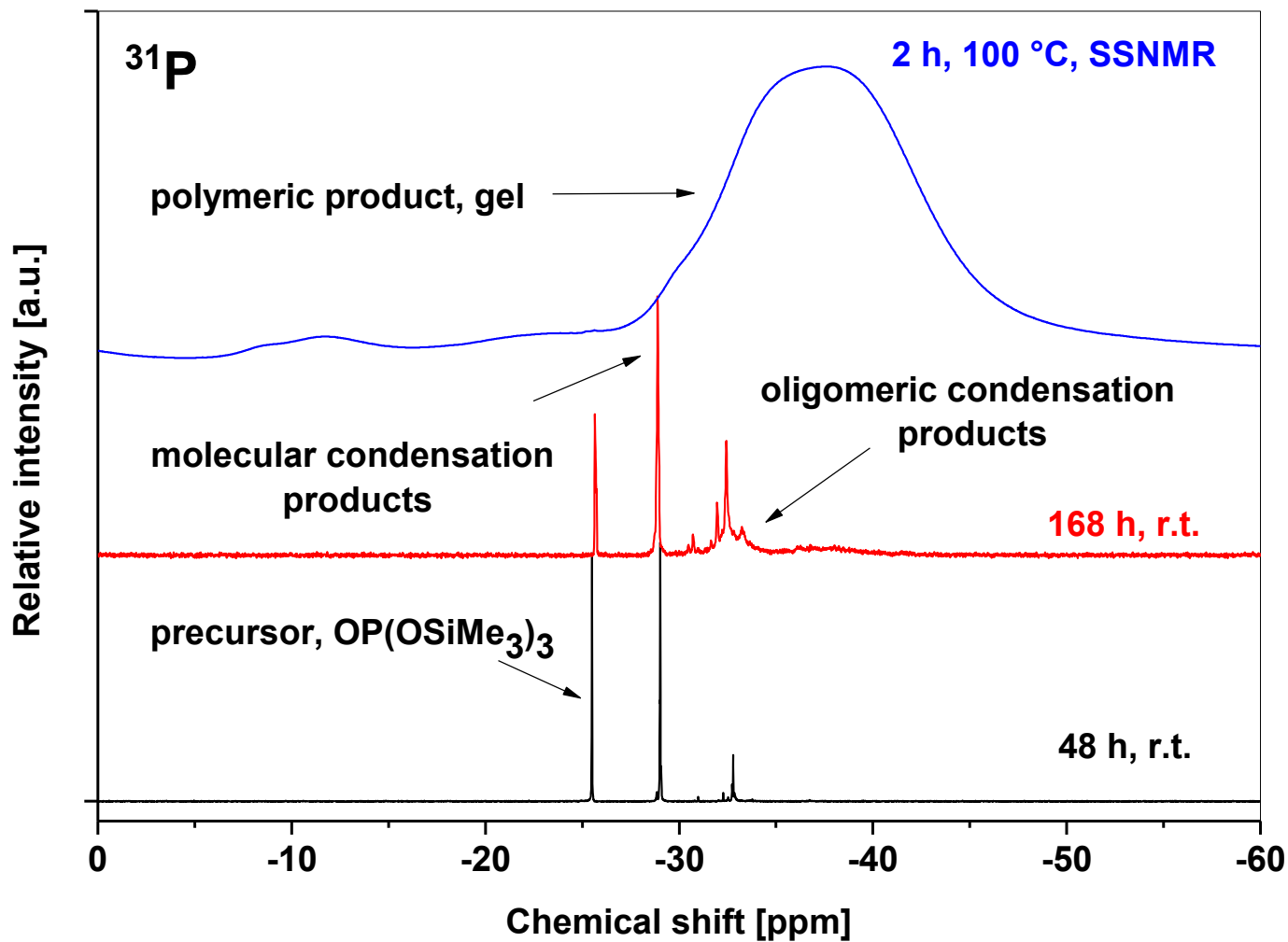
- Magic angle spinning (MAS)
- Multiple quantum experiments (MQ)
- Dynamic nuclear polarization (DNP)

Solid state NMR techniques

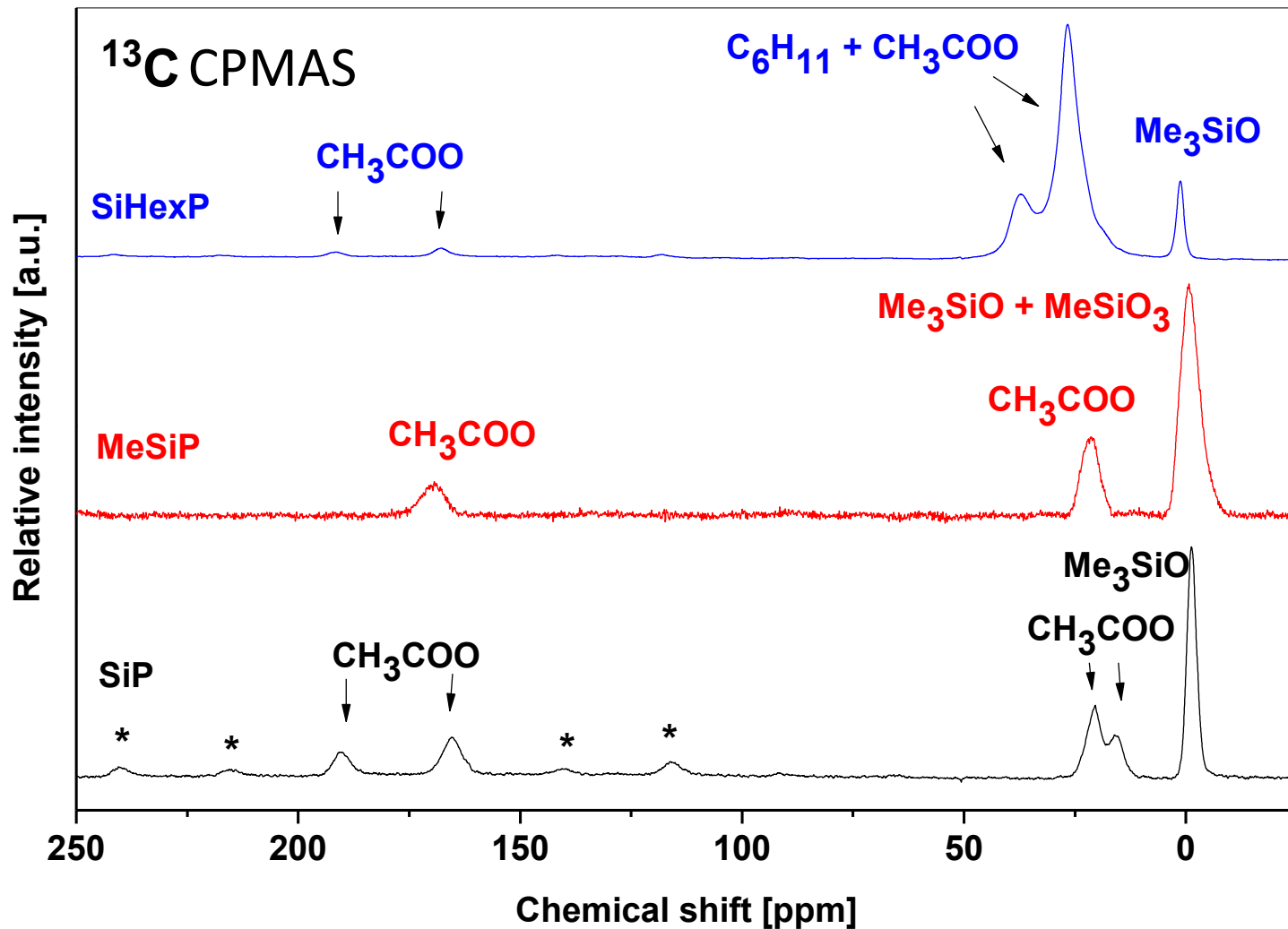
- Magic angle spinning (MAS)
 - 54.74°
 - Spinning averages dipolar interaction, chemical shift anisotropy, and quadrupolar interaction
 - “Residues” of these interactions are observed as spinning sidebands
 - Static and double rotation NMR possible



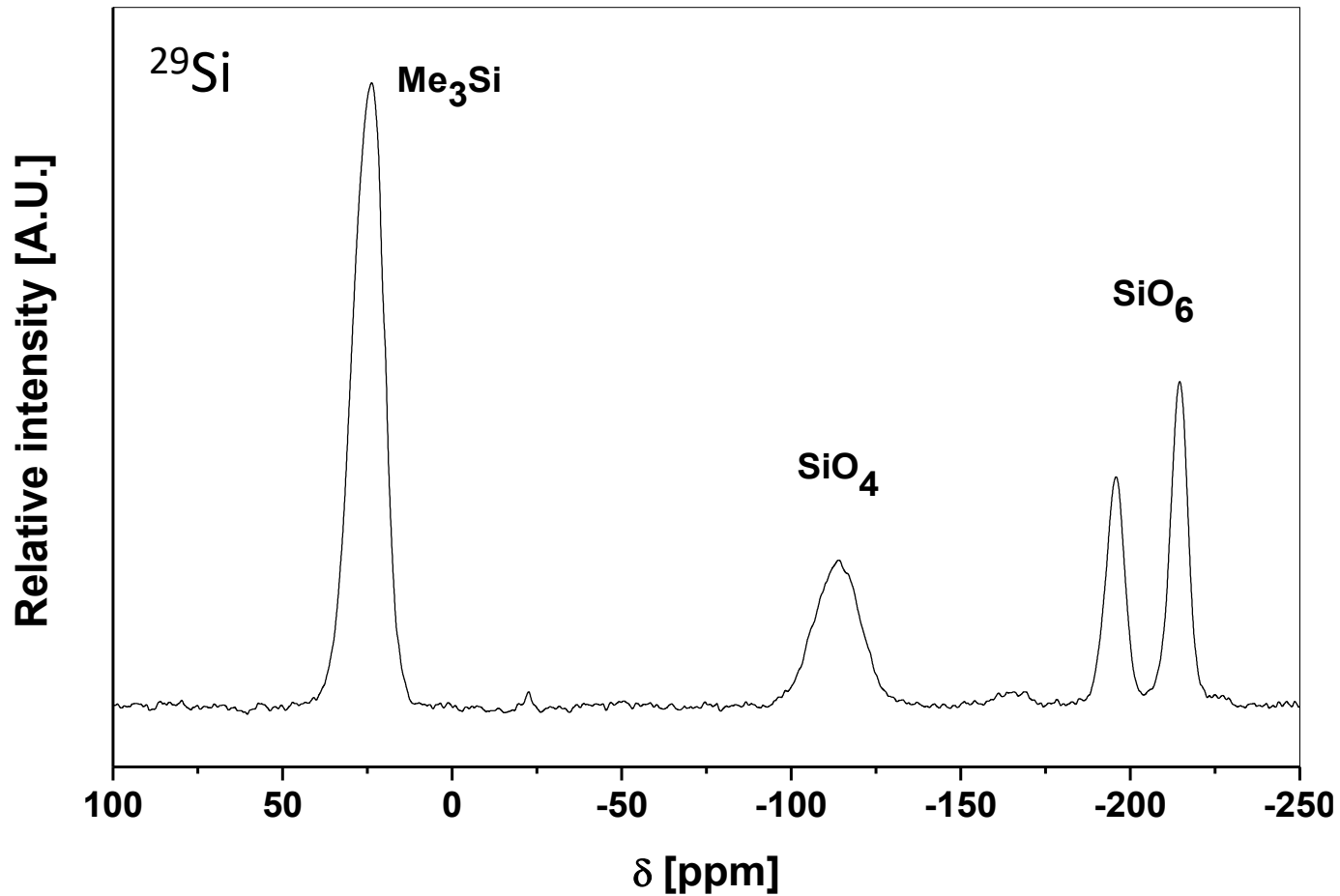
Solid state NMR techniques



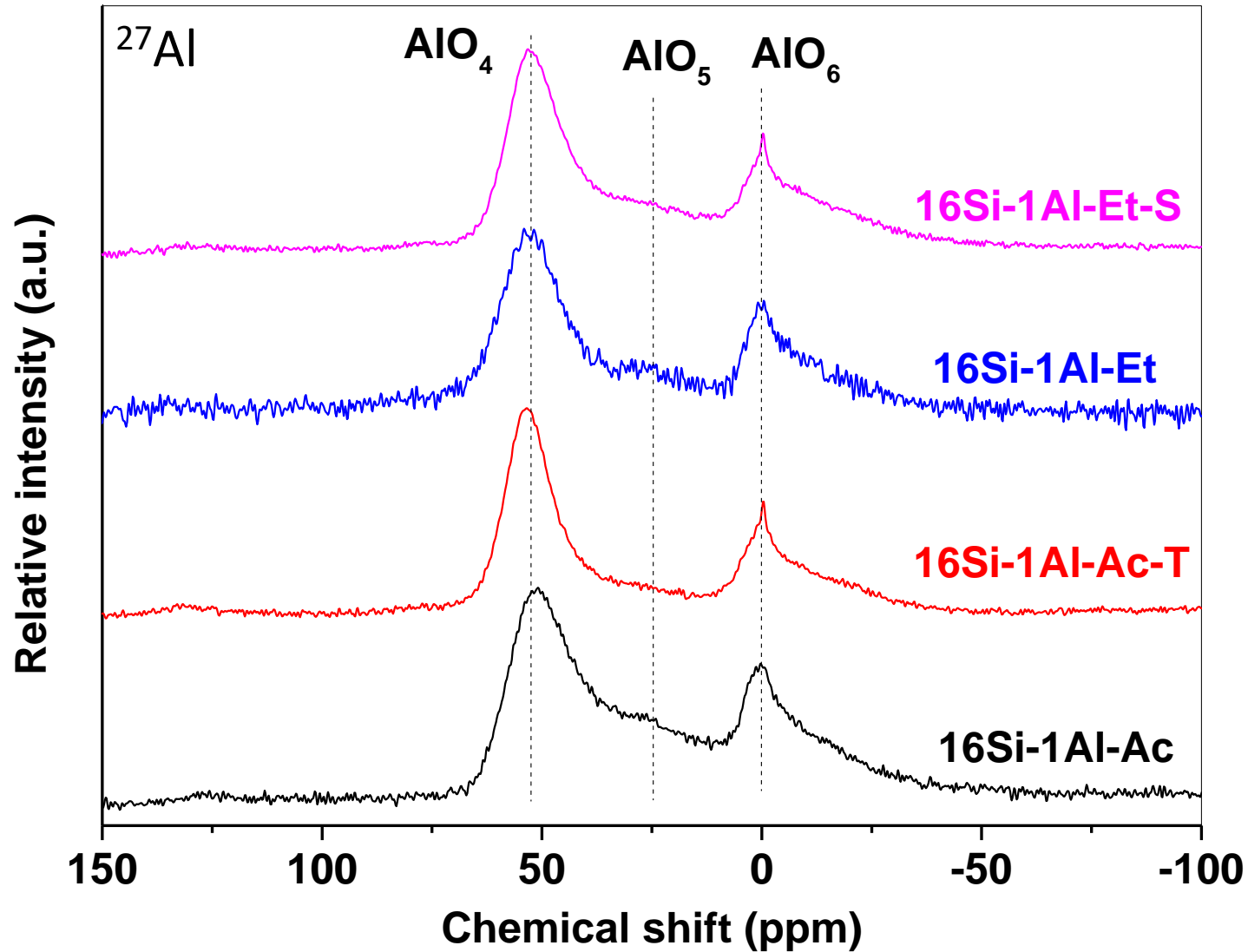
Solid state NMR techniques



Solid state NMR techniques

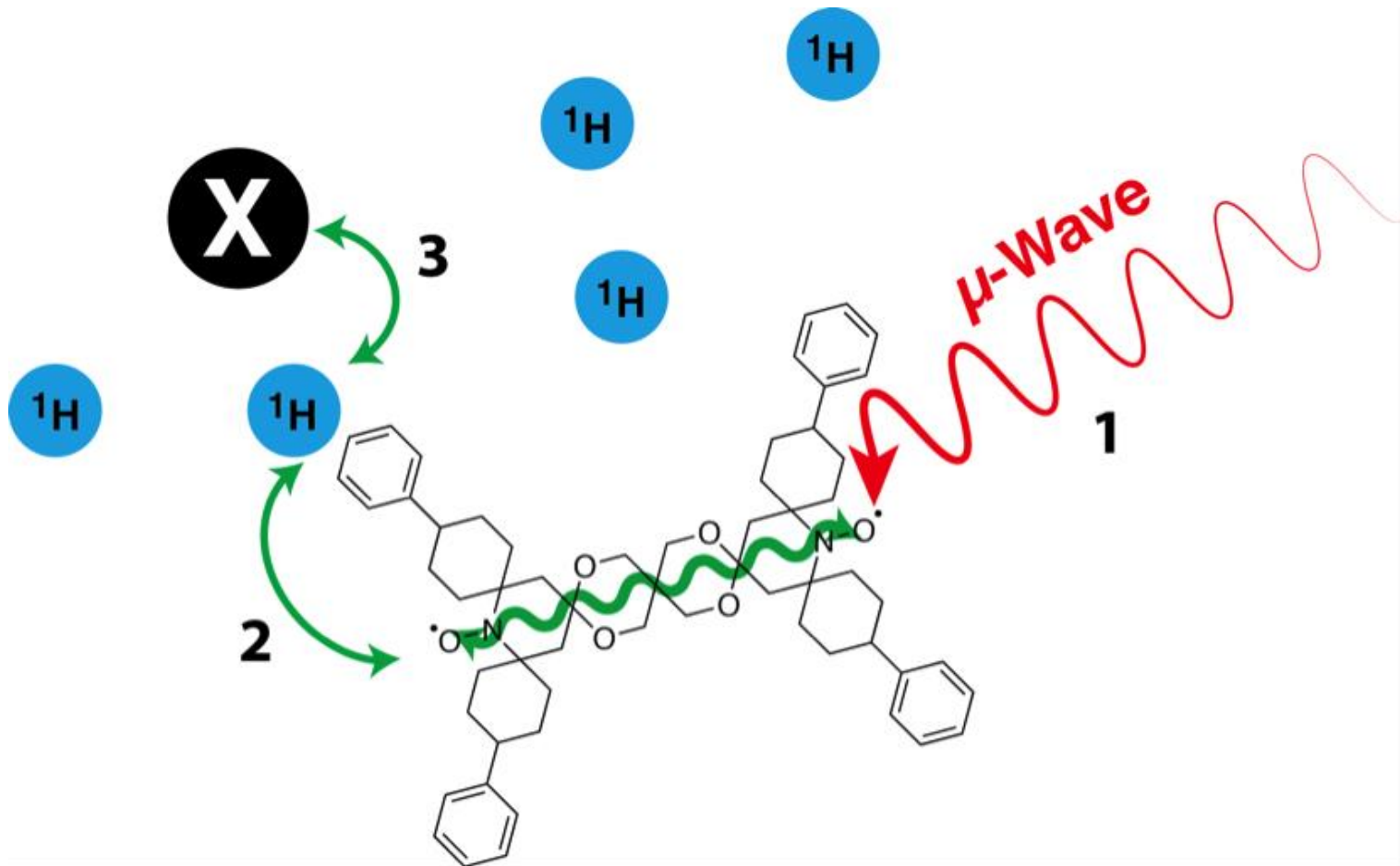


Solid state NMR techniques



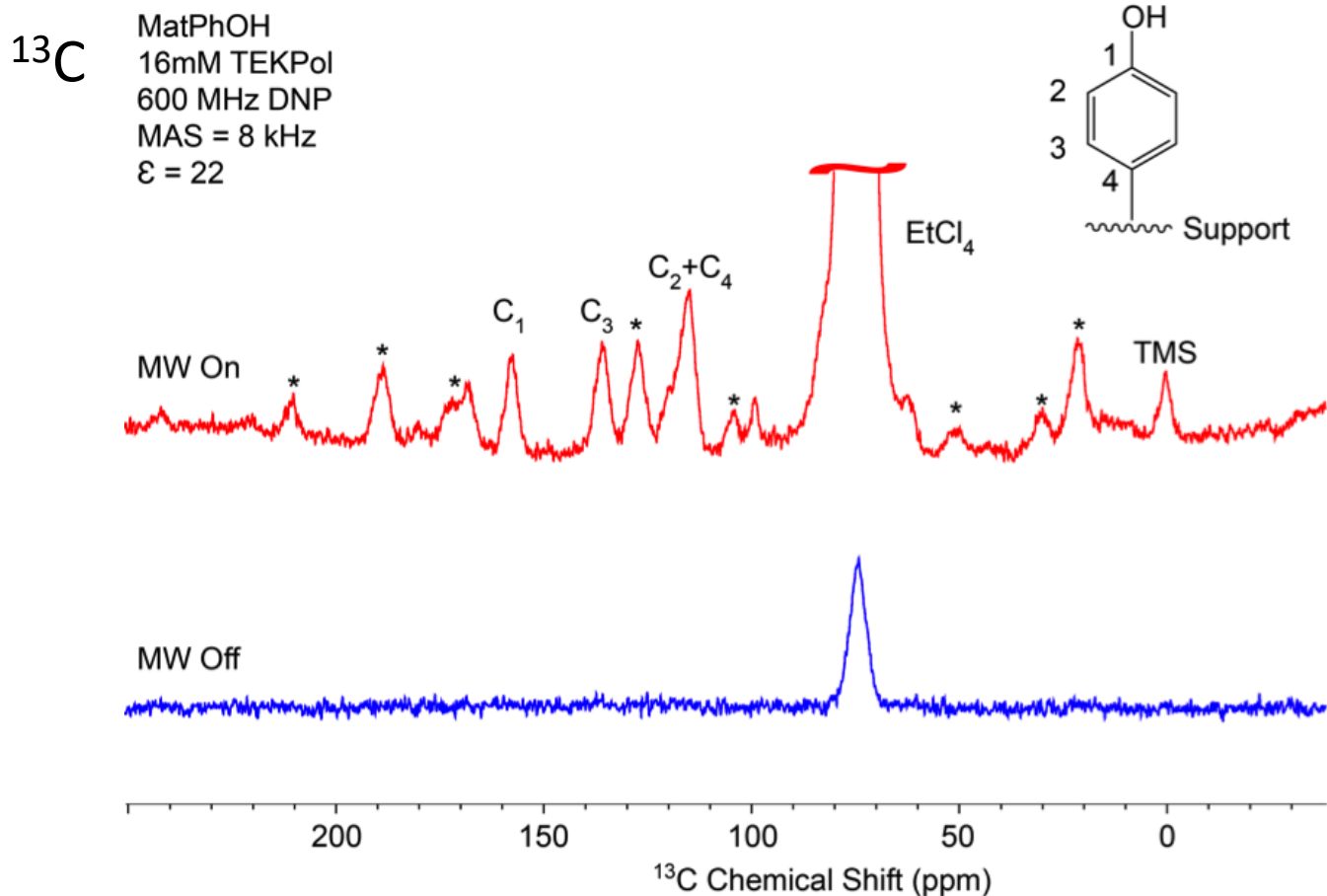
Solid state NMR techniques

- Dynamic nuclear polarization (DNP)



Solid state NMR techniques

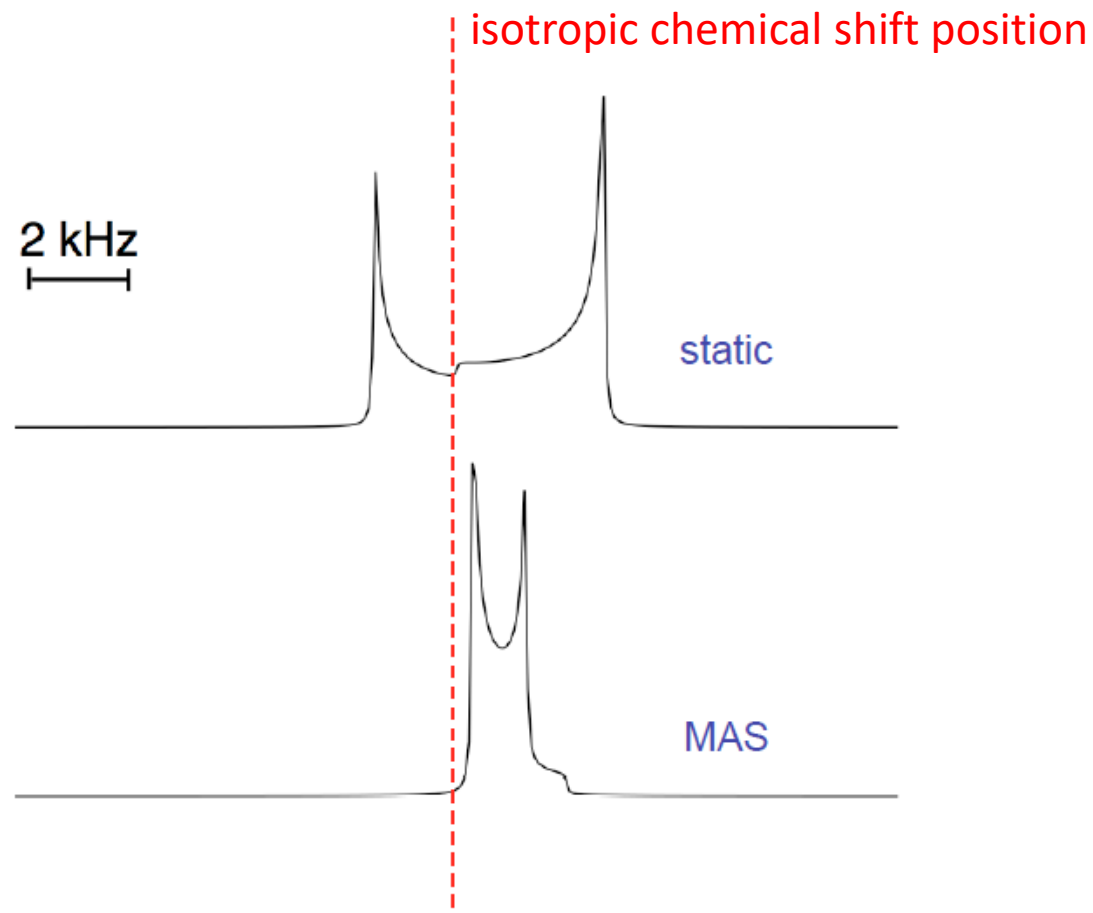
- Dynamic nuclear polarization (DNP)



Solid state NMR techniques

- Static NMR (=no spinning)

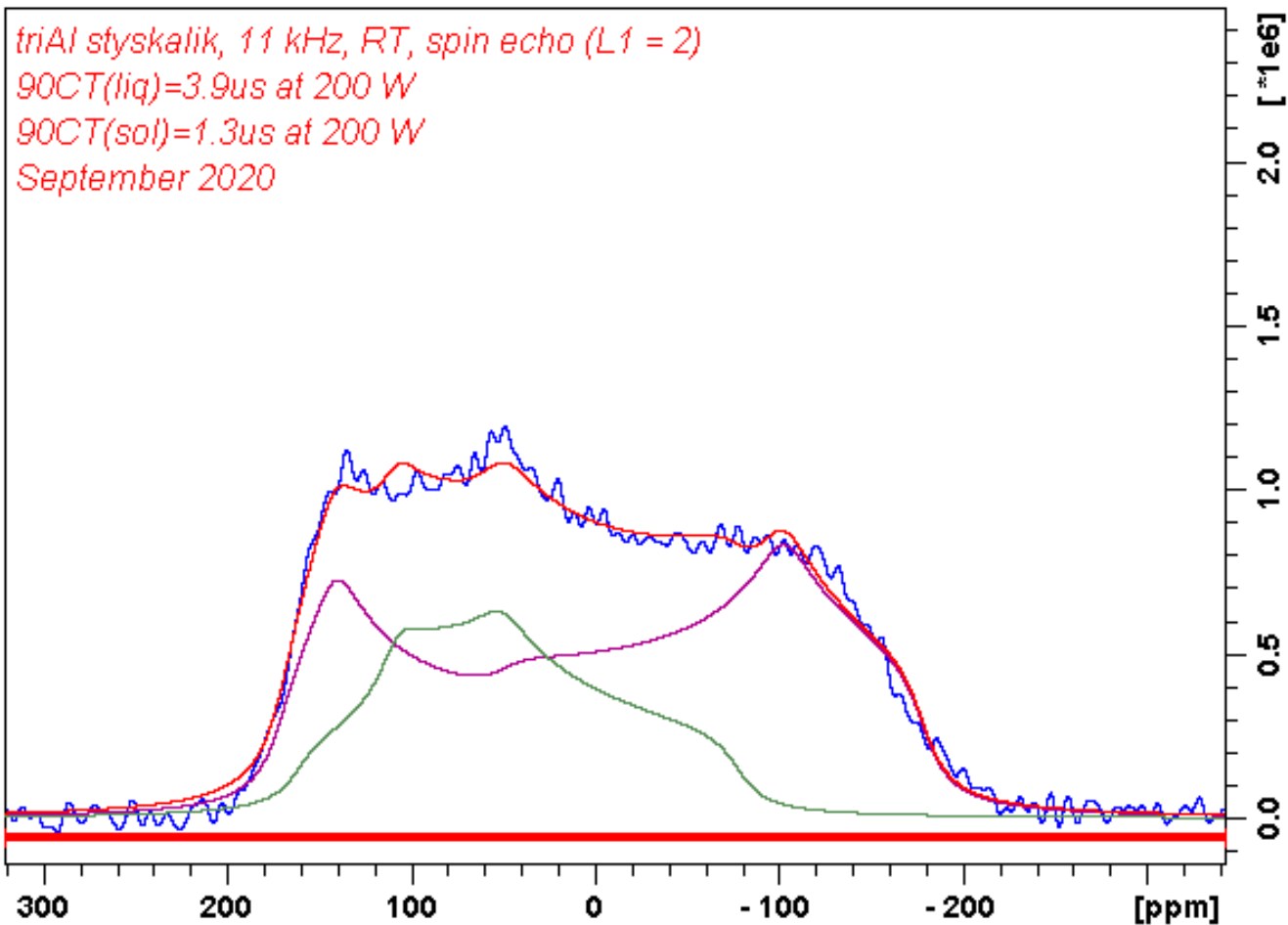
^{27}Al



Solid state NMR techniques

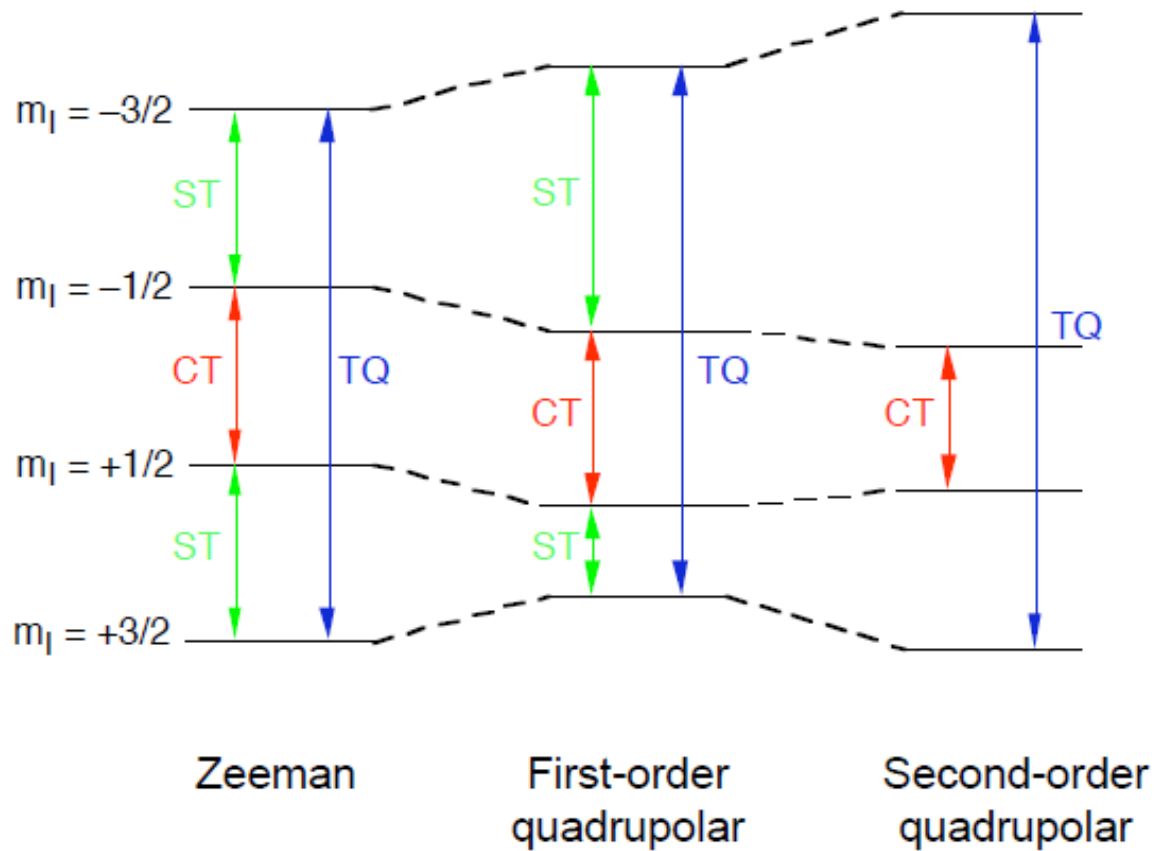
- Static NMR (=no spinning)

^{27}Al



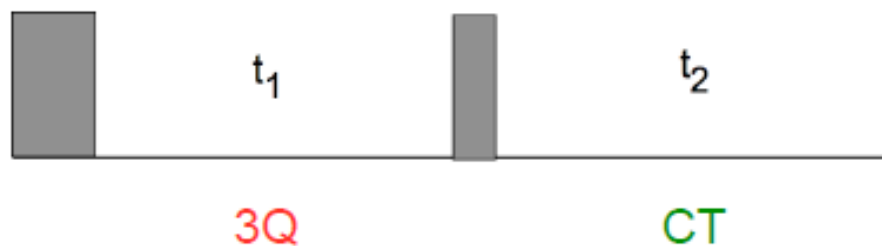
Solid state NMR techniques

- Multiple quantum MAS (MQMAS)



Solid state NMR techniques

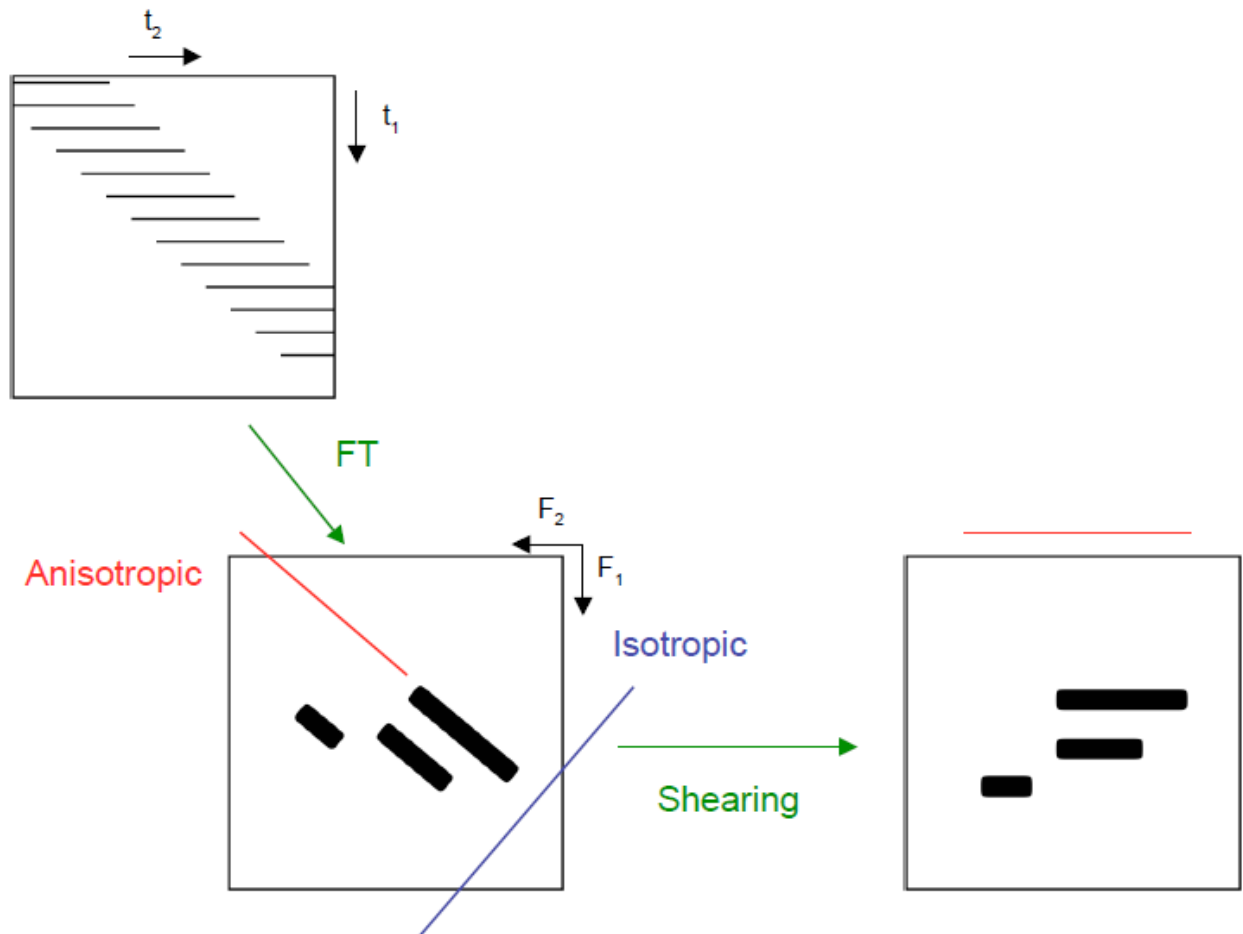
- Multiple quantum MAS (MQMAS)



Frydman et al., J. Am. Chem. Soc. **117**, 5367 (1995)

Solid state NMR techniques

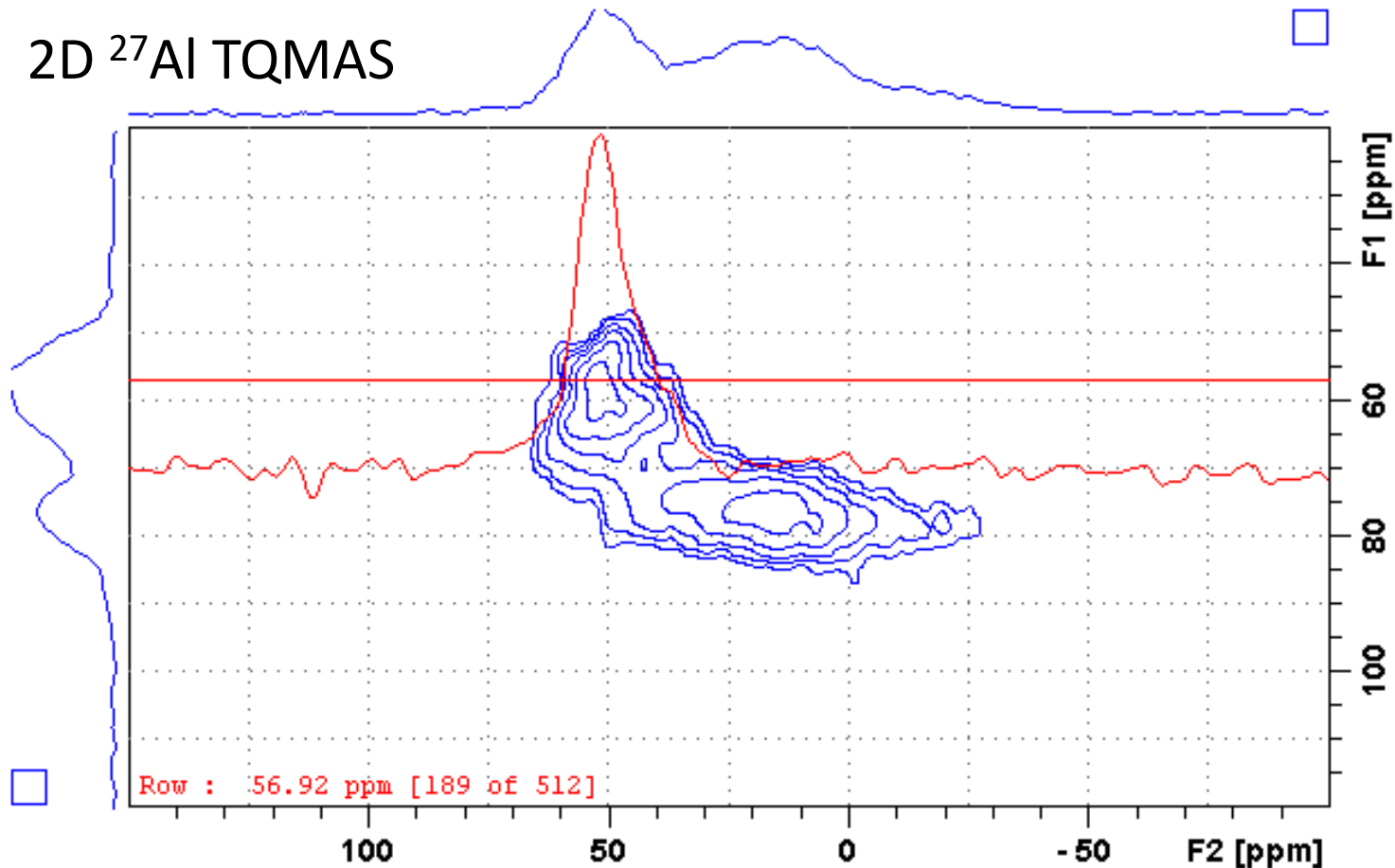
- Multiple quantum MAS (MQMAS)



Solid state NMR techniques

- Multiple quantum MAS (MQMAS)

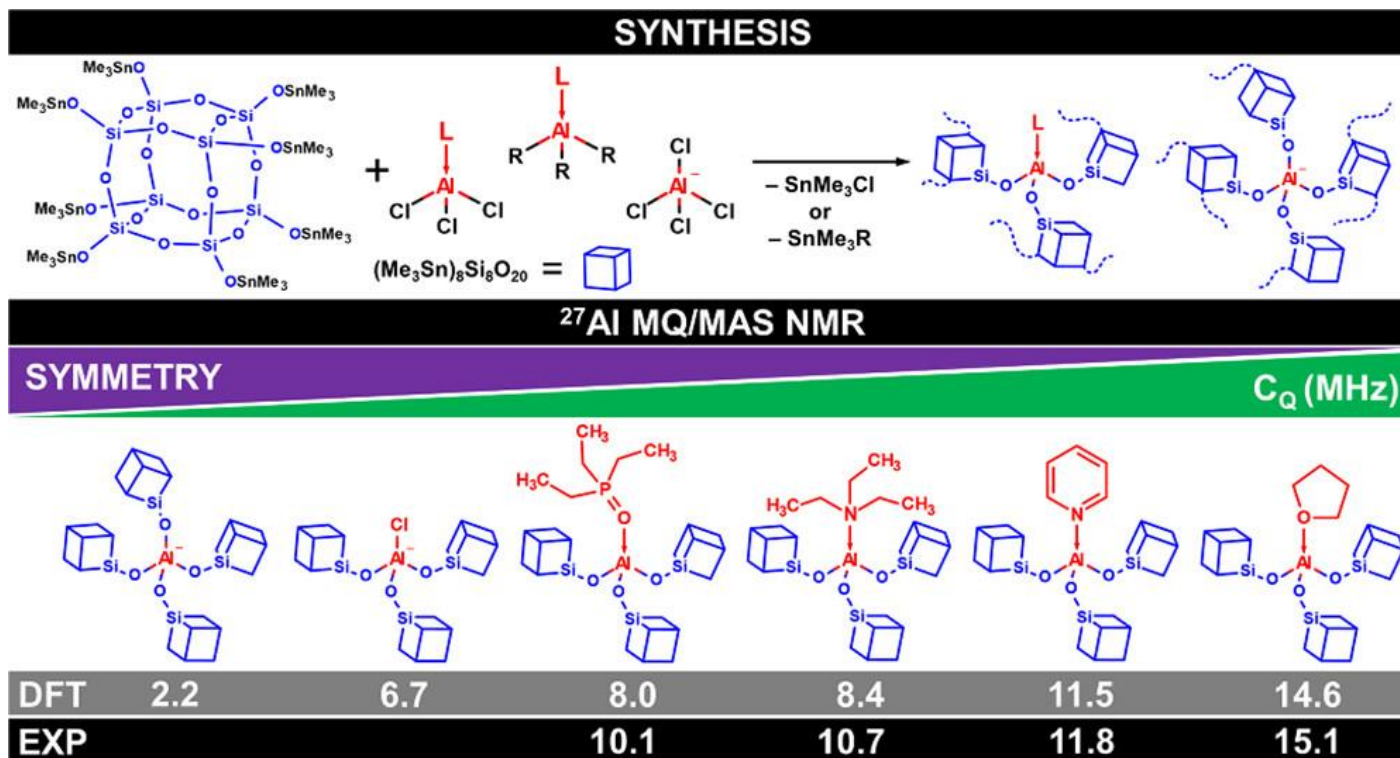
2D ^{27}Al TQMAS



Solid state NMR techniques

- Multiple quantum MAS (MQMAS)

2D ^{27}Al TQMAS



Hydrophilicity vs. Hydrophobicity

- Why?
 - Remember: Adsorption/desorption step in cata as important as cata rxn itself (@Lecture 1)
 - Moreover: strong efforts nowadays put forward bio-sources instead of fossil fuels
 - Compare
 - Oil – long hydrocarbon chains (hydrophobic)
 - Wood – cellulose, lignin = sugar based materials = a lot of oxygen, OH groups (hydrophilic)

Hydrophilicity vs. Hydrophobicity

- Possibilities:
 - Water sorption
 - Dynamic vapor sorption/Quartz crystal microbalance
 - Inverse gas chromatography
 - Microcalorimetry

Hydrophilicity vs. Hydrophobicity

- Problems:
 - What is a measure of hydrophobicity?
 - A material can have high/low affinity to both water and organic molecules (i.e. if a sample is hydrophilic, it does not necessarily mean it is hydrophobic!)
 - % of pore volume filled with water at certain p/p_0
 - Hydrophobic index (water/toluene competitive sorption)
 - Heat of adsorption, heat of immersion
 - ...
 - Chemisorption: Do chemisorbed molecules account for hydrophobicity/philicity?

Hydrophilicity vs. Hydrophobicity

- Water sorption
 - Similar to N_2 physisorption
 - Known volume and pressure in the cell, known mass of the sample

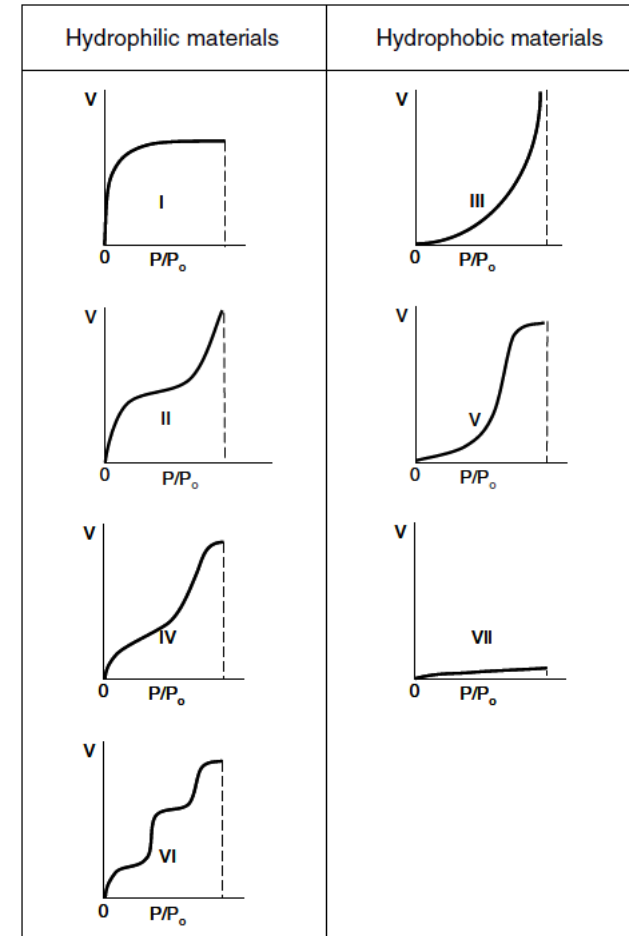
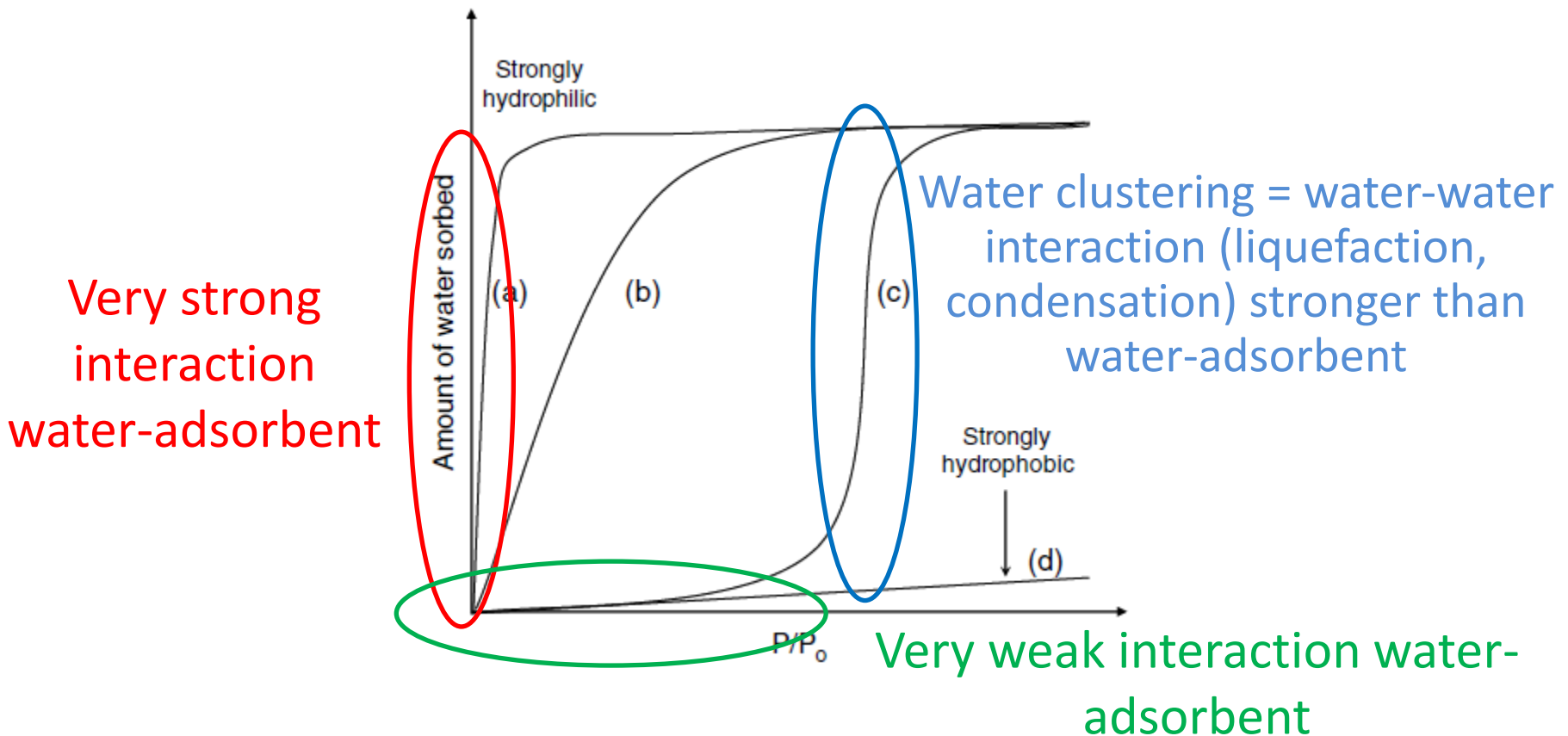


Fig. 1. Adsorption isotherms classified according to IUPAC: type I: very hydrophilic material, type II: hydrophilic material, type III: hydrophobic/low hydrophilic material with weak sorbent-water interactions, type IV: hydrophilic material, type V: hydrophobic/low hydrophilic material with weak sorbent-water interactions, type VI: hydrophilic material with multiple sorbent-water interactions and stepwise sorption, type VII: very hydrophobic material.

Hydrophilicity vs. Hydrophobicity

- Water sorption



Hydrophilicity vs. Hydrophobicity

- Water sorption
 - Plotting $\ln p$ against $1/T$ at constant adsorption uptake gives a straight line with a slope equal to H_{iso}/R
 - H_{iso} = isosteric heat of adsorption
 - Isotherms at multiple temperatures needed!

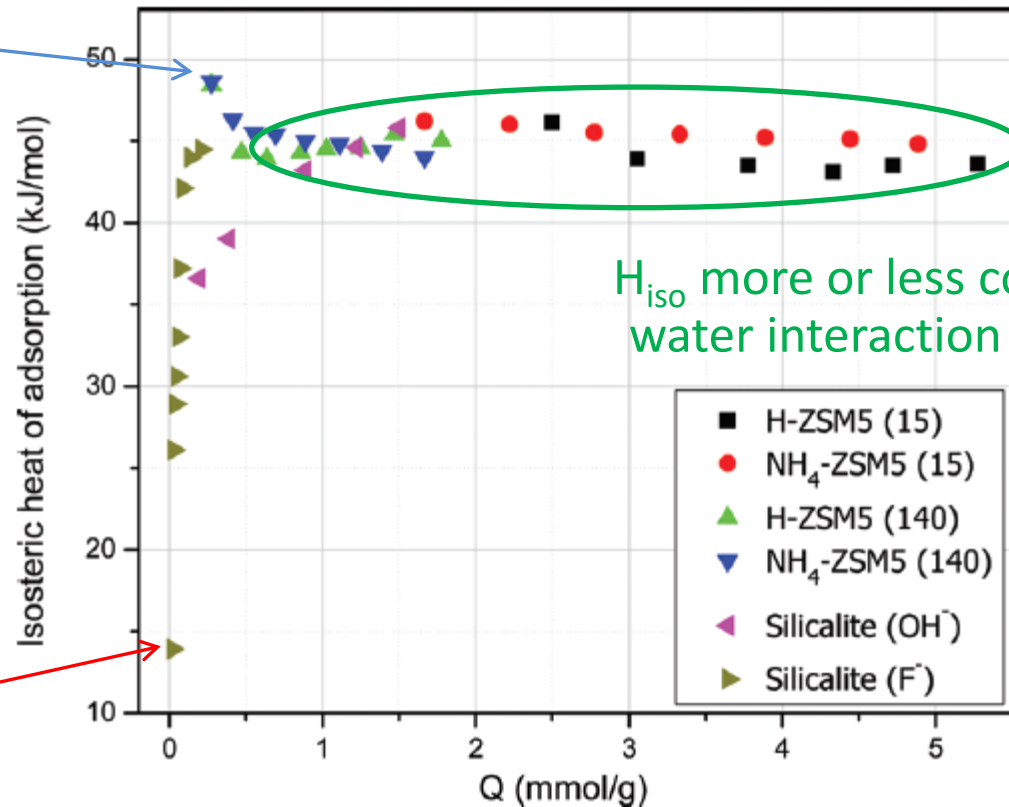
Hydrophilicity vs. Hydrophobicity

- Water sorption

- H_{iso} = isosteric heat of adsorption

Hydrophilic material = very high H_{iso} = strong interaction water-adsorbent

Hydrophobic material = very low H_{iso} = negligible interaction water-adsorbent



H_{iso} more or less constant = water-water interaction (condensation)

Hydrophilicity vs. Hydrophobicity

- Dynamic vapor sorption/Quartz crystal microbalance
 - You deposit your material on an accurate microbalance
 - You expose it to vapors of different gases/liquids (water, alcohols, hydrocarbons,...)
 - You follow the uptake by changes of mass
 - If we can control/follow pressure, then isotherms can be obtained similar to a classic physisorption

Hydrophilicity vs. Hydrophobicity

- Inverse gas chromatography
 - You pack column (≈ 50 cm) with the material you want to test (≈ 0.5 g)
 - You inject series of gases/liquids (e.g. methane, ethane,...hexane, heptane; methanol, ethanol,...; benzene, toluene, xylene...)
 - You follow retention time (you directly see „affinity“ of your material to selected liquids)
 - Models (math) can give surface energy,...

Hydrophilicity vs. Hydrophobicity

- Immersion microcalorimetry
 - Evacuated sample sealed in a bulb with brittle end
 - Bulb immersed in a testing liquid, sealed
 - Bulb broken (rod pushed down)
 - Liquid gets into the bulb, adsorbs, heat of immersion released and measured

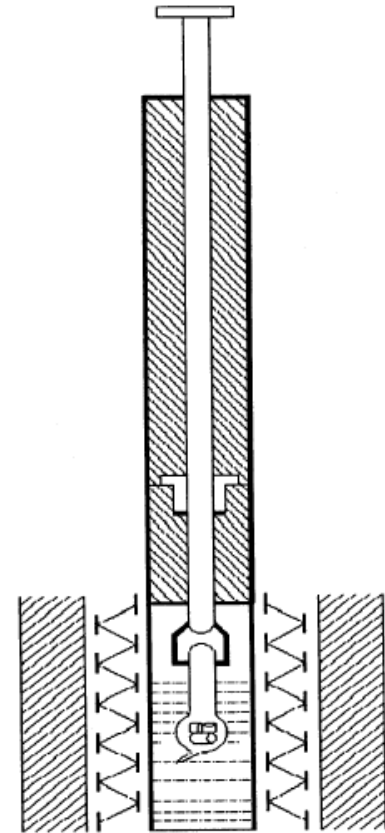


Fig. 1. Set-up for immersion calorimetry.

Hydrophilicity vs. Hydrophobicity

- Immersion microcalorimetry
 - Ti-MCM-41, pure inorganic vs. increasing degree of surface silylation (increasing carbon content)

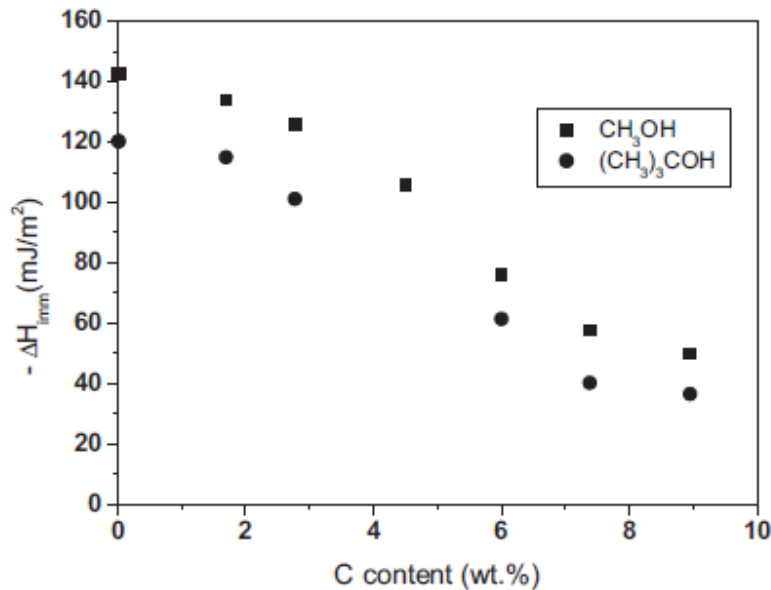


Fig. 7. Evolution of the areal enthalpy of immersion (mJ/m^2) in methanol and 2-methyl-2-propanol as a function of carbon content for the different silylated Ti-MCM-41 samples.

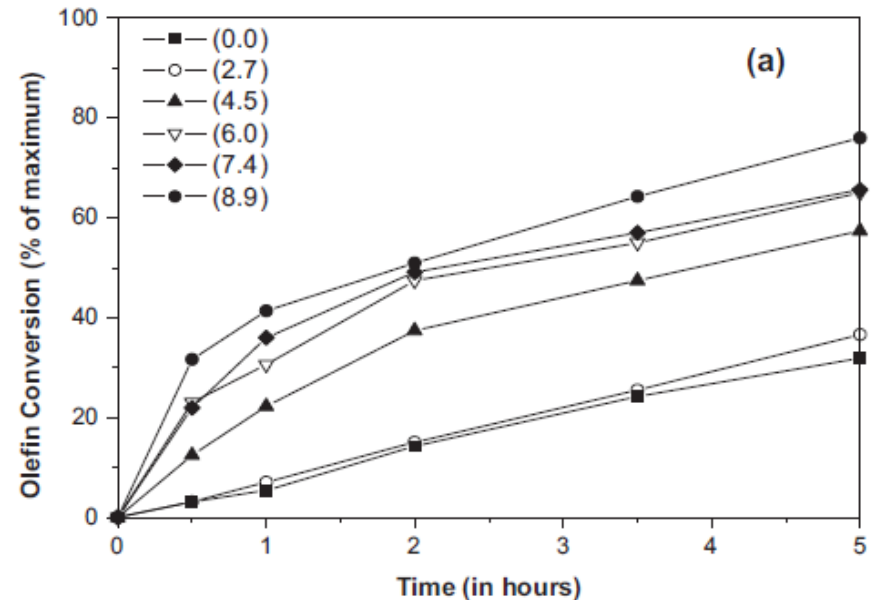


Fig. 8. Catalytic activity of Ti-MCM-41 materials with different silylation degrees in the cyclohexene epoxidation with TBHP (and 0.5 wt% of catalyst) at 333 K during 5 h. (a) Cyclohexene Conversion (%Mol.); (b) Epoxide Selectivity (%Mol.).

In situ and operando techniques

- In situ = online analysis of a working catalyst
- Operando = online analysis of a working catalyst at relevant conditions (p, T, WHSV)

| | Technique | Information | Suitable to <i>operando</i> charac. |
|---------|---|---|---|
| v → v | IR spectroscopy (FTIR/DRIFTS/ATR/IRAS) | Adsorbate nature, adsorption site | ✓ |
| | Raman spectroscopy (SERS/TERS) | Solid structure, adsorbate nature | ✓ |
| | XAS (EXAFS/XANES/QXAS) | Local environment, oxidation state | ✓ |
| | SAXS (GISAXS) | NP size and morphology | ✓ |
| | XRD (PXRD/SXRD/HEXRD) | Crystal phase and dimension | ✓ |
| v → e- | XPS (NAP-XPS) | Chemical composition, oxidation state | Low pressure, large volume |
| e- → e- | TEM (Environmental TEM) (<i>In situ</i> TEM) | Atomic structure, chemical distribution | Low pressure, large volume Low catalyst amount |
| | SPM (STM, AFM...) | Surface structure | Planar model catalysts |

In situ and operando techniques

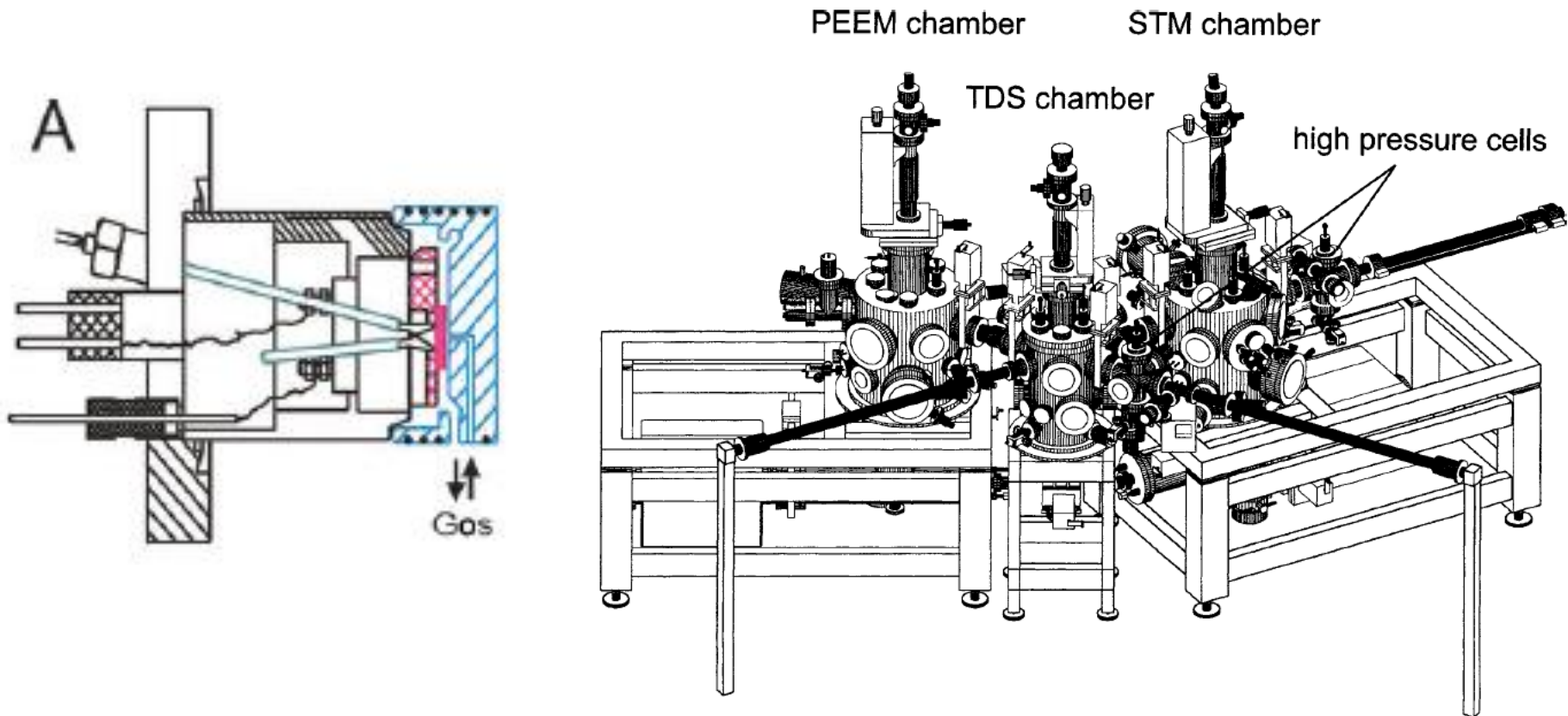
- We already know these techniques, let's look at the examples!
 - Low energy electron diffraction (LEED, gives similar results to x-ray diffraction = analysis of crystal structures)
 - X-ray absorption near edge structure (XANES)
 - Diffusive reflectance infrared Fourier transform spectroscopy (DRIFTS)

In situ and operando techniques

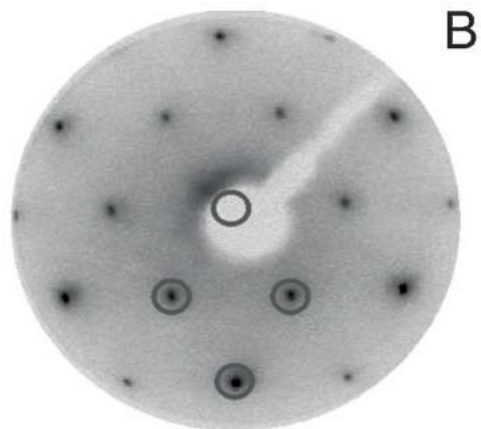
- Example: LEED – low energy electron diffraction
- Ethylbenzene dehydrogenation to styrene
 - 600 °C, 1 atm, 10-fold excess water vapor
 - Over Fe₂O₃ epitaxially grown on Pt(111)
- Flow reactor located in a high pressure cell
 - Heated by lasers
 - GC-MS analysis of catalytic products
 - LEED analysis enabled by gate valve (high pressure/ultra high vacuum) and sapphire window

In situ and operando techniques

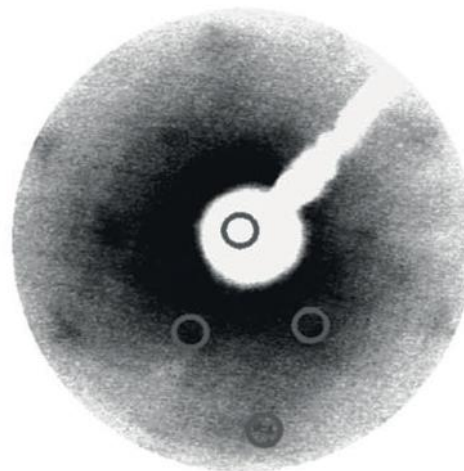
- Example: LEED – low energy electron diffraction



Before catalytic reaction

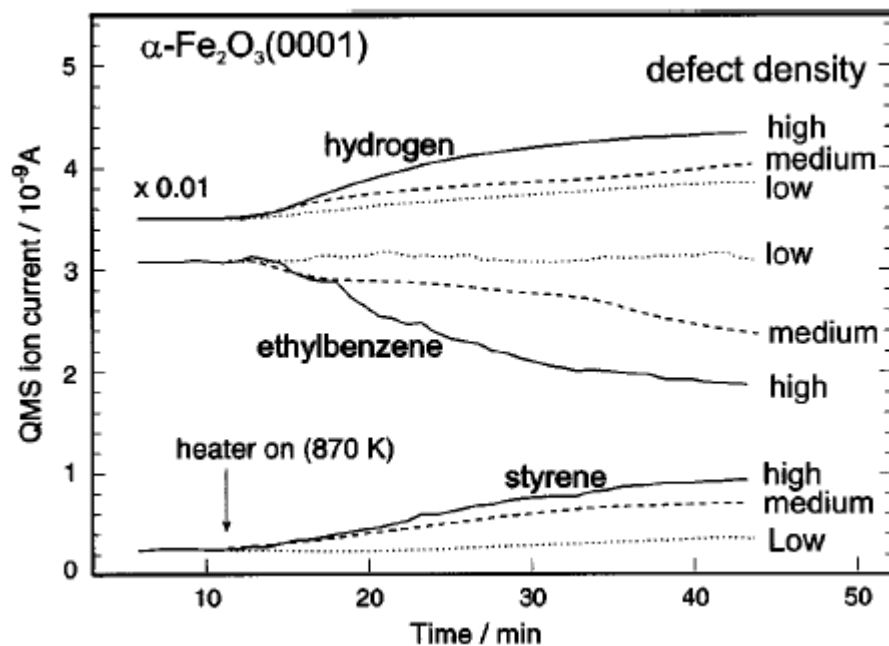


After catalytic reaction



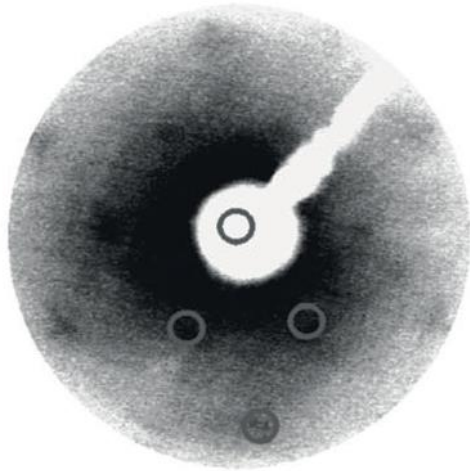
Initial period, no catalytic activity

Working catalyst



In situ and operando techniques

- Example: LEED – low energy electron diffraction



Active catalyst

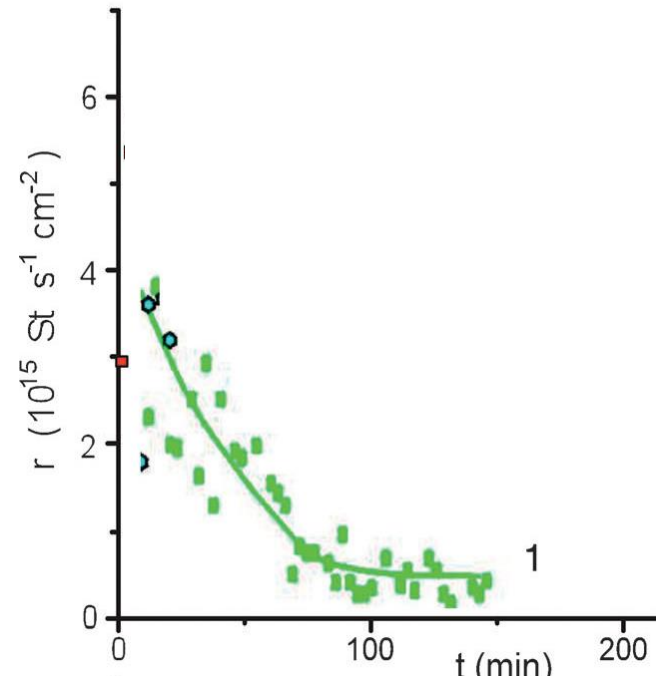
Non-crystalline Fe_2O_3

Metastable (Reduction!)

Carbon deposition (Styrene!)

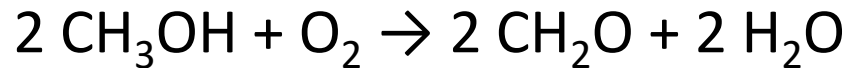
Loss of catalytic activity

0.5 eq O_2 addition



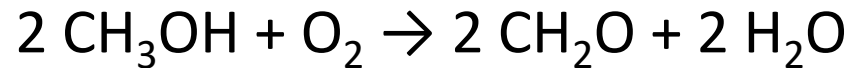
In situ and operando techniques

- Example: XANES – x-ray absorption near edge structure
- Methanol oxidation to formaldehyde over Cu
 - 25–450 °C, 1 mbar
 - Cu in the form of polycrystalline foil

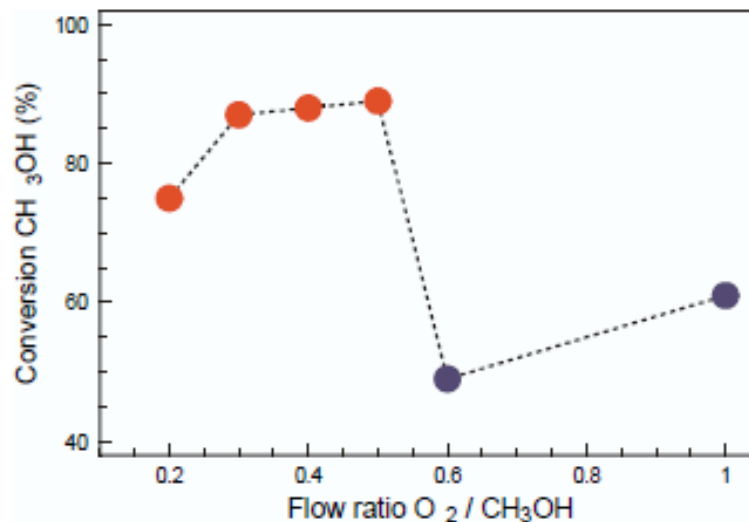


In situ and operando techniques

- Example: XANES – x-ray absorption near edge structure

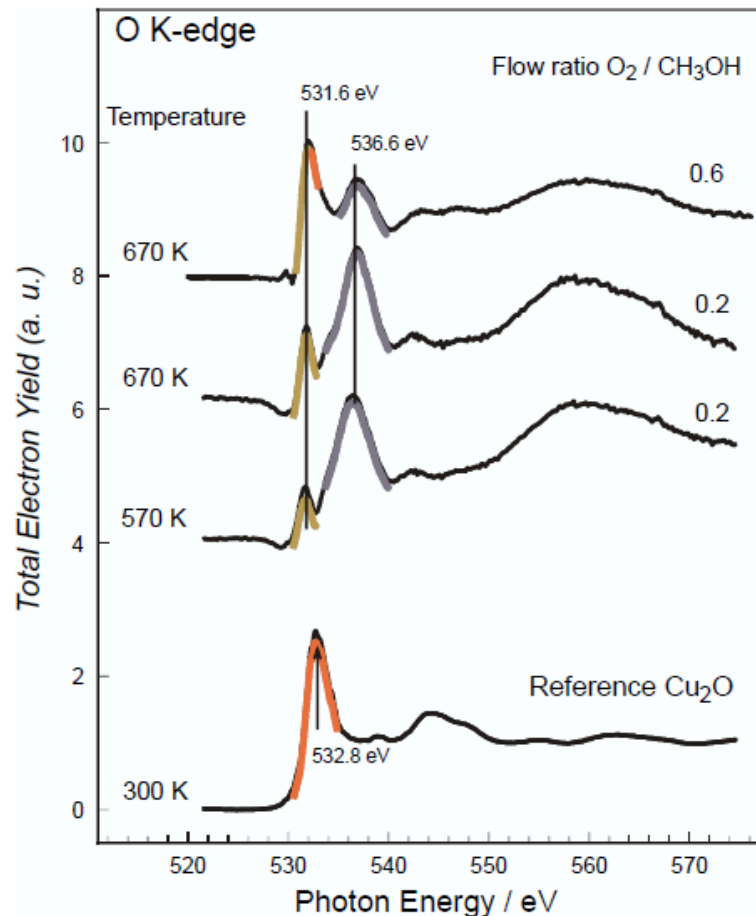


Catalytic Activity



In situ and operando techniques

- Example: XANES – x-ray absorption near edge structure



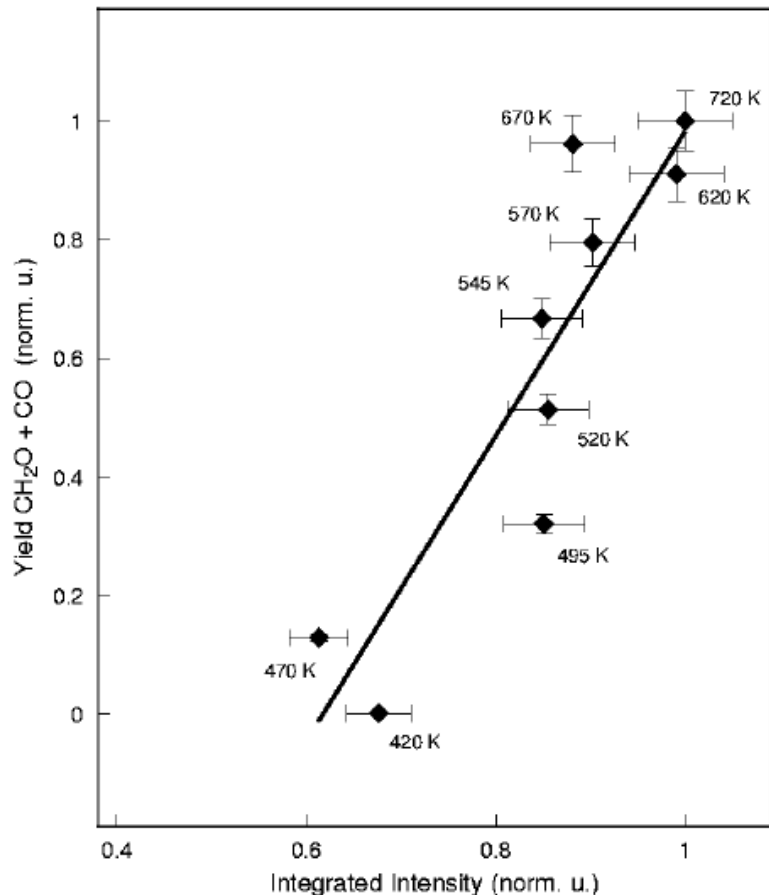
less active

very active

2 oxide and 1 suboxide species

In situ and operando techniques

- Example: XANES – x-ray absorption near edge structure



- Positive correlation between catalytic activity and Cu suboxide species
- Explanation/idea/description of active species: Subsurface oxygen cover by a strained layer of copper atoms
- Results confirmed by near ambient pressure XPS (NAP XPS)

In situ and operando techniques

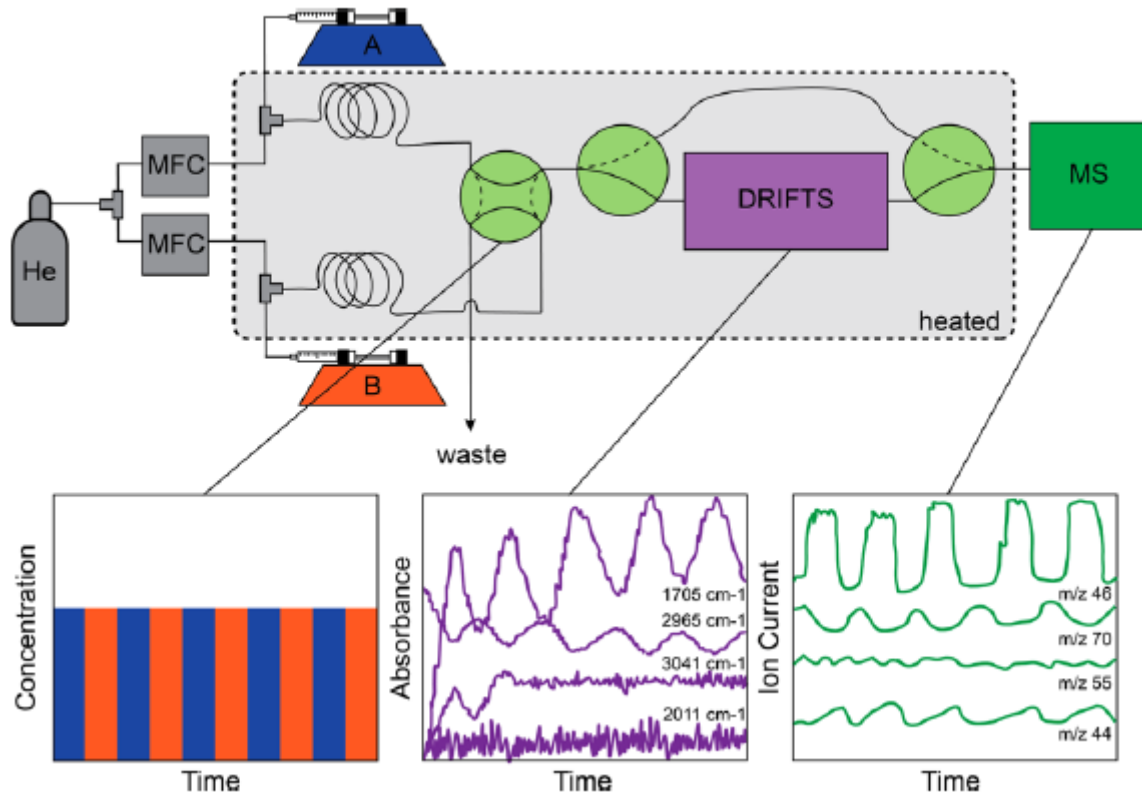
- Example: DRIFTS – diffuse reflectance infrared Fourier transform spectroscopy
- Coupling of ethanol and acetaldehyde to 1,3-butadiene over Ta doped zeolite
 - 300 °C, 1 atm
 - Well dispersed (virtually isolated) Ta sites



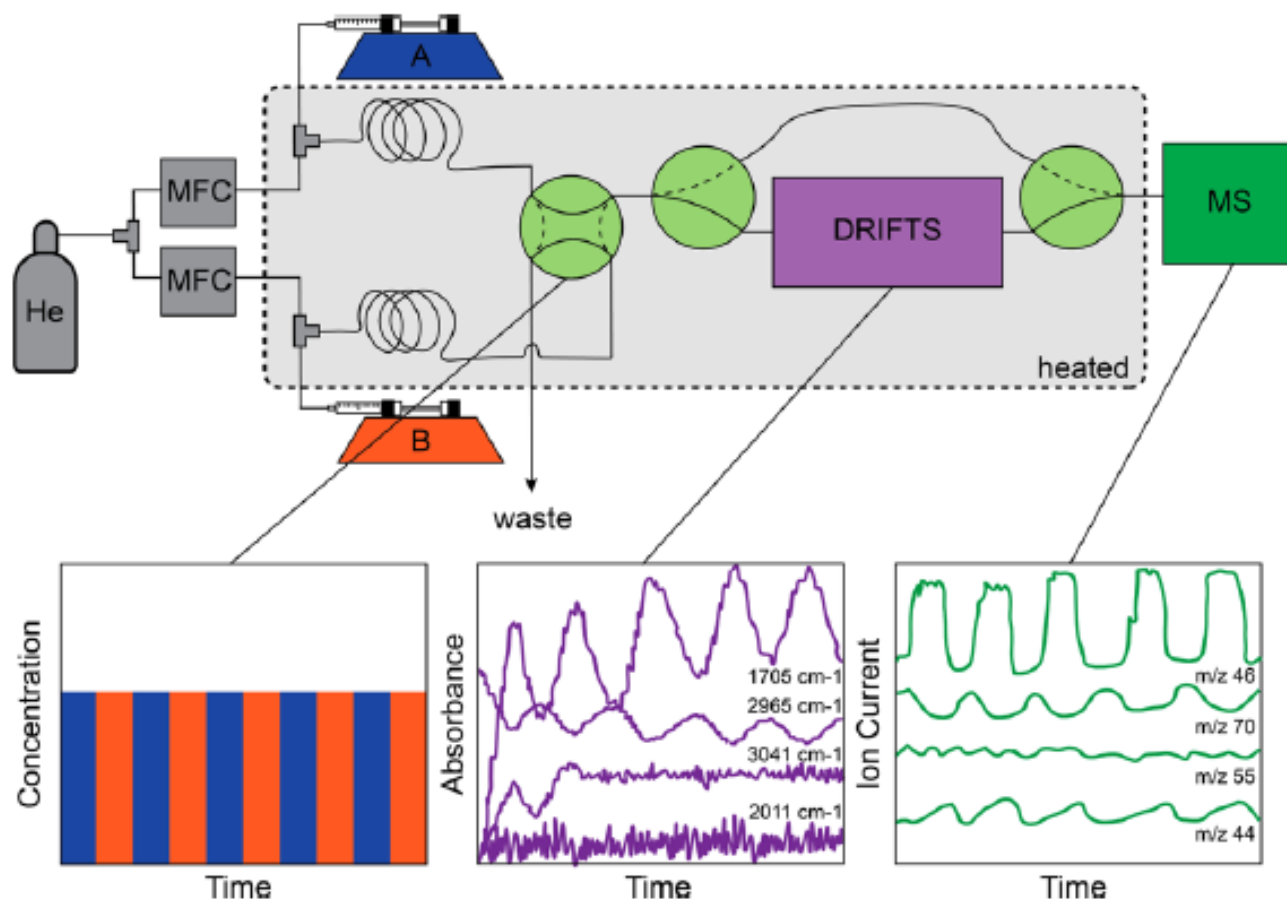
In situ and operando techniques

- Example: DRIFTS – diffuse reflectance infrared Fourier transform spectroscopy

Scheme 2. Schematic Overview of the DRIFTS-MS Setup^a



Scheme 2. Schematic Overview of the DRIFTS-MS Setup^a



^aBy using two syringe pumps and heated coils that enter a two-position-four-way valve, the gas-phase composition can be modulated between substrates A and B. Depending on the chosen position, either flow A or B flows through the DRIFTS accessory followed by the online mass spectrometer. In addition, two three-way valves allow the substrates to bypass the DRIFTS accessory to ensure steady gas-phase concentrations prior to measurements.

In situ and operando techniques

- Example: DRIFTS

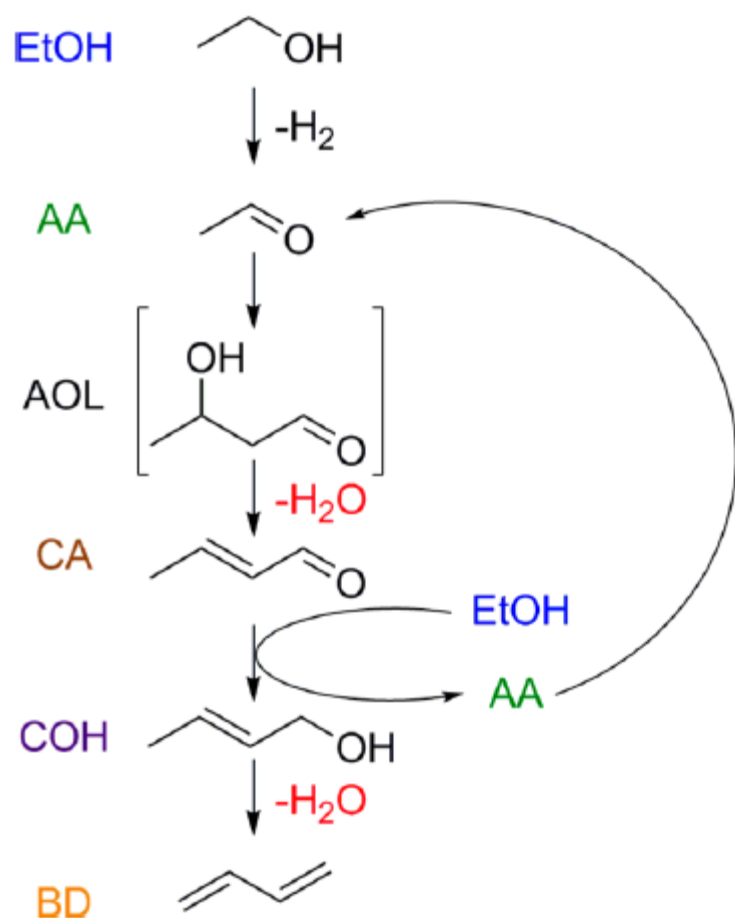


Table 1. Chosen m/z Signals^a and Some IR Vibrational Frequencies^b for the Most Important Intermediates

| Molecule | m/z | C-H stretch [cm^{-1}] | C=O stretch [cm^{-1}] | C=C stretch [cm^{-1}] | other vibrations [cm^{-1}] |
|----------|-------|-------------------------------------|-------------------------------------|-------------------------------------|--|
| EtOH | 46 | 2988, 2970, 2900 | | | 1065 |
| AA | 44 | 2820, 2725, 2700 | 1750 | | |
| BD | 54 | 3108, 3090, 3045 | | 1605, 1588 | |
| CA | 70 | 2935, 2820, 2730 | 1722, 1710 | 1640 | |
| COH | 57 | 3025, 2937, 2880 | | 1676 | 1450, 1440 |

In situ and operando techniques

- Example: DRIFTS – diffuse reflectance infrared Fourier transform spectroscopy

