

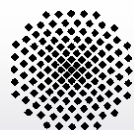


Contributions of Solid-state NMR Spectroscopy to the Development and Understanding of Solid Catalysts

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**Seminar at the Key Laboratory of
Advanced Energy Materials Chemistry**
College of Chemistry
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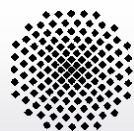




Most abundant isotopes with nuclear spin

spin $I = 1/2$																															
integer spin																															
quadrupolar nuclei with spin $I = 3/2, 5/2$ etc.																															
H																	He														
Li	Be															B	C	N	O	F	Ne										
Na	Mg															Al	Si	P	S	Cl	Ar										
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr														
Rb	Sr	Y	Zr	Nb	Mo	Tc	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	I	Xe														
Cs	Ba	La	Hf	Ta	W	Re	Os	Ir	Pt	Au	Hg	Tl	Pb	Bi	Po	At	Rn														
Fr	Ra	Ac																Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
																		Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr





Isotopes interesting for solid-state NMR in heterogeneous catalysis

interesting isotopes (nuclear spin, relative sensitivity in comparison with ^1H)

^1H (1/2, 1.00), ^2H (1, 1.4×10^{-6})

^{13}C (1/2, 1.8×10^{-4}), ^{15}N (1/2, 3.8×10^{-6})

^{31}P (1/2, 6.6×10^{-2})

^7Li (3/2, 0.27)

^{11}B (3/2, 0.13)

^{17}O (5/2, 1.1×10^{-5})

^{23}Na (3/2, 9.2×10^{-2})

^{27}Al (5/2, 0.21)

^{29}Si (1/2, 3.7×10^{-4})

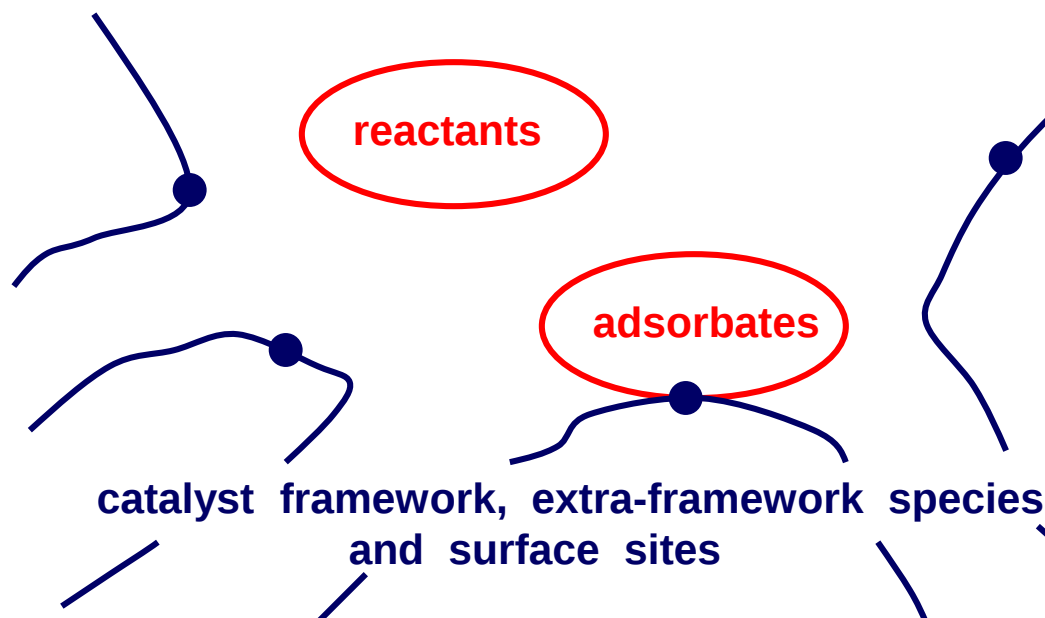
^{31}P (1/2, 6.6×10^{-2})

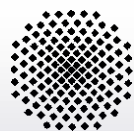
^{51}V (7/2, 0.38)

^{67}Zn (5/2, 1.2×10^{-2})

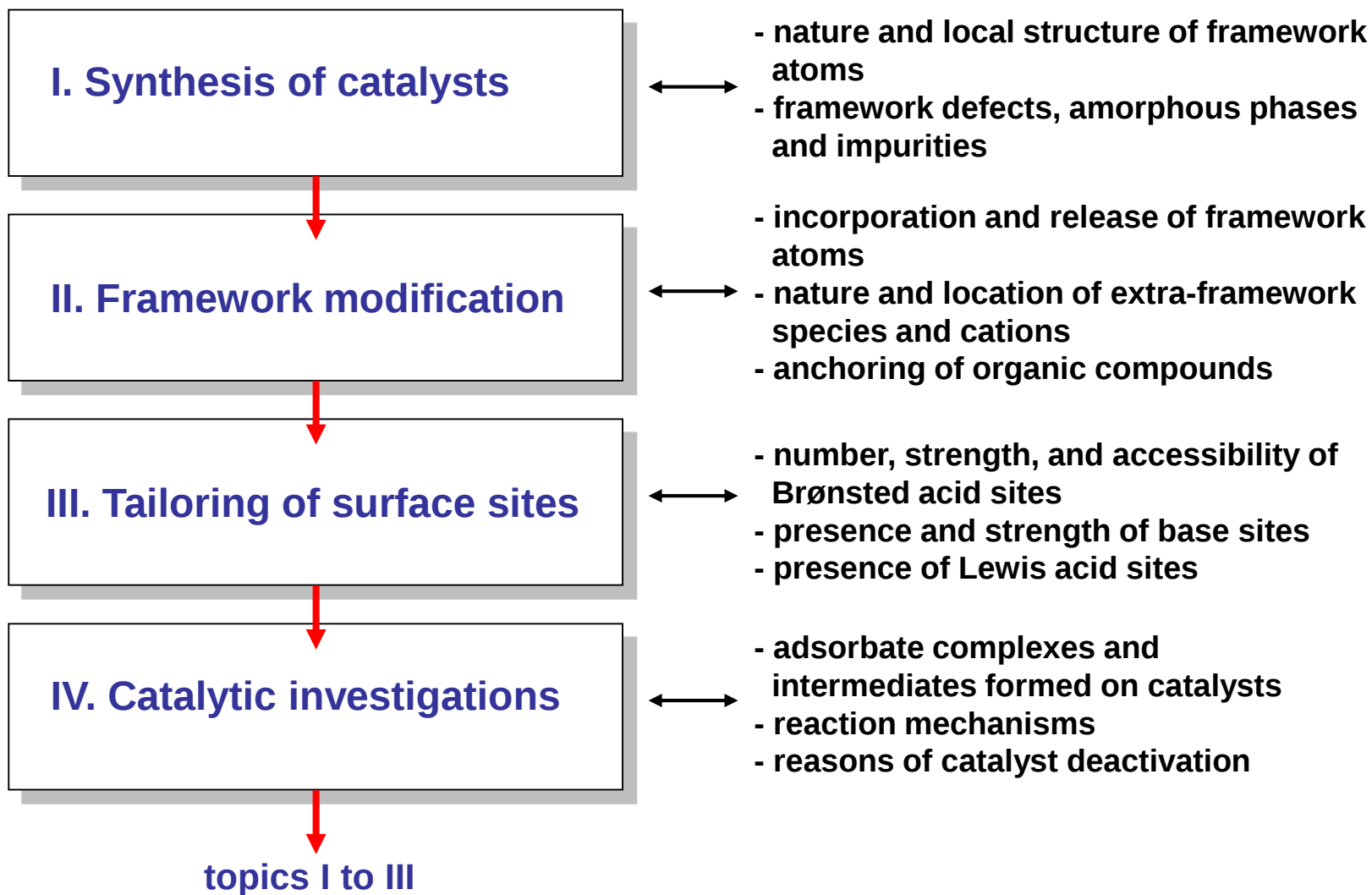
^{71}Ga (3/2, 5.6×10^{-2})

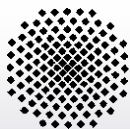
^{133}Cs (7/2, 4.7×10^{-2})





Solid-state NMR of solid catalysts





Specific problems of solid-state NMR spectroscopy

Magnetization M_0 , Curie's law:

$$M_0 = \frac{N \gamma^2 h^2 I (I + 1) B_0}{(2\pi)^2 3 k_B T}$$

detection limit of solid-state NMR is ca. $N = 10 \mu\text{mol}$ spins per gram for ^1H nuclei at room temperature

magnetization M_0 decreases with increasing temperature T

broadening of NMR signals due to internal solid-state interactions

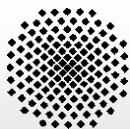
most important solid-state interactions:

H_{CSA} : anisotropic shielding of the magnetic field due to the anisotropic electron density in the local structure of the resonating nuclei; $\mathcal{F} \{LP^{(2)}\}$
 $\Delta \nu_{\text{CSA}}$ up to 50 kHz

H_{DI} : dipolar interaction with magnetic dipole moments of neighboring spins in the local structure; $\mathcal{F} \{LP^{(2)}\}$
 $\Delta \nu_{\text{DI}}$ up to 100 kHz

H_{Q} : quadrupolar interaction of the electric quadrupole moment of nuclei with spin $I = 3/2, 5/2$ etc. with electric field gradients; $\mathcal{F} \{LP^{(2)} + LP^{(4)}\}$
 $\Delta \nu_{\text{Q}}$ up to 20 MHz





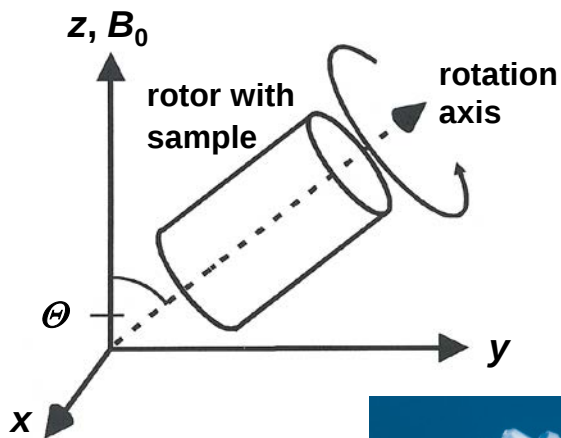
Techniques of solid-state NMR spectroscopy

spin $I = 1/2$:

magic angle spinning (**MAS**)

$$\nu_{\text{CSA, DI, 1QI}} = \mathcal{F} \{ (3\cos^2\Theta - 1)/2 \}$$

$$\rightarrow \Theta = 54.7^\circ$$



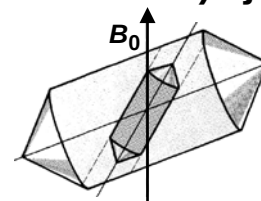
spin $I = 3/2, 5/2$ etc. :

double oriented rotation (**DOR**)

$$\nu_{2\text{QI}} = \mathcal{F} \{ (35\cos^4\Theta - 30\cos^2\Theta + 3)/8 \}$$

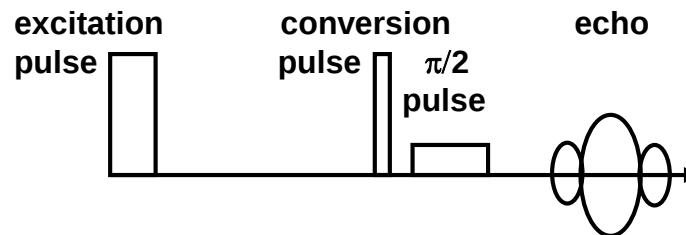
$$\rightarrow \Theta = 30.6^\circ$$

$$\Theta = 70.1^\circ$$



multiple-quantum MAS NMR (**MQMAS**)

- excitation of three- and five-quantum transitions
- recording of a spin-echo free of anisotropic contributions



J. Amoureux et al., J. Magn. Reson A 123 (1996) 116





I. Synthesis of catalysts

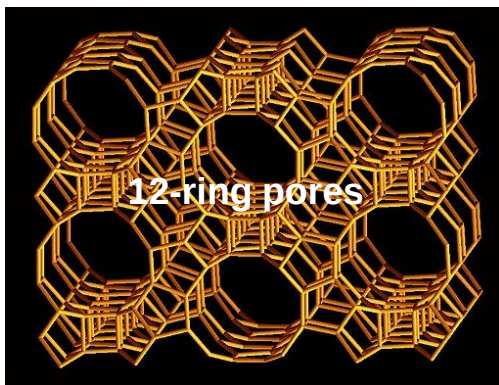




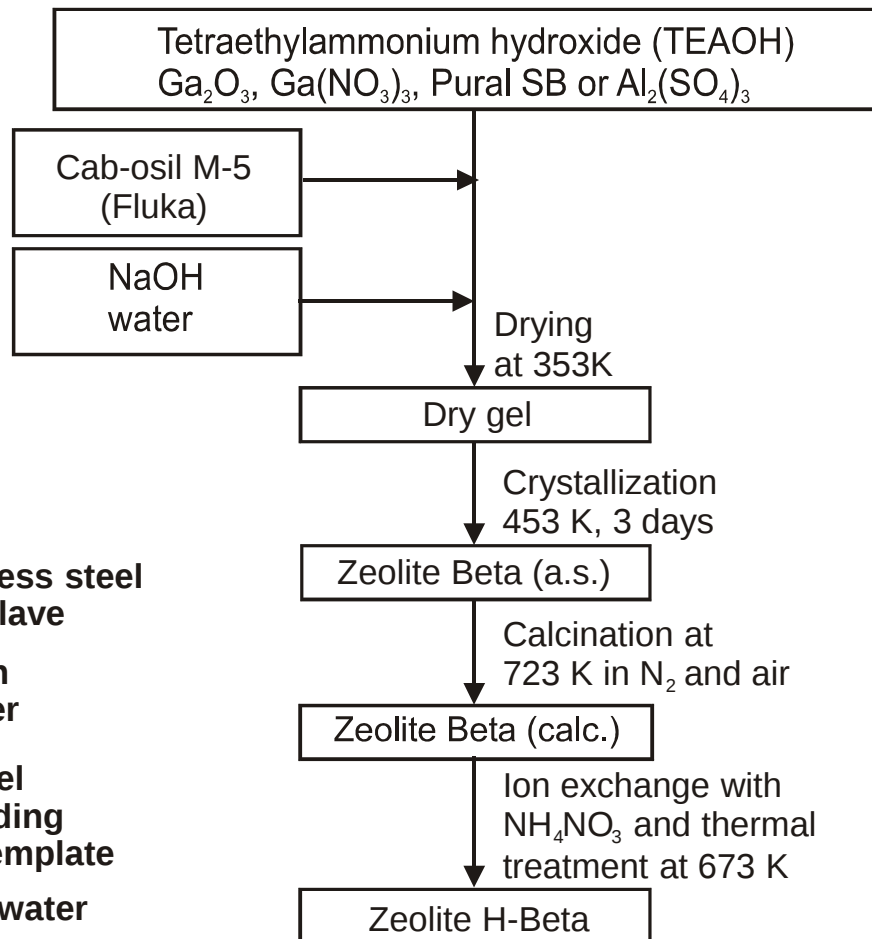
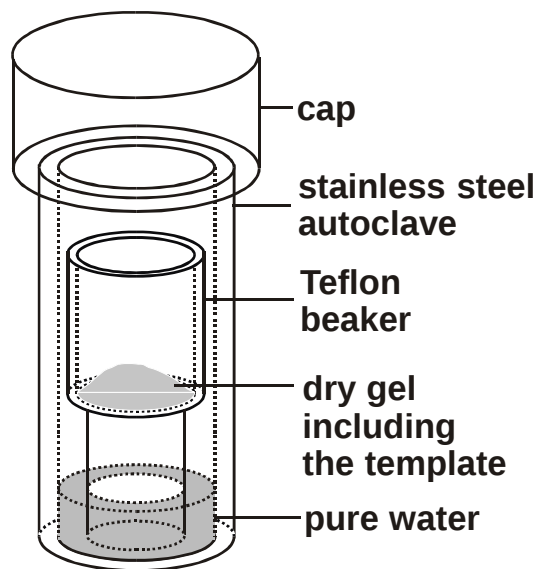
I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

structure of zeolite Beta (BEA)



autoclave volume:
110 cm³
Teflon beaker:
14.5 cm³

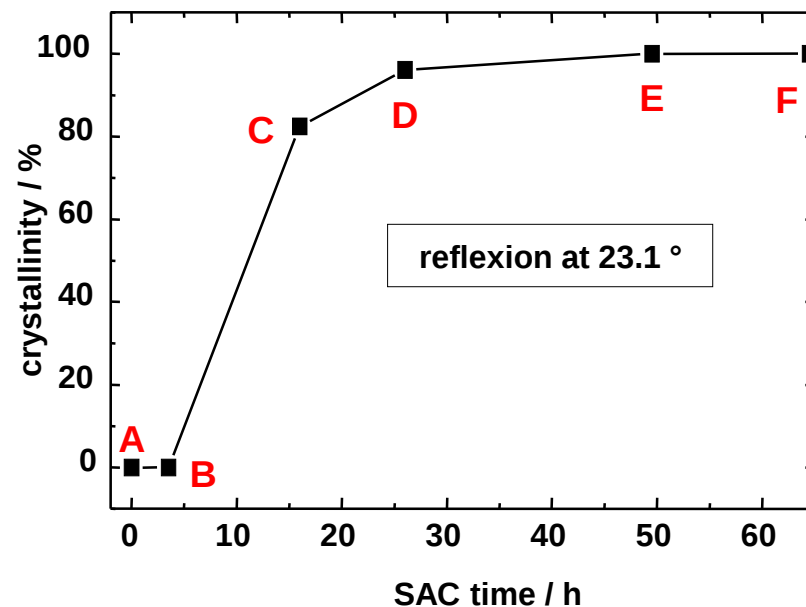
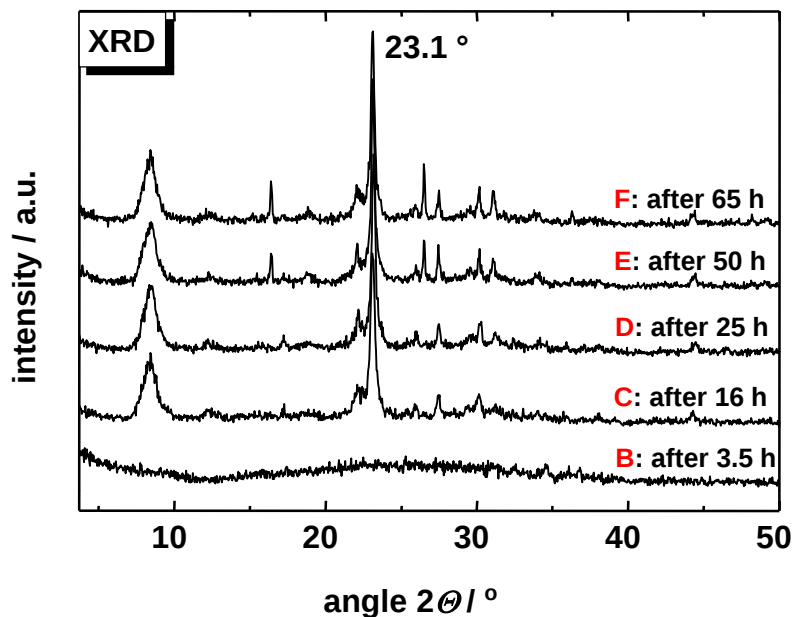




I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

powder X-ray diffractograms and crystallinity as a function of the dry-gel conversion time



sample **A**:

dry gel taken before conversion

samples **B** to **F**:

materials taken after conversion for 3.5 to 65 h





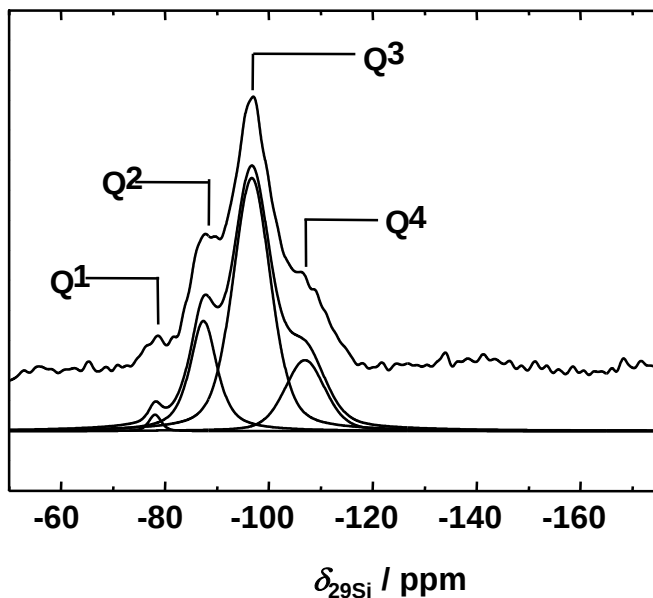
I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

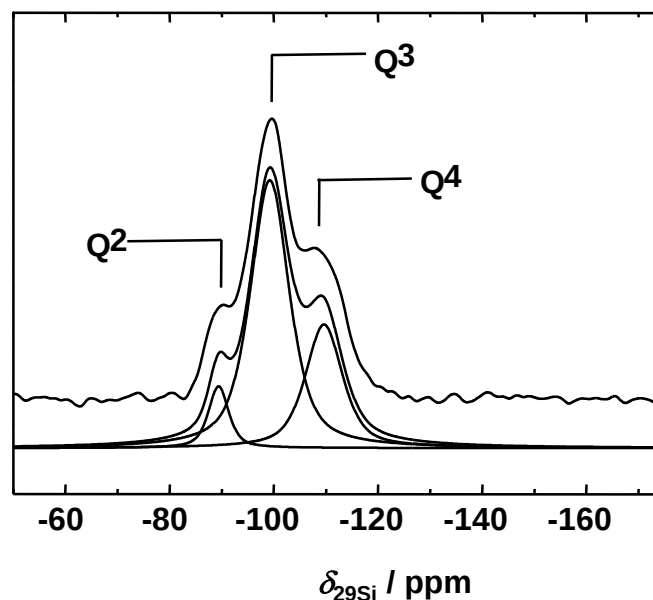
^{29}Si MAS NMR spectra recorded after different SAC times

$B_0 = 9.4 \text{ T}$, $\nu_0 = 79.5 \text{ MHz}$

A: fresh dry-gel particles (0 h)

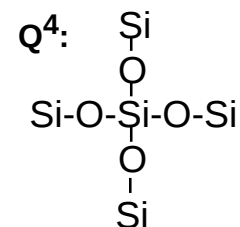
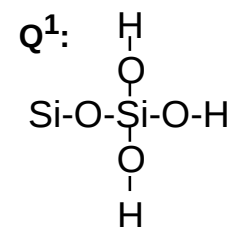


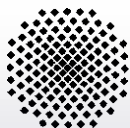
B: SAC time of 3.5 h



Q¹: silicon with one Si-O-Si bond, Si(1Si,3OH), at ca. -78 ppm

Q⁴: silicon with four Si-O-Si bonds, Si(4Si), at ca. -107 ppm





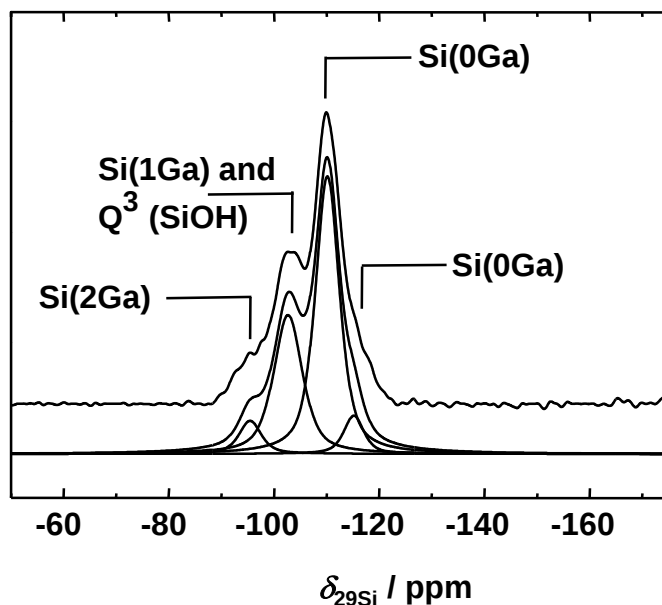
I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

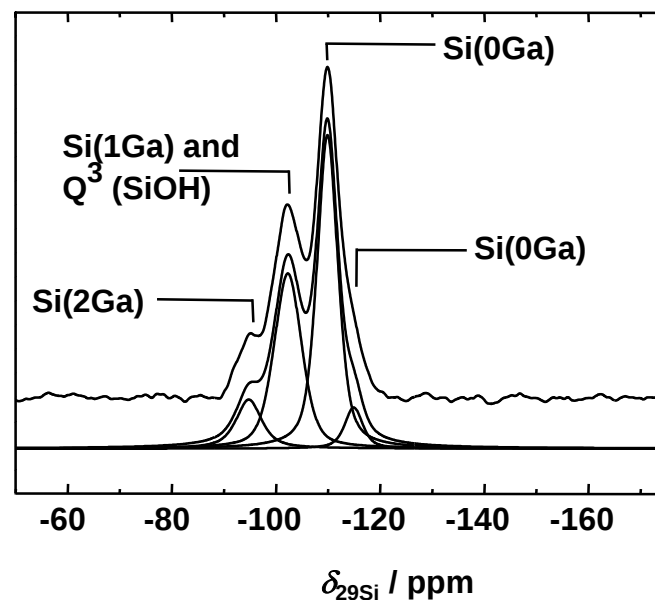
^{29}Si MAS NMR spectra recorded after different SAC times

$B_0 = 9.4 \text{ T}$, $\nu_0 = 79.5 \text{ MHz}$

C: SAC time of 16 h



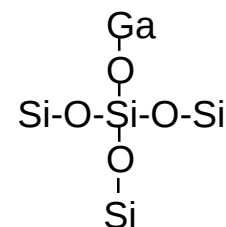
F: SAC time of 65 h

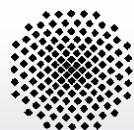


Si(1Ga): Q^4 framework silicon with one Si-O-Ga and three Si-O-Si bonds, Si(3Si,1Ga), at -101 ppm is overlapped by the Q^3 signal, Si(3Si,1OH), at ca. -103 ppm

Si(2Ga): framework silicon with two Si-O-Ga and two Si-O-Si bonds, Si(2Si,2Ga), at -95 ppm

Q^4 : Si(3Si,1Ga)

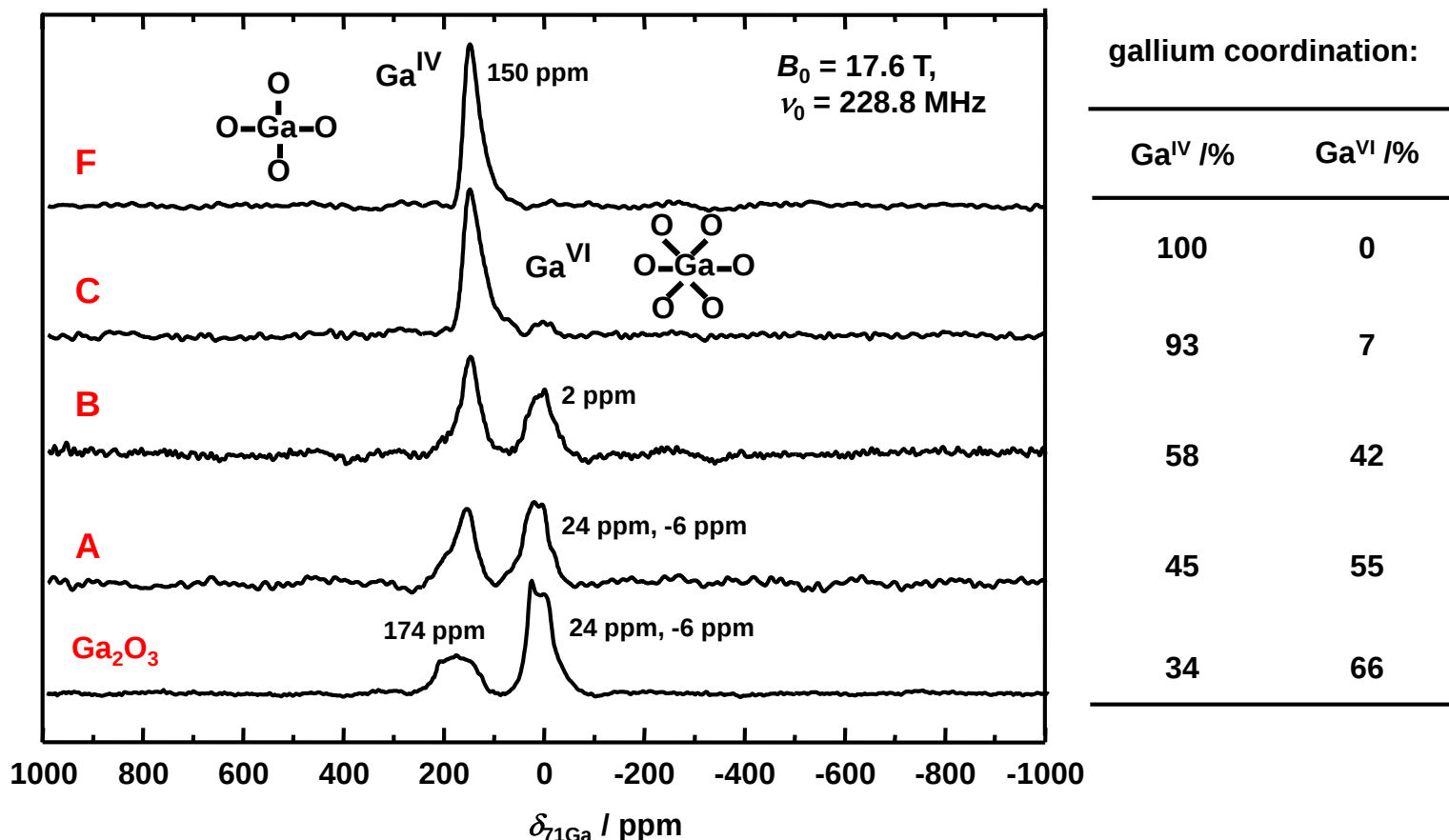




I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

^{71}Ga MAS NMR spectra (spin $I = 3/2$) recorded after different SAC times



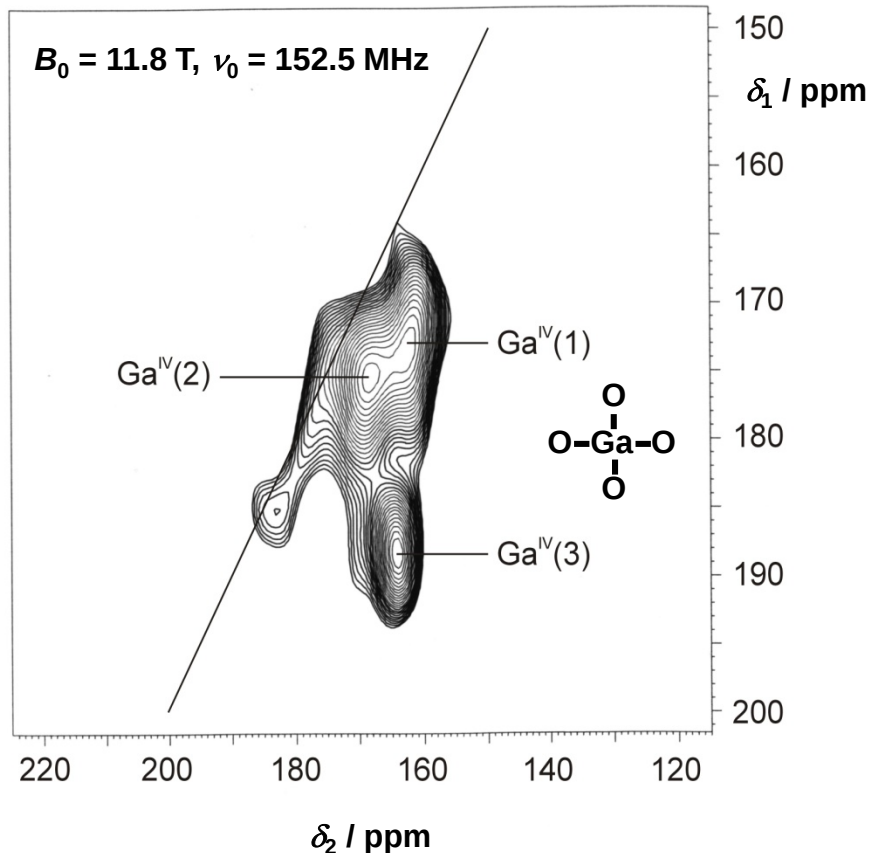
complete tetrahedral oxygen coordination of gallium atoms after a SAC time of 65 h



I. Synthesis of catalysts

synthesis of zeolite [Ga,Al]Beta by steam-assisted dry-gel conversion (SAC)

^{71}Ga MQMAS NMR spectroscopy (spin $I = 3/2$) of [Ga]Beta sample **F** (65 h)



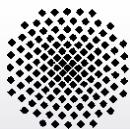
$\text{Ga}^{\text{IV}}(1, 2)$:
framework gallium species,
 $\text{SOQE} = 2.2$ and 1.8 MHz ,
 $\delta_{\text{iso}} = 172$ and 168 ppm

$\text{Ga}^{\text{IV}}(3)$:
extra-framework gallium
species or amorphous
phases,
 $\text{SOQE} = 4.7 \text{ MHz}$,
 $\delta_{\text{iso}} = 176 \text{ ppm}$

SOQE: second-order quadrupole
effect parameter,
strength of quadrupolar interaction

A. Arnold et al., Microporous Mesoporous
Mater. 62 (2003) 97.





I. Synthesis of catalysts

synthesis of flame-derived silica-alumina with different aluminum contents

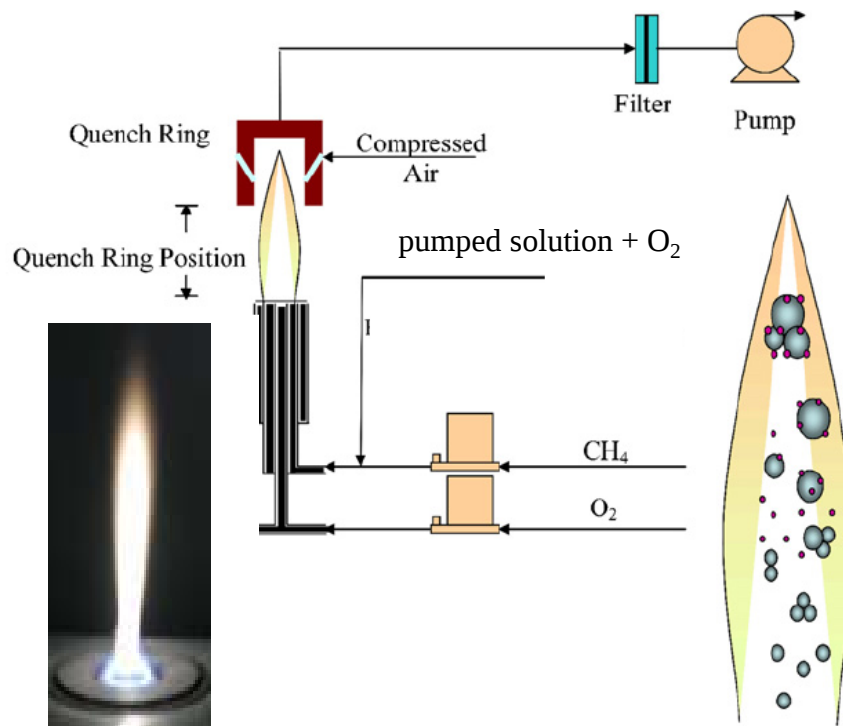
preparation of flame-made silica-alumina (SA) catalysts:

dissolving aluminum(III) acetylacetonate and tetraethoxysilane in a 1:1 (vol%) mixture of acetic acid and methanol

this solution was pumped through a capillary with 5 ml/min and nebulized with 5 l/min O₂

the spray was ignited by a methane/oxygen flame (1.5/0.9 L/min) resulting in an approximately 6 cm long flame

particles (ca. 20 nm) were collected on a cooled Whatman GF6 filter





I. Synthesis of catalysts

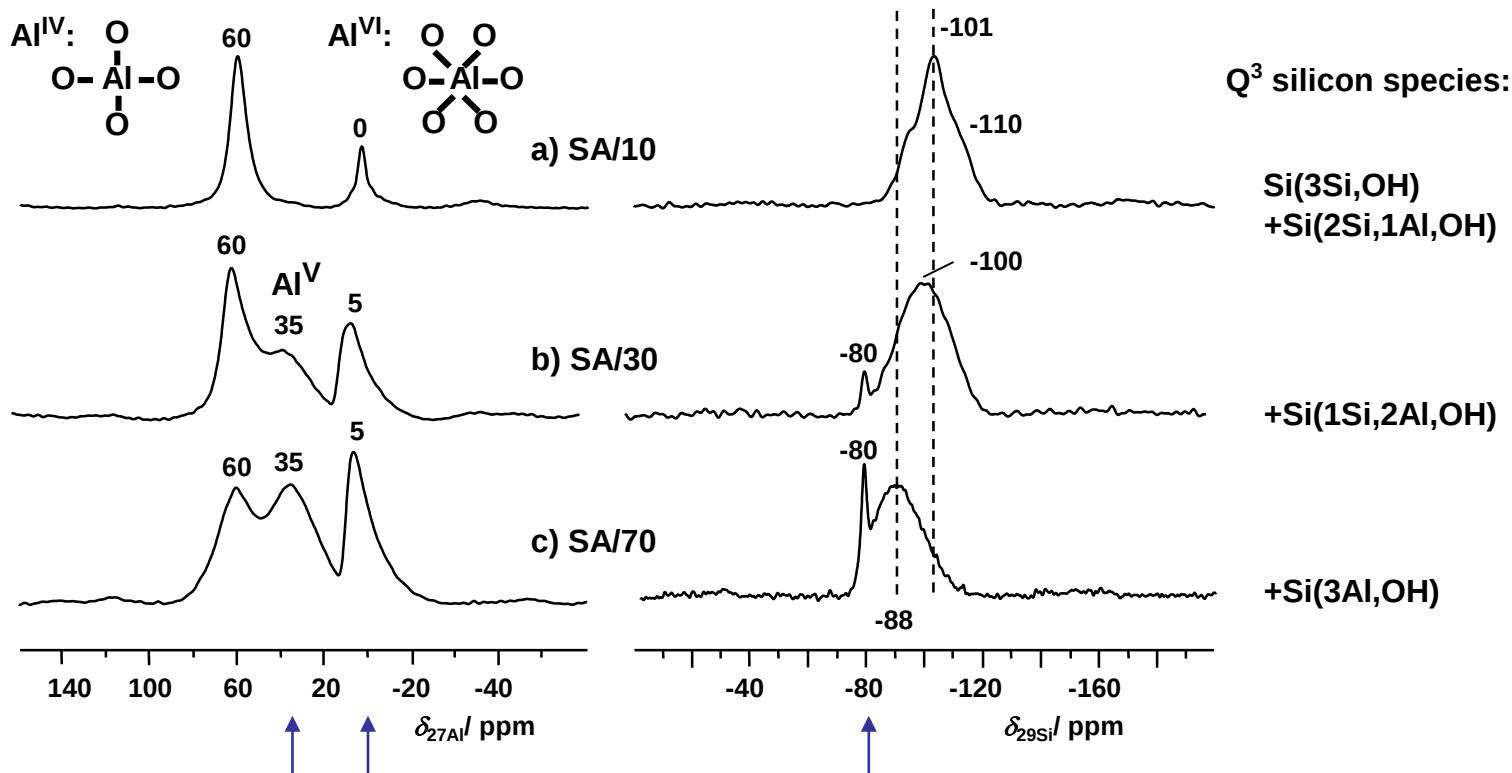
synthesis of flame-derived silica-alumina with different aluminum contents

^{27}Al MAS NMR (spin $I = 5/2$)

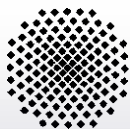
$B_0 = 9.4 \text{ T}$, $\nu_0 = 104.4 \text{ MHz}$

^{29}Si MAS NMR

$B_0 = 9.4 \text{ T}$, $\nu_0 = 79.5 \text{ MHz}$



formation of andalusite phases (Al_2SiO_5) with $\delta_{^{29}\text{Si}} = -79.8 \text{ ppm}$ (M. Mägi *et al.*, J. Phys. Chem. 88 (1984) 1518) and octahedrally ($\delta_{^{27}\text{Al}} = 5 \text{ ppm}$) as well as pentacoordinated aluminum atoms ($\delta_{^{27}\text{Al}} = 35 \text{ ppm}$)



I. Synthesis of catalysts

synthesis of flame-derived silica-alumina with different aluminum contents

^1H MAS NMR

upon dehydration at 723 K

SA/0, pure silica

SiOH

1.8

$B_0 = 9.4 \text{ T}$

$\nu_0 = 400.1 \text{ MHz}$

SA/0 + NH_3

NH_4^+

1.8

SA/10

1.8

SA/10 + NH_3

7.0

1.8

SA/30

1.8

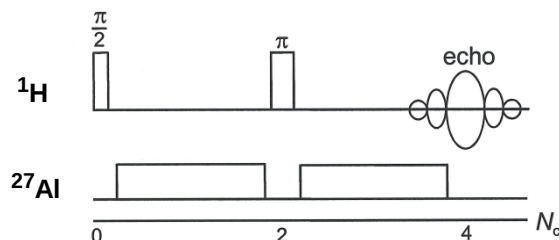
SA/30 + NH_3

7.0

1.8

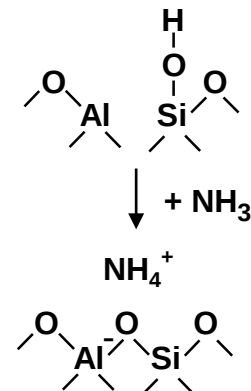
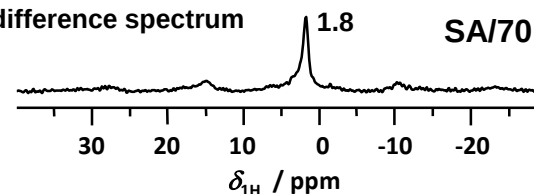
increases with increasing
molar aluminum content

$^1\text{H}/^{27}\text{Al}$ TRAPDOR MAS NMR



difference spectrum

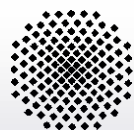
SA/70





II. Modification of catalysts

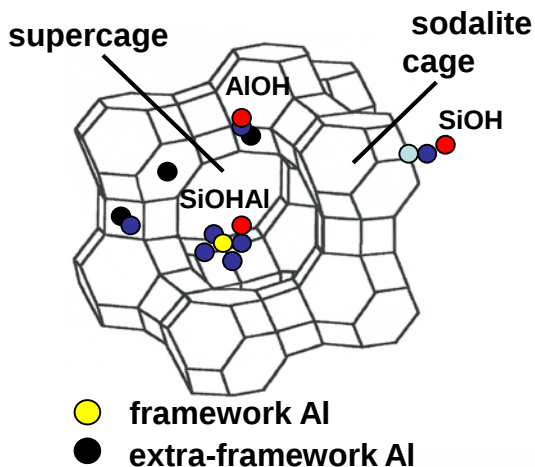




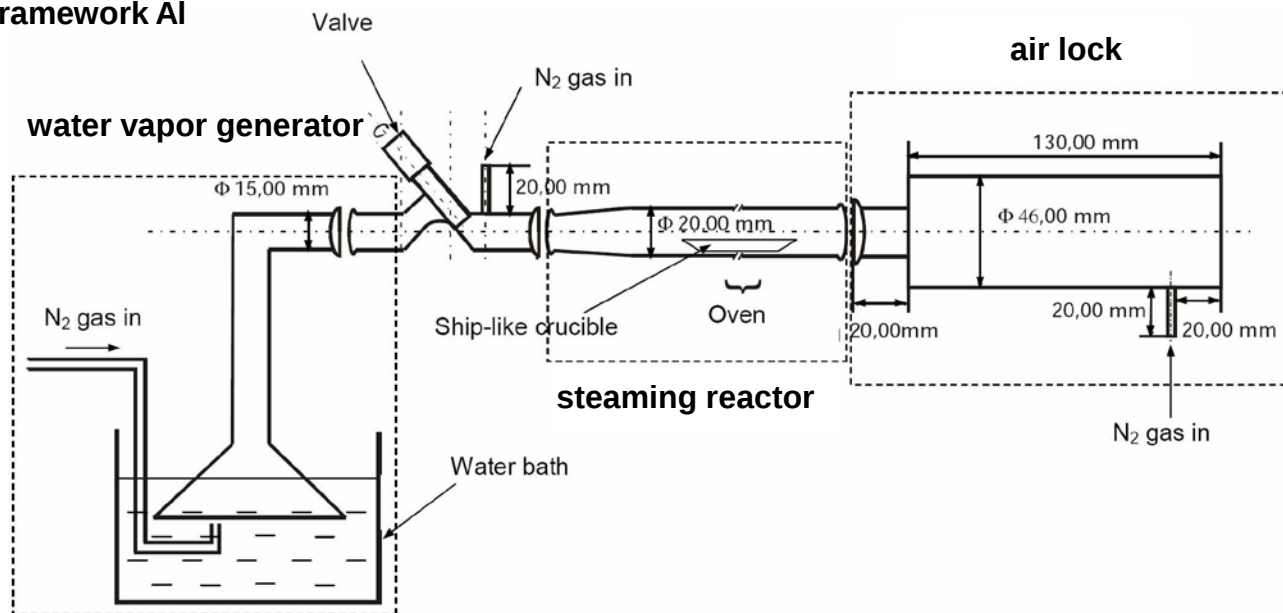
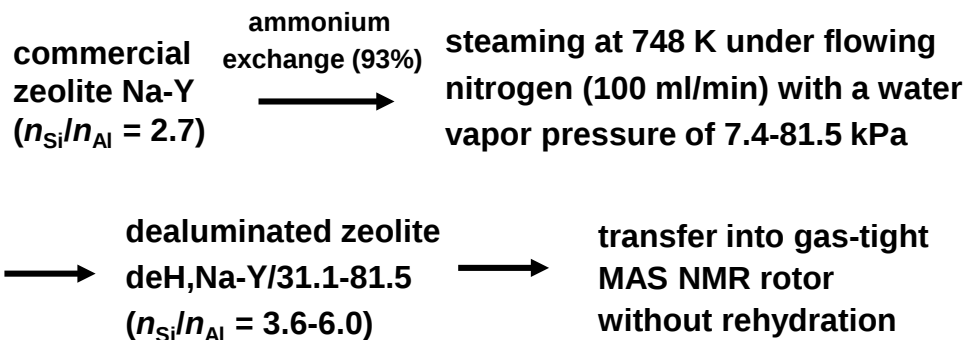
II. Framework modification

dealumination of zeolite H,Na-Y by steaming

structure of zeolite Y (FAU)



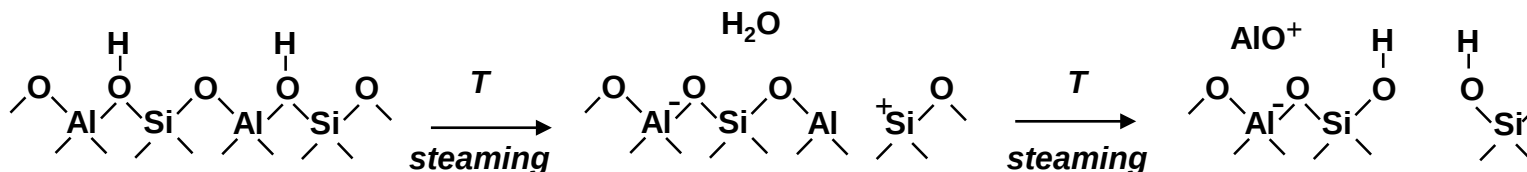
dealumination procedure



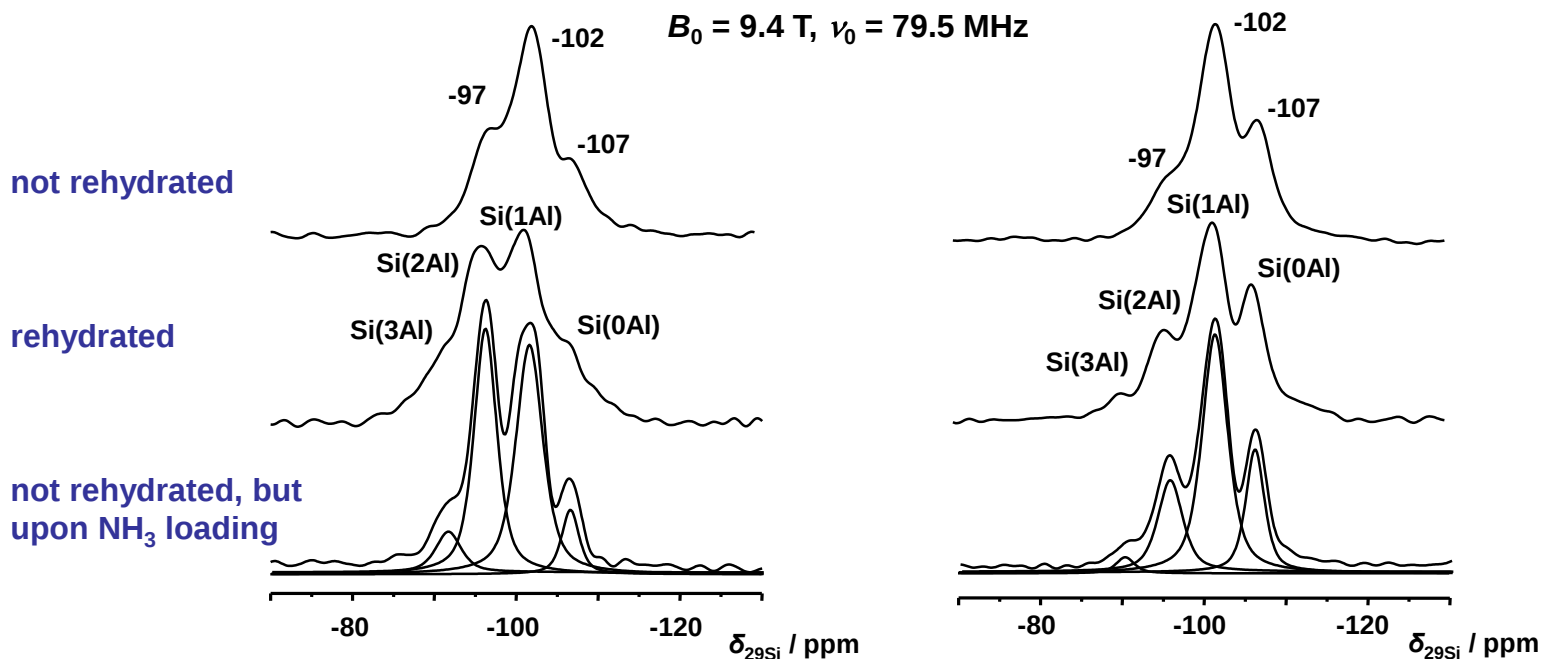


II. Framework modification

dealumination of acidic zeolites by steaming

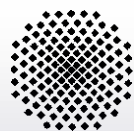


²⁹Si MAS NMR spectroscopy of zeolites H,Na-Y (left) and deH,Na-Y/31.1 (right)



$$\frac{n_{\text{Si}}}{n_{\text{Al}}} = \frac{100}{\sum_n 0.25 \cdot n \cdot I_{\text{Si}(n\text{Al})}} : 2.7 \text{ (51.9 Al}^{\text{fr}} / \text{u.c.)}$$

$$3.8 \text{ (40.0 Al}^{\text{fr}} / \text{u.c.} + 11.9 \text{ Al}^{\text{ex}} / \text{u.c.)}$$

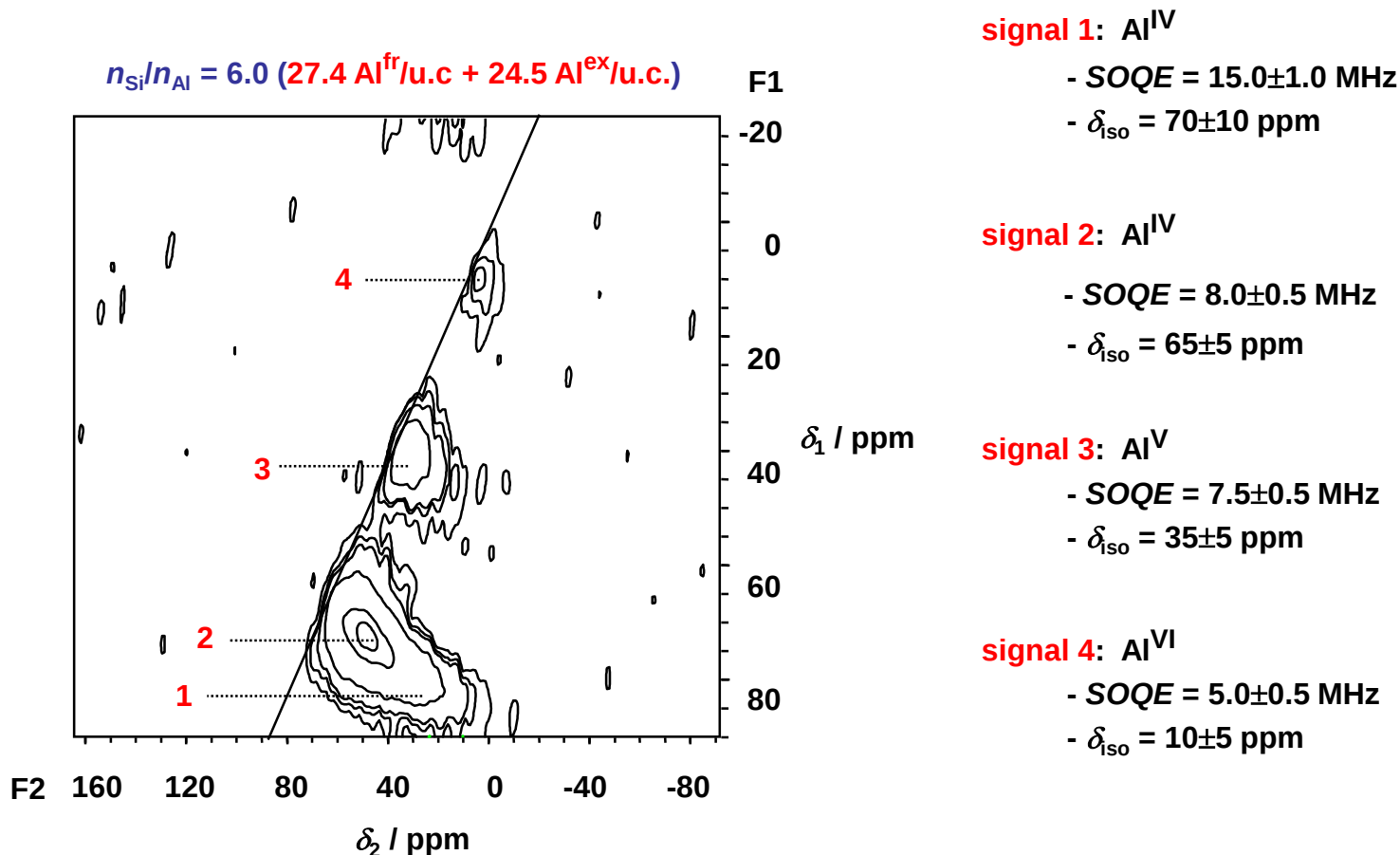


II. Framework modification

dealumination of zeolite H,Na-Y by steaming

^{27}Al MQMAS NMR spectroscopy (spin $I = 5/2$) of not rehydrated zeolite deH,Na-Y/81.5

$B_0 = 17.6 \text{ T}$, $\nu_0 = 195.5 \text{ MHz}$, $\nu_{\text{rot}} = 30 \text{ kHz}$





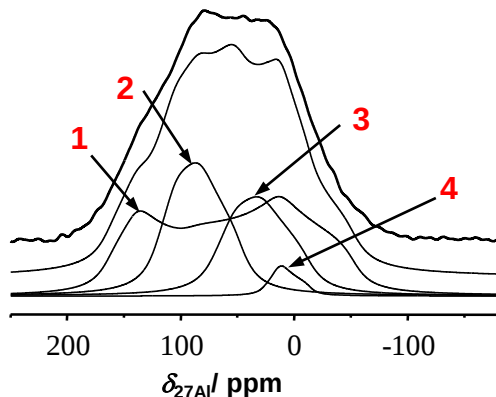
II. Framework modification

dealumination of zeolite H,Na-Y by steaming

^{27}Al solid-state NMR spectroscopy (spin $I = 5/2$) of zeolite deH,Na-Y/81.5 (51.9 Al/u.c.)

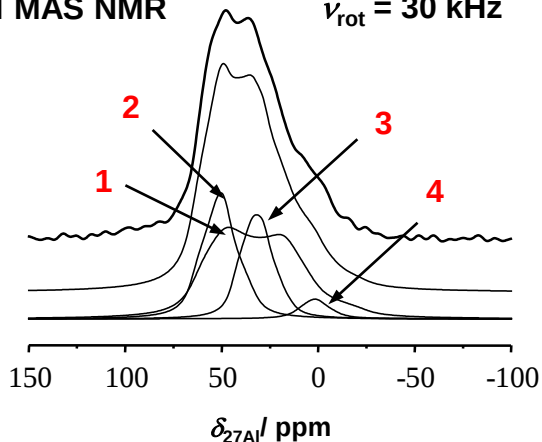
$B_0 = 17.6 \text{ T}$, $\nu_0 = 195.5 \text{ MHz}$

^{27}Al spin-echo NMR



^{27}Al MAS NMR

$\nu_{\text{rot}} = 30 \text{ kHz}$



Signals	1	2	3	4
assignments	$\text{Al}^{\text{IV}}/\text{H}^+$ $\text{Al}^{\text{IV}}/\text{Al}^{\text{cat},\text{x}+}$	$\text{Al}^{\text{IV}}/\text{Na}^+$ cluster Al^{IV}	$\text{Al}^{\text{V},\text{x}+}$ cat.	cluster Al^{VI}
$\delta_{\text{iso}} / \text{ppm}$	70 ± 10	65 ± 5	35 ± 5	10 ± 5
$C_{\text{QCC}} / \text{MHz}$	15.0 ± 1.0	8.0 ± 0.5	7.5 ± 0.5	5.0 ± 0.5
η_{Q}	0.3	0.8	0.7	0.7
I in %	48 (27+21)	27 (7+20)	21	4

extra-framework
Al cations ($\text{Al}^{\text{V},\text{x}+}$):

10.9 $\text{Al}^{\text{V},\text{x}+}/\text{u.c.}$

extra-framework
 Al_2O_3 clusters:

12.5 $\text{Al}^{\text{cluster}}/\text{u.c.}$

in comparison with 24.5 $\text{Al}^{\text{ex}}/\text{u.c.}$ calculated using results of
 ^{29}Si MAS NMR spectroscopy

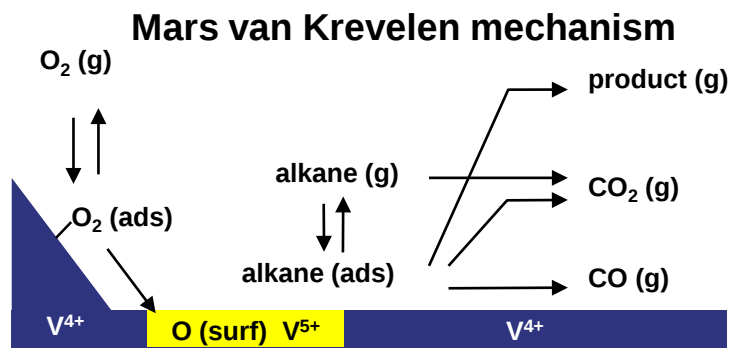




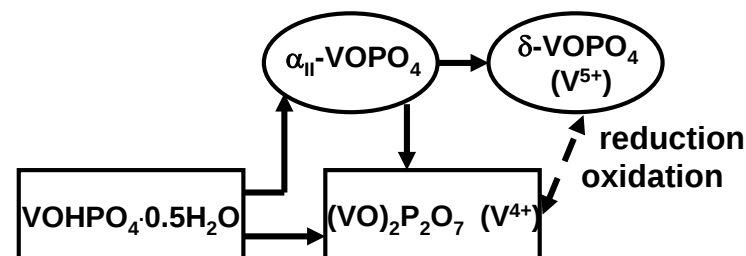
II. Modification of catalysts

VPO catalysts supported on mesoporous SBA-15 materials

selective oxidation of *n*-butane



discussed phase transitions



preparation procedure of X.-K. Li et al., J. Catal. 238 (2006) 232:

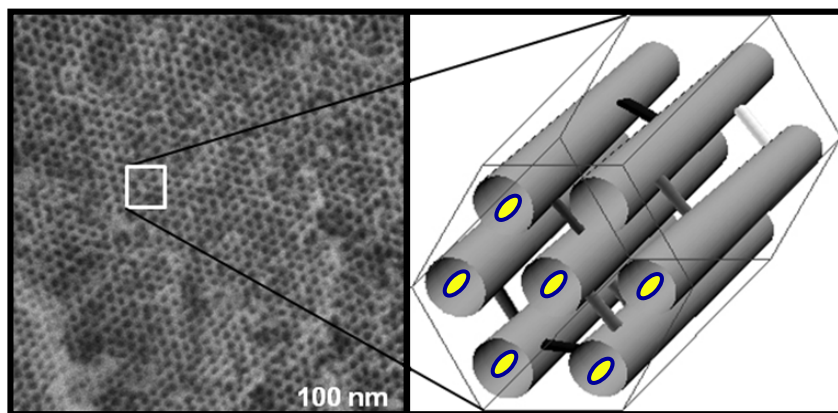
- siliceous SBA-15 is added to isobutyl/benzyl alcohols (1 : 1) with V_2O_5 and H_3PO_4
- activation in a flow of 1.5 % *n*-butane, 17.5 % O_2 and balance N_2 (100 ml/min) at 673 K



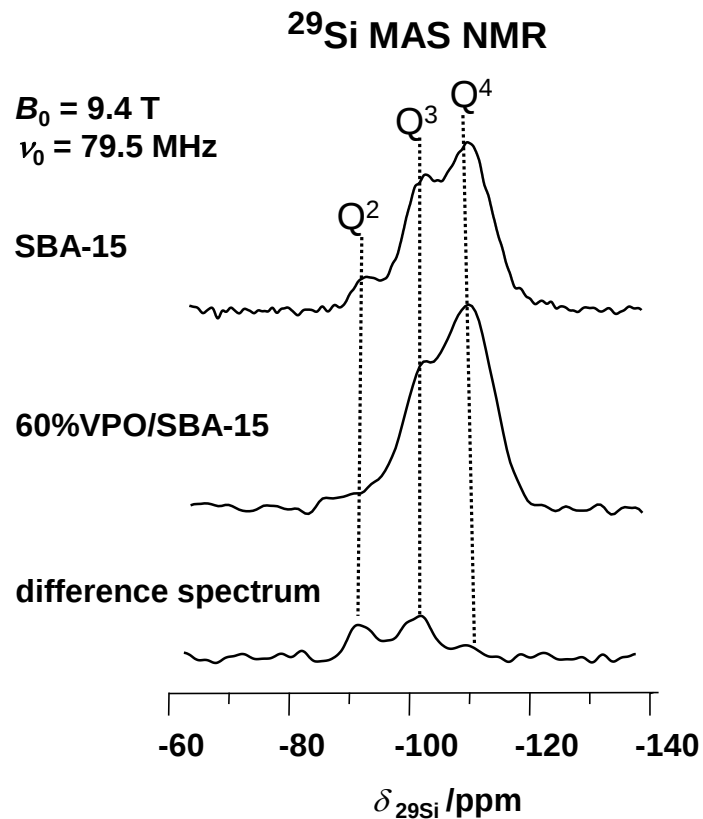


II. Framework modification

VPO catalysts supported on mesoporous SBA-15 materials

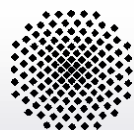


Materials	P / V	BET surface m ² / g	Pore volume cm ³ / g
SBA-15	-	1164	1.25
20%VPO/SBA-15	1.09	662	0.80
60%VPO/SBA-15	1.04	456	0.54



VPO is bound at former silanol groups on the mesoporous surface of SBA-15

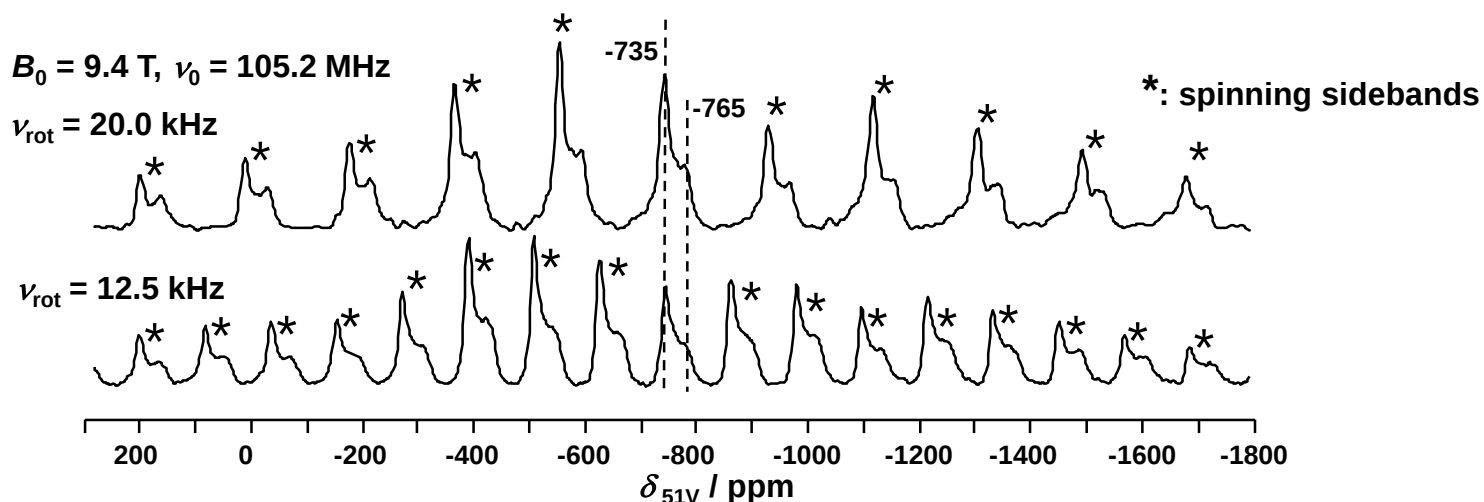




II. Framework modification

VPO catalysts supported on mesoporous SBA-15 materials

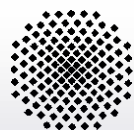
^{51}V MAS NMR spectroscopy (spin $I = 7/2$) of 60%VPO/SBA-15 after activation



Materials	$\delta_{\text{iso}} / \text{ppm}$	$\Delta\sigma_{\text{CSA}} / \text{ppm}$	η_{CSA}	C_Q / MHz	η_Q
60%VPO/SBA-15	-735	900	0.10	2.00	0.60
	-765	950	0.10	1.90	0.60
$\alpha_{\text{II}}\text{-VOPO}_4$ ¹⁾	-755	992	0.08	0.63	0.09
$\beta\text{-VOPO}_4$ ¹⁾	-735	818	0.05	1.45	0.44

¹⁾ R. Siegel et al., Mag. Res. Chem. 42 (2004) 1022.
 O.B. Lapina et al., Prog. Nucl. Magn. Reson. Spectrosc. 53 (2008) 128.





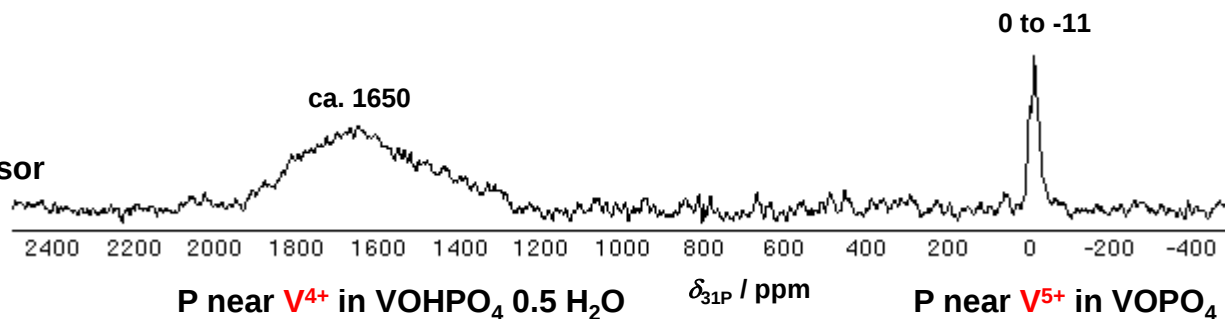
II. Framework modification

VPO catalysts supported on mesoporous SBA-15 materials

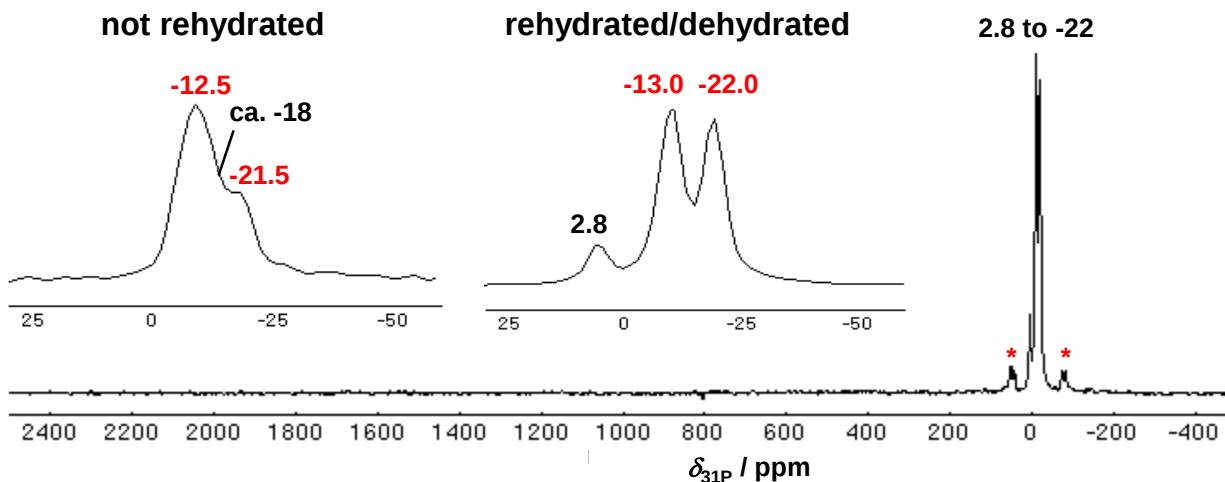
^{31}P MAS NMR spectroscopy of 60%VPO/SBA-15 before and after activation

$B_0 = 9.4 \text{ T}$
 $\nu_0 = 162.0 \text{ MHz}$
 $\nu_{\text{rot}} = 10 \text{ kHz}$

non-activated precursor



activated at 673 K
for 15 h



P near V^{5+} in VOPO_4 phases, such as $\beta\text{-VOPO}_4$ (-13 ppm) and $\alpha_{\text{II}}\text{-VOPO}_4$ (-22 ppm)

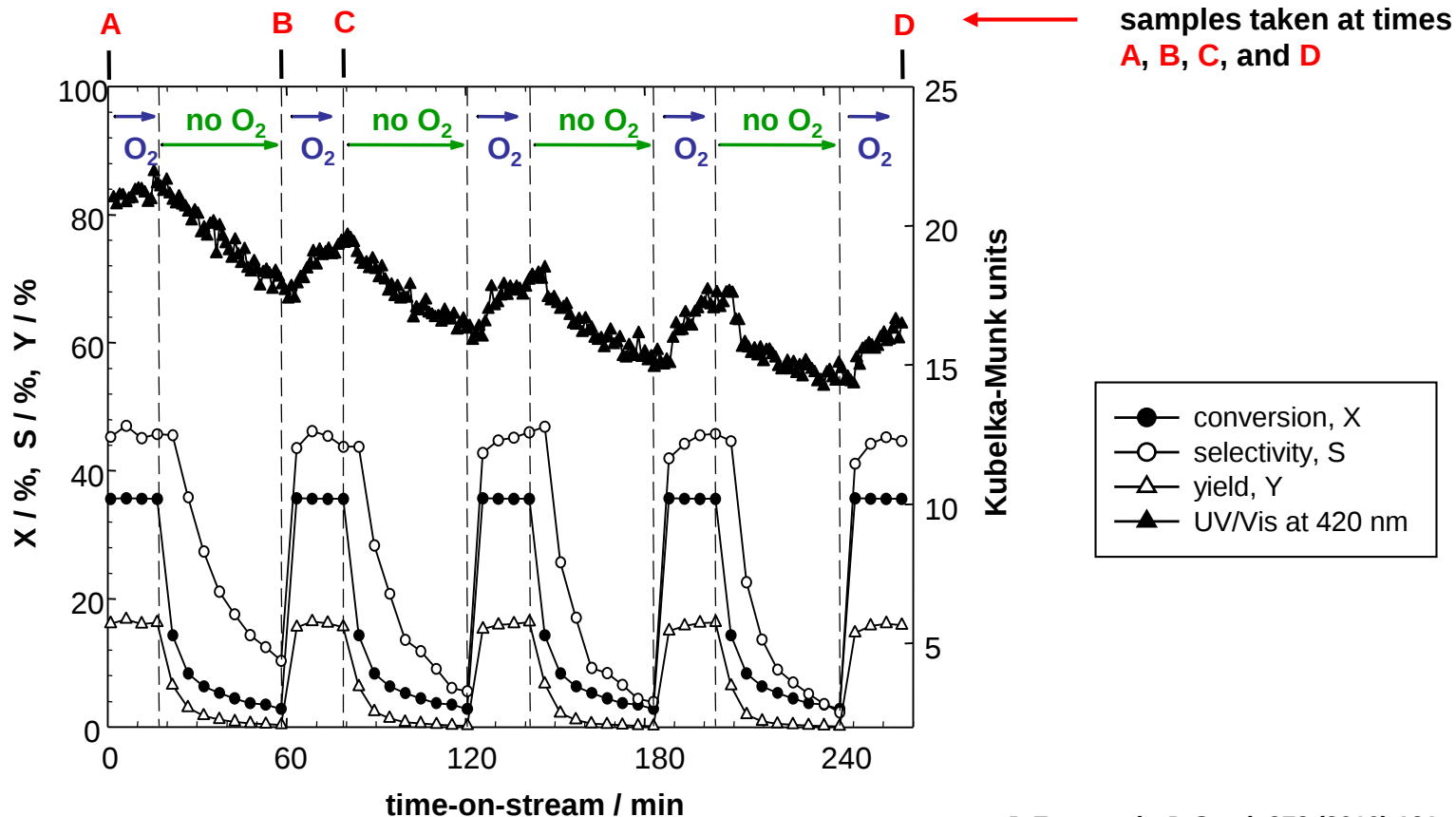


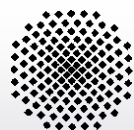


II. Framework modification

VPO catalysts supported on mesoporous SBA-15 materials

selective oxidation of *n*-butane to maleic anhydride at 678 K with sequential switching on (15 min) and off (40 min) of the gaseous oxygen



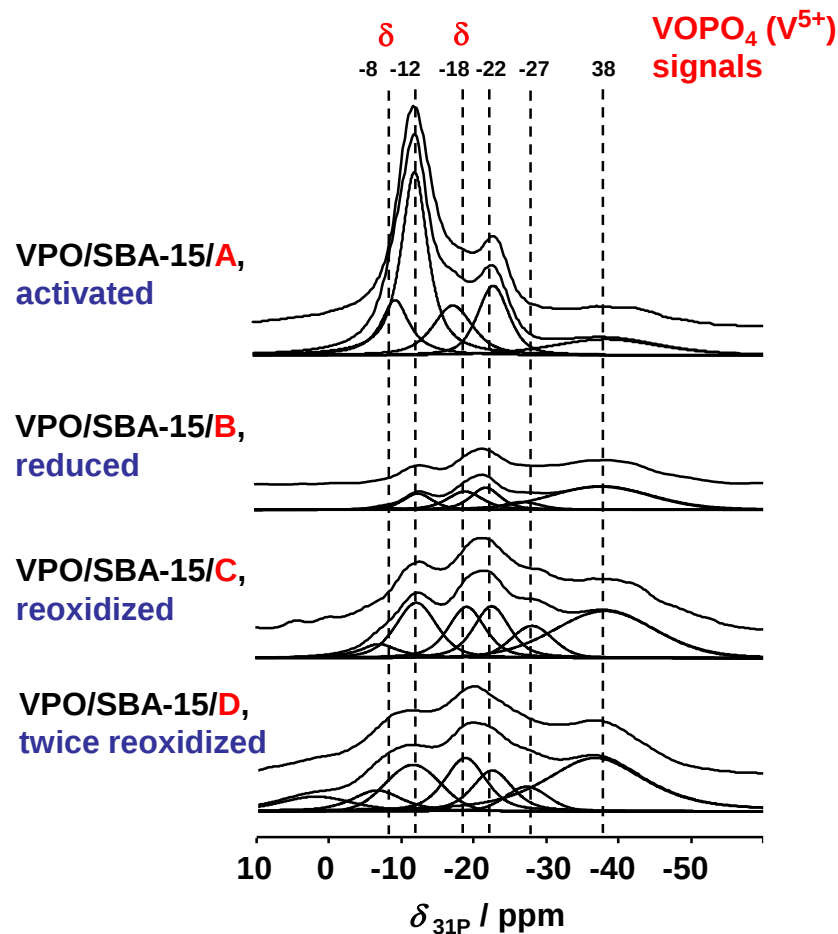


II. Framework modification

VPO catalysts supported on mesoporous SBA-15 materials

^{31}P MAS NMR spectra of VPO/SBA-15/**A** to VPO/SBA-15/**D**

$B_0 = 9.4 \text{ T}$, $\nu_0 = 162.0 \text{ MHz}$



Catalysts	Samples	Contents of P/ V^{5+}
VPO/bulk	A	0.5 %
	B	0 %
	C	0.35 %
	D	0.4 %
VPO/SBA-15	A	2.6 %
	B	0.3 %
	C	0.9 %
	D	1.3 %





III. Tailoring of surface sites



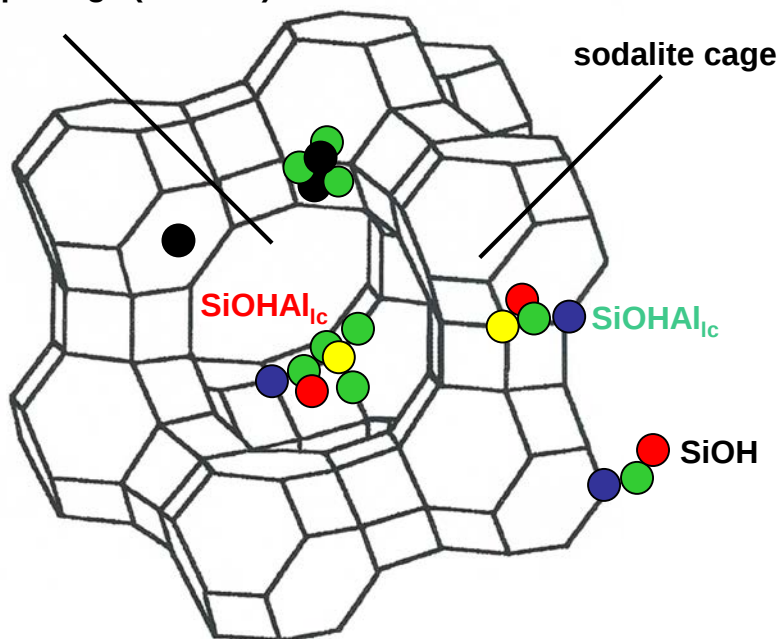


III. Tailoring of surface sites

nature of acid sites on zeolites

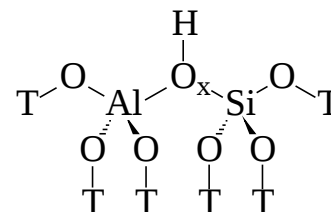
zeolite Y (FAU, faujasite): supercages connected via 12-ring windows (0.74 nm)

supercage (1.23 nm)



- | | | |
|---|---|--|
| ● Al atom | ● O atom | ● extra-framework cations (Na ⁺ , Al ₃ ⁺ etc.) |
| ● Si atom | ● H atom | |

Brønsted acid sites:



bridging OH group (SiOHAl)

Lewis acid sites:

- framework defects,
- extra-framework species (Al oxide clusters)





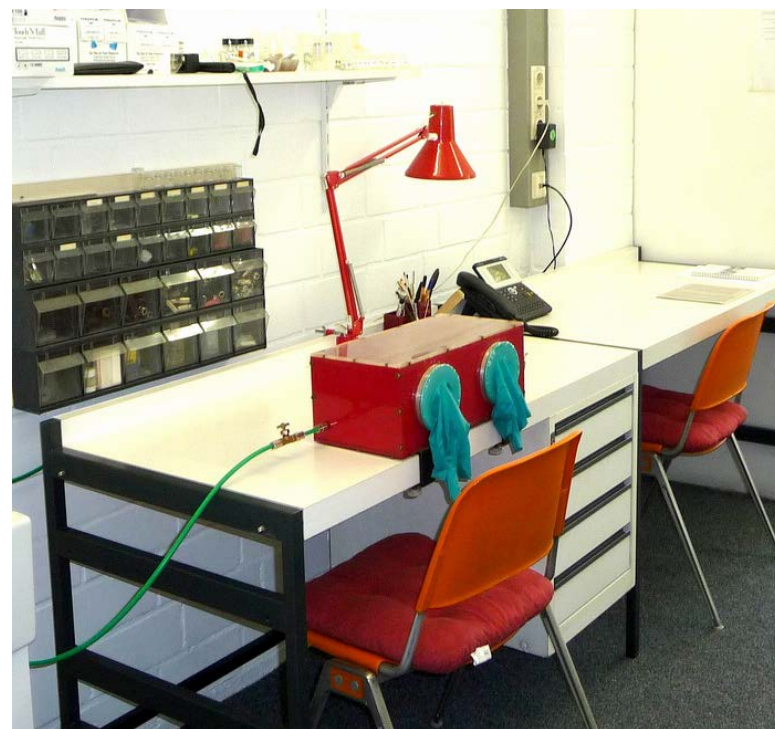
III. Tailoring of surface sites

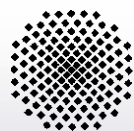
preparation of dehydrated catalyst samples



vacuum line for dehydration of catalyst samples in glass tubes at elevated temperatures (often 673 K)

transfer of the dehydrated catalyst samples from glass tubes into MAS rotors inside a glove box (dry nitrogen)



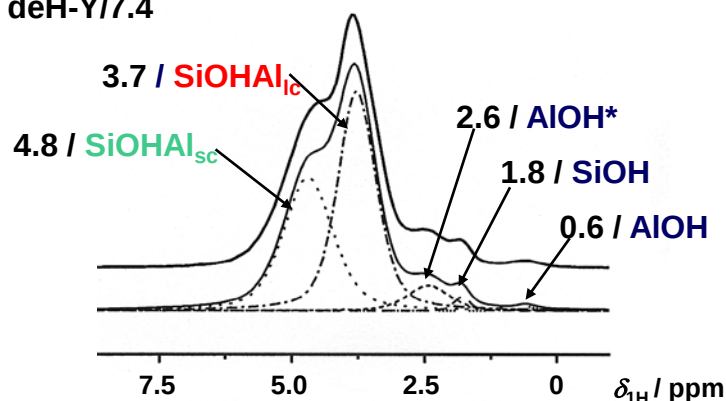


III. Tailoring of surface sites

modification of Brønsted acid sites in zeolites *via* steaming

^1H MAS NMR

deH-Y/7.4



undisturbed metal OH (**AlOH**) groups:
-0.5 to 0.5 ppm

defect **SiOH** groups:
1.2 to 2.2 ppm

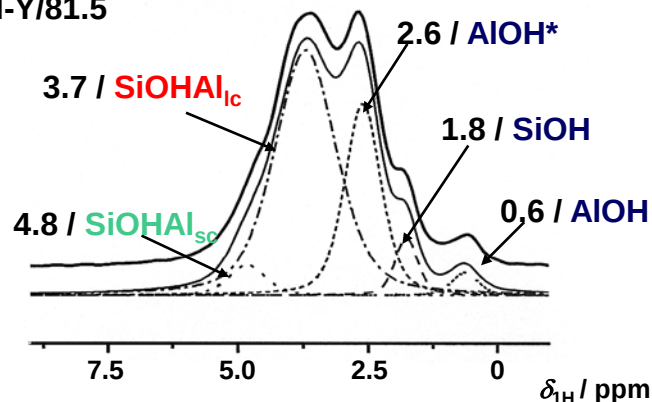
AlOH* groups at extra-framework Al
clusters:
2.8 to 3.6 ppm

bridging OH groups in large cages
and pores (**SiOHAl_{ic}**):
3.6 to 4.3 ppm

bridging OH groups in small cages
(**SiOHAl_{sc}**):
4.6 to 5.2 ppm

(hydrogen bonded SiOH and SiOHAl
groups: 5.2 to 13 ppm)

deH-Y/81.5



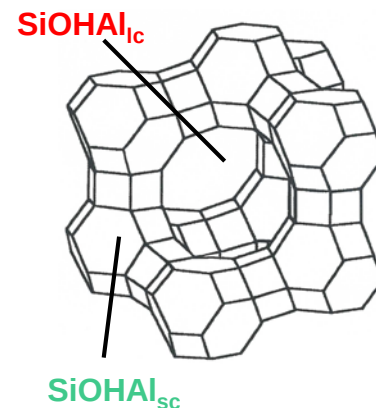
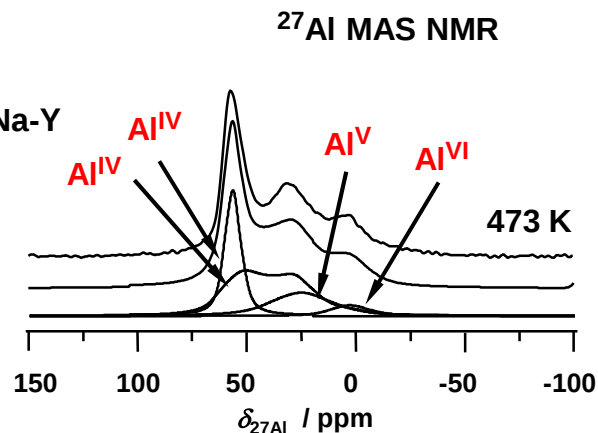
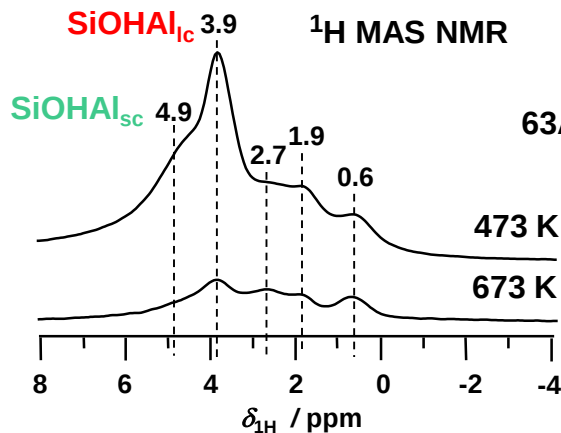
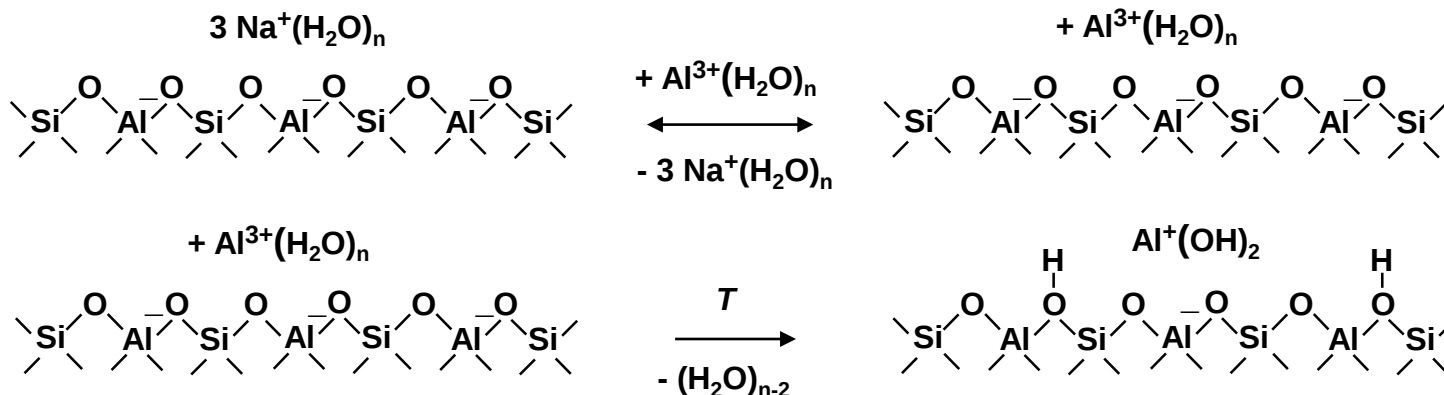


III. Tailoring of surface sites

preparation of Brønsted acid sites in zeolites *via* exchange with multivalent cations

exchange of zeolite Na-Y ($n_{\text{Si}}/n_{\text{Al}} = 2.7$) in 0.1 M $\text{Al}(\text{NO}_3)_3$ solution

Hirschler Plank mechanism



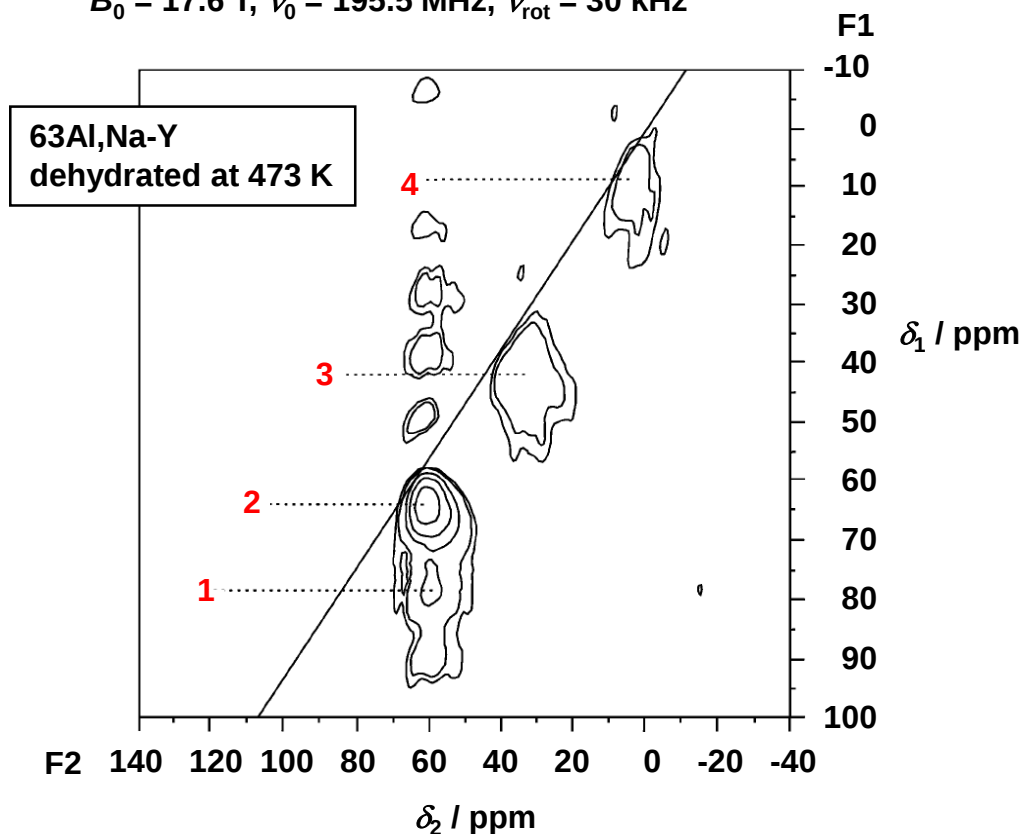


III. Tailoring of surface sites

preparation of Brønsted acid sites in zeolites *via* exchange with multivalent cations

^{27}Al MQMAS NMR (spin $I = 3/2$)

$B_0 = 17.6 \text{ T}$, $\nu_0 = 195.5 \text{ MHz}$, $\nu_{\text{rot}} = 30 \text{ kHz}$



signal 1: Al^{IV}

- $\text{SOQE} = 10.1 \text{ MHz}$

- $\delta_{\text{iso}} = 71 \text{ ppm}$

signal 2: Al^{IV}

- $\text{SOQE} = 3.6 \text{ MHz}$

- $\delta_{\text{iso}} = 62 \text{ ppm}$

signal 3: Al^{V}

- $\text{SOQE} = 7.6 \text{ MHz}$

- $\delta_{\text{iso}} = 39 \text{ ppm}$

signal 4: Al^{VI}

- $\text{SOQE} = 6.2 \text{ MHz}$

- $\delta_{\text{iso}} = 6 \text{ ppm}$

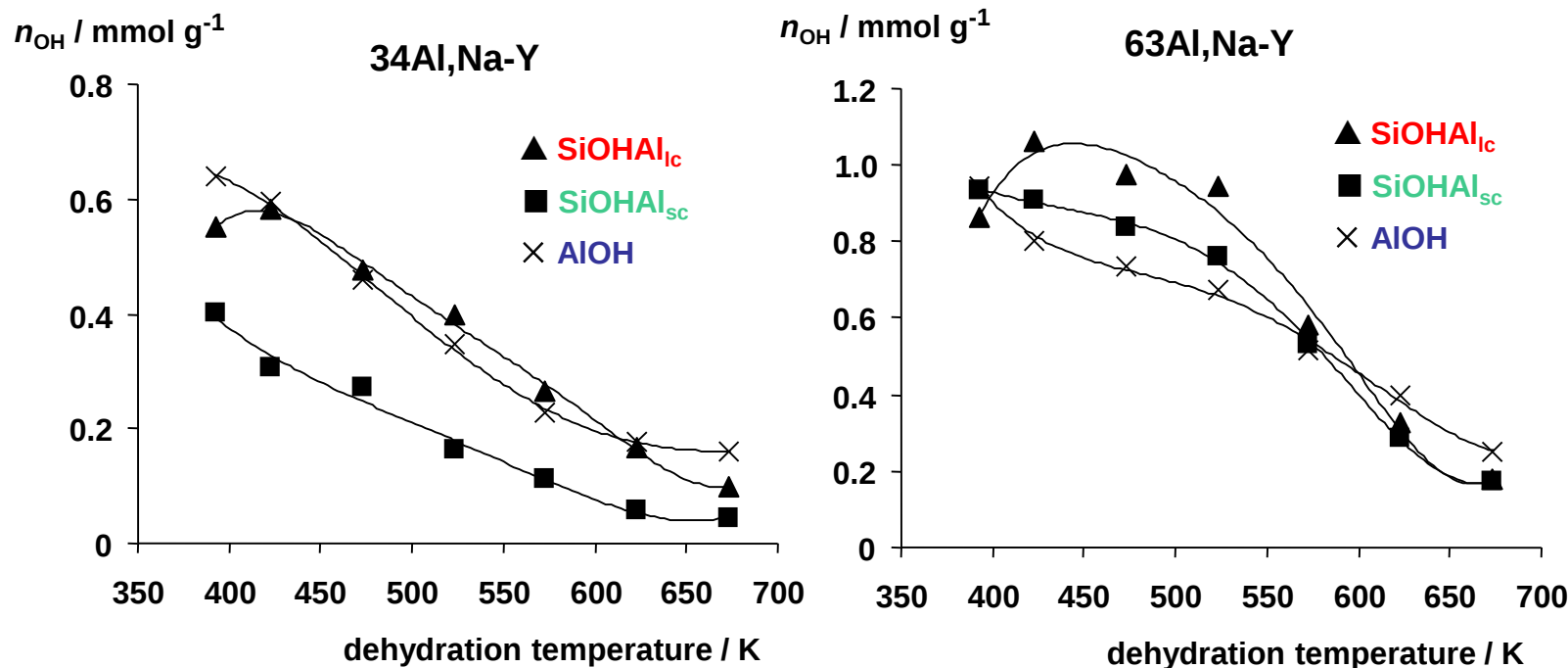




III. Tailoring of surface sites

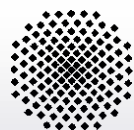
Brønsted acid sites in zeolites prepared via exchange with multivalent cations (Al^{3+})

determination of the concentration of OH groups by evaluation of the ^1H MAS NMR intensities



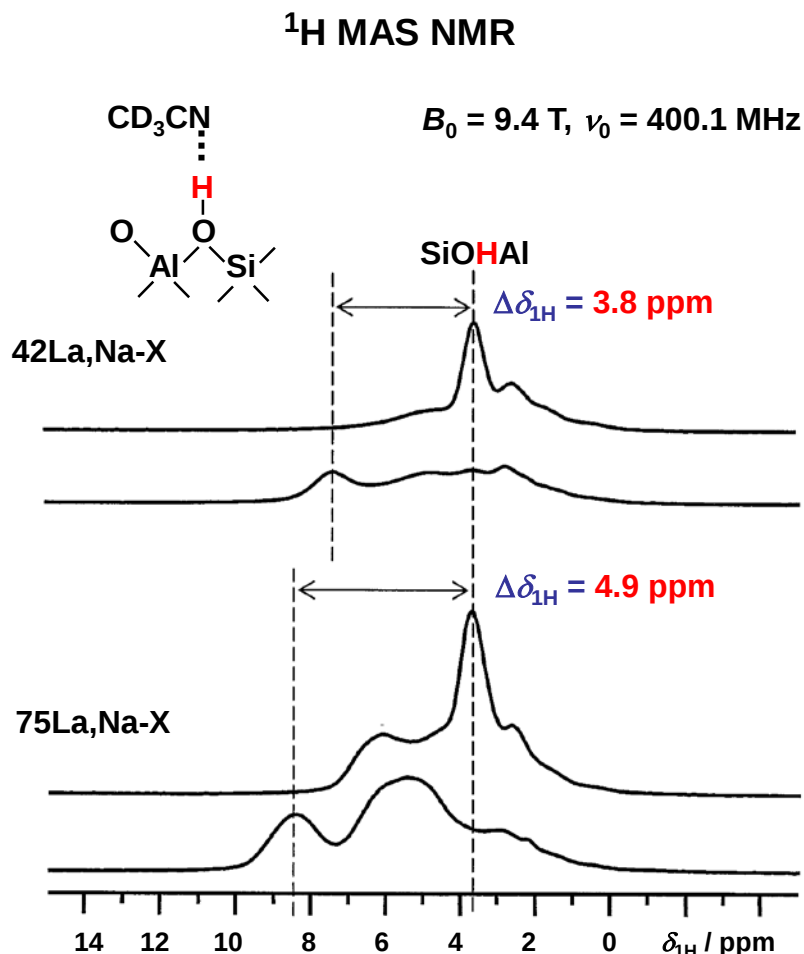
cation-exchange degree and dehydration temperature affect the number of Brønsted sites





III. Tailoring of surface sites

characterization of the Brønsted acid strength via acetonitrile-induced low-field shift $\Delta\delta_{1H}$



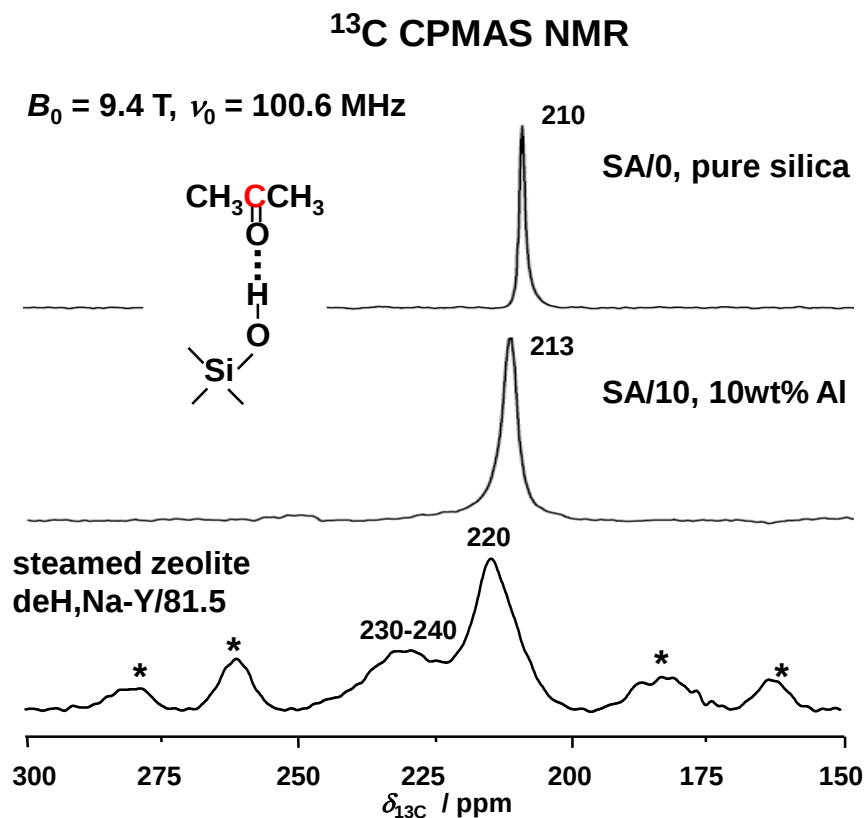
Low-field shift $\Delta\delta_{1H}$	Adsorbent and type of OH group
1.2 ppm	AlOH in MIL-53(Al)
3.6 ppm	H,Na-X (Si/Al = 1.3)
3.8 ppm	42La,Na-X and 32Al,Na-X (Si/Al = 1.4)
4.9 ppm	75La,Na-X (Si/Al = 1.4)
5.1 ppm	H,Na-Y (Si/Al = 2.7)
5.3 ppm	34Al,Na-Y and 63Al,Na-Y (Si/Al = 2.7)
5.7 ppm	42La,Na-Y and 75La,Na-Y (Si/Al = 2.7)
6.5 ppm	steamed deH,Na-Y/81.5 (Si/Al = 6)
6.7 ppm	H-MOR (Si/Al = 10)
7.0 ppm	dealuminated H,Na-Y (Si/Al = 18)
7.9 ppm	H-ZSM-5 (Si/Al = 26)





III. Tailoring of surface sites

characterization of Brønsted and Lewis acid sites *via* ^{13}C -2-acetone as probe molecule



Materials	$\delta_{^{13}\text{C}}$	$\Delta\delta_{^{13}\text{C}}$
CDCl_3	205 ppm	0 ppm
SA/0	210 ppm	5 ppm
SA/10	213 ppm	8 ppm
H,Na-X	215 ppm	10 ppm
SA/70	216 ppm	11 ppm
H,Na-Y	220 ppm	15 ppm
ZSM-5	223 ppm	18 ppm
Lewis sites	230-240 ppm	25-35 ppm

J.F. Haw et al., Accounts of Chemical Research 29 (1996) 259.

Lewis sites formed by steaming



Brønsted sites with different acid strengths (see Tab.)

Y. Jiang et al., Solid State Nucl. Magn. Reson. 39 (2011) 116-141.

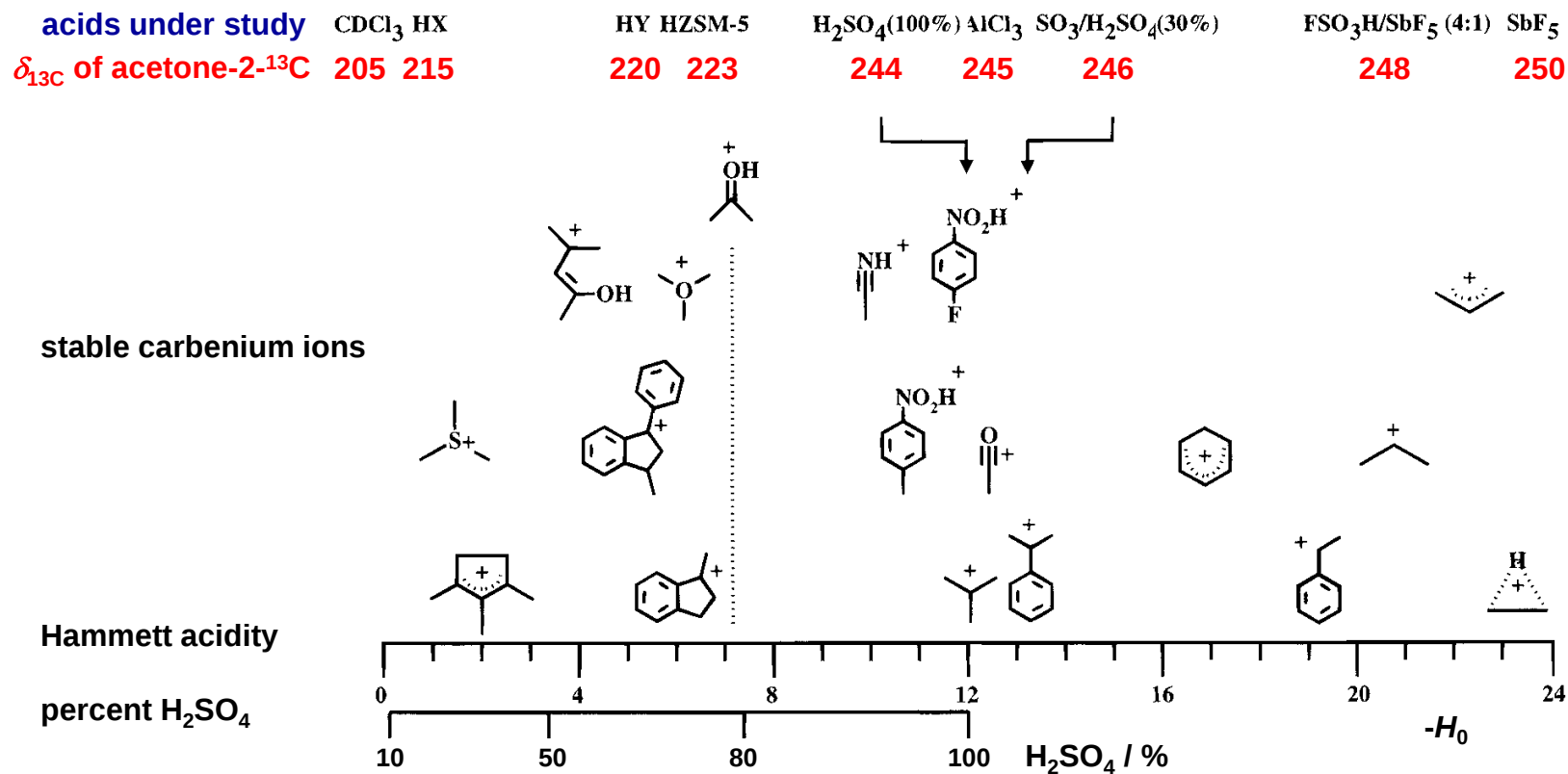
H. Fang et al., J. Phys. Chem. C 114 (2010) 12711-12718.





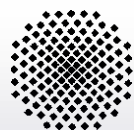
III. Tailoring of surface sites

comparison of different acidity scales



J.F. Haw et al., Acc. Chem. Res. 29 (1996) 259.



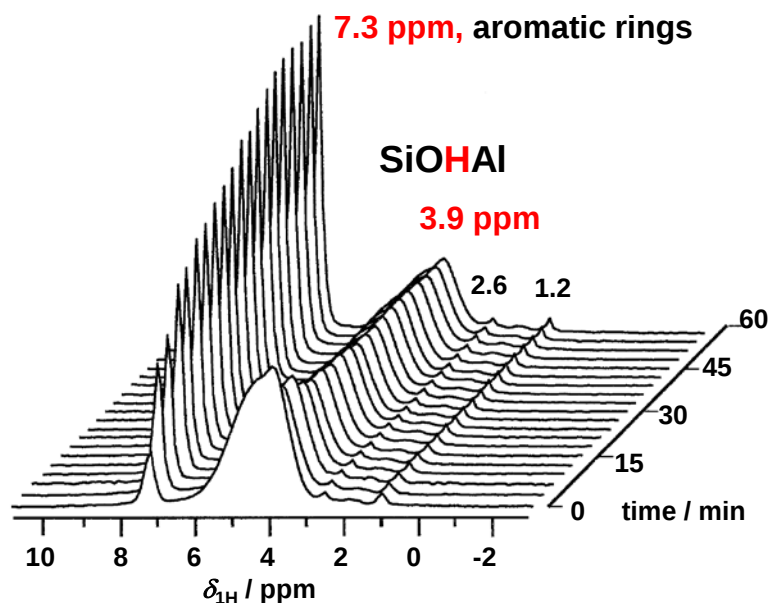


III. Tailoring of surface sites

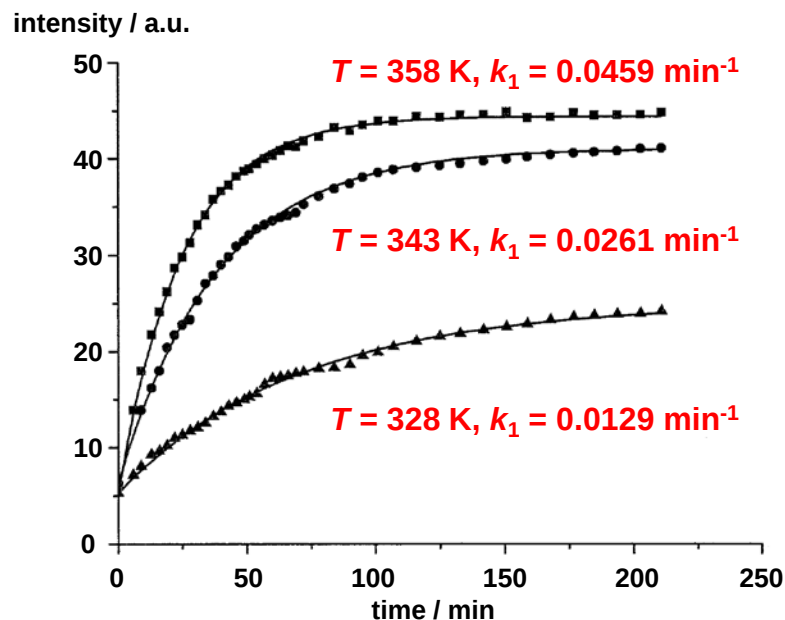
H/D exchange kinetics as an acidity scale

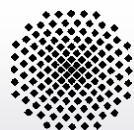
in situ ^1H MAS NMR studies of H/D exchange in zeolite $\text{H}_2\text{Na-Y}$ loaded with ethylbenzene ($\text{C}_6\text{D}_5\text{C}_2\text{H}_5$)

stack plot of ^1H MAS NMR spectra recorded at $T = 358\text{ K}$
 $B_0 = 9.4\text{ T}$, $\nu_0 = 400.1\text{ MHz}$



H/D exchange rates at $T = 328 - 358\text{ K}$



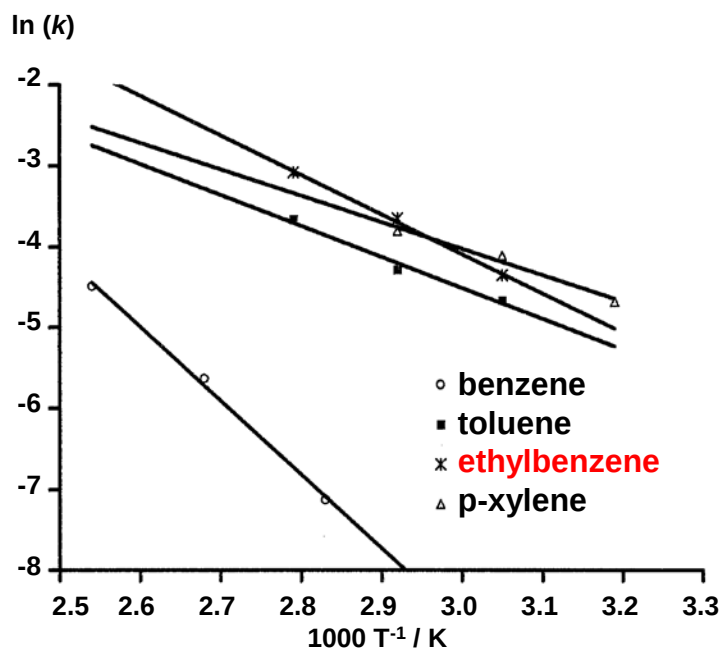


III. Tailoring of surface sites

H/D exchange kinetics as an acidity scale

in situ ^1H MAS NMR spectroscopy of H/D exchange in zeolites H,Na-Y ($n_{\text{Si}}/n_{\text{Al}} = 2.7$), 75La,Na-Y ($n_{\text{Si}}/n_{\text{Al}} = 2.7$), and H-ZSM-5 ($n_{\text{Si}}/n_{\text{Al}} = 26$)

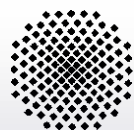
deuterated aromatics on zeolite H,Na-Y



activation energies E_A of H/D exchange and low-field shifts $\Delta\delta_{1\text{H}}$ upon adsorption of CD_3CN :

catalyst	molecule	E_A / kJ mol ⁻¹	$\Delta\delta_{1\text{H}}$ / ppm
H,Na-Y	benzene	76	
	ethylbenzene	41	
	toluene	32	
	p-xylene	27	
H,Na-Y	benzene	76	5.1
La,Na-Y	benzene	67	5.7
H-ZSM-5	benzene	46	7.9



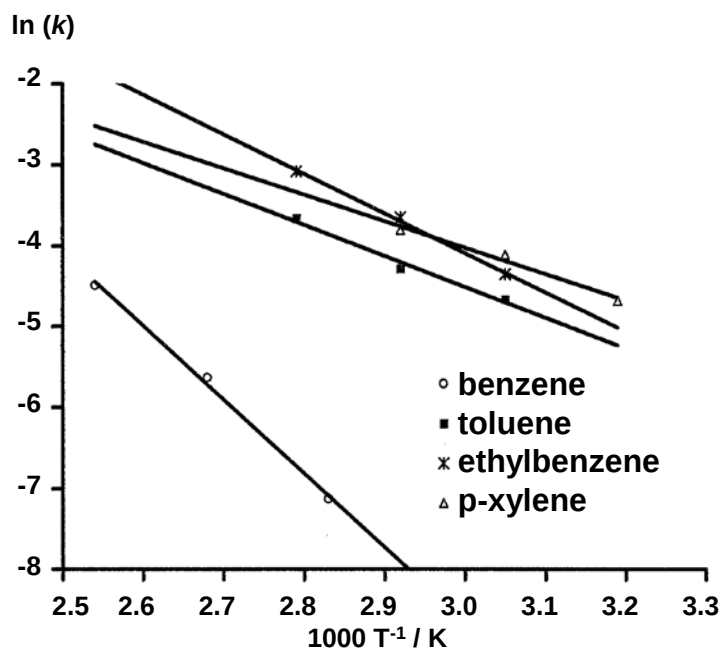


III. Tailoring of surface sites

H/D exchange kinetics as an acidity scale

in situ ^1H MAS NMR spectroscopy of H/D exchange in zeolites H,Na-Y ($n_{\text{Si}}/n_{\text{Al}} = 2.7$), 75La,Na-Y ($n_{\text{Si}}/n_{\text{Al}} = 2.7$), and H-ZSM-5 ($n_{\text{Si}}/n_{\text{Al}} = 26$)

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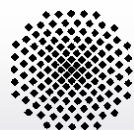
catalyst	molecule	E_A / kJ mol ⁻¹	$\Delta\delta_{1\text{H}}$ / ppm
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	p-xylene	27	
H,Na-Y	benzene	76	5.1
La,Na-Y	benzene	67	5.7
H-ZSM-5	benzene	46	7.9





IV. Catalytic investigations

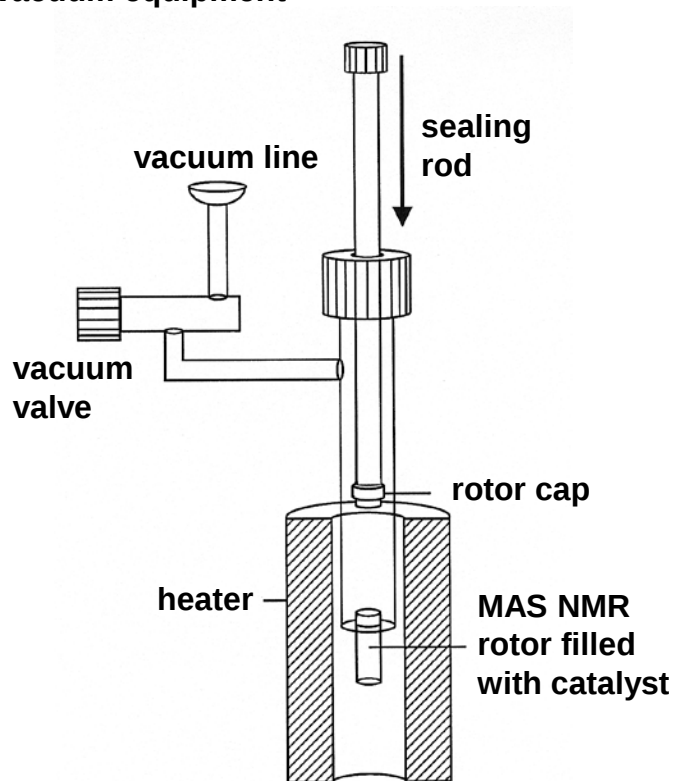




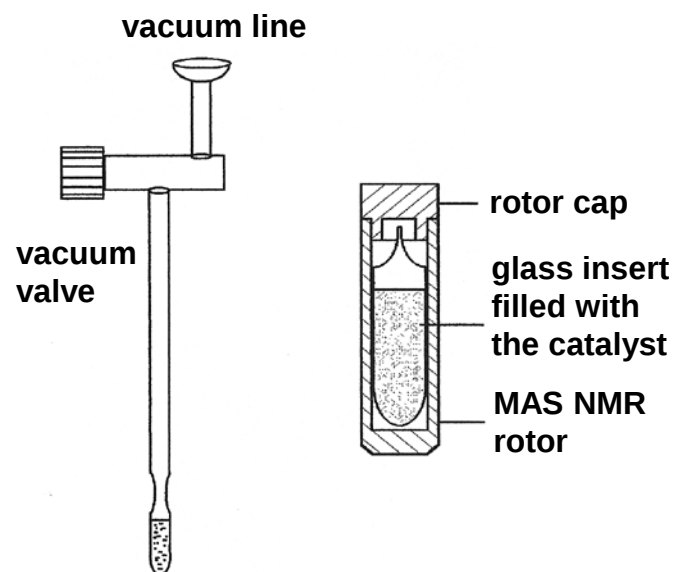
IV. Catalytic investigations

preparation of catalyst samples for *in situ* MAS NMR studies under **batch** conditions

dehydration, loading, and sealing of the catalyst filled in an MAS NMR rotor inside a vacuum equipment



dehydration and loading of the catalyst inside the glass insert (e.g. commercial Wilmad MAS NMR inserts for 4 mm and 7 mm rotors)

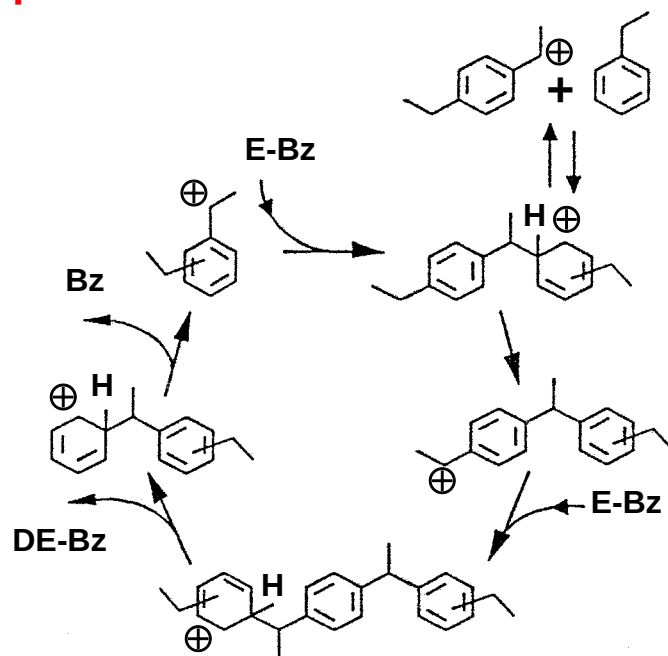




IV. Catalytic investigations

mechanisms of the ethylbenzene disproportionation

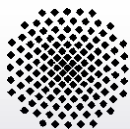
Streitwieser-Reif mechanism for the homogeneously catalyzed ethylbenzene disproportionation



suggested reaction mechanisms for the heterogeneously catalyzed reaction on zeolites:

- via diphenylethane intermediates in **large-pore zeolites**
- dealkylation/realkylation with free ethene or alkoxy species in **medium-pore zeolites**





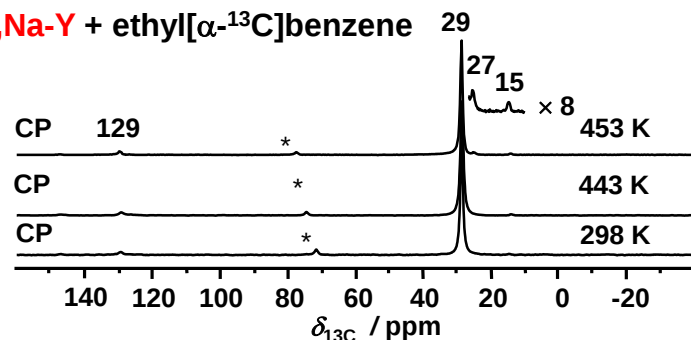
IV. Catalytic investigations

mechanisms of the ethylbenzene disproportionation

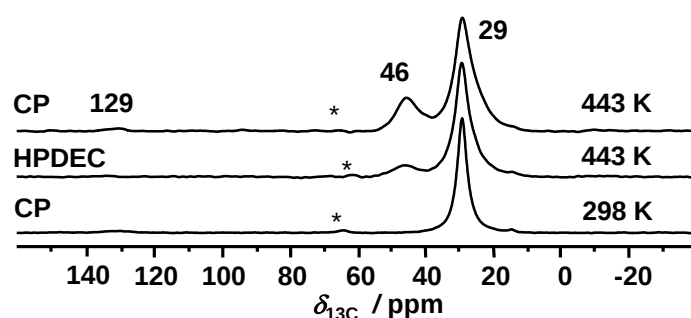
^{13}C MAS NMR investigation of the ethylbenzene conversion on large-pore zeolites H,Na-Y and 63Al,Na-Y

^{13}C MAS NMR

H,Na-Y + ethyl[α - ^{13}C]benzene



63Al,Na-Y + ethyl[α - ^{13}C]benzene



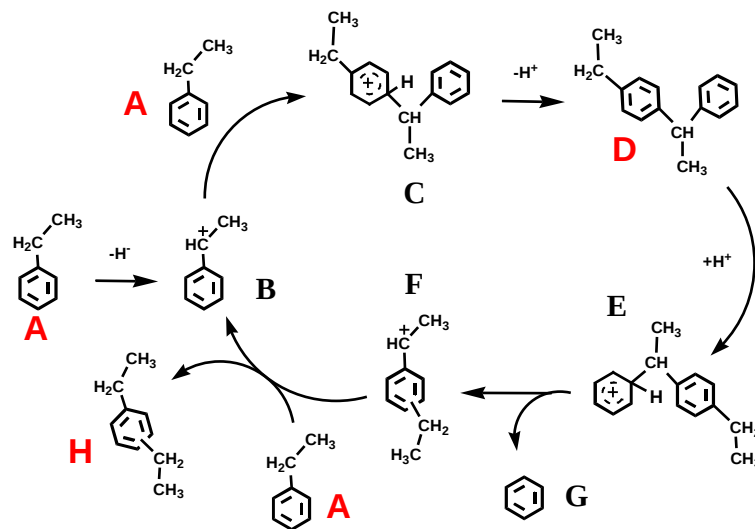
assignment: 27 ppm
29 ppm

diethylbenzene (H)
ethylbenzene (A)

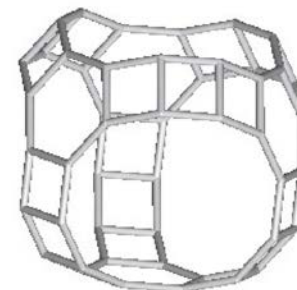
46 ppm
129 ppm

diphenylethane (D)
aromatic carbons

reaction mechanism



in supercages with
diameter of 1.2 nm

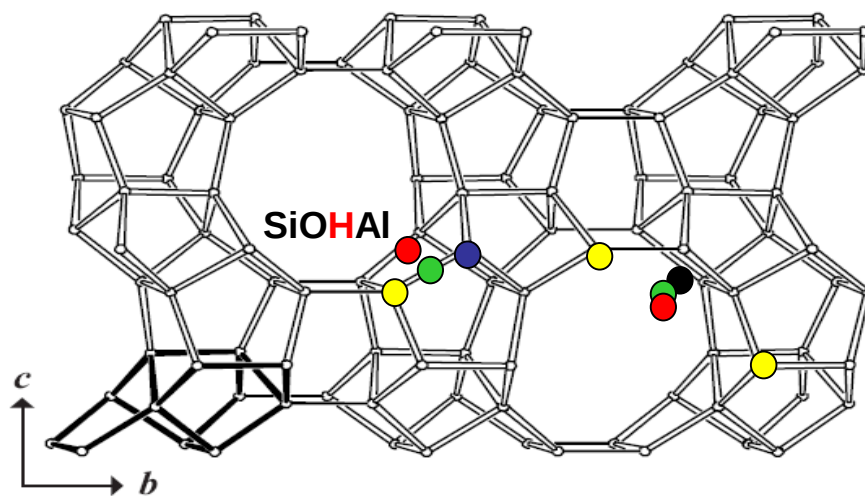




IV. Catalytic investigations

mechanisms of the ethylbenzene disproportionation

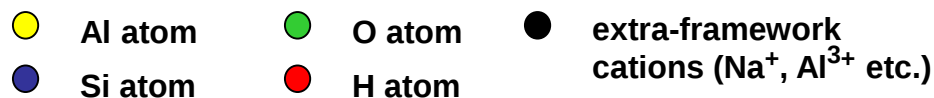
structure of medium-pore zeolite ZSM-5:

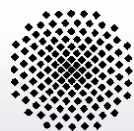


crossing intersections with
interconnecting 10-ring
channels

[100] 0.51 nm x 0.55 nm

[010] 0.53 nm x 0.56 nm





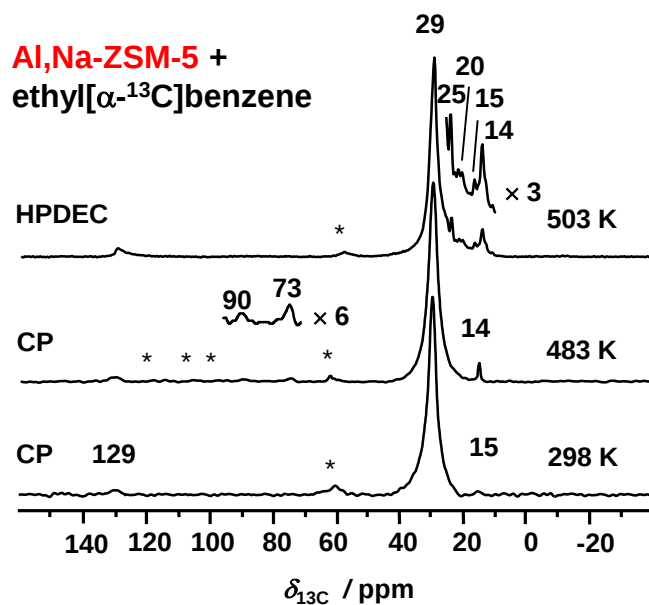
IV. Catalytic investigations

mechanism of the ethylbenzene disproportionation

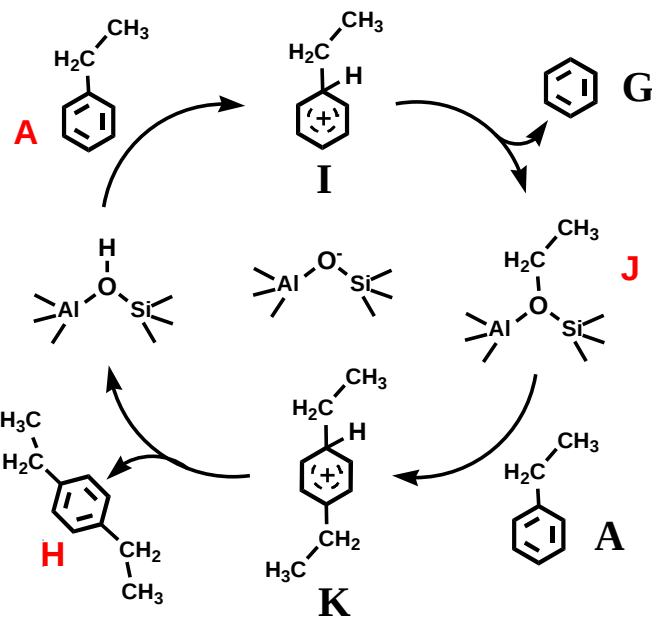
^{13}C MAS NMR investigation of the ethylbenzene conversion on medium-pore zeolite ZSM-5

^{13}C MAS NMR
mechanism

Al,Na-ZSM-5 +
ethyl[α - ^{13}C]benzene



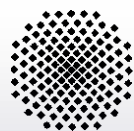
reaction



assignment: 25 ppm diethylbenzene (**H**)
29 ppm ethylbenzene (**A**)

73 ppm surface ^{13}C -1-ethoxy groups (**J**)
90 ppm oligomeric alkoxy groups
129 ppm aromatic carbons

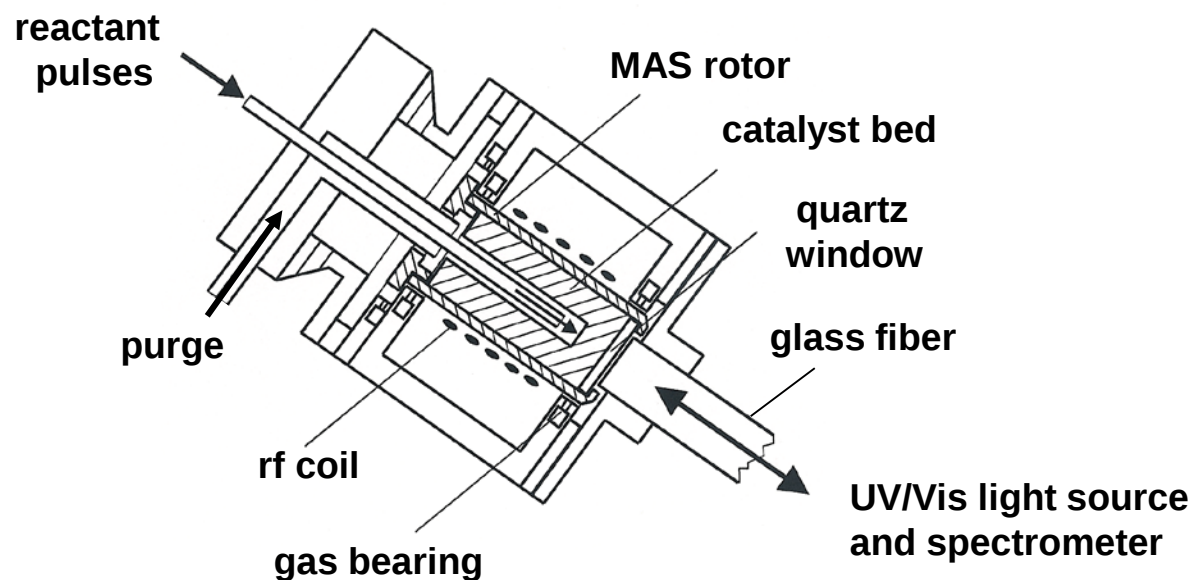




IV. Catalytic investigations

continuous-flow (CF) and pulsed-flow (PF) MAS NMR spectroscopy

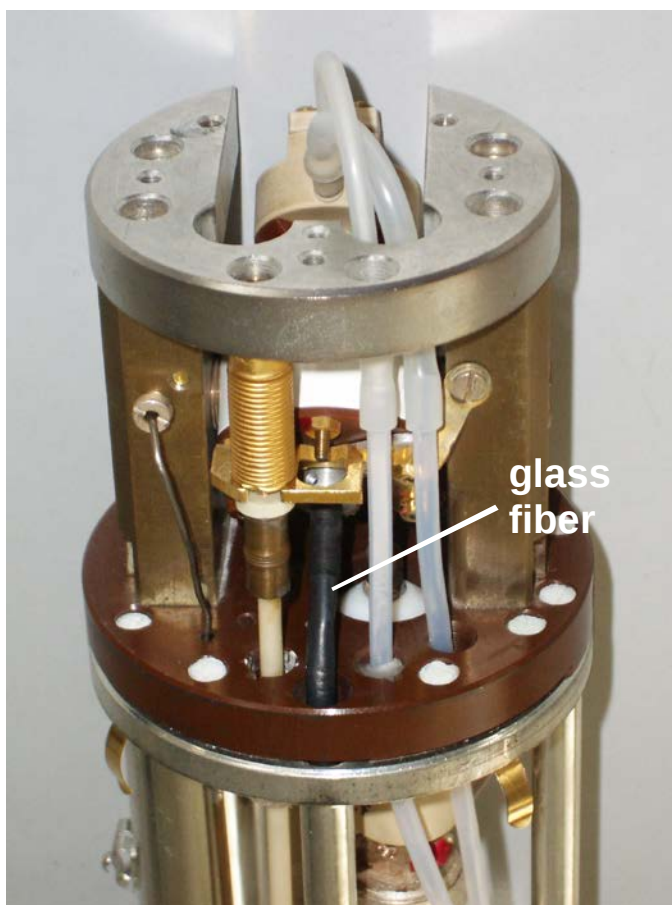
coupling of MAS NMR and UV/Vis spectroscopy by installation of a quartz fiber at the bottom of the MAS NMR stator





IV. Catalytic investigations

continuous-flow (CF) and pulsed-flow (PF) MAS NMR spectroscopy



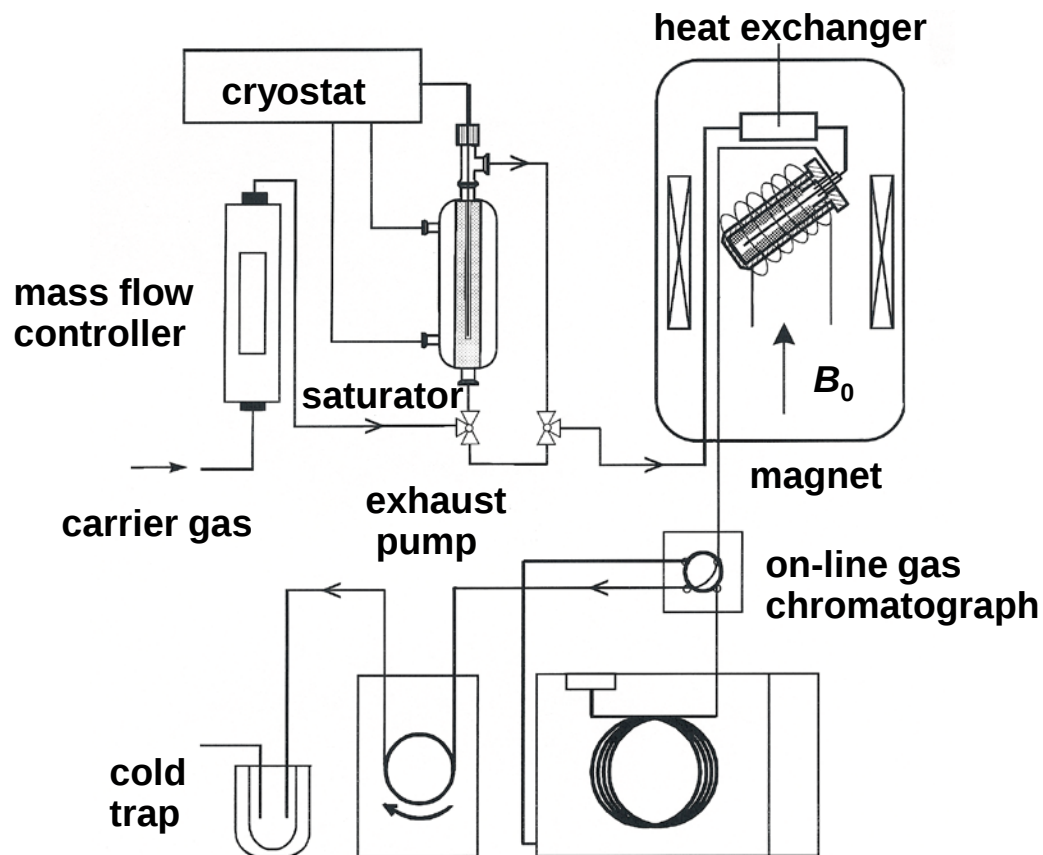
7 mm Bruker MAS NMR probe
equipped with a glass fiber (left) and
UV/Vis light source and spectrometer
of Avantes (bottom)





IV. Catalytic investigations

continuous-flow (CF) MAS NMR spectroscopy

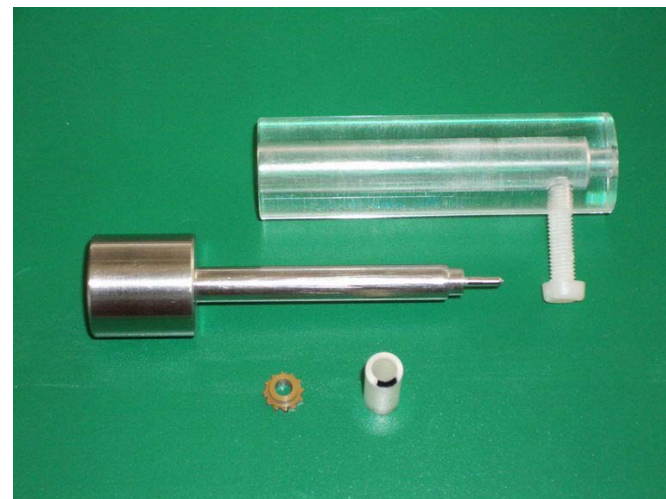
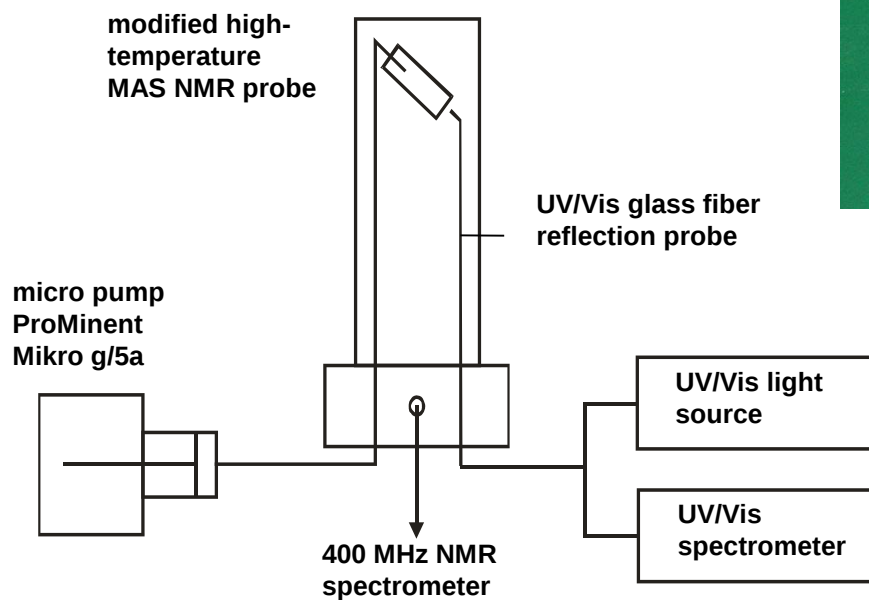




IV. Catalytic investigations

pulsed-flow (PF) MAS NMR spectroscopy

connection of a micro pump with the *in situ* MAS NMR probe



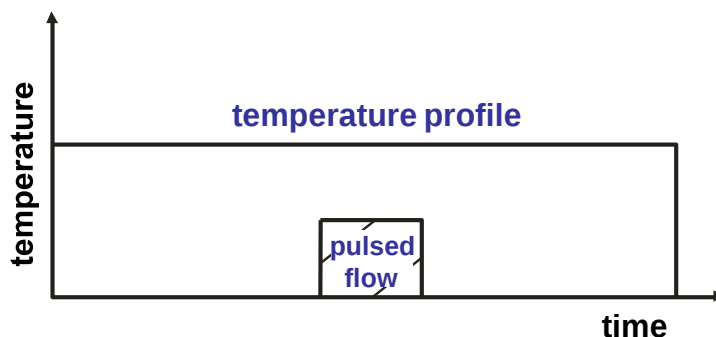


IV. Catalytic investigations

pulsed-flow (PF) MAS NMR-UV/Vis spectroscopy

injection of liquid reactants into the spinning MAS NMR rotor *via* a micro-pulse pump

pulsed-flow experiment



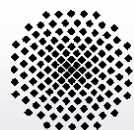
pulsed-flow experiments:

- study of the time dependence of the conversion of reactants
- study of the isotopic exchange of reactants at high temperatures with well-defined starting time



pump Mikro g/5 of Fa. ProMinent, Germany, for single pulses with volumes of 2 to 50 μl

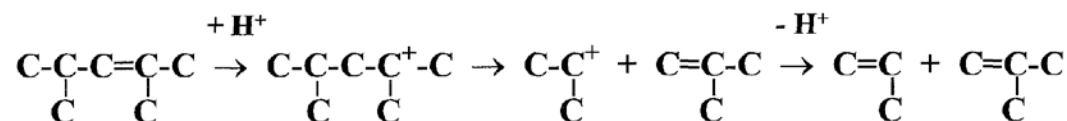
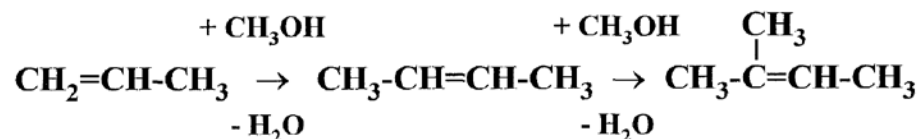




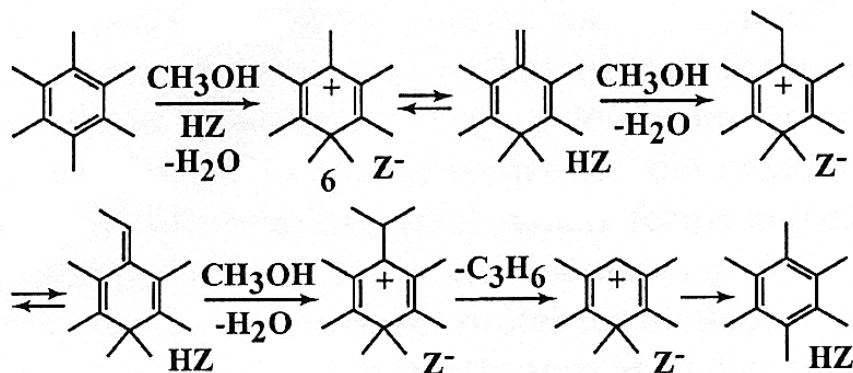
IV. Catalytic investigations

mechanism of the methanol-to-olefin (MTO) conversion on acidic zeolites

via olefinic compounds (H. Schulz, M. Wei, Microporous Mesoporous Mater. 29 (1999) 205)



via polyalkylaromatics (J.F. Haw et al., Acc. Chem. Res. 36 (2003) 317)





IV. Catalytic investigations

mechanism of the methanol-to-olefin (MTO) conversion on acidic zeolites

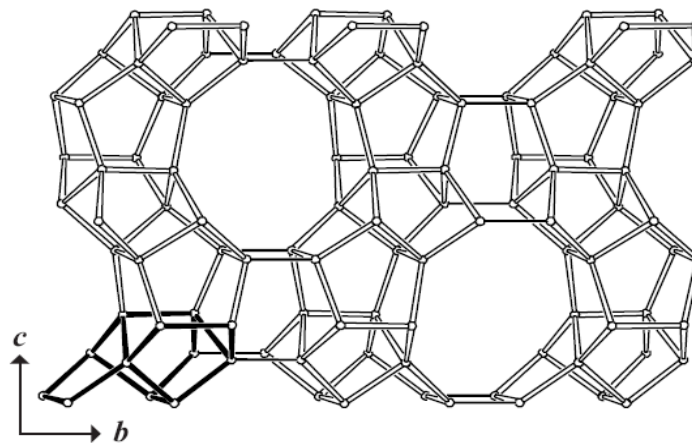
H-ZSM-5: Structure type MFI



crossing intersections at
interconnecting 10-ring channels

[100] 0.51 nm x 0.55 nm

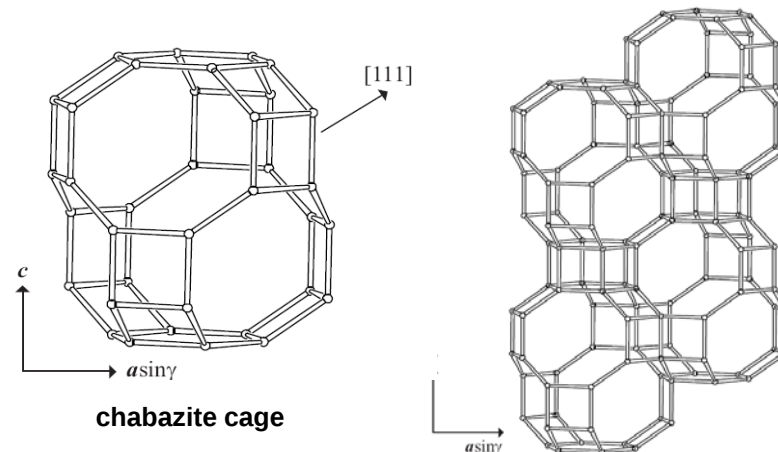
[010] 0.53 nm x 0.56 nm



H-SAPO-34: Structure type CHA



chabazite cages accessible by
8-ring windows perpendicular to
[001] 0.38 nm x 0.38 nm

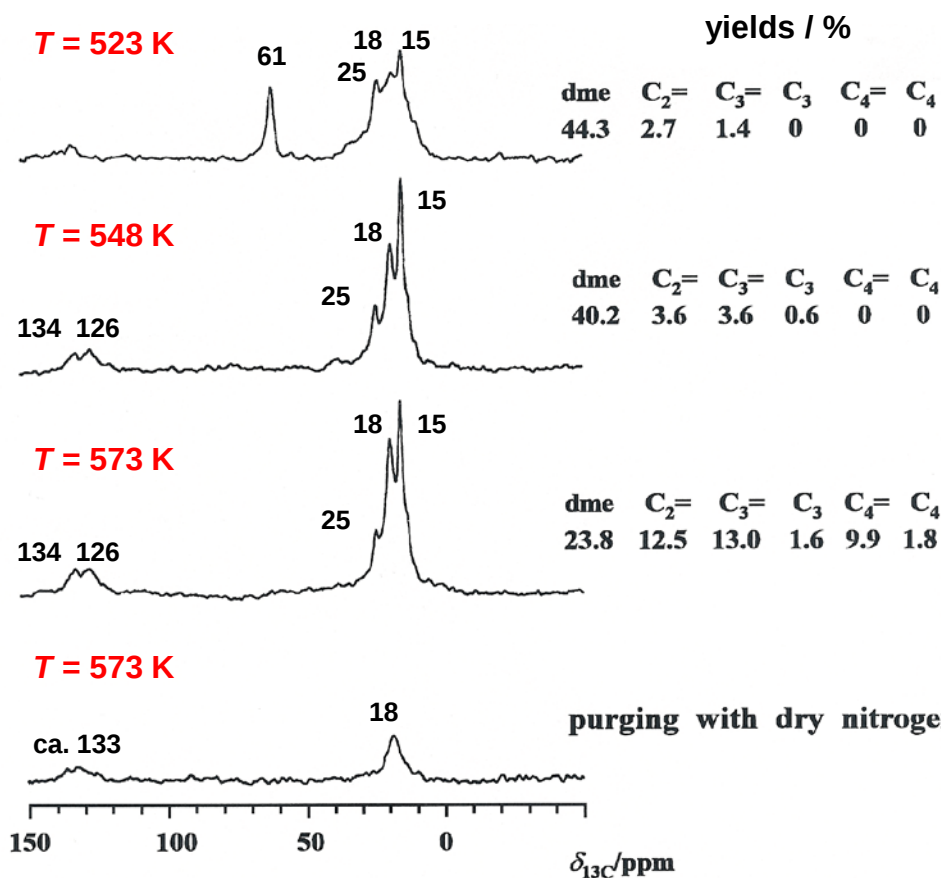




IV. Catalytic investigations

mechanism of the methanol-to-olefin (MTO) conversion on acidic zeolites

in situ **CF** ^{13}C MAS NMR study of ZSM-5 ($W_{\text{cat}}/F_{\text{me}} = 25 \text{ gh/mol}$)



$T \leq 548 \text{ K}$:

mixture of olefinic and aromatic compounds, such as:

3-hexene (14.4, 25.9, 131.2 ppm)

2,4-hexadiene (17.2, 126.2, 132.5 ppm)

....

hexamethylbenzene (17.6, 132.1 ppm)

$T = 573 \text{ K}$:

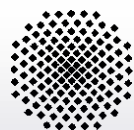
polymethylaromatics remain after purging, such as:

toluene (20.3, 128.5, 129.0 ppm)

....

hexamethylbenzene (17.6, 132.1 ppm)



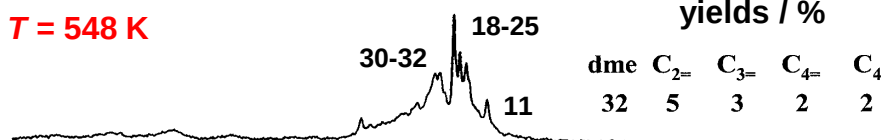


IV. Catalytic investigations

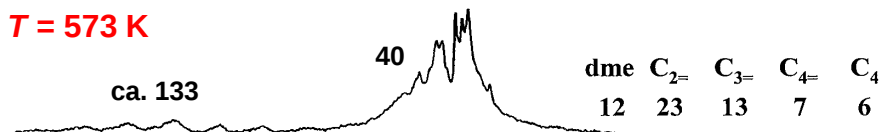
mechanism of the methanol-to-olefin (MTO) conversion on acidic zeolites

in situ **CF** ^{13}C MAS NMR study of SAPO34 ($W_{\text{cat}}/F_{\text{me}} = 25 \text{ gh/mol}$)

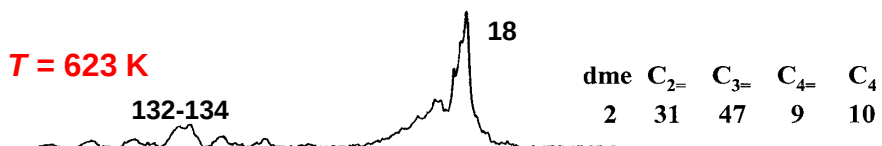
$T = 548 \text{ K}$



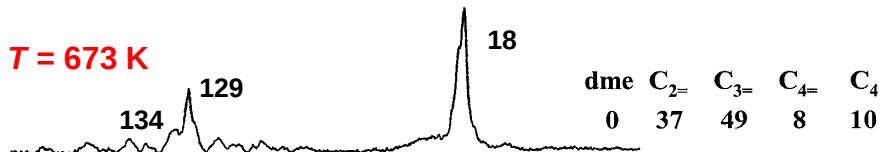
$T = 573 \text{ K}$



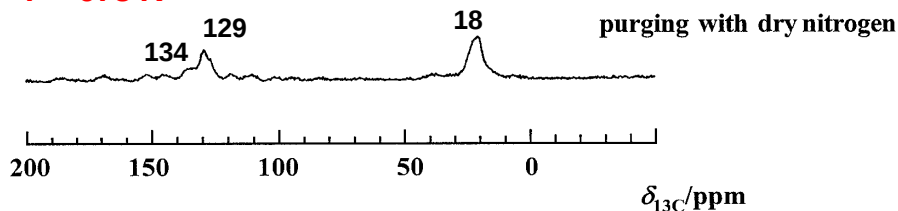
$T = 623 \text{ K}$



$T = 673 \text{ K}$



$T = 673 \text{ K}$



$T \leq 573 \text{ K}$:

mixture of olefinic and aromatic compounds, such as:

2-methyl-2-butene (9.3-22.5, 118.8, 131.8 ppm)

2,4-hexadiene (19.5, 132.5 ppm)

....

tetramethylbenzene (18.9, 131.1, 134.0 ppm)

$T > 573 \text{ K}$:

domination of polymethyl-aromatics, such as:

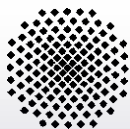
toluene (20.3, 128.5, 129.0 ppm)

....

trimethylbenzene (21.2, 127.4, 137.6 ppm)

hexamethylbenzene (17.6, 132.1 ppm)



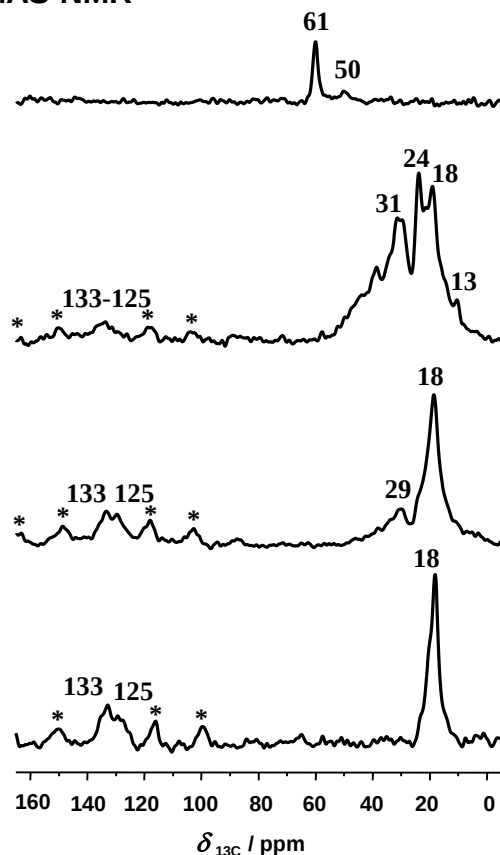


IV. Catalytic investigations

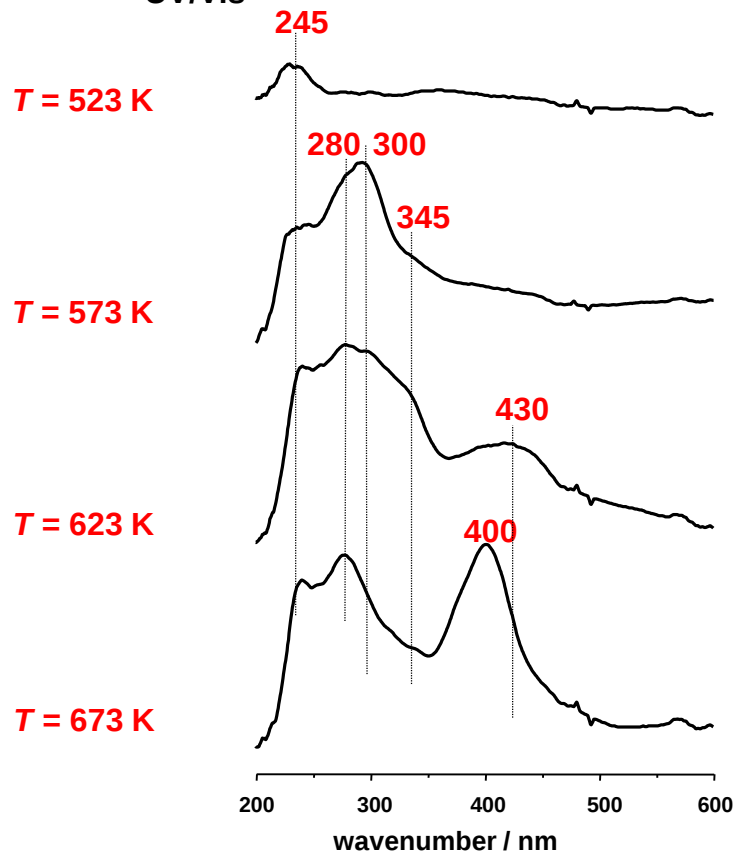
mechanism of the methanol-to-olefin (MTO) conversion on acidic zeolites

comparing study of organic deposits formed on H-SAPO-34 ($W_{\text{cat}}/F_{\text{me}} = 25 \text{ gh/mol}$)

^{13}C MAS NMR



UV/Vis



245 nm: dienes

280 nm: polyalkylaromatics

300 nm: monoenylic carbenium

345 nm: dienylic carbenium

400 nm: polycyclic coke

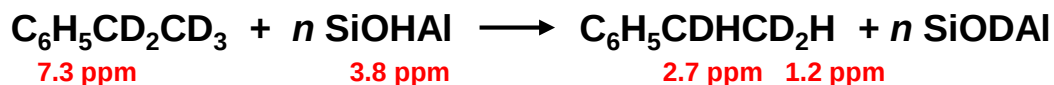
430 nm: trienylic carbenium





IV. Catalytic investigations

side-chain H/D exchange of ethylbenzene on steamed zeolite deH,Na-Y/81.5



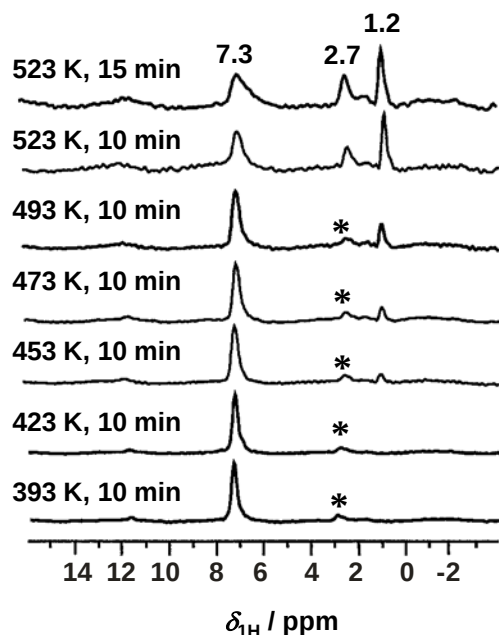
in situ **PF** ^1H MAS NMR experiments:

- pulses of 7.8 mg ethyl- d_5 -benzene
- 32 scans per spectrum with repetition time of 10 s at 9.4 T
- sample spinning rate of ca. 2 kHz

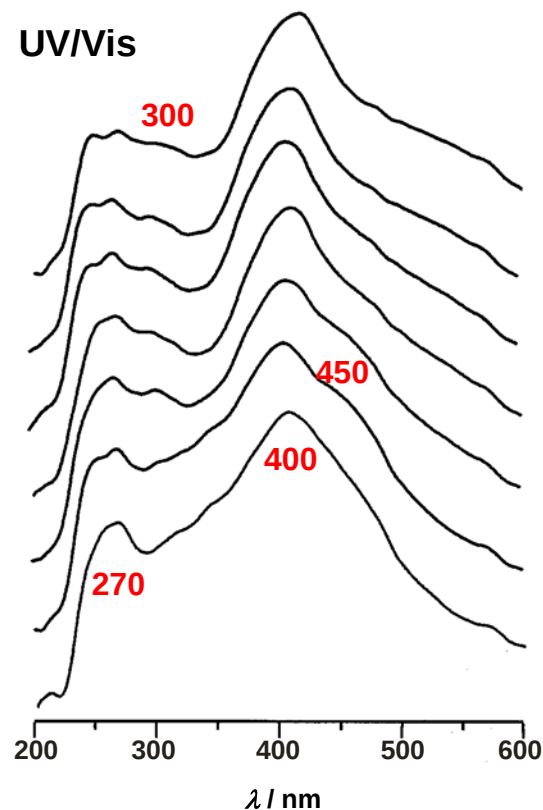
message:

- regioselective H/D exchange at 443 to 463 K (^1H MAS NMR)
- different types of carbenium ions (UV/Vis)

^1H MAS NMR



UV/Vis



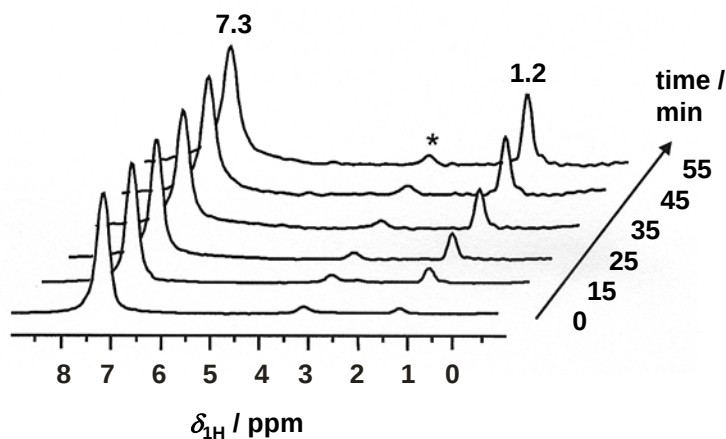


IV. Catalytic investigations

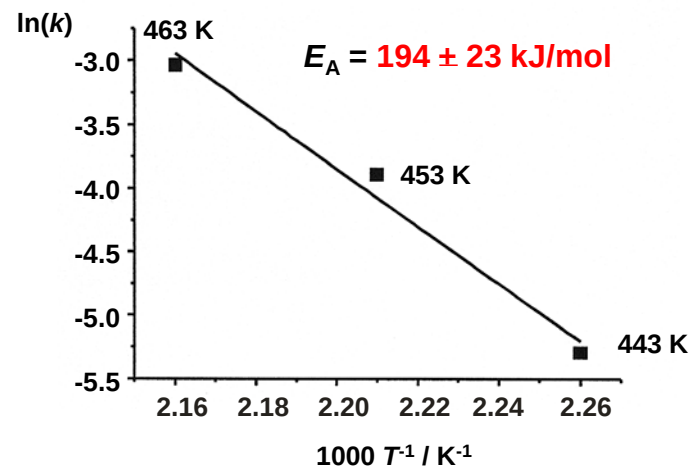
side-chain H/D exchange of ethylbenzene on steamed zeolite deH,Na-Y/81.5

$\text{C}_6\text{H}_5\text{CD}_2\text{CD}_3$ on dealuminated zeolite deH,Na-Y/81.5 (24.5 Al^{ex}/u.c, 10.9 SiOHAl /u.c)

in situ ^{1}H MAS NMR

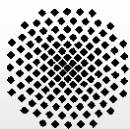


Arrhenius plot



activation energy of the regioselective H/D exchange (194 kJ/mol) indicates that a hydride transfer reaction is the rate determining step





IV. Catalytic investigations

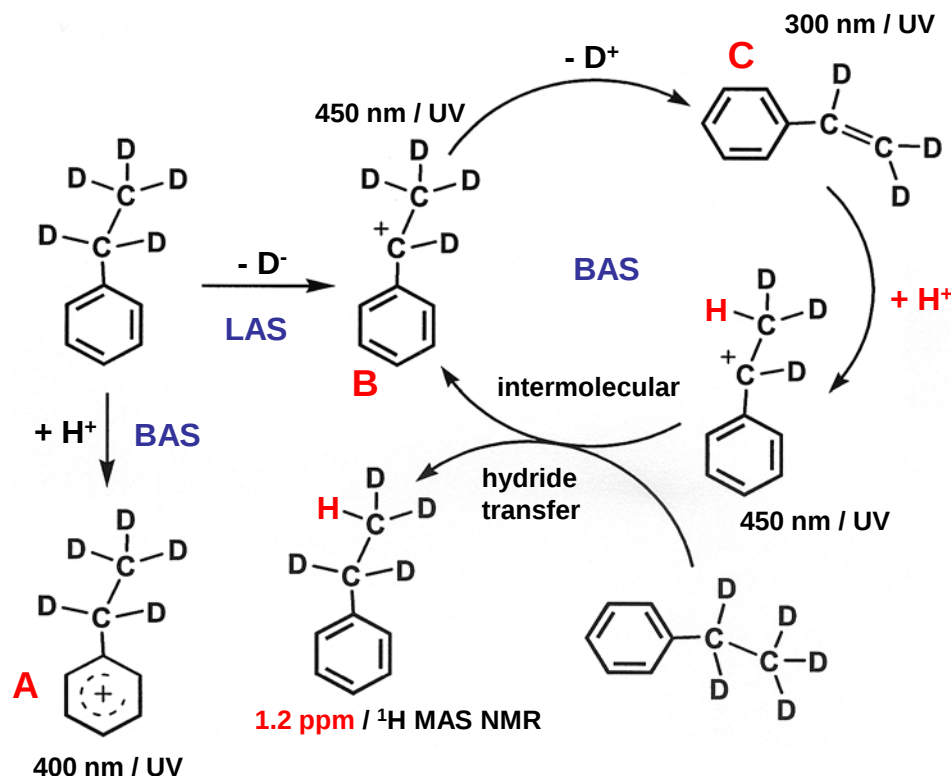
side-chain H/D exchange of ethylbenzene on steamed zeolite deH,Na-Y/81.5

^1H MAS NMR:

- selective H/D exchange of methyl groups (**1.2 ppm**)
- activation energy of **194 kJ/mol** indicates hydride transfer

UV/Vis:

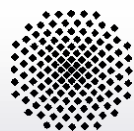
- ethylcyclohexadienyl carbenium ions at **BAS** (400 nm), **A**
- sec-ethylphenyl carbenium ions at **LAS** (450 nm), **B**
- styrene at **BAS** (300 nm), **C**



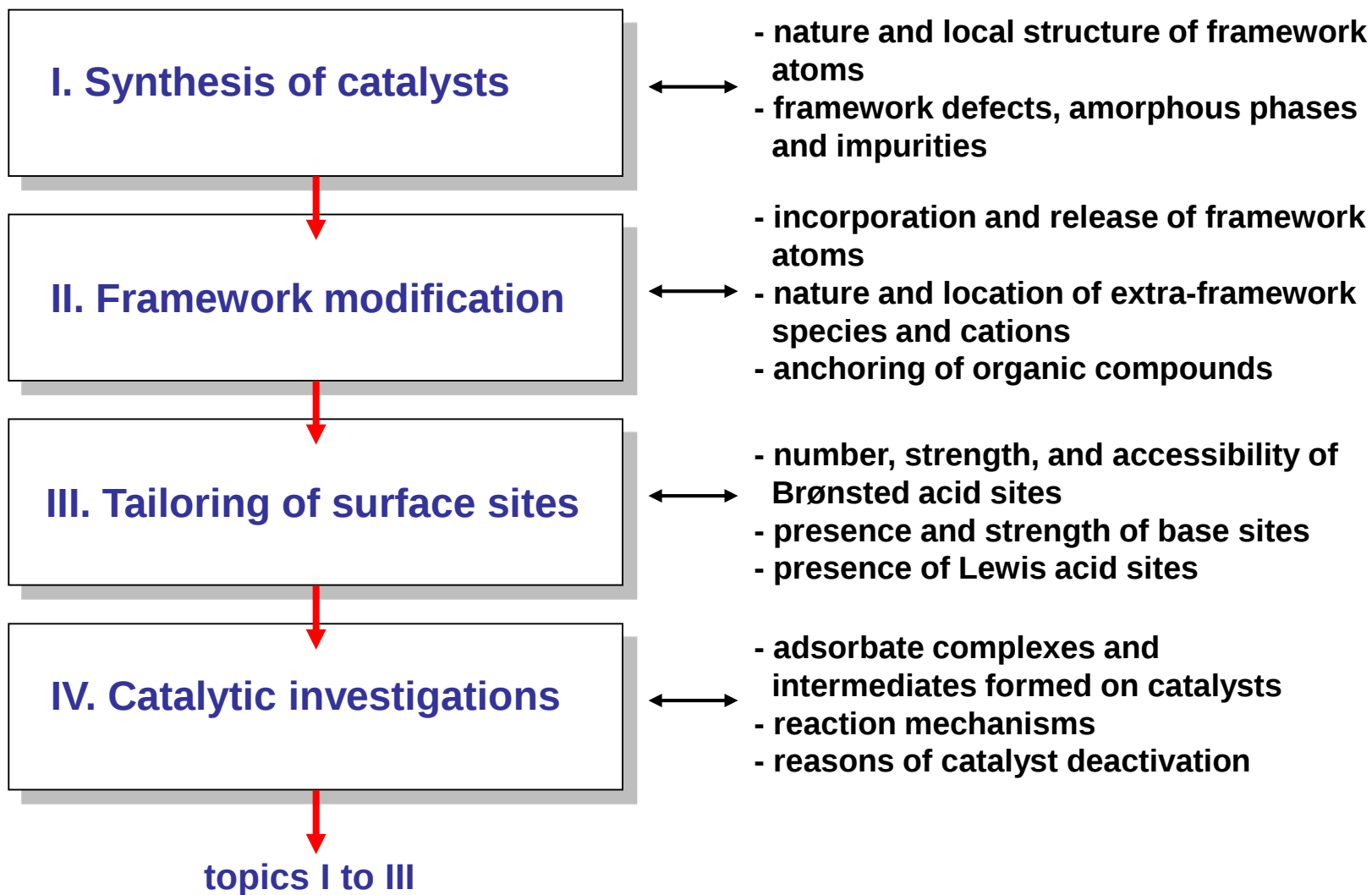
BAS: Broensted acid site

LAS: Lewis acid site





Solid-state NMR of solid catalysts





Thanks to

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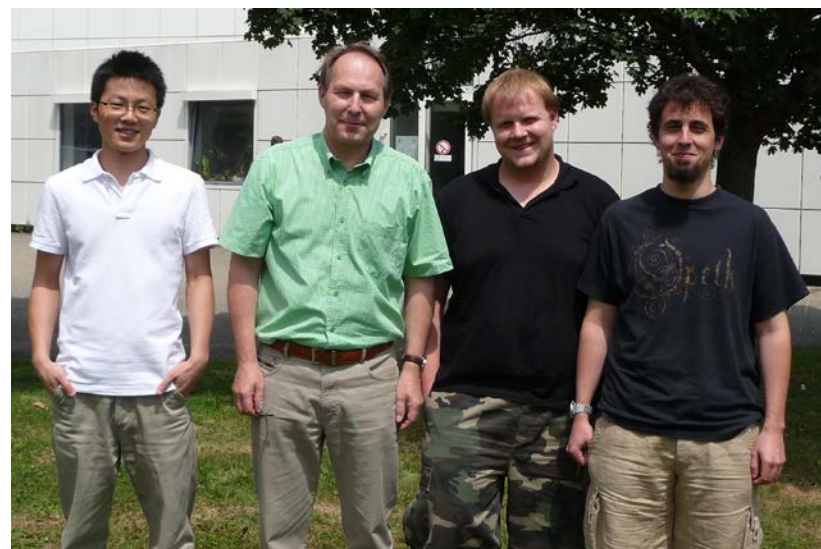
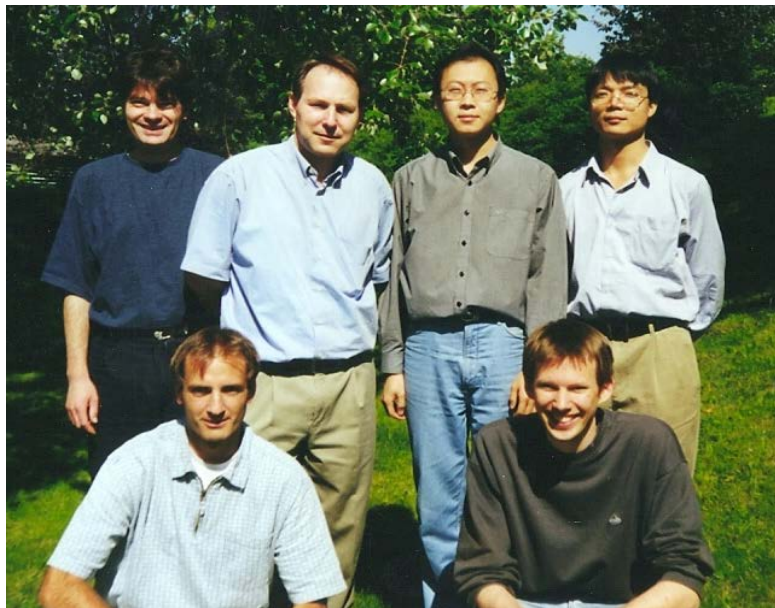
**Alexander von Humboldt-
Foundation**

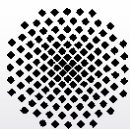
u.a.





Thanks to

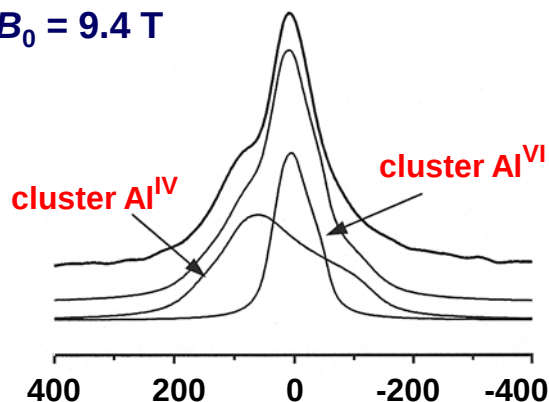




²⁷Al spin-echo NMR studies of reference materials

dehydrated γ -Al₂O₃

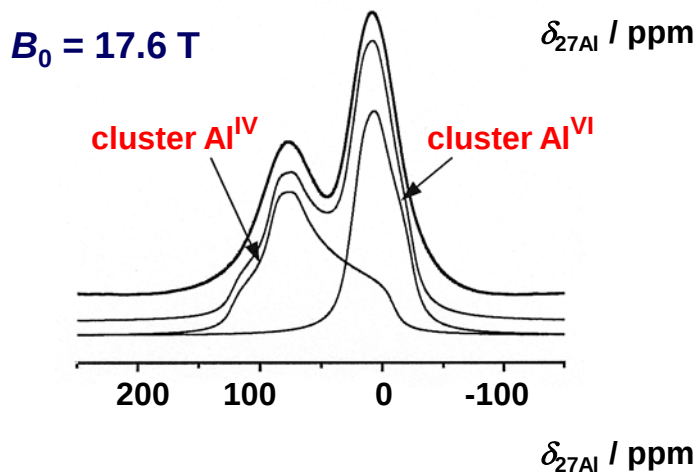
$B_0 = 9.4$ T



- spectroscopic parameters of **cluster Al^{IV}**:

- QCC = 8.5 MHz, $\eta = 0.8$
- $\delta_{\text{iso}} = 68 \pm 5$ ppm
- $I_{\text{rel}} = 60$ %

$B_0 = 17.6$ T



- spectroscopic parameters of **cluster Al^{VI}**:

- QCC = 5.5 MHz, $\eta = 0.7$
- $\delta_{\text{iso}} = 12 \pm 5$ ppm
- $I_{\text{rel}} = 40$ %

