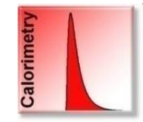




Microcalorimetry beyond Adsorption:

Basics and Applications in Heterogeneous Catalysis



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Lecture Series: Modern Methods in Heterogeneous Catalysis;

10:45 - 12:15



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- 1. Introduction
- 2. Adsorptive microcalorimetric setup
- Tian-Calvet calorimeter
 Evolved adsorption heat
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- Volumetric-Barometric System calibration & measurement of adsorbed amount
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 - Propane & ethane ads. on MoVTeNb oxide, V/SBA15, MoV oxide and P/oCNT at 40°C
 - Propylene adsorption on MoO_x/SBA-15 at T_{reaction}=50°C
 - 1-Hexyne ads. on CeO₂/TiO₂ at T_{reaction}=80°C
 - CO on IrOx at $T_{reaction} = r.t.$
 - CO on Ni/MgAl oxide at 30°C



Introduction



- ➤ adsorption steps, surface reaction processes, and desorption steps
- ➤ 1st step in the catalytic cycle: activation of the reacting molecules by adsorption (strength of chemisorption bond can effect the activation energy)
- adsorption phenomena (bond strength between adsorbate and surface) play an important role in heterogeneous catalysis

For a detailed understanding of complex reaction networks we need:

- thermodynamic data of high accuracy
- information about the nature of the catalyst surface
- →quantitatively study the adsorption, activation, and reaction phenomena close to the reaction parameters
- → simulate reactants-induced responses of the surface
- > since perhaps only a minor fraction of all surface atoms form active centers

Ads. Isothermal Microcalorimetry / Microcalorimetry beyond Ads. direct method to determine number, strength and energy distribution of the adsorption sites

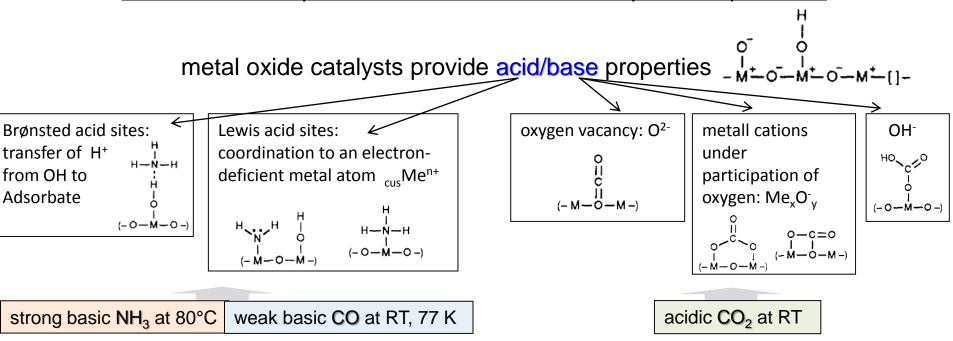
key to the effective use of adsorptive microcalorimetry is the careful choice of probe molecules and adsorption temperature to study



surface sites



careful choice of probe molecules and the adsorption temperature



Focus

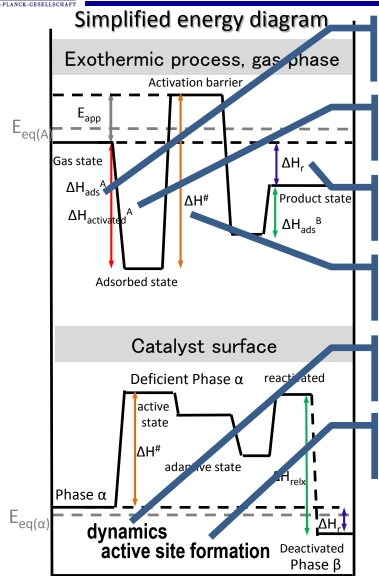
use of probe molecules such as educt, intermediate, product or molecules closely related to the reactants is an elegant method to study the surface sites relevant for catalytic reaction

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T_{adsorption} < T_{reaction} \rightarrow study of the pure ads. process or the activation of the reactant T_{reaction} \rightarrow study of the surface chemical events during the reaction
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Introduction





Adapted van Santen, R. A., Modern Heterogeneous Catalysis, Wiley, 2017
using Schlögl, R., Introduction to Heterogeneous Catalysis, Lecture, FHI Berlin, 2017

CO₂ ads._{chemisorption} on CeO₂ for DEACON reaction ⁵

MoV oxide catalyst in oxidative dehydrogenation of alkanes

Ir-based catalysts for the oxygen evolution reaction at r.t. ¹

Vanadium Oxide-based and Metal-free Catalysts in the ODH of Ethane and Propane ²

Ni based catalysts for the dry reforming of methane ³

Propylene Metathesis over MoOx/SBA-15 ⁴

- V. Pfeifer, T. Jones, S. Wrabetz, C. Massué, J. Velesco-Velez, R. Arrigo, M. Scherzer, S. Piccinin, M. Haevecker, A. Knop-Gericke and R. Schlögl, Chem. Sci., 2016,7, 6791-6795.
- P. Kube, B. Frank, S. Wrabetz, J. Kröhnert, M. Hävecker, J. Valasco-Vélez, J. Noack, R. Schlögl, A. Trunschke, ChemCatChem 9 (2017) 1-14.
- 3 K. Mette, St. Kühl, A. Tarasov, M. G. Willinger, J. Kröhnert, S. Wrabetz, A. Trunschke, M. Scherzer, F. Girgsdies, H. Düdder, K. Kähler, K. Friedel Ortega, M. Muhler, R. Schlögl, M. Behrens, T. Lunkenbein, ACS Catal., 2016, 6 (10), pp 7238–7248.
- 4 Amakawa, K., Wrabetz, S., Kröhnert, J., Tzolova-Müller, G., Schlögl, R., Trunschke, A.; J. Am. Chem. Soc., 134 (28) (2012) 11462-11473.
- Farra, R., Wrabetz, S., Schuster, M. E., Stotz, E., Hamilton, N., Amrute, A. P., Pérez-Ramírez, J., López, N., Teschner, D., Phys. Chem. Chem. Phys., 15 (2013) 3454 3465.



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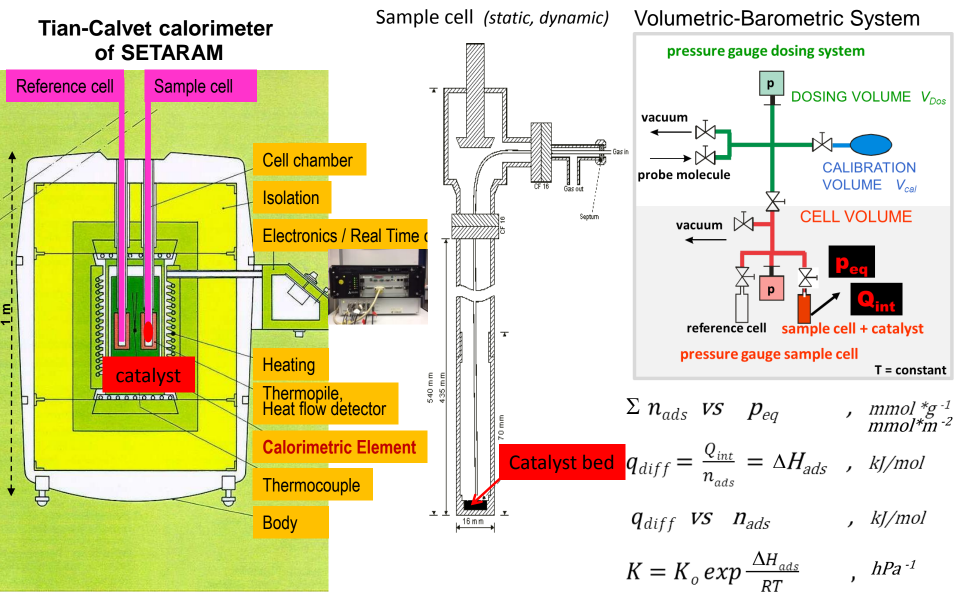


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Adsorptive microcalorimetric setup

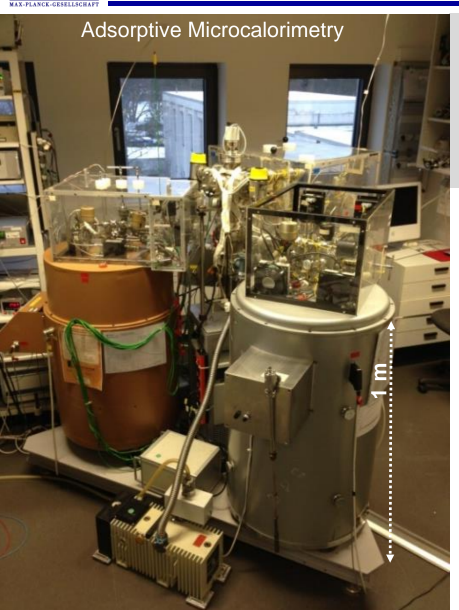






Adsorptive microcalorimetric setup





HT1000 (rt-1000°C), MS 70 (rt-100°C) BT 2.15 (200°C-77K)

Tian-Calvet calorimeter of SETARAM combined with a custom-designed high vacuum and gas dosing apparatus.

The history of modern calorimetry began at the University of Provence in Marseilles.



Tian described his compensation microcalorimeter for the first time in 1922, when he and his colleague COTTE used it to study the metabolism of insects. He brought further improvements to this thermocouple instrument in 1924 and 1926.

Prof. Albert Tian



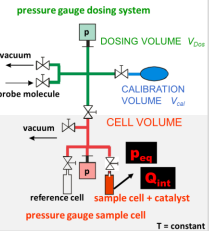
Prof. Edouard Calvet (1895–1966)

CALVET introduced the differential setup (1948) and a rational construction of the **two twinned calorimetric elements**, transforming Tian's apparatus into a true laboratory instrument that today is commercialized by Setaram.

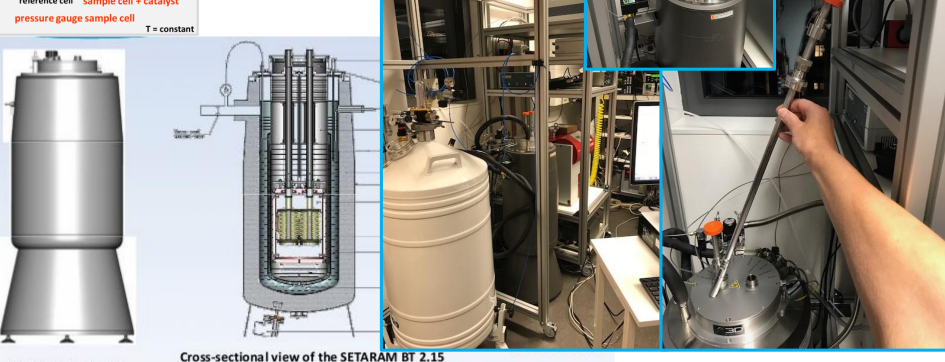


Low Temperature Calorimeter (from -196°C to 200°C) Setaram BT 2.15





Combination of LT-Calorimetry and LT-FTIR
 CO, CO₂, CH₄ ads. on e.g. Cu catalysts,
 Ni-catalysts, MgO



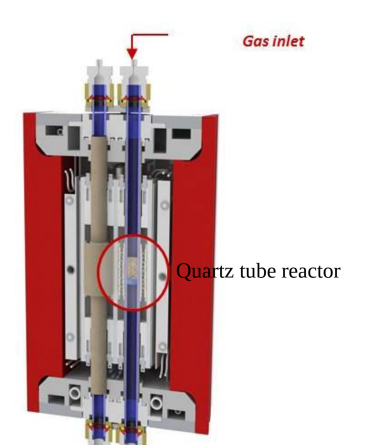


in-situ DSC setup (Differential scanning calorimetry)

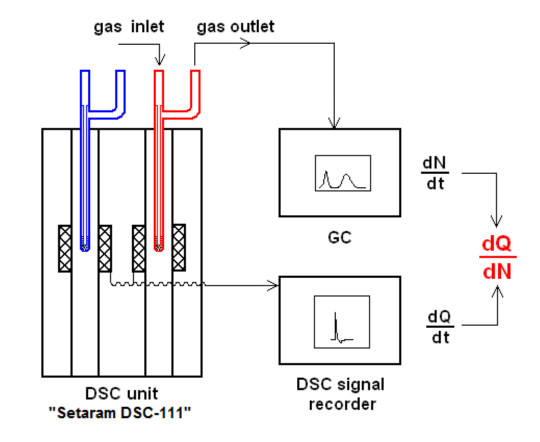
Setaram instrument SENSYS EVO.

Dr. Andrey Tarasov





- Direct measurement of reaction enthalpy
- Precise product analysis



- Calvet calorimetric element
- 3D sensor totally surrounds the catalyst

Gas outlet

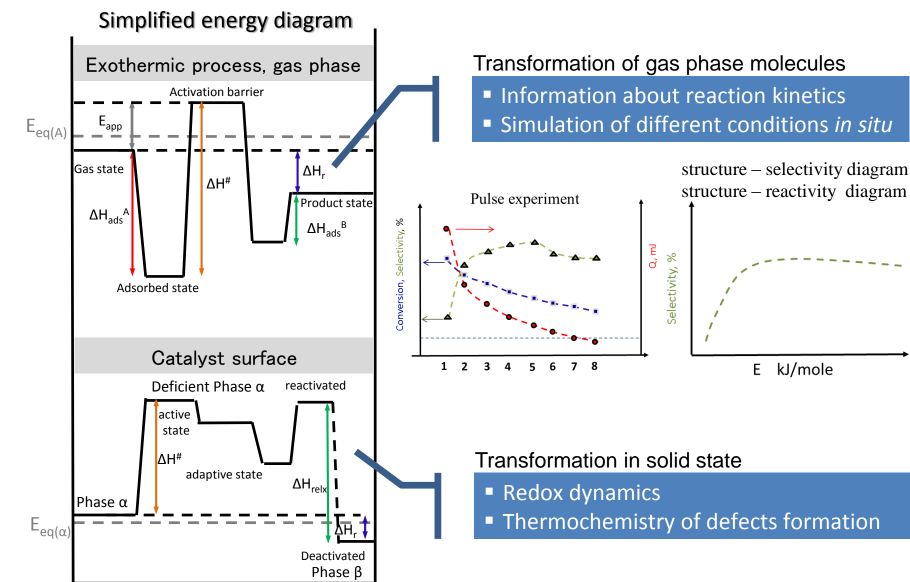


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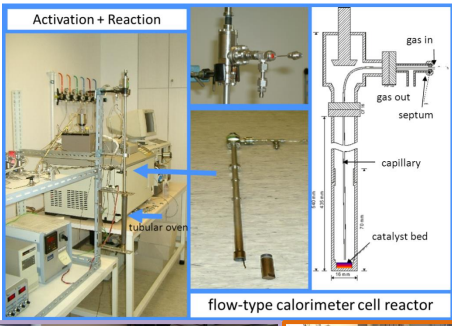


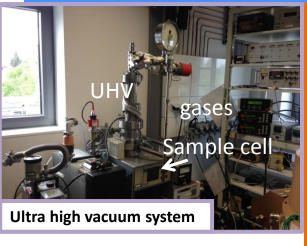




Experimental procedure









Activation:

UHV (10⁻⁸hPa), gases (H₂, O_{2}), rt - 600°C

Reaction:

Calorimeter cell can used as a flowtype reactor.

Catalyst is used in the selected reaction until stady-state performance, rt - 600°C

S. Wrabetz, F.C. Jentoft etal, J. Catal. 269 (2010) 351-358

Transfer

of the sample cell into the calorimeter and degassing/equilibration at T_{ads}

Adsorptive microcalorimetric experiment:

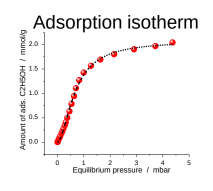
Stepwise adsorption, desorption and re-adsorption of the selected probe molecule at the selected temperature.

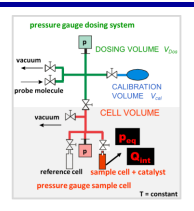


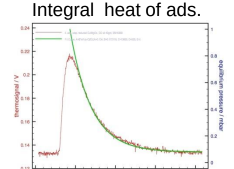
Adsorption Calorimetry - Standard procedure

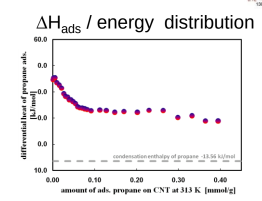


- The probe molecule must be introduced stepwise at constant temperature, the pressure is increased slowly
- For each adsorption step, the adsorbed amount must be determined (isotherm)
- For each adsorption step, the evolved heat must be determined (integral heat of adsorption)
- The differential heat (ΔH_{ads}) can then be determined by dividing the evolved heat through the number of molecules adsorbed in a particular step











Introduction



	Physisorption	Chemisorption
Type of interaction and heat of adsorption (negative enthalpy of adsorption)	van der Waals force $0 < \Delta_p H < 50 \text{ kJ/mol}$ Noble gases, CH_4 , N_2 Dipole-dipole interaction $20 - 25 \text{ kJ/mol}$ H_2O , NH_3	chemical bonding, electron transfer $60 < \Delta_{c}H < 400 \text{ kJ/mol}$ CO on metals (Pt, Pd) Dissociative adsorption
		$(O_2, H_2 \text{ on Pt, } H_2O \text{ on oxides})$
Reversibility	reversible	reversible, irreversible or partially irreversible
Speed	fast	can be slow (e.g. activated adsorption, dissociative adsorption)
Coverage	non-specific and weak	specific (chemisorbed molecule blocks surface sites)
	multilayers possible	monolayer only



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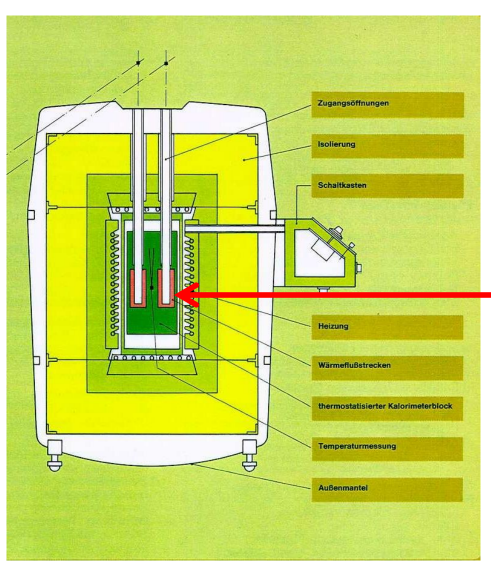


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The Calorimetric Element



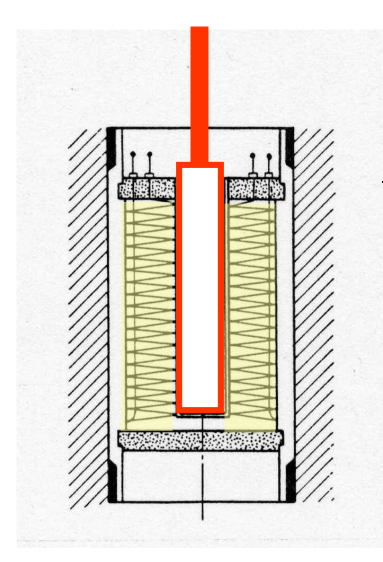


Calorimetric Element



The Calorimetric Element

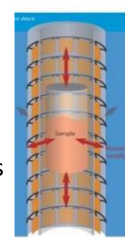




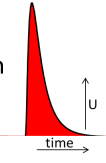
The sample cell is placed into a calorimeter element

3D Calvet heat flow sensor

The cell is totally surrounded by a thermopile made of more than 400 conductive thermocouples in series



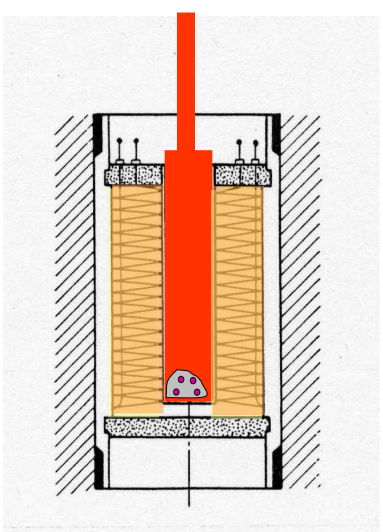
- Thermopile has 2 functions:
 - 1. heat transfer
 - 2. signal generation





Heat and Heat Flow





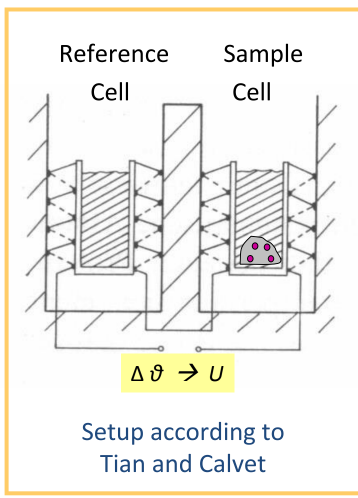
- The **heat** produced by the adsorption/activation/reaction of a dosed **probe molecule** on/with the **catalyst** surface is consumed by 2 processes
 - 1. Increase of the temperature of the sample cell
 - 2. Once there is a temperature gradient between cell and surrounding block, heat flow through the thermopile



Reference cell



ti<u>me</u> 、



The calorimetric block consists of a sample cell and a reference cell.

The reference cell compensate external temperature fluctuations and it provides a good stability of the baseline.

Measurement of the temperature difference $\Delta \vartheta$

The heat-flow detector gives an **electrical signal "U"** which is proportional to the heat transferred per time unit.

N. C. Cardona-Martinez and J.A. Dumesic, Advances in Catalysis 38 150-243.



Power balance, measuring signal and the Tian equation



The power P [W] necessary to heat the cell by $d\theta$ is proportional to the heat capacity C [J/K] of the cell

$$P = C \frac{d\theta}{dt}$$

The heat flow Φ [power] is proportional to the temperature gradient $\Delta\theta$ between cell and block and to the thermal conductivity G [W/K]

$$\Phi = G \left(\theta_{cell} - \theta_{block} \right) = G \Delta \theta$$

Total thermal power of calorimetric element

$$P_{total} = P + \Phi = C \frac{d\theta}{dt} + G \Delta \theta$$

The electrical signal is proportional to the temperature difference; (proportionality factor g=f (number and type of the thermocouple))

$$U = g \Delta \theta$$

The relation between power and electrical signal is then

$$P_{total} = \frac{C}{g} \frac{dU}{dt} + \frac{G}{g} U$$

G [W/K] is constant and if C [J/K] can be considered constant, then C/G is a constant with units of time

$$\tau = \frac{C}{G}$$

The Tian equation shows that the power is not proportional to the temperature difference, the power is delayed with respect to the signal U produced by the cell

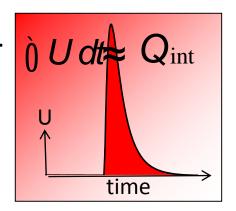
$$P_{total} = \frac{G}{g} \left(U + \tau \frac{dU}{dt} \right)$$



Evolved Heat & Differential Heat of Adsorption



◆ If heat is released in the cell for a limited period of time, e.g. through adsorption, then an electrical signal U with an exponential decrease is obtained.



◆ The integral under the curve is proportional to the evolved heat

$$Q_{\rm int} = \frac{G}{g} \int U \, dt = \int A$$

A: area under curve [Vs]

f: calibration factor [J/(Vs)]

The heat signal of the calorimeter can be calibrated by:

- using an Ohm resistance which produces a certain amount of heat

Q Ohm resistance =
$$U*I*t$$

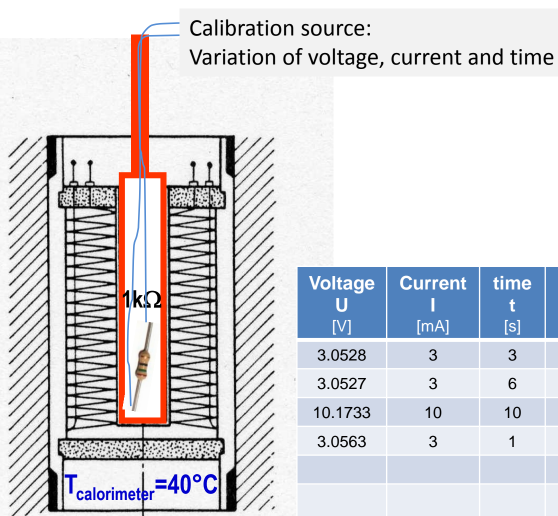
 $f = Q$ Ohm resistance $A_{Ohm resistance}$ [Ws/Vs]

- chemical reaction



Heat signal calibration at 40°C





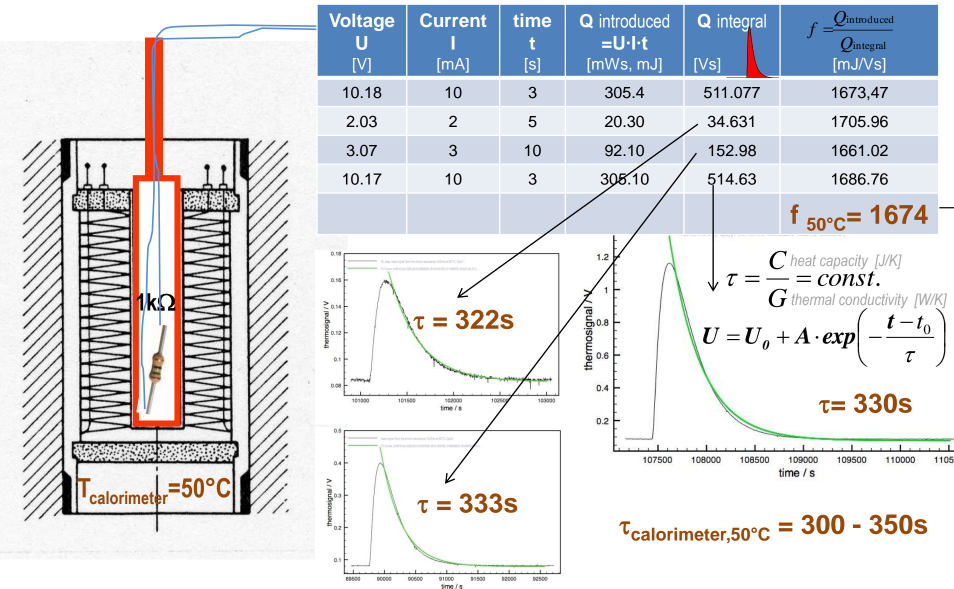


Voltage U [∀]	Current I [mA]	time t [s]	Q introduced =U·I·t [mWs, mJ]	Q integral [Vs]	$f=rac{Q_{ m introduced}}{Q_{ m integral}} \ [ext{mJ/Vs]}$
3.0528	3	3	27.48	45.544	1657.64
3.0527	3	6	54.95	91.143	1658.69
10.1733	10	10	1017.33	1692.075	1663.25
3.0563	3	1	9.16	14.978	1635.30
					•
					f _{40°C} = 1659



Heat signal calibration, time constant τ at 50°C

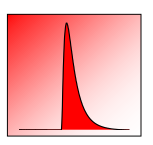






Evolved Heat &Differential Heat of Adsorption





$$Q_{\rm int} = \frac{G}{g} \int U \, dt = \int A$$

A: area under curve [Vs]

f: calibration factor [J/(Vs)]

 Differential heats of adsorption as a function of coverage can be determined:

$$q_{diff} = (\delta Q_{int} / \delta n)_{T,V,p(equ.)}$$

Calculation of the adsorbed amount



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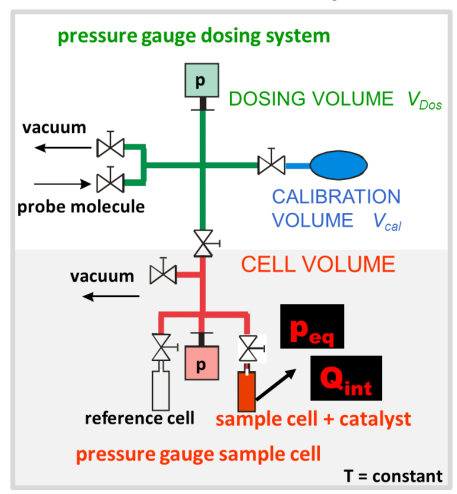
Measurement of the adsorbed amount



- adsorption via stepwise dosing of probe molecule into the sample cell
- Q_{int}, p, T are measured
- $q_{diff} = \Delta H_{cond} \rightarrow saturation of the surface$
- The probe molecule is distributed into three partitions: gas phase, wall adsorption, sample adsorption
- V_{dos} V_{cal} $n_{introduced}$ calibration isotherm $n_{inner\,wall\,+\,gas} = f\left(p_{equ.}\right)$ adsorbed amount

on the surface of the catalyst

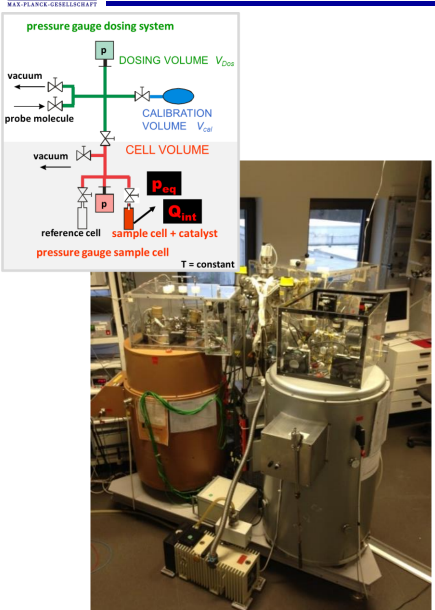
Volumetric-barometric system



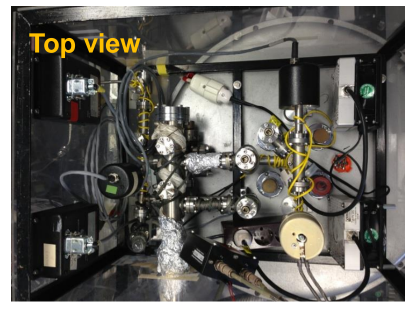


Volumetric-Barometric System







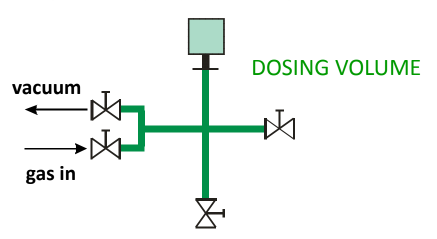




The Dosing Volume V



pressure gauge dosing system



- p, T can be easily measured
- V needs to be determined

Volume Calibration

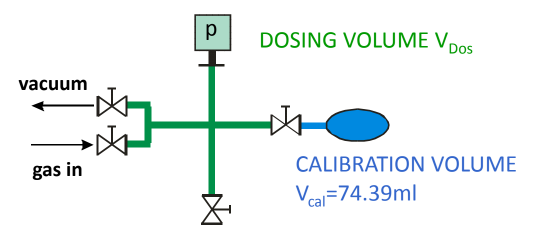
An unknown **volume** of any shape can then be determined through expansion from gas (an ideal gas that does not stick much to the walls) from one volume to the other and pressure measurement before and after the equilibration.

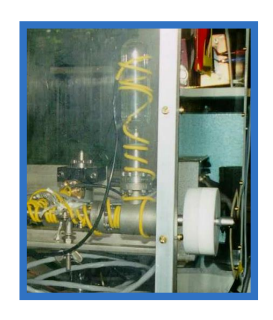


Calibrating the Dosing Volume



pressure gauge dosing system





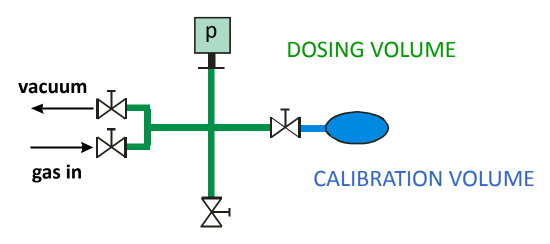
- ❖ fill V_{Cal} and V_{Dos}, same pressure
- close valve between V_{Cal} and V_{Dos}
- ❖ set pressure in V_{dos} to p_{Dos}
- open valve, equilibrate



Calibrating the Dosing Volume



pressure gauge dosing system



Initial situation:

$$nRT = p_{Cal}V_{Cal} + p_{Dos}V_{Dos}$$

❖ After opening valve:

$$nRT = p_{fin} \left(V_{Cal} + V_{Dos} \right)$$

n, T are constant

$$V_{Dos} = \frac{p_{Cal} - p_{fin}}{p_{fin} - p_{Dos}} V_{Cal}$$



Example Data



Propane at 40° C (Project: V_xO_y /SBA15 for ODH of propane)

Nr	p _{Cal} [mbar]	p _{Dos} [mbar]	p _{fin} [mbar]	V(Dos) [ml]
	before	before	after equilibration	
1	5,314	9,467	8,683	137
2	8,683	7,648	7,843	137
3	7,638	6,341	6,585	137
4	6,585	4,768	5,111	137
5	5,111	4,712	4,787	137
6	4,786	3,197	3,496	137
7	3,469	2,701	2,846	136

$$V_{Dos} = \frac{p_{Cal} - p_{fin}}{p_{fin} - p_{Dos}} V_{Cal}$$

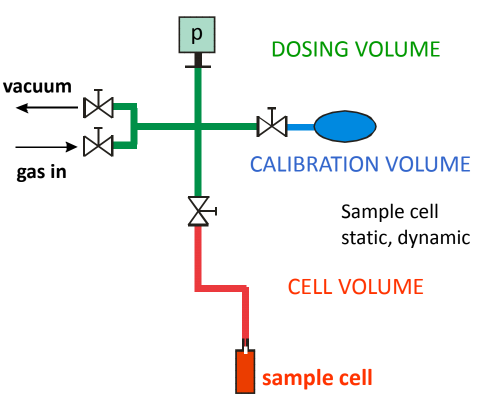
It is important that the entire system is at the same constant temperature!



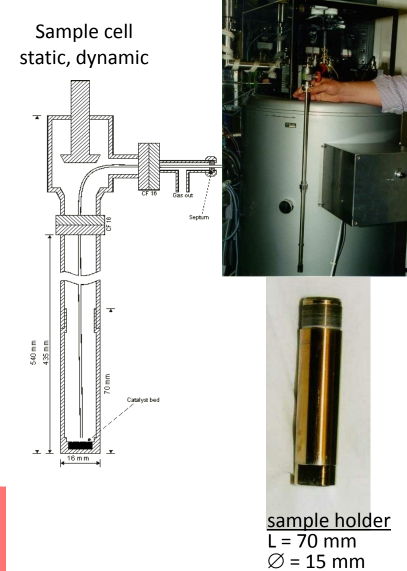
Determining the Dosed Amount $n_{int, i}$







$$n_{\text{int,i}} = \frac{(p_{\textit{Dos,bef}} - p_{\textit{Dos,aft}})V_{\textit{Dos}}}{RT}$$





Total Number of Molecules in Sample Cell



Total number of molecules accumulated in cell

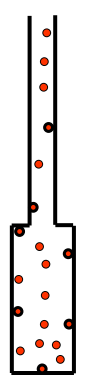
$$n_{SC,tot,i} = \sum_{i} n_{\text{int},i} = n_{SC,tot,i-1} + n_{\text{int},i}$$

i.e. the sum of the number of molecules already in the cell the number of molecules introduced in the ith step

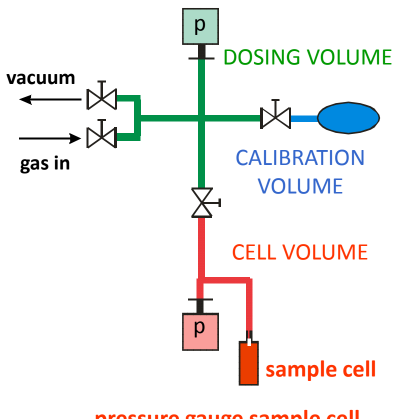


Empty Cell: The Wall Adsorption





- Molecules are in the gas phase but also adsorbed on the inner wall surface
- Only the gas phase molecules contribute to the measured pressure
- The number of molecules adsorbed on the walls depends on the pressure
 - → wall adsorption isotherm



pressure gauge sample cell

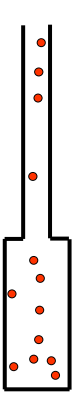


Inner wall Adsorption Blank Measurement



- Measure the pressure in the cell as a function of the total number of molecules introduced into the cell
- Without wall adsorption and without sample, the relation between pressure and number of molecules in the sample cell would be given by the ideal gas law

$$p_{SC,i} = \frac{n_{SC,tot,i}RT}{V_{SC}}$$





Inner wall Adsorption Blank Measurement



❖ With wall adsorption and with or without a sample, the relation between number of molecules in the gas phase + on the walls and the pressure can be written as a polynomial expression

$$n_{SC,w+g,i} = a(p_{SC,i}) - b(p_{SC,i})^2 + c(p_{SC,i})^3 - d(p_{SC,i})^4 \dots$$

Without a sample, the coefficients can be determined $n_{SC,w+g,i} = n_{SC,tot,i}$



Example Blank Measurement



Nr	p _{Dos,bef} /mbar	p _{Dos,aft} /mbar	p _{sc,i} /mbar	n _{int,i} /μmol	n _{SCtot,i} /μmol
1	9,682	9,674	0,007	0,04	0,04
2	9,653	9,641	0,026	0,06	0,10
3	9,607	9,591	0,054	0,08	0,18
4	9,562	9,544	0,081	0,09	0,27
5	9,499	9,481	0,117	0,09	0,36
6	9,442	9,428	0,147	0,07	0,43
7	9,392	9,371	0,181	0,11	0,54
9	9,230	9,195	0,283	0,18	0,72
10	9,128	9,091	0,344	0,19	0,91
11	9,024	8,988	0,403	0,19	1,10
13	8,814	8,758	0,537	0,29	1,39
14	8,644	8,582	0,640	0,32	1,71
15	8,466	8,386	0,755	0,42	2,13
16	8,209	8,122	0,909	0,45	2,58
17	7,783	7,612	1,208	0,90	3,48
19	6,672	6,487	1,870	0,97	3,45
20	5,893	5,526	2,436	1,93	5,38
21	4,477	4,041	3,314	2,30	7,68
22	9,146	8,409	4,056	3,88	11,56

Isobutane at 40°C

without catalyst

(Project:

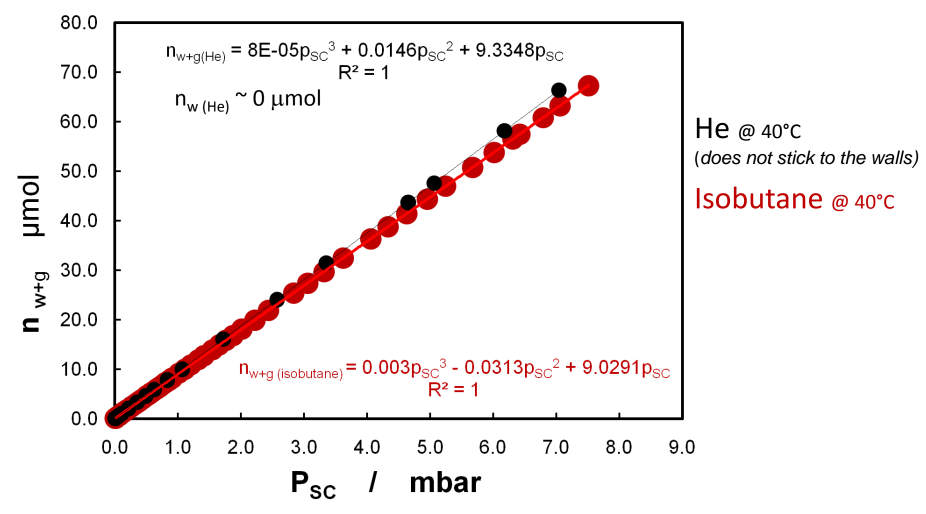
isomerisation of n-butane to isobutane over sulf. ZrO₂)

$$n_{\text{int},i} = \frac{(p_{Dos,bef} - p_{Dos,aft})V_{Dos}}{RT} \qquad n_{SC,tot,i} = \sum_{i} n_{\text{int},i} = n_{SC,tot,i-1} + n_{\text{int},i}$$



Correction for Wall Adsorption



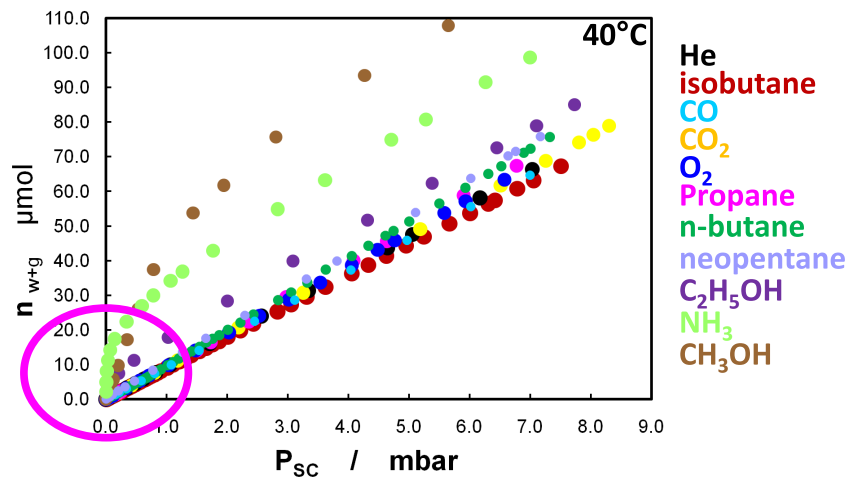


The adsorption of isobutane on the walls is insignificant



Correction for Wall Adsorption



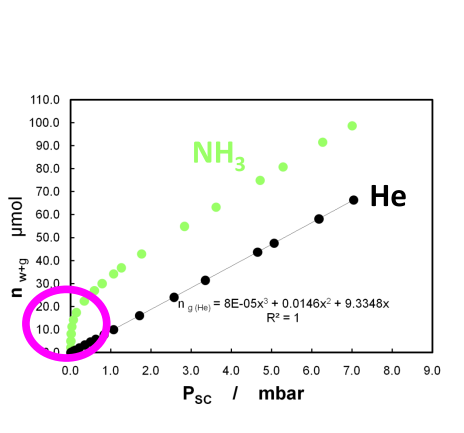


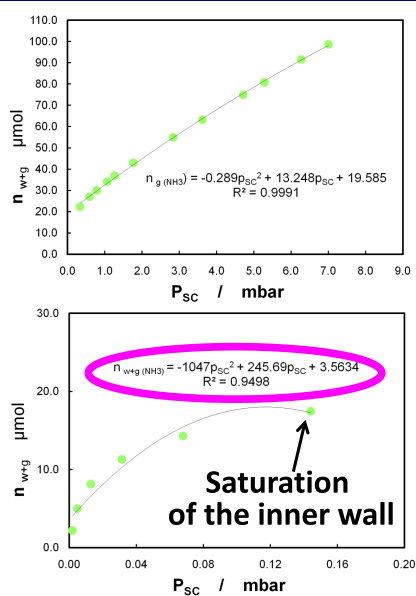
The adsorption of e.g. C₂H₅OH NH₃ CH₃OH on the walls is significant



Correction for Wall Adsorption









Calculation of Adsorbed Amount

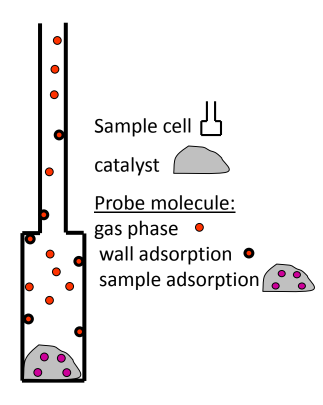


Total number of molecules in sample cell after the ith step

$$n_{SC,tot,i} = n_{ads,tot,i} + n_{SC,w+g,i}$$

Total number of molecules in sample cell after the (i+1)th step

$$n_{SC,tot,i+1} = n_{ads,tot,i} + n_{SC,w+g,i} + n_{ads,i+1} + n_{SC,w+g,i+1}$$



❖ The difference in number of molecules between ith and (i+1)th step is the number of molecules introduced in the (i+1)th step



Calculation of Adsorbed Amount



❖ The number of molecules adsorbed in the (i+1)th step is then

$$n_{ads,i+1} = n_{int,i+1} + n_{SC,w+g,i} - n_{SC,w+g,i+1}$$

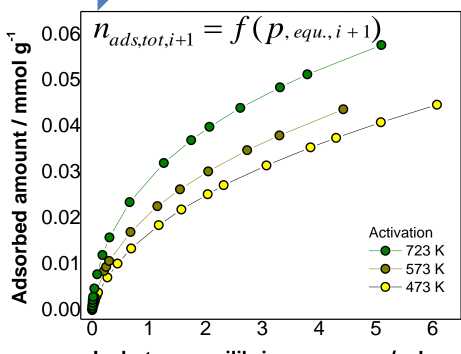
❖ The total number of molecules adsorbed after (i+1) steps is

$$n_{ads,tot,i+1} = n_{ads,tot,i} + n_{ads,i+1}$$
 Adsorption isotherm

Isobutane adsorption on different activated sulf. ZrO₂ at 40°C.

(Project: isomerisation of n-butane to isobutane over sulf. ZrO₂

[S. Wrabetz, F.C. Jentoft etal, J. Catal. 269 (2010) 351-358.])



Isobutane equilibrium pressure / mbar



Contents



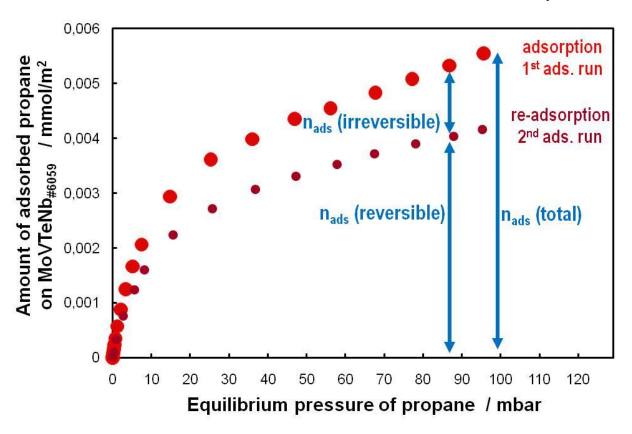
- 1. Introduction
- 2. Adsorptive microcalorimetric setup
- Power balance of Tian-Calvet calorimeter & Evolved adsorption heat
 Differential heats of adsorption
- 4. Volumetric-Barometric System calibration & measurement of adsorbed amount
- Obtained physical quantities & evaluation criteria of the calorimetric results
- 6. Applications of microcalorimetry in heterogeneous catalysis
 - H₂ and CO adsorption on 2%Pt / Al₂O₃ at 40°C
 - NH₃ adsorption on pure-phase MoVTeNb oxide at 80°C
 - CO₂ ads. on CeO₂ at 40°C
 - O₂ adsorption on supported CeO₂ at 200°C
 - Ethanol adsorption on VO_x/γ-Al₂O₃ at 40°C
 - Propane ads. On MoVTeNb oxide at 40°C
 - Propylene adsorption on MoO_x/SBA-15 at T_{reaction}=50°C
 - 1-Hexyne ads. on CeO₂/TiO₂ at Treaction=80°C



Obtained physical quantities & evaluation criteria of the calorimetric result







 n_{ads} (total) : overall adsorbed amount under an equilibrium pressure of 95 mbar n_{ads} (irrev.) : chemisorbed amount



Analysis of the Adsorption Isotherm



Adsorption Isotherm

of propane on 10%V/SBA15 catalyst active in oxidation of propane amount of adsorbed propane [mmol / g] 0,14 **Analysis** 0,12 0,10 0,08 activation 0,06 0,04 $\Delta_a H (q_{diff}) = 40kJ/mol = const$ 0,02 10 20 0 30 50 60

Freundlich isotherm:

The enthalpy of adsorption $\Delta_a H$ per site $N_{ads} = \frac{A p^n}{\frac{1}{1 + A p^n}}$ decreases exponential with coverage Θ decreases exponential with coverage Θ

$$N_{ads} = \frac{A p^{\frac{1}{n}}}{1 + A p^{\frac{1}{n}}}$$

Tempkin isotherm: The enthalpy of adsorption $\Delta_a H$ per site decreases linear with coverage Θ

equilibrium pressure [mbar]

$$heta = rac{RT}{q_{\mathsf{ads}}^{(heta=0)} - q_{\mathsf{ads}}^{(heta=1)}} \mathsf{In} \left(rac{1 + rac{p}{p^{st}} \mathsf{exp} rac{q_{\mathsf{ads}}^{(heta=0)}}{RT}}{1 + rac{p}{p^{st}} \mathsf{exp} rac{q_{\mathsf{ads}}^{(heta=1)}}{RT}}
ight)$$

Langmuir model: The enthalpy of adsorption $\Delta_{a}H$ per site is constant with coverage Θ

Langmuir model for dissociative adsorption

$$\theta_H = \frac{n_{ads}}{n_m} = \frac{\sqrt{p_{H_2}K_{ads}}}{1+\sqrt{p_{H_2}K_{ads}}} \quad \begin{array}{l} \text{e.g., dissociative ads. of} \\ \text{H}_2 \text{ on Pt supported on} \\ \text{Al}_2\text{O}_3 \text{ at 40°C} \end{array}$$

Higher order Langmuir model

$$N_{ads} = \frac{N_{mono}(Kp)^{\frac{1}{n}}}{1 + (Kp)^{\frac{1}{n}}}$$
 e.g., activated ads. of n-butane (educt) on sulf. ZrO₂ at 40°C < $T_{reaction} = 60$ °C [S. Wrabetz et.al., Catal. 2010, 269, 351 – 358.]

ightharpoonup Langmuir model $\theta = \frac{Kp}{1 + Kn}$

e.g., propane adsorption (313 K) on the oxygen surface groups of CNT and B2O3-CNT catalysts used in ODH of **propane** [B. Frank, S. Wrabetz et.al., ChemPhysChem 2011, 12, 2709 – 2713]

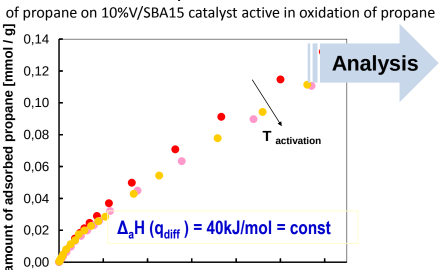
CO Adsorption on supported Gold nanoparticle catalysts [Ch. J. Pursell et.al., J. Phys. Chem.C, 2012, 116(20),11117]

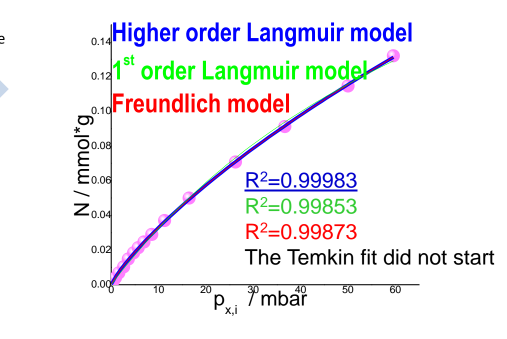


Analysis of the Adsorption Isotherm



Adsorption Isotherm





$$Specific \ surface \ area = N_{mono} \ * \ Avogadro \ constant \ * \ S_{1:1, \ assumed} \ * \ cross-section \ area_{39 \text{\AA}2 \ for \ propane} \ [\text{A. L. McClellan et.al., J. of Colloid and Interface Science, 23 (1967) 577}]$$

$$N_{ads} = \frac{N_{mono} (Kp)^{\frac{1}{n}}}{1 + (Kp)^{\frac{1}{n}}}$$

10

0

20

30

equilibrium pressure [mbar]

n = 1 non-activated ads.n > 1 activated ads.

10% V/SBA15 dehydration temperature	N_{mono} $\mu mol *g^{-1}$	n	R ²	Spropane m ^{2*} g ⁻¹	BET S _{N2} m ^{2*} g ⁻¹
373 K	0.9 (2)	1.20 (2)	0.99983	226 (10)	329 (4)
573 K	1.3 (4)	1.22 (2)	0.99982	304 (10)	
673 K	1.2 (3)	1.22 (2)	0.99905	290 (10)	

60

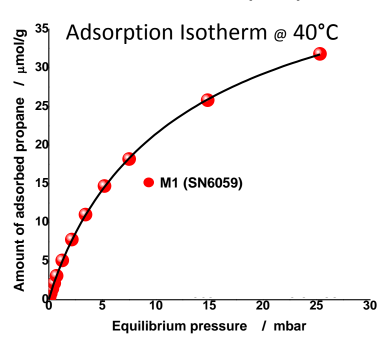
50

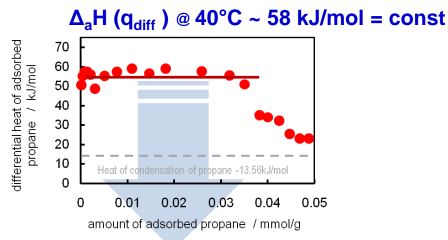


Analysis of the Adsorption Isotherm



Propane adsorption on pure-phase M1 catalyst (MoVTeNb oxide #6059) active in selective oxidation of propane





$$N_{ads} = \frac{N_{mono} (Kp)^{\frac{1}{n}}}{1 + (Kp)^{\frac{1}{n}}}$$

Langmuir model for activated ads. : The enthalpy of adsorption $\Delta_a H$ per site is constant with coverage Θ

Specific surface area = N_{mono} * Avogadro constant * $S_{1:1, assumed}$ * cross-section area $_{39\text{Å}2 \text{ for propane}}$ [A. L. McClellan et.al., J. of Colloid and Interface Science, 23 (1967) 577]

Catalyst (ID)	$N_{monolayer} \ \mu mol \ ^*g^{-1}$	\mathbb{R}^2	$S_{propane} m^2 g^{-1}$ $A_{propane} \sim 39 \text{ Å}^2$	BET m ² *g ⁻¹ A _{N2} =16.2 Å ²
M1 (5630)	34.9 (2)	0.99951	8.2 (1)	6.6
M1 (6059)	48.5 (8)	0.99991	11.3 (1)	8.8
M1 (5737)	42.3 (8)	0.99986	9.9 (1)	13.4

N_{ads} - coverage with certain equilibrium pressure

 N_{mono} - monolayer coverage

p - equilibrium pressure

- equilibrium pressure - adsorption order

VA - adsorption equilibrium constant

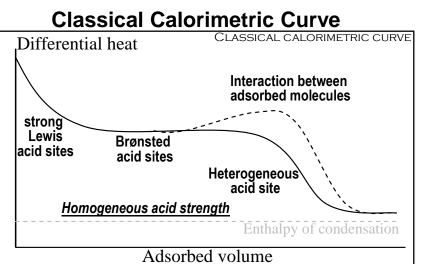
- correlation coefficient; goodness of fit

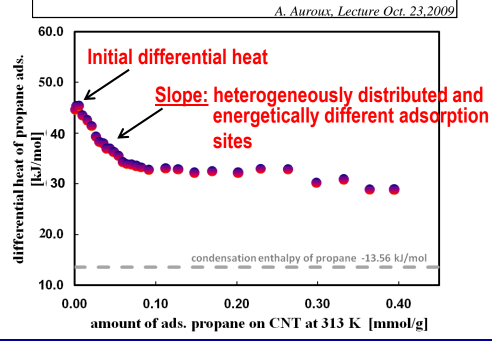
S - stoichiometry

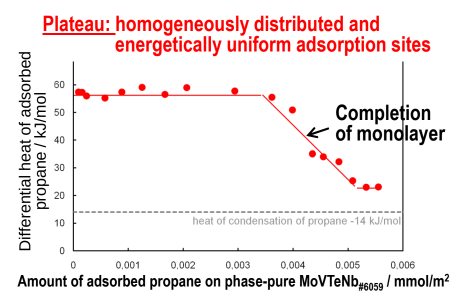


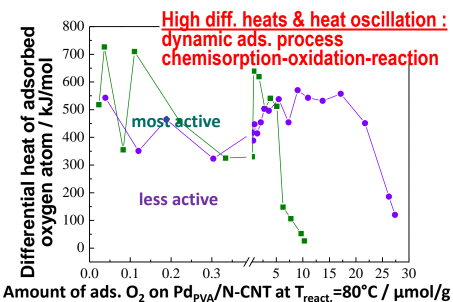
Obtained physical quantities & evaluation criteria of the calorimetric result











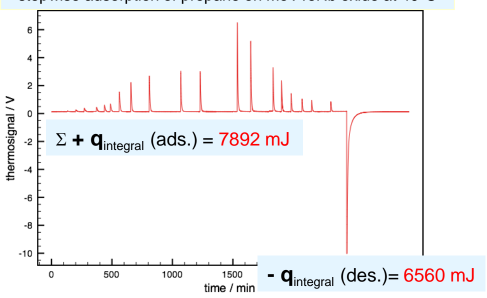


Evaluation criteria of the calorimetric result



Integral heat signal of adsorption and desorption

stepwise adsorption of propane on MoVTeNb oxide at 40°C



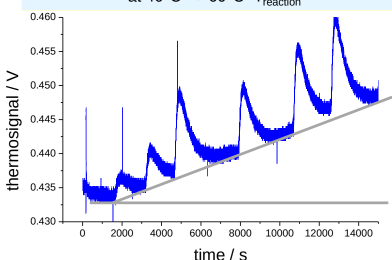
 Σq_{int} (ads.) > q_{int} (des.) \rightarrow partially irreversible ads.; activated ads. process

 Σq_{int} (ads.) < q_{int} (des.) \rightarrow instability of the catalyst in the presence of probe molecule

 Σq_{int} (ads.) = q_{int} (des.) \rightarrow reversible ads. process

Background of the thermo signal during the stepwise adsorption

stepwise n-butane ads. on sulf. ZrO_2 at $40^{\circ}C < 60^{\circ}C = T_{reaction}$



Heat signal deviates from the base-line

- → Adsorption process is accompanied by secondary processes
 - e.g. during n-butane ads. a partial isomerization of n-butane to isobutane in the calorimeter cell was observed

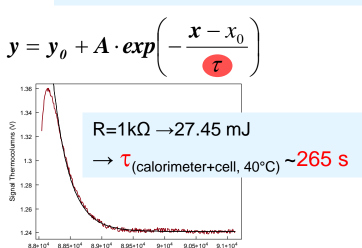


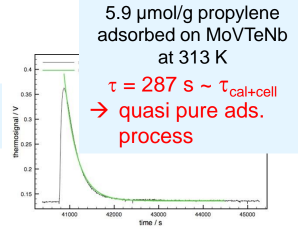
Evaluation criteria of the calorimetric result

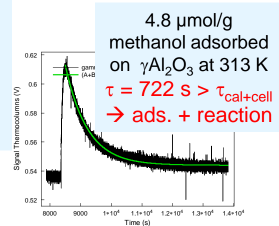


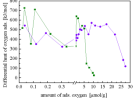
Determination of the time constant τ of the integral heat signal

heat capacity C thermal conductivity G

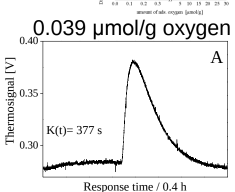


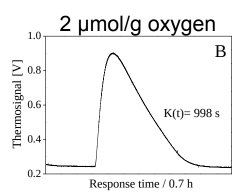


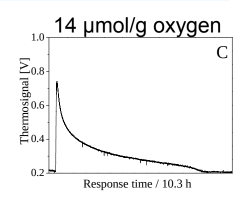




Shape of the integral heat signalO₂ adsorption on 2%Pd/N-CNT473K at 353K=T_{reaction}







At low coverage quasi pure dissociative oxygen chemisorption on Pd

At higher coverage oxygen chemisorption is accompanied by secondary processes





Specific surface area of Pt: H₂ and CO ads. on 2%Pt/Al₂O₃ at 40°C

- NH₃ ads. on pure-phase MoVTeNb oxide at 80°C Titration of acid or basic surface sites:

> - NH₃ ads. on H-ZSM5 at 80°C; Validation of the TPsurface chemical probe reaction of n-propylamin

- CO₂ ads. on CeO₂ at 40°C

Investigation of the oxidation process: - O₂ ads. on supported CeO₂ at 200°C

Study of catalytic relevant sites via calorimetry close to the reaction conditions:

- Ethanol ads. on VO_x/γ -Al₂O₃ at 40°C

- Propane ads. on MoVTeNb oxide at 40°C

- Propane and EB adsorption on the oCNT

Study of catalytic relevant sites via calorimetry under reaction conditions:

- Propylene chemisorption on MoO_x/SBA-15 at T_{reaction}=50°C

- O₂ ads. on Ag for ethylene epoxidation at T_{reaction}=230 °C - 1-Hexyne chemisorption on CeO₂/TiO₂ at T_{reaction}=80°C

- CO chemisorption on Ir based catalysts for OER at r.t.

Reactants induced dynamic responses of catalyst surface:

- CO chemisorption cycles (30°C) on Ni/MgAI oxide catalyst for dry reforming of methane (DRM)

- propane and ethane ads./des. cycles on MoV oxide at 40°C

Estimation of the enthalpy of formation of the transition state (activation barrier):

- ethane & ethylene and propane & propylene ads. on the Vanadium oxide-based and Metal-free catalysts for ODH



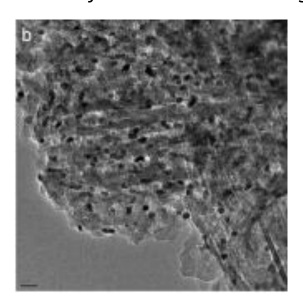
VALIDATION

H₂ chemisorption on 2 wt.% Pt_{H2-673K-2h}/Al₂O₃ at 40°C

Reference Material from



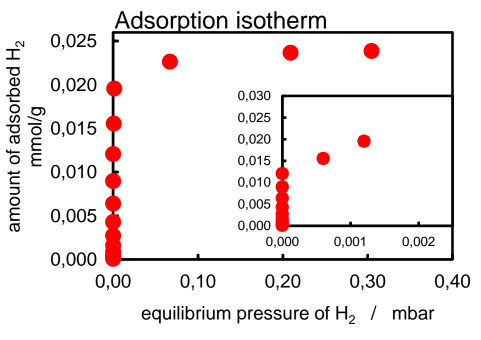
Active Metal Surface Area = $1.146 \text{ m}^2 \text{ g}^{-1}$

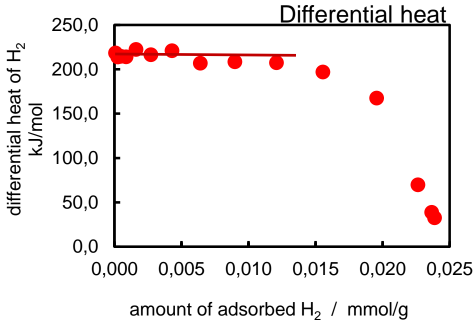




H₂ adsorption on Pt_{H2-673K-2h}/Al₂O_{3 Quantachrome} at 40°C







Heat plateau:

homogeneously distributed and **energetically uniform** adsorption sites

 \underline{q}_{diff} = 215 kJ/mol : chemisorption ! $\Delta_{C}H$ = 215 kJ/mol Hydrogen is dissociative adsorbed on platinum $H_2 + Pt + Pt \rightarrow Pt-H + Pt-H \Delta_C H = 266 \text{ kJ/mol}$ 2.5% Pt/silica J.A. Dumesic et.al., Catal. Letters 45 (1997) 155-163.

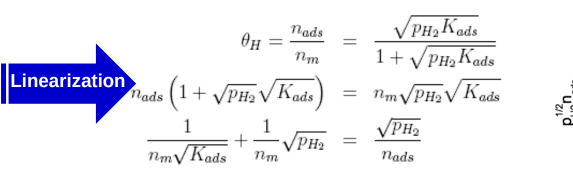
Langmuir isotherm for dissociative adsorption

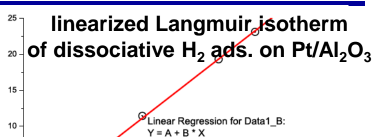
$$\theta_H = \frac{n_{ads}}{n_m} = \frac{\sqrt{p_{H_2} K_{ads}}}{1 + \sqrt{p_{H_2} K_{ads}}}$$



H₂ adsorption on Pt/Al₂O_{3 Quantachrome} at 40°C







Value Error

0.15128 0.48555

		• H2				
registered $\frac{p_{H_2}}{mbar}$	10^{-6}	$6 \cdot 10^{-4}$	$1.2\cdot 10^{-3}$	$6.69 \cdot 10^{-2}$	$2.09 \cdot 10^{-1}$	$3.05 \cdot 10^{-1}$
calculated from the calibrated volumetric system $\frac{n_{ads}}{(mmol \cdot g^{-1})}$	$1.208 \cdot 10^{-2}$	$1.555 \cdot 10^{-2}$	$1.956 \cdot 10^{-2}$	$2.264 \cdot 10^{-2}$	$2.366 \cdot 10^{-2}$	$2.387 \cdot 10^{-2}$
calculated $rac{\sqrt{p_{H_2}}}{\sqrt{mbar}}$	10^{-3}	$24.5 \cdot 10^{-3}$	$34.6\cdot 10^{-3}$	$259 \cdot 10^{-3}$	$457\cdot 10^{-3}$	$552\cdot 10^{-3}$
calculated $\frac{\sqrt{{}^{p}H_{2}}}{\sqrt{{}^{n}ads}}$	0.083	1.58	1.77	11.44	19.32	23.13

$$\frac{1}{n_m} = 42 \frac{g}{mmol} \Rightarrow n_m = 24 \cdot 10^{-3} \frac{mmol}{g}$$
 Specific surface area of Pt m^2_{Pl}/g_{cat}

$$\frac{1}{n_m \sqrt{K_{ads}}} = 0.4 \frac{\sqrt{mbar} \cdot g}{mmol} \implies K_{ads} = \left(\frac{1 \cdot g \cdot mmol}{24 \cdot 10^{-3} \ mmol \cdot 0.4 \sqrt{mbar} \cdot g}\right)^2$$

$$\Rightarrow K_{ads} = 10851 \ mbar^{-1}$$



H₂ adsorption on Pt/Al₂O_{3 Ouantachrome} at 40°C



Specific surface area of Pt m^2_{Pt}/g_{cat}

$$S_{Pt} = \frac{n_m \cdot \text{Avogadro constant}}{\text{Surface sites density } \Gamma_{\text{fcc lattice, Pt}}}$$

$$S_{Pt} = \frac{24 \cdot 10^{-6} \text{ mol } \cdot 6.022 \cdot 10^{23} \text{ particles} \cdot \text{cm}^2}{\text{g} \cdot 1.5 \cdot 10^{15} \text{ atoms} \cdot \text{mol}}$$

$$S_{Pt} = 9667 \text{ cm}^2/\text{g} \approx 1 \text{ m}^2/\text{g}}$$

$$S_{Pt; \text{certified by Quantachrome}} = 1.146 \text{ m}^2/\text{g}}$$

Final Results

Kinetic parameters determined for the H₂ ads. on Pt/Al₂O₃

$$n_{mono} = 24 * 10^{-6} mol \cdot g^{-1}$$
 $K_{ads} = 10851 mbar^{-1}$
 $\Delta H_{ads.} = 215 \pm 4 kJ \cdot mol^{-1}$
 $S_{Pt} \approx 1.0 m^2 \cdot g^{-1}$

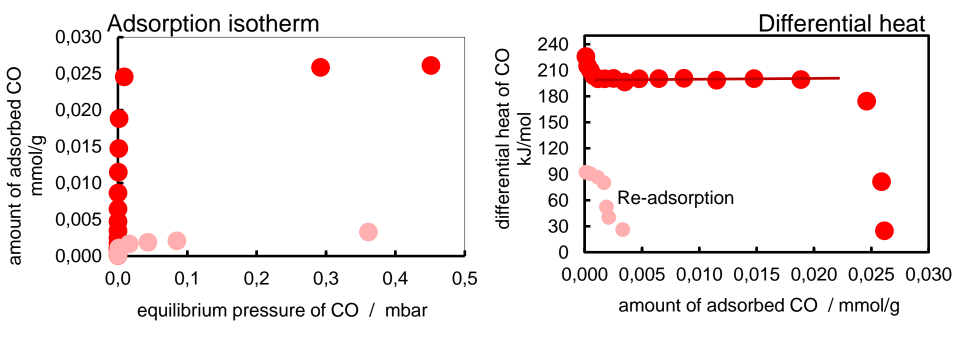


CO adsorption on Pt_{H2-673K-2h}/Al₂O_{3 Quantachrome} at 40°C



CO adsorption on Pt_{H2-673K-2h}/Al₂O_{3 Quantachrome} at 40°C





Heat plateau:

homogeneously distributed and **energetically uniform** adsorption sites

Reversibility:

CO is mainly irreversible adsorbed on Pt at 40°C

 \underline{q}_{diff} = 200 kJ/mol : chemisorption ! $\Delta_{C}H$ = 200 kJ/mol

Pt-CO

J. Therm. Anal. Cal., 82, 2005 105

terminal CO $\Delta_{\rm C}$ H= 209 kJ/mol $\nu_{\rm CO}$ ~ 2055 cm⁻¹ $(Pt)_{n; n>1}$ -CO bridged CO $\Delta_CH=94$ kJ/mol $\nu_{CO}\sim1822$ cm⁻¹ higher order Langmuir model for activated adsorption

$$V_{ads} = \frac{N_{mono}(Kp)^{\frac{1}{n}}}{1 + (Kp)^{\frac{1}{n}}}$$



CO adsorption on Pt_{H2-673K-2h}/Al₂O_{3 Quantachrome} at 40°C





$$N_{ads} = \frac{N_{mono} (Kp)^{\frac{1}{n}}}{1 + (Kp)^{\frac{1}{n}}}$$

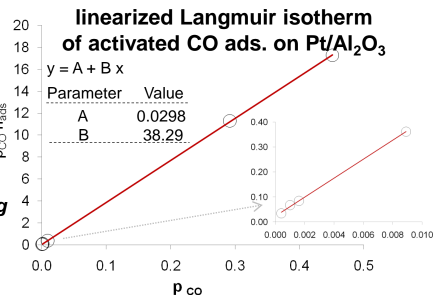
$$1/(n_{mono}K_{ads}^{1/n}) + (1/n_{mono}) p_{CO} = p_{CO} / n_{ads}$$

$$1/n_{\text{mono}} = 38.29 \ g/mmol \rightarrow n_{\text{mono}} = 26.11 \ \text{x} 10^{-3} \ mmol/g$$

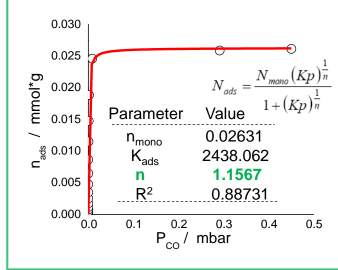
 $1/(n_{\text{mono}}K_{\text{ads}}^{1/n}) = 0.0298 \text{ mbar} \cdot g/\text{mmol} \rightarrow K_{\text{ads}} = (1/n_{\text{mono}}A)^{n}$ $K_{\text{ads}} = = (1 \text{ g mmol} / 26.11x10^{-3} \text{ mmol} 0.0298 \text{ mbar g})^{1.1567}$

$$K_{ads} = 3945,96 \text{ mbar}^{-1}$$

$$S_{Pt} = 10563 \text{ cm}^2/\text{g} \approx 1.1 \text{ m}^2/\text{g}$$
;









Comparison



Kinetic parameters determined for the CO ads. on Pt/Al₂O₃

```
n_{mono} = 26 * 10^{-6} mol \cdot g^{-1}
K_{ads} = 3946 mbar^{-1}
\Delta H_{ads.} = 200 \pm 4 kJ \cdot mol^{-1}
S_{Pt} \approx 1.1 m^2 \cdot g^{-1}
```

Kinetic parameters determined for the H_2 ads. on Pt/Al_2O_3

Validation

of the SETARAM calorimeter combined with a custom-designed high vacuum and gas dosing apparatus.

```
J. Therm. Anal. Cal., 82 (2005)105 : Pt-CO \Delta_CH = 209 \text{ kJ/mol} \Delta_CH = 200 \pm 4 \text{ kJ/mol} (Dept. AC FHI) S_{Pt}; certified by Quantachrome = 1.146 m²/g
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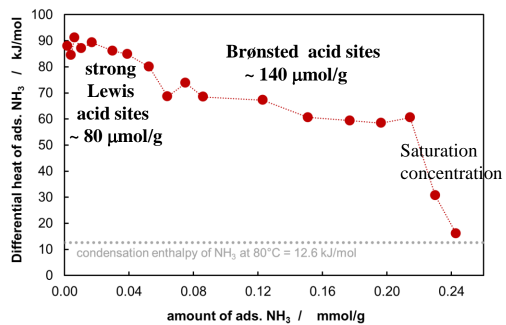


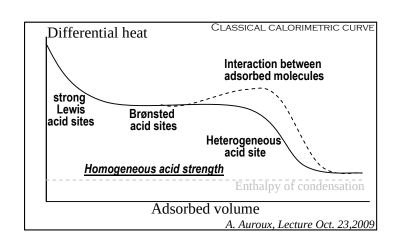
Titration of acid sites on the surface of MoVTeNb oxide catalyst #6059 using NH₃ at 80°C



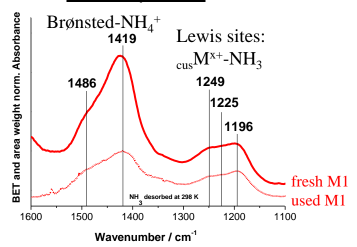
Titration of acid sites on the surface of M1 #6059 using NH₃ at 80°C







FTIR spectra



Determination of the amount of Brønsted and Lewis acid sites using the extinction coefficients from the literature:

Brønsted acid sites: ~ 40 µmol/g Lewis acids sites: ~ 90 µmol/g





Development of a NEW METODE:

Quantification of Brønsted Acid Sites on Catalyst Surfaces via Temperature Programmed Surface Reaction of n-Propylamine

VALIDATION via Calorimetry NH₃ ads. on H-ZSM5 zeolite at 80°C



NEW METODE of Quantification of Brønsted acid sites on catalyst surfaces by temperature programmed surface reaction of n-Propylamin. VALIDATION via Calorimetry & FTIR

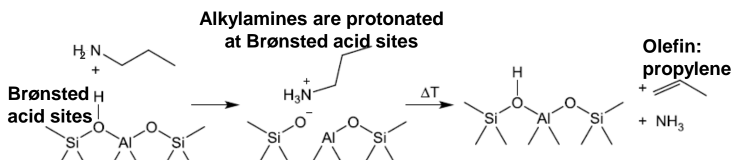


Motivation:

Development of a standard characterization method for the quantification of acid sites on surfaces.

Gorte et al. developed a method that uses alkylamines as reactive probe molecules for TPSR experiments on zeolites.

It was investigated if the needed information can be readily obtained by a commercial TPD machine. - BasCat



Combination of TPSR and TPD/MS: quantify **Brønsted** acid sites is the

amount of desorbed olefin

Samples:

- H-ZSM5-30 (H-ZSM5 zeolite with a SiO₂/Al₂O₃ ratio of 30)
- H-ZSM5-80 (H-ZSM5 zeolite with a SiO₂/Al₂O₃ ratio of 80)

Zeolite



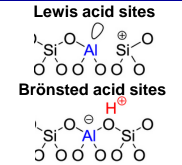
Microcalorimetry <u>Differential heat of ads. NH₃ at 80°C</u>



H-ZSM5-30

SiO₂/Al₂O₃ ratio of 30

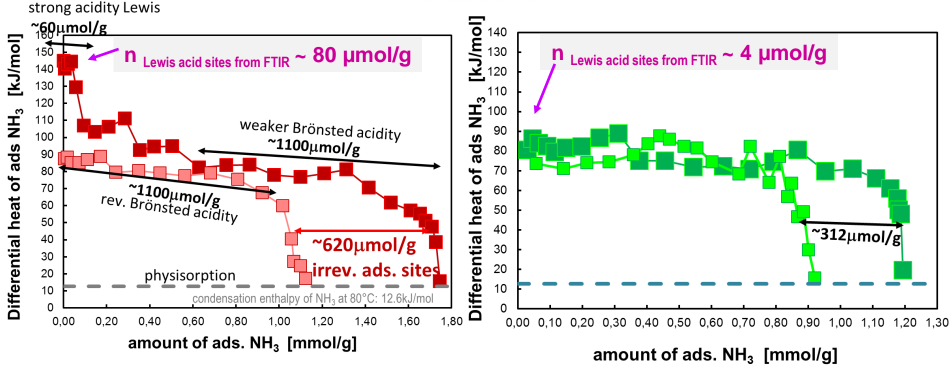
n Brønsted acid sites = 560µmol/g



H-ZSM5-80

 SiO_2/Al_2O_3 ratio of 80

n _{Brønsted acid sites} = 312μmol/g



n relevant Brønsted acid sites = n irrev. ads sites - n Lewis acid sites





Stoichiometric and measured acidities for H-ZSM5 zeolites TPSR/TPD & microcalorimetry & FTIR

Quantification of Brønsted Acid Sites

Sample Stoichiometric acidity ^[5] µmol/g		FTIR ^[6] µmol/g	Microcalorimetry ^[7] Alkylamine TP μmol/g μmol/g	
H-ZSM5-30	525	437	560	557
H-ZSM5-80	204	269	312	297

> The comparison shows that the obtained results from the TPSR are consistent with the microcalorimetry and IR measurements.

^[5] A. S. Al-Dughaither, H. de Lasa, Ind. Eng. Chem. Res. 2014, 53, 15303–15316.

^[6] O. Bortnovsky, Z. Melichar, Z. Sobalík, B. Wichterlová, Micropor. Mesopor. Mat. 2001, 42, 97–102.

^[7] S. Wrabetz, X. Yang, G. Tzolova-Müller, R. Schlögl, F. C. Jentoft, J. Catal. 2010, 269, 351–358.; Klaus Dieter Friedel Ortega, Dissertation, "6.3.4. Investigation of gasphase acidity by NH3 microcalorimetry and NH3-TPD, S. Wrabetz, page102.



Titration of basic sites on the surface of ceria using CO₂ at 40°C



Titration of basic sites on the surface of supported ceria using acidic CO₂ at 40°C ³



DEACON reaction

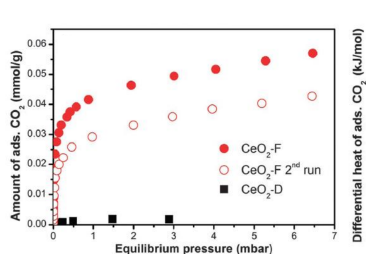
 $4 \text{ HCI} + O_2$

Oxidation of HCl to Cl₂ over CeO₂

$$T = 430$$
°C, HCl/O₂/N₂, 100ml/min 2 Cl₂ + 2 H₂O

Intention: Understanding CeO₂ as a Deacon catalyst, characterize ceria in its fresh CeO₂-F and

post-reaction states CeO₂-D



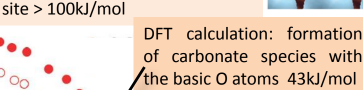
Structural defects such as edges in nanoparticles ¹ or vacancies in more open facets such as (110) ²

DFT calculation: single vacancy

0.03

Amount of ads. CO, (mmol/g)

0.04



DFT calculation: partially hydroxylated surfaces or oxygen vacancies are filled by one of the O atoms of the CO2 molecules 18kJ/mol

Adsorption sites

on the CeO₂(111) surface

The basic character of the ceria surface has been eliminated upon reaction, indicating that most of the basic lattice O sites are exchanged by chlorine and that the OH groups formed are rather acidic.

NH₃ ads. 40°C fresh: 30 kJ/mol 75µmol/g acid sites used: 50 kJ/mol 75µmol/g acid sites

> increased acidity of the chlorinated surface

0.00

0.01

⁽¹⁾ G. N. Vayssilov, M. Mihaylov, P. S. Petkov, K. I. Hadjiivanov and K. M. Neyman, J. Phys. Chem. C, 2011, 115, 23435–23454.

⁽²⁾ M. Nolan, S. C. Parker and G. W. Watson, Phys. Chem. Chem. Phys., 2006, 8, 216–218.

⁽³⁾ Farra, R., Wrabetz, S., Schuster, M. E., Stotz, E., Hamilton, N., Amrute, A. P., Pérez-Ramírez, J., López, N., Teschner, D., Phys. Chem. Chem. Phys., 15 (2013) 3454 - 3465 .

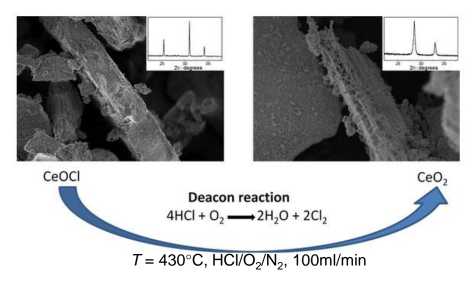


Investigation of the oxidation process: Oxidation of HCl to Cl₂ over bare and supported CeO₂

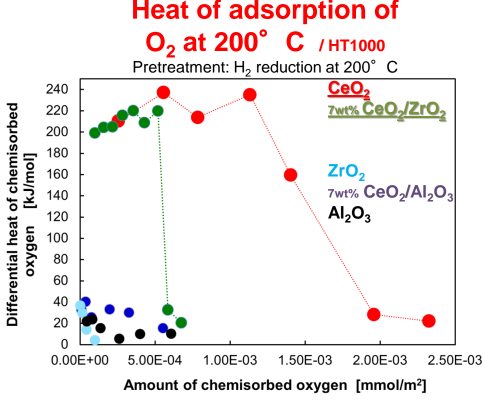


Oxidation of HCl to Cl₂ over bare and supported CeO₂ DEACON reaction





- **Best reactivity** was obtained from bare CeO₂ and CeO₂ /ZrO₂.
- One essential parameter seems to be the reducibility of the catalyst in order to create catalytically important oxygen vacancies.



Reactivity seems to be linked with the reducibility of the surface.

Farra, R., Wrabetz, S., Schuster, M. E., Stotz, E., Hamilton, N., Amrute, A. P., Pérez-Ramírez, J., López, N., Teschner, D., Phys. Chem. Chem. Phys., 15 (2013) 3454 - 3465 .
M. Mosera C. Mondellia T. Schmidtb F. Girgsdiesc M.E. Schusterc R. Farra, L. Szentmiklósi, D. Teschner, J. Pérez-Ramírez, Applied Catalysis B: Environmental 132–133 (2013) 123–131.



Study of catalytic relevant sites via calorimetry close to the reaction conditions:

Oxidation of alcohol over vanadium supported Al₂O₃



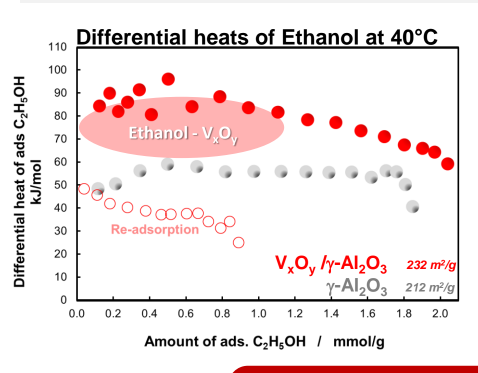
Oxidation of alcohol over vanadium supported Al₂O₃

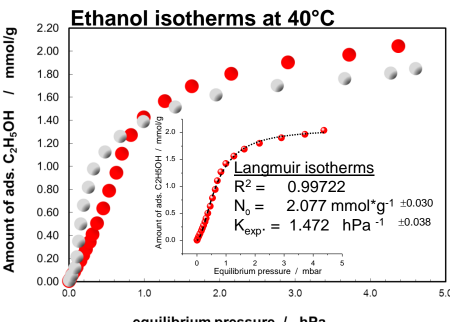


$$C_2H_5OH + \frac{1}{2}O_2 \longrightarrow C_2H_4O + H_2O \quad \Delta_RH = -178,5 \text{ kJ/mol}$$

Catalyst: $V_xO_v / \gamma - Al_2O_3$ $T_{reaction} = 140 - 200 \,^{\circ}C$

Intention: Determination of kinetic data: N_{ads} , K_{ads} and ΔH_{ads} of Ethanol on $V_x O_y / Al_2 O_3$





equilibrium pressure / hPa

Kinetic parameters determined for the ethanol ads. on V_xO_y/Al₂O₃

= 2.077mmol*q⁻¹

hPa⁻¹

 $\Delta H_{ads.}$ 90 ±10 kJ*mol⁻¹

A. Dinse, S. Khennache, B. Frank, C. Hess, R. Herbert, S. Wrabetz, R. Schlögl and R. Schomäcker, J. of Mol. Catal. A: Chem. 307 (2009) 43-50.

Study of catalytic relevant sites via calorimetry close to the reaction conditions:

New insights on active sites of oxygen functionalized carbon nanotube (oCNT) for oxidative dehydrogenation (ODH)



Oxidative dehydrogenation over oxygen functionalized carbon nanotube - oCNT

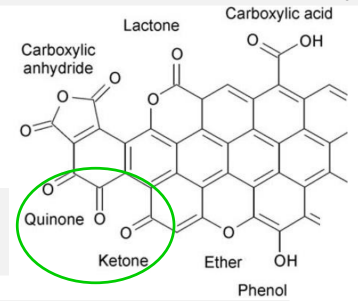


Carbon based materials are active catalysts for the oxidative dehydrogenation (ODH) reaction [1]



Quinone groups are believed to be the active sites. These **nucleophilic oxygen** species can selectively abstract hydrogen atoms ^[2-5]

The well defined active surface oxygen sites will be created by oxygen functionalization of the CNTs with HNO₃.



Intention:

To establish a mechanistic model for carbon catalyzed ODH.

Ads. properties are important input for kinetic modeling.

The research project was performed in a multidisciplinary approach (quasi *in situ* adsorption microcalorimetry, *in situ* XPS, kinetics, DFT).

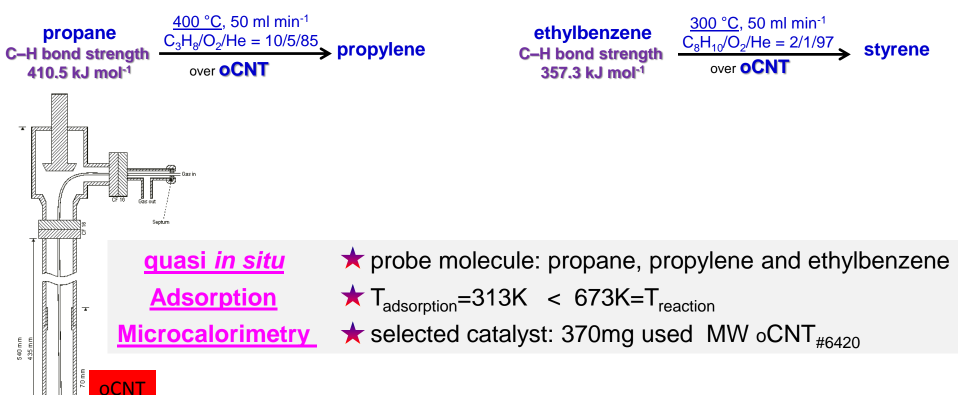


Catalytic Performance



We choose the ODH of propane and ethylbenzene (EB) as the model reactions.

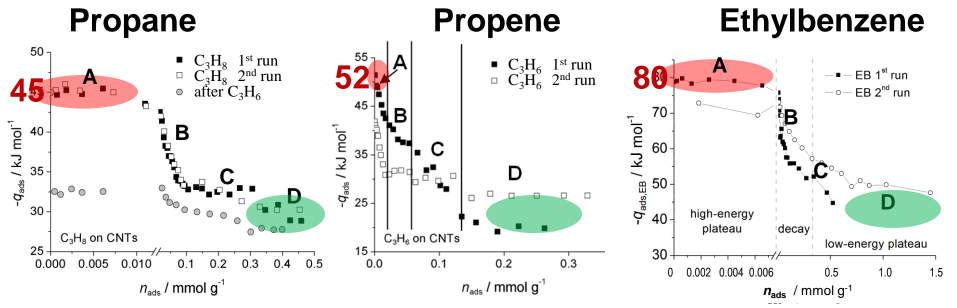
Model reaction





Differential heats of propane, propene and EB adsorption at 40°C [1]





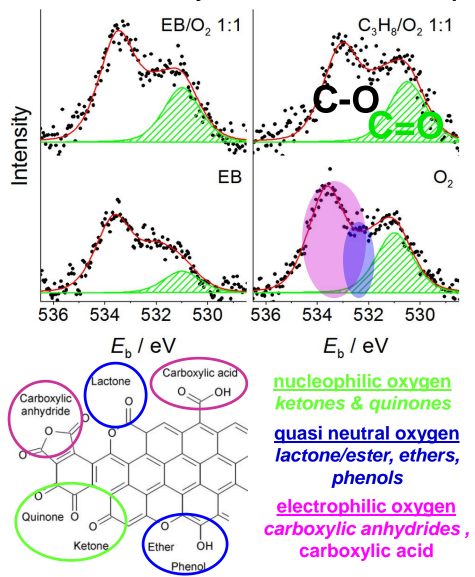
- ~8 μmol/g active sites equilibrated in EB and propane ODH reaction
- > 3-5% of the surface is covered by high-energy adsorption sites which can be correlated to 4-5% of surface oxygen determined by XPS [2]
- ➤ EB reacts stronger (80 kJ/mol) than propane (45 kJ/mol).
- Graphite (free of oxygen) shows a constant low level of diff. heats for propane (32kJ/mol) and propylene (40kJ/mol); reversible

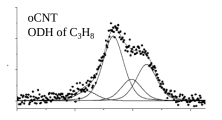


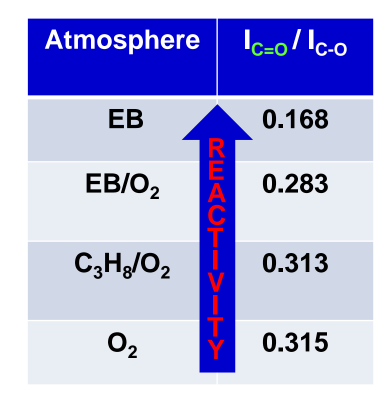
O1s ranges of XP spectra



MW oCNT catalyst under different atmospheres @ 350°C.



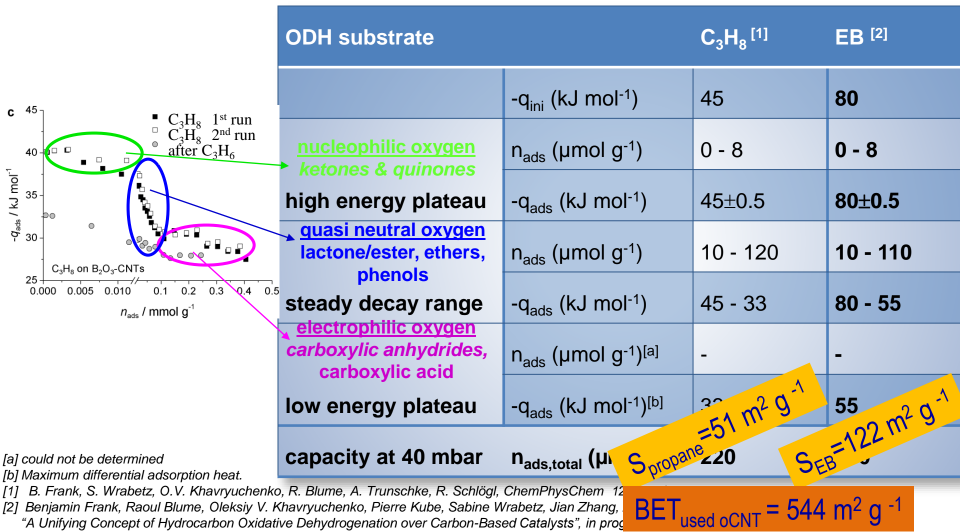








Site classification & quantification from microcalorimetric analyses, XPS, and TPDnot shown of active surfaces of oCNT catalysts in ODH of propane and EB [1,2]



[2] Benjamin Frank, Raoul Blume, Oleksiy V. Khavryuchenko, Pierre Kube, Sabine Wrabetz, Jian Zhang, "A Unifying Concept of Hydrocarbon Oxidative Dehydrogenation over Carbon-Based Catalysts", in prod



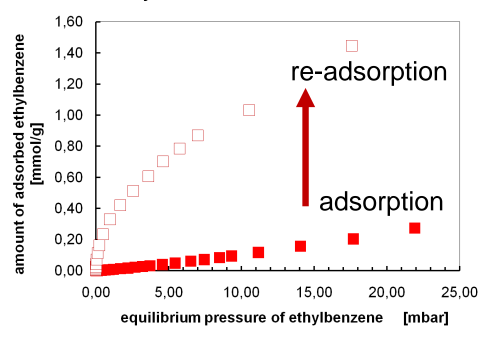
Adsorption Isotherm 40°C



Explanation:

- high energy ads. sites Ketone reacts with irreversibly to C-OH groups
- → formation of H₂O during initial reaction → blocked active sites → turns the surface more hydrophilic → hindering the ads. of non-polar EB molecule
- → reversible 2nd ads. run of EB without reaction (consumed C=O groups) and hence no formation of H₂O
- → more weak ads, sites available for EB

Ethylbenzene on used oCNT



➤ 2nd run of EB ads. reveals a drastic increase of the total adsorption capacity

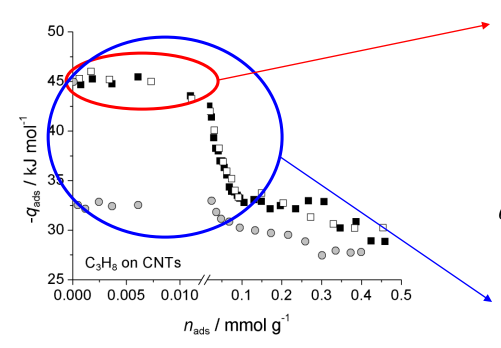


Analyzing of Adsorption Isotherm 40°C



Note: - Adsorption Isotherms only for oxygenated sites which are relevant for catalysis

- Adsorption isotherms of propylene and EB have not been evaluated because of irreversible processes and surface reactions.



Energetically uniform sites

→ Langmuir model

 $\theta = \frac{K p}{1 + K p}$ K: Langmuir adsorption constant

$$\theta = \frac{RT}{q_{ads}^{(\theta=0)} - q_{ads}^{(\theta=1)}} \ln \left(\frac{1 + \frac{p}{p^*} \exp \frac{q_{ads}^{(\theta=0)}}{RT}}{1 + \frac{p}{p^*} \exp \frac{q_{ads}^{(\theta=1)}}{RT}} \right)$$

p*: is a constant combining surface and molecular properties

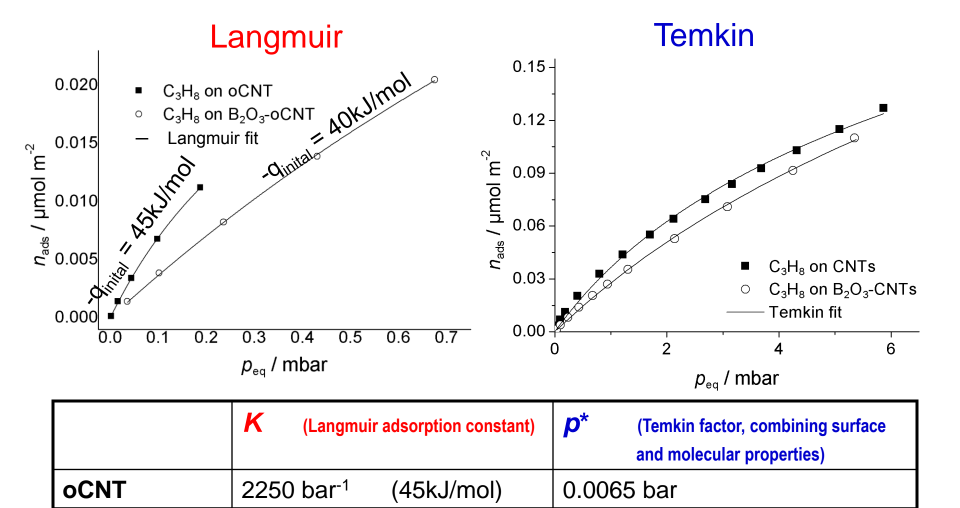
Energetically non-uniform sites (decay of q_{ads} with increasing coverage) \rightarrow Temkin model



B₂O₃-oCNT

Analyzing of Adsorption Isotherm 40°C





(40kJ/mol)

0.048 bar

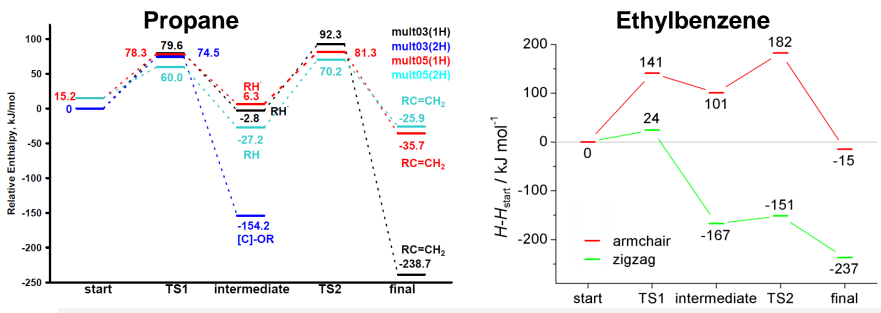
370 bar-1



Electronic structure theory calculations by DFT [1]



Energy Diagram for ODH of propane and EB (1st H-abstraction) over zigzag-termination of the carbon cluster



- the barrier for activation of propane is higher than for EB
- the zigzag-termination is much more active than the armchair geometry



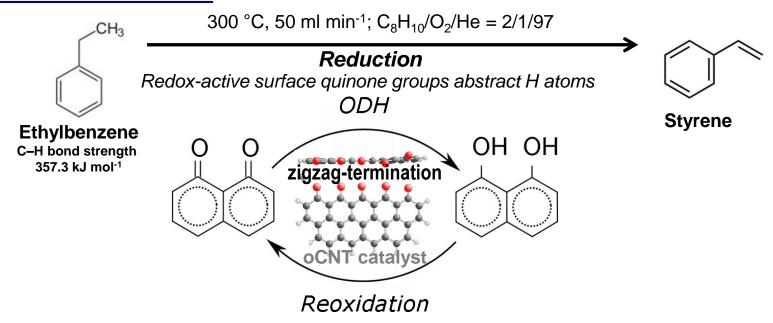


Conclusion



- Classification and quantification of the of the carbon surface under reaction conditions.
 - ➤ EB: lower activation barriers and stronger irreversible adsorption well correlate with the lower stability of the benzylic C-H bond over the aliphatic C-H bond in propane

two-site redox kinetics:



The redox cycle of surface C=O and C-OH groups is the key process, which includes only a small fraction of surface O species (2.4 m² g⁻¹, 8 μmol/g, 3-5% calo/XPs) and favorably occurs at the zigzag-termination of sp2 carbon planes.

Applications of microcalorimetry in heterogeneous catalysis



Study of catalytic relevant sites via calorimetry close to the reaction conditions:

Selective oxidation of propane to acrylic acid over MoVTeNb oxide catalyst



Selective oxidation of propane to acrylic acid over M1



study of the post-reaction state of the surface "used catalyst" Intention: in comparison with the prepared state of the surface "fresh catalyst" in order to describe a structure-selectivity relationship

Adsorption

propane as probe molecule

Microcalorimetry:

 \star T_{adsorption}=313K < 673 K=T_{reaction}

selected catalysts: different selective catalysts

pure-phase MoVTeNb $S_{aa} = 53\%$ $_{\text{modified}}$ MoVTeNb by an oxidizing agent $S_{aa} = 37\%$

MoV oxide

S₂₂ = 1.8 %



N₂ with a cooling rate of

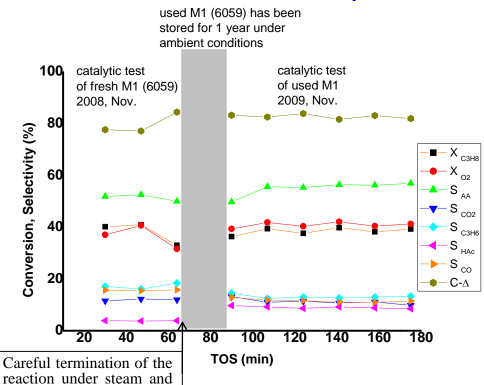
5 K/min.

Active catalyst surface stable under ambient conditions?



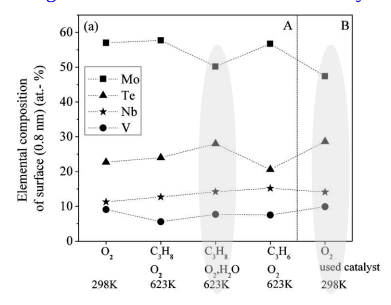
CATALYTIC TEST (fresh/used catalyst)

propane oxidation data obtained from fresh and used MoVTeNb catalyst



SURFACE STUDY (fresh/used catalyst)

in situ XPS: surface element composition during and after PO over MoVTeNb catalyst



Surface composition of the used M1 is similar to the composition measured in the *in situ* experiment in the presence of steam.

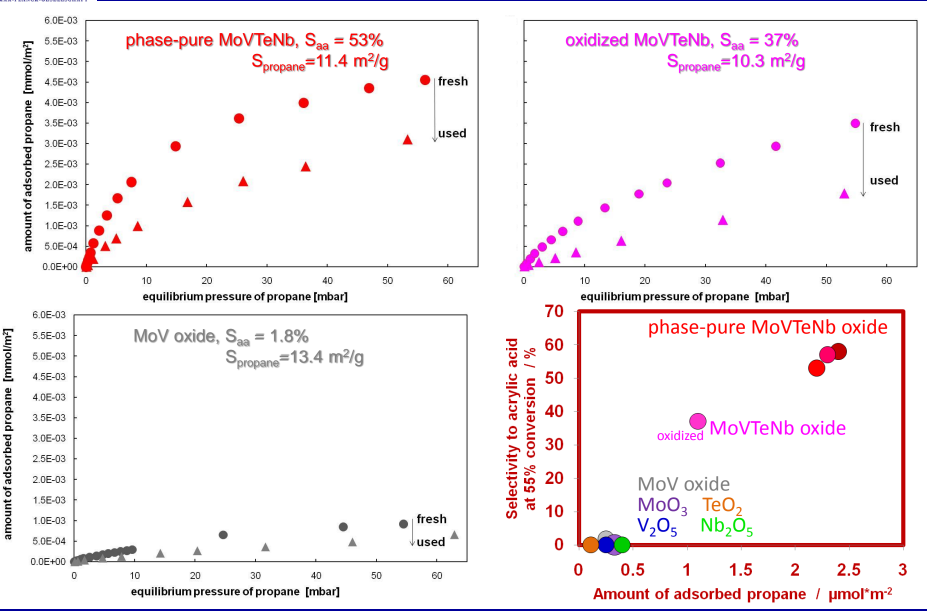
[A. Celaya Sanfiz et.al., J. Phys. Chem. C, 2010, 114, pp 1912]

ex situ surface analysis of the used catalyst can be done



Propane adsorption isotherms of different selective catalysts

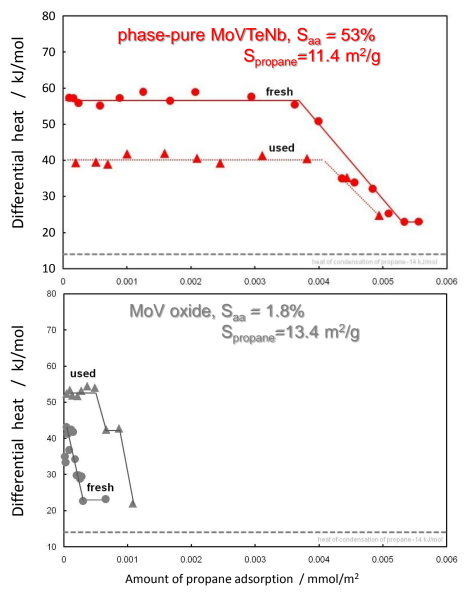


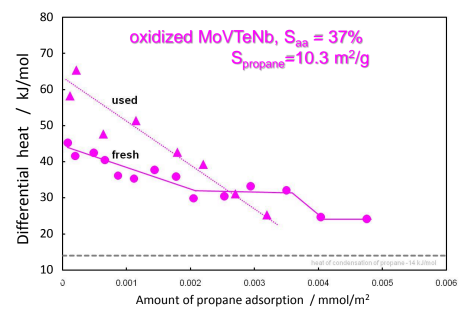




Differential heats of adsorbed propane of different selective catalysts







- Strength of interaction of propane with the surface changes during reaction.
- S_{aa} is correlated with energetically homogenous distributed propane adsorption sites
- S_{aa} is associated with reduced interaction of propane and active surface sites
- \bigstar CO & CO₂ ads. exp. : weak acid/basic sites



Structure-selectivity relationship





the prepared state of the surface is different from the post-reaction state of the surface

dynamic surface during reaction



remarkable selective (S_{aa} = 53%) MoVTeNb oxide surface is characterized by high density of energetically uniform propane adsorption sites with weak acid-base character



The weaker interaction of propane with the post-reaction state of the surface is apparently favorable for the catalytic performance; perhaps because of facile product desorption.

Applications of microcalorimetry in heterogeneous catalysis

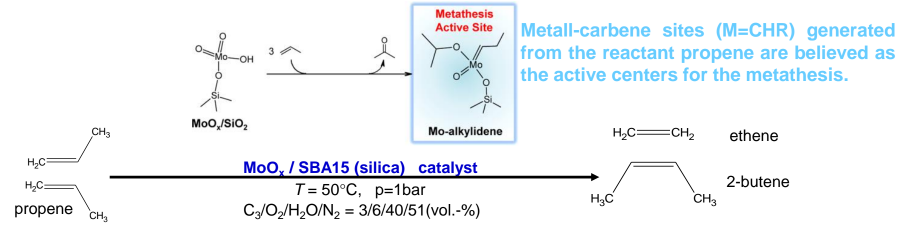


Study of catalytic relevant sites via calorimetry under reaction conditions:

Active Site Quantification on Propylene Metathesis over MoOx/SBA-15







<u>Intention:</u> Studying of the catalytically active surface sites - quantification

Adsorption

*

propylene as probe molecule

Microcalorimetry:



 $T_{adsorption} = 50^{\circ}C = T_{reaction}$



selected catalysts: different activity

13 % MoOx/SBA15 M 10 % MoOx/SBA15 M

 $Metathesis \ rate = \ \ \textbf{3.0} \ \mu mol/m^{2*}h$

0 % MoOx/SBA15 Metathesis rate = **13.0** μ mol/m^{2*}h 5 % MoOx/SBA15 Metathesis rate = **1.5** μ mol/m^{2*}h

SBA15 Metathesi

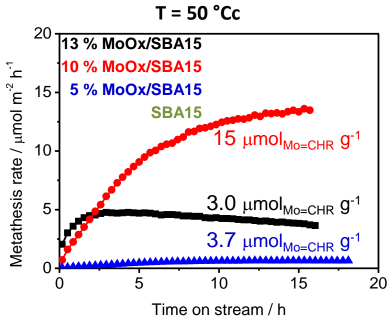
Metathesis rate = $0.0 \, \mu \text{mol/m}^{2*}$

K. Amakawa, S. Wrabetz, R. Schlögl et.al.; J. Am. Chem. Soc., 134 (28) (2012) 11462-11473.





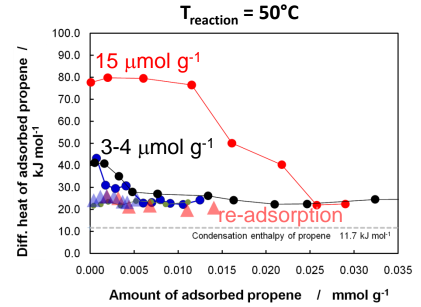
Propene Metathesis Activity



Activity strongly depends on the Mo loading

Microcalorimetry

Differential heat of propene adsorbed on MoOx/SBA15



➤ Correlation between amount & strength of C₃H₆ adsorption sites and catalytic activity.

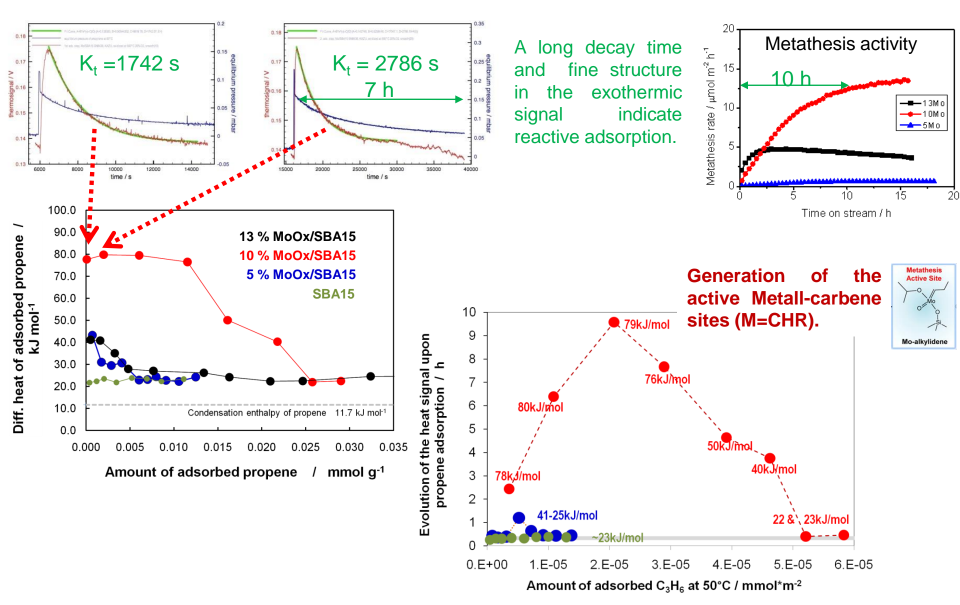
Post-reaction carbene titration

Active site counting was performed after metathesis using post-reaction ethene- d_4 (CD₂=CD₂) metathesis to titrate the formed metal-carbene (Mo=CH-CH₂) sites.

$$Mo=CH-CH_3 + CD_2=CD_2 \rightarrow Mo=CD_2 + CD_2=CH_2-CH_3$$



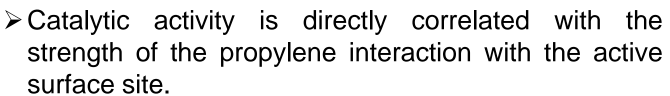


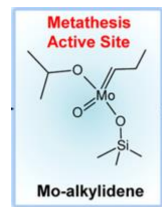






- > The active catalyst is characterized by:
 - higher amount of ads. sites for propylene
 - strong and irreversible adsorption of propylene on MoOx sites
 - energetically homogenously distributed active sites





> ca. 1% of Mo atoms formed active sites



Applications of microcalorimetry in heterogeneous catalysis



Study of catalytic relevant sites via calorimetry under reaction conditions:

Silver as a catalyst for the ethylene epoxidation - Ag-O system in catalysis -

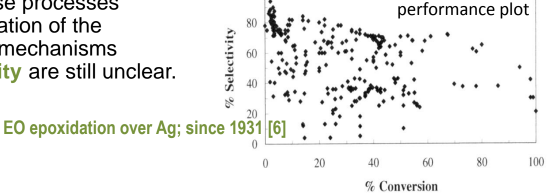


The Ag-O system in catalysis



Ag is used as a catalyst in two important large scale processes in the chemical industry, the ethylene epoxidation and methanol oxidation.

The **economic relevance** of these processes motivated the extensive investigation of the Ag-O system in past years. The mechanisms behind the **remarkable selectivity** are still unclear.



- ➤ The current understanding of oxidation reactions on Ag catalysts rely on the knowledge about the oxygen species formed on silver [1].
- Final goal is to safely translate the electronic signatures obtained by X-ray spectroscopies into structural information, which can be used to construct reaction mechanisms [4,5].

^[1] C. Hess, R. Schlögl, A. T. Bell, A. Trunschke, A. Knop-Gericke, Nanostructured Catalysts: Selective Oxidations, Royal Society Of Chemistry, 2011.

^[2] T. C. R. Rocha, A. Knop-Gericke, R. Schlögl, The Journal of Physical Chemistry C 2012, 116, 11408-11409.

^[3] R. Reichelt, S. Gunther, J. Wintterlin, Journal of Physical Chemistry C 2011, 115, 17417–17428.

^[4] T. C. R. Rocha, A. Oestereich, D. V Demidov, M. Hävecker, S. Zafeiratos, G. Weinberg, V. I. Bukhtiyarov, A. Knop-Gericke, R. Schlögl, Phys. Chem. Chem. Phys. 2012, 14, 4554–64.

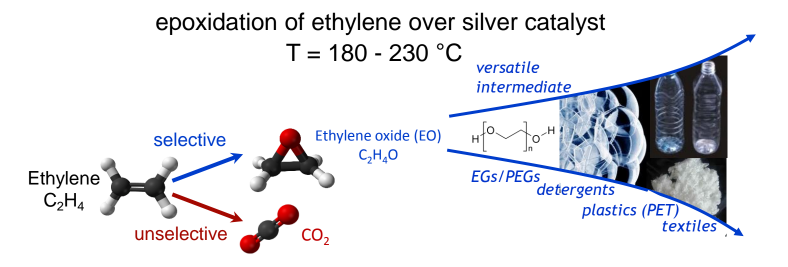
^[5] V. I. Bukhtiyarov, M. Hävecker, V. V. Kaichev, A. Knop-Gericke, R. W. Mayer, R. Schlögl, Phys. Rev. B 2003, 67, 235422.

^[6] B. K. Hodnet, Heterogeneous Catalytic Oxidation, John Wiley & Sons, New York, 2000.



The Ag-O system in catalysis





<u>Intention:</u> to construct reaction mechanisms. The energetic data provided by calorimetry is essential to any reaction mechanism. It will provide additional information to interpret the spectroscopic measurements and it will also be used as reality check for the predictions of the computational calculations.

quasi in situ

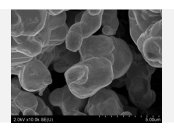
Oxygen

Adsorption:

 \star T_{reaction} = 230 °C

Microcalorimetry

* selected catalyst: Ag powder <45micron





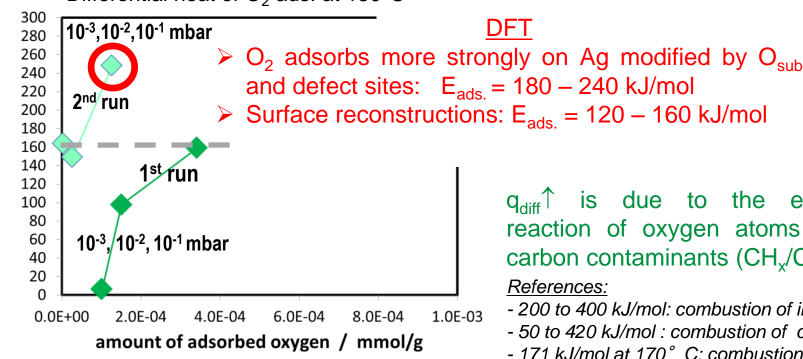
differential heat of oxygen / kJ/mol

The Ag-O system in catalysis



Initial State

Cleaning of the silver surface observed by calorimetry Differential heat of O₂ ads. at 150°C



Note: mild pre-treatment (150° C, 0.1 mbar, 2h) is enough to clean the samples also observed by NAP-XPS q_{diff} is due to the exothermic reaction of oxygen atoms with the carbon contaminants (CH_x/CO₃).

References:

- 200 to 400 kJ/mol: combustion of impurities [4]
- 50 to 420 kJ/mol: combustion of contaminants [5]
- 171 kJ/mol at 170° C: combustion of impurities [6]

Auroux, A. & Gravelle, P.C., 1981. Comparative study of the bond energy of oxygen at the surface of supported silver catalysts and of the activity of these catalysts for ethylene epoxidation. Thermochimica Acta, 47, pp.333–341.

Anderson, K.L., Plischke, J.K. & Vannice, M., 1991. Heats of Adsorption of Oxygen, Ethylene, and Butadiene on Al₂0₃- Supported Silver. J. of Catalysis, 160, pp.148–160.

Anderson, K.L., Plischke, J.K. & Vannice, M., 1991. Heats of Adsorption of Oxygen, Ethylene, and Butadiene on Al203- Supported Silver. Journal of Catalysis, 160, pp.148–160.1



oxygen / kJ/mol

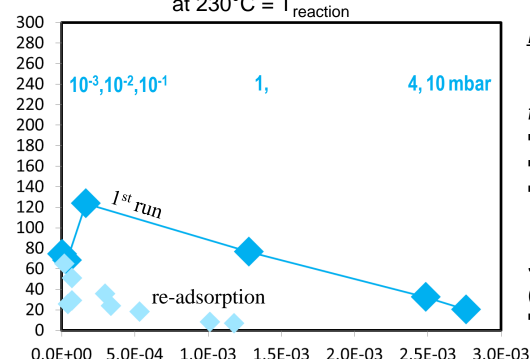
differential heat of

The Ag-O system in catalysis



Steady State

Differential heat of O_2 ads. on the clean Ag surface at $230^{\circ}C = T_{reaction}$



References:

[12]

H is the standard enthalpy of formation for model adsorbates at 298.15 K

■ $O_{ads.}$: H = -63.0 kJ/mol

■ $O_{2 \text{ ads.}}$: H = -44.5 kJ/mol

• $O_{ads.}/O_{surface}$: H = -103.0 kJ/mol

[13]

Standard enthalpies of formation at 298 K (ΔH°_{f}) of O adatoms on Pt(111):

■ 109 – 99 kJ/mol

amount of adsorbed oxygen / mmol/g

Note: The enthalpy is insignificantly affected by the temperature (DFT)

^[12] C. Stegelmann, N.C. Schiødt, C.T. Campbell, and P. Stoltze, J. Of Catal. 221 (2004) 630-649.

^[13] Eric M. Karp, Charles T. Campbell , Felix Studt, Frank Abild-Pedersen and Jens K. Nørskov; SLAC-PUB-15339.



oxygen / kJ/mo

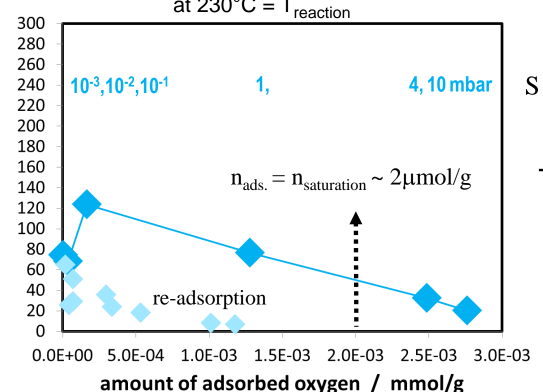
differential heat of

The Ag-O system in catalysis



Steady State

Differential heat of O_2 ads. on the clean Ag surface at $230^{\circ}C = T_{reaction}$



Specific surface area of Ag for O_2 at $T_{reaction} = 230$ °C

$$S_{Ag} = \frac{n_{ads.} \cdot Avogadro constant}{Surface sites density \Gamma_{fcc lattice, Ag}}$$

$$\frac{2 \cdot 10^{-6} \text{ mol} \cdot 6.022 \cdot 10^{23} \text{ particles} \cdot \text{cm}^2}{\text{g} \cdot 1.4 \cdot 10^{15} \text{ atoms} \cdot \text{mol}}$$

$$S_{Ag-O} = 0.86 \text{ m}^2/\text{g}$$

BET_{N2.77K} = 0.72 m²/g

$$\Delta S = 0.14 \text{ m}^2/\text{g}$$

The excess can be due to dissolving of oxygen into the Ag subsurface.



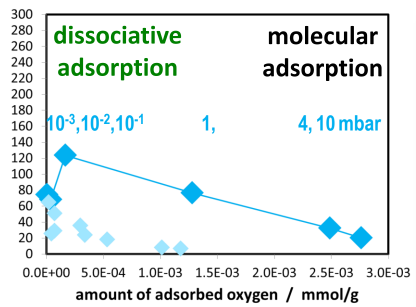
differential heat of oxygen / kJ/mol

The Ag-O system in catalysis



Experiment

Theory



DFT calculations

system	Eads per 1/2 O ₂ [kJ/mol]	Eads per O ₂ [kJ/mol]	
reconstructions O _{ads} stronger on Ag(110)	60-80	120-160	
unreconstructed O _{ads} on hollow sites of Ag(111)	40-60 80-12		
on subsurface O	110-120	220-240	
surface defects	90-120	180-240	
subsurface	<10	10	
grain boundaries	10-20	20-40	
O2 clean surface	5-10	10-20	
O2 on sub O	10-20	20-40	
O2 on defects	20-30	40-60	

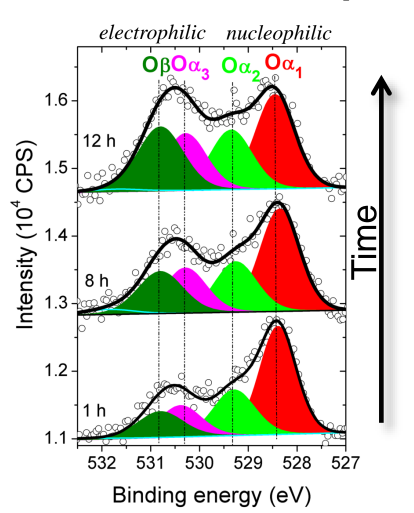


The Ag-O system in catalysis



O1s XPS spectra for Ag powders

measured in situ under 0.5 mbar O₂ at 180 °C



The distribution of O species on the Ag surface dynamically changes with time as the catalyst surface equilibrates with the gas phase at mbar pressure and temperatures typical of alkene (180-230 °C).

→ Dynamics at 180 °C



Ag-O dynamics depend on time / temperature and pressure

Summary



					−
T ° C	p₀₂ mbar	n _{ads.} μmol _{O2} /g _{Ag}	q _{diff} kJ/mol	Ag-O dynamics	DFT Agsub O
T _{cleaning} 150	< 10 ⁻¹	0.5	>120 - 260	 dissociative adsorption cleaning the surface from CH_x/CO₃ initial formation of the surface reconstructions 	Ag(110)reconstruction
T _{reaction} 230	10 ⁻¹ - 1	1.5	>50 - 120	 extensive structural changes * oxygen begins to dissolve to subsurface ** oxide-like structures = 60 -80 kJ/mol [8] chemisorbed oxygen modified by a subsurface oxygen *** 	
T _{reaction} 230	>1 - 10	1	< 50	 molecular oxygen adsorbed on surface - vacancies **** formation of electrophilic oxygen ***** O₂ on subsurface oxygen 	> () () () () () () () () () (

^{*} Ag atoms moving from defects and edges to form the reconstructions. Island formation at low coverages.

^{**} Formation of oxide like reconstruction and/or surface oxide layer.

^{***} Depends on time/temperature because its formation is limited by oxygen diffusion to subsurface.

^{****} Proposed by theory and calorimetry. In this case the time/temperature dependence is related to the defect formation.

The active site for epoxidation! Two interpretations are believed: O_2 stabilized on defects on O covered surface (DFT) or surface oxygen modified by sub-surface species (XPS)

Applications of microcalorimetry in heterogeneous catalysis



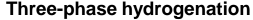
Study of catalytic relevant sites via calorimetry under reaction conditions:

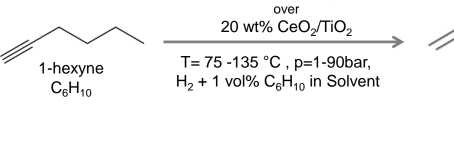
1-hexyne adsorption at $T_{\rm react.}$ on supported ceria , being unexpectedly active in the hydrogenation of 1-hexyne

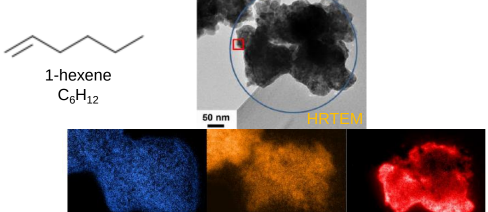


1-hexyne adsorption at T_{react} = 80°C on supported ceria









Intention: Does calorimetry offers the possibility to distinguish between catalysts of similar characteristic properties_{IR. XRD.TEM.XPS.BET}), but different catalytic activity?

Adsorption Microcalorimetry:



1-hexyne as probe molecule



 $T_{adsorption} = 80^{\circ}C \leftarrow as the Arrhenius plot of 1-hexyne hydrogenation$ is smooth in the whole 75 - 135 °C range

FETEM maps



selected catalysts: different active catalysts

20 wt% CeO₂/TiO₂ -WI very active

(wet impregnation; # 17380)

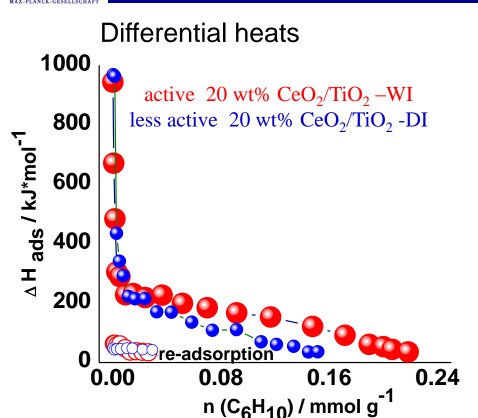
20 wt% CeO₂/TiO₂ -DI less active

(dry impregnation; # 17378)



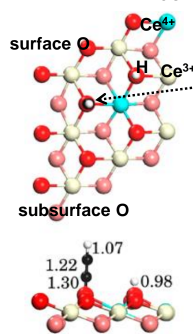
1-hexyne adsorption at T_{react} = 80°C on supported ceria







C₂H



The initial step of the C_2H_2 hydrogenation to C_2H_4 is the dissociative homolytic adsorption of C_2H_2 . This process is strongly exothermic. (-1.77eV)

 $\Delta H_{
m ads.}$, initial stage :

< 200 - 900 kJ/mol

10μmol/g

- → very strong irreversible ads.
- → multiple dehydrogenation steps [1] formation of stable surface intermediates [2]

 $\Delta H_{
m ads.}$, plateau

~ 200 kJ/mol

- 60μ mol/g \rightarrow dissociative homolytic adsorption of C₆H₁₀
 - → single dehydrogenation step [1]

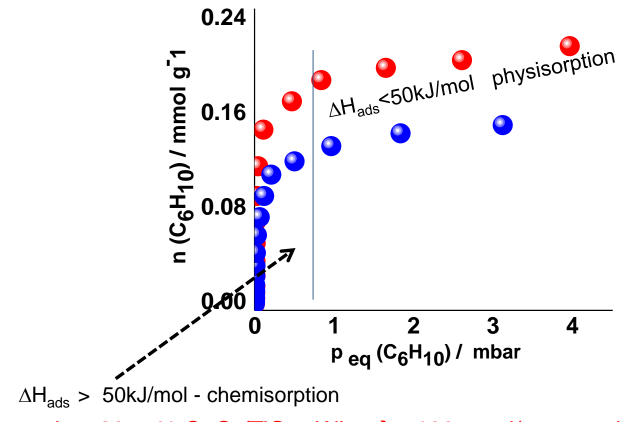
[1] J. Carrasco, G. Vilé, D. Fernández-Torre, R. Pérez, J. Pérez-Ramírez, M. V. Ganduglia-Pirovano, J. Phys. Chem. C 2014, 118, 5352. [2] Claire L. Pettiette-Hall, Donald P. Land, Robert T. McIver, Jr., and John C. Hemminger, J. Am. Chem. SOC. 1991, 113, 2755-2756.



1-hexyne adsorption at T_{react} = 80°C on supported ceria



Adsorption Isotherm



very active 20 wt% CeO₂/TiO₂ –WI \rightarrow 190 μmol/g reacted 1-hexyne molecules

less active 20 wt% CeO_2/TiO_2 -DI \rightarrow 130 µmol/g reacted 1-hexyne molecules

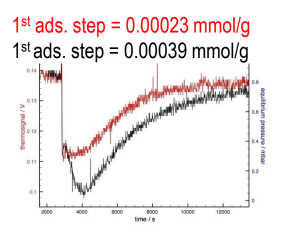


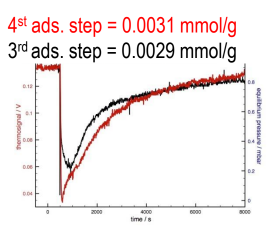
1-hexyne adsorption at T_{react} = 80°C on supported ceria

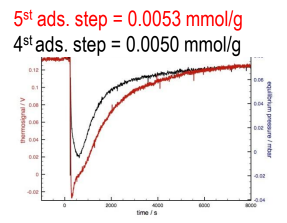


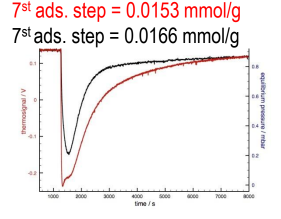
Corresponding integral heats

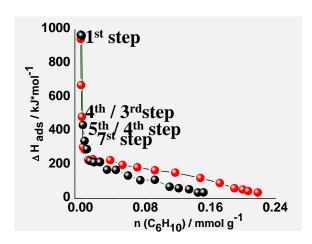
The temporal evolution of the thermo signal during 1-hexyne ads.











The adsorption signature is composed of two main contributions:

- physisorption / fast-signal
- reaction / delayed signal

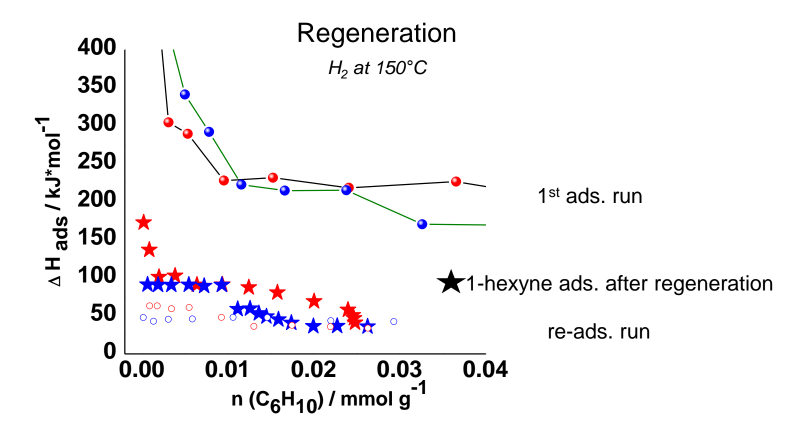
The **most active sample** has such sites that will hold 1-hexyne very weakly until the molecule chemisorb stronger. **dynamic**

In the **less active sample** the physisorbed molecule undergoes much faster side reactions or strong chemisorption.



1-hexyne adsorption at T_{react.} = 80°C on supported ceria





The regeneration of the catalyst surface using H₂ (at 150°C) has been somewhat more successful for the more active catalyst.



Summary



Yes, calorimetry is a useful tool to distinguish between catalysts of similar characteristic properties.

Key-note:

under reaction conditions a significant portion of the surface sites is covered by dehydrogenated species and is not available for hydrogenation. Nevertheless, the remaining <u>small number of surface sites is active and selective</u> in alkyne hydrogenation.

The ads. of 1-hexyne at T_{react} is composed of 2 main processes:

- 1) unspecific adsorption (physisorption <50kJ/mol)
- 2) time-consuming secondary process is due to single or multiple dehydrogenation steps and potentially oligomerization. [10] (>50 900 kJ/mol)

The most active catalyst is characterize by:

- higher amount of adsorption places for 1-hexyne 190 μmol/g
- slightly easier regeneration of the surface after 1-hexyne contact at T_{react.}
- pronounced trapped and phys. state of the adsorbates to find the most suitable place for reaction → dynamic surface is apparently favorable for the catalytic performance

Applications of microcalorimetry in heterogeneous catalysis



Reactants induced responses of catalyst surface:

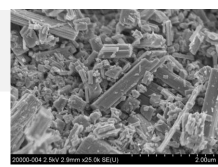
Propane and ethane ads./des. cycles (40°C) on MoV oxide model catalyst for oxidative dehydrogenation of alkanes.



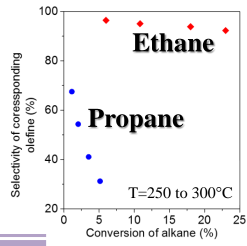
1st MoV oxide model catalyst in oxidative dehydrogenation of alkanes



MoV oxide¹ has been studied as a **model system** for a better understanding of the **complex V-containing bulk MoVTeNb** oxide catalyst



→ <u>different dynamic behavior</u> of the MoV oxide surface under reaction conditions due to the different chemical potential of the feeds while the chemical potential of the solid remains the same



Microcalorimetry

- specific ads. phenomena of PROPANE and ETHANE at 40°C
- reactants induced response of the MoV oxide surface via ads./des. cycles.



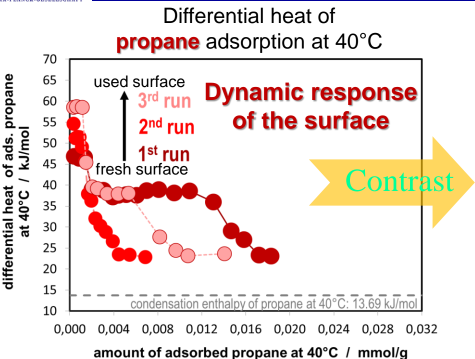
PROPANE and ETHANE adsorption on MoV oxide at 40°C

of adsorbed ethane

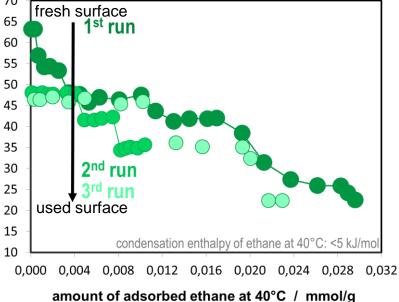
differential heat

at 40°C / kJ/mol





Differential heat of ethane adsorption at 40°C



S specific for propane $\sim 3.6 \text{ m}^2/\text{g}$ $BET = 25.52 \text{ m}^2/\text{g}$

S specific for ethane
$$\sim 2.9 \text{ m}^2/\text{g}$$

 $BET = 25.52 \text{ m}^2/\text{g}$



PROPANE and ETHANE adsorption on MoV oxide at 40°C



propane

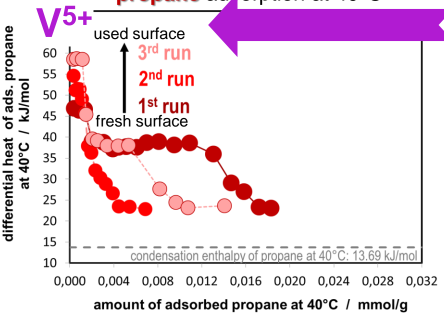
ethane

20

10

Conversion of alkane (%)

Differential heat of propane adsorption at 40°C



In-situ photoelectron spectroscopy

with TOS (presents of H₂O) the surface is progressively enriched in V5+, which leads to a decrease 100 in selectivity ¹ Selectivity of coressponding olefine (%)

20

- ✓ segregation of V^{5+} occurs already at r.t.
 - ✓ alkanes are already activated at r.t.
 - **✓** dynamic nature of the surface
- very strong interaction of propane with used surface explains the decrease in selectivity caused by V-segregation

Applications of microcalorimetry in heterogeneous catalysis



Reactants induced dynamic responses of catalyst surface:

CO chemisorption cycles (30°C) on Ni/MgAl oxide catalyst for dry reforming of methane (DRM)



2nd Ni based catalysts for the dry reforming of methane

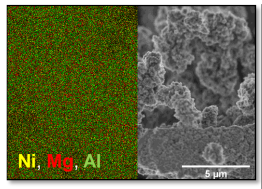


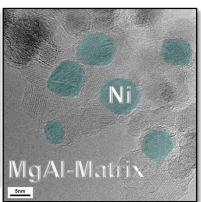
<u>Task:</u> study the influence of structural and compositional properties of nickel catalysts on the catalytic performance during DRM

Dry reforming (DRM):
$$CO_2 + CH_4 \stackrel{\text{Ni}}{=_{900} \, ^{\circ}\text{C}} 2 \text{ CO} + 2 \text{ H}_2$$
 $\Delta H^0 = 247 \text{ kJ/mol}$

Catalyst preparation pathway:

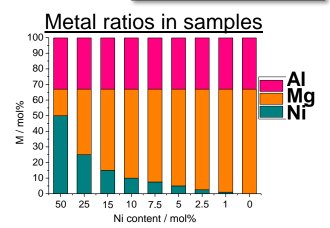
- synthesized by constant pH coprecipitation with Ni contents between 0 and 55 wt.-%.
- decomposition to mixed oxides by calcination at 600°C





Obtained Ni/MgAl oxide catalyst

- reduction at 1000°C
- $d_{P. Ni} = 7 9$ nm in all samples
- $d_{P, Ni} = 7 20$ nm only in the 50wt% sample



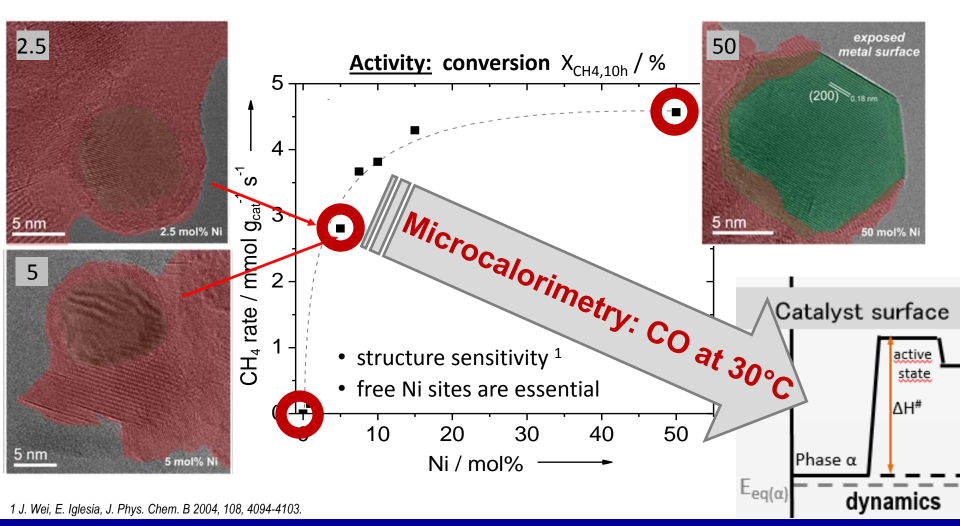
K. Mette, St. Kühl, A. Tarasov, M. G. Willinger, J. Kröhnert, S. Wrabetz, A. Trunschke, M. Scherzer, F. Girgsdies, H. Düdder, K. Kähler, K. Friedel Ortega, M. Muhler, R. Schlögl, M. Behrens, T. Lunkenbein, ACS Catal., 2016, 6 (10), pp 7238–7248.



catalysis / overgrowth / reactivity



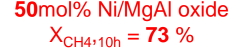
- HR-TEM images of 2.5, 5 and 50 mol% Ni/MgAl oxide
- EDX, NEXAFS & FTIR/CO/77K: overgrowth might be interpreted as a NiAl₂O₄ spinel

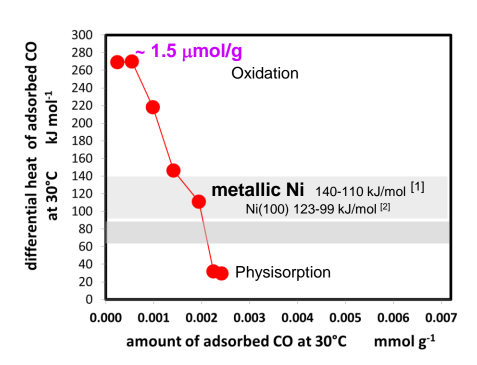


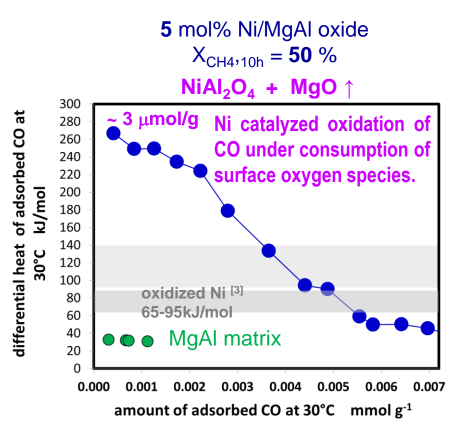


CO adsorption at 30°C using microcalorimetry









The differential heat profiles are dominated by the presence of nickel

^[1] A. Tanksale, J.N. Beltramini, J.A. Dumesic, G.Q. Lu, Journal of Catalysis 258 (2008) 366–377.

^[2] J. T. Stuckless, N. Al-Sarraf, C. Wartnaby, D. A. King, J. Chem. Phys. 1993, 99, 2202-2212.

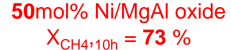
^[3] M. Cerro-Alarcó, B. Bachiller-Baeza, A. Guerrero-Ruiz, I. Rodríguez-Ramos, J. Mol. Catal. Chem. 2006, 258, 221-230.

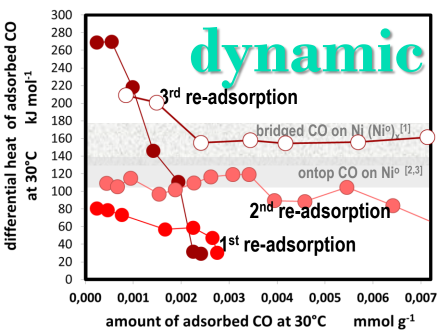


Reactants induced dynamic responses of the catalyst surface was simulated via ads./des. cycles of CO at 30°C using microcalorimetry

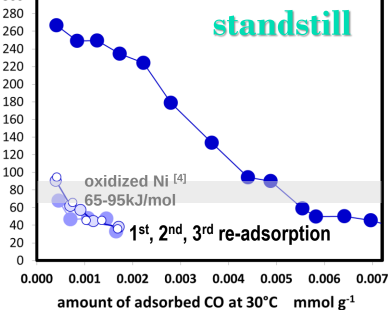
differential heat







5 mol% Ni/MgAl oxide $X_{CH4,10h} = 50 \%$ 300 of adsorbed CO at 280 260 240



Catalytically relevant Ni⁰ is back again!

[1] R. S. Bordoli, J. C. Vickerman, J. Wolstenholme, Surf. Sci. 1979, 85, 244-262.

[2] A. Tanksale, J.N. Beltramini, J.A. Dumesic, G.Q. Lu, J. of Catalysis 258 (2008) 366-377.

[3] J. T. Stuckless, N. Al-Sarraf, C. Wartnaby, D. A. King, J. Chem. Phys. 1993, 99, 2202-2212 [4] M. Cerro-Alarcó, B. Bachiller-Baeza, A. Guerrero-Ruiz, I. Rodríguez-Ramos, J. Mol. Catal. Chem. 2006, 258, 221-230.

Applications of microcalorimetry in heterogeneous catalysis



Estimation of the enthalpy of formation of the transition state (activation barrier):

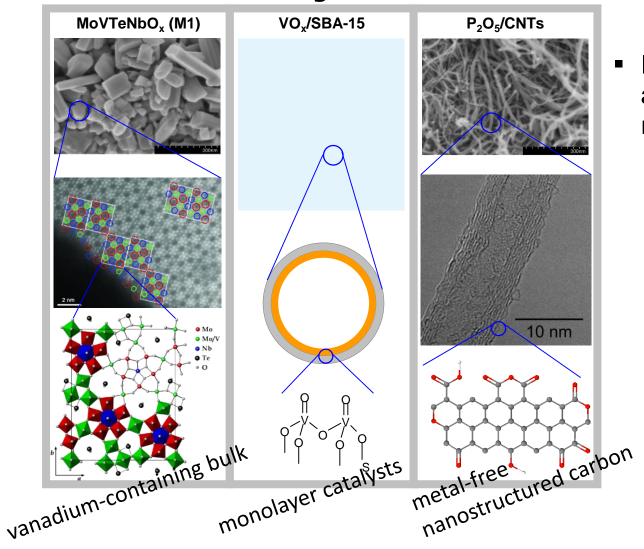
ethane & ethylene and propane & propylene ads. on the Vanadium oxide-based and Metal-free catalysts for ODH



Vanadium Oxide-based and Metal-free Catalysts in the ODH of Ethane and Propane ¹



Catalysts



 Identification of differences and similarities in the reaction network.

> structurally similar functional groups: V-OH / C-OH V=O / C=O V-O-V / C-O-C



Catalytic Performance



➤ Bulk MoVTeNb oxide, 6V/SBA-15 and P/oCNT have been compared in the ODH of C₂H₆ and C₃H₈ under identical conditions. ¹

Similarities and Differences in the reaction network

Strong temperature dependence → different transition states

enthalpy of formation of the transition state

Note: $\Delta H_{ads.} \neq T_{ads.}^2$; ads. in quasi-equilibrium ³; intermediates occure in pseudo-steady-state

E_{app}

Gas state
ΔH#

Adsorbed state

¹ P. Kube, B. Frank, S. Wrabetz, J. Kröhnert, M. Hävecker, J. Valasco-Vélez, J. Noack, R. Schlögl, A. Trunschke, ChemCatChem 9 (2017) 1-14.

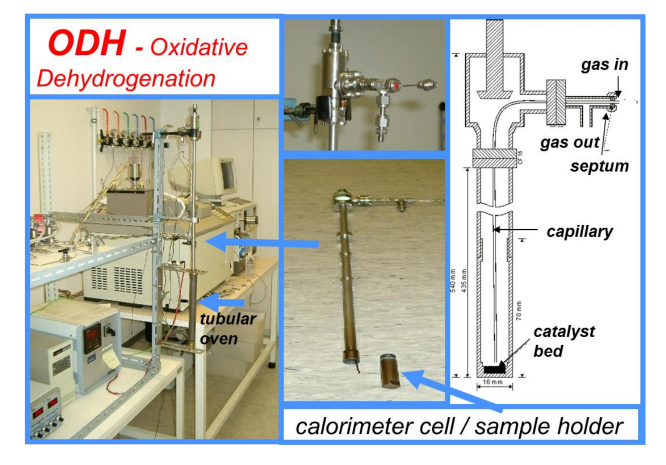
 ² a) D. C. Tranca, N. Hansen, J.A. Swisher, B. Smit, F. J. Keil, J. Phys. Chem. C 2012, 116,23408 – 23417; b) E.J. Maginn, A. T. Bell, D. N. Theodorou, J. Phys. Chem. 1995, 99,2057 –2079.
 3 K. Chen, A. T. Bell, E. Iglesia, J. Phys. Chem. B 2000, 104, 1292–1299



Creation of an active catalyst surface



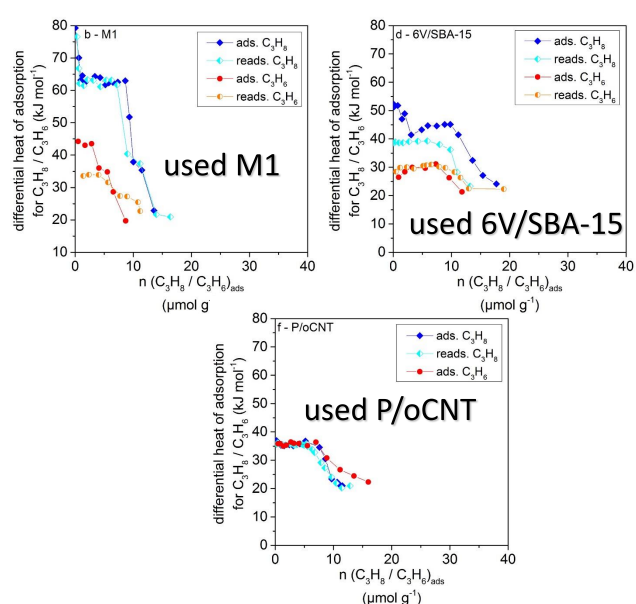
- Feed of 10% hydrocarbon (C₃H₈ or C₂H₆) and 5% oxygen in helium with a total flow rate of 20 mLmin⁻¹.
- T_{reaction} = 400°C for 6V/SBA-15, 360°C for P/oCNT and 350°C for M1
- The reaction was performed at a steady state for 20 h, subsequently, the cell was cooled down to RT in pure helium.

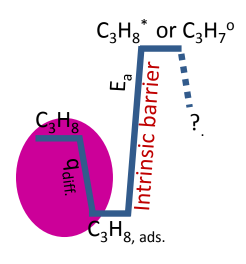




Adsorption Experiments C₃H₈ & C₃H₆ Adsorption at 40°C



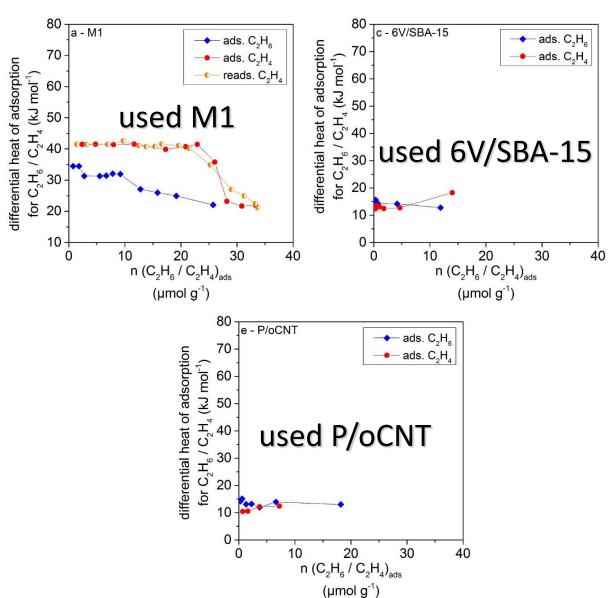


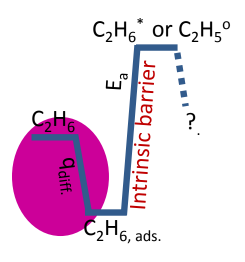




Adsorption Experiments C₂H₆ & C₂H₄ Adsorption at 40°C









Data Discussion

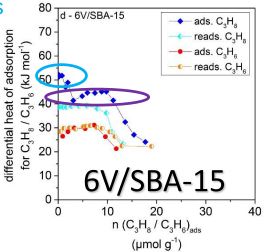


MAX-PLANCK-GESELLSCHAFT						
	M1	6V/SBA-15	P/oCNT			
S _{BET} m ² g ⁻¹	10.6	355	229			
V _{surf.} ; O _{surf.}	(30)	1113	1063			
$\Gamma_{ m V}$ atoms nm- 2	1.7	1.9	2.8			
Γ _{C3H8} molecules nm²	0.44 (63 kJ/mol)	0.0114 (52 kJ/mol)	0.013 (36 kJ/mol)			
V _{surf.} /C ₃ H _{6 ads.}	~4	~150				
C3H8,consumtion, 0	5.3 ±0.2	0.45 ±0.006	0.21 ±0.004			
r _{C2H6,consumtion, 0}	5.4 ±0.2	0.055 ±0.001	0.03 ±0.001			
q _{diff} (C ₃ H ₈)	79 / 63) rev. ads.	52 44 rrev. ads.	36 rev. ads.			
q _{diff} (C ₃ H ₆)	44 irrev. ads.	30 rev. ads.	36 rev. ads.			
q _{diff} (C ₂ H ₆)	34 rev. ads.	14 *	14 *			
q _{diff} (C ₂ H ₄)	41 rev. ads.	14 *	14 *			
E _{a, propane}	80 ±3	110 ±2	103 ±7			
E _{a, ethane}	90 ±2	121±2	110 ±7			
intrinsic barrier	143 (63 kJ/mol)	162 154	139			
intrinsic barrier	124					

6V/SBA-15:

- $-\Gamma_{\rm C3H8} = 0.0014 \text{ molecules nm}^{-2}$
- intrinsic barrier for C₃H₈ activation = 162 kJ/mol
- DFT calculations by Rozanska ² = 160 kJ/mol due to monomeric species
- \rightarrow ~0.1% of all V atoms.

~ 1%



- intrinsic barrier for C₃H₈ activation = 154 kJ/mol
- DFT calculations by Rozanska ² = 148 kJ/mol due to silica-supported vanadium oxide dimers
- \rightarrow ~ 0.6% of all V atoms

Surface concentration of V (M1 and 6V/SBA-15) and O (P/oCNT) atoms (µmol g-1)

Ea: propane/ethane consumption,0 (kJ/mol); initial rate

^{*} Differential heat near the condensation enthalpy of reaction molecules at 40°C intrinsic barrier= E_a+q_{diff} (kJ mol⁻¹)

Hävecker, M.; Wrabetz, S.; Kröhnert, J.; Csepei, L.-I.; Naumann d'Alnoncourt, R.; Kolen'ko, Y. V.; Girgsdies. F.; Schlöal. R.; Trunschke, A. Journal of Catalysis 2012, 285, 48.

X. Rozanska, R. Fortrie, J. Sauer, J. Am. Chem. Soc. 2014, 136, 7751 –776.
P. Concepcijn, P. Botella, J. M. L. Nieto, Appl. Catal. A 2004, 278, 45–56; f) R. Coast, M. Pikus, P. N. Henriksen, G. A. Nitowski, J. Phys. Chem. 1996, 100, 15011-15014



Conclusion



- Microcalorimetry and DFT: quantification of propane adsorption sites in vanadium oxide monolayer catalysts
 - resolves different degrees of V_xO_y oligomerization

V_xO_y	Microcalorimetry Intrinsic barrier for C_3H_8 activation	Density functional theorie DFT Energy barriers	Microcalorimetry % of the total amount of V _{surf}	
	kJ/mol	kJ/mol	Suii.	
Monomer (a)	162	160	0.1	0/
Dimer (b)	154	148	0.6	7 0
trimer		143		
tetramer	143	139		

ODH of propane over M1 phase: tetramers of vanadium oxide species have been postulated to be the required ensemble size ²

- Structure Activity Relationship Higher activity correlates with:
 - higher density of C₃H₈ adsorption sites
 - lower intrinsic barrier for C₃H₆ formation



Conclusion



- Microcalorimetry and DFT: quantification of propane adsorption sites in vanadium oxide monolayer catalysts
 - resolves different degrees of $V_x O_y$ oligomerization

	<u>Microcalorimetry</u>		netry	Density functional	1
V_xO_v				theorie DFT ¹	
·	Intrinsic barrier for			Energy barriers	
	C ₃ H ₈ activation		tion		
	kJ/mol			kJ/mol	
Monomer (a)		162		160	
Dimer (b)		154		148	
trimer				143	
tetramer		143		139	

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Size-Dependent Catalytic Activity of Supported Vanadium Oxide Species: Oxidative Dehydrogenation of Propane

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3 Supporting Information

ABSTRACT: Possible reaction pathways for the oxidative dehydrogenation of propane by vanadium oxide catalysts supported on silica are examined by density functional theory. Monomeric and dimeric vanadium oxide species are both considered and modeled by vanadyl-substituted silsesquioxanes. The reaction proceeds in two subsequent steps. In a first step, hydrogen abstraction from propane by a vanadyl (O=V) group yields a propyl radical bound to a HOVIV surface site. Propene is formed by a second hydrogen abstraction, either at the same vanadia site or at a different one. V^VV^W redox cycles are preferred

over V^V/V^{III} cycles. Under the assumption of fast reoxidation, microkinetic simulations show that the first step is rate-determining and yields Arrhenius barriers that are lower for dimers (114 k]/mol at 750 K) than for monomers (124 k]/mol). The rate constants predicted for a mixture of monomers and dimers are 14% larger (750 K) than for monomers only, although the increase remains within experimental uncertainty limits. Direct calculations of energy barriers also yield lower values for dimeric species than for monomeric ones. Reactivity descriptors indicate that this trend will continue also for larger oligomers. The size distribution of oligomeric species is predicted to be rather statistical. This, together with the small increase in the rate constants, explains that turnover frequencies observed for submonolayer coverages of vanadia on silica do not vary with the loading within the experimental uncertainty limits.

1. INTRODUCTION

Supported transition metal oxides are an important class of

Here, we use density functional theory (DFT) to provide information that cannot be easily obtained from experiments.

monomeric over oligomeric to polymeric transition metal oxide clusters anchored on the surface of the supporting oxide, to nanocrystallites of the active component with the structure

but proving this appears difficult. 10,11 For example, it has been shown that V-O-V bonds, which would be absent in monomeric species, cannot be identified in IR or Raman spectra because of overlap with bands of the supporting oxide. 7 UV-vis absorption spectra are also not size-discriminating, although there is no doubt that the O 2p-V 3d charge transfer transitions will shift to lower energies with increasing particle size. 8,12

Here, we use density functional theory (DFT) to provide information that cannot be easily obtained from experiments. We construct models for monomeric, dimeric, and polymeric supported species, and we examine them in comparison to surfaces of the bulk crystal. We study the reaction mechanisms

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Applications of microcalorimetry in heterogeneous catalysis



Limitations of method:

Propane ads. on vanadium-phosphorus-oxide / VPO at $T_{reaction} = 100 - 400$ °C



vanadium-phosphorus-oxide - VPO



Natural gas as raw material

Methane Ethane Propane Butane

OCM Oxidative Dehydrogenation Selective Catalytic Oxidation

Olefines, Oxygenates

Ethane, Ethylene Ethylene, Acetic acid Propylene, Acrylic acid Butylenes, Butadiene, Maleic anhydride

<u>Intention</u>: Determination of kinetic data: N_o , K and ΔH_{ads} of alkanes

on VPO at T_{reaction}

Adsorption

★ propane, ethane and n-butane as probe molecule

Microcalorimetry:

 $T_{\text{adsorption}} = 400, 300, 200, 100 \text{ and } 40^{\circ}\text{C}$

selected catalyst: VPO #10449

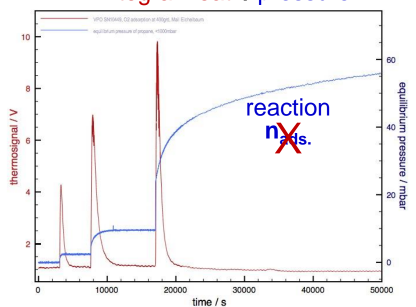


vanadium-phosphorus-oxide - VPO

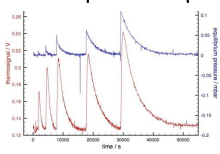


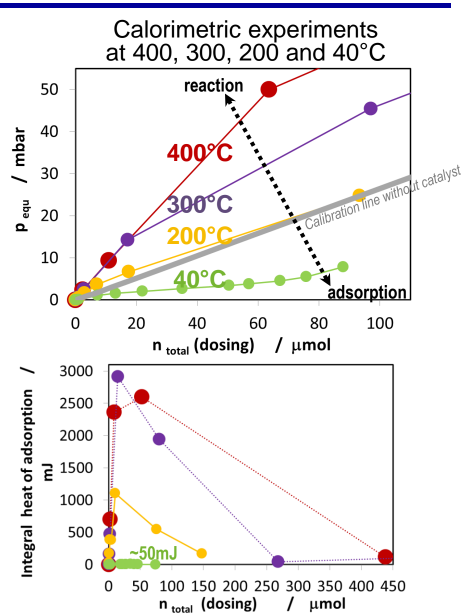
The temporal evolution of the thermo signal during propane ads. at 400°C Catalyst was oxidized at 400°C for 2h

Catalyst was oxidized at 400°C for 2h Integral heat / pressure



Classical pressure profile





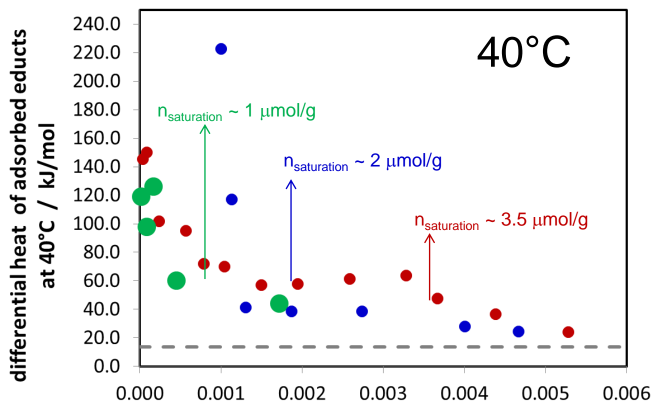


vanadium-phosphorus-oxide - VPO



Differential heat of propane / ethane / n-butane

Catalyst was oxidized at 400°C for 2h



amount of adsorbed educts at 40°C / mmol/g



The End



- Microcalorimetry alone or combined with to other techniques is a very powerful/sensitive tool to probe catalytically active surfaces quantitatively.
- Quantitative data (reversibility, heat of adsorption, number of adsorption sites, equilibrium constant) provide a basis for theoretical modeling and can contribute to a better understanding of the complex microkinetics.
- Microcalorimetry can applied under or close to reaction conditions!

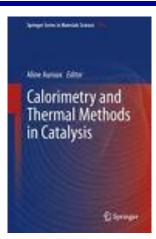
 → investigation of the <u>catalytic relevant surface sites</u>



Literature



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