**Davide Ferri :: Paul Scherrer Institut** 

# Infrared spectroscopy

Molecular aspects of catalysts and surfaces :: ETHZ

# The electromagnetic spectrum



source: Andor.com

# Importance of IR spectroscopy in catalysis



Number of publications containing in situ, catalysis, and respective method Source: ISI Web of Knowledge (Sept. 2008)

- pros ٠
- economic
- non-invasive
- versatile (e.g. solid, liquid, gas, interfaces)
- very sensitive (concentration)
- fast acquisition (down to ns!)
- no atomic resolution

### Infrared spectroscopy



# Why IR spectroscopy

- 'quality control': identification of compounds according to their fingerprint spectrum
  - also inorganic materials, e.g. metal oxides
  - ex situ, but also after degassing in cell (vacuum)
- Identification of surface sites Detailed characterization of surface
  - use of molecular probes
  - in situ experiments, controlled dosage of probe
- Identification of surface sites under reaction conditions
  - in situ/operando experiments to obtain molecular reaction mechanism, exposure to reaction conditions

# **Vibrational spectroscopy**

- Interaction with matter
  - energy causes vibration of molecular bonds
  - energy is absorbed in correspondence of vibrational modes
  - an absorption band is generated



#### **Vibrations**



# **Vibrations**

• Why do vibrations appear in the IR spectrum?

selection rule  $\left(\frac{\partial \mu}{\partial O}\right) \neq 0$ 



molecular dipole moment  $\mu$  must change due to vibration or rotation along its coordinate (so called, normal mode or normal coordinate, **Q**)

# $H_2O$

#### N=3, non-linear, 3 fundamental modes



# Gas and liquid phase H<sub>2</sub>O



# **Vibrations**

- Harmonic oscillator
  - The stretching frequency of a bond can be approximated by Hooke's law. Two atoms and the connecting bond are treated as a harmonic oscillator composed of two masses (atoms) joined by a spring.



#### **Vibrations**





# The spectrometer



# **Dispersive vs. FT**

FT-IR spectrometer has significant advantages over dispersive one

#### Multiplex (Fellgett) advantage

All source wavelengths are measured simultaneously

#### Throughput (Jacquinot) advantage

For the same resolution, the energy throughput in an interferometer can be higher  $\rightarrow$  the same S/N as a dispersive-IR in a much shorter time

#### Precision (Connes) advantage

The wavenumber scale of an interferometer is derived from a HeNe laser that acts as an internal reference for each scan

### The IR spectrum



#### The IR spectrum



#### The IR spectrum



## The background

DRIFT spectra of V-W-TiO<sub>2</sub> catalysts after adsorption of NH<sub>3</sub>
 – aspect of spectra changes with background



- The spectrum contains information on – terminal O-H bonds | 3800-3600 cm<sup>-1</sup>
  - bridge hydroxyls Brønsted acidity
  - H-bonded hydroxyls
  - M-O and M=O bonds, bulk and surface
    fundamental (n) and overtone (2×n) modes
  - other groups, e.g. C-H, carbonates, carboxylates...





Busca, in Metal Oxide Catalysis, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim (2009) 95 Uslamin, Zeolite-based catalysis for sustainable aromatics production, Technische Universiteit Eindhoven (2019)



Busca, in Metal Oxide Catalysis, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim (2009) 95

• Also other disciplines...



Moros et al., Anal. Bioanal. Chem. 385 (2006) 708

# **Adsorbates by FTIR**



Morterra, Catal. Today 27 (1996) 497

# **Adsorbates by FTIR**



#### **Techniques, sample form, environment**



• Lambert-Beer law



**T**: transmittance, **A**: absorbance, ε: molar absorption (extinction) coefficient, **c**: concentration, **d**: path length

#### Solid samples

Large solid particles generally absorb too much IR light, therefore particles should be small and also special preparations are often necessary.

Most popular sample preparation methods (for mid-IR):

#### Alkali halide disk method

- Typically solid samples are diluted in KBr and ground
- Then pressurized to form a disk

#### Mull method

- Most common one is Nujol (liquid paraffin)
- Samples are ground and suspended in one or two drops of a mulling agent
- Followed by further grinding until a smooth paste is obtained

#### Film method

By solvent casting or melt casting

# **NOT** FOR IN SITU/OPERANDO EXPERIMENTS

- Sample preparation
  - self supporting wafers (few mg)
- · Controlled exp. conditions
  - vacuum and controlled dosages
- Baseline
  - slope increases at high frequency (beam scattering increases with increasing frequency)
  - slope depends on particle size (very steep for powders with large particles, ca. 1 µm)
  - T @ 4000 cm<sup>-1</sup> is ca. 0 for large particle size oxides



Busca, in Metal Oxide Catalysis, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim (2009) 95

• Quantification Molar absorption coefficient  $\epsilon$ 







- $\epsilon$ , integrated molar absorption coefficient
- *l*, disc thickness (optical path)
- n, amount of adsorbed molecule
- S, disc area

### **Diffuse reflection**



Chalmers and Dent, Industrial analysis with vibrational spectroscopy, 1997, 153

# **Diffuse reflection**

- Qualitative analysis
  - very sensitive to surface species due to its diffuse reflective nature
  - the detected light is reflected multiple times at powder surfaces
- Quantitative analysis
  - can be very complicated
  - the spectra are largely influenced by various experimental parameters, e.g. particles shape and size, refractive index of particles, absorption characteristics of particles, and porosity of the powder bed
  - a popular method is to use the Kubelka-Munk (K-M) function to transform reflectance to a sort of absorbance (K-M) unit
  - solid (approximated) theory
  - applicability and accuracy for highly absorbing and non-absorbing samples is questionable

# **Kubelka-Munk function**

• Infinitely thick medium

K/S =  $(1-R_{\infty})^2/2R_{\infty}$ K, absorption coeff. S, scattering coeff.

Adsorbate on infinitely thick medium

 $(K+\epsilon C)/S = (1-R)^2/2R$ 



 $F(R) = J(1/R-R_{\infty}) = 2\varepsilon C/S$ 

**Kubelka-Munk function** 

optical length (d in L-B Law) much larger than in transmission  $\rightarrow$  more sensitivity

Matyshak et al., Catal. Today 25 (1995) 1

#### **Diffuse reflection**



Sirita et al., Anal. Chem. 79 (2007) 3912

# **Diffuse reflection vs Transmission**

- Advantages of diffuse reflection
  - easier sampling
  - applicability to powders that scatter too much in transmission, assuming the surface area is sufficiently high to detect surface
  - vibrations with a sufficiently high signal-to-noise ratio
  - slightly lower sensitivity to bulk conduction phenomena, because of a higher surface-to-bulk sensitivity ratio
  - ideally suited for in situ/operando studies
- Disadvantages of diffuse reflection
  - less obvious optical setup
  - work in flow rather than in vacuum
  - more difficult sample activation (i.e. water removal from highly porous materials)

#### **Diffuse reflection vs Transmission**

- Comparison between techniques with different sensitivity (bulk/surface) should be careful
- DRIFTS more sensitive than TIRS
- Band assignment depends on surface sensitivity of the technique



Rödel et al., PCCP 10 (2008) 6190

### **Diffuse reflection**

• Typical mirror unit, sample holder and in situ cell




IRE, internal reflection element; high refractive index material





• Stable films needed for in situ investigations



# **Materials for internal reflection elements**

Material	Useful range / cm <sup>-1</sup>	n	$d_{ m p}$	Properties
ZnSe	20000-700	2.43	long	soluble in strong acid; usable up to ca. 300°C
Ge	5000-900	4.02	short	good chemical resistance; hard and brittle; becomes opaque at 250°C
Si	9400-1500; 350-FIR	3.42	short	excellent chemical resistance; hard; usable up to ca. 300°C
KRS-5 (Thallium bromoiodide)	14000-330	2.45	long	toxic; slightly soluble in water and soluble in base; usable up to ca. 200°C





- Solid (powders, crystals, foils, plates, seeds...) and liquid samples
- Ex situ
  - structure assignment
  - identification
  - quality control







manual mirrors • Sample compartment 0 00 员 cell holder liquid out to/from thermostat liquid in





ethyl cyanoacetate

ethyl  $\alpha$ -cyanocinnamate

- γ-aminopropyl modified SiO<sub>2</sub> (APS-SiO<sub>2</sub>)
   1.5 mmol/g NH<sub>2</sub>
   202 m<sup>2</sup>/g
   deposited on ZnSe from
   CCl<sub>4</sub> slurry
   toluene/PE slurry prep. 80°C

  - dried in *vacuum*
- toluene, 60°C





toluene, 60°C, 20 mM



toluene, 60°C, 20 mM

- Consecutive dosage of reactants
- Time dependence





Wirz et al., Langmuir 22 (2006) 3698



toluene, 60°C, 20 mM



Wirz et al., Langmuir 22 (2006) 3698





metal donates electrons back to the anti-bonding  $\pi$  orbital of CO

- Low CO coverage: v<sub>CO</sub> depends on the geometry of adsorption site (face order: terrace – corner – edge) – BD is strong
- High CO coverage: v<sub>CO</sub> depends on dipoledipole interactions – BD is weak



- Model studies Surface science
  - stainless steel UHV setup with flanges, pumps, pressure gauges, etc.
  - 10<sup>-10</sup> to 10<sup>-11</sup> mbar base pressure
  - tools and components for preparation, characterization, sample manipulation, resistive

heating





MS

XPS

etecto

**FTIR** 

manipulator

LEED





Szanyi et al., J. Vac. Sci. Technol. A 11 (1993) 1969

• Powders



- the larger the particles, the less CO adsorbs (intensity)
- the larger the particles, the less the available defects (nr. of signals)

How does the CO stretching frequency shift when a Pt surface is covered with hydrogen or oxygen?







Lear et al., J. Chem. Phys. 123 82005) 174706





Y. Soma-Noto et al., J. Catal. 32 (1974) 315

#### The surface selection rule



Fig. 1. (a) The lines of force and the electrical "image" resulting from a positive charge over the surface of a conductor (the metal surface is the upper line above the hatched area). (b) The changes during the vibration of a dipole parallel to the surface of the metal; the "image" dipole change is in the opposite direction to the original. (c) The changes during the vibration of a dipole perpendicular to the surface; the "image" dipole change is in the same direction as the original.

Pearce and Sheppard, Surf. Sci. 59 (1976) 205

#### The surface selection rule

Carboxylate groups



Greenler et al. Surf. Sci. 118 (1982) 415

#### **Reflection-absorption (IRRAS)**



Haq et al., J. Phys. Chem. 100 (1996) 16957; Preuss et al., Phys. Rev. B 73 (2006) 155413

#### **Reflection-absorption (IRRAS)**



Haq et al., J. Phys. Chem. 100 (1996) 16957

# **Reflection-absorption (IRRAS)**

• Adsorption of ethylene



Vibrational assignments of ethylidyne

Mode	$Pt(111) (cm^{-1})$	Co <sub>3</sub> (CO) <sub>9</sub> CCH <sub>3</sub>	
	FT-RAIRS [this work]	EELS	(cm <sup>-1</sup> ) [8]
		[5]	
$\overline{\nu_{as}(CH_3)}$	Not allowed	2950 (impact)	2924
$\nu_{\rm s}(\rm CH_3)$	2884	2895	2882
$\delta_{as}(CH_3)$	Not allowed	1420 (impact)	1432
$\delta_{s}(CH_{3})$	1341	1350	1359
VC-C	1124	1130	1161
$\rho(CH_3)$	Not allowed	980 (impact)	1006
$\nu_{\rm C-Pt}$	Below detector cut off	435	~ 600

Chesters et al., Surf. Sci. 187 (1987) L639

# **Phase-modulation IRRAS (PM-IRRAS)**

- Generation of 2 polarizations (photoelastic modulator)
  - excellent gas-phase compensation
  - non-UHV experiments pssible
  - highly sensitive, time-resolved studies possible



# **Acid sites**

- Quality and quantity of acid sites Criteria
  - unequivocal analysis of intermolecular interaction
  - selective interaction with acidic or basic sites
  - sufficient accuracy in frequency shift determination
  - high (and available) extinction coefficients of adsorbed probe
  - appropriate acid (base) strength to induce interaction
  - high specificity (allow discrimination between sites with different strength) Use different molecules !
  - small molecular size Use different molecules !
    - pyridine (smaller channels) and picoline (larger channels or surface only)
  - low reactivity under exp. conditions
- Examples
- acidity of zeolites with different channel sizes
- acid sites located in all channels

#### **Probe molecules**

• Adsorption of NH<sub>3</sub>



#### **Probe molecules**

• Adsorption of NH<sub>3</sub>




• Adsorption of NH<sub>3</sub>  $-V_2O_5/WO_3$ -TiO<sub>2</sub>



 Probe molecules for the study of localization of active sites in microporous materials





Liu et al., J. Phys. Chem. C 121 (2017) 23520