# Infrared spectroscopy

Instrumentation Techniques Examples

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### **Double beam spectrometer**



## **Fourier transform 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

# **Sampling techniques**



**TIRS**: transmission infrared spectroscopy

**IRES**: infrared emission spectroscopy

**PA-IRS**: photoacoustic infrared spectroscopy

**VCD**: vibrational circular dichroism

**ATR-IRS**: attenuated total reflection infrared spectroscopy

**IRRAS**: infrared reflectionabsorption spectroscopy

**PM-IRRAS**: polarizationmodulation IRRAS

**DRIFTS**: diffuse reflectance infrared Fourier transform spectroscopy

Straight' IR light absorption



- Popular for detections of gas and liquid samples
- Solids have to be diluted or shaped in a very thin film
- Quantification is more straightforward than other IR techniques

### In heterogeneous catalysis

- Popular for *in situ* investigations
- Typically a very thin self-supporting catalyst disk is used
- Powder sample dispersed on transparent grid (W)
- Mass transfer can be an issue



IR transparent window

### Gas samples









Temperaturecontrolled cell [ca. 200 – 500 K]



#### In situ cells for heterogeneous catalysis studies



FIG. 1. In situ high-pressure and temperature IR cell. 1:  $CaF_2$  window; 2: KBr rod (1); 3: KBr rod (2); 4: catalyst disk; 5: Viton O-ring; 6: cooling water inlet; 7: cooling water outlet; 8: gas inlet; 9: gas outlet; 10: thermocouple; 11: swagelok fitting; 12: heater; 13: asbestos molding; 14: Teflon<sup>®</sup> window holder.

#### Arakawa et al. Appl. Spec., 40, 884 (1986)

heating up to 500°C pressure up to 10 MPa



FIG. 1. The high-temperature IR cell: 1, gas inlet; 2, electrical connection; 3, insulating material; 4, Viton O-ring; 5, thermocouple; 6, gas outlet; 7, heating element; 8, ceramic; 9, nut and bolt; 10, Vycor tubing; 11,  $CaF_2$  window; 12,  $Al_2O_3$  tubing; 13, sample; 14, stainless steel sample holder.



Mawhinney et al., Langmuir 15 (1999) 4617; Basu et al., Rev. Sci. Instrum. 59 (1988) 1321

In situ cells



Rasmussen et al., PCCP 14 (2012) 2171

### 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

Quantification: most straightforward than other techniques

Lambert-Beer law



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

### Common window materials

Material	Useful range / cm <sup>-1</sup>	Refractive index	Properties
NaCl	40'000-600	1.52	Soluble in water; slightly soluble in alcohol; low cost
KBr	43'500-400	1.54	Soluble in water; slightly soluble in alcohol; hygroscopic
CaF <sub>2</sub>	77'000-900	1.40	Insoluble in water; chemically resistant; mechanically strong
BaF <sub>2</sub>	66'666-800	1.45	Low water solubility; soluble in acids and NH <sub>4</sub> Cl
CsI	42'000-200	1.74	Soluble in water and alcohol; hygroscopic
ZnSe	20'000-500	2.43	Soluble in strong acid

## **Reflection based techniques**

### Aim for heterogeneous catalysis studies

study events occurring at interfaces and maximize signals related to catalysts and active species on surfaces, especially during reactions



## **Specular vs. diffuse reflection**



Specular reflection (smooth surface)

Surface smoothness like mirror = reflection and incident angles are equal



Diffuse reflection (rough surface)

Incident light is reflected to a number of directions due to surface roughness

Very popular for *in situ* measurements of physicochemical processes at **gas**solid interfaces using realistic **powder catalysts** 





The praying mantis (very popular, highly efficient light collection)

In situ DRIFTS cell

#### qualitative analysis

Very sensitive to surface species due to the diffuse reflective nature of the method; the detected light can be multiply reflected at powder surfaces

#### quantitative analysis

It can be very complicated; the spectra are largely influenced by a number of experimental parameters such as shape and size of particles, refractive index of particles, absorption characteristics of particles, and porosity of the powder bed

A popular quantitative analysis method is using Kubelka-Munk (K-M) function to transform reflectance to a kind of absorbance (K-M) unit

Kubelka-Munk function  
$$f(R) = \frac{(1-R)^2}{2R} = \frac{k}{s}$$

*k*: molar absorption coefficient, *k*=2.303 $\alpha$ c,  $\alpha$ : absorptivity, *c*: concentration,  $\sigma$ : diffusion (scattering) coefficient

There is a solid (approximated) theory behind and the K-M function is widely used; however its applicability and accuracy for highly absorbing and non-absorbing samples is questionable

recent discussion on this topic: Sirita et al., Anal. Chem. 79 (2007) 3912



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



Meunier, Chem. Soc. Rev. 39 (2010) 4602

### Model system investigation

- single crystals
- well-defined nano-particles



### 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.

## The surface selection rule

Carboxylate groups



## The surface selection rule

### Also valid for small (nm) metal particles?



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

### Also RAIRS; specular/external reflection method



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



### Adsorption of ethylene



√/cm-1

Vibrational assignments of ethylidyne

Mode	$Pt(111) (cm^{-1})$		Co <sub>3</sub> (CO) <sub>9</sub> CCH <sub>3</sub> (cm <sup>-1</sup> ) [8]
	FT-RAIRS [this work]	EELS [5]	
$\nu_{\rm s}(\rm CH_3)$	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
$\nu_{C-Pt}$	Below detector cut off	435	~ 600

### Also RAIRS; specular/external reflection method



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

### Generation of 2 polarizations (photoelastic modulator)

- excellent gas-phase compensation
- non-UHV experiments possible
- highly sensitive, time-resolved studies possible



### Attenuated total reflection infrared spectroscopy

The most rapidly developing IR method over the last years in bioscience, organic-, inorganic chemistry, and catalysis



occurs

#### **Total internal reflection** (attenuated total reflection)





Calculate the critical angle at ZnSe/air interface n(ZnSe) = 2.4, n(air) = 1.0

### How does it work?

Light travels through a waveguide



Assuming that  $n_1$  and  $n_2$  are constant, how much difference do you expect in the penetration depth at 400 and 4000 cm<sup>-1</sup>?



 $\theta$  : angle of incidence

$$\lambda_1 = \frac{\lambda}{n_1} \qquad n_{21} = \frac{n_2}{n_1}$$

 $d_p$ : penetration depth; defined as the distance from interface where the electric field has decayed to 1/e of its value E<sub>0</sub> at the interface

Very powerful method for investigations of (catalytic) solid-liquid interfaces

### Common window materials

Material	Useful range / cm <sup>-1</sup>	Refractive index	Properties
ZnSe	20 000-700	2.43	Soluble in strong acid; usable up to ca. 573 K
Ge	5000-900	4.02	Good chemical resistance; hard and brittle; becomes opaque at 400 K
Si	9400-1500; 350-FIR	3.42	Excellent chemical resistance; hard; usable up to ca. 573 K
KRS-5 (Thallium bromoiodide)	14 000-330	2.45	Toxic; slightly soluble in water and soluble in base; usable up to ca. 473 K

### Stable films needed for in situ investigations



#### Quantification

Similar to Lambert-Beer law but replacing d by wavelength dependent thickness, i.e. effective thickness  $d_e$  (Harrick)

$$A = -\log(T) = -\log(\frac{I}{I_0}) = \varepsilon c d_e$$
$$d_e = \frac{n_{21} E_0^2 d_p}{2\cos\theta}$$

### Cells







thermostatting plates







cyclohexane, 50° C, Pd/Al<sub>2</sub>O<sub>3</sub>



Ferri et al. J. Phys. Chem. B 110 (2006) 22982



# **Comparison of techniques**



- Comparison between techniques with different sensitivity (bulk/surface) should be careful
- Band assignment depends on surface sensitivity of the technique
- PM-IRRAS suitable for investigation of powder samples...



## **Adsorbed molecules**

Orientation on surfaces

#### **Powders**

qualitative

 adsorption mode, coordination to surface (e.g., mono-, bidentate, bridging, tilted...)

#### Metallic surfaces (e.g. single crystals)

- more accurate
- surface selection rule
- orientation information from dynamic dipole moment direction
- group theory
- combination with theory (Density Functional Theory DFT)

# **IR spectroscopy for catalysis**

### Ex situ experiments

structure determination (M-O bonds, OH groups), nature of adsorbates

#### In situ experiments

- structure determination (M-O bonds, OH groups), nature of adsorbates
- determination of adsorption sites (probe molecules)
- determination of acidity and basicity (probe molecules)
- typically, vacuum experiments and low T (LN<sub>2</sub>)
- other atmospheres and T possible

### Operando experiments

Combination of

- spectroscopic measurement (not only IR!)
- simultaneous detection of reactants and products (MS, GC, ...)
- Evolution of adsorbates with changes in reaction conditions (e.g. T)
- simulation of reactor studies (pay attention to reactor design issue)
- adsorption-desorption experiments under relevant conditions (not vacuum)

- 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 Hard–Soft classification of sites and probes
- high specificity (allow discrimination between sites with different strength) Use different molecules !
- small molecular size Use different molecules !
- Iow reactivity under exp. Conditions
- **.**..
- Example
  - acidity of zeolite with different channel sizes
  - acid sites located in all channels
  - use of pyridine (smaller channels) and picoline (larger channels or surface only)





 $SiO_2-AI_2O_3 > AI_2O_3 > SiO_2$ ; next issue: coordination environment of acid site

## Molar absorption coefficient of adsorbates



- *l*, disc thickness (optical path)
- n, amount of adsorbed molecule
- S, disc area

 $H_3C$ 

### Molar absorption coefficient of adsorbates



 $\epsilon_{1605} = 1.9$ 

nd: not determined;  $\varepsilon = cm \mu mol^{-1}$ 

Onfroy et al., Micropor. Mesopor. Mater. 82 (2005) 99

### Indirect characterization of supported metal oxides



### Carbon monoxide (CO)

• Widely used as a *sensor* to investigate the electronic state of catalytic active sites

![](_page_47_Figure_3.jpeg)

![](_page_47_Figure_4.jpeg)

### Carbon monoxide (CO)

![](_page_48_Figure_2.jpeg)

![](_page_49_Figure_1.jpeg)

The larger the particles, the less CO adsorbs (intensity)

The larger the particles, the less defects available (nr. of signals)

![](_page_50_Figure_1.jpeg)

Winkler et al., Appl. Catal. B 93 (2009) 177; Matam et al., Appl. Catal. B 129 (2013) 214

### Carbon monoxide (CO)

![](_page_51_Figure_2.jpeg)

### Carbon monoxide (CO)

Q

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

![](_page_52_Figure_3.jpeg)

![](_page_53_Figure_1.jpeg)

Pd/Ag alloy on SiO<sub>2</sub>

Y. Soma-Noto, W.M.H. Sachtler, J. Catal. 32 (1974) 315

![](_page_54_Figure_1.jpeg)