5.12 Vibrational Surface Spectroscopy

One of the most powerful methods for surface chemical compositional analysis - produces vibrational spectrum of adsorbed species. Can:

- fingerprint adsorbed species
- use selection rules to deduce bonding orientation

Based on absorption of energy from incident particle (electron - EELS, IR photon - RAIRS) - (won't discuss Raman or SHG)

- measure intensity vs. energy distribution of particle following absorption

---

**Fig. 1.5.** Schematic illustration of infrared absorption. The HCl molecule whose vibrational frequency is $8.67 \times 10^{13}$ Hz increases its vibrational energy by absorbing the energy of an infrared photon which has this same frequency.

---

**Fig. 1.6.** Forces generated on a dipole by an oscillating electric field. These forces tend to alternately increase and decrease the dipole spacing.
5.12.1 Introduction to IR Methods

Can be performed using conventional (non-dispersive) FTIR instrument

- sample all channels simultaneously (Jaquinot advantage)
- high signal level at detector (Fellgett advantage)
- high accuracy (use of accurate internal standard HeNe laser)

in transmission or reflectance mode

In transmission mode

\[ I = I_0 \exp(k \cdot c \cdot l) \quad \frac{I}{I_0} = T = \exp(k \cdot c \cdot l) \quad \text{Transmittance} \]

\[ \ln \frac{I}{I_0} = k \cdot c \cdot l \quad A = \varepsilon \cdot c \cdot l \quad \text{Absorbance} \]

where \( k \) is the absorption coefficient, \( \varepsilon \) is absorptivity, \( c \) is surface concentration and \( l \) is pathlength

It can be shown that \( k \) is also the imaginary part of the refractive index

\[ n = n + ik \]

In a non-absorbing medium (dielectric), refractive index \( n = n \). In absorbing medium, refractive index contains a portion corresponding to absorption \( k \).

Transmission experiments must be performed twice and difference calculated:

- reference spectrum with no absorbed material
- sample spectrum with adsorbed material

Surface transmission experiments less common

- large bulk contribution unless sample very thin
- many interesting samples not sufficiently transparent to IR
5.12.2 Reflection, Refraction and Absorption

Reflection, refraction and absorption all related through Maxwell's equations for electromagnetic radiation for interface of different $n$.

The reflection and refraction angles are given by Snell's Law:

\[
\Theta_I = \Theta_R \quad \text{and} \quad \frac{n_1}{n_2} = \frac{\sin \Theta_I}{\sin \Theta_T}
\]

for $\Theta_I < \Theta_{\text{crit}} = \sin^{-1}\left[\frac{n_2}{n_1}\right]$

Where the plane containing the incident and reflected (and transmitted) waves is called the plane of incidence.

The intensities of the reflected waves are given by Fresnel equations and are dependant on their polarizations:

When $n_1 = 1.00$ (vacuum/air) and $n_2 = n + ik$:

For $p$-polarized light (E-field $\parallel$ to plane of incidence)

\[
R_p = \frac{\cos^2 \Theta_T - 2n \cos \Theta_I \cos \Theta_T + \left(n^2 + k^2\right) \cos^2 \Theta_I}{\cos^2 \Theta_I + 2n \cos \Theta_I \cos \Theta_T + \left(n^2 + k^2\right) \cos^2 \Theta_I}
\]

and for $s$-polarized light (E-field $\perp$ to plane of incidence)

\[
R_p = \frac{\cos^2 \Theta_I - 2n \cos \Theta_I \cos \Theta_T + \left(n^2 + k^2\right) \cos^2 \Theta_2}{\cos^2 \Theta_I + 2n \cos \Theta_I \cos \Theta_T + \left(n^2 + k^2\right) \cos^2 \Theta_2}
\]

Such equations allow us to predict the reflectivity of materials ($n_2$) with incidence angle
For very thick films on reflective surface (metal) $\Theta_I$ can be normal (0°) - double-pass transmission experiment

For very thin films on reflective surface $\Theta_I$ should be large (~75-85°) - grazing incidence experiment (RAIRS)

5.13 Reflection-Absorption Infrared Spectroscopy (RAIRS)

also known as Infrared Absorption-Reflection Spectroscopy (IRAS)

The intensity of light reflected from a surface (Fresnel equations) related to phase shift $\delta$ upon reflection

- $s$-polarized light almost cancelled by reflection at grazing incidence
- $p$-polarized light almost doubled by reflection at grazing incidence (until very grazing situations)

![Figure 7.10](image)

Figure 7.10: The phase shift for light reflected from a metal surface as calculated for light polarised both parallel to ($p$) and perpendicular to ($s$) the plane of incidence, after Greenler [9]
Only $p$-polarized component light can be reflected from surface at high incidence angles.

Only vibrations with component dynamic dipole moment aligned $\perp$ to surface plane can interact with ($p$-polarized) incident light - "surface selection rule".

Group theory tells us which vibrations have components in x, y and z directions.

Since absorbance $A$ is proportional to $E^2$ and area of surface sampled increases as $1/\cos \Theta_I$, sensitivity of RAIRS increases as $E^2/\cos \Theta_I$. 

**Fig. 1.7.** Dipole moment changes in certain molecular vibrations.
Best sensitivity for observing an absorption feature in RAIRS is

- \( p \)-polarized light
- grazing incidence
- molecule with transition dipole arranged along surface normal
- molecule with large transition moment
Using high quality FTIR spectrometers spectral acquisition (reference and sample) with high S/N can be acquired in few minutes with 2-4 cm\(^{-1}\) resolution.

Width of absorption peak usually related to inhomogeneity of surface.

Fig. 7.12. A schematic representation of the FT-RAIRS experiment as described by Chesters [10].

Fig. 9.7 Comparison of spectrum of CO on a polycrystalline Cu film with band shapes and positions found on individual single crystal planes (Pritchard, 1979b).
Frequency of internal vibrational mode often close to gas phase (red-shifted 200-800 cm$^{-1}$)

Frequency used to infer nature of bonding with surface
- $\nu$(X-Y) decreases as coordination to surface atoms increases
- occupation of sites often varies with coverage - high symmetry/coordination at low coverages $\rightarrow$ low symmetry/coordination at high coverages
- compared with organometallic compounds

"Softened" modes (by 100-200 cm$^{-1}$) sometimes appear due to substantial interactions with surface

$\Delta R/R$ often very small - absorbance 0.1 to $10^{-3}$!
- sufficient for submonolayer sensitivity for molecule with strong dynamic dipole moment

$\Delta R/R$ approximately linear with coverage (unless molecular orientation change)

BUT relative intensity of band not good indicator of population
- dynamic dipoles of molecules in ordered array couple - modulates dynamic dipole moment and hence modulates absorption
- orientation affects absorbance

Cannot observe vibrations with $\nu < 800$ cm$^{-1}$ - detector cut-off

5.13.1 Example Application of RAIRS: CO/NO on Metals

CO and NO have similar electronic structure
- HOMO is filled 5 $\sigma$ and LUMO is 2 $\pi^*$ MO
Both considered $\sigma$ donors and $\pi^*$ acceptors

Coadsorption of CO and NO on Rh/Pt/Ni extremely important in catalysis (vehicle emissions)

- CO most intensively studied adsorbate by vibrational spectroscopy

<table>
<thead>
<tr>
<th>Wavenumbers (cm(^{-1}))</th>
<th>1600</th>
<th>1700</th>
<th>1800</th>
<th>1900</th>
<th>2000</th>
<th>2100</th>
<th>2200</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hollow Bridge Top CO(g)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni(100) c(2x2)-CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu(100) c(2x2)-CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pd(100) (2\sqrt{2}x\sqrt{2})R45-2CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ru(0001) (√3x√3)R30-CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rh(111) (√3x√3)R30-CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rh(111) (2x2)-3CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rh(111) c(2\sqrt{3}x4)rect-C(_6)H(_6)+CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pt(111) c(4x2)-2CO</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$\nu$(CO(g)) = 2140 \text{ cm}^{-1}$

CO on Ni(111):
- 2-fold bridge at $\sim$1840 cm\(^{-1}\) at low coverage
- additional on-top sites at $\sim$2040 cm\(^{-1}\) at coverages $>0.5$ ML

NO on Ni(111):
- tilted 2-fold bridge at $\sim$1480 cm\(^{-1}\) at very low coverages
- shifts to more perpendicular 2-fold bridge up to $\sim$1580 cm\(^{-1}\) at low coverages
When NO is added to CO covered surface:

- on-top sites appear - CO displaced by NO
- bridge CO shifts to higher $\nu$ - strengthened CO internal bond
- bridge CO broadens into doublet - multiple adsorption sites?

(Recent results have suggested that this interpretation is not correct - CO and NO on Ni(111) occupy fcc and hcp sites at all coverages!)
5.14 Electron Energy Loss Spectroscopy (EELS)
also known as high resolution EELS (HREELS)

Based on inelastic scattering of monoenergetic beam of low energy electrons ($E_i = 1$-$10$ eV) from surface

- kinetic energy analysis of inelastically-scattered electrons gives information about energy transfer to surface vibrational modes

- $E_s = E_i - h\nu \leftarrow$ energy loss

$\text{H}_2$ adsorbed on TM expect $h\nu \approx 4560$ cm$^{-1}$ but for H on TM expect $h\nu \approx 800$-$1000$ cm$^{-1}$

- minimum resolution $\Delta E_i$ (FWHM of incident beam) of EELS > RAIRS - best $< 8$ cm$^{-1}$, typically 20-$30$ cm$^{-1}$ ($8$ cm$^{-1} \approx 1$ meV)

![Diagram of EELS and RAIRS](image)

**Figure 7.13.** RAIRS spectrum of a monolayer of CO adsorbed on a Cu (111) surface at 95 K in comparison with analogous data obtained using electron energy loss spectroscopy, after Chesters et al. [10]
- sufficient for submonolayer sensitivity, even for weak dynamic dipoles (best $10^{-3}$ ML)

5.14.1 Electron Scattering Mechanisms in EELS

Three basic scattering modes for electron, each with own selection rules:

(1) **Dipole scattering**

(2) **Impact scattering**

(3) **Negative ion resonance** scattering

- gives EELS added versatility over RAIRS
- all three modes operative (one usually dominates under certain experimental conditions)

(1) Dipole scattering

Electron is scattered at long range (100's Å) by oscillating Coulombic field of surface

Electron momentum $\perp$ to surface is conserved

Energy loss results in small angle deflections around specular direction - dipole scattering lobe

- Elastically-scattered electrons appear at $\Theta_i = \Theta_s$ (specular)
- Inelastically-scattered electrons appear at $\Theta_i \approx \Theta_s$ (sharply peaked near specular)
small momentum loss $q_{||} = h\nu$

Only modes with component of vibration perpendicular to surface can be excited (not parallel modes) - *surface selection rule*

same selection rules as RAIRS

Total dipole = 2\(\mu\)

Large cross-section for small $E_i$'s (<20 eV) - dominant mechanism near specular

Well-developed/straightforward theory

(2) Impact scattering:

Electron is "kinematically" scattered at short range (few Å) by atomic core potentials of surface - modulated by nuclear motion (vibration)

Electron scattered over wide range of angles (in and out of plane of incidence)
Probability of scattering depends on surface dipole amplitude/direction (symmetry of mode) and electron energy (momentum)

Neither electron momentum || nor ⊥ conserved

- can see vibrational modes with dynamic dipoles both || and ⊥ to surface plane (only off-specular)

Much lower cross-section than dipole scattering (weak loss features)

Poorly-developed/complex theory

(3) Negative Ion Resonance

Incident electron becomes transiently trapped in empty, high lying MO of adsorbate - Rydberg state

Vibrational feature intensity strongly depends on incident energy - resonances

Very small cross-section off-resonance (rare)

Regardless of mechanism, most electrons scattered *elastically* - loss spectrum usually very weak

5.14.2 Instrumentation for HREELS
Monochromation of electrons emitted by thermionic emission achieved by passing through spherical-sector electron energy analyzer

- typical radius 5 cm, typical pass energy <500 meV (in XPS, radius ~20-30 cm, pass energy 50-100 eV)

Electrons retarded before entry into sector, then accelerated to sample

Second spherical-sector used to analyze electron kinetic energy

Entire instrument magnetically shielded - no magnetic materials for construction

In very high resolution instruments, two sectors for monochromator and two for analyzer
Space-charge effects always tend to produce divergent beam

Optimization of lens and sector potentials very difficult - computer controlled

In best HREELS, analyzer rotatable for off-specular (impact scattering) analysis

E_i variable 1-50 eV or more - can study vibrational and electronic loss processes (plasmons)

- observe electronic excitation from 0->50 eV
- observe vibrational modes from 0->4000 cm\(^{-1}\)

![Diagram](image)

Fig. 9.11 Set of imaginary adsorption states showing the expected loss peaks associated with each structure.
5.14.3 Example Application of HREELS: CO adsorption on W(100)

Fig. 9.13 Electron energy loss spectra of CO on W[100] at 300 K. The spectra are recorded with a primary electron beam energy of 5 eV incident at an angle of 75°; $\Delta \phi$ is the measured increase in work function (Froitzheim et al., 1976).

(gas exposure units 1 L = 1 Langmuir = $10^{-6}$ torr·s)

At low CO exposures
- 70 meV (565 cm$^{-1}$) W-C (hollow site)
- 78 meV (630 cm$^{-1}$) W-O (hollow site)

At high $p_{CO}$
- 45 meV (363 cm$^{-1}$) W-CO (on-top)
- 258 meV (2081 cm$^{-1}$) -C≡O (CO(g) = 2140 cm$^{-1}$)
5.14.4 Example Application of HREELS: Ethylene adsorption on Rh(111)

Ethylene is an important starting material in many industrial catalytic processes

How does it bond to transition metal catalyst surface?

\[ \begin{align*}
\pi \text{-bonded} & \\
\text{di } \sigma \text{-bonded}
\end{align*} \]

<table>
<thead>
<tr>
<th>Wavenumber (cm(^{-1}))</th>
<th>(\pi)-bonded</th>
<th>(\text{di } \sigma)-bonded</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\nu_s(\text{CH}_2))</td>
<td>3075-2990</td>
<td>3000-2910</td>
</tr>
<tr>
<td>(\nu(\text{C-C})/\text{wag(\text{CH}_2)})</td>
<td>1560-1500</td>
<td>1170-830</td>
</tr>
<tr>
<td>(\nu(\text{C-M}))</td>
<td>&lt;400</td>
<td>480-400</td>
</tr>
</tbody>
</table>

Low temp (<150 K)-

\(\pi\)-bonded: Cu(100), Pd(111), Pd(100), Pd(110)

\(\text{di } \sigma\)-bonded: Fe(111), Fe(110), Ni(111), Ni(100), Ru(001), Pd(100), Pt(111), Pt(100)

At 77 K, ethylene non-dissociatively chemisorbs on Rh(111) with C=C approximately parallel to surface in π-bonded configuration.
At 320 K-

- (i) $v(\text{C-C}) (1120 \text{ cm}^{-1})$ more like C-C than C=C
- (ii) appearance of CH$_3$-type stretches
- (iii) 1/4 of total hydrogen desorbed between 100 and 300 K

Internal rearrangement to produce C$_2$H$_3$ - ethylidyne

At higher temperatures, sequential dehydrogenation $\rightarrow$ H-terminated C polymers $\rightarrow$ graphite

Also observe adsorbate-induced reconstruction:

![Diagram](image.png)

**Figure 2.27.** Ethylidyne-chemisorption-induced restructuring of the Rh(111) crystal face. Note the expansion of metal atoms around the adsorption site.

Before HREELS (and LEED) analysis, ethylidyne structure never proposed
Acetylene + H₂ or ethylene adsorption at room temperature produce ethylidyne on many transition metal surfaces.
5.15 Summary

"Fingerprint technique" - can identify functional groups through vibrational frequency analysis

Use of selection rules/group theory allows determination of bond orientation

Difficult to quantify species (RAIRS easier than HREELS)

RAIRS:

Excellent energy resolution (<2 cm\(^{-1}\)) - useful for separating multiple peaks, phase transitions, lateral interactions, dynamics of coupling

Straightforward instrumentation

Not restricted to surfaces in vacuum - can be used in "real world" conditions from several atm to UHV

Theory well-developed

Rapid - 30 s-10 minutes - time-resolved spectra possible

BUT

Less sensitive than HREELS

Cannot observe modes <800 cm\(^{-1}\)

Only dipole active modes - component perpendicular to surface

Need background (reference spectrum) subtraction

HREELS:

Multiple scattering mechanisms allow observation of modes parallel \(\text{and}\) perpendicular to surface
Observe modes 0->4000 cm\(^{-1}\) - low frequency modes (molecule-surface) visible

Can perform vibrational and electronic loss spectroscopy

Can induce and probe electron-induced reactions with variable \(E_i\)

No background subtraction

Most common technique for surface and bulk phonon measurement

BUT

Poor resolution (6-8 cm\(^{-1}\) best)

Only works in UHV

Very expensive, complex, delicate instrumentation

Slow - 15-60 minutes per spectrum

Electron spectroscopy - surface charging on insulators

Difficult theory