



Marianne Liebi– Material Science at Large Scale Facilities

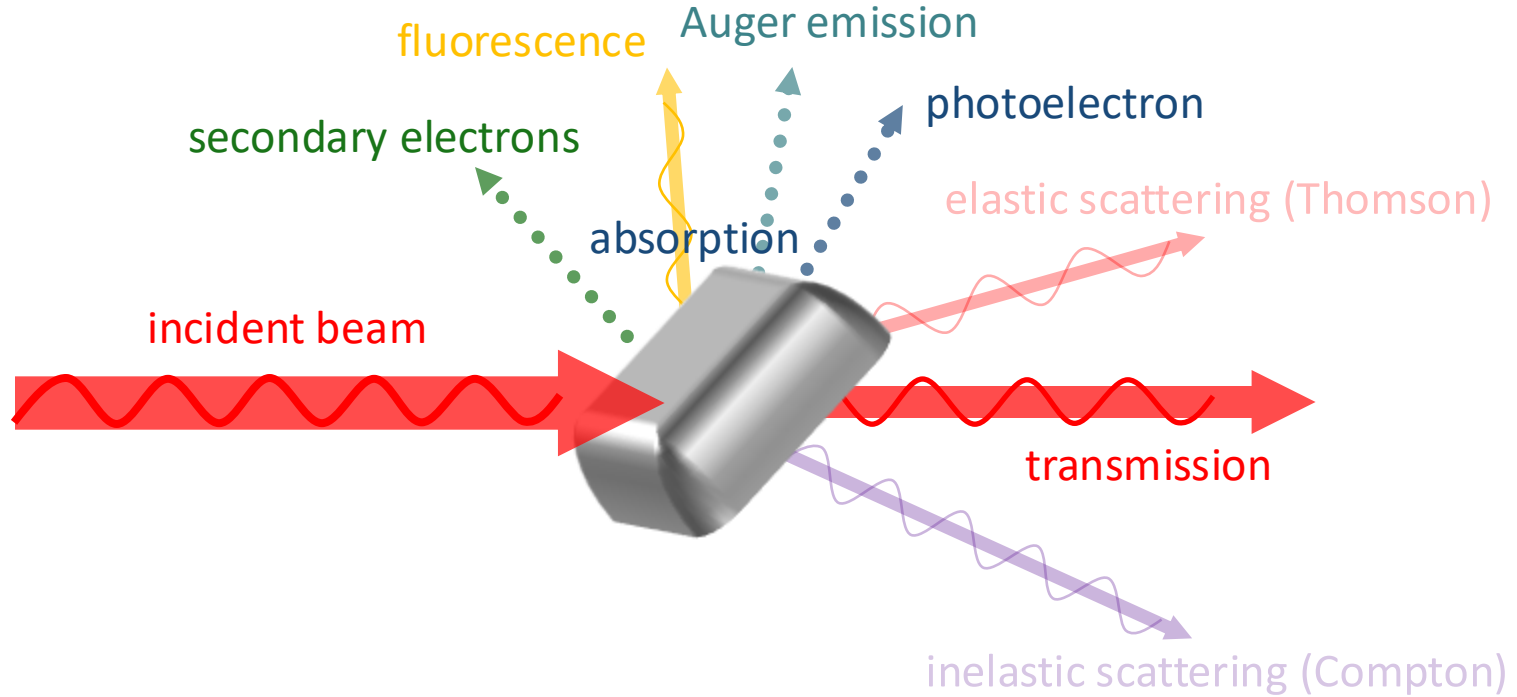
# X-ray Absorption Fine Structure(XAFS): XANES/NEXAFS and EXAFS

EPFL Master Course 2025 MSE435

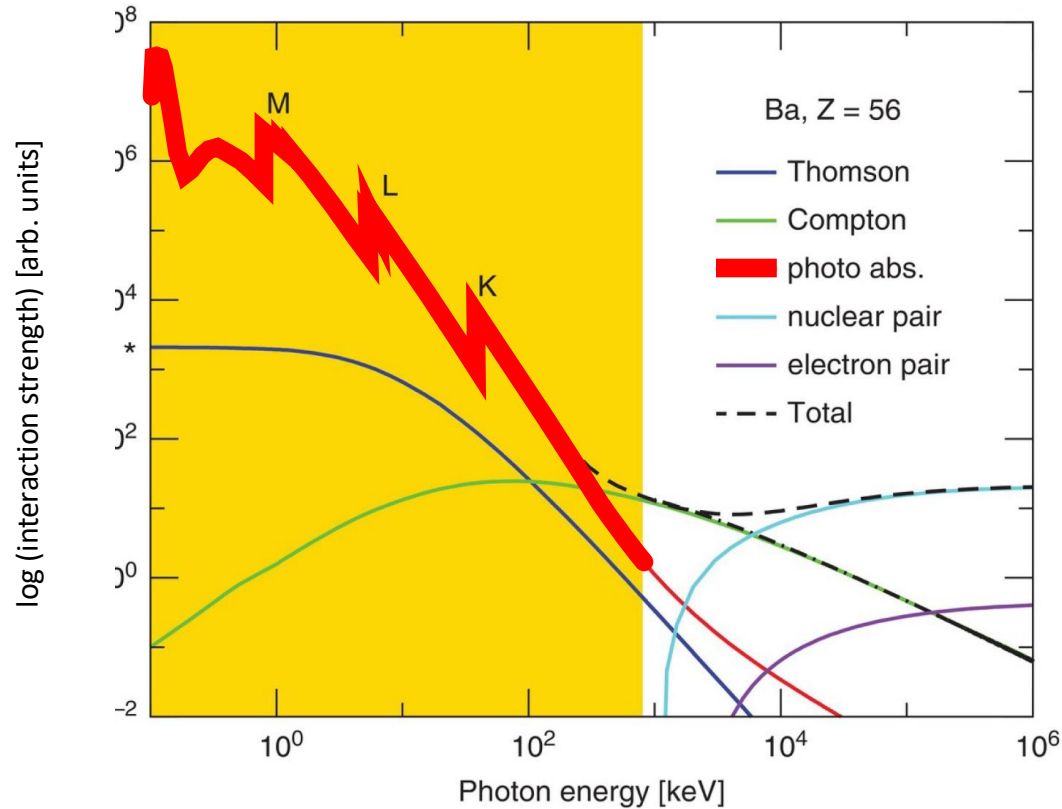
# Course program

<b>Dates</b>	<b>Content</b>	<b>Lecturer</b>
08.09.25	Introduction, sources, beamlines, detectors	Steven Van Petegem
15.09.25	Excursion to PSI	Marianne Liebi
22.09.25	Holiday	
29.09.25	Interaction with matter	Steven Van Petegem
06.10.25	Fluorescence	Marianne Liebi
13.10.25	X-ray absorption spectroscopy	Marianne Liebi
20.10.25	Break	
27.10.25	Diffraction I	Steven Van Petegem
03.11.25	Small angle scattering	Marianne Liebi
10.11.25	Diffraction II	Steven Van Petegem
17.11.25	Phase contrast / Tomography	Steven Van Petegem
24.11.25	Coherent imaging	Marianne Liebi
01.12.25	Neutron imaging	Steven Van Petegem
08.12.25	PEEM / Magnetic scattering	Steven Van Petegem
15.12.25	Case study presentations	Steven Van Petegem / Marianne Liebi

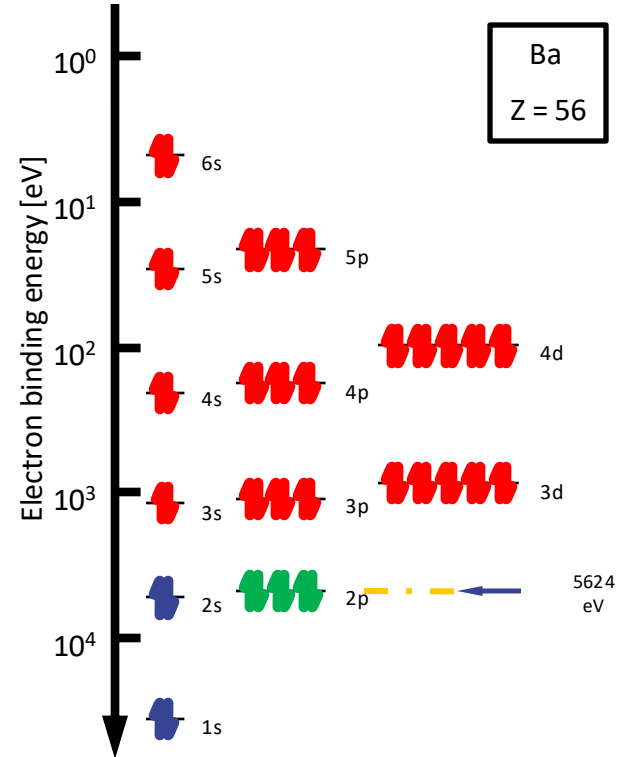
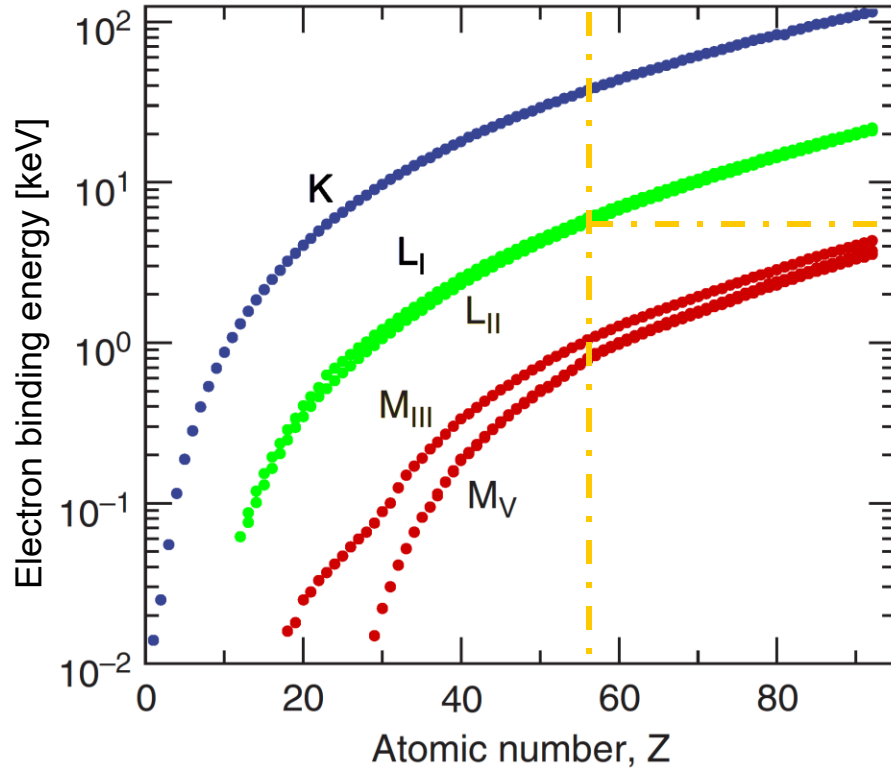
# Interaction of X-ray with matter



# Interaction strengths of x-rays with matter

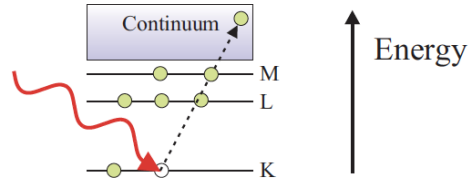


# Electron binding energies of the elements

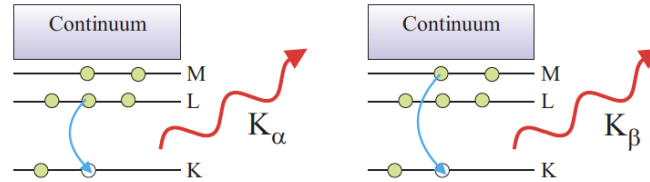


# Absorption

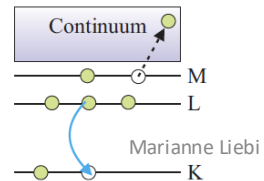
(a) Photoelectric absorption



(b) Fluorescent X-ray emission



(c) Auger electron emission



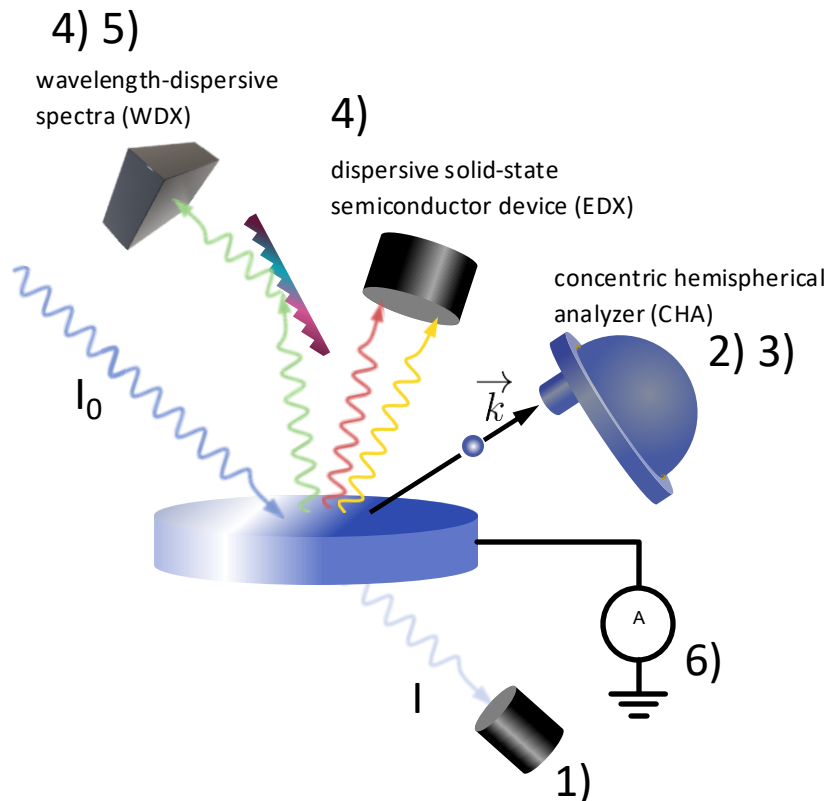
# Overview of spectroscopy techniques

Technique	Abbreviation	Scanned $h\nu_{in}$ ?	Detection methods
X-ray absorption near-edge spectroscopy	XANES	✓	1, 2, 4, 6
Photoemission electron microscopy	PEEM	✓	6
Scanning transmission x-ray microscopy	STXM	✓	1, 3
Extended x-ray absorption fine structure	EXAFS	✓	1, 2, 4, 6
X-ray fluorescence	XRF		4
Resonant inelastic x-ray scattering	RIXS	✓	5
Angle-resolved photoelectron spectroscopy	ARPES	(✓)	2
X-ray photoelectron spectroscopy	XPS	(✓)	2, 3

Variations of XAS

not covered in this course

# Detection methods

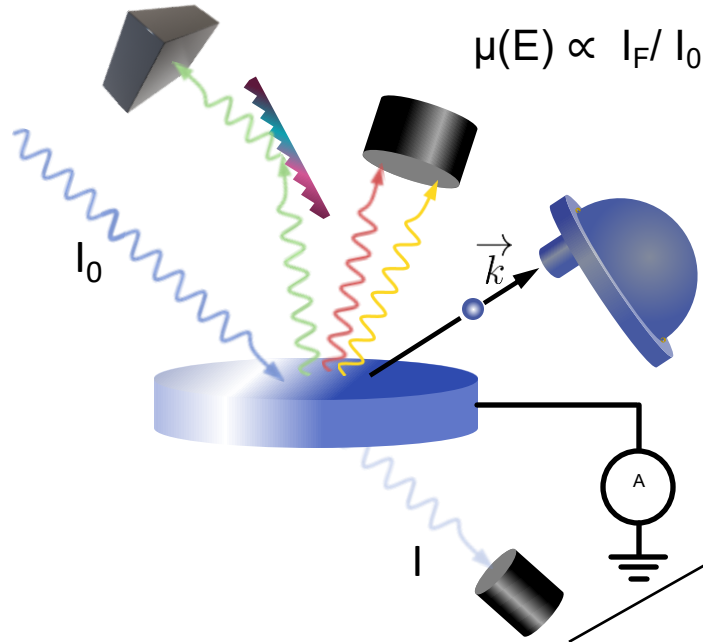


## Form of detection

- 1) Transmitted x-radiation
  - 2) Emitted photoelectrons  
Also as function of  $\vec{k}$
  - 3) Auger electrons
  - 4) Emitted fluorescence
  - 5) Inelastically scattered x-radiation
- } Spectrally resolved
- 6) Secondary electrons/total electron yield
    - Not spectrally resolved
    - Used as a measure of absorption strength

# Detection methods

for thick samples or low concentrations  
(down to ppm level), fluorescence is  
preferred

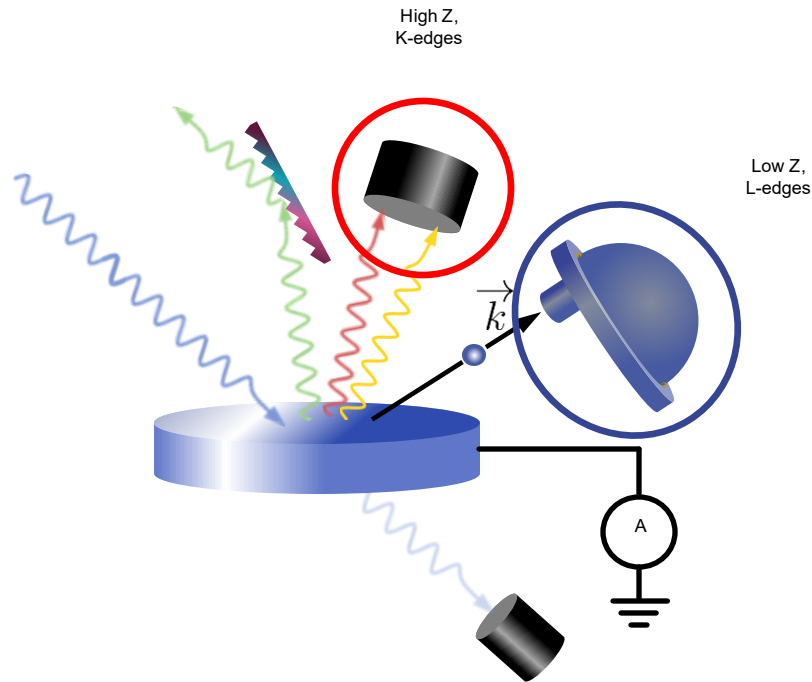
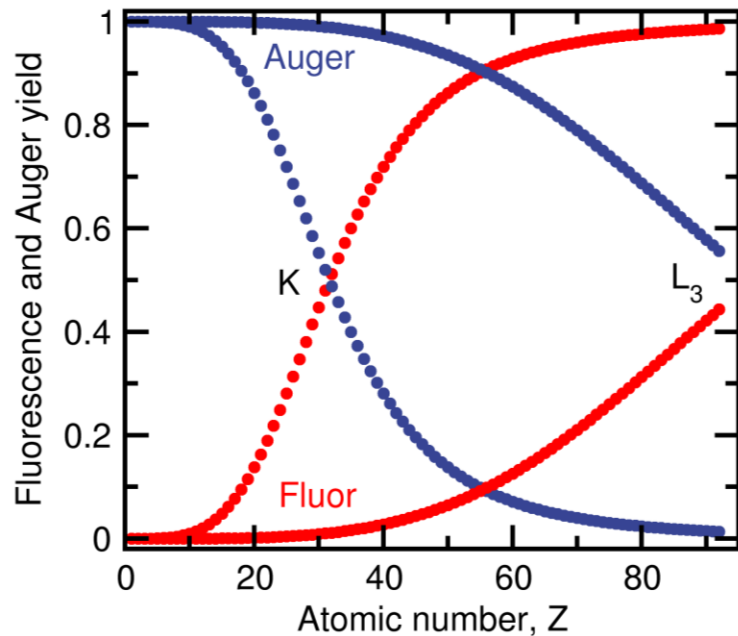


$$\mu(E) \propto I_F / I_0$$

$$I = I_0 e^{-\mu(E)t}$$

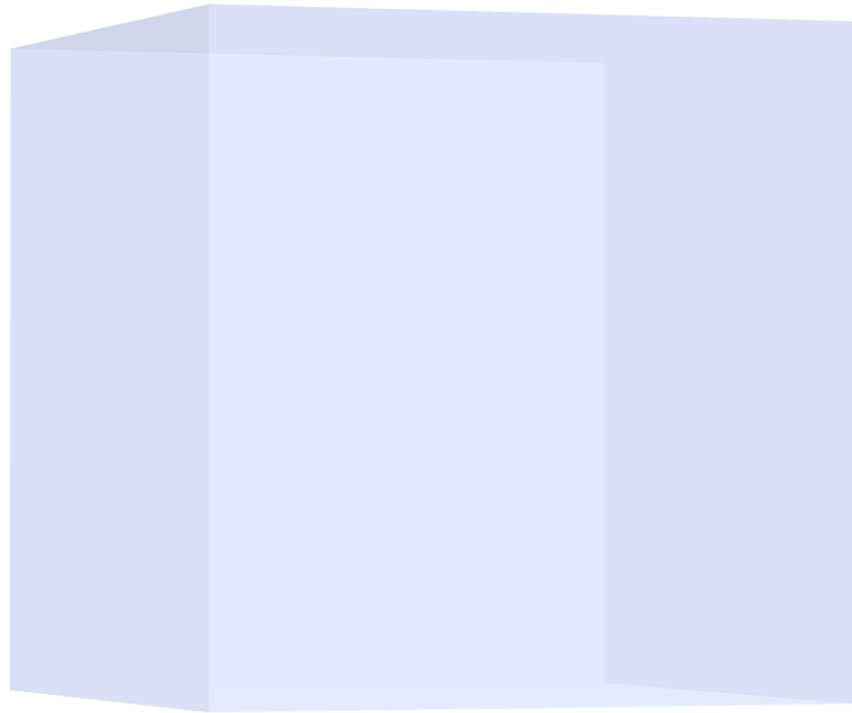
$$\mu(E)t = -\ln(I/I_0)$$

# Detection methods

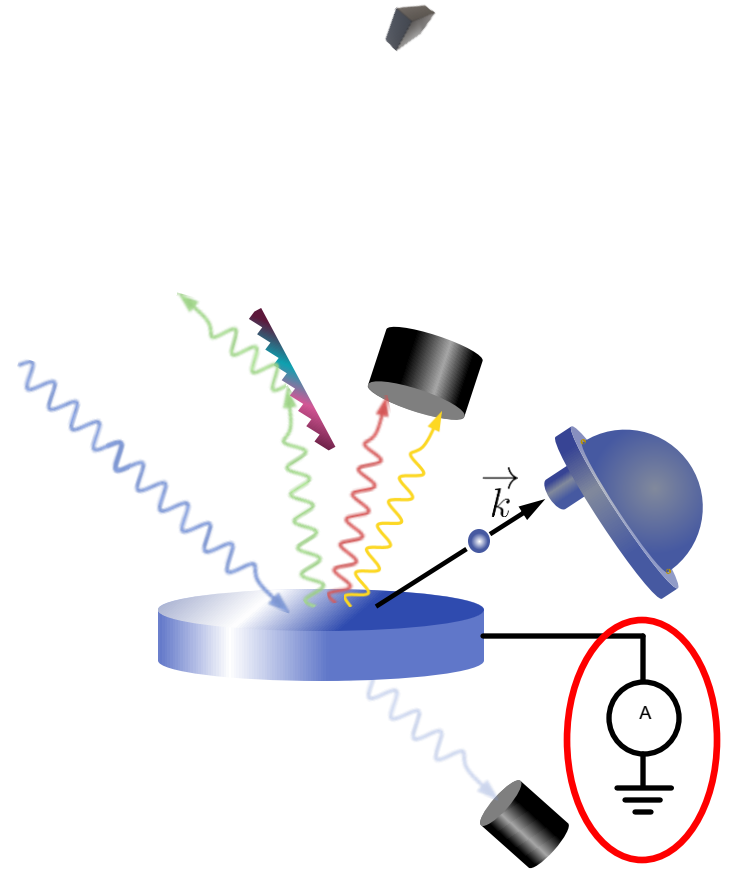


N.B. Detection via Auger-electron yield provides high surface sensitivity!!

# Detection methods



50 Å

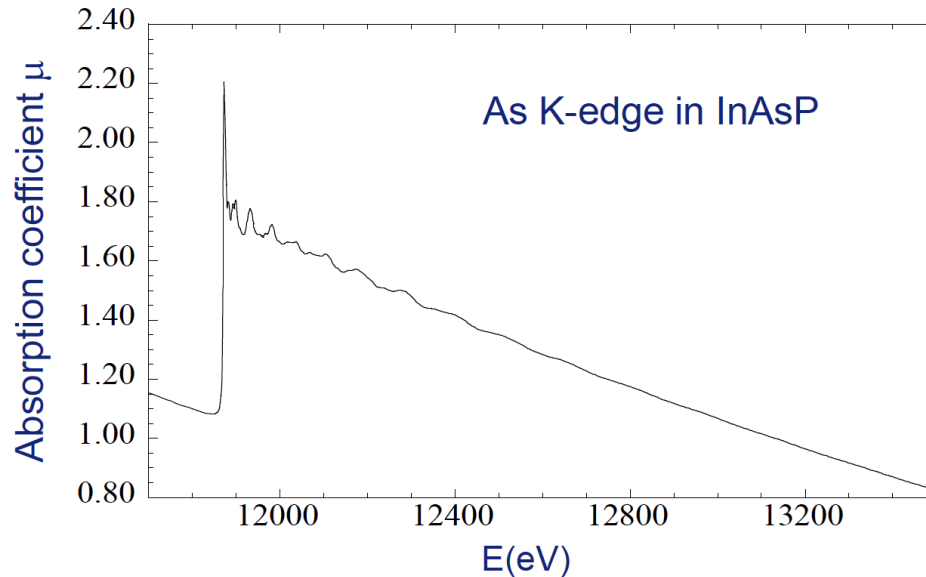


# X-ray absorption fine structure XAFS

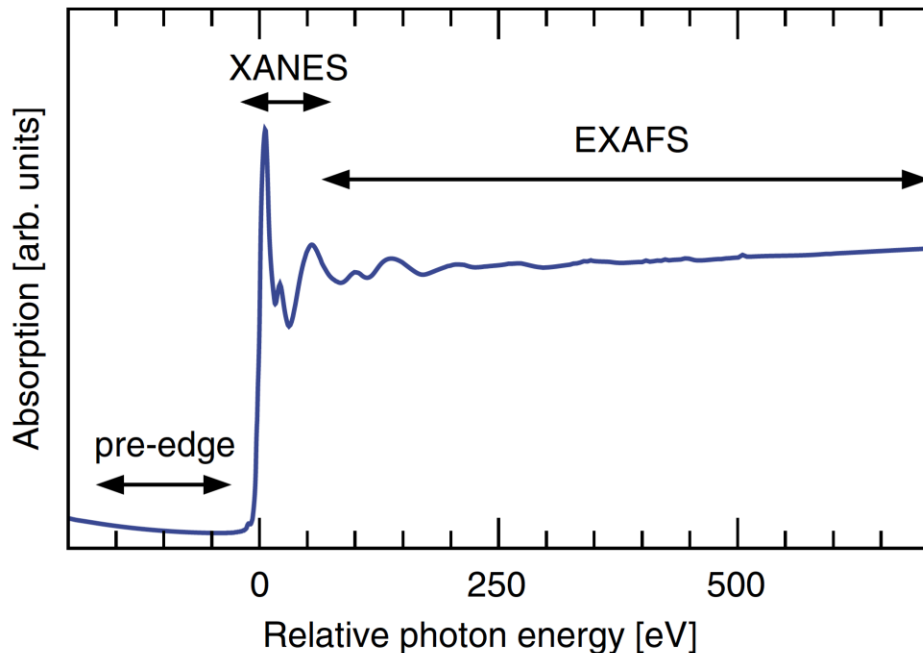
Oscillatory variations of the X-ray absorption as a function of photon energy beyond an absorption edge

Proximity of neighboring atoms strongly modulates the absorption coefficient

Determine the chemical state and local atomic structure for one selected element (element specific)



# XAFS – XANES and EXAFS



## XANES

X-ray absorption near-edge spectroscopy  
(alternatively, NEXAFS)

Pre-edge to ca. 50 eV above edge

## EXAFS

Extended x-ray absorption spectroscopy  
ca. 50 – 1000 eV above edge

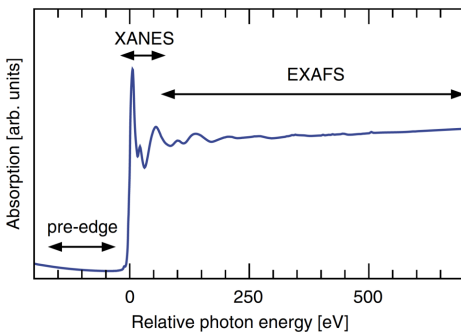
# XAFS – XANES and EXAFS

## XANES

X-ray absorption near-edge spectroscopy (alternatively, NEXAFS)

Pre-edge to ca. 50 eV above edge

transitions to unfilled bound states, nearly bound states, continuum: low energy photoelectrons



## EXAFS

Extended x-ray absorption spectroscopy

ca. 50 – 1000 eV above edge

transitions to continuum: high energy photoelectrons

# XAFS – XANES and EXAFS

## XANES

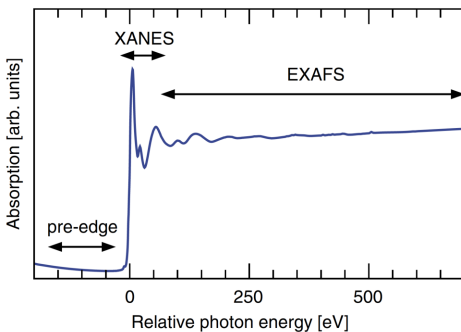
X-ray absorption near-edge spectroscopy (alternatively, NEXAFS)

Pre-edge to ca. 50 eV above edge

transitions to unfilled bound states, nearly bound states, continuum: low

energy photoelectrons

local site symmetry, charge state, valence, oxidation state, orbital hybridization, coordination geometry



## EXAFS

Extended x-ray absorption spectroscopy

ca. 50 – 1000 eV above edge

transitions to continuum: high energy photoelectrons

local structure (bond distance, number, type of neighbors, static and thermal relative displacements)

# XAFS – XANES and EXAFS

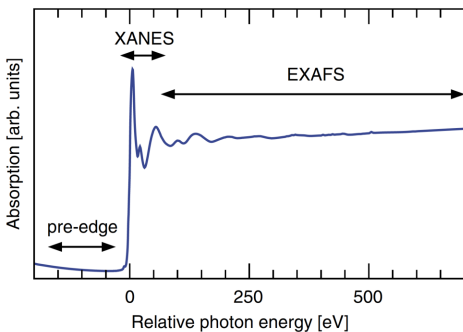
## XANES

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Pre-edge to ca. 50 eV above edge

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## EXAFS

Extended x-ray absorption spectroscopy

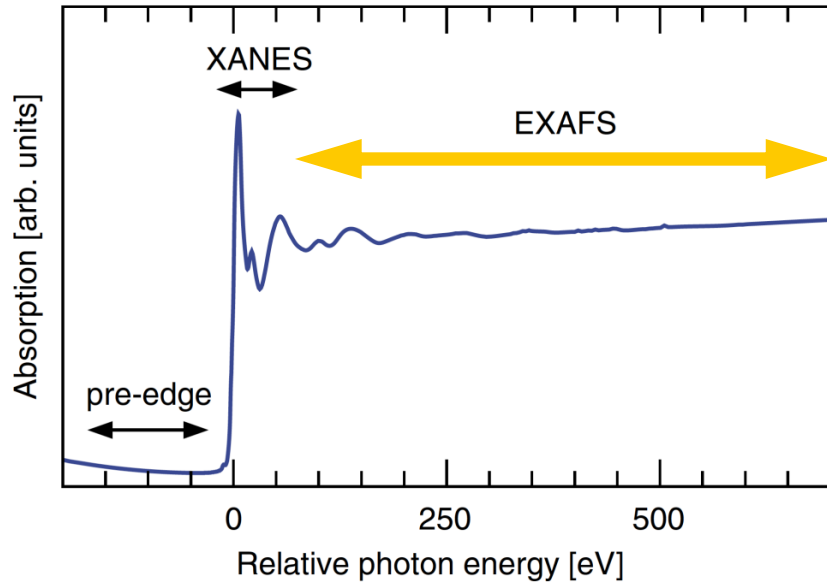
ca. 50 – 1000 eV above edge

transitions to continuum: high energy photoelectrons

local structure (bond distance, number, type of neighbors, static and thermal relative displacements)

approximations can be used to interpret EXAFS, that are not valid for XANES → different analysis

# EXAFS



ca. 50 – 1000 eV above absorption edge

Probes immediate neighbourhood around absorbing atom

Single-scattering dominates

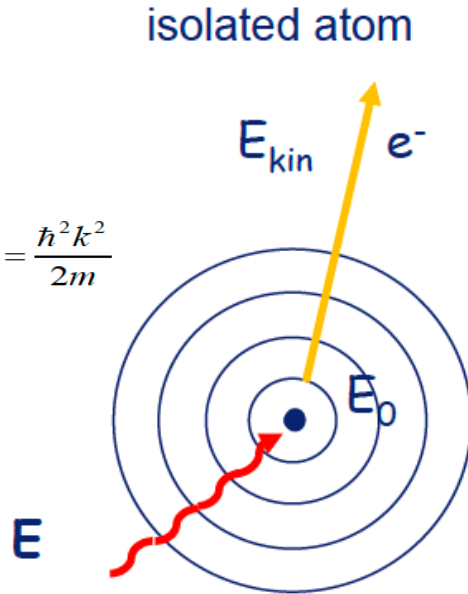
Oscillations due to interference of backscattered electron with outgoing photoelectron

Sensitivity down to ca. 50 ppm

# EXAFS

$$E_{kin} = E - E_0 = \frac{p^2}{2m} = \frac{\hbar^2 k^2}{2m}$$

$$\lambda = 2\pi/k$$



The absorption coefficient  $\mu$

→ probability of photon absorption

→ probability of electron presence at origin (is there an available state for the photo-electron?)

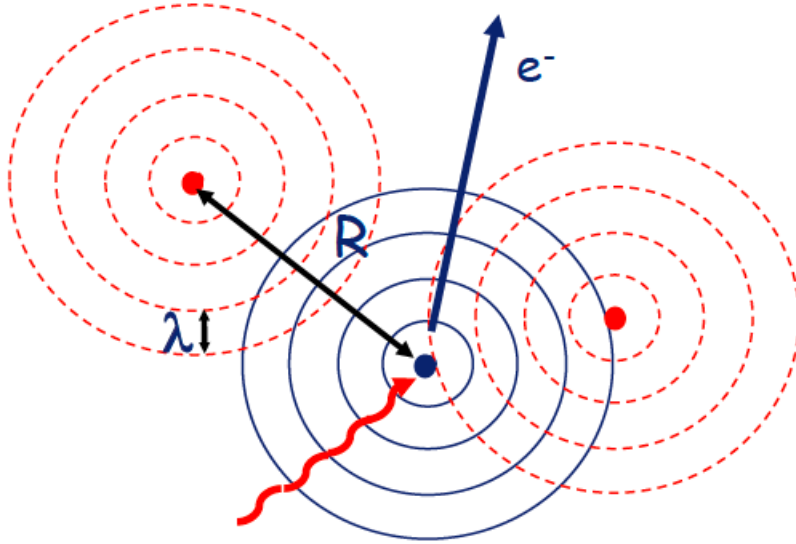
→ "amount of wave" of the ejected photoelectron at origin

kinetic Energy of the ejected photoelectron changes if the Energy E is scanned

# Origin of the oscillations in EXAFS

$$E_{kin} = E - E_0 = \frac{p^2}{2m} = \frac{\hbar^2 k^2}{2m}$$

$$\lambda = 2 \pi / k$$

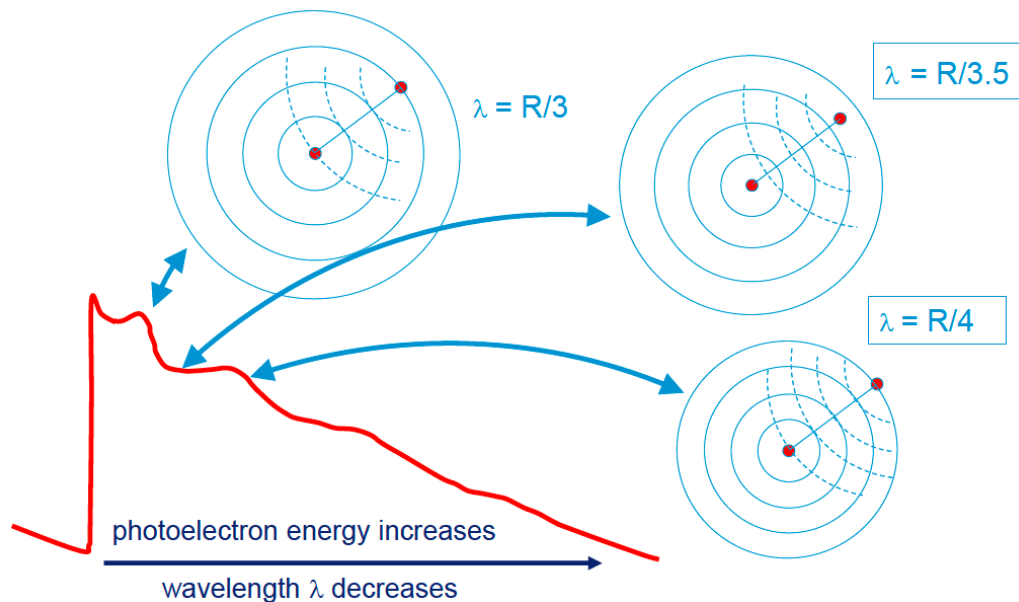


Oscillations due to interference of backscattered electron with outgoing photoelectron

The outgoing and backscattered parts of the wave interfere either constructively or destructively, depending on the ratio between  $\lambda$  and  $R$

$\lambda$  changes when the energy of the incoming X-rays is scanned

# Origin of the oscillations in EXAFS



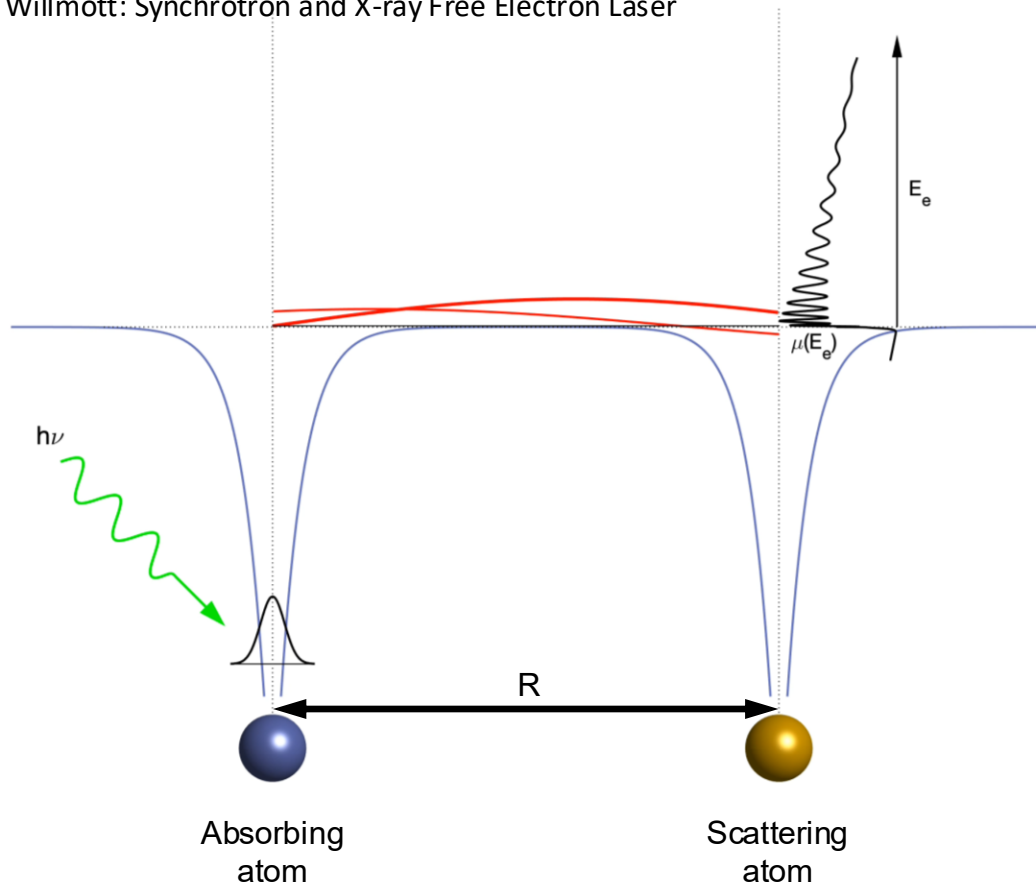
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# EXAFS signal

Philip Willmott: Synchrotron and X-ray Free Electron Laser



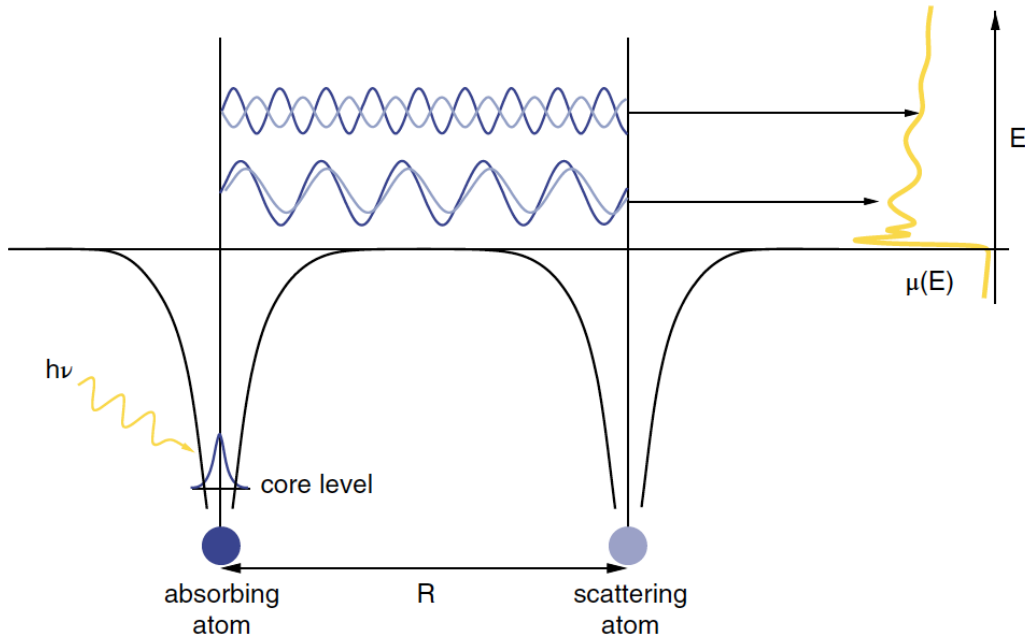
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$\lambda$  changes when the energy of the incoming X-rays is scanned

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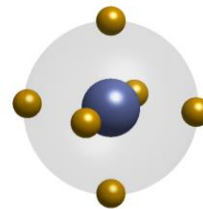
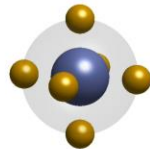
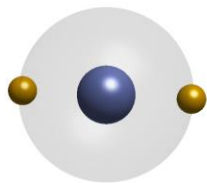


Oscillations due to interference of backscattered electron with outgoing photoelectron

The outgoing and backscattered parts of the wave interfere either constructively or destructively, depending on the ratio between  $\lambda$  and  $R$

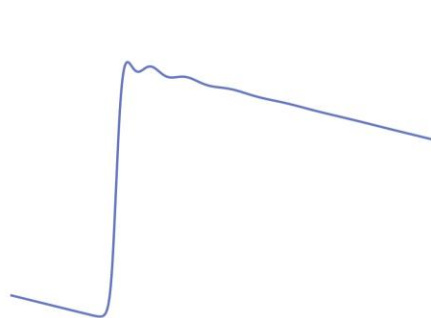
$\lambda$  changes when the energy of the incoming X-rays is scanned

# The EXAFS signal – rules of thumbs

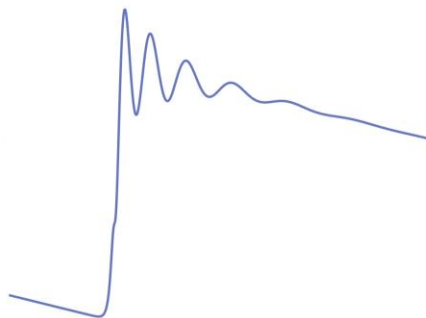


frequency  $\rightarrow$  distance from neighbors

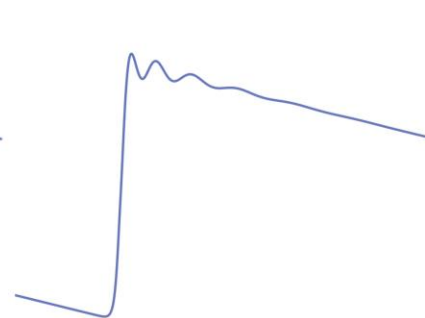
amplitude  $\rightarrow$  number and type of neighbors



Number of scatterers



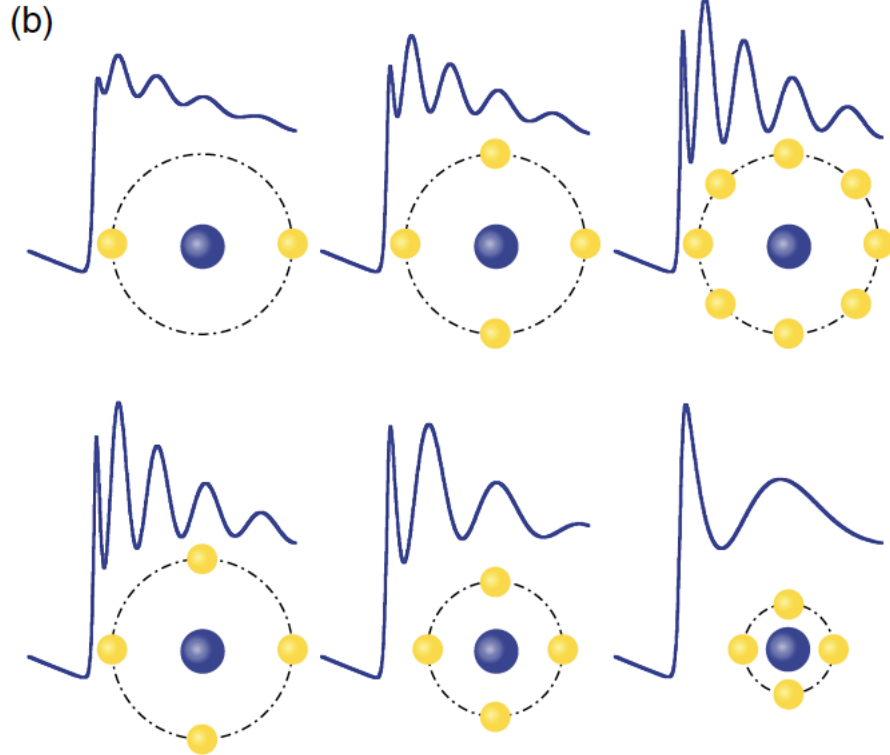
Absorber-scatterer distance



Scatterer atomic number

# The EXAFS signal – rules of thumbs

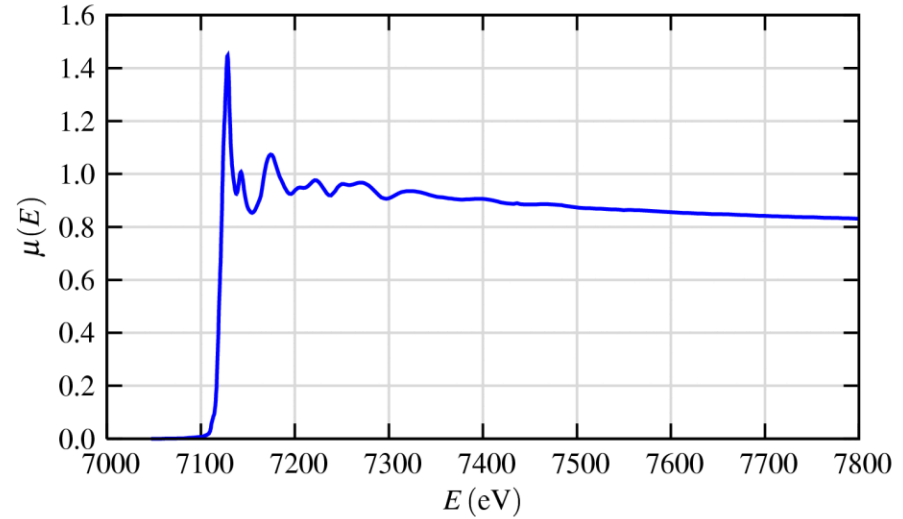
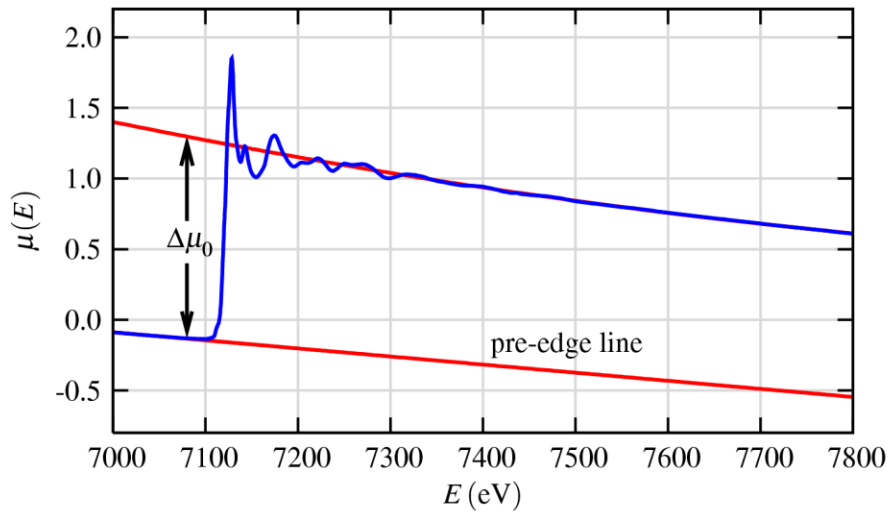
Philip Willmott: Synchrotron and X-ray Free Electron Laser



amplitude  $\rightarrow$  number and type of neighbors  
frequency  $\rightarrow$  distance from neighbors

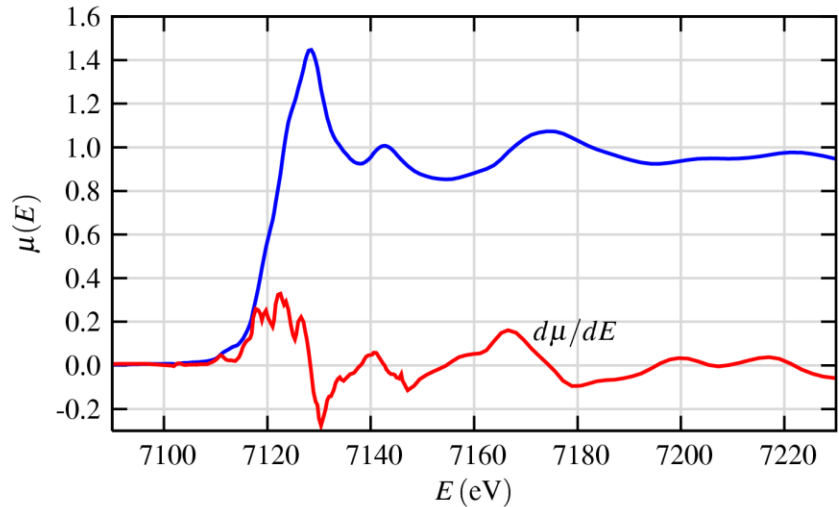
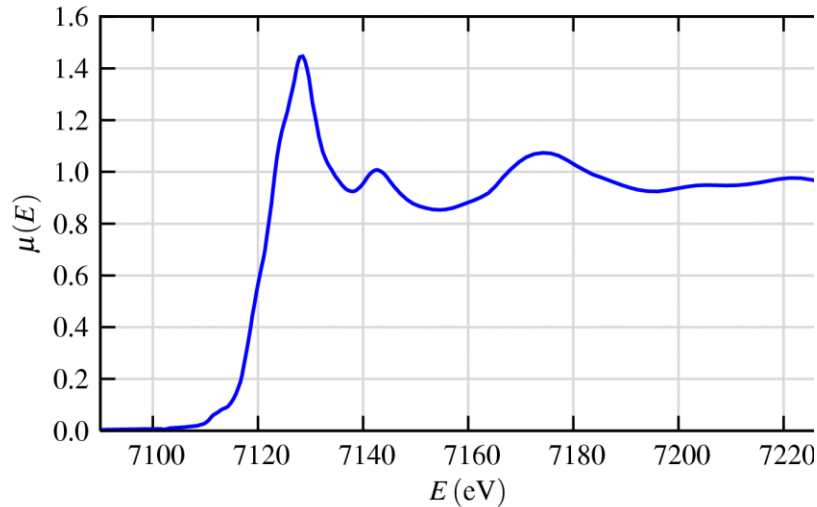
# XAFS – data reduction steps

1. Convert measured intensities to  $\mu(E)$ , possibly correcting systematic measurement errors such as self-absorption effects and detector dead-time.
2. Subtract a smooth pre-edge function from  $\mu(E)$  to get rid of any instrumental background and absorption from other edges.



# XAFS – data reduction steps → XANES

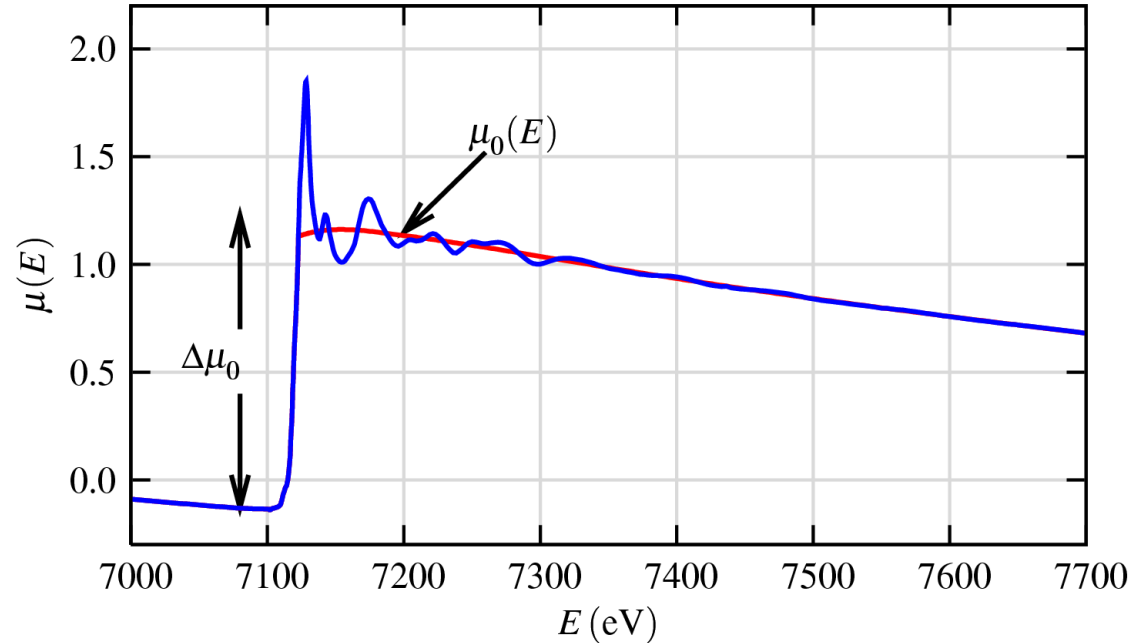
- Identify the threshold energy  $E_0$ , typically as the energy of the maximum derivative of  $\mu(E)$ .
- Normalize  $\mu(E)$  to go from 0 to 1, so that it represents the absorption of 1 x-ray. The normalized spectra are useful for XANES analysis.



# XAFS – data reduction steps → EXAFS

5. Remove a smooth post-edge background function to approximate  $\mu_0(E)$ .

$$\chi(E) = \frac{\mu(E) - \mu_0(E)}{\Delta\mu_0(E)}$$



# EXAFS

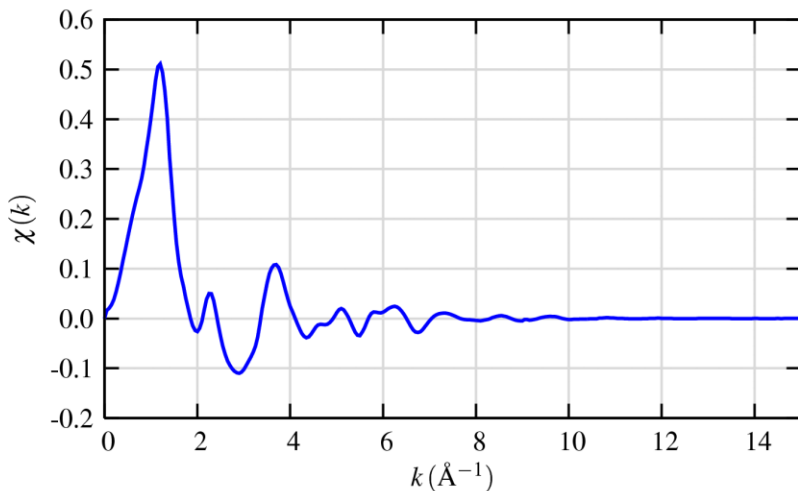
EXAFS is best understood in terms of the wave behavior of the photo-electron created in the absorption process. Because of this, it is common to convert the x-ray energy to  $k$ , the wave number of the photo-electron, which has dimensions of 1/distance and is defined as

$$k = \sqrt{\frac{2m(E - E_0)}{\hbar^2}}$$

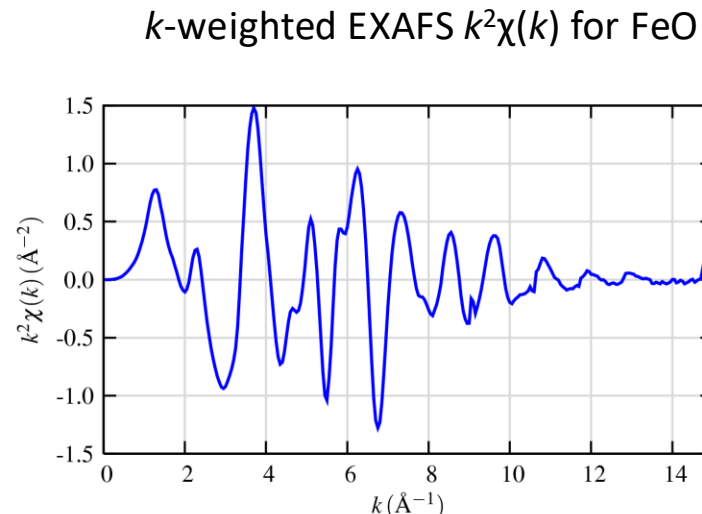
# XAFS data reduction steps → EXAFS

6. Isolate the XAFS  $\chi(k)$ , where  $k = \sqrt{\frac{2m(E - E_0)}{\hbar^2}}$

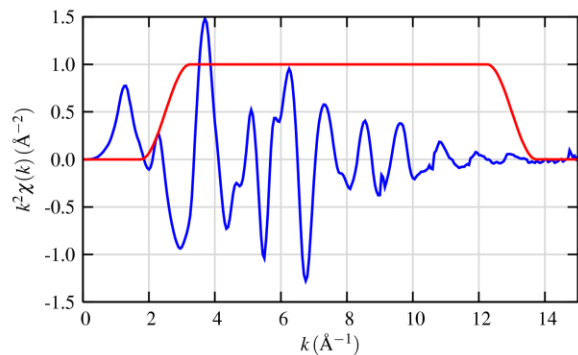
k-weight the XAFS



isolated EXAFS  $\chi(k)$  for FeO  
oscillatory, decays quickly with  $k$

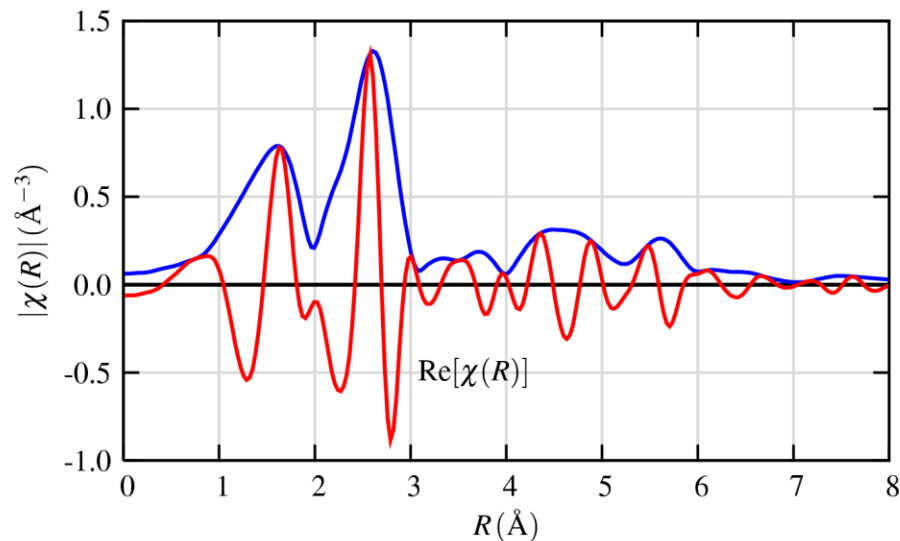


## 8. Fourier Transform to $\chi(R)$



Window function before doing a Fourier transform

## Fourier transformed EXAFS, a complex function



magnitude  $|\chi(R)|$

real part  $\text{Re}[\chi(R)]$

# EXAFS equation

Can be derived by a formal description of X-ray absorption as a transition probability between two quantum states

## initial state

an X-ray  
a core electron  
no photo-electron



## final state

no X-ray  
core hole  
a photo-electron

core electron tightly bound, not altered by neighboring atoms

photo-electron “sees” the neighboring atoms

$$\mu(E) = \mu_0(E)[1 + \chi(E)]$$

“bare atom absorption”, only depends on the absorbing atom

fine structure  $\chi(E) \propto \langle i | H | \Delta f \rangle$

# EXAFS equation

Mean-square displacement of distance (disorder in neighbor distance)

Number of neighboring atoms

Amplitude

Phase shift

$$\chi(k) = \sum_j \frac{|N_j f_j(k) e^{-2k^2 \sigma_j^2}|}{k R_j^2} \sin[2k R_j + \delta_j(k)]$$

EXAFS fine structure function

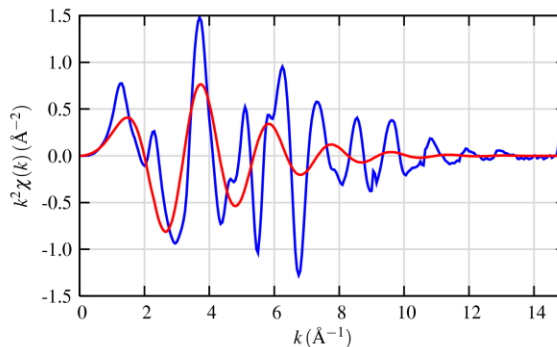
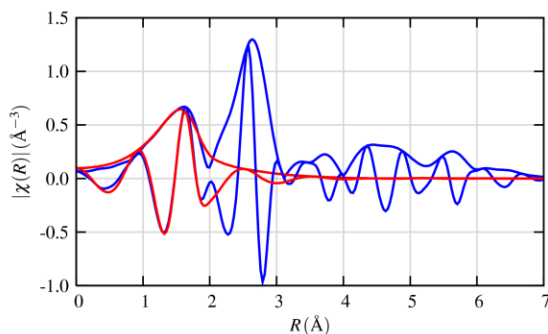
distance to neighboring atom ( $\pm 0.03 \text{ \AA}$ )

$f(k)$  and  $\delta(k)$  are scattering properties of the neighbouring atoms to the excited atom  
 → EXAFS is sensitive to the atomic species of the neighboring atom

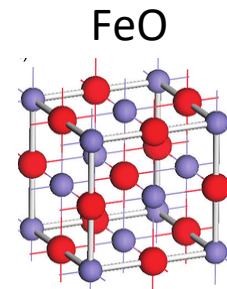
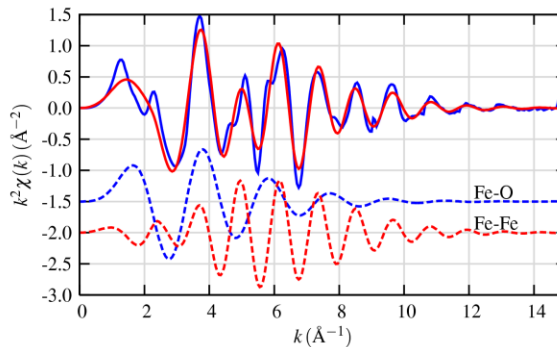
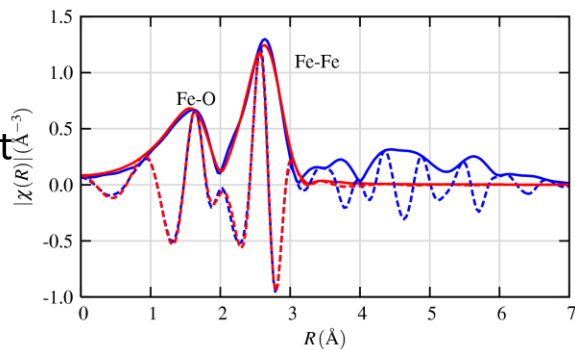
# EXAFS data fitting

calculation of the theoretical scattering factors  $f(k)$  and  $\delta(k)$   
 refine structural parameters  $R$ ,  $N$  and  $\sigma^2$  by fitting the data,  
 either fit  $\chi(k)$  or the Fourier transformed, which has the advantage of fitting one shell at a time

first shell fit



second shell fit



# EXAFS equation

Can be derived by a formal description of X-ray absorption as a transition between two quantum states

## initial state

an X-ray  
a core electron  
no photo-electron



## final state

no X-ray  
core hole  
a photo-electron

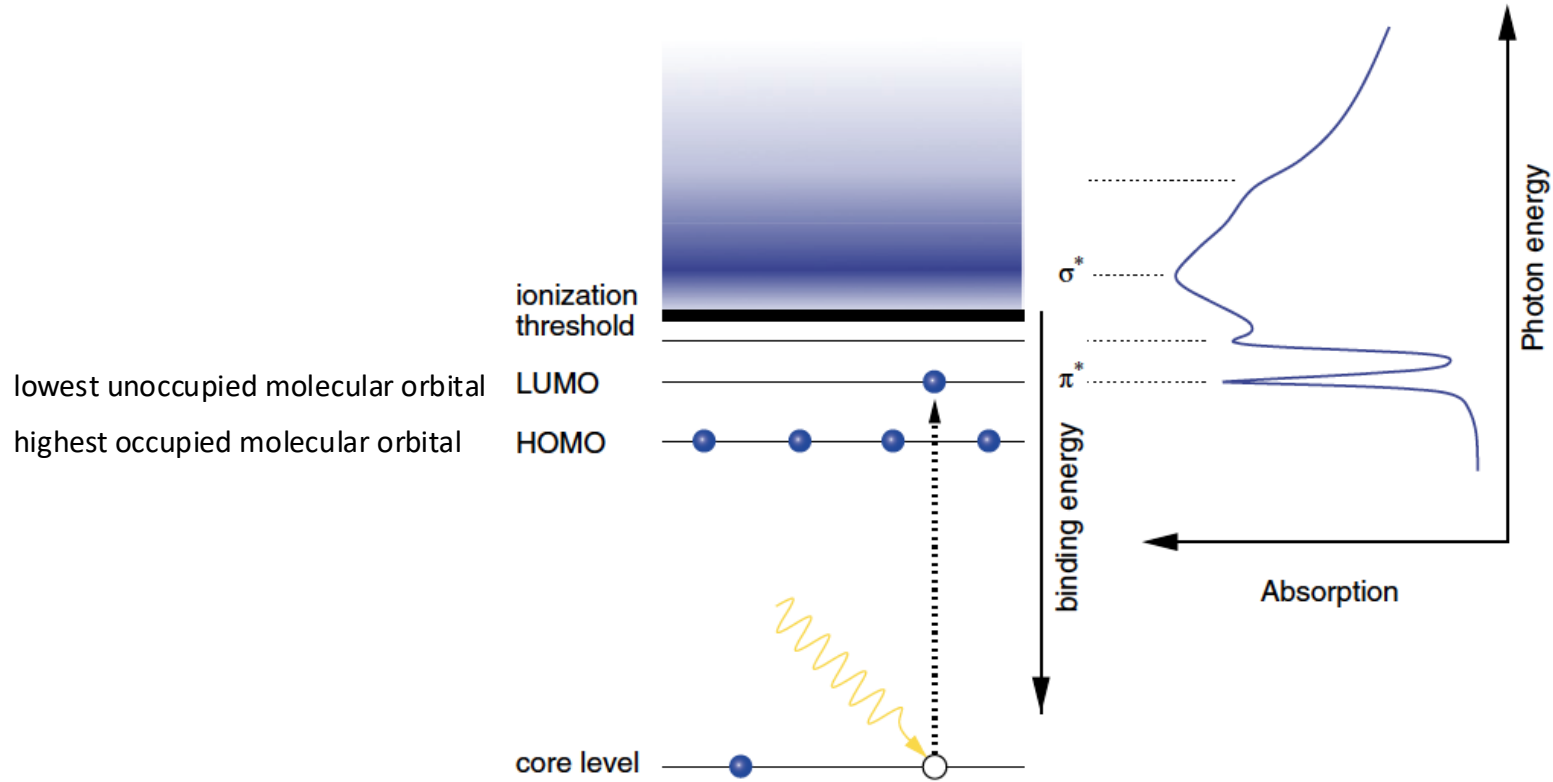
approximation of neglecting multiple scattering is valid in the EXAFS regime, but **not for the XANES part**, the EXAFS equation breaks down: interpretation of XANES is complicated as there is not a simple analytical (or even physical) description of XANES

But the edge position and shape is sensitive to valence state, ligand type and coordination environment → fingerprint

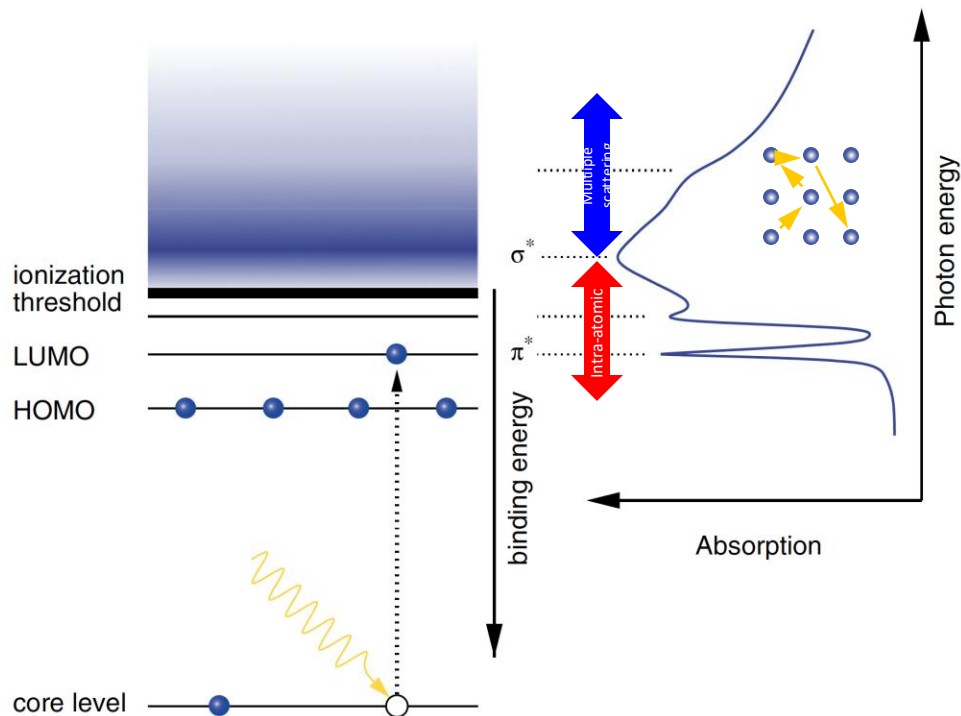
# XANES

- XANES is a much larger signal than EXAFS
  - XANES can be done at lower concentrations, and less-than-perfect sample conditions.
- XANES is harder to fully interpret than EXAFS
  - The exact physical and chemical interpretation of all spectral features is still difficult to do accurately, precisely, and reliably.
- XANES is easier to crudely interpret than EXAFS
  - For many systems, the XANES analysis based on linear combinations of known spectra from “model compounds” is sufficient: compositional fraction of these components
  - often Principle Component Analysis and Factor Analysis are used
- key factors influencing XANES: ligands/charge, symmetry, bond length

# XANES



# Transitions in XANES



Core electron promoted to unoccupied excited state (both bound and unbound)

**“White lines”**: unoccupied bound states

LUMO

e.g.  $\pi^*$ ,  $\sigma^*$

Partially empty valence electron shell

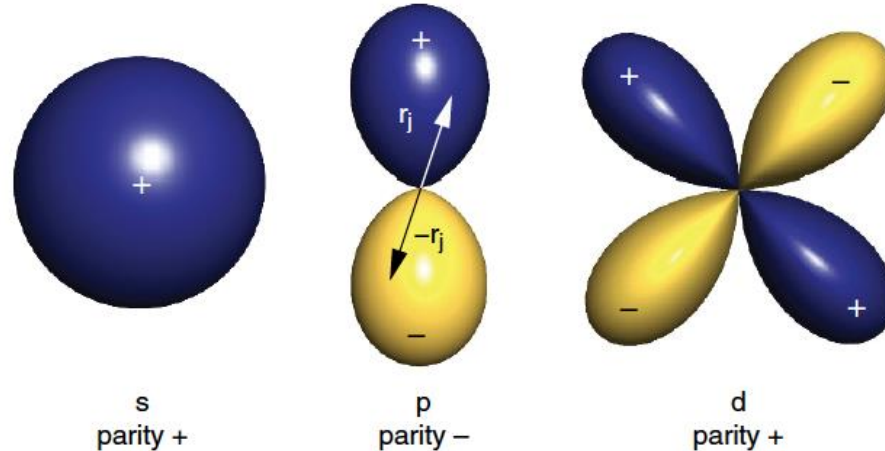
e.g. 5d shell of noble metals

Probes large local region around absorbing atom

Multiple elastic-scattering events

⇒ difficult to model

allowed transitions determined by the symmetry of the local environment  
initial and final states must have opposite symmetries

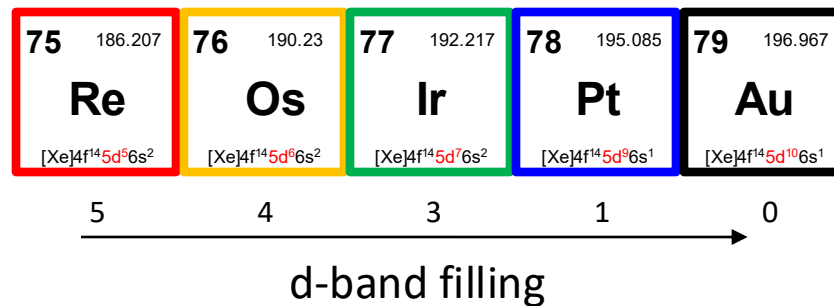
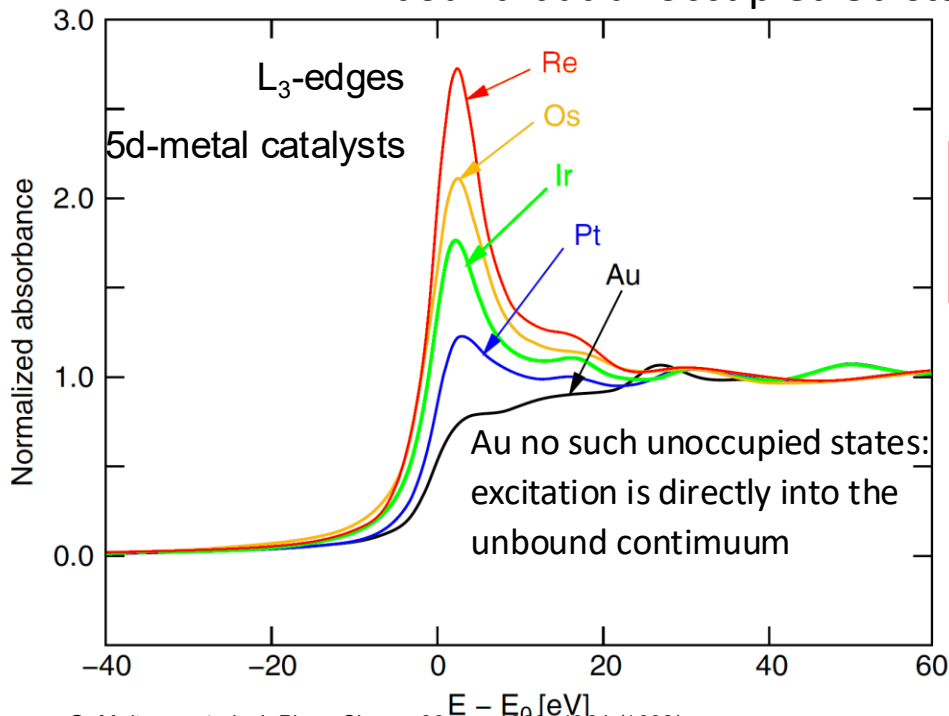


**Figure 7.12** Parity and symmetry. The parity of an atomic electron orbital is either positive or negative, depending on how it is transformed when moving all the elements  $j$  of the orbital's amplitude wavefunction from  $r_j$  to  $-r_j$ . So, for example,  $p$  orbitals are antisymmetric with negative parity, while  $d$  orbitals are symmetric and have positive parity.

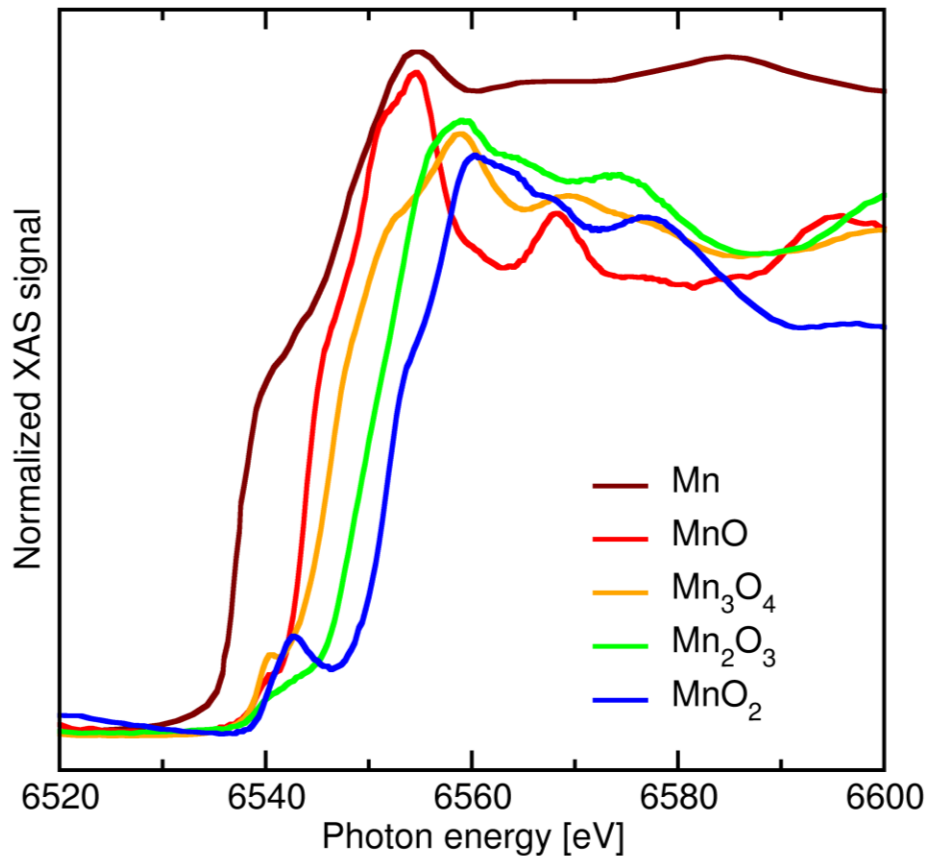
# X-ray absorption fine structure XAFS

“white line”: bound excited state features here:

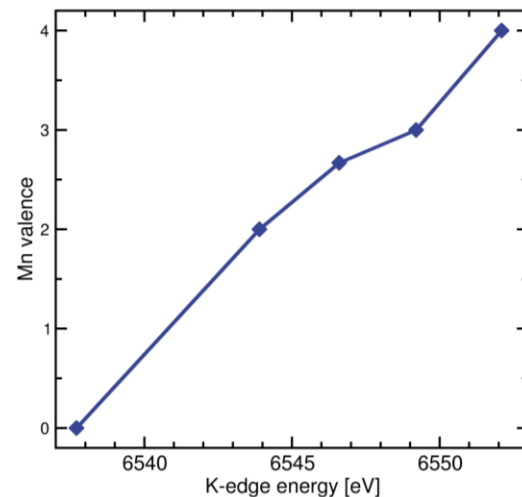
bound but unoccupied 5d states, below the ionization threshold



# XANES: Oxidation state



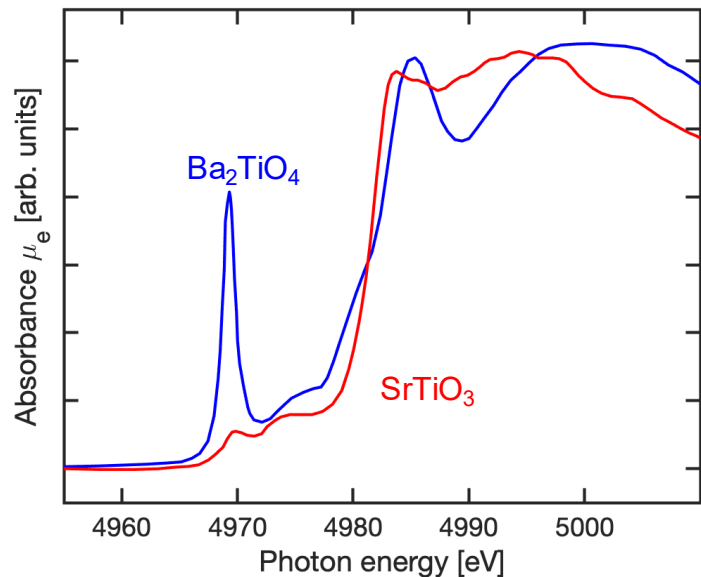
Contains information about the oxidation state:  
shift in edge energy



# XANES: coordination chemistry/bond geometry

Contains information about coordination chemistry/bond geometry

## XANES on Ti K-Edge



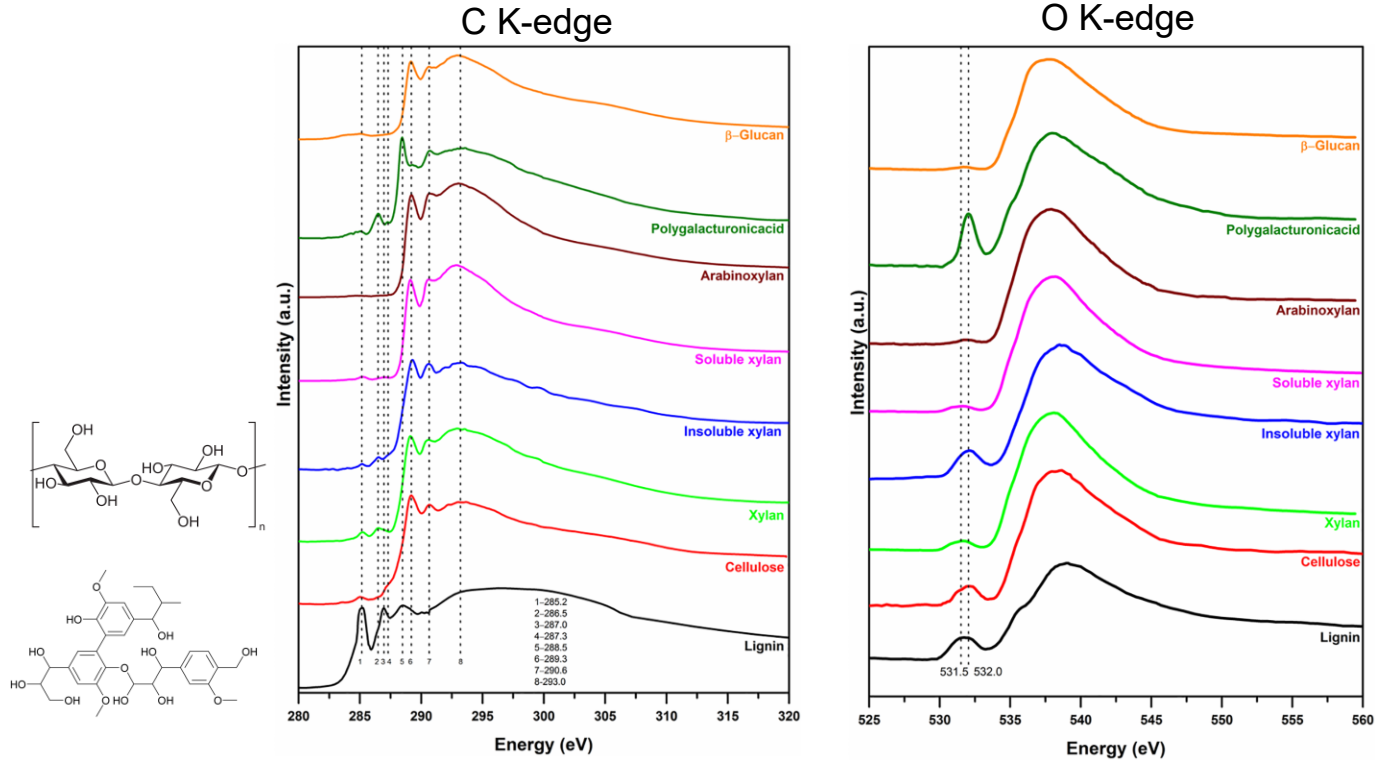
### $\text{TiO}_6$ octahedra in $\text{SrTiO}_3$

- $t_{2g}$ : LUMO with octahedral inversion symmetry
- Cannot be accessed by dipole from 1s core state
- Very weak signal via quadrupole (2-electron) perturbation

### $\text{TiO}_4$ tetrahedra in $\text{Ba}_2\text{TiO}_4$

- $3d^34p$  mixed LUMO states
- p-character of these MOs can be accessed from 1s core state via direct dipole perturbation ( $\Delta l = 1$ )

# XANES of Bio-Polymers



Karunakaran C, et al. (2015) Introduction of Soft X-Ray Spectromicroscopy as an Advanced Technique for Plant Biopolymers Research.

# XANES interpretation

XANES can be described qualitatively (and nearly quantitatively ) in terms of

- coordination chemistry regular, distorted octahedral, tetrahedral, . . .
- molecular orbitals p-d orbital hybridization, crystal-field theory, . . .
- band-structure, the density of available electronic states
- multiple-scattering, multiple bounces of the photoelectron

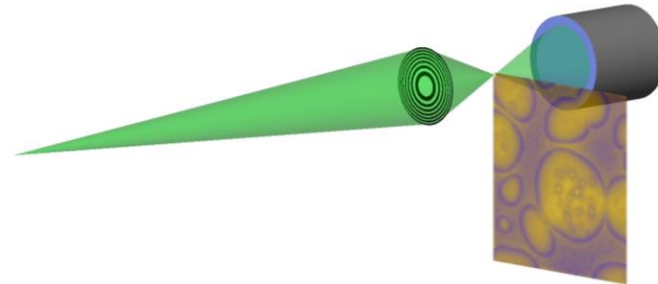
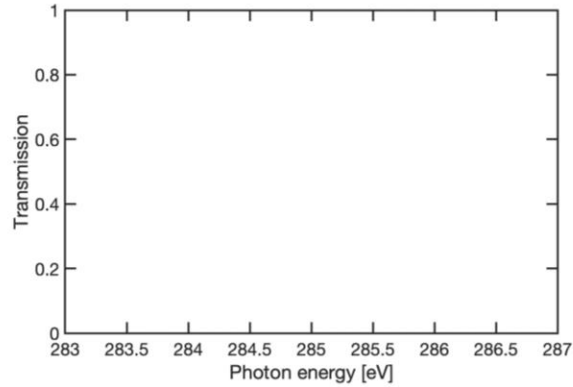
These chemical and physical interpretations are all related, of course:

What electronic states can the photoelectron fill?

XANES calculations are becoming reasonably accurate and simple. These can help explain what bonding orbitals and/or structural characteristics give rise to certain spectral features.

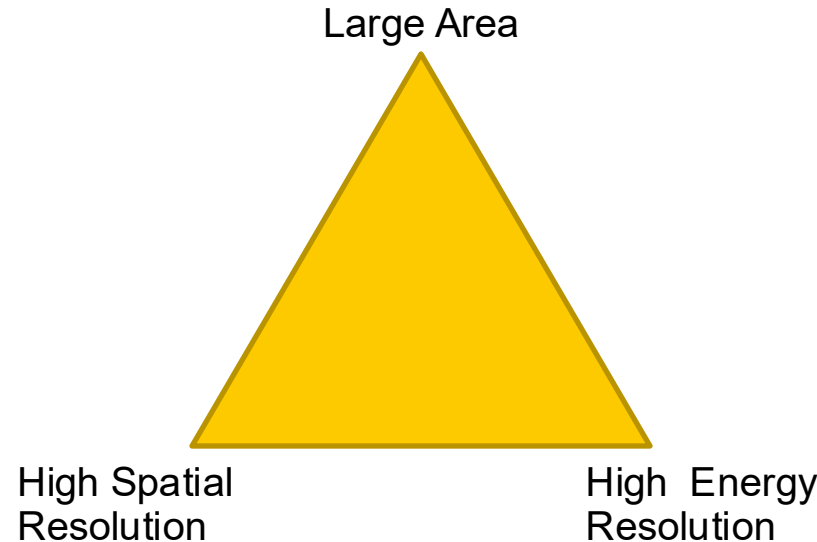
Quantitative XANES analysis using first-principles calculations are still rare, but becoming possible...

# STXM: scanning transmission X-ray microscopy

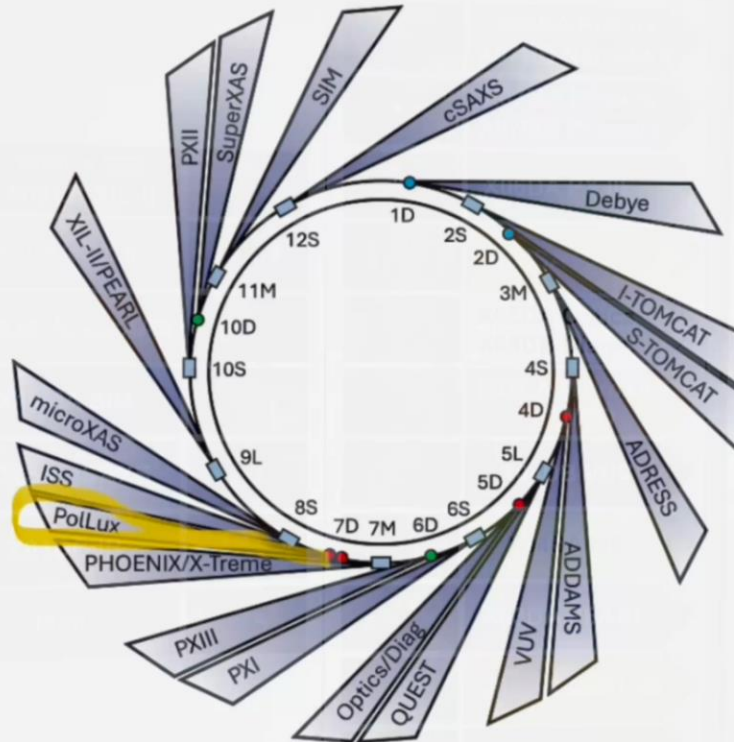


# STXM: scanning transmission X-ray microscopy

- full energy scan at highest resolution for large areas take a long time
- additional problem: radiation damage (multiple exposure, high flux density)
- Combination of two approaches:
  - Energy stacking
    - full Energy range over edge e.g. from 280-350 eV (Carbon edge) with a resolution of  $>0.1$  eV
    - Gives full energy spectra containing information of the chemical compounds present
    - Slow
    - Small Areas or Large stepsize
  - Scan at resonance energies
    - Only predetermine energies
    - Fast
    - Allows high spatial resolution over extended areas



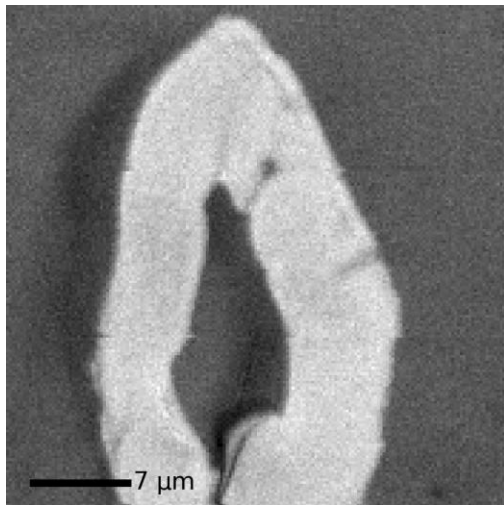
# STXM XAN



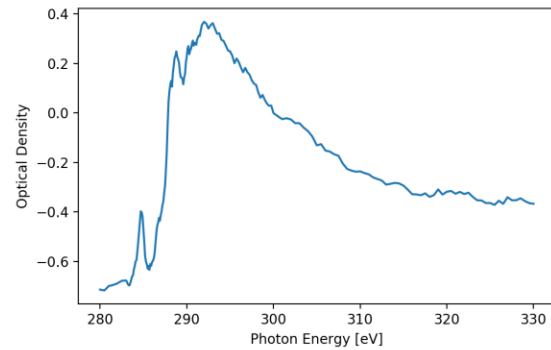
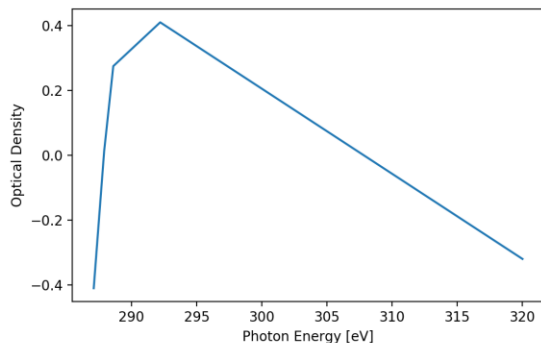
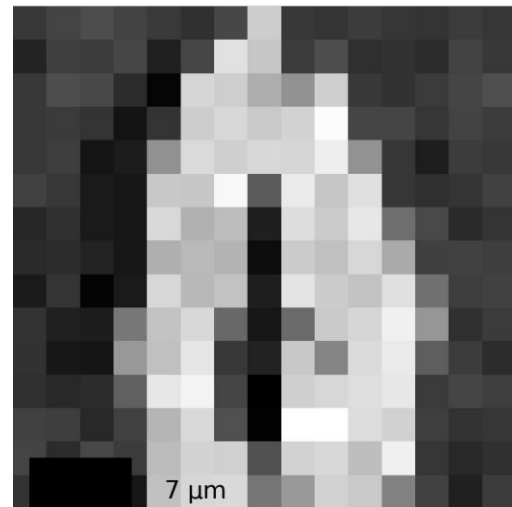
# STXM: high spatial resolution vs. high energy resolution

cellulose fibre

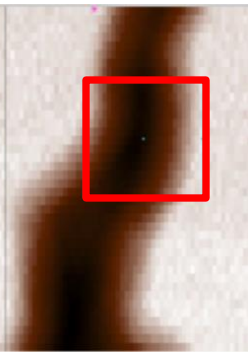
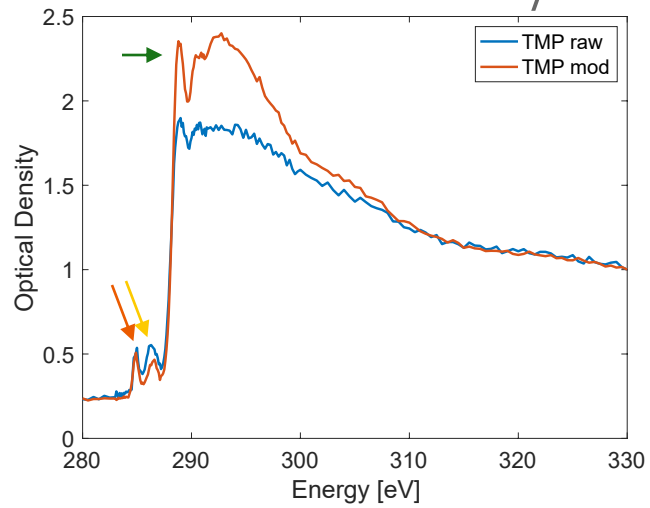
- Trade off between high spatial resolution and high energy resolution



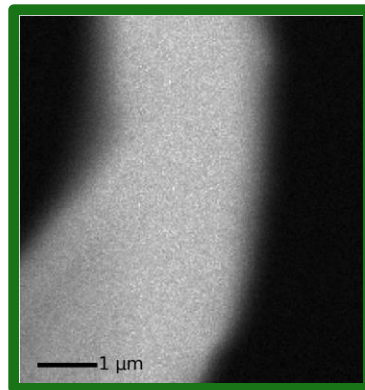
Low spatial resolution scan at 287.1 eV



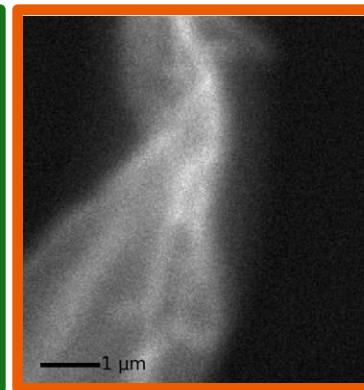
## STXM/XANES lignin on fibers



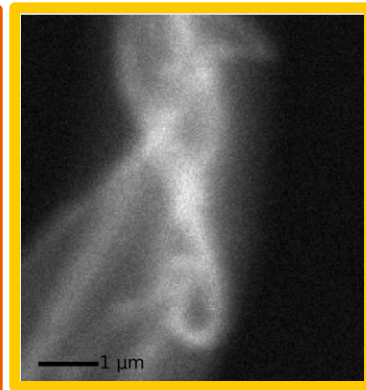
288.9 eV (cellulose)



284.9 eV (lignin)



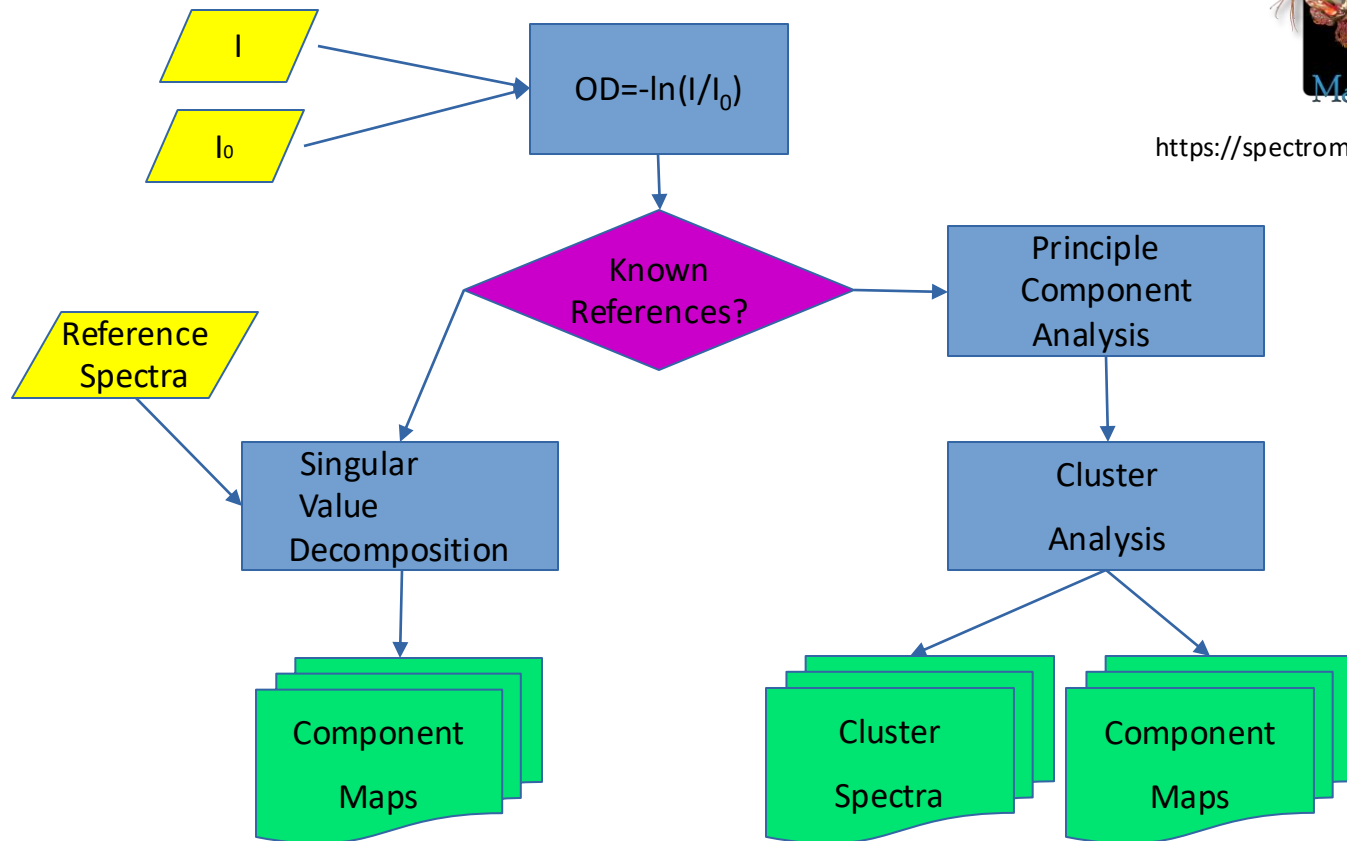
286.2 eV (lignin)



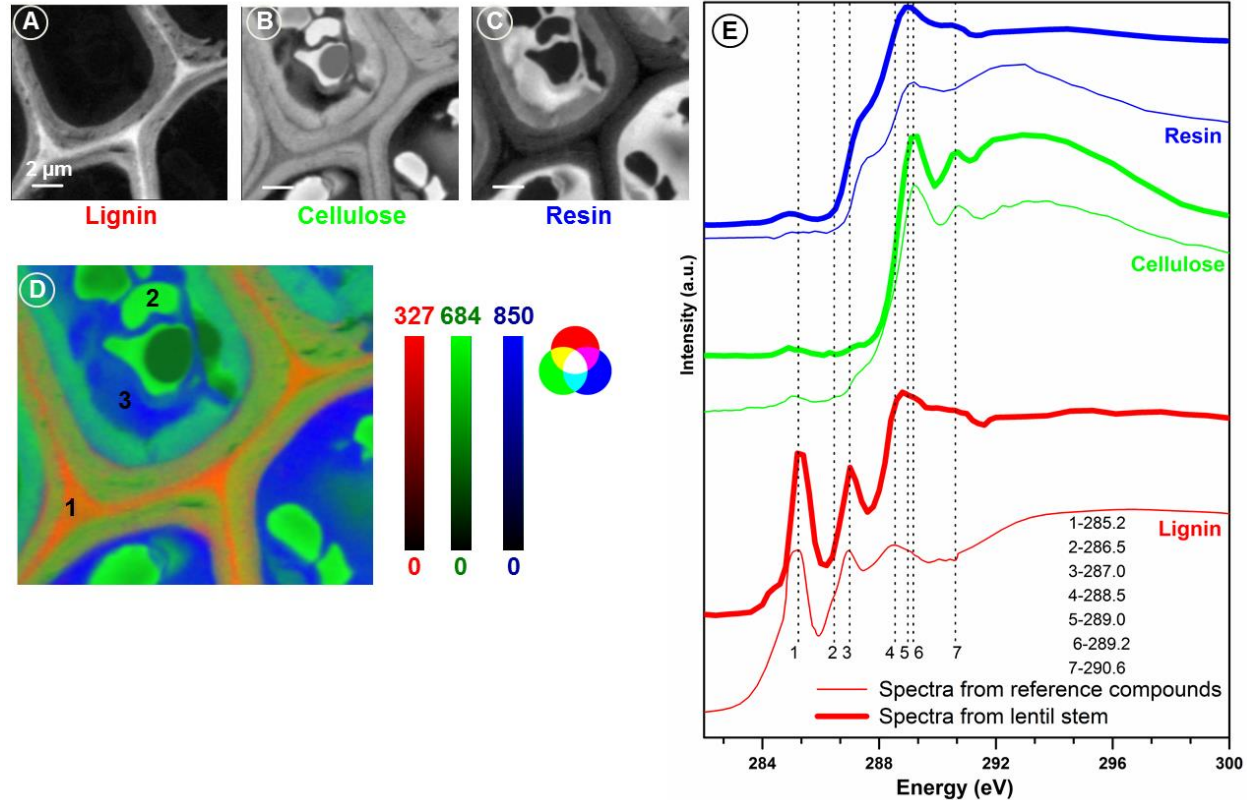
# XANES Data analysis: SVD and PCA



<https://spectromicroscopy.com/>

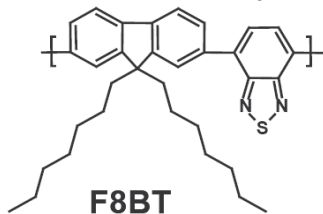


# SVD Analysis of Lentil Stem Section



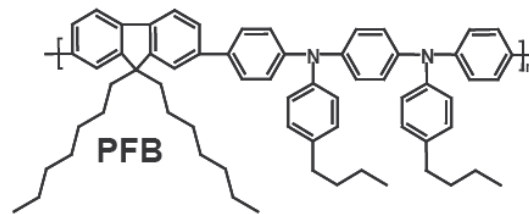
# Model Conjugated Polymers

Electron Acceptor

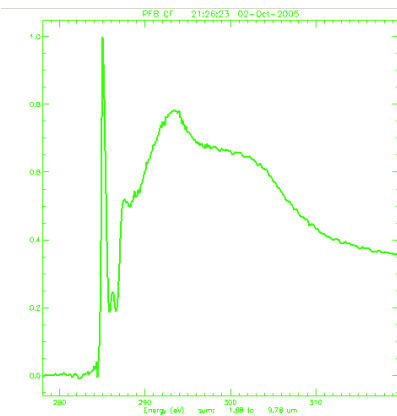
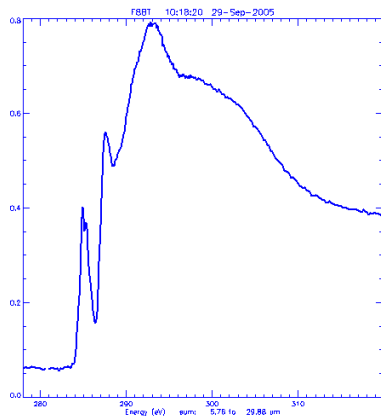


Poly(9,9'-dioctylfluorene-co-benzo-thiadiazole)

Electron Donor



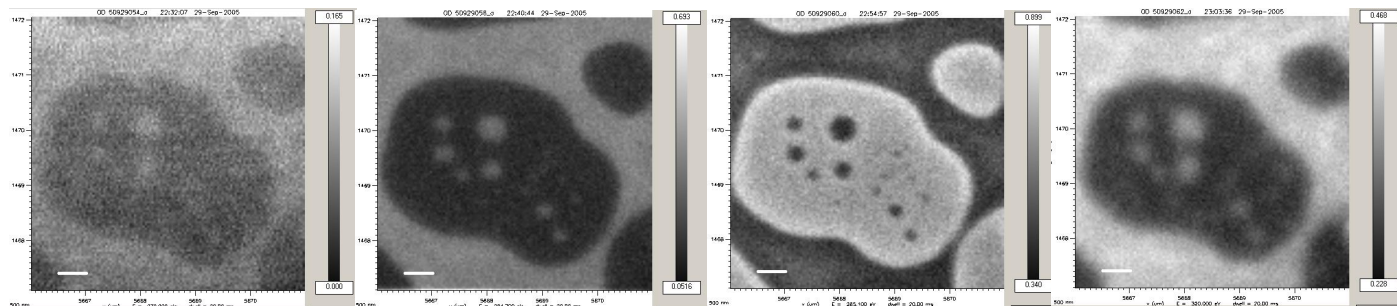
Poly(9,9'-dioctylfluorene-co-bis-N,N'-(4-butylphenyl)-bis-N,N'-phenyl-1,4-phenylene-diamine)



- Absorption step due to photoemission
- Resonance peaks due to core to anti-bonding transitions

reference spectra

# STXM Contrast and Composition Mapping

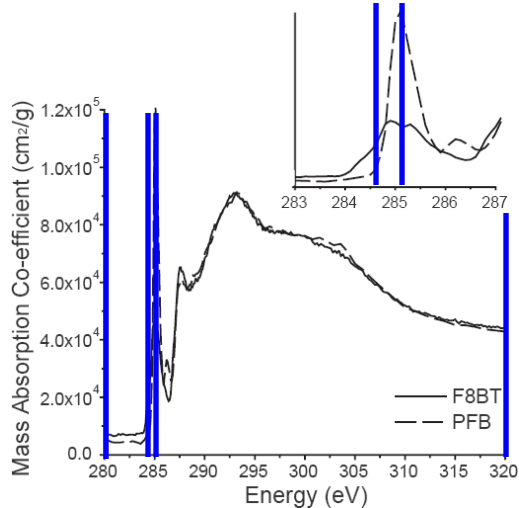


280 eV

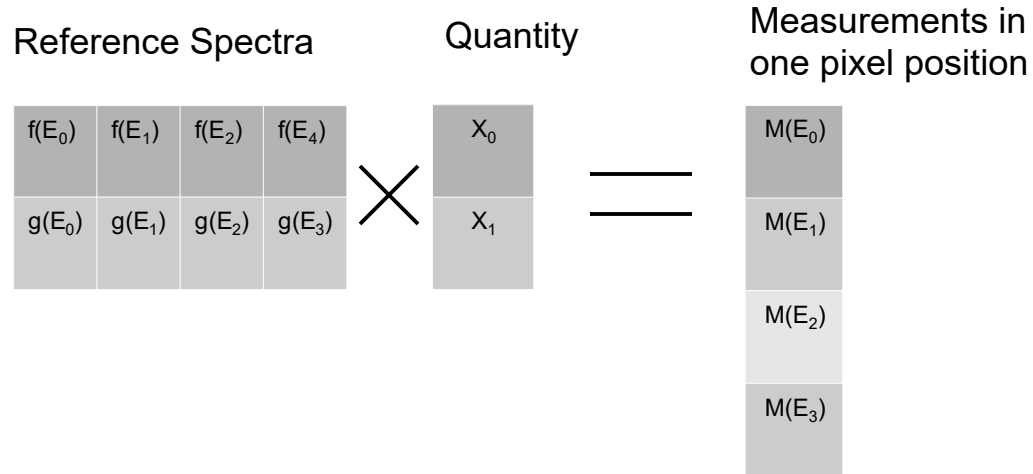
284.7 eV

285.1 eV

320 eV



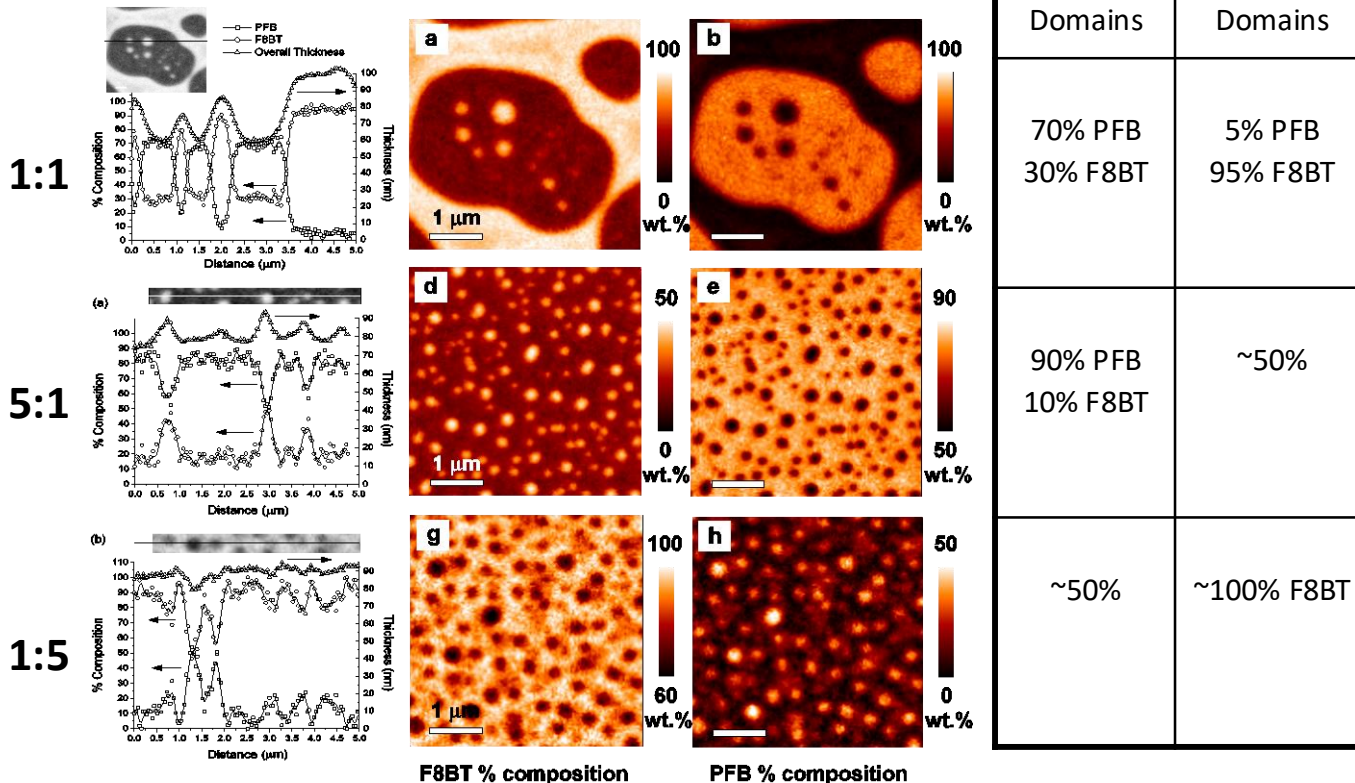
# Singular Value Decomposition



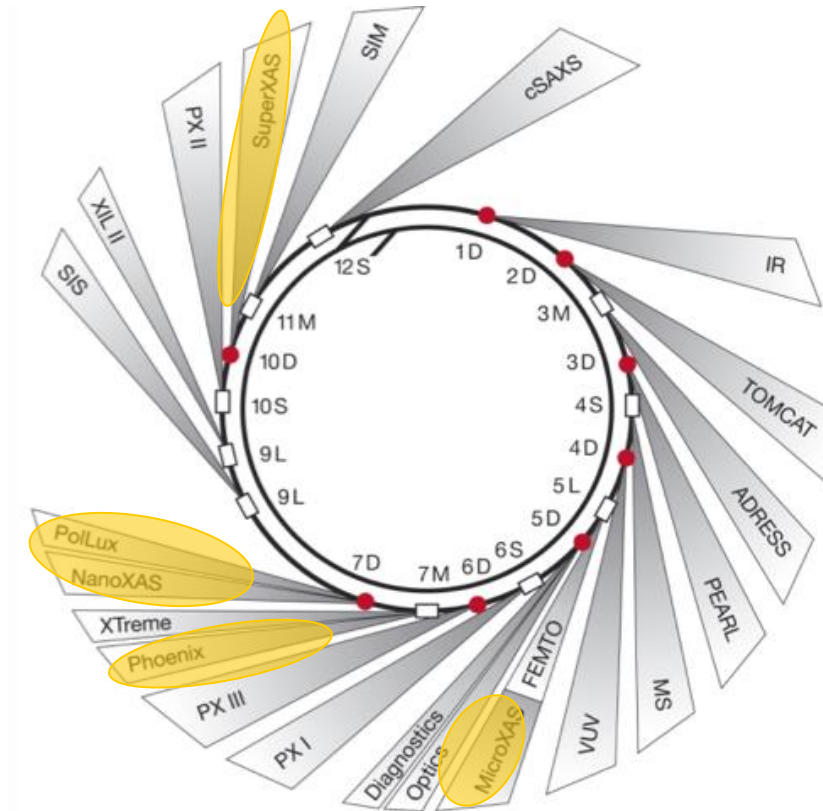
- Solve for X!
- Need at least as many energy measurements as there are material components to solve for.
- Oversampling helps to reject noise.

# Composition maps from SVD

segregated domains are not as pure as it was expected



# XANES/EXAFS @ PSI



# XANES/EXAFS@PSI

C, N, O K-edges, up to Si/P

L-edges  $Z = 22 - 30$ ,  
 important 1<sup>st</sup>-row  
 transition metals

S, Cl, Ca K-edges  
 (biology)

L-edges 2<sup>nd</sup>-row  
 transition metals

K-edges above  $Z = 20$  (Ca)

L-edges above  $Z = 48$  (Cd)



EUV <125 eV  
 VUV

soft  
 X-ray

2keV

tender  
 X-ray

6keV

hard  
 X-ray



# XANES/EXAFS@PSI

C, N, O K-edges, up to Si/P

L-edges  $Z = 22 - 30$ ,  
important 1<sup>st</sup>-row  
transition metals

S, Cl, Ca K-edges  
(biology)

L-edges 2<sup>nd</sup>-row  
transition metals

K-edges above  $Z = 20$  (Ca)

L-edges above  $Z = 48$  (Cd)



EUV <125 eV  
VUV

soft  
X-ray

2keV

tender  
X-ray

6keV

hard  
X-ray  
temporal resolution

nanoXAS/Pollux

PHOENIX

microXAS

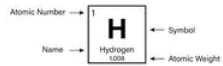
SuperXAS

spatial resolution

# Accessible Elemental X-ray Absorption Edges at Phoenix

## Periodic Table of the Elements

1 <b>H</b> Hydrogen 1.008																	2 <b>He</b> Helium 4.002602						
3 <b>Li</b> Lithium 6.94	4 <b>Be</b> Beryllium 9.0121831																	5 <b>B</b> Boron 10.81	6 <b>C</b> Carbon 12.011	7 <b>N</b> Nitrogen 14.007	8 <b>O</b> Oxygen 15.999	9 <b>F</b> Fluorine 18.99840323	10 <b>Ne</b> Neon 20.1797
11 <b>Na</b> Sodium 22.98976928	12 <b>Mg</b> Magnesium 24.305																	13 <b>Al</b> Aluminum 26.9815386	14 <b>Si</b> Silicon 28.086	15 <b>P</b> Phosphorus 30.973762	16 <b>S</b> Sulfur 32.06	17 <b>Cl</b> Chlorine 35.45	18 <b>Ar</b> Argon 39.948
19 <b>K</b> Potassium 39.0983	20 <b>Ca</b> Calcium 40.078	21 <b>Sc</b> Scandium 44.955912	22 <b>Ti</b> Titanium 47.88	23 <b>V</b> Vanadium 50.9415	24 <b>Cr</b> Chromium 51.9961	25 <b>Mn</b> Manganese 54.938044	26 <b>Fe</b> Iron 55.845	27 <b>Co</b> Cobalt 58.933194	28 <b>Ni</b> Nickel 58.6934	29 <b>Cu</b> Copper 63.546	30 <b>Zn</b> Zinc 65.38	31 <b>Ga</b> Gallium 69.723	32 <b>Ge</b> Germanium 72.630	33 <b>As</b> Arsenic 74.9216	34 <b>Se</b> Selenium 78.96	35 <b>Br</b> Bromine 79.904	36 <b>Kr</b> Krypton 83.798						
37 <b>Rb</b> Rubidium 85.468	38 <b>Sr</b> Strontium 87.62	39 <b>Y</b> Yttrium 88.90584	40 <b>Zr</b> Zirconium 91.224	41 <b>Nb</b> Niobium 92.90638	42 <b>Mo</b> Molybdenum 95.94	43 <b>Tc</b> Technetium (98)	44 <b>Ru</b> Ruthenium 101.07	45 <b>Rh</b> Rhodium 102.90550	46 <b>Pd</b> Palladium 106.42	47 <b>Ag</b> Silver 107.8682	48 <b>Cd</b> Cadmium 112.414	49 <b>In</b> Indium 114.818	50 <b>Sn</b> Tin 118.710	51 <b>Sb</b> Antimony 121.757	52 <b>Te</b> Tellurium 127.60	53 <b>I</b> Iodine 126.905	54 <b>Xe</b> Xenon 131.29						
55 <b>Cs</b> Cesium 132.90545196	56 <b>Ba</b> Barium 137.327	57 - 71 Lanthanoids	72 <b>Hf</b> Hafnium 178.49	73 <b>Ta</b> Tantalum 180.94788	74 <b>W</b> Tungsten 183.84	75 <b>Re</b> Rhenium 186.207	76 <b>Os</b> Osmium 190.23	77 <b>Ir</b> Iridium 192.222	78 <b>Pt</b> Platinum 195.084	79 <b>Au</b> Gold 196.966569	80 <b>Hg</b> Mercury 200.592	81 <b>Tl</b> Thallium 204.38	82 <b>Pb</b> Lead 207.2	83 <b>Bi</b> Bismuth 208.9804	84 <b>Po</b> Polonium 209	85 <b>At</b> Astatine 210	86 <b>Rn</b> Radon 222						
87 <b>Fr</b> Francium 223	88 <b>Ra</b> Radium 226	89 - 103 Actinoids	104 <b>Rf</b> Rutherfordium 261	105 <b>Db</b> Dubnium 268	106 <b>Sg</b> Seaborgium 269	107 <b>Bh</b> Bohrium 270	108 <b>Hs</b> Hassium 270	109 <b>Mt</b> Meitnerium 276	110 <b>Ds</b> Darmstadtium 281	111 <b>Rg</b> Roentgenium 282	112 <b>Cn</b> Copernicium 285	113 <b>Nh</b> Nihonium 286	114 <b>Fl</b> Flerovium 289	115 <b>Mc</b> Moscovium 289	116 <b>Lv</b> Livermorium 293	117 <b>Ts</b> Tennessine 294	118 <b>Og</b> Oganesson 294						



57 <b>La</b> Lanthanum 138.90547	58 <b>Ce</b> Cerium 140.12	59 <b>Pr</b> Praseodymium 140.90766	60 <b>Nd</b> Neodymium 144.242	61 <b>Pm</b> Promethium (147)	62 <b>Sm</b> Samarium 150.36	63 <b>Eu</b> Europium 151.964	64 <b>Gd</b> Gadolinium 157.25	65 <b>Tb</b> Terbium 158.92535	66 <b>Dy</b> Dysprosium 162.500	67 <b>Ho</b> Holmium 164.93032	68 <b>Er</b> Erbium 167.259	69 <b>Tm</b> Thulium 168.93047	70 <b>Yb</b> Ytterbium 173.045	71 <b>Lu</b> Lutetium 174.967
89 <b>Ac</b> Actinium 227	90 <b>Th</b> Thorium 232.0377	91 <b>Pa</b> Protactinium 231.03688	92 <b>U</b> Uranium 238.02891	93 <b>Np</b> Neptunium 237	94 <b>Pu</b> Plutonium 244	95 <b>Am</b> Americium 243	96 <b>Cm</b> Curium 247	97 <b>Bk</b> Berkelium 247	98 <b>Cf</b> Californium 251	99 <b>Es</b> Einsteinium 252	100 <b>Fm</b> Fermium 257	101 <b>Md</b> Mendelevium 258	102 <b>No</b> Nobelium 259	103 <b>Lr</b> Lawrencium 260

K-edge

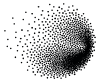
L-edge

M-edge

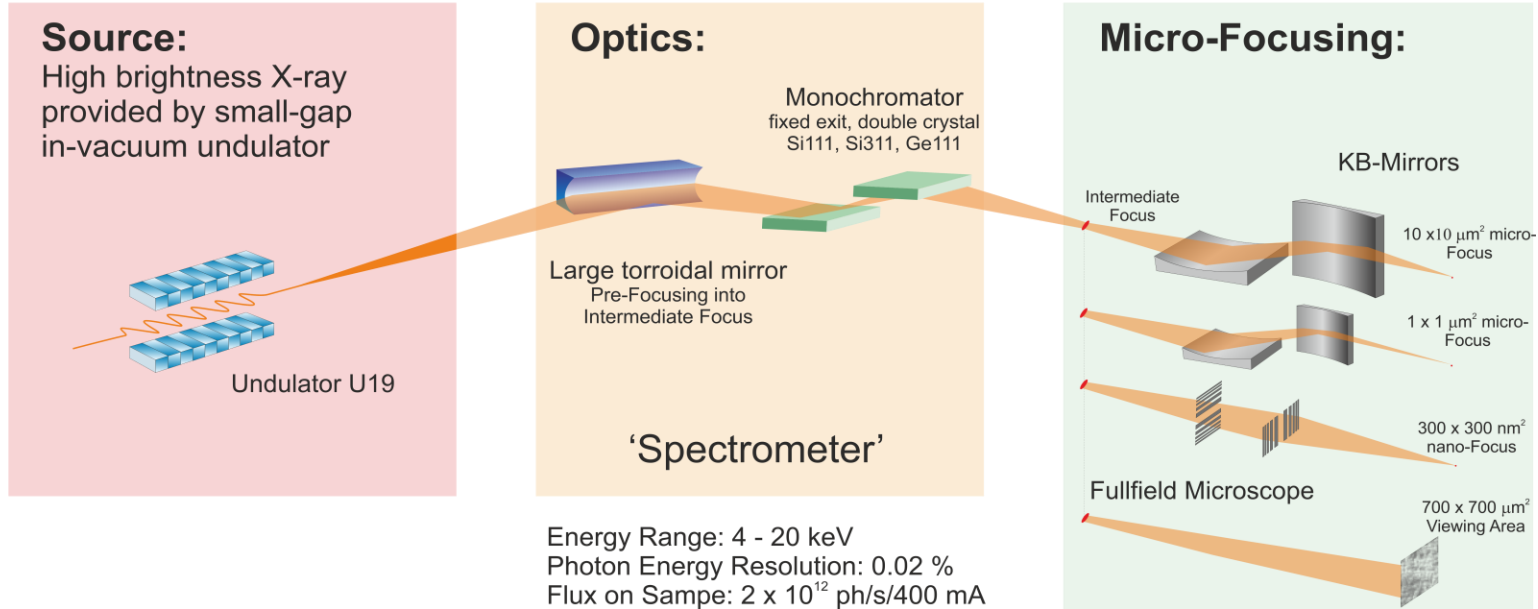
# MicroXAS



MSE435 - Marianne Liebi



# MicroXAS



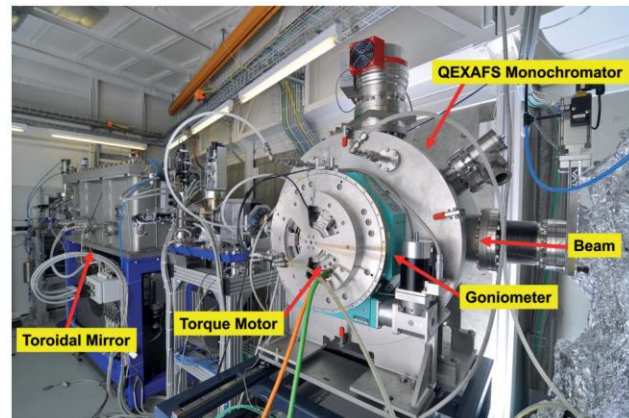
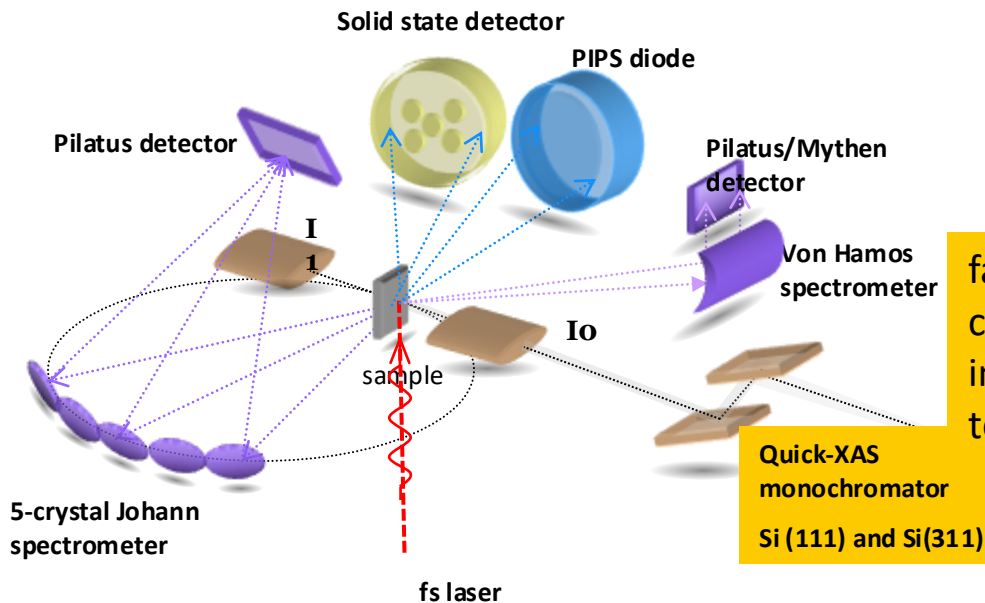
# SuperXAS: Hard X-ray spectroscopy

SuperXAS

Energy range: 4.5-35 keV

Flux : up to  $1 \times 10^{12}$  ph/s (@ 12 keV)

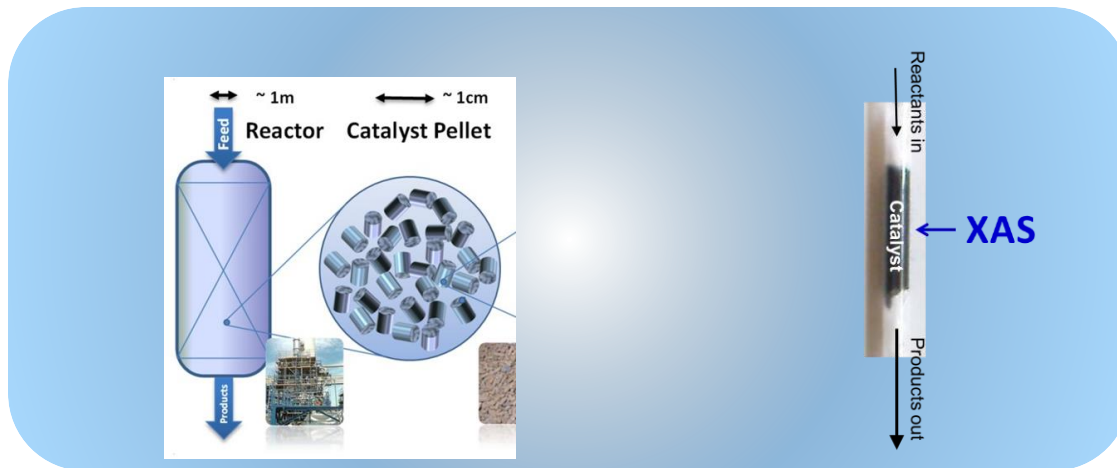
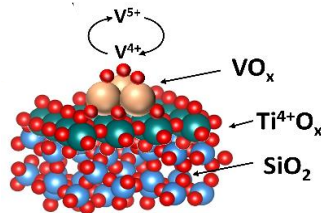
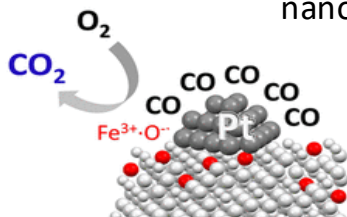
Beam size: from  $100 \times 100 \mu\text{m}^2$  to  $5000 \times 500 \mu\text{m}^2$



fast energy scan! (full spectra in s)  
chemical local structure transformed  
in process, in-situ, in-operando  
temperature, gas atmosphere, etc.

# Catalytic Materials

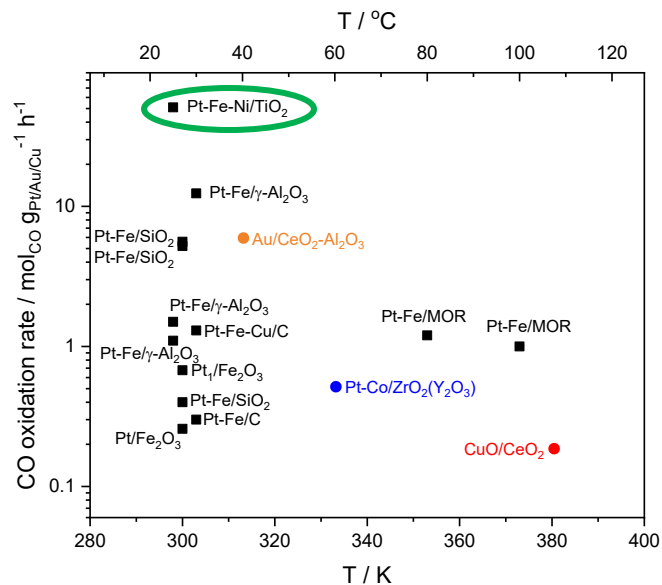
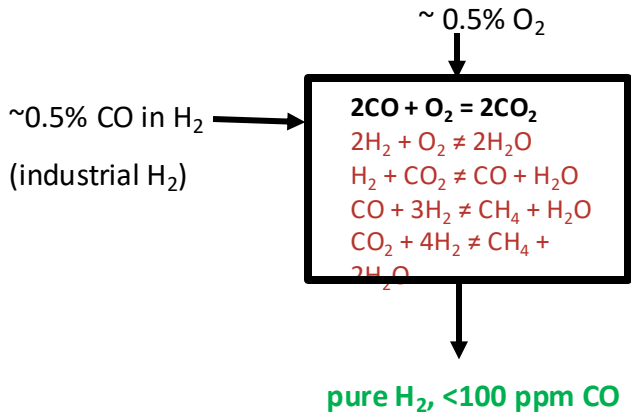
nanoparticles, supported oxides, interfaces...



catalysis: complex materials with several elements: which part of the material is working and in which oxidation state

# Supported Pt-FeO<sub>x</sub> Catalysts for Preferential CO Oxidation (PROX) for H<sub>2</sub> Purification

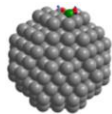
Hydrogen stream: oxidation of CO without oxidation of H<sub>2</sub>



why are some much more active? what is the active site

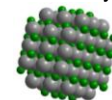
# Active Sites in Pt-FeO<sub>x</sub> System Proposed in the Literature

Atomically-dispersed Fe<sup>+3</sup>(OH)<sub>x</sub> / Pt



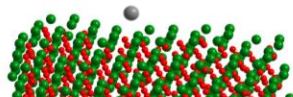
Cao L. et al, *Nature*, **2019**, 565

Pt-Fe<sup>0</sup> alloy



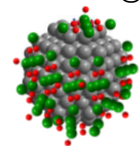
Yin J. et al, *Catal. Lett.*, **2008**, 325

Atomically-dispersed Pt / Fe<sup>+3</sup><sub>2</sub>O<sub>3</sub>



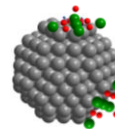
Fu Q. et al, *Science*, **2010**, 328

Core-shell Pt@FeO<sub>x</sub>



Da Silva T.L. et al, *Mater. Res. Expr.*,  
**2018**, 6

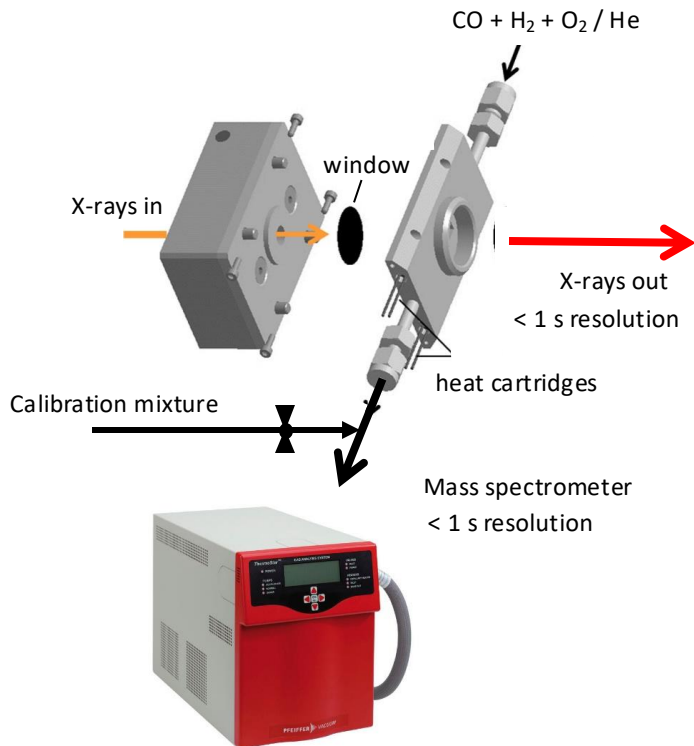
Fe<sup>+2</sup>O islands on Pt



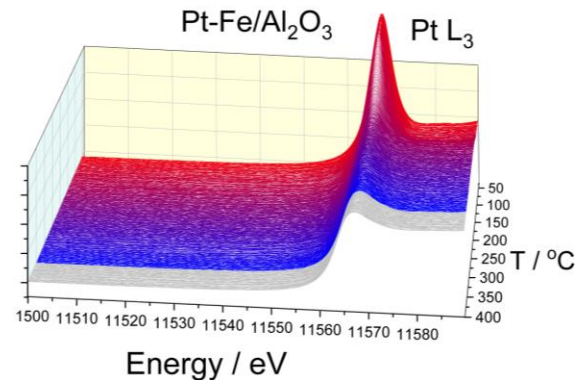
Kudematsch W. et al, *ACS Nano*,  
**2015**, 9

?

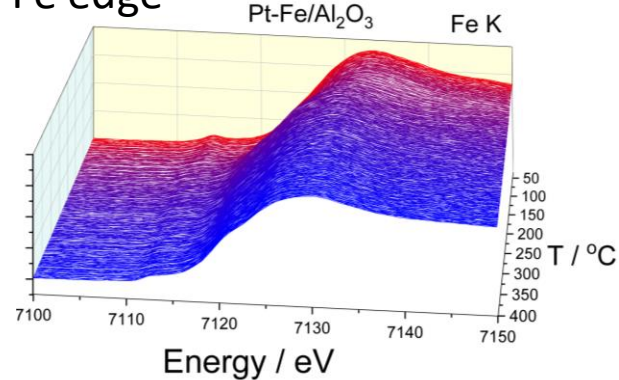
# Operando XAS Study of Pt-FeO<sub>x</sub> catalysts



Pt edge

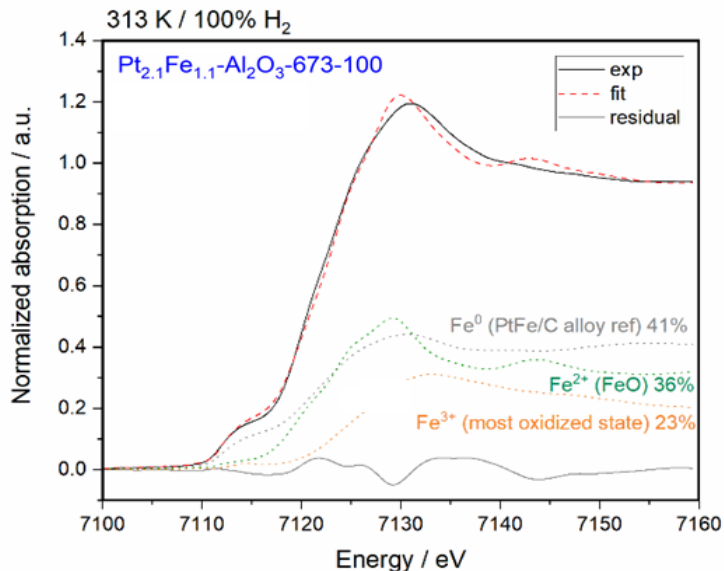


Fe edge



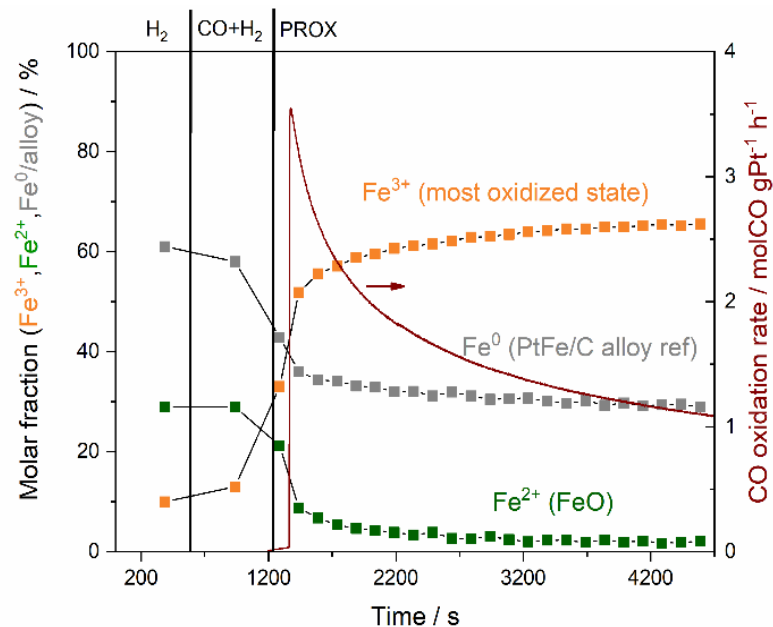
# Supported Pt-FeO<sub>x</sub> Catalysts: Operando XAS

Operando Fe K-edge XAS

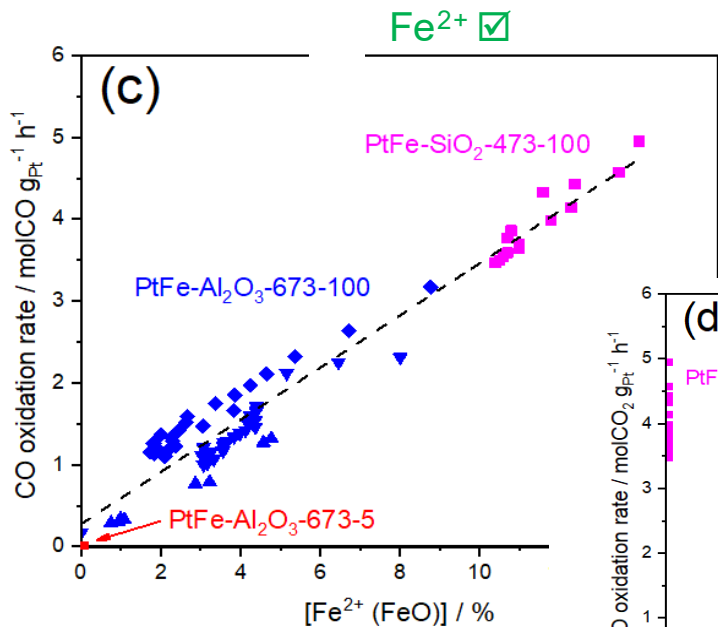


fit: linear combination of components

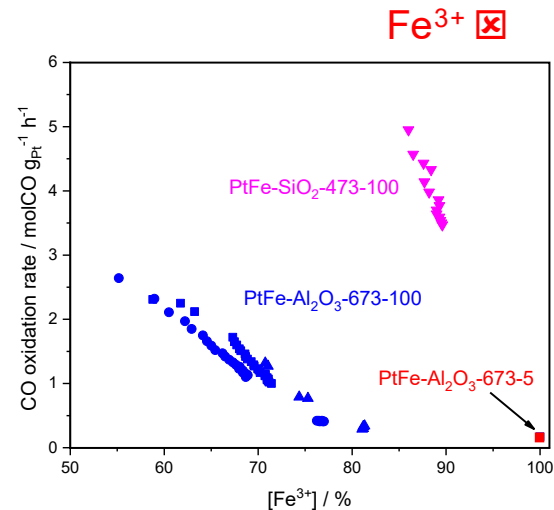
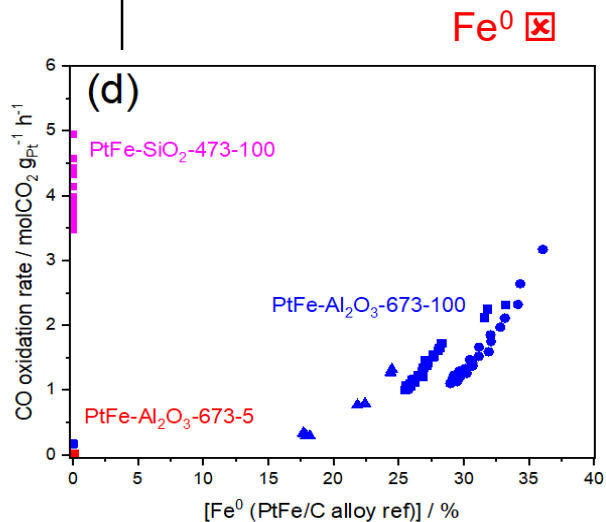
Fe speciation - PROX activity correlation (mass spectrometer)



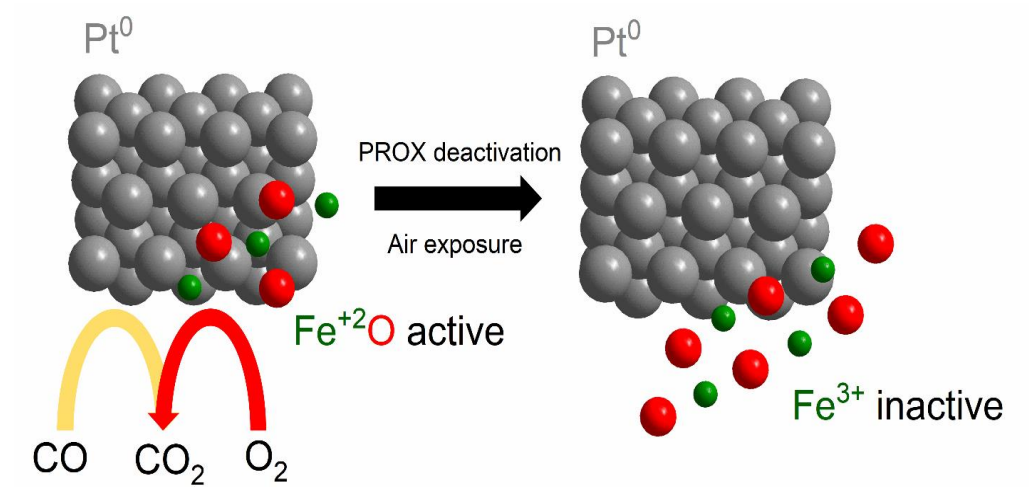
# Supported Pt-FeO<sub>x</sub> Catalysts: Activity vs. Fe species



different catalysts with different support  
→ clear correlation with Fe<sup>2+</sup>, negative correlation with Fe<sup>3+</sup>

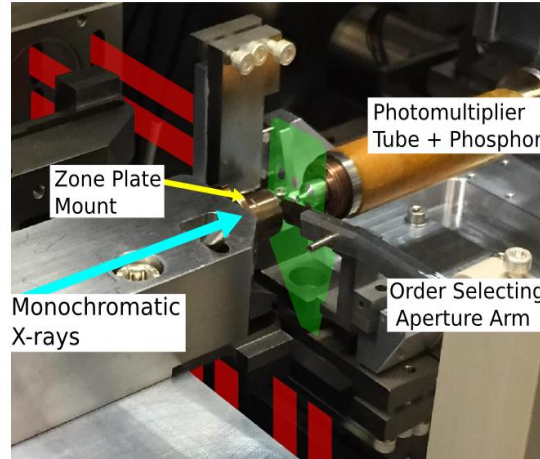
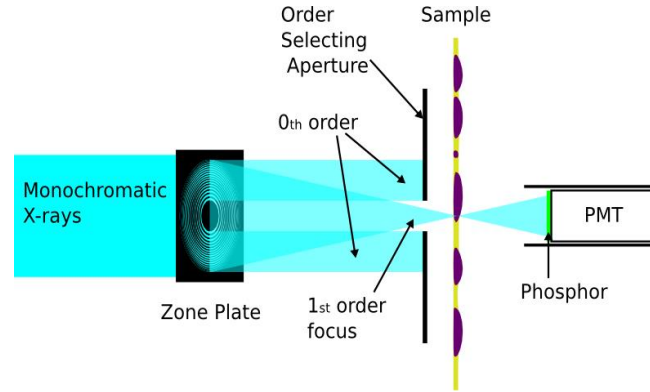


# Supported Pt-FeO<sub>x</sub> Catalysts: Active site and its deactivation



*I. Sadykov et al. Angew. Chem., 2023, 135, e2022140*

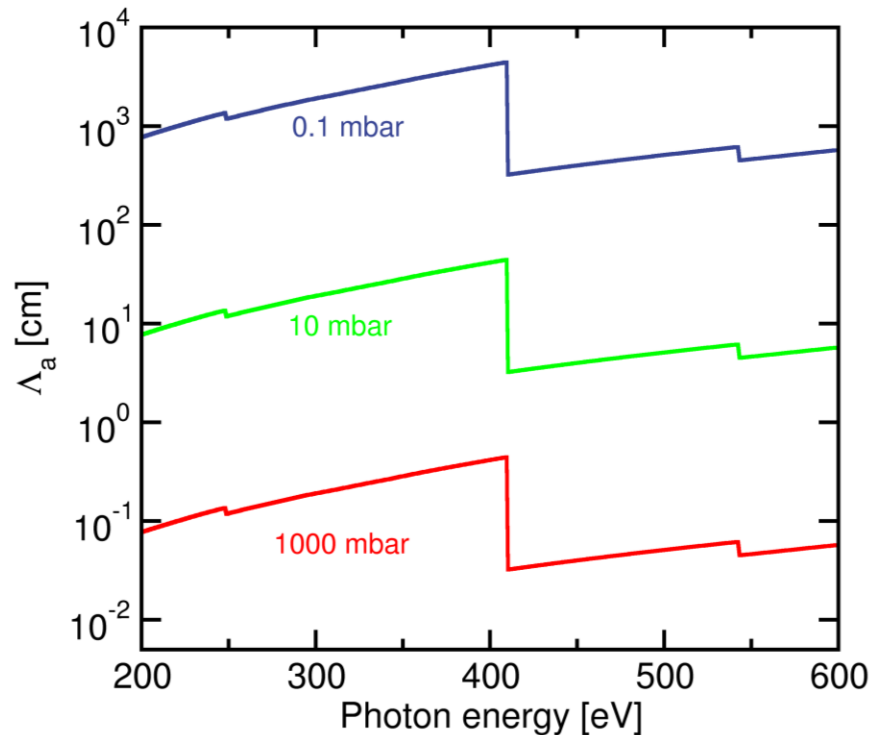
# Pollux



# Accessible Elemental X-ray Absorption Edges at Pollux

1 H																	2 He																
3 Li	4 Be													5 B	6 C	7 N	8 O	9 F	10 Ne														
11 Na	12 Mg													13 Al	14 Si	15 P	16 S	17 Cl	18 Ar														
19 K	20 Ca	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr																
37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe																
55 Cs	56 Ba			72 Hf	73 Ta	74 W	75 Re	76 Os	77 Ir	78 Pt	79 Au	80 Hg	81 Tl	82 Pb	83 Bi	84 Po	85 At	86 Rn															
87 Fr	88 Ra			104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111 Rg	112 Cn	113 Uut	114 Fl	115 Uup	116 Lv	117 Uus	118 Uuo															
																			57 La	58 Ce	59 Pr	60 Nd	61 Pm	62 Sm	63 Eu	64 Gd	65 Tb	66 Dy	67 Ho	68 Er	69 Tm	70 Yb	71 Lu
																			89 Ac	90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr

K-edge  
L-edge  
M-edge



Synchrotron-only technique

Scan photon energy

Strict vacuum requirements between source and sample/detector

Transmission for 1-mm air = 6% above N-edge @ 410 eV

Typically requires 0.1 mbar or better

Long-term accumulation of carbon contamination on optics (cryo worse!)

Compromises “real” sample C-XANES

Incident radiation “cracks” CO<sub>2</sub> on surface

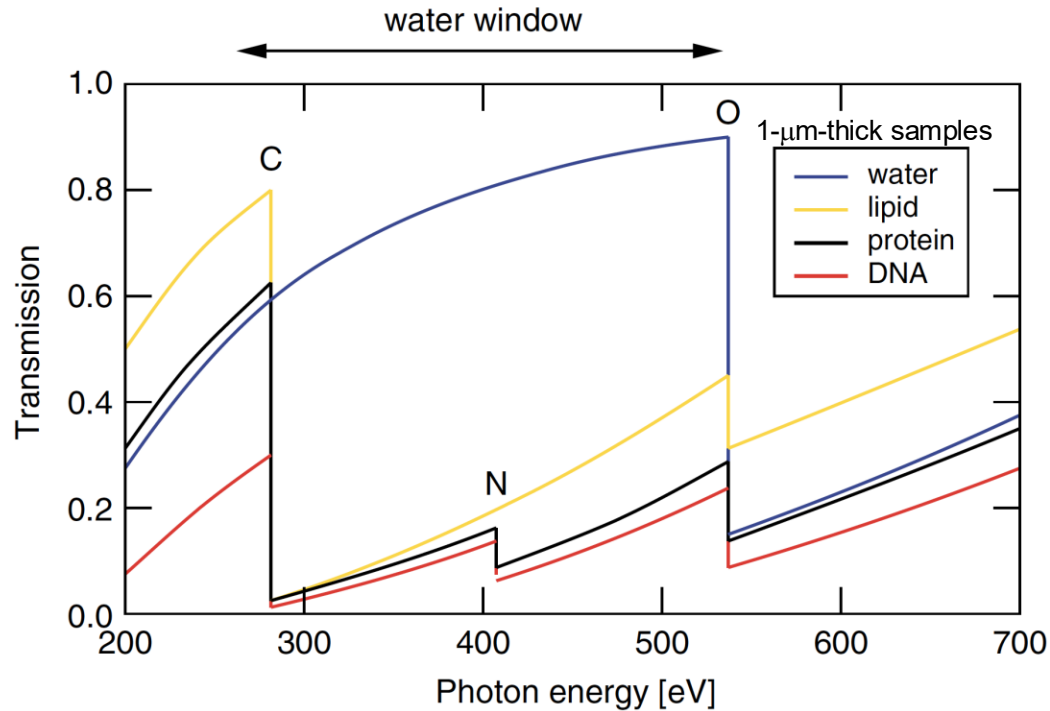
Remove using low-pressure O<sub>2</sub> leak

FZP

Focal length  $\propto h\nu$ : requires axial scanning

OSA

# The water window



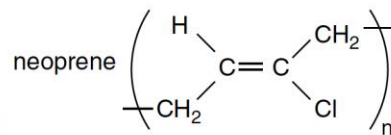
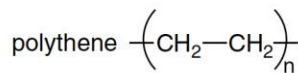
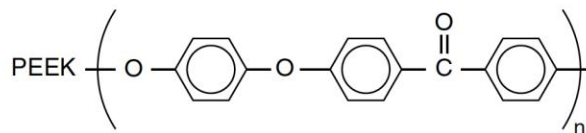
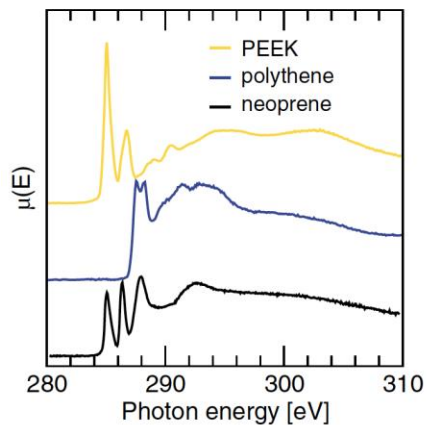
Water transparent below oxygen K-edge down to ca. 200 eV

At lower photon energies, transmission drops due to  $L_a \propto (hn)^3$

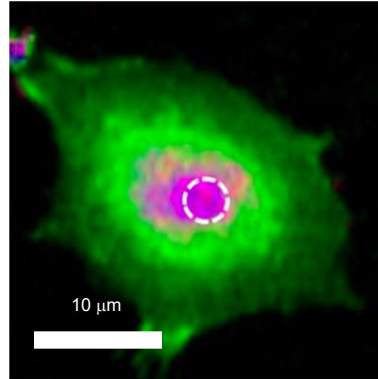
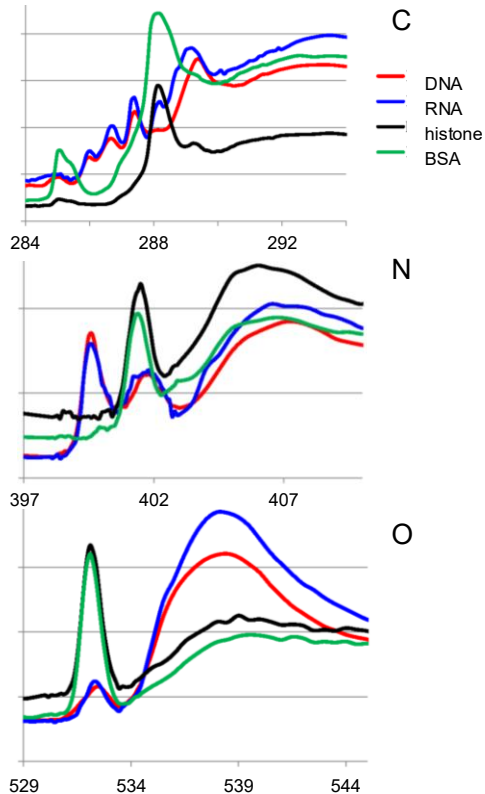
Organic compounds containing C, N, O have high absorption contrast

L-edges of other biologically relevant elements accessible, especially K, Ca

# The water window: polymers



# The water window: cells/biological samples

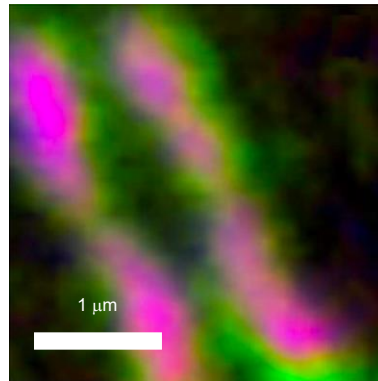


In-vivo or cryo-experiments possible

DNA, RNA, histone, protein

Easily distinguishable (c.f. hard x-ray phase-contrast methods)

cell: DNA, proteins other than histone, and histone are displayed as red, green, and blue, respectively



chromosome: DNA, RNA, and histone are displayed as red, green, and blue, respectively.