

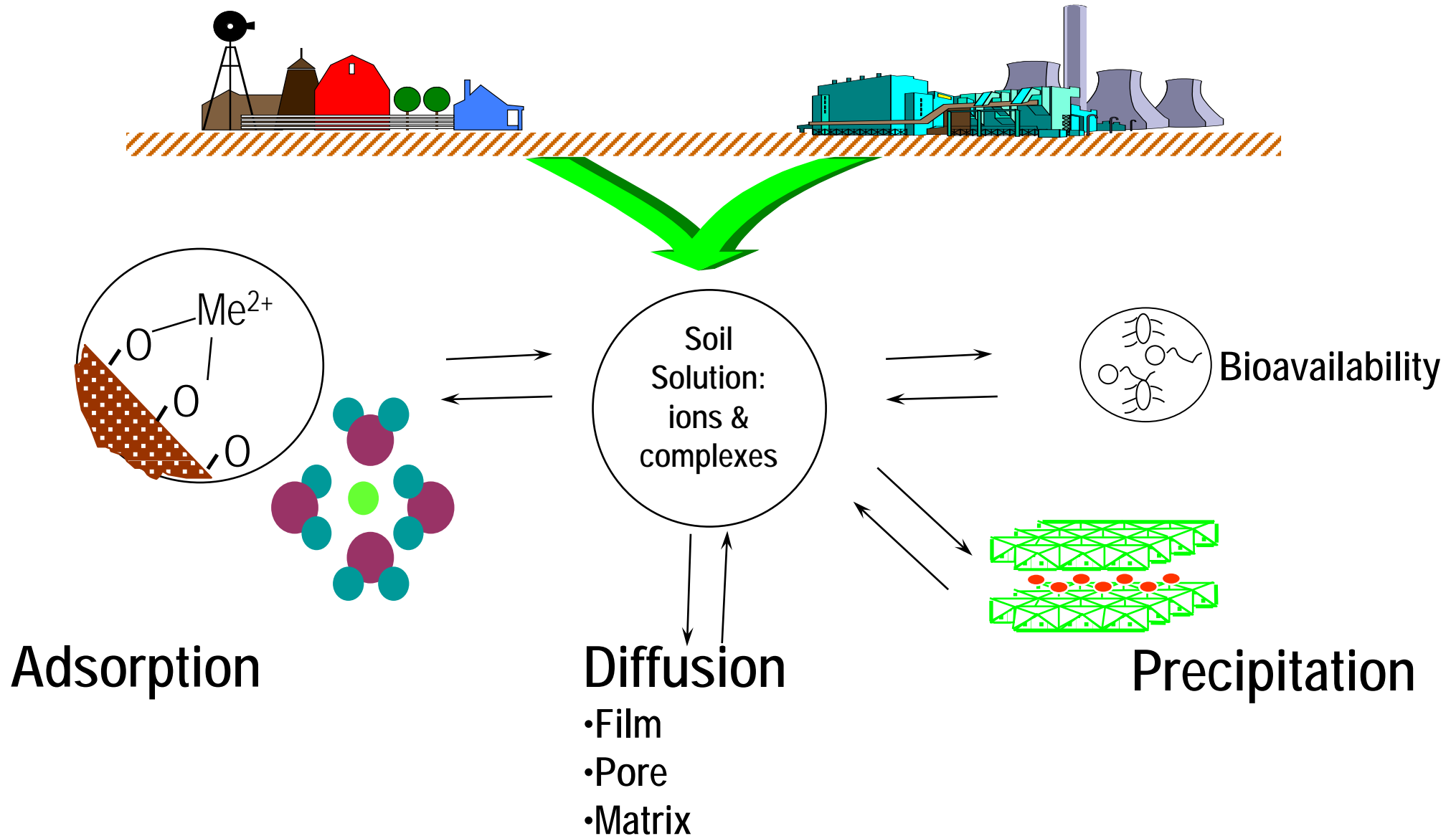


Wir schaffen Wissen – heute für morgen

Introduction to EXAFS practical – scientific problem to be analysed

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Paul Scherrer Institute

Introduction of Metals into Soil Environments: Primary minerals, Agriculture, Industry, Sewage sludge, etc



- Iron (Fe) - abundant in earth crust, mostly as iron (hydr)oxides
- 16 different (hydr)oxides, mostly formed as weathering products
- Often nano-sized crystals with high surface area – most reactive sorbents for contaminants in the environment



We will look at the structure of:

Ferrihydrite

Very common iron hydroxide

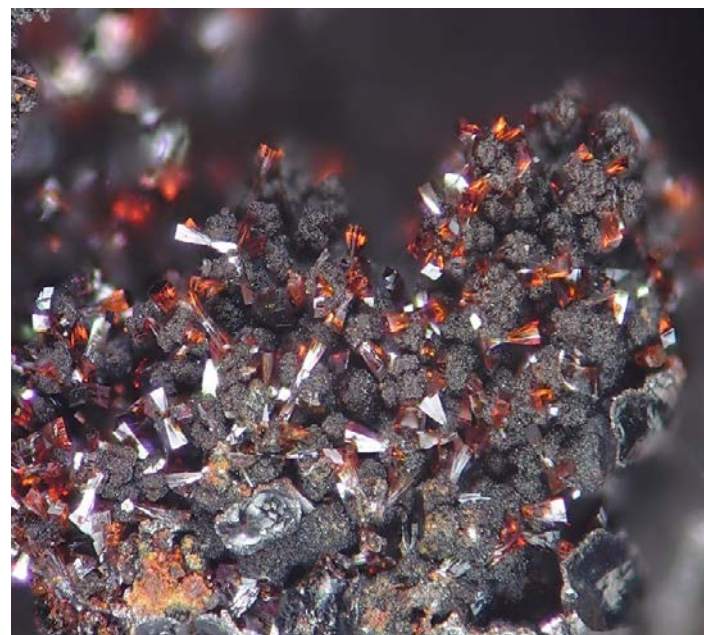
Poorly ordered



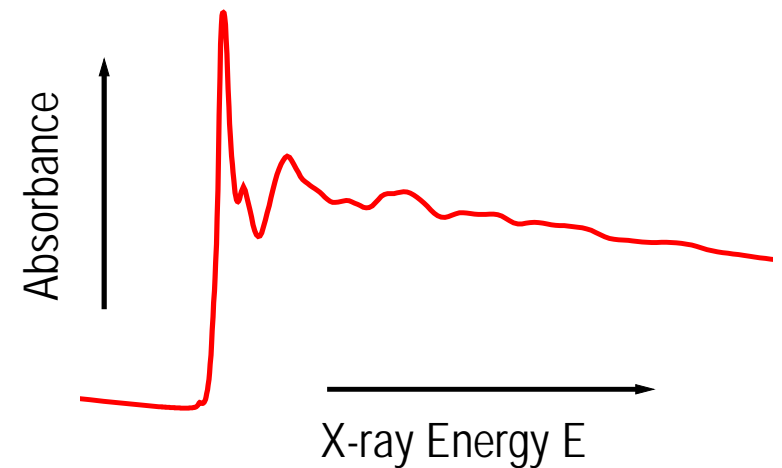
Lepidocrocite

Orthorhombic crystal structure

Well-crystalized



- ✓ Detailed chemical und structural information (oxidation state, coordination numbers, bond distances, system disorder)
 - Solution species
 - Crystalline and amorphous solids
 - Surface complexes
- ✓ in-situ, non destructiv
- ✓ minimal sample preparation
- ✓ high selectivity/sensitivity (few ppm)



X-ray absorption spectroscopy

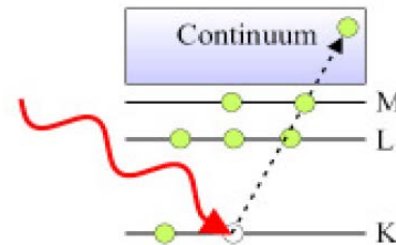
Visible light
 $\lambda \sim 0.5 \mu\text{m}$
 $E \sim 2 \text{ eV}$



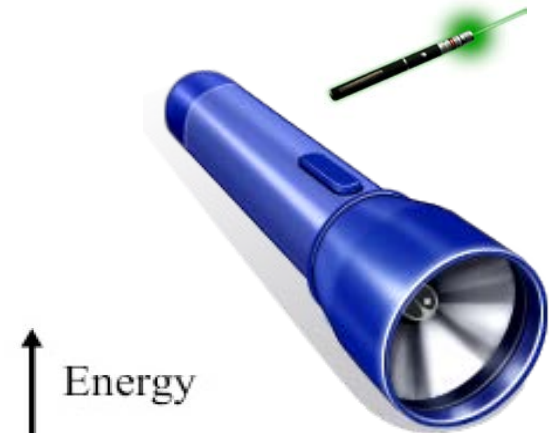
X-ray light
 $\lambda \sim 1 \text{ \AA} (=0.1 \text{ nm})$
 $E \sim 10 \text{ keV}$



valence
electrons



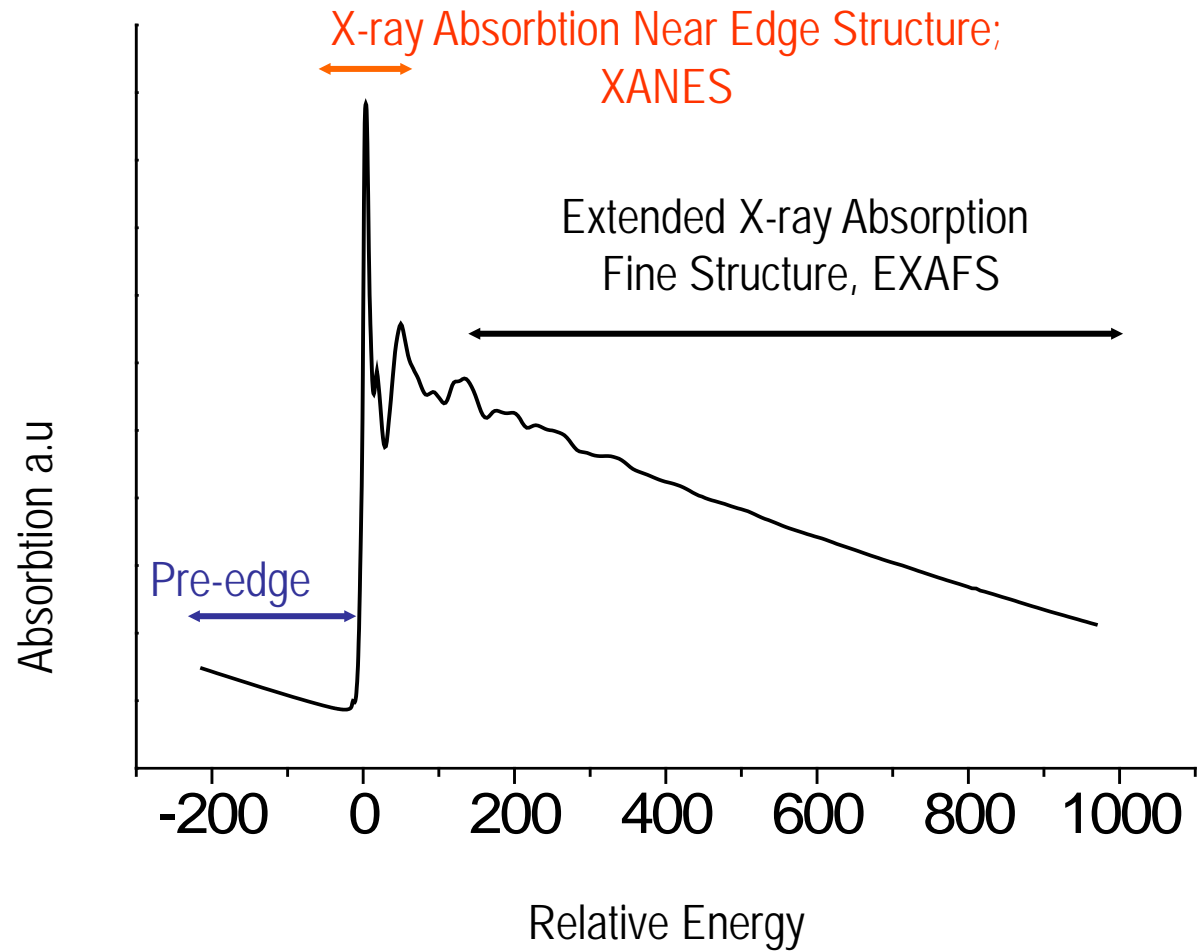
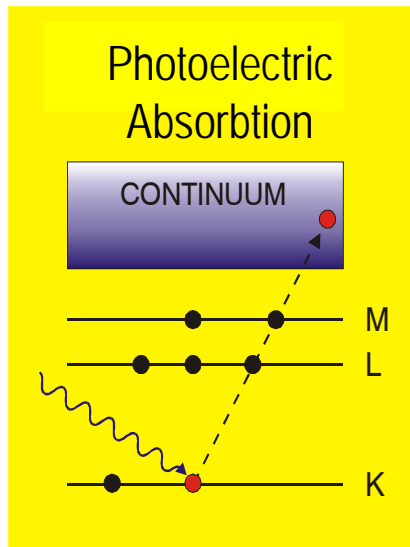
core
electrons



Energy ↑



Main Principle



Pre-edge

- $E < E_b$
- transition of e^- from ground state (e.g., $1s, 2s$) to empty or partly filled, excited states (nd orbital)

- selection rules for e^- transitions

- speciation of Cr(VI) and Cr(III)

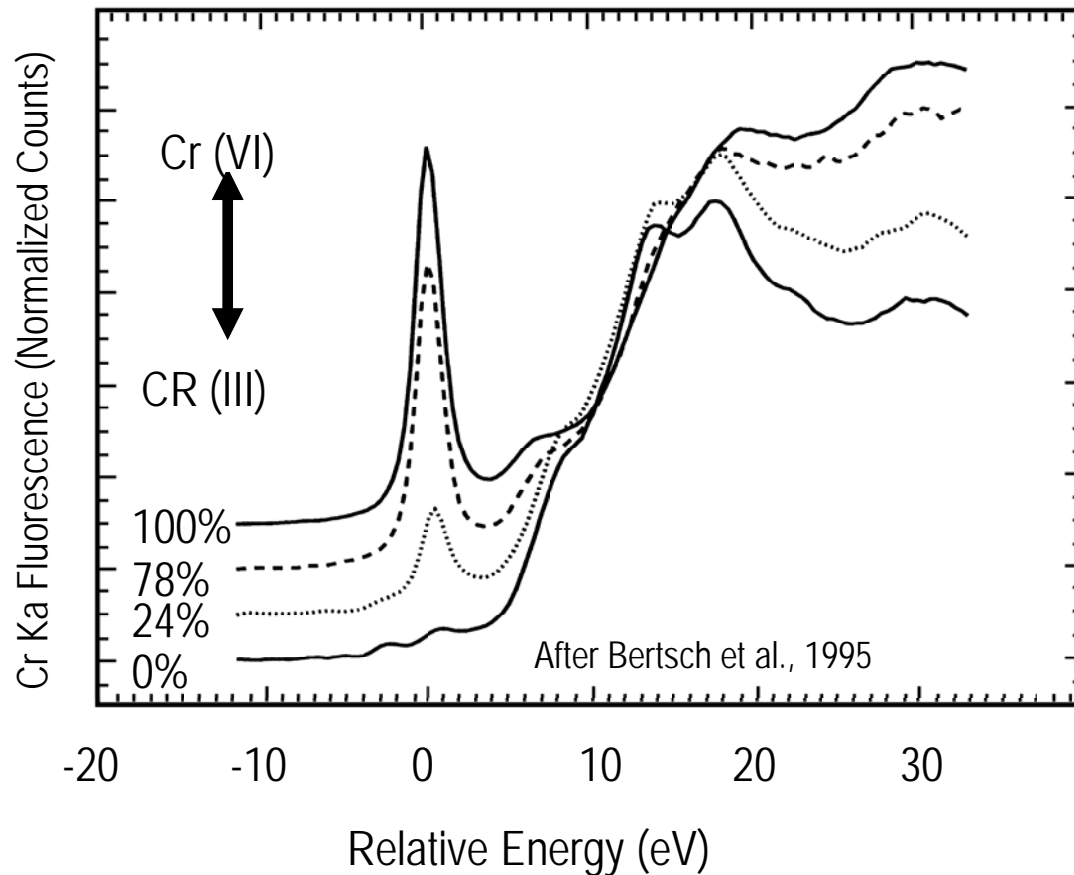
- Cr(VI)

- toxic and mobil

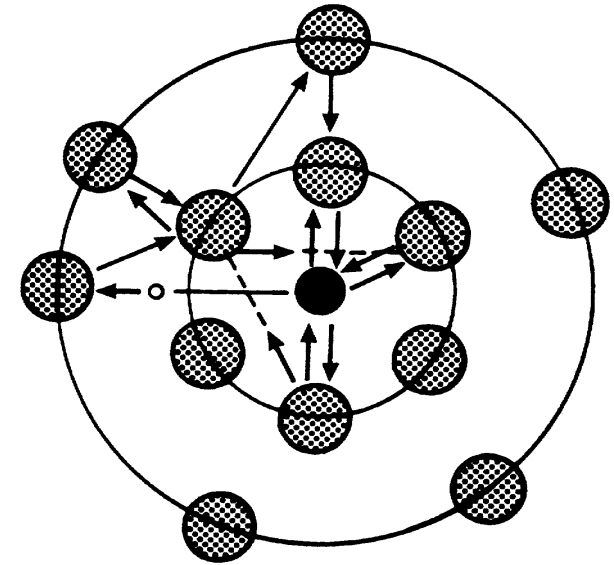
- Cr(III)

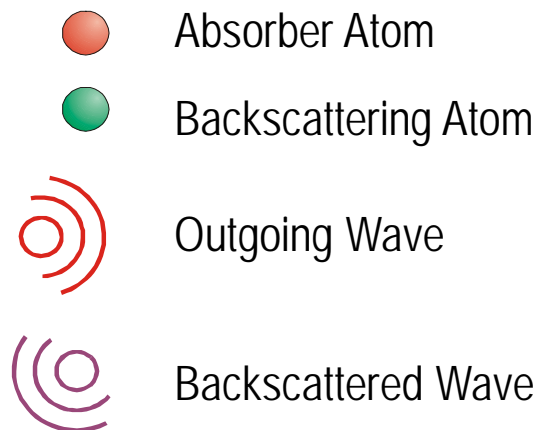
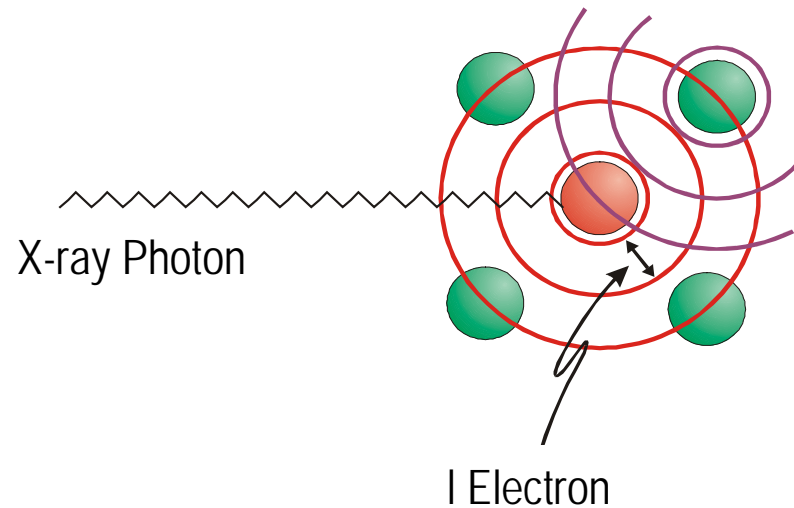
- hardly toxic

- sorbed or incorporated into mineral phases



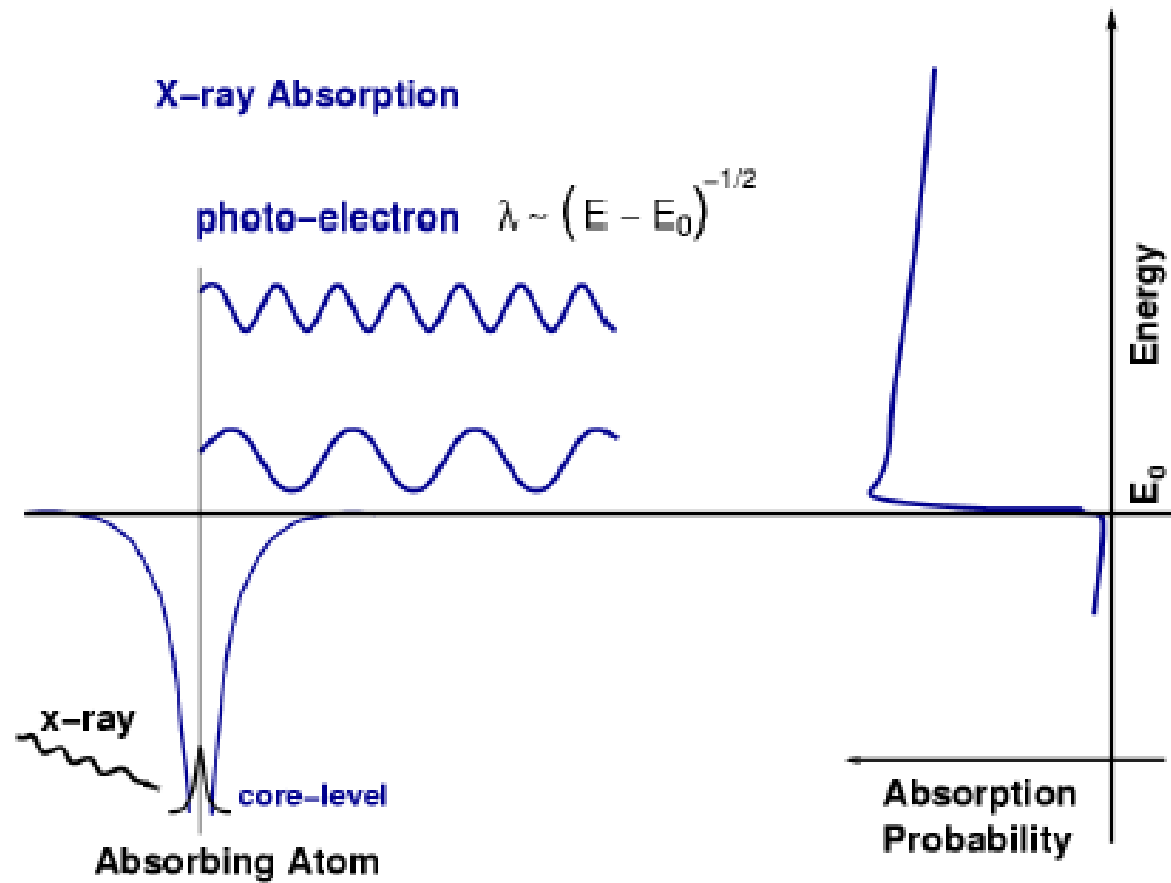
- $E \sim E_b$
- Multiple scattering of photoelectron
- Very intense and complicate resonance features: fingerprinting and theoretial calculations
- Energy of absorption edge depends on oxidation state
 - o Chemical shift of 1-3 eV for each withdrawing e^-



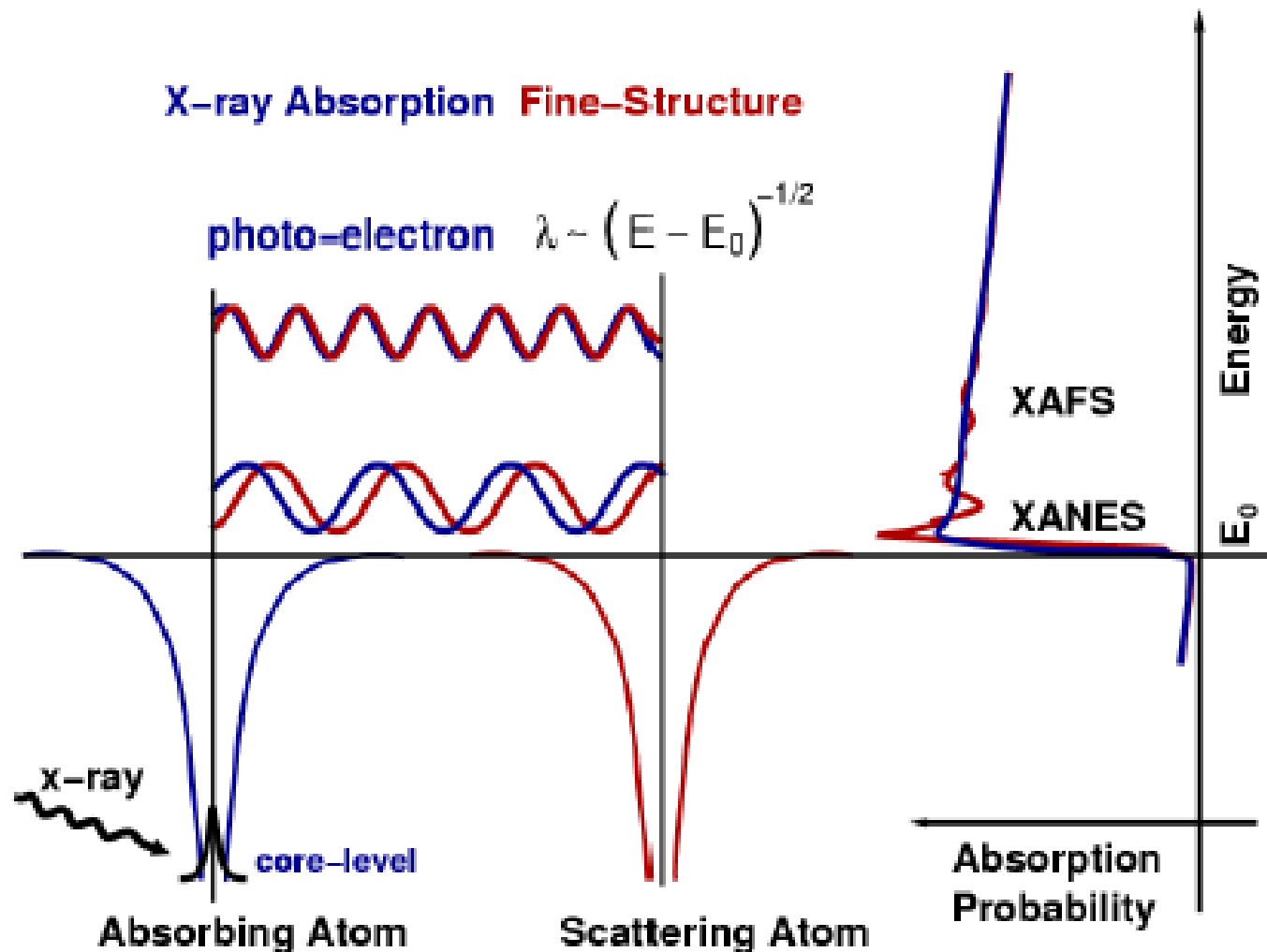


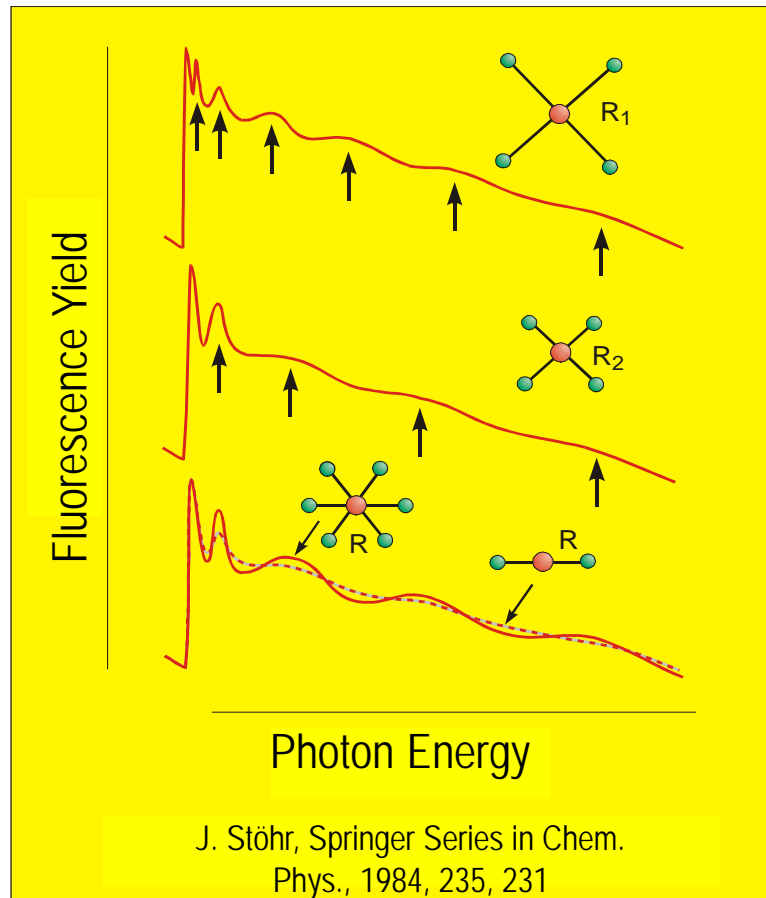
- $E > E_b$
- 50 - 1000 eV above absorption edge
- simplified illustration: constructive & destructive frequencies from the outgoing photoelectron

Free atom



Cluster of atoms

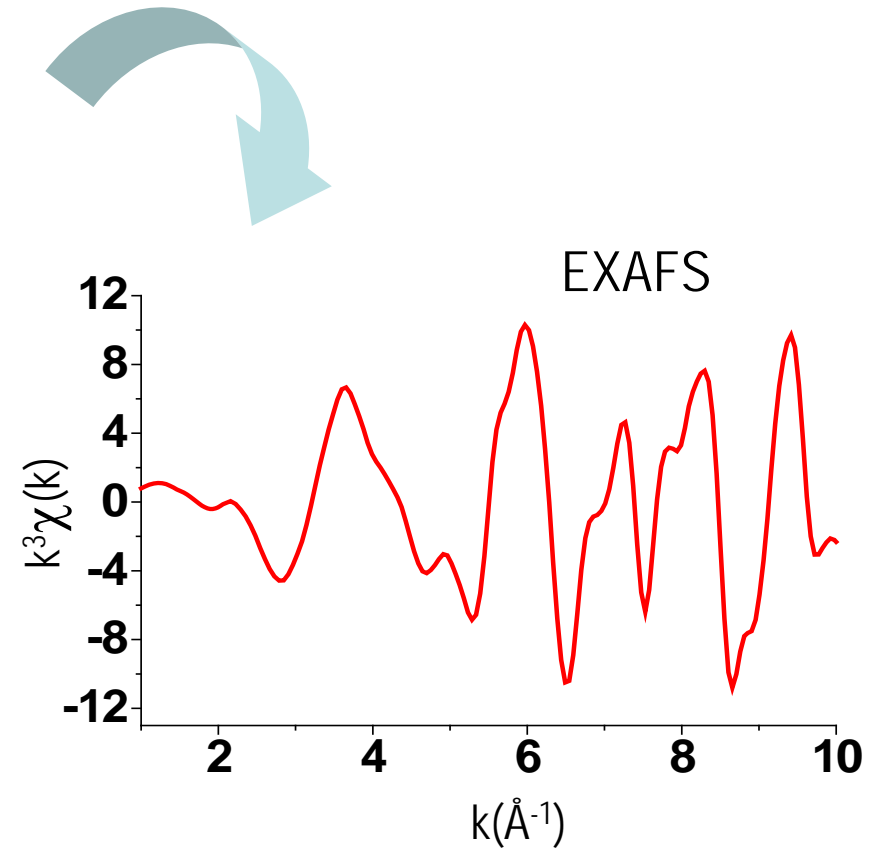
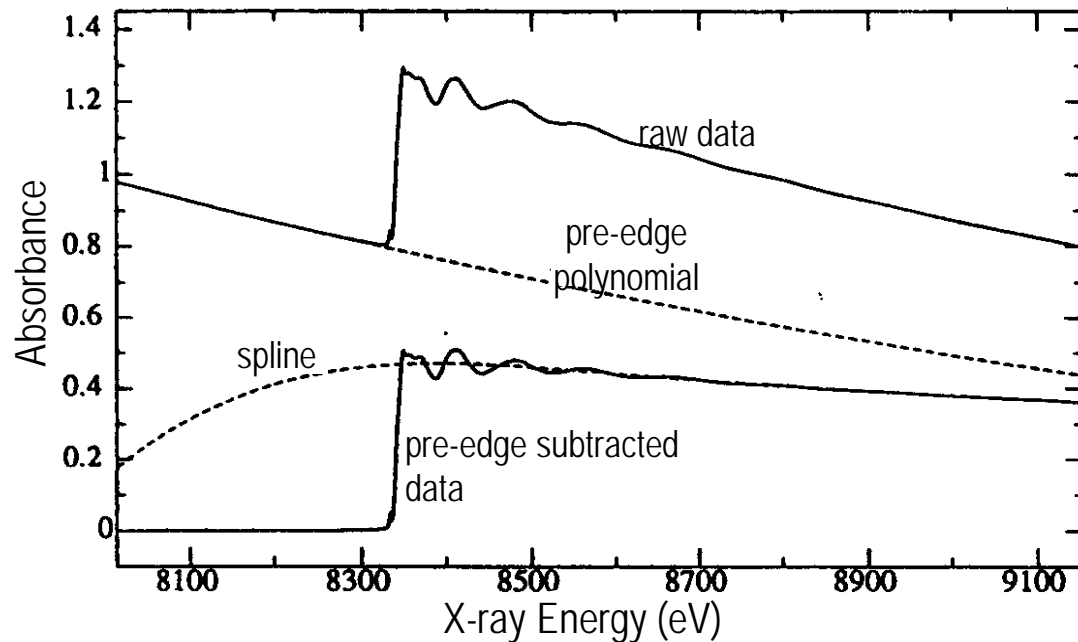




➤ Interference pattern (EXAFS)

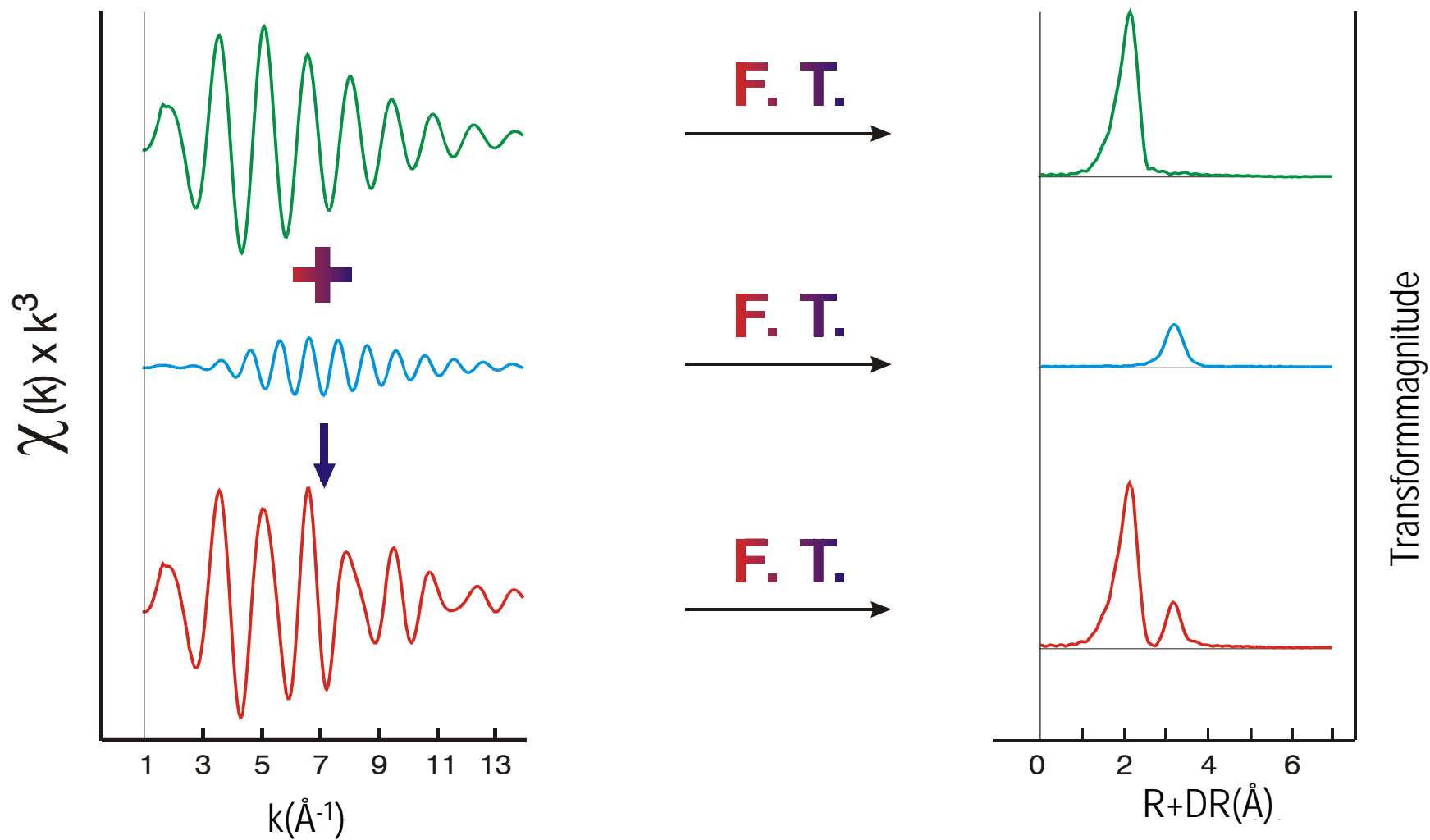
- Frequency correlated to bond distance
- Amplitude correlated to coordination number and identity

$$\chi(k) = \sum a(k) \sin[2kR + \alpha(k)]$$



- extraction of EXAFS
 - pre-edge & spline, normalization
 - conversion to wave vector $k^2 = 2m (E - E_0) / \hbar^2$
 - normally weighted by k^3

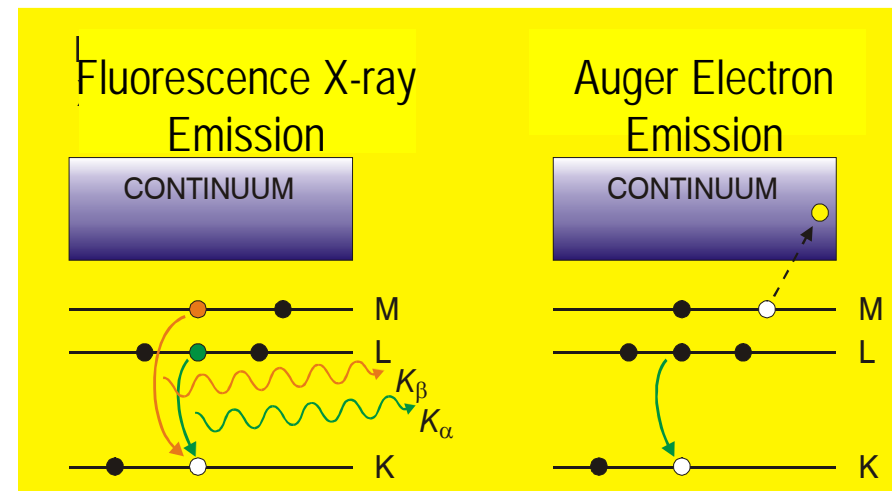
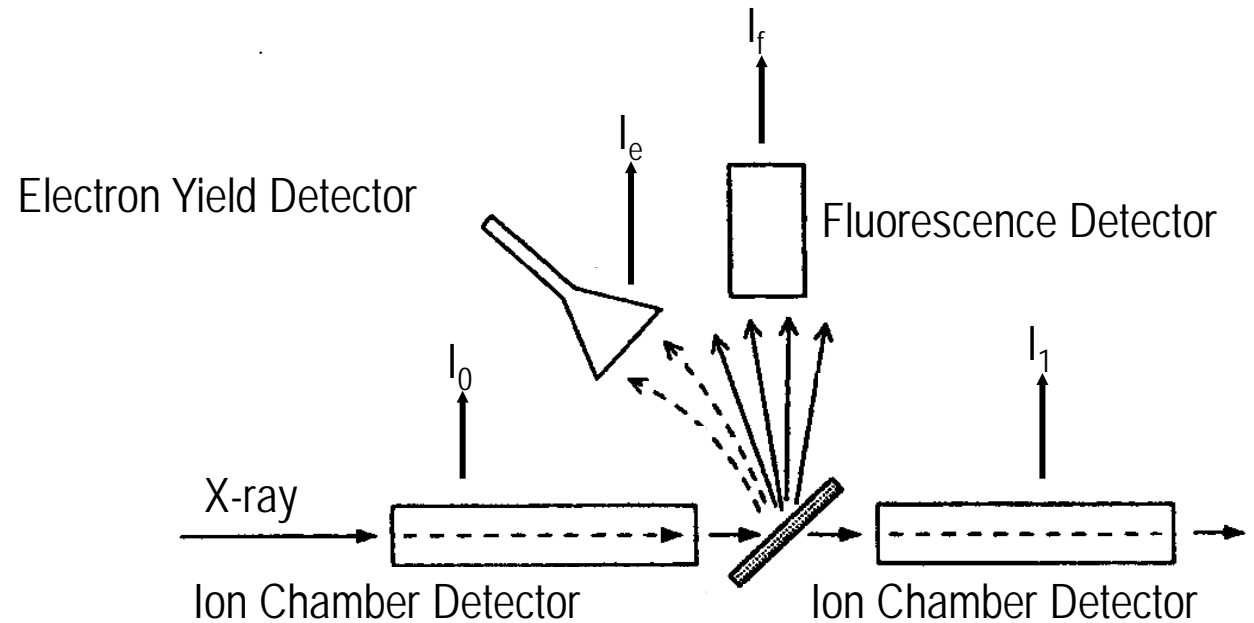
Fourier Transformation

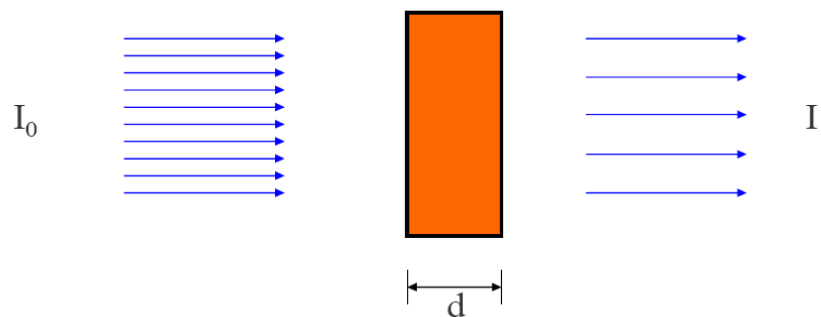


$$\chi(k) = \sum a(k) \sin[2kR + \alpha(k)]$$

Experimental set-up

- Transmission
 - $A = mx = \ln(I_0/I_1)$
 - concentrated samples
- Fluorescence
 - $A = mx = I_f/I_0$
 - dilute samples
- Electron-yield
 - $A = mx = I_e/I_0$
 - surface sensitive

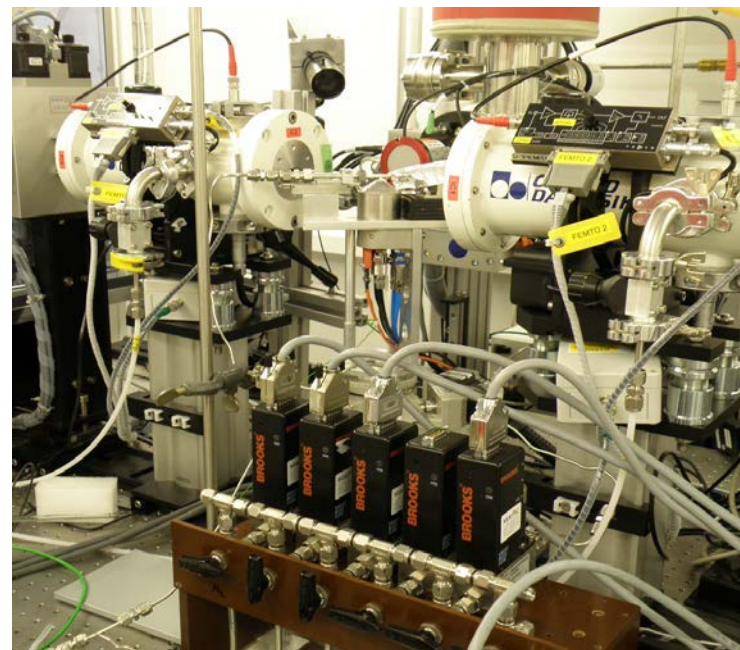




Lambert Beer's Law: $I = I_0 \cdot e^{-\mu d}$

μ depends strongly on x-ray energy E and atomic number Z , and on the density ρ and Atomic mass A :

$$\mu \approx \frac{\rho Z^4}{A E^3}$$



What do you measure?

- Absorption as function of energy

Need a Synchrotron...

SLS



Diamond



Soleil

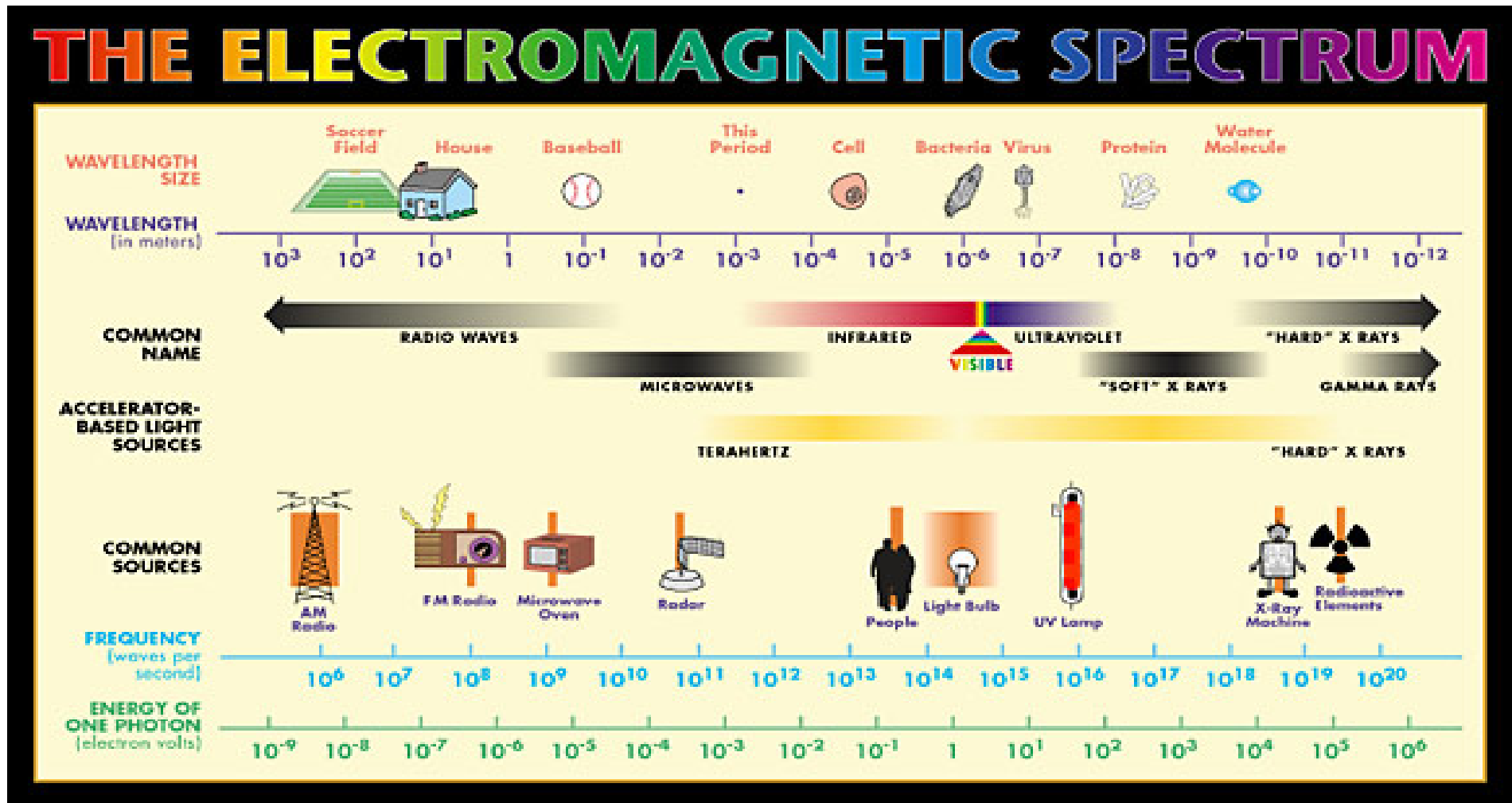


ESRF

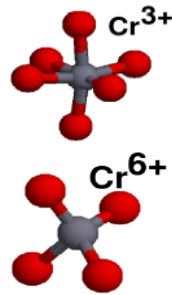
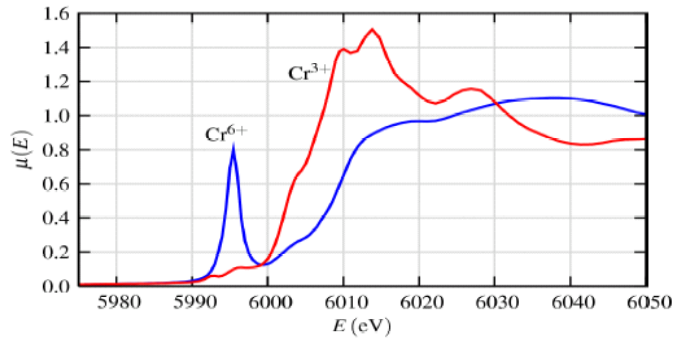


and many more...

Synchrotrons produce **bright** light

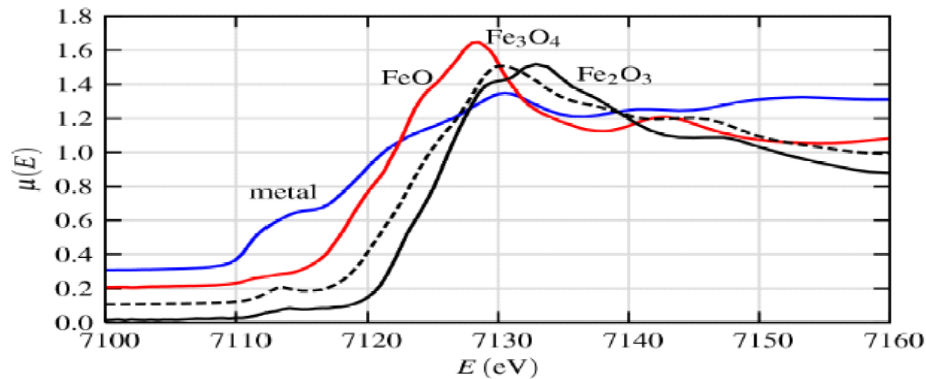


Summary so far:



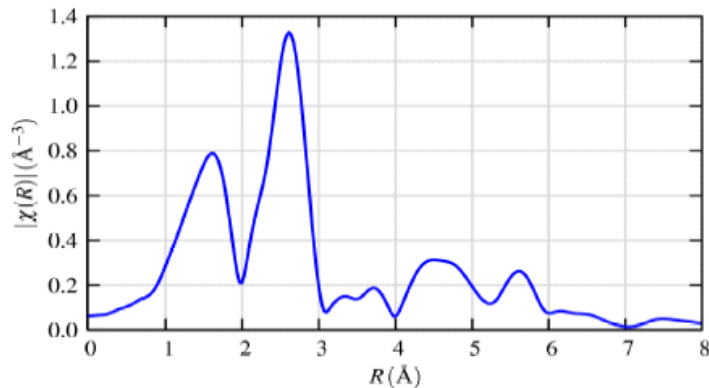
Pre-edge:

- localized electronic states
- coordination chemistry



XANES: (DOS)

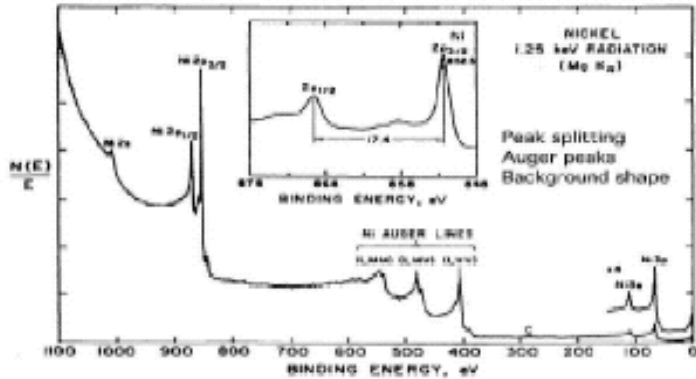
- oxidation state
- band structure
- multiple scattering



EXAFS:

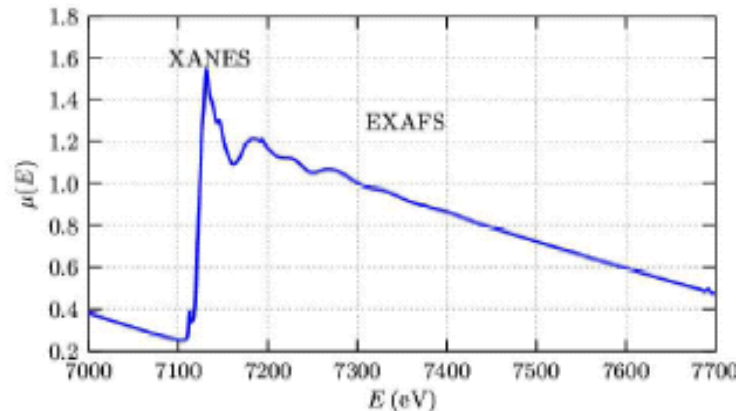
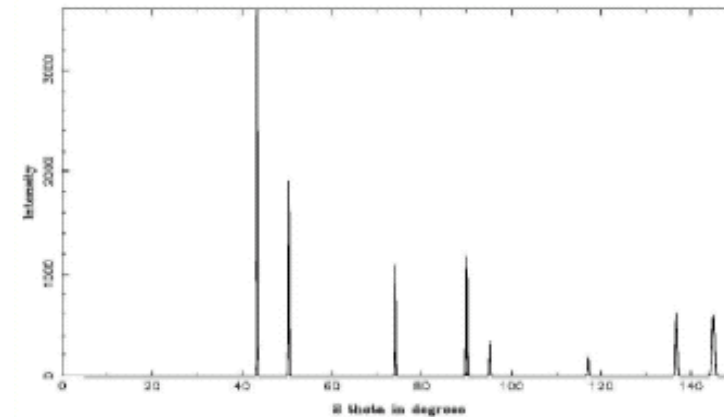
- Identity of nearest neighbors
- Bond distances
- Coordination numbers
- Amount of disorder

Comparison: XPS, XRD, XAS



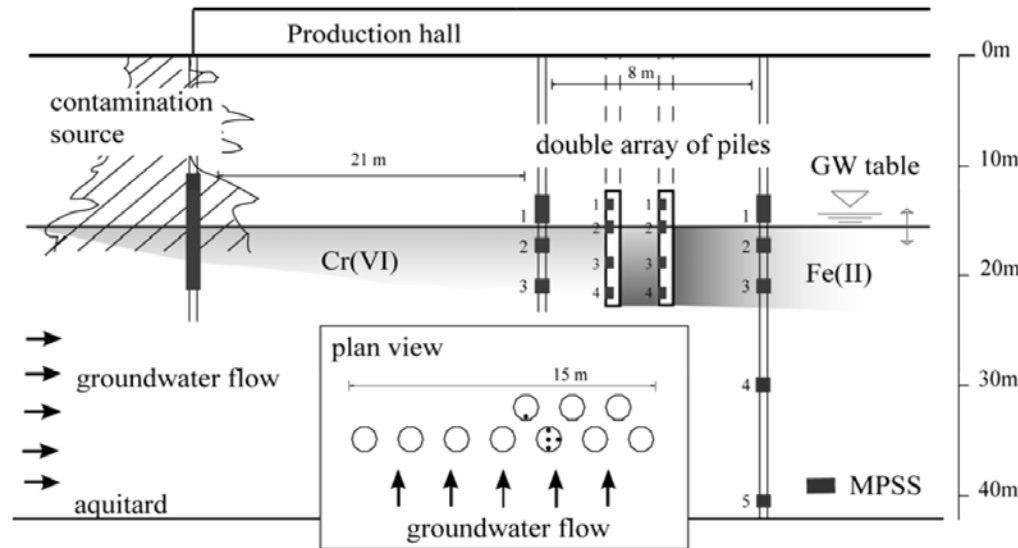
- electronic information
- surface sensitive
- In-situ application difficult
- UHV needed

- structural information
- bulk technique
- In-situ
- long range order



- electronic and structural information
- bulk and surface sensitive
- amorphous materials
- In-situ
- synchrotron needed

Example: Fe-oxides as contaminant sorbent



$E_H = 150 \text{ mV}$
 $[\text{SO}_4^{2-}] = 30 \text{ mg/L}$
 $[\text{NO}_3^-] = 11 \text{ mg/L}$
 $\text{pH} = 7$

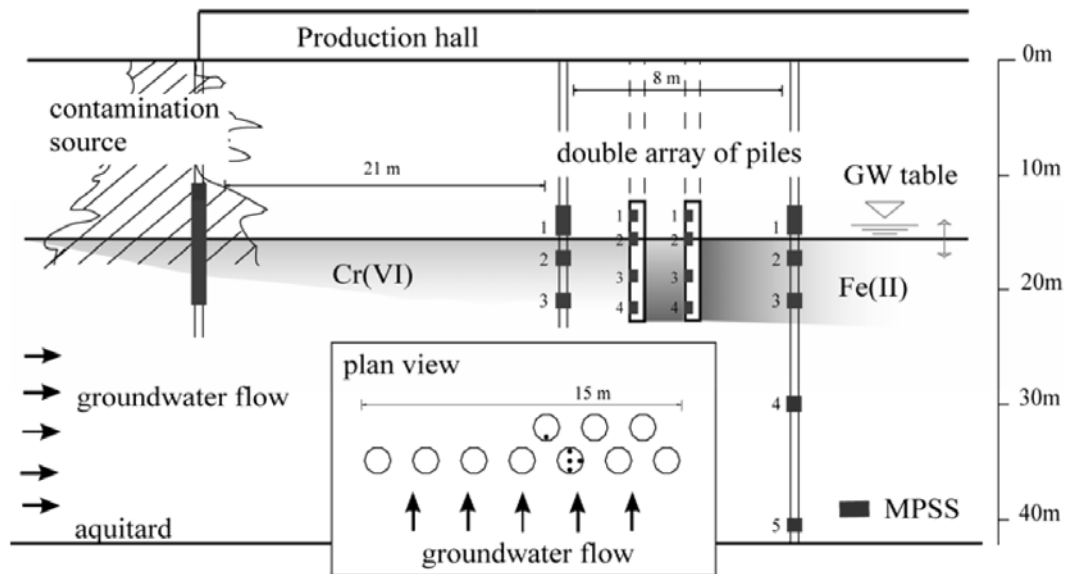


- Contaminants Cr(VI), Cu
- ~1 t Cr at a depth of 3-12 m
- Groundwater protection zone
 $[\text{Cr(VI)}]_{\text{max}} = 0.01 \text{ mg/L}$



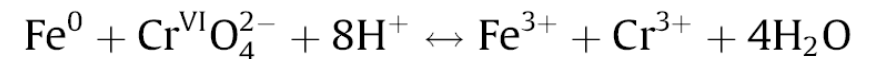
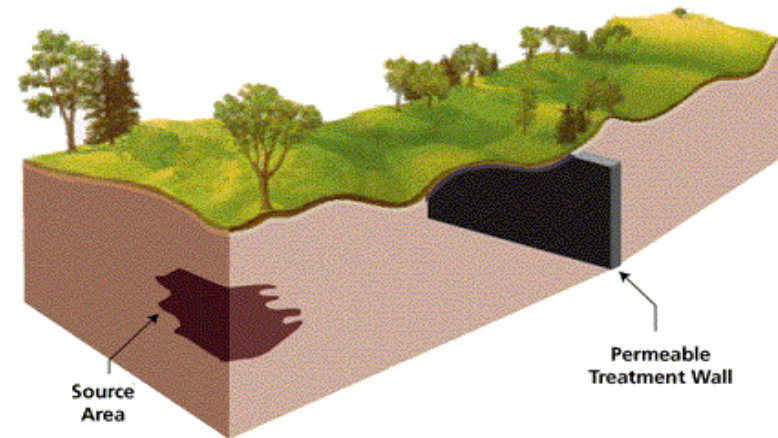
Permeable reactive barrier

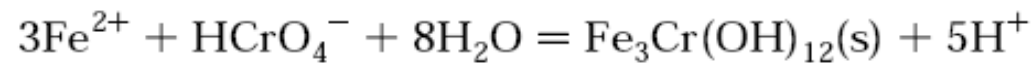
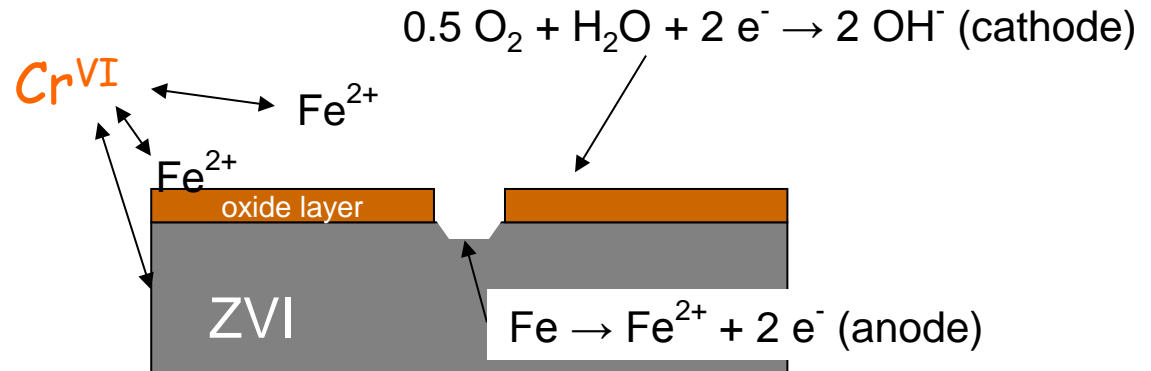
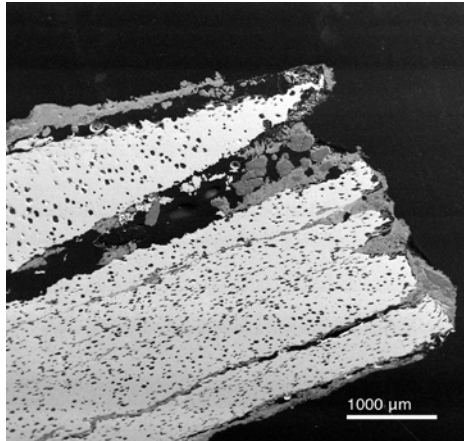
24
Cr
52.00



- Contaminants Cr(VI), Cu
- ~1 t Cr at a depth of 3-12 m
- Groundwater protection zone
([Cr(VI)]_{max} = 0.01 mg/L)

Permeable Reactive Barrier (PRB):
couple the oxidation of Fe(0) with
the reduction of Cr(VI)





- homogeneous redox reaction
(Buerge & Hug, 1997)
- heterogeneous redox reaction
(Buerge & Hug, 1999)
- ZVI-Cr^{VI}-direct reaction
(Liu et al., 2008)

—————>

Molecular Cr/Fe ratio:

1/3

Hansel et al., 2003

—————>

>1/3 (Cr clusters)

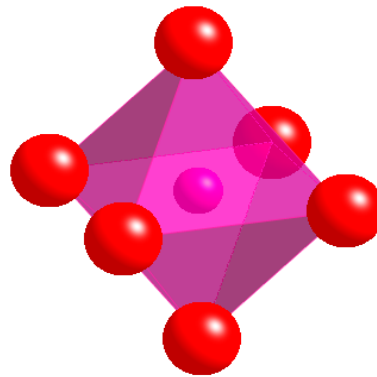
Grolimund et al., 1999

—————>

?

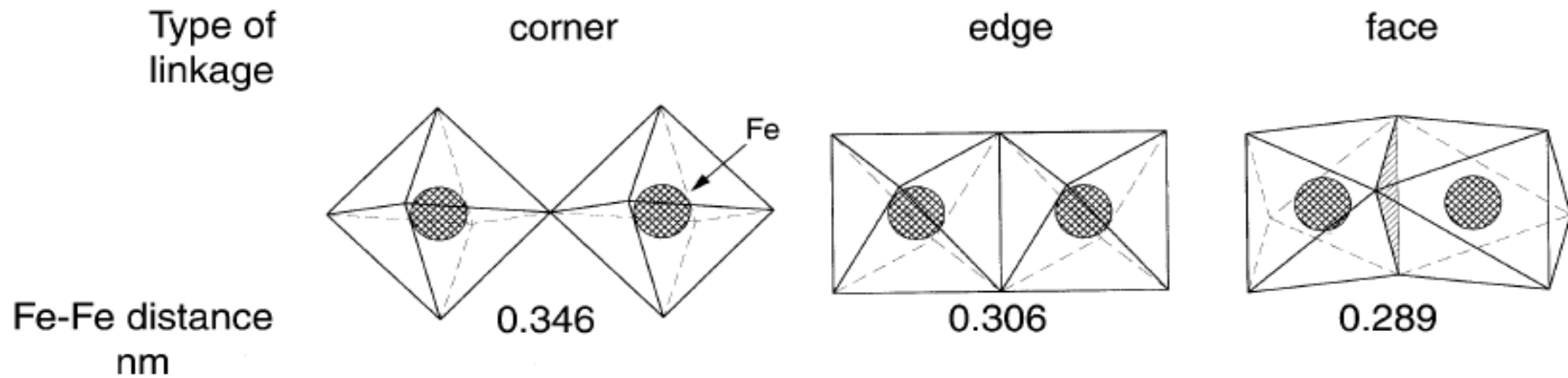
Local structure ↔ Mechanism

The basic structural unit
of Fe^{III} and oxides:

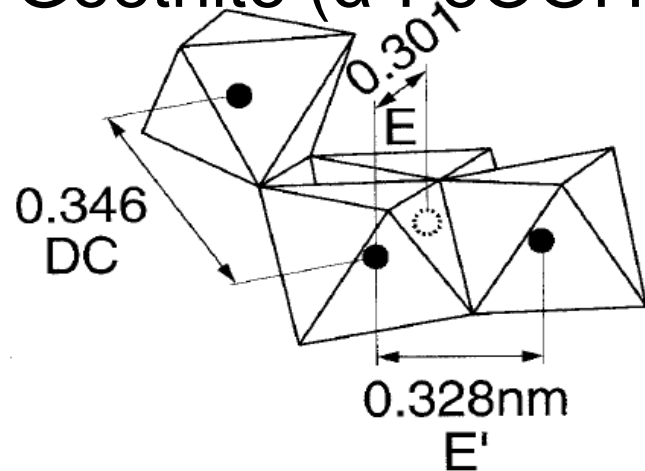


FeO_6 octahedron
(also: $\text{CrO}_3(\text{OH})_3$ $\text{FeO}_3(\text{OH})_3$)

Local structure



Goethite (α -FeOOH)



- interatomic distances
- 3D-arrangement

Why study local (~5Å) structure ?

Nucleation, growth, aggregation of mineral phases

Sorption complexes

Important properties e.g. color
($\alpha\text{-Cr}_2\text{O}_3 - \alpha\text{-Al}_2\text{O}_3:\text{Cr}^{3+}$)

Relation to molecular Fe/Cr
(-> mechanism)

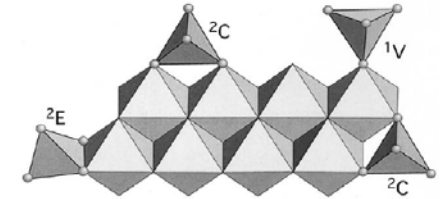
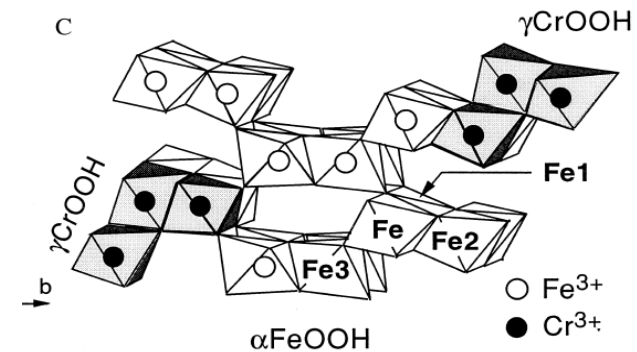
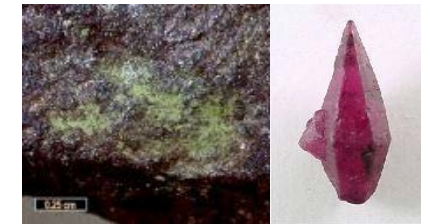
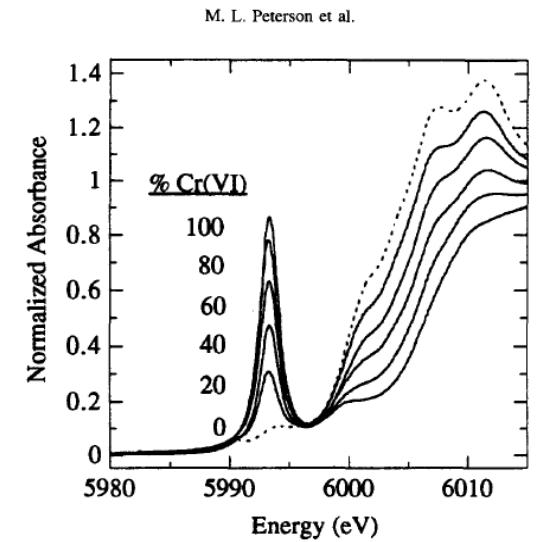
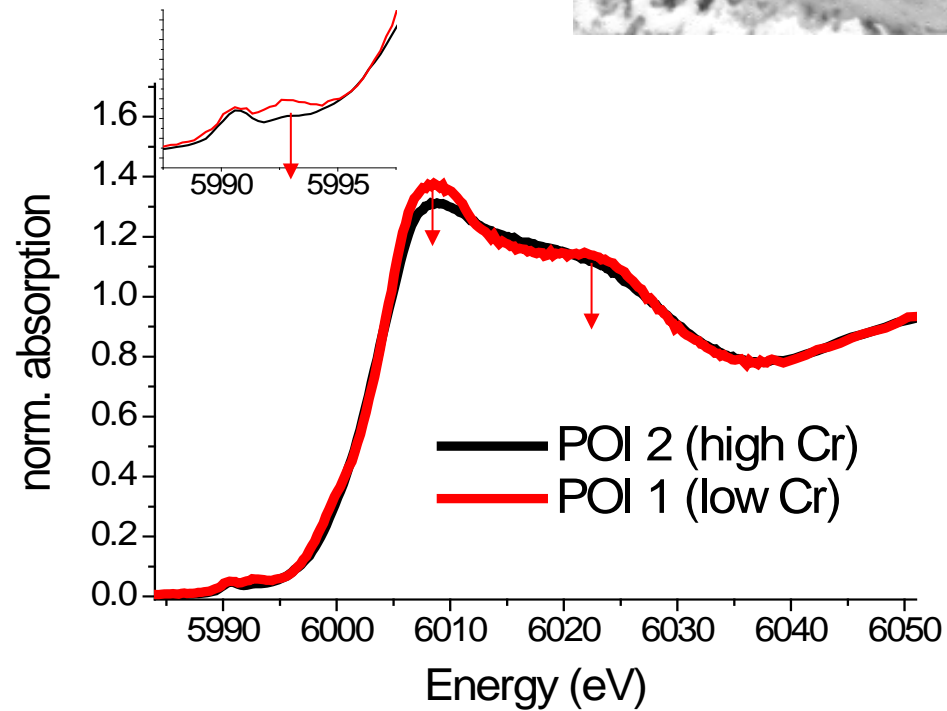
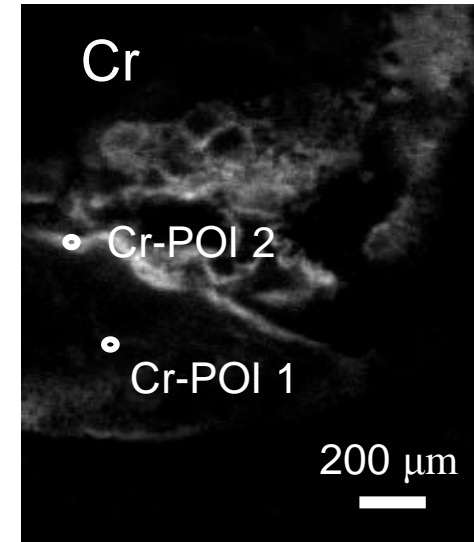
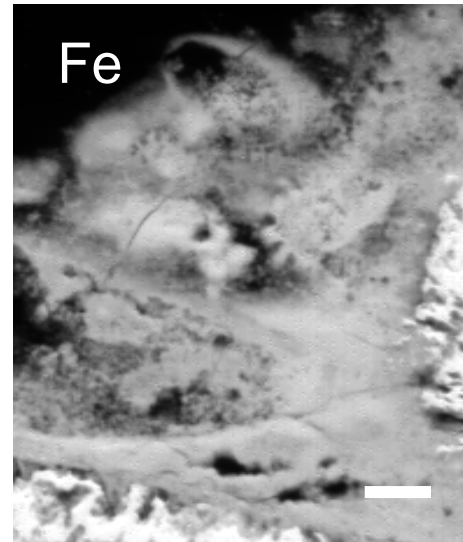
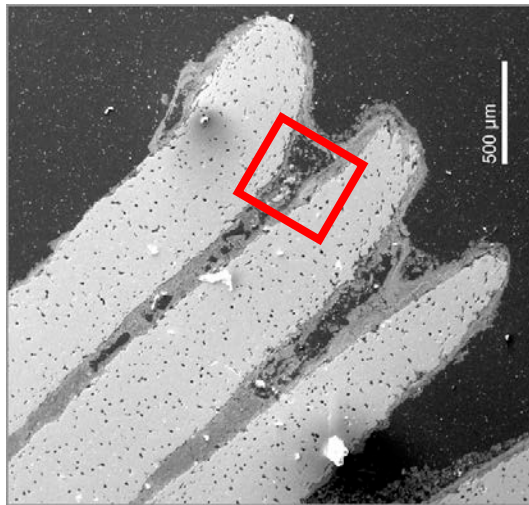


Fig. 1. Possible surface complexes of AsO_4 tetrahedra on goethite.

Sherman & Randall, GCA, 67 (2003), 4223.

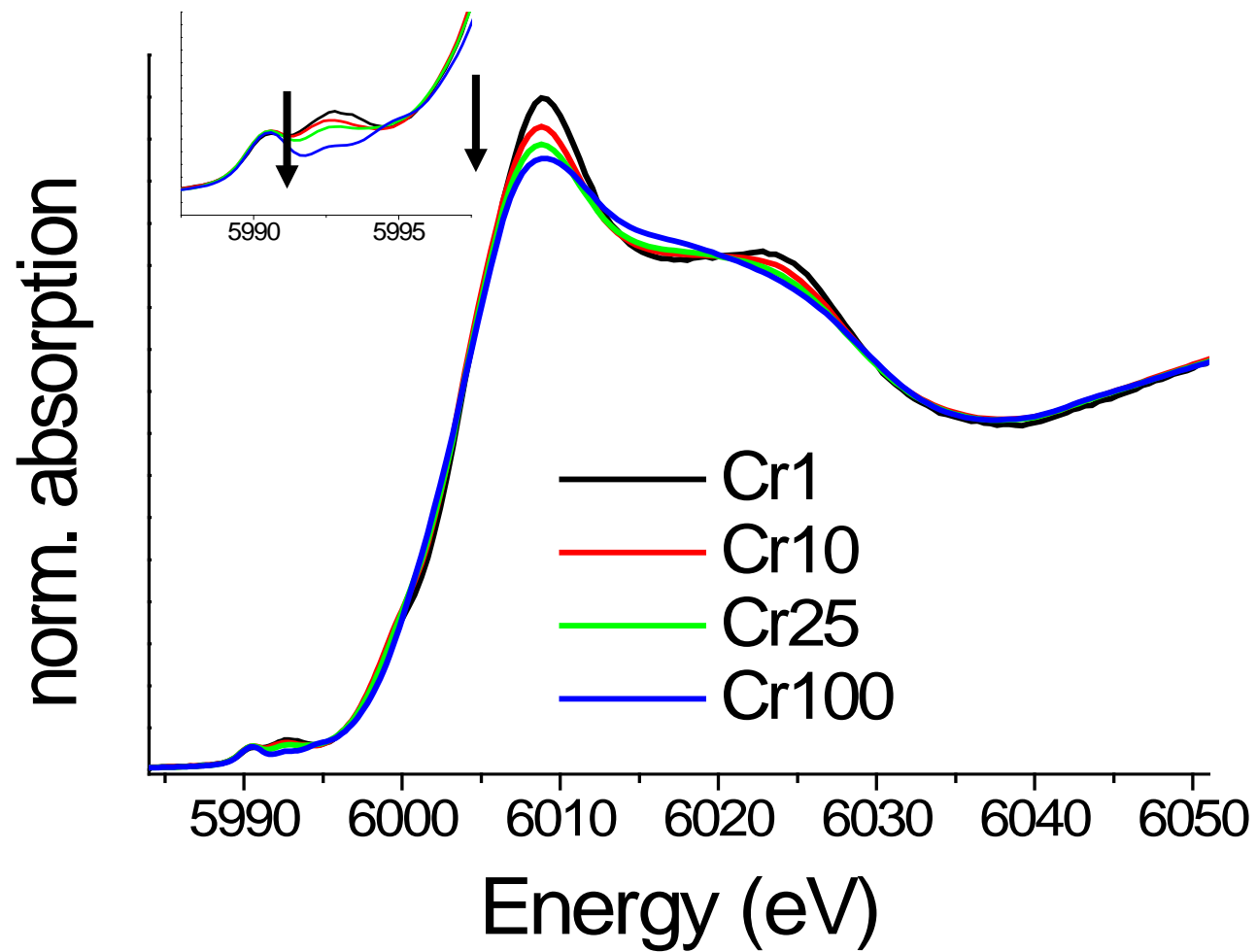


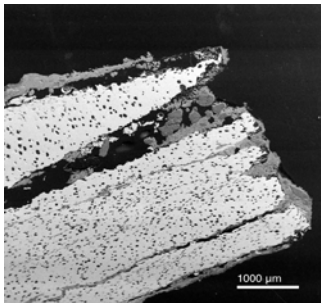
Permeable reactive barrier



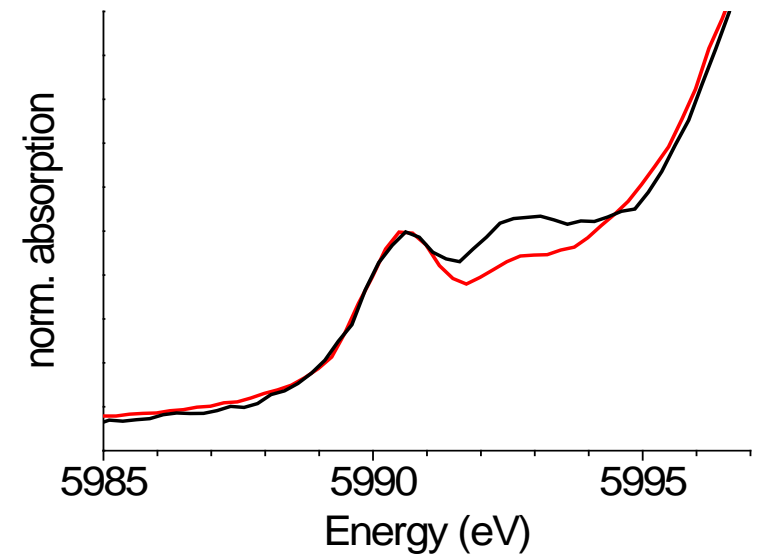
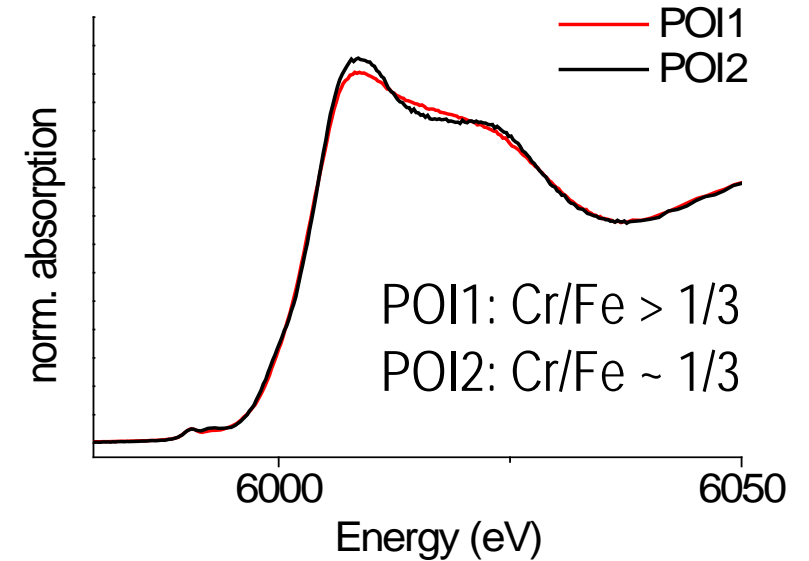
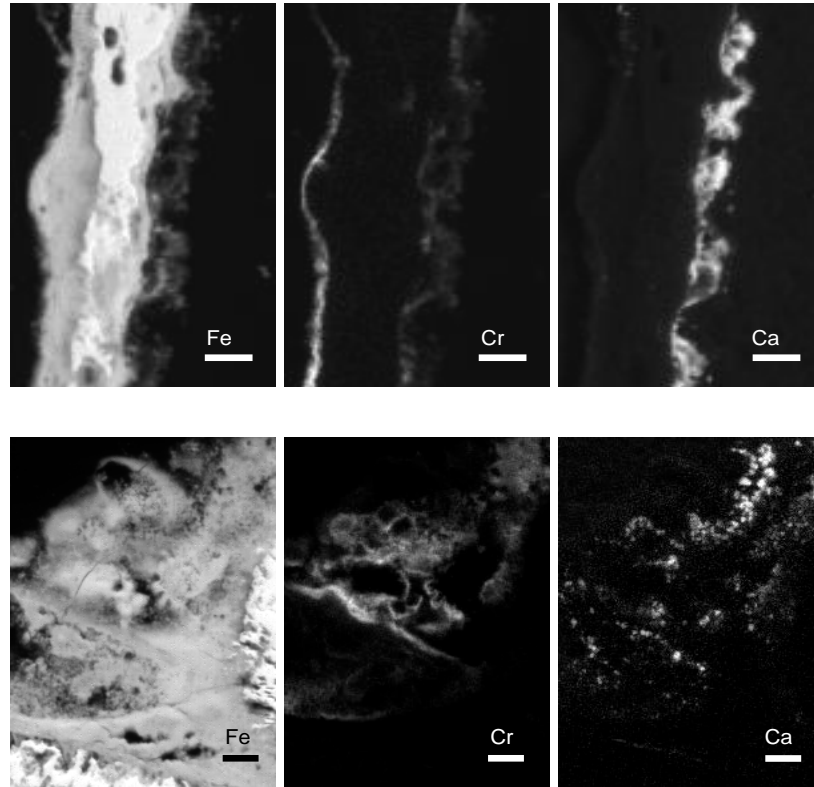
Peterson et al., *GCA* 61: 3399, 1997.

XANES





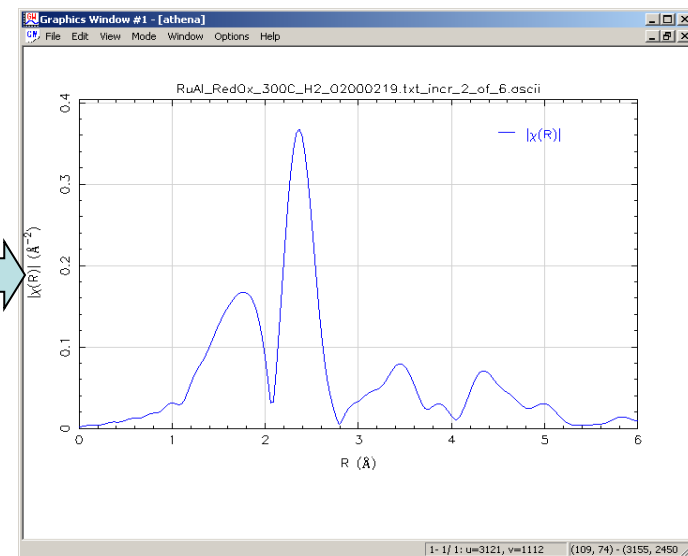
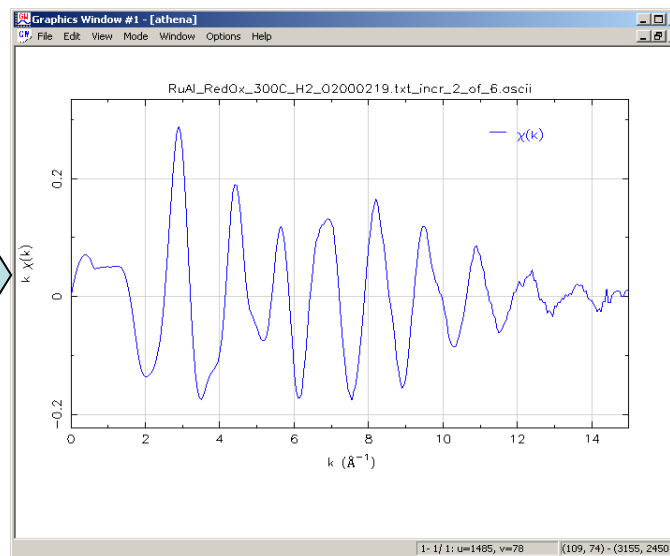
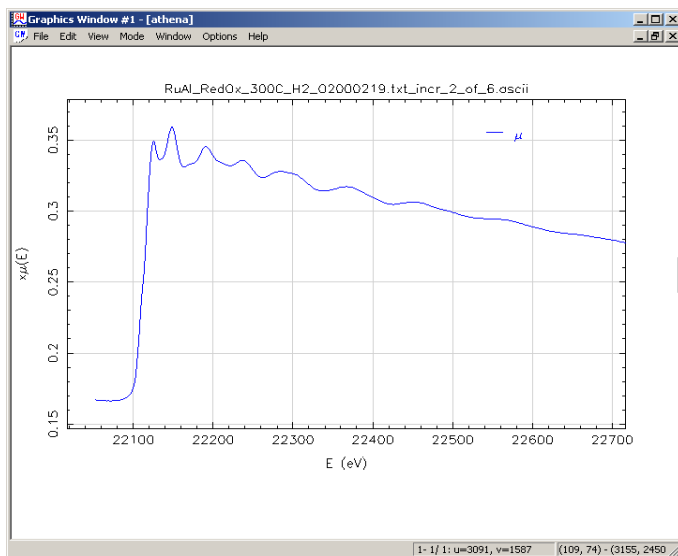
Cr/Fe < 1/10



homogeneous & heterogeneous
redox reactions

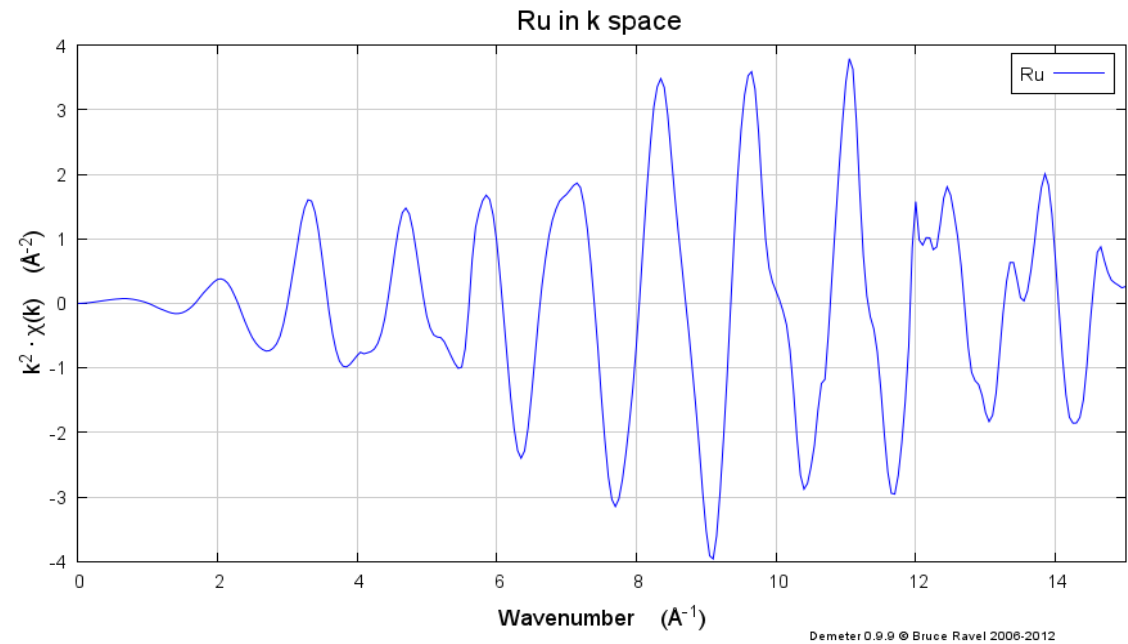
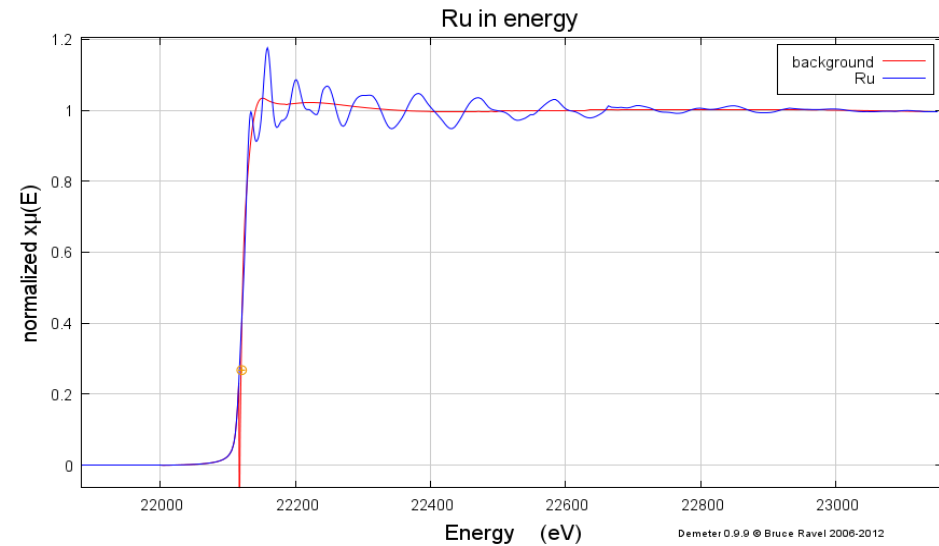
Basic data reduction steps of

- compare data that is measured in different modes (trans. / fluo.) and correct for different absorbances
- extract the EXAFS signal ($\chi(k)$) and the Fourier-transformed EXAFS signal
- look at radial distribution function, and fit the first shell neighbour



Excellent tutorial of Demeter (by Bruce Ravel): <http://www.diamond.ac.uk/Beamlines/Spectroscopy/Techniques/XAS.html>

Extraction of the EXAFS signal – $\chi(k)$



1. Extract EXAFS signal

- Subtract smooth background

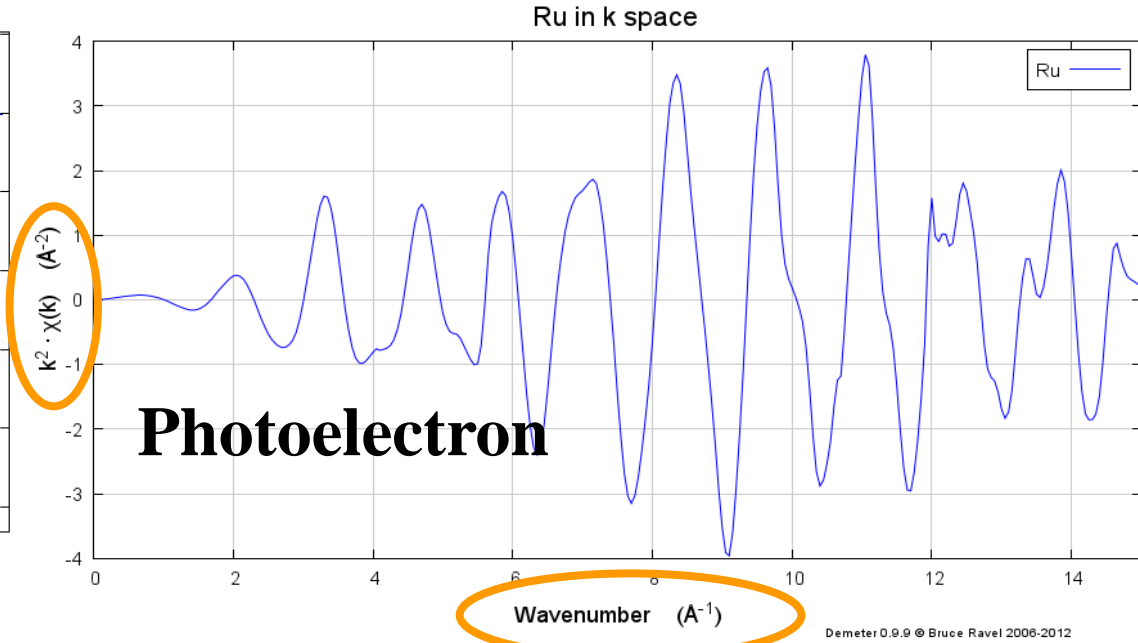
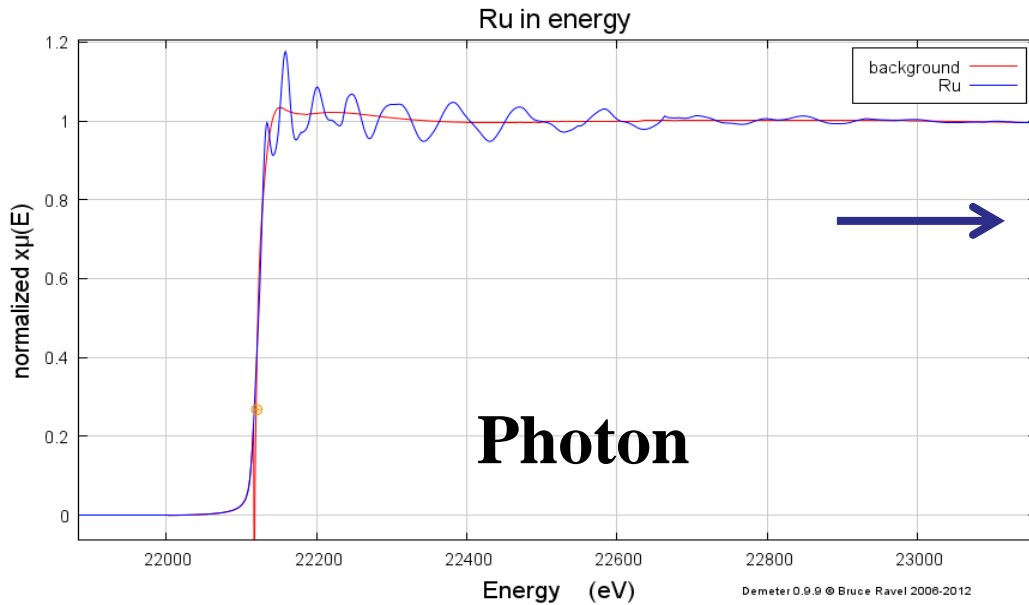
Normalized oscillatory part of absorption coefficient

Measured Absorption coefficient

Bkg: Absorption coefficient without contribution from neighboring atoms (Calculated)

$$\chi(E) = \frac{\mu(E) - \mu_0(E)}{\Delta\mu(E)} \sim \frac{\mu(E) - \mu_0(E)}{\Delta\mu(E_0)}$$

Evaluated at the Edge step (E_0)



2. Transformation from E-space to k-space

- Scattering of photoelectron on neighboring atoms
- Unit of k-space: inverse Angstroms
- „stretch“ the x-axis
- amplify signal at high energies (k-weight)

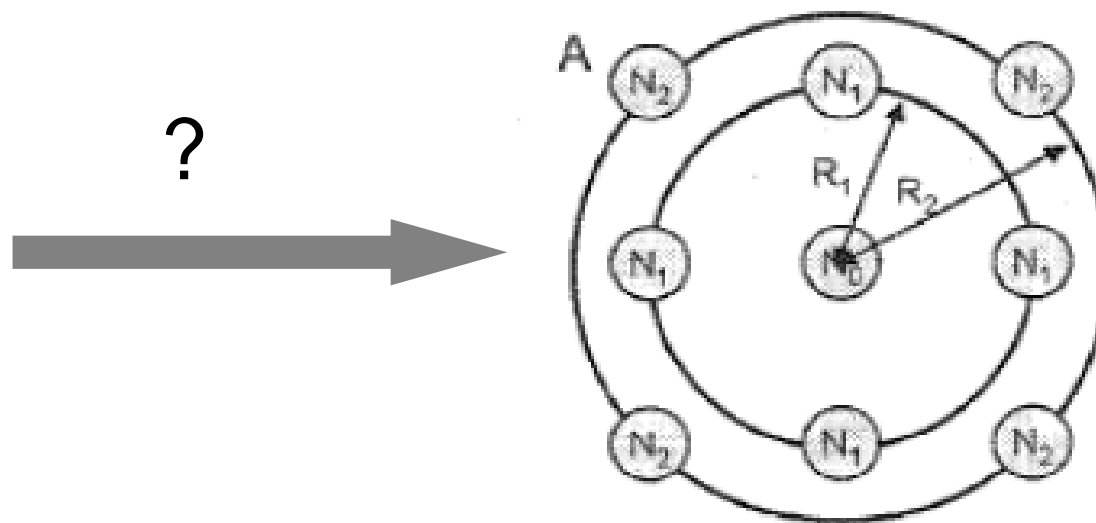
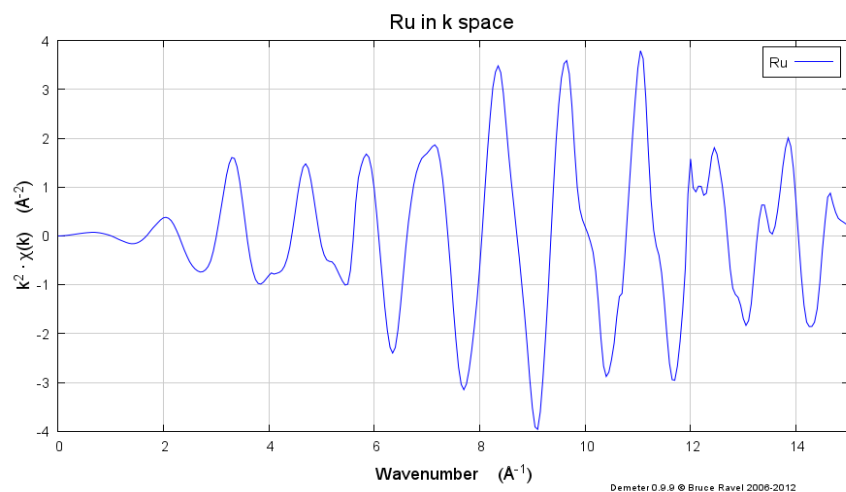
$$k^2 = \frac{2m_e(E-E_0)}{\hbar^2} \sim \frac{\Delta E}{3.81}$$

Mass of electron

Edge Energy

Plank's constant

To understand and visualize the geometric structure of our sample we need to Fourier transform the EXAFS signal from inverse distance into the distance domain



N_0 X-ray absorbing atom

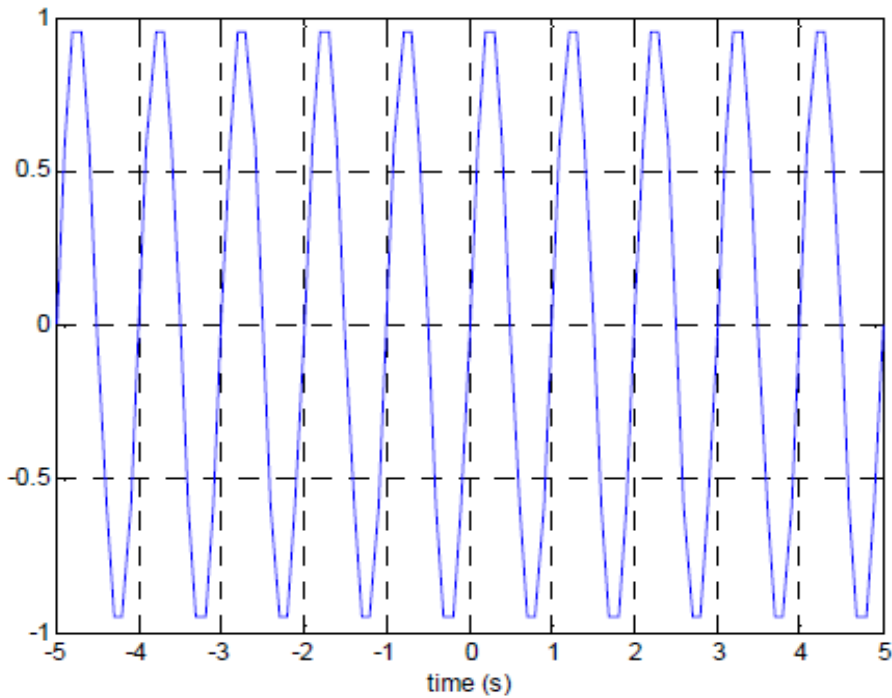
N_1 1st neighboring scatterer (shell)

N_2 2nd neighboring scatterer (shell)

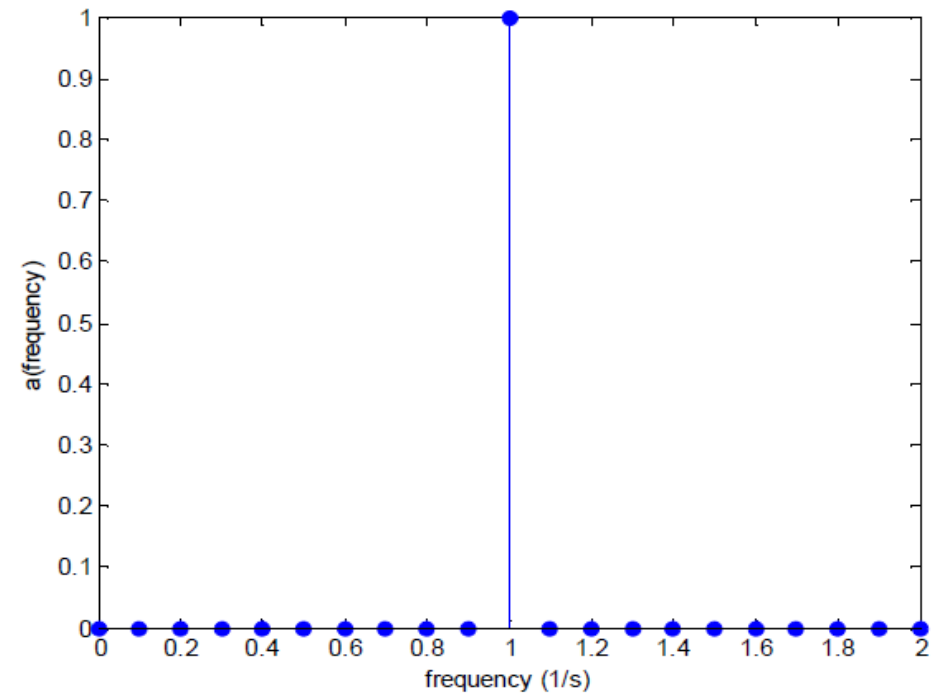
R_x distance between absorber and scatterer

Fourier transformation crash course

- FT transforms data from one dimension into its reciprocal one
- example: transformation from the time into the frequency domain



Wave function with period length of 1 s



Frequency of 1 Hz

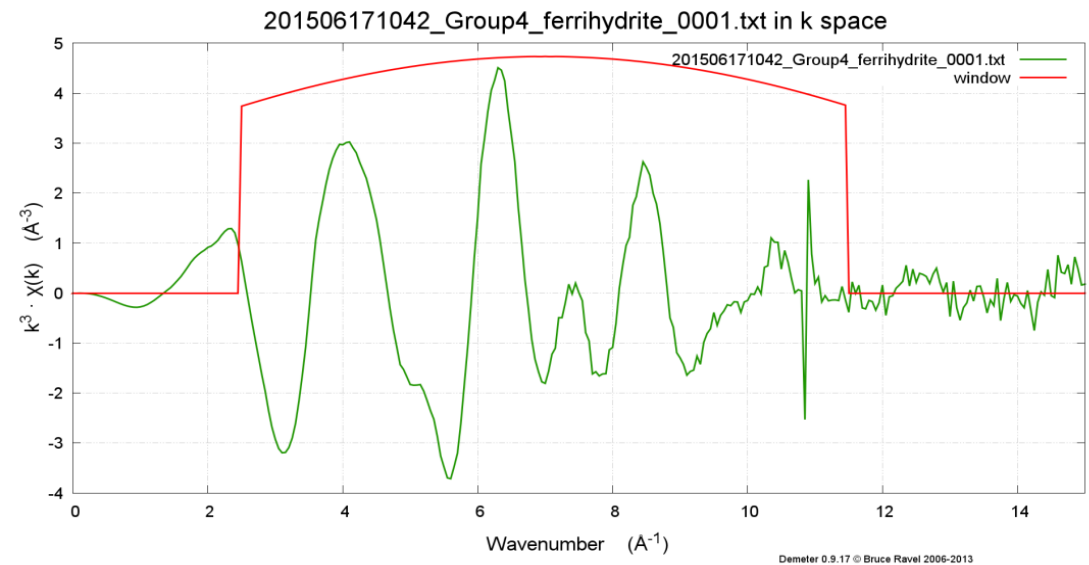
Define k-range for Fourier transformation

Lower limit: 2-3 \AA^{-1} ; upper limit: as far as reasonable (consider signal to noise)

Fourier Transform Button

Forward Fourier transform parameters
 k-range 2.000 to 13.211 dk 1 window Hanning
 arbitrary k-weight 0.5 phase correction

Plot in k-space
 x(E) Window



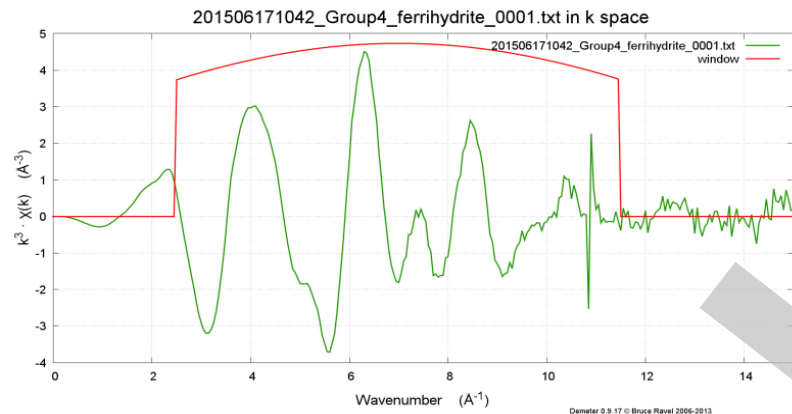
Recover the distance of scattering atoms from the frequency of the scattered wave

→ Fourier transform the EXAFS function (from \AA^{-1} to \AA)

Fourier transformed of the EXAFS signal

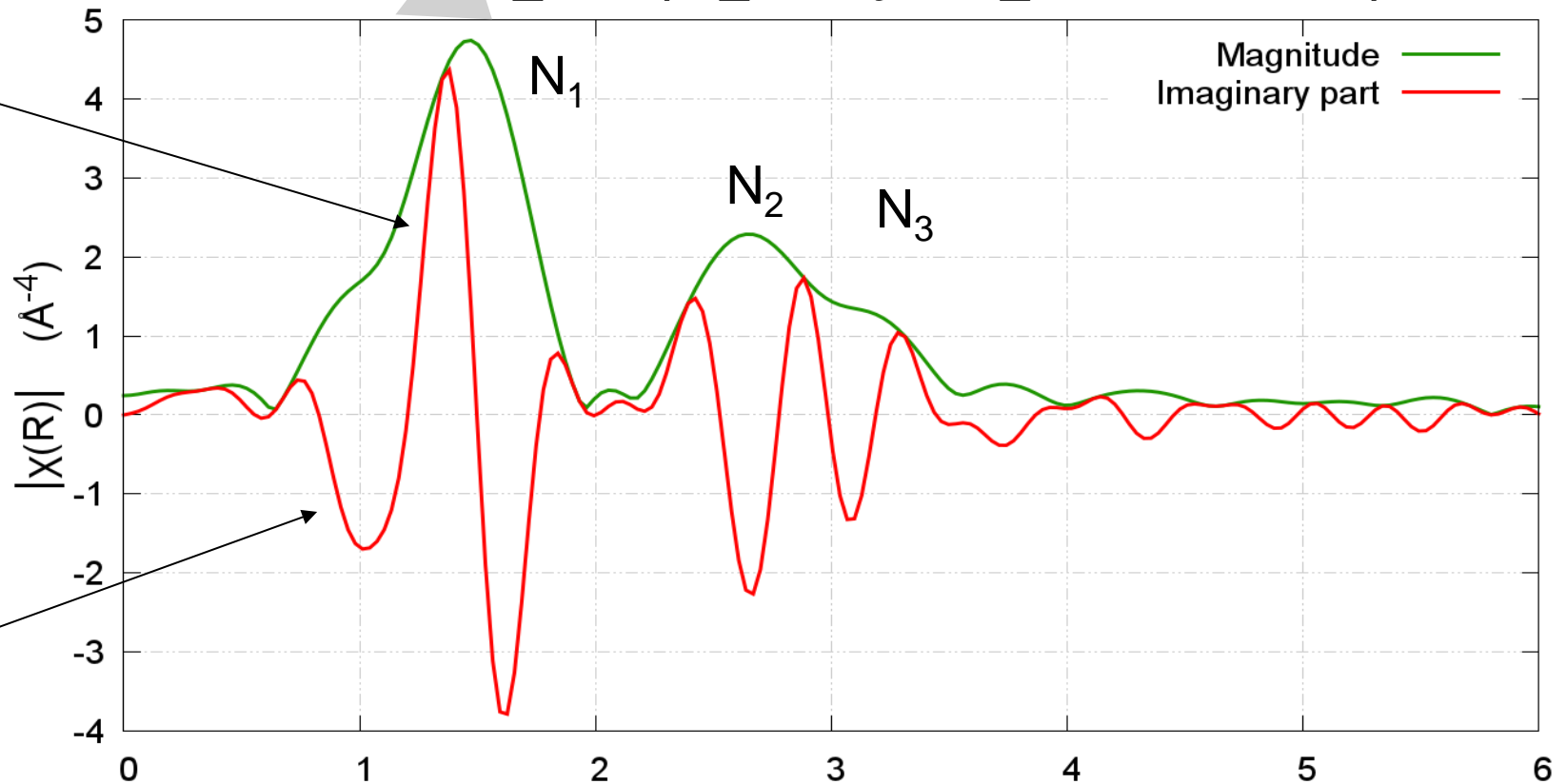
Remember: FT creates a complex function

→ always show magnitude AND real or imaginary part



201506171042_Group4_ferrihydrite_0001.txt in R space

magnitude

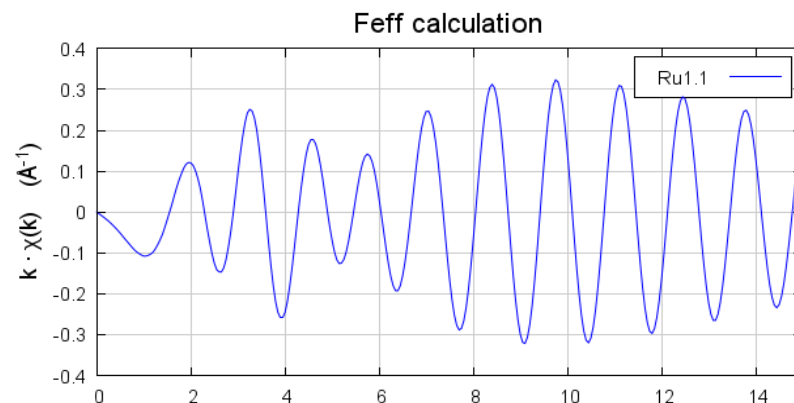
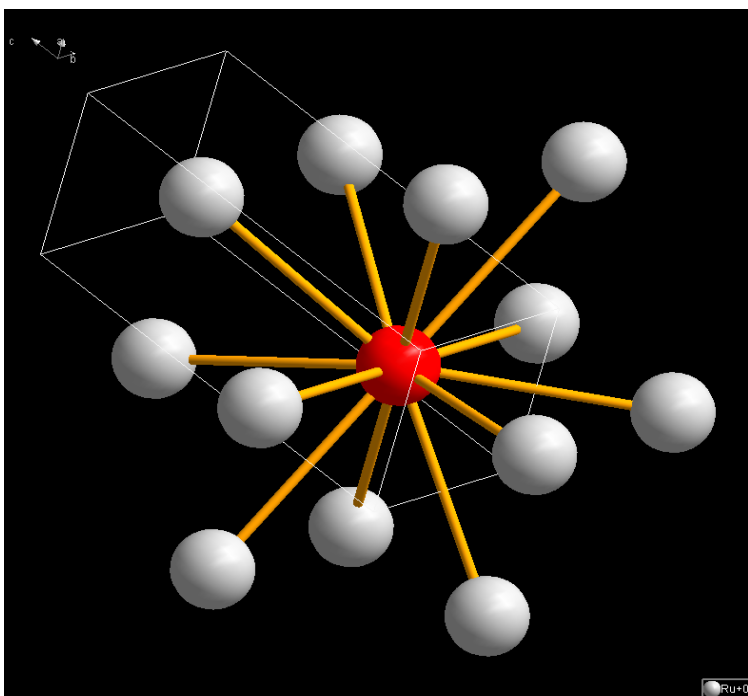


Imag part

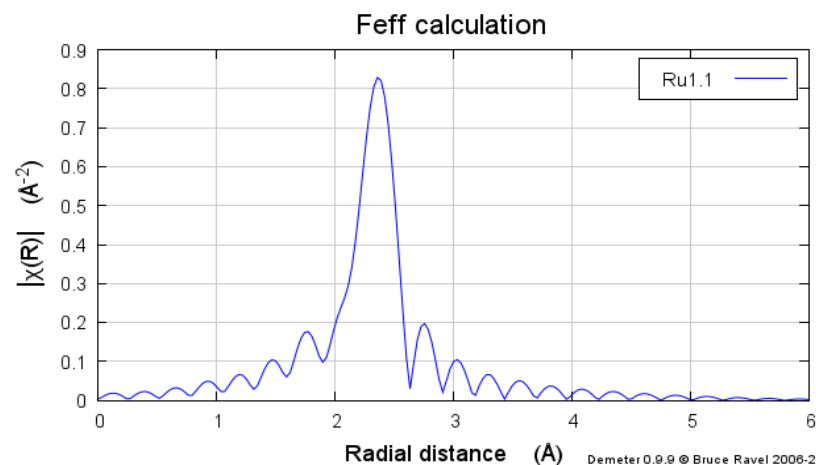
Radial distance (Å)

The pseudo RDF created in the previous step is now analyzed by fitting with a set of model structures

- ➔ Create a model of the assumed structure
- ➔ Calculate the EXAFS function of this theoretical structure (ATOMS @ ifeffit)
- ➔ Fit the model to the data, extract structural parameters



Calculated
scattered
wave



Corresponding
RDF

What information can we extract from a fit?

EXAFS function

$$\chi(k) = \sum_i \frac{N_i F_i(k) S_0^2}{k R_i^2} e^{-\frac{2R_i}{\lambda}} e^{-2\sigma_i^2 k^2} \sin(2kR_i + \phi_i(k))$$

Sum of damped sine functions with a pre-factor

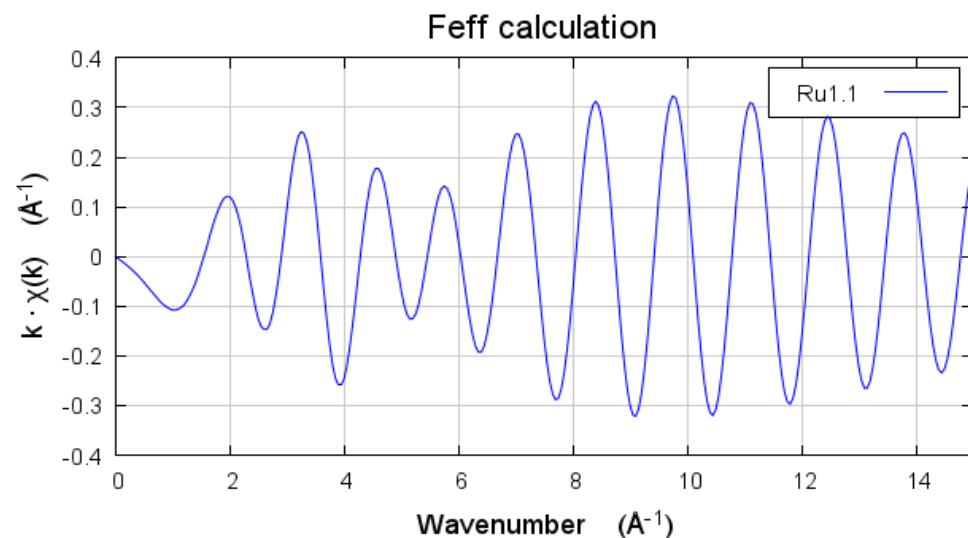
Structural Parameters:

N: coordination number → amplitude

R: radial distance → frequency

σ^2 : pseudo Debye waller factor → damping

Theoretical first Ru-Ru shell of metallic ruthenium

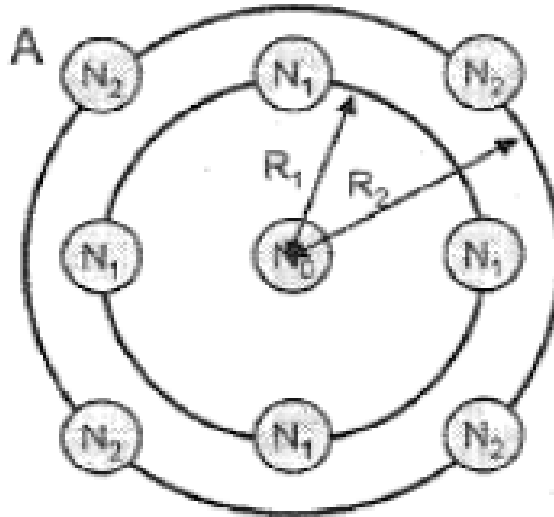


Demeter 0.9.9 © Bruce Ravel 2006-2012

Calculate EXAFS signal of model compound! → FEFF

Reminder: what's a coordination shell?

Every shell of atoms has a specific distance from the absorber and a specific coordination number



From fitting of EXAFS data information about

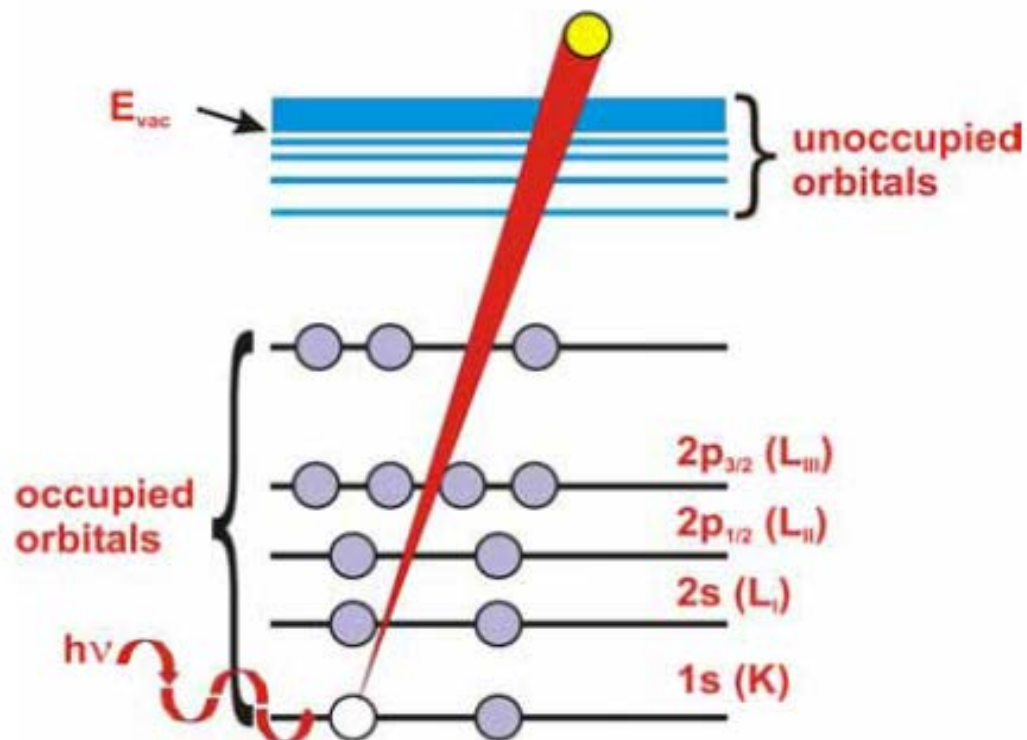
- the distance R of each shell from the absorbing atom
- the number of atoms in each shell (coordination number) is obtained.

Beamline	Energy Range (keV)	Spot Size (μm)	K-Edges	Absorption	Fluorescence	Electron yield	Speciality
PHOENIX	0.8 - 8	2 x 2	Na - Co	x	x	x	Energy Range Variable Polarization
MicroXAS	5 - 23	1 x 1	Ti - Mo	x	x		Femto Diffraction radioactive samples
SuperXAS	6 - 40	5 x 10	Cr - La	x	x		Absorption, Fluorescence, Quick, Chemical environments
Pollux	0.2 - 1.2	0.04 x 0.04	C - Mg	x	(x)		Nano focus, STXM, absorption spectroscopy, phase contrast
nanoXAs	0.2 - 1.2	0.04 x 0.04	C - Mg	??	??	x	Combine AFM with STXM



Thank you for your attention! Questions?

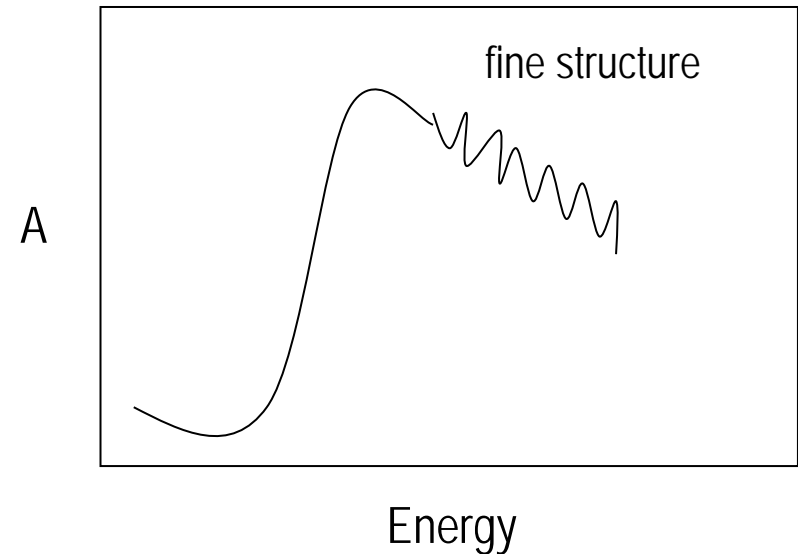
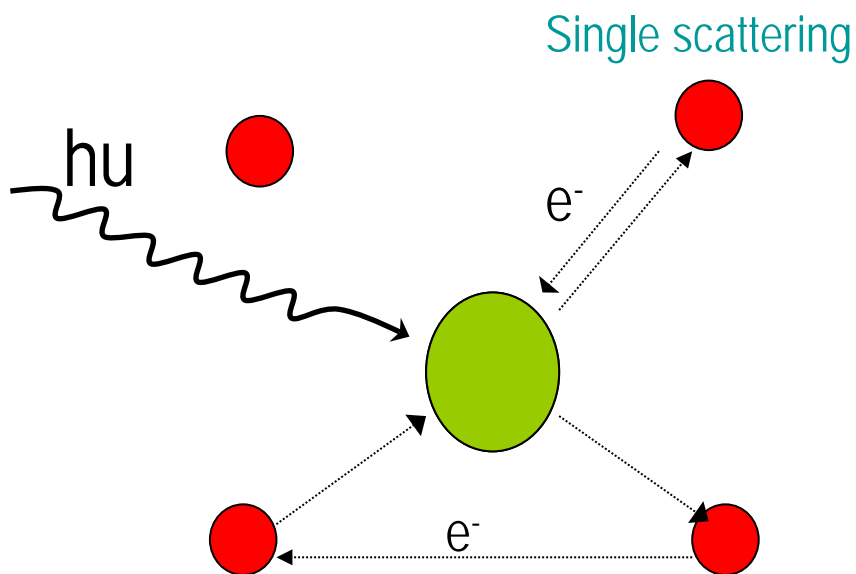
The basic concept of XAFS

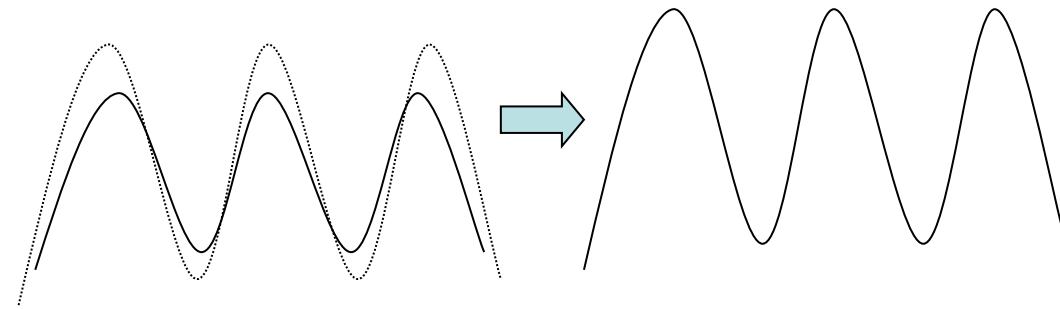
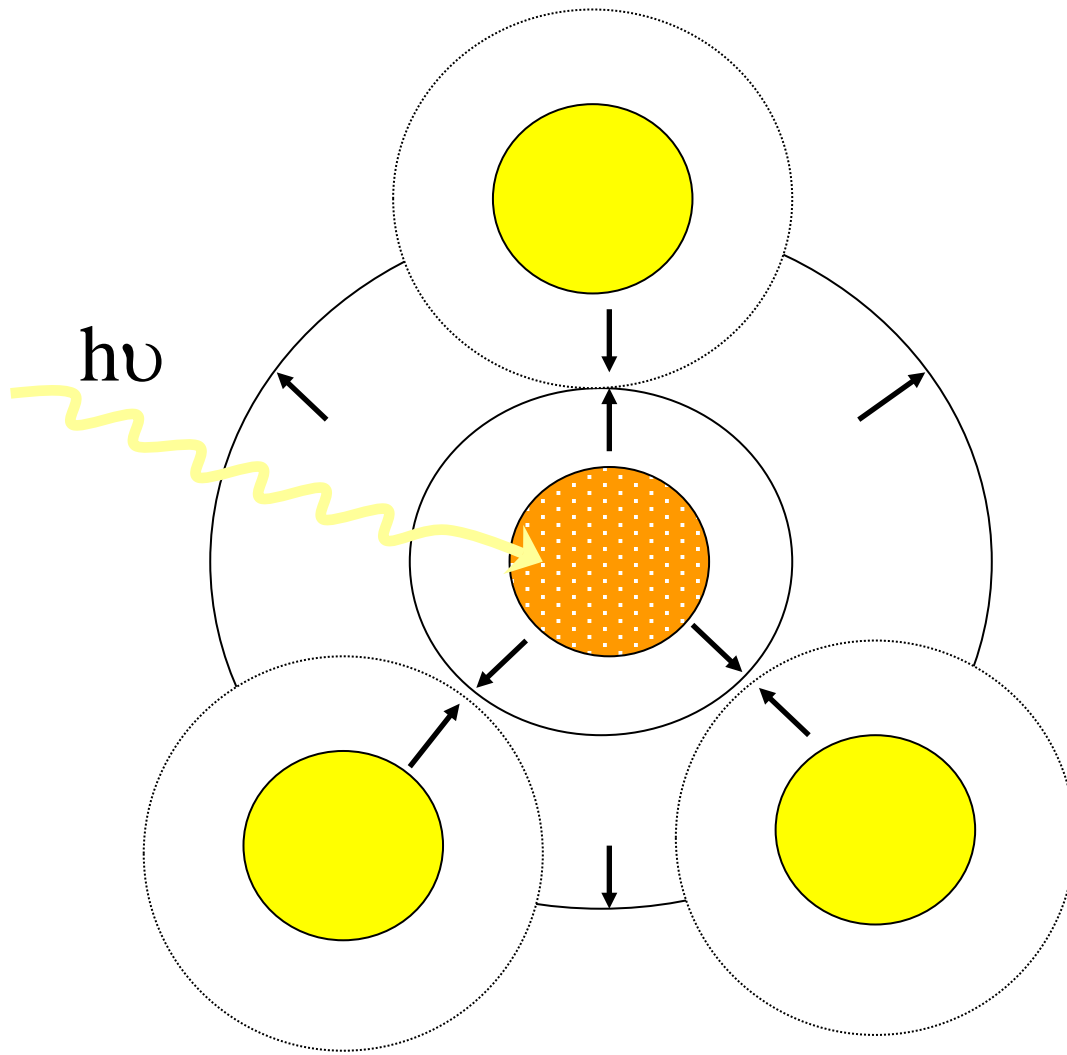


X-rays (light with wavelength 0.06- 12 Å or energy 1-200 keV) are absorbed by all matter through the **photo-electric effect**:

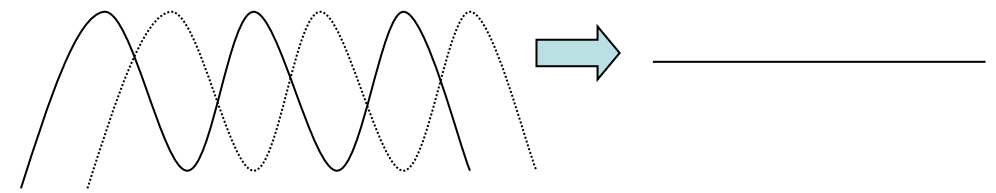
An X-ray is absorbed by an atom when the energy of the X-ray is transferred to a core level electron (K, L, or M shell) which is subsequently ejected from the atom. Any excess energy from the X-ray is given to the ejected photo-electron.

Electrons have a particle and wave nature. The photoelectron wave propagates away from the central atom (absorber), and it may scatter off neighboring atoms and finally return to its point of origin.





In phase



Out of phase

The consequence of these scattering phenomena and wave interactions is that the intensity of X-ray absorption oscillate with a dependence on the structural environment of the absorber. Mathematically modeling these oscillations provides precise local structural information.