XANES Measurements and Interpretation

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Acronyms

XANES

- X-ray Absorption Near Edge Structure

NEXAFS

- Near-Edge X-ray Absorption Fine Structure

The two acronyms should be interchangeable but over the years NEXAFS has become terminology for “low Z” elements - C, N, O...
What Is XANES?

- XANES is the region of x-ray absorption spectrum within ~50eV of the absorption edge.
- Suggested that division is that at which wavelength of excited electron is equal to distance between absorbing atom and its nearest neighbor. \( \lambda (\text{\AA}) \approx \frac{12}{[\text{e(eV)}]^{\frac{1}{2}}} \).

X-ray absorption spectrum of molybdenum metal from 19.8 to 21.5 keV
What Is XANES?

XANES = Pre-edge + Edge + XANES

(X-ray absorption spectrum of Ti K-edge of Ba$_2$TiO$_4$)
Why Are We Interested In XANES?
Local Coordination Environment

- Ti K-edge XANES shows dramatic dependence on the local coordination chemistry.

Both Ti$^{4+}$

Ba$_2$TiO$_4$

K$_2$TiSi$_3$O$_9$
Why Are We Interested In XANES?  
Oxidation State

- Many edges of many elements show significant edge shifts (binding energy shifts) with oxidation state.
What Is XANES and Why Are We Interested?

XANES is strongly sensitive to the chemistry (formal oxidation state and geometry) of the absorbing atom.

<table>
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<tr>
<th>Region</th>
<th>Transitions</th>
<th>Information Content</th>
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<tbody>
<tr>
<td>Pre-edge</td>
<td>Features caused by electronic transitions to empty bound states. Transition probability controlled by dipolar selection rules.</td>
<td>Local geometry around absorbing atom. Dependence on oxidation state and bonding characteristics (chemical shift).</td>
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<tr>
<td>Edge</td>
<td>Defines ionization threshold to continuum states.</td>
<td>Dependence on oxidation state (chemical shift), main edge shifts to higher energy with increased oxidation state. (As much as 5 eV per one unit change).</td>
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<tr>
<td>XANES</td>
<td>Features dominated by multiple-scattering resonances of the photoelectrons ejected at low kinetic energy. Large scattering cross section.</td>
<td>Atomic position of neighbors: interatomic distances and bond angles. Multiple scattering dominates but ab initio calculations providing accessible insight (e.g. FEFF8).</td>
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**XANES Transitions**

- XANES directly probes the angular momentum of the unoccupied electronic states: these may be bound or unbound, discrete or broad, atomic or molecular.

- Dipole selection rules apply: $\Delta l = \pm 1$, $\Delta j = \pm 1$, $\Delta s = 0$.

- Primary transition will be:
  - $s \rightarrow p$ for K (1s core electron) and L\textsubscript{1} (2s core electron initial state) edges
  - $p \rightarrow d$ for L\textsubscript{2} (2p\textsubscript{1/2}) and L\textsubscript{3} (2p\textsubscript{3/2}) edges

- But…..final state usually not atomic-like and may have mixing (hybridization) with other orbitals. This is often the interesting part of the XANES!
**XANES Interpretation**

- The EXAFS equation breaks down at low-\(k\), which complicated XANES interpretation.

- **We do not have a simple equation for XANES.**

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

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

- These chemical and physical interpretations are all related:

  **What electronic states can the photoelectron fill?**
Advantages of XANES vs. EXAFS

- Spectra simpler to measure than EXAFS: features intense, concentrated in small energy region.
- Weak temperature dependence (Debye-Waller), so spectra can be recorded at reaction temperature (in situ).
- Faster to measure than full spectrum: <msec demonstrated.
- Sensitive to chemical information: valence, charge transfer.
- Probes unoccupied electronic states: important in chemistry.
- Often used as simple “fingerprint” to identify presence of a particular chemical species.
- Beamlines with micro-probe capabilities can also scan energy and obtain XANES spectra with elemental distribution.
**XANES Analysis: Oxidation State Sulfur**

Sulfur K-edge XANES used to identify and quantify the form of sulfur in heavy petroleum, coals, soils etc.

11 eV edge shift from $S^{2-}$ to $S^{6+}$.

Spectra of S in similar environments similar: benzothiophene, dibenzothiophene.

Can be used as fingerprint.

XANES Analysis: Oxidation State

Many, many examples in the literature……

Mo K-edge

V K-edge

Ref: Cramer et al., JACS, 98 (1976) 1287

**Local Site Symmetry in Ti-containing Compounds**

- Symmetry around absorbing atom strongly affects pre-edge transition: ability to differentiate 4, 5, 6-fold coordination.
Local Site Symmetry in Ti-containing Compounds

- Correlation between **absolute position and peak height** of pre-edge peak: all 4-fold, 5-fold and 6-fold coordinated Ti compounds fall into separate domains.

- Ability to distinguish Ti coordination from pre-edge peak information.

XANES of 3d Transition Metals: Coordination

- 1s to 3d pre-edge peak sharp and intense from Ti-Mn, decreases Fe-Cu, absent for Zn.
- Decrease in intensity due to progressive filling of the 3d band.
- Oh symmetry shows only a small pre-edge peak throughout series.

“White line” Intensity of Group VIII Metals

- Peak historically called a “white line” as when it was detected by x-ray film it showed up as a white line due to the strong absorption.

L₃ edge XANES for 5d metals

- Transition from 2p3/2 to 5d states.
- Absence of peak for Au - 5d states almost completely occupied.
- For others Pt<Ir<Os<Re, corresponding to increase in number of unoccupied 5d states on the atoms.

Quantification of “White Line”

- Fit to combination of Lorentzian and arctangent functions.
- Determine: area, fwhm, position.


- Linear correlation between white line area and number of 5d-holes for Au-Re
“White Line” Intensity

Re L₃-edge - Transition from 2p3/2 to 5d states.

- Intensity of Re L₃ white line probes Re LDOS

Re metal (Re⁰) - 5d⁵
ReO₂ (Re⁴⁺) - 5d¹
NH₄ReO₄ (Re⁷⁺) - 5d⁰
**Pt L₃ and L₂ Edge XANES**

- Significant difference in L₃ and L₂ edge XANES: 2p to 5d transitions.

- Pt 5d³/₂ filled, so no white line.

- Pt 5d₃/₂ filled, so no white line.

- Same l=2 final density of states but because of selection rule, Δj = ±1, different total quantum number probed.

- j=3/2 probed by L₂-edge, j=5/2 probed by L₃-edge.
XANES to Probe Charge Transfer in Alloys

- Transition is $2p$ to $5d$: Pt $d$-band full, so “no” intensity at edge.
- PtGe intermetallics: charge transfer from $d$-band of Pt to Ge, resulting in significant intensity at edge.
- Use as signature of Pt-Ge intermetallic formation.
Effect of Adsorbed Hydrogen on Pt L₃ XANES

- White-line intensity decreases and spectra broaden to higher energies as H is added.
- Difference signal typically leads to broad structure ~8 eV above absorption edge.
- Several different interpretations in the literature.
Symmetry: Ni K-edge

- Only square planar Ni compounds exhibit intense pre-edge peak at 5 eV.
- Ni in feed predominantly square planar Ni\(^{2+}\). This converted to different species during reaction.

Ref: Barkigia et al, JACS, 115 (1993) 3627
Which Edge to Choose: Energy Resolution

Mo K-edge XANES of Na$_2$MoO$_4$

- Mo K-edge at 20.00 keV, effective resolution of 10 eV dominated by core-hole lifetime.
Which Edge to Choose: Energy Resolution

- Comparison of normalized Mo L\textsubscript{3}-edge (2.5 keV) XANES of Na\textsubscript{2}MoO\textsubscript{4} with that of Mo K-edge (20.0 keV).

- Mo L\textsubscript{3}-edge at 2.5 keV, 0.5 eV spectral resolution!
Which Edge to Choose: Energy Resolution

Sodium Molybdate, NaMoO$_4$
Mo atom tetrahedral coordination

Cobalt Molybdate, CoMoO$_6$
Mo atom octahedral coordination

Mo L$_3$-edge XANES

- $\Delta E$ tetrahedral = 2.2-2.5 eV; $\Delta E$ octahedral = 3.25-4.2 eV
Time Evolution of XANES

In situ temperature programmed reduction of Re$_2$O$_7$/Al$_2$O$_3$

Re L$_3$-edge XANES

Heat in H$_2$ flow
Q-XANES & D-XANES

Quick XANES

- Slew monochromator continuously to obtain a XANES spectrum in few seconds.
- All modes of detection.

Dispersive XANES

- Polychromatic beam dispersed onto linear detector.
- XANES spectrum in msec.
- Transmission only.
- Need extremely uniform samples.
Micro-XANES

• Specialized application of XANES.
• Focus x-ray beam to <10\(\mu\)m diameter.

• Applications:
  – Speciation of metals in soils, sediments and organisms
  – Grazing incidence studies of cations and anions on surfaces
  – Time-resolved studies of reactions on surfaces and interfaces
  – High temperature studies (trace elements in melts)
  – Oxidation states of planetary material
  – High pressure phases (diamond anvil cell)

• See http://www.bnl.gov/x26a/ for information.
Analysis of Mixtures

• XANES useful technique to quantitatively determine composition of a mixture of species.

• Useful for following time evolution of species during a chemical reaction.

• Two most common methods:
  – Least squares linear combination fitting
  – Principal component analysis
Least Squares Linear Combination Fitting

• Use a linear combination of spectra of various reference samples.

• Allows quantification of species in multiple-component mixture from their fingerprint in the XANES region.

• Use a least-squares algorithm to refine the sum of a given number of reference spectra to an experimental spectrum.

• Simple method, easy to implement.

• Must have spectra of the reference compounds.
Linear Combination Fitting

Fit experimental data to linear combination of known reference compounds

TPR-XANES of *in situ* reduction of Ce$_{0.7}$Zr$_{0.3}$ oxide

LC-XANES fit to determine amount of Ce(III) and Ce(IV) present as function of temperature
**Principal Component Analysis**

- Used for many years in other chemical spectroscopy. First published reference in XANES 1992*.

- Traditional approach: choose pure model standard, fit edges to these standards, but…

- How many standards are needed?

- How do we know models are reasonable?

- If you have wrong group of standards…there is no way to get the correct answer…

Principal Component Analysis

• PCA estimates number of distinct species in a series of spectra.

• Used as a first stage of analysis.

• Based on linear algebra - each spectrum represented as a vector.

• Goal is to find number of components that can reproduce the experimental spectra to within experimental error.

• Growing popularity in XANES spectroscopy.
## Summary

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<tr>
<th>XANES is a much larger signal than EXAFS</th>
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<td>XANES can be done at lower concentrations, and less-than-perfect sample conditions.</td>
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<th>XANES is easier to crudely interpret than EXAFS</th>
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<td>For many systems, the XANES analysis based on linear combinations of known spectra from “model compounds” is sufficient.</td>
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<td>More sophisticated linear-algebra techniques, such as principal component analysis can be applied to XANES spectra.</td>
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<th>XANES is harder to fully interpret than EXAFS</th>
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<td>The exact physical and chemical interpretation of all spectral features is still difficult to do accurately, precisely, and reliably.</td>
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<td>This situation is improving…..</td>
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