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### Sample preparation and sample environments

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### Why talking about sample preparation?

- 1. Over a half of lost time during synchrotron beamtime is due to the sample quality deficiencies
- 2. If you do not recognize how sample quality affects the XAS data, you will collect and potentially publish misleading results

This presentation does not include math: see, e.g., <u>http://gbxafs.iit.edu/training/XAFS\_sample\_prep.pdf</u> for more quantitative explanation of the underlying phenomena





# Speak with beamline staff!

### Sample preparation – victim of XAS popularity

XAS is versatile: almost anything can be measured (solids, liquids, gases, soil, dust, batteries, fuel cells, meteorites, wastewater, colloidal suspensions, protein solutions etc)

### Your **specimen** must be transformed into a **sample!**

What are the characteristics of a good sample?

- 1. Concentration compatible with one of the common detection methods
- 2. Doesn't introduce spectra distortion
- 3. Can be handled \*

Oftentimes, the samples are just what they are...

### Detection methods: transmission vs. fluorescence

Transmission: photon flux detected upstream of the sample and downstream of the sample



Fluorescence detector

Fluorescence: emitted photons are collected by the detector placed on the side of the sample

For transmission absorbance (per Beer-Lambert law) is  $\mu = -log(I_t/I_0)$ 

For fluorescence, in thin dilute sample approximation, intensity is proportional to  $\mu$ 

### Sample size

Considerations:

- 1. Beam size at a specific beamline
  - ISS: 1 mm (H) x 1 mm(V) or 50-100 µm (FWHM).
  - QAS: 10 mm(H) 1.5 mm(V) x 10 mm(H), or ~0.5 mm (FWHM),
  - BMM: 8 mm (H) by 1 mm (V) or 300 µm (FWHM).
- 2.Standard holder
- 3.Ease of handling

### Uniformity is the key!!

Basic rule of sample preparation: sample shall be made **homogenous** so that every X-ray photon interacts identically no matter what area on the sample it hits.

#### Attenuation Length:

The depth into the material measured along the beam where the X-ray flux falls to 1/e of its value at the surface.



To be homogenous enough means particles shall be smaller than the attenuation length

Disclaimer: sometimes you cannot make smaller particles, e.g., certain sieve fraction of a catalyst to prevent gas flow blockage.

### Uniformity is the key!! (cont.)

Imagine your sample looks like that

There are regions of varying density and pinholes: the X-ray leakage problem leads to spectra distortion:

- 1. White line intensity decreases
- Photons seeing different effective thicknesses create a "disorder" that manifests itself in artificially exaggerated <sup>2</sup>
- 3. Noise in the data increases

#### Takeaway

Particles shall be smaller than attenuation length and uniformly dispersed over the sample

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Simulated spectra for particle sizes from 30 um to 0.3



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### Sample preparation for transmission experiments

It is recommended that the sample for transmission measurements has an edge step of 1-1.5 (however, 0.1-1 can be acceptable if the sample is of good quality)



### Sample preparation for transmission experiment

Finding right amount of sample with Hephaestus

Hephaestus Help Hephaestus Formulas: compute total cross sections of materials Density Materials Absorption Formula CoFe2O4 List of typical Xvlene a/cm^3 Density | 🛊 | 5 0 Xenon materials Formulas Eneray YAG 7800 Energy (100 Zerodur Compute Ion chambers Zinc eV above the Results Acetone Air edge energy) Data element number barns/atom cm^2/am Alcohol (Ethyl) ۲ Co 1.000 33808.991 345.519 Alcohol (Methyl) Fe 325.464 2.000 30183.487 Transitions Alcohol (Propyl) 0 4.000 333.532 12.553 Absorption length Aluminum This weighs 234.633 amu. Edae finde Argon Absorption length = 8.2 micron at 7800 eV. Beryllium Boron Nitride A sample of 1 absorption length with area of 1 square Line finde Carbon (Diamond) cm requires 4.079 milligrams of sample at 7800.00 eV. Carbon (Graphite) Unit edge step length at c ←edge (7709.0 eV) is 28.3 Standards Copper microns Fluorite 2-The Elam database and the full cross-sections we F' and F' Gold used in the calculation. Helium Amount of This calculation uses the Elam data resource and full cross sections material/cm<sup>2</sup>

Chemical formula

for the edge

jump of 1

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### Making pellets

Most reliable ex situ sample preparation method is making pellets made using a hydraulic (or hand) press (similar to IR pellets)

Inert binder/dilutant may be added to maintain correct concentration

Rule-of-thumb 1 Total amount of sample+binder >100 mg (for ease of handling)

Rule of thumb 2 Grind as long as you can... and then some

Rule of thumb 3 Sample+binder mixture color shall be uniform

Note on binders

BN, cellulose, zeolite, polyethylene glycol (Inert is a relative term: BN catalyzed methane oxidation, zeolites are catalysts, PEG polymerizes under pressure...)

### Making pellets

Equipment



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#### Mind binder absorption





### Other sample prep methods

Capillaries

Typical materials – borosilicate glass, quartz, Kapton

Mind the wall thickness: attenuation can be strong Even though *in situ* comes later:

If heating is involved:

- 1. Glass up to 600 °C
- 2. Quartz up to ~ 1300 °C (becomes brittle)
- 3. Kapton up to 300 °C





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## Capillaries can be sealed with epoxy, quartz wool or wax

### Other powder sample prep methods

• Fill the wells in the sample holder with powder

• Sample on adhesive tape My personal opinion: It shall not be done

If you have **absolutely** no choice, the powder shall be ground and spread as thin as possible, and the tape folded

- Always take the spectrum of the tape you might be surprised
- Mind the absorption of the tape may be critical at lower side of the spectra





### When to measure in fluorescence

- 1. Sample has low loading yielding edge jump of 0.1 or less
- 2. Sample is supported on opaque substrate (thin film on Si wafer)
- 3. It is undesirable or impossible to manipulate the sample (art object, meteorite etc.).
- 4. Low energy edges cannot be measured in transmission

A good fluorescence sample:

 Homogenous – while effects of leakage and non-uniformity are not as severe as in transmission, it is still desirable.

### Self-absorption in fluorescence measurement

Spectral distortion (reduced white line and EXAFS wiggles) is the result of changing penetration depth of X-ray beam when scanning through the edge





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### Fluorescence measurements best practices

This is particularly important for dilute sample

- Avoid sample holders/sample environments made of same or neighbor elements (and mind overlap between 3d K edges and 4f and 5d L edges )
- Check windows/tapes/capillary sealing materials for the element of interests (very low levels of Mn, Zn in Mylar, Fe in Kapton have been found)
- Avoid fluorescence from in-line calibration foil getting fluorescence detector. Use apertures upstream of the reference foil.
- When compounds containing lighter elements are used to support heavier elements (e.g., Pd/CuO), non resonance fluorescence from the support obscures the signal from the element of interest (energy discriminating detector is not always a solution). Use AI foil for reducing lower energy signal.



In the past lead tape was used extensively for *ad hoc* shielding. It's banned now due to health and safety reasons. Tin foil ca be used instead – not as blocking but as pliable.

### Thin films

Thin films are supported on a substrate Measured in fluorescence mode Films are positioned at grazing or near-grazing incidence angle to the beam to spread the beam over a larger surface and increase surface sensitivity

Substrate and films are often crystalline, leading to Bragg peaks hitting the detector. We often spin the samples to "smear" Bragg peaks





### Liquid samples

### Naturally homogenous (unless they are suspensions) Appropriate containment

- 1. Capillaries sample
- 2. Sample holders with wells or holes sealed with adhesive tape



#### Beware of beam damage (more on it later)

### Drop casting

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A method to deposit catalyst slurry onto a porous flat substrate, e.g., carbon paper.

Common in electrocatalysis/fuel cells

Usually produces highly inhomogeneous samples due to "coffee stain" effect



### Sample environments

What we want to expose the sample to:

- 1. Temperature
- 2. Reactive gases /pressure
- 3. Reactive liquids
- 4. Pressure
- 5. Voltage/current
- 6. Magnetic field

We also want to keep the sample intact (limit exposure to X-rays/minimize X-ray beam damage

### Klausen (capillary) flow cell

#### Advantages:

- Compatible with different capillaries
- True plug flow catalytic reactor
- Fast cooldown

#### Disadvantages

- Hard to do operando low catalysis amount, and low gas flow rates are needed to observe conversion
- Temperature profile is poorly defined



### Another capillary flow cell

#### Advantages:

- Compatible with different capillaries
- True plug flow catalytic reactor
- Better temperature stability
- Fast cooldown

#### Disadvantages

- Hard to do operando low catalysis amount, and low gas flow rates are needed to observe conversion
- Temperature profile is poorly defined



### Other temperature cells (commercia

Commercial designs originally for optical microscopy

- Temperatures up to 1500 C
- Some design allow cooling
- While OK for gas treatment, not suitable for catalysis due to large dead volume.
- Some design allow cooling to -195 C



### Other temperature cell

Users are allowed to design their own cells, as long as they are safe and have windows/openings for the all/preferred detection methods.





### Electrochemical cells

Typically user designed:

- Accommodate variable number of electrodes
- Location of electrode being investigated is important (due to electrolyte absorption)
- Bubble formation is sometimes inevitable
- These cells leak!!



Electrochem. Comm., 94, 2018, 14-17

### **Battery holders**

While not sample environments by themselves, they make battery cells handling much easier.

Allow for high throughput measurements

Side note: adding X-ray transparent windows to a coin cell may change field distribution and chemistry





### Cryostats

## Allow for low temperature measurements down to a few K

- Thermal motion decreases, and EXAFS can be recorded up to higher k.
- Improves sample radiation sensitivity
- Cooling is slow (can be >1h)
- Sample handling is difficult, especially if samples need to be mounted frozen



### High pressure gas and liquid cells

When catalytic (or other) experiment requires reactant pressures above ambient, custom cell are required.



### Really high pressure cell

Diamond anvil cells allow to create pressures up to 700 Gpa (usually below 200 Gpa)

- Recreate the pressure existing deep inside planets and to synthesize materials and phases not observed under normal ambient conditions.
- Diffraction from diamonds can (and does) lead to very intense spikes in the spectra



### Liquid jet

To prevent sample damage in liquids, the sample needs to be constantly exchanged.

The liquid is pumped with syringe or/peristaltic pump through a nozzle

One can protect air sensitive liquids by enclosing it within another fluid in a "coaxial" jet







### Conclusions

- 1. Sample preparation should be considered in the early stages of experiment planning
- 2. Discuss detection method(s) with beamline staff and tailor the samples and sample environment(s) to them.
- 3. Discuss how to mount the sample environment if you bring your own.



# Speak with beamline staff!