

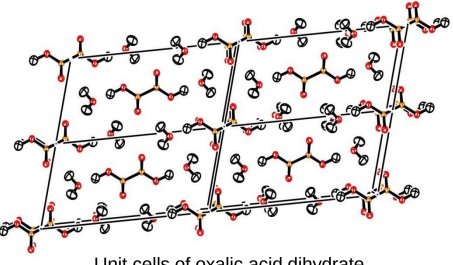
# **Single Crystal Diffraction**

Arthur J. Schultz Argonne National Laboratory

National School on Neutron and X-Ray Scattering August, 2013



#### What is a crystal?



Unit cells of oxalic acid dihydrate

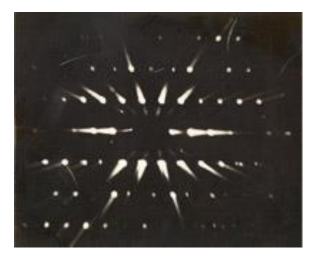
- Atoms (molecules) pack together in a regular pattern to form a crystal.
- Periodicity: we superimpose (mentally) on the crystal structure a repeating lattice or unit cell.
- A lattice is a regular array of geometrical points each of which has the same environment.



Quartz crystals

# Why don't the X-rays scatter in all directions?





X-ray precession photograph (Georgia Tech, 1978).

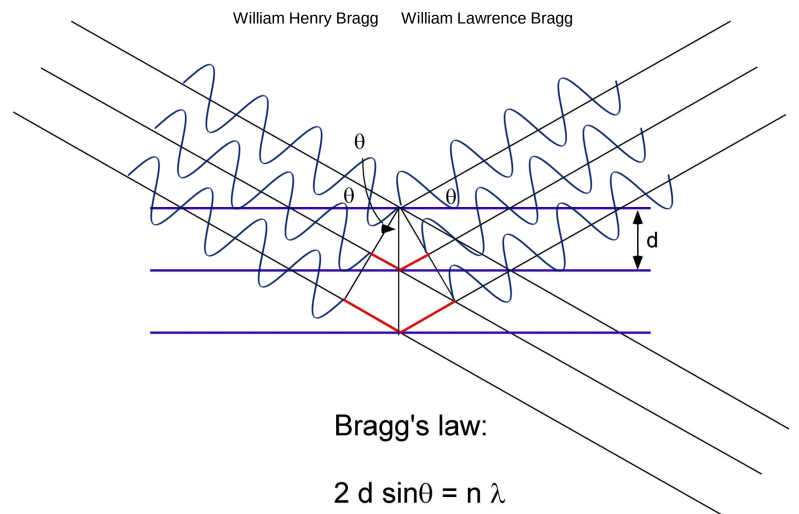
- X-rays (and neutrons) have wave properties.
- A crystal acts as a diffraction grating producing constructive and destructive interference.

# Bragg's Law



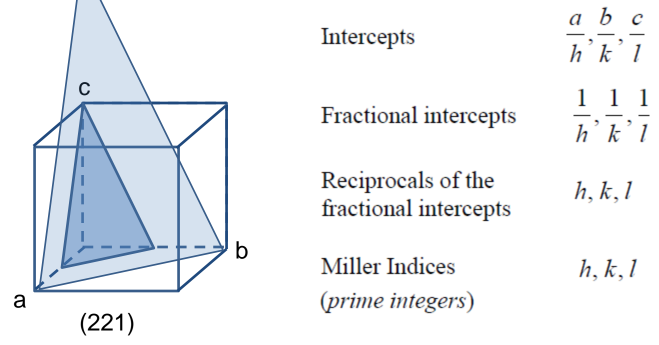


Jointly awarded the 1915 Nobel Prize in Physics

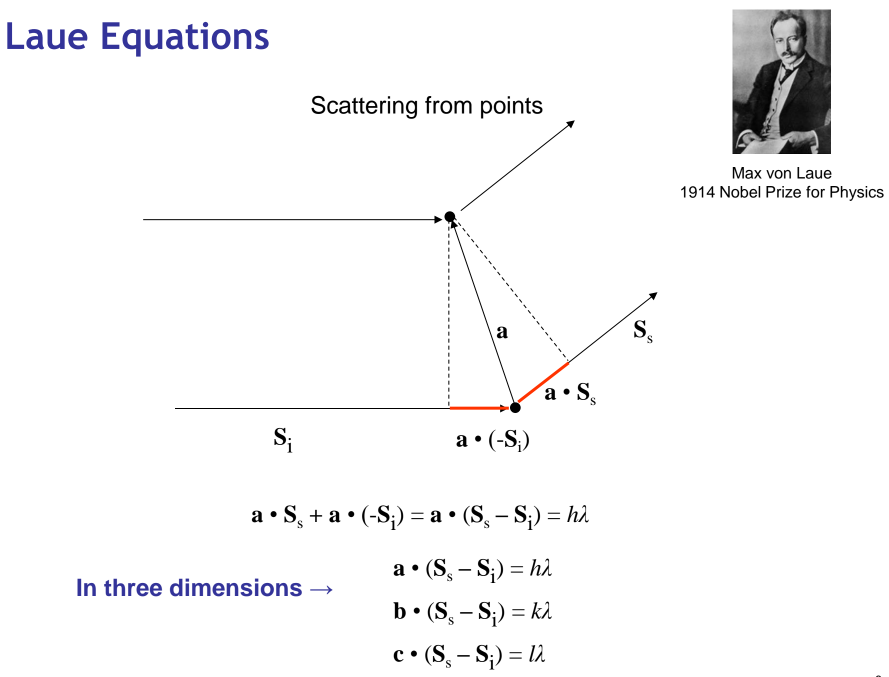


### **Crystallographic Planes and Miller Indices**

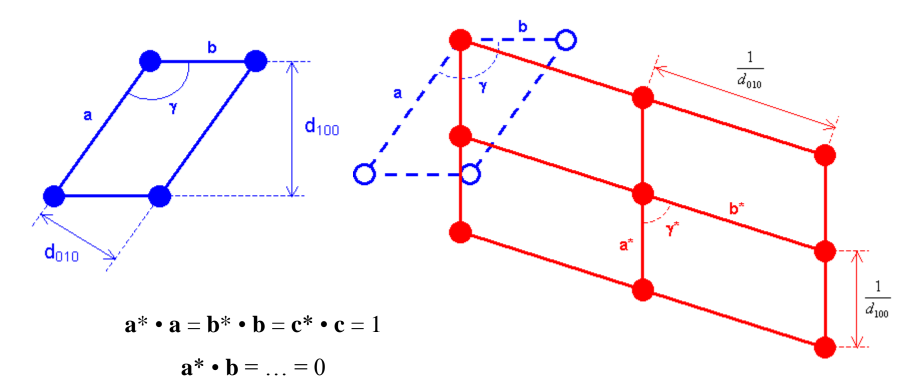
The (*hkl*) plane intercepts a/h, b/k, c/l on crystallographic axes X, Y, Z, where h, k, l are relatively prime integers.



*d*-spacing = spacing between origin and first plane or between neighboring planes in the family of planes.



## **Real and reciprocal Space**

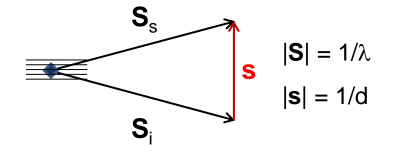


Laue equations:

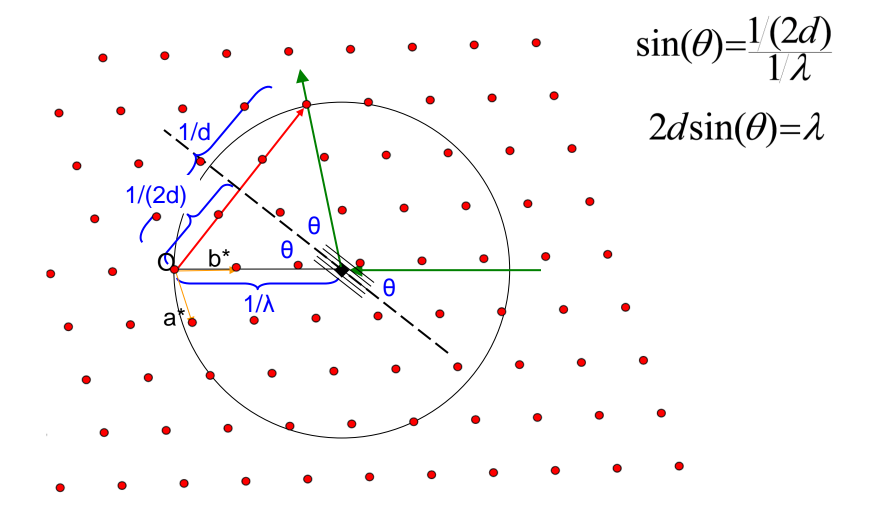
 $\mathbf{a} \cdot (\mathbf{S}_{s} - \mathbf{S}_{i}) = h\lambda, \text{ or } \mathbf{a} \cdot \mathbf{s} = h$  $\mathbf{b} \cdot (\mathbf{S}_{s} - \mathbf{S}_{i}) = k\lambda, \text{ or } \mathbf{b} \cdot \mathbf{s} = k$  $\mathbf{c} \cdot (\mathbf{S}_{s} - \mathbf{S}_{i}) = l\lambda, \text{ or } \mathbf{c} \cdot \mathbf{s} = l$ 

where

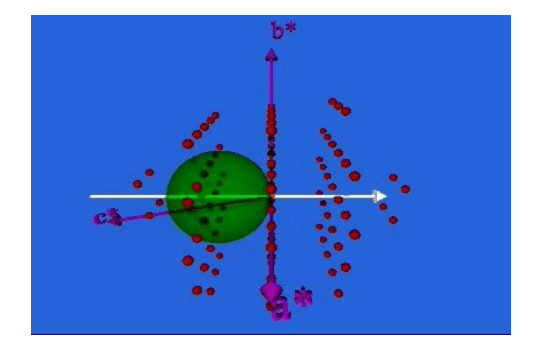
$$\mathbf{s} = (\mathbf{S}_{s} - \mathbf{S}_{i})/\lambda = h\mathbf{a}^{*} + k\mathbf{b}^{*} + l\mathbf{c}^{*}$$



# The Ewald Sphere

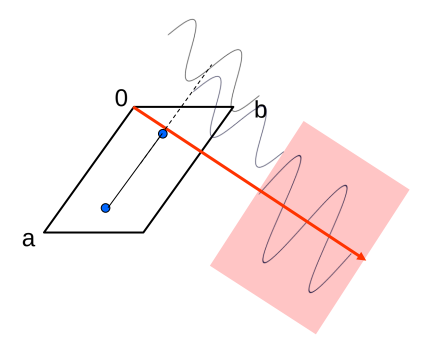


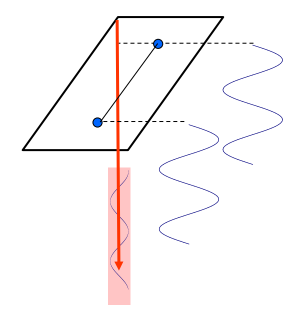
#### The Ewald sphere: the movie



Courtesy of the CSIC (Spanish National Research Council). http://www.xtal.iqfr.csic.es/Cristalografia/index-en.html

#### **Bragg Peak Intensity**





Relative phase shifts related to molecular structure.

$$F_{hkl} = \sum_{i} b_{i} \exp(2\pi i \mathbf{s} \cdot \mathbf{r})$$
$$F_{hkl} = \sum_{i} b_{i} \exp[2\pi i (hx + ky + lz)]$$

$$\rho(xyz) = \frac{1}{V} \sum_{hkl} F_{hkl} e^{-2\pi i (hx+ky+lz)}$$

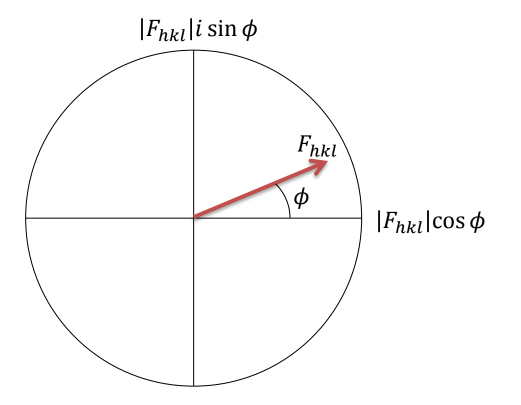
 $b_{i}$  is the neutron scattering length. It is replaced by  $f_{i}$ , the x-ray form factor.

#### The phase problem

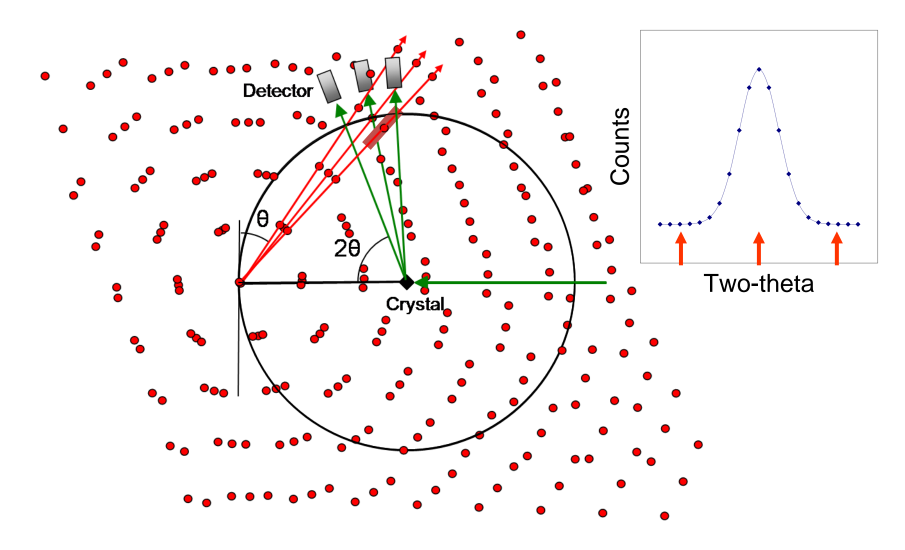
$$I_{hkl} = F_{hkl}\overline{F_{hkl}} = |F_{hkl}|e^{i\phi}|F_{hkl}|e^{-i\phi} = |F_{hkl}|^2$$

Euler's formula:

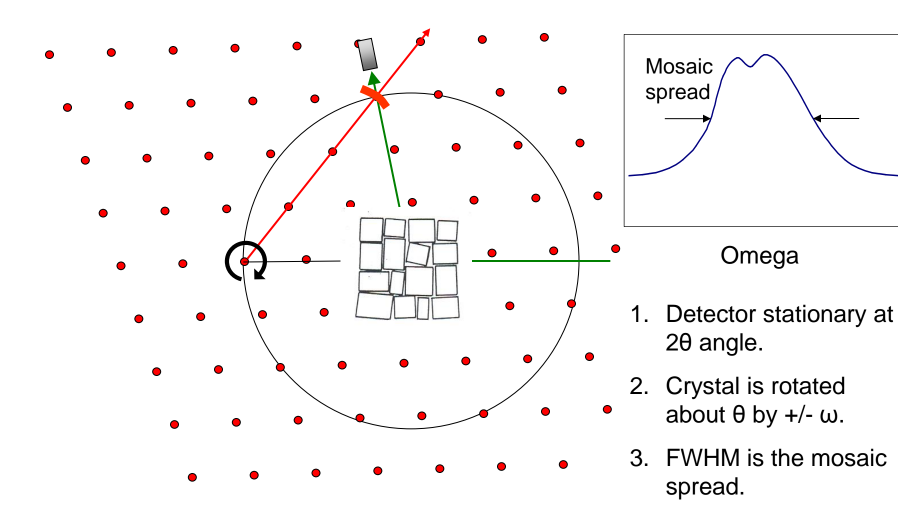
$$F_{hkl} = |F_{hkl}|e^{i\phi} = |F_{hkl}|(\cos\phi + i\sin\phi) = A + iB$$
$$I_{hkl} = (A + iB)(A - iB) = A^2 + B^2$$



# $\theta$ -2 $\theta$ Step Scan

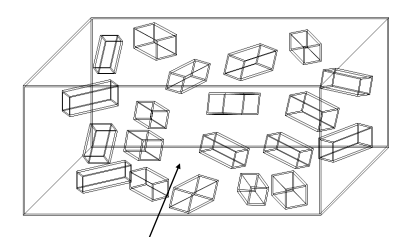


# **Omega Step Scan**



Something completely different - polycrystallography

What is a powder? - polycrystalline mass



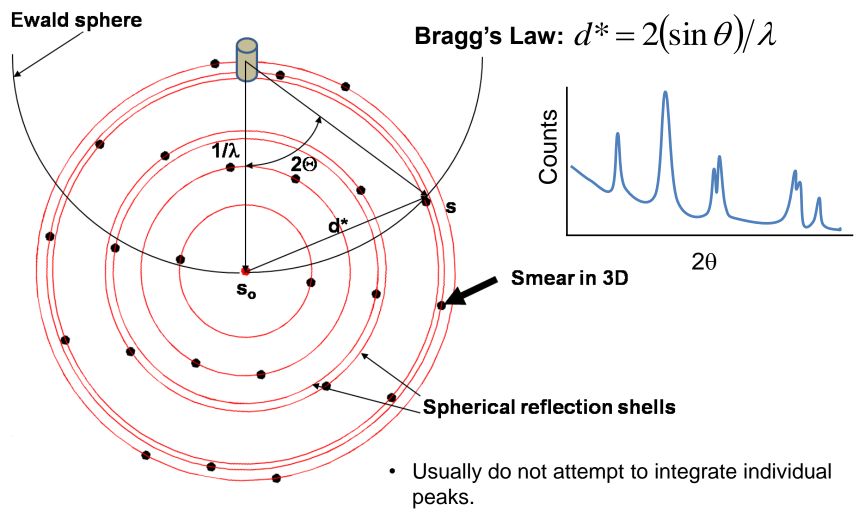
All orientations of crystallites possible

Sample:  $1\mu$ l powder of  $1\mu$ m crystallites - ~10<sup>9</sup> particles

Packing efficiency – typically 50% Spaces – air, solvent, etc.

Single crystal reciprocal lattice - smeared into spherical shells

# **Powder Diffraction**



 Instead, fit the spectrum using Rietveld profile analysis. Requires functions that describe the peak shape and background.

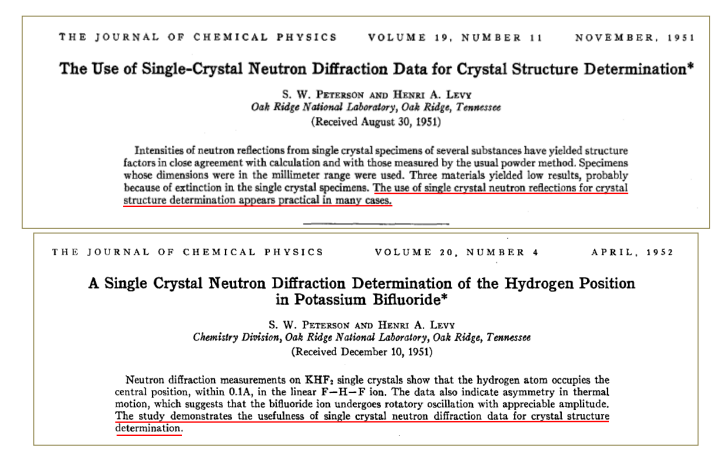
# Why do single crystal diffraction (vs. powder diffraction)?

- Smaller samples
  - neutrons: 1-10 mg vs 500-5000 mg
  - x-rays: μg vs mg
- Larger molecules and unit cells
- Neutrons: hydrogen is ok for single crystals, powders generally need to be deuterated
- Less absorption
- Fourier coefficients are more accurate based on integrating wellresolved peaks
- Uniquely characterize non-standard scattering superlattice and satellite peaks (commensurate and incommensurate), diffuse scattering (rods, planes, etc.)

#### But:

- Need to grow a single crystal
- Data collection can be more time consuming

#### Some history of single crystal neutron diffraction



- 1951 Peterson and Levy demonstrate the feasibility of single crystal neutron diffraction using the Graphite Reactor at ORNL.
- 1950s and 1960s Bill Busing, Henri Levy, Carroll Johnson and others wrote a suite of programs for singe crystal diffraction including ORFLS and ORTEP.
- 1979 Peterson and coworkers demonstrate the single crystal neutron timeof-flight Laue technique at Argonne's ZING-P' spallation neutron source.

#### **The Orientation Matrix**

Acta Cryst. (1967). 22, 457

Angle Calculations for 3- and 4- Circle X-ray and Neutron Diffractometers\*

BY WILLIAM R. BUSING AND HENRI A. LEVY

Chemistry Division, Oak Ridge National Laboratory, Oak Ridge, Tennessee 37830, U.S.A.

(Received 13 June 1966)

Methods are derived for calculations useful in the operation of 3- and 4-circle X-ray or neutron singlecrystal diffractometers. These include: (1) establishing the sample orientation from the cell parameters and the observed angles for two reflections, or from the observed angles for three reflections only, (2) calculating the angles for observing a given reflection either in a special setting or at a specified azimuthal angle, (3) obtaining the vectors needed for calculating absorption corrections, and (4) using observations of several reflections to refine cell and orientation parameters by the method of le ast squares.

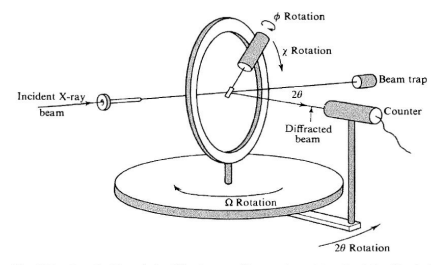


Fig. 5.29. A typical four-circle diffractometer. The counter rotates about the 2 $\theta$  axis in one plane and the crystal may be orientated in any way by the three axes of rotation  $\phi$ ,  $\chi$  and  $\Omega$ .

$$\mathbf{B} = \begin{pmatrix} b_1 & b_2 \cos \beta_3 & b_3 \cos \beta_2 \\ 0 & b_2 \sin \beta_3 & -b_3 \sin \beta_2 \cos \alpha_1 \\ 0 & 0 & 1/a_3 \end{pmatrix}$$

1

**U** is a rotation matrix relating the unit cell to the instrument coordinate system.

The matrix product **UB** is called the *orientation matrix*.

# **Picker 4-Circle Diffractometer**



# Kappa Diffractometer

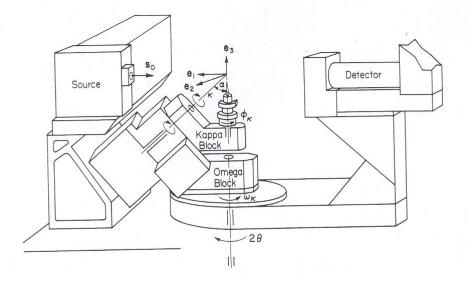


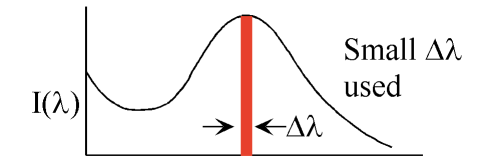
FIGURE 6-13. Kappa geometry. Adapted from operating manual for ENRAF-NONIUS CAD 4 diffractometer (angles  $\omega$ ,  $\phi$ , and  $\varkappa$  are opposite in sign to those of Enraf-Nonius). (By permission of ENRAF-NONIUS Service Corp., Bohemia, New York.)



Brucker AXS: KAPPA APEX II

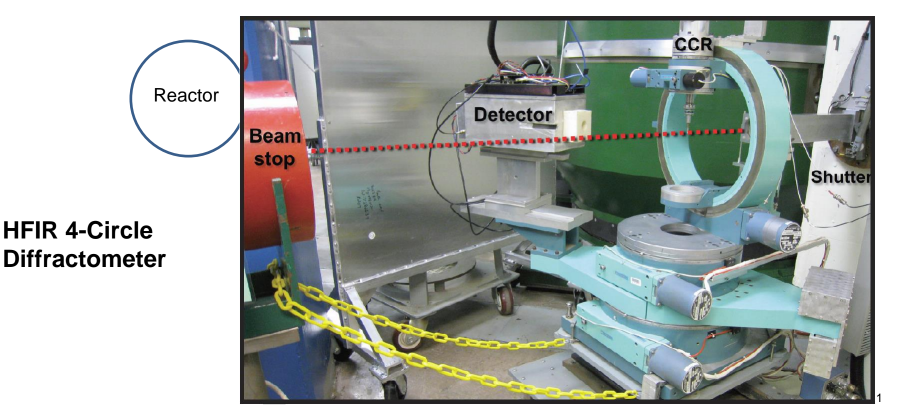
- Full 360° rotations about  $\omega$  and  $\phi$  axes.
- Rotation about κ axis reproduces quarter circle about χ axis.

### Monochromatic diffractometer

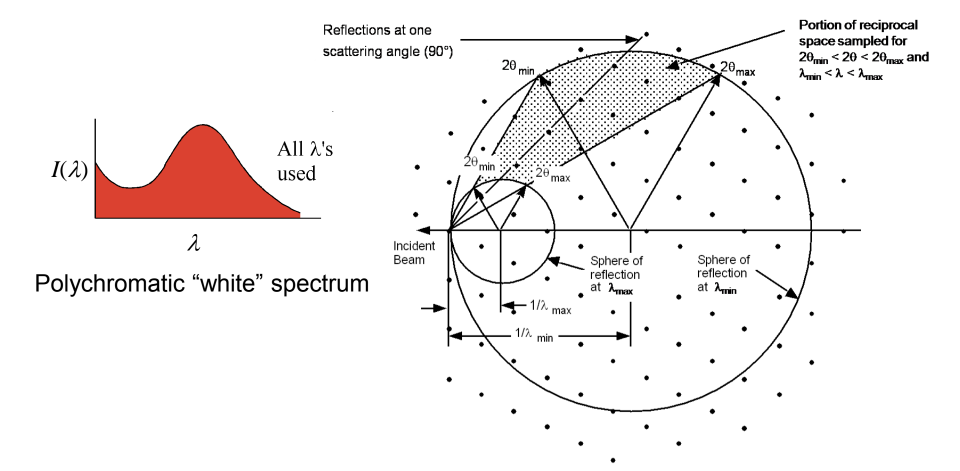


- Rotating crystal
- Vary  $\sin\theta$  in the Bragg equation:

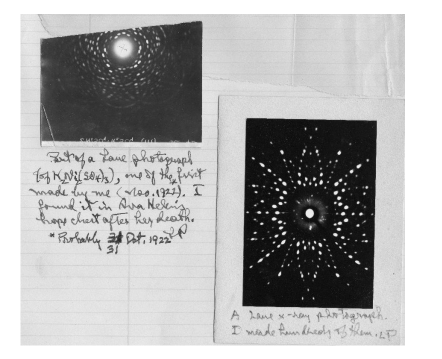
 $2d\sin\theta = n\lambda$ 



# Laue diffraction



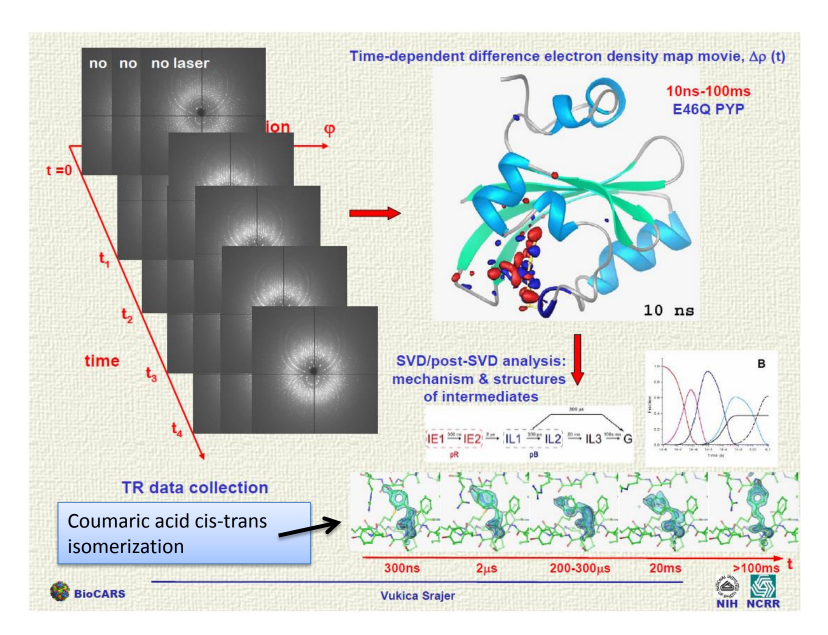
# Laue photo from white radiation



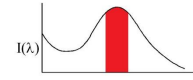
X-ray Laue photos taken by Linus Pauling



# Time-resolved X-ray Laue diffraction of photoactive yellow protein at BioCARS using pink radiation



# **Quasi-Laue Neutron Image Plate Diffractometer**

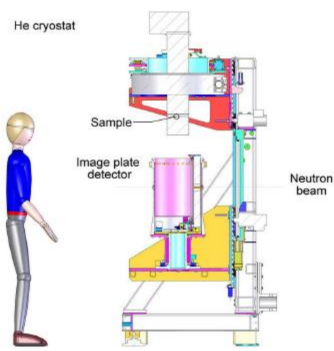


Select  $\Delta\lambda/\lambda$  of 10-20%

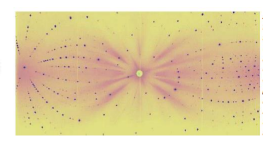
#### 2012 at HFIR: IMAGINE



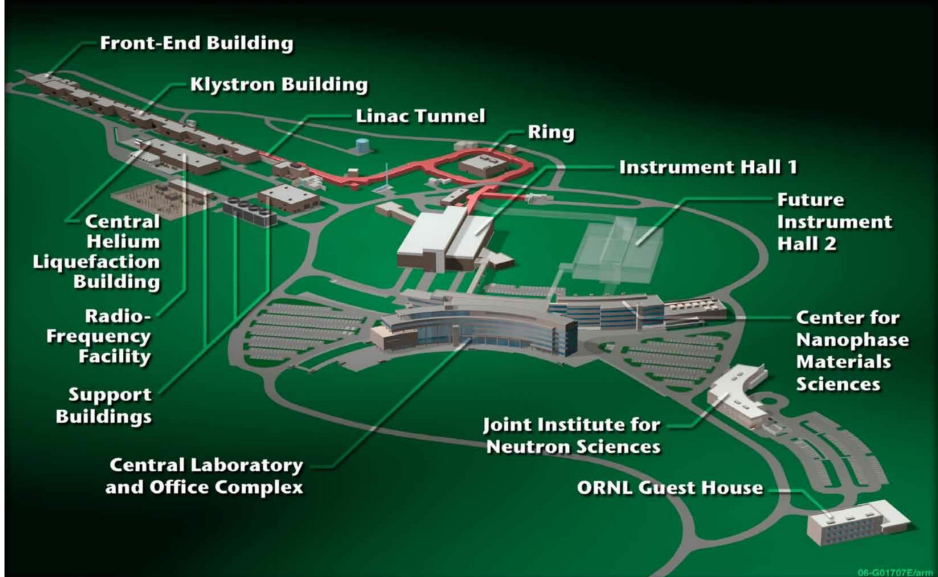
General view of the QLD



QLD schematic (open position)

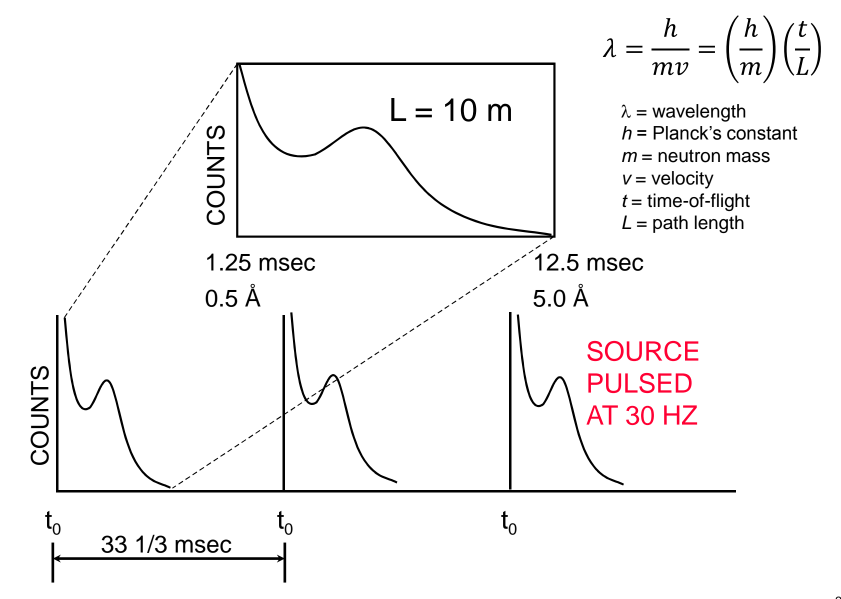


A typical Laue diffraction pattern from FeTa<sub>2</sub>O<sub>6</sub> just above the 3-D ferroelectric ordering temperature (Chung et al. J. Phys.: Condens. Matter, 16 (2004) 1-17). The faint cross of radial streaks about the central hole, which allows passage of the transmitted neutron beam, arises from 2-D magnetic ordering. Results from the Laue diffractometer VIVALDI at the ILL

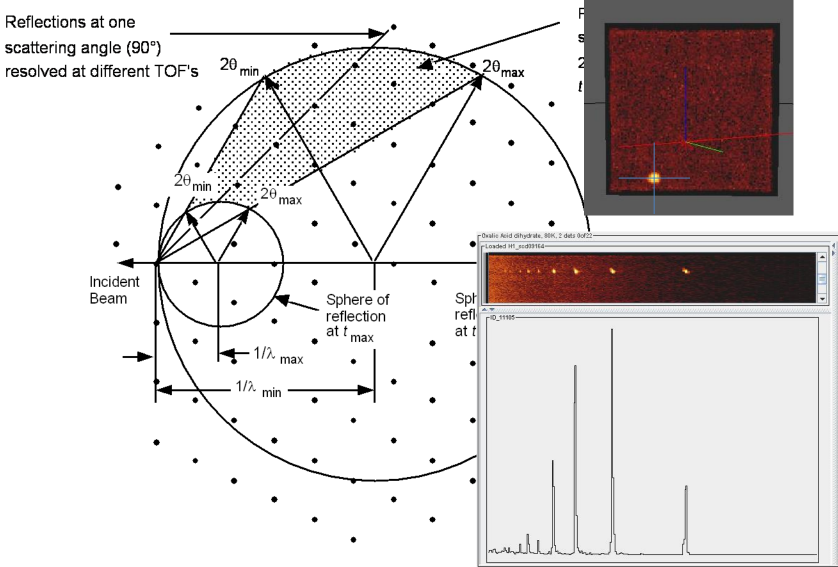


derrorean

## **Pulsed Neutron Incident Spectrum**



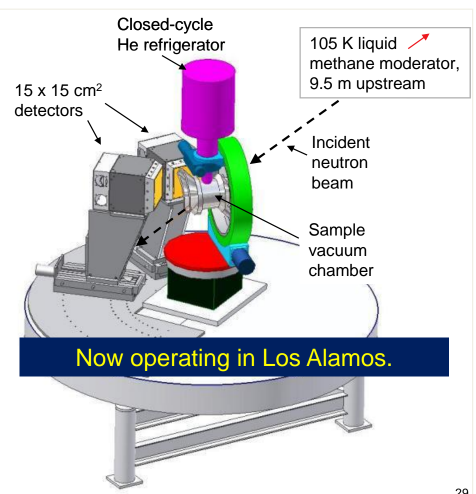
# **Time-of-Flight Laue Technique**



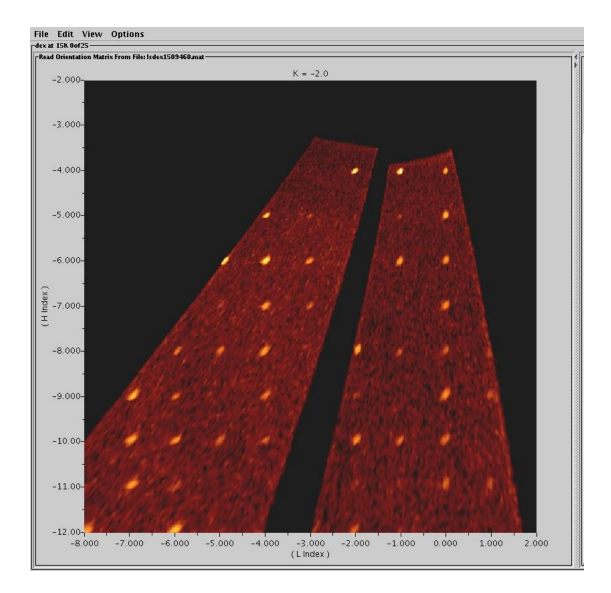
#### **SCD Instrument Parameters**

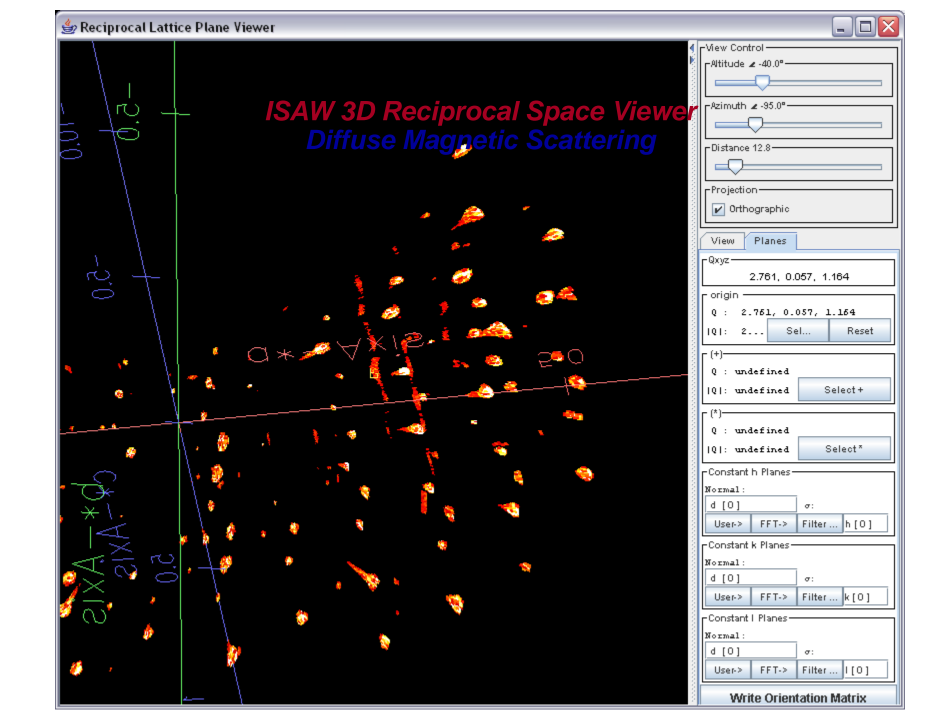
Moderator	liq. methane at 105
Source frequency	30 Hz
Sample-to-moderator dist.	940 cm
Number of detectors	2
Detector active area	155 x 155 mm <sup>2</sup>
Scintillator	GS20 <sup>6</sup> Li glass
Scintillator thickness	2 mm
Efficiency @ 1 Å	0.86
Typical detector channels	100 x 100
Resolution	1.75 mm
Detector 1:	
angle	75°
sample-to-detector dist.	23 cm
Detector 2:	
angle	120°
sample-to-detector dist.	18 cm
Typical TOF range	1–25 ms
wavelength range	0.4–10 Å
d-spacing range	~0.3–8 Å
TOF resolution, $\Delta t/t$	0.01
Sample Environments	
Hot-Stage Displex: 4-900 K	
Displex Closed Cycle Helium Refrigerator: 12–473 K	
Furnaces: 300–1000 K	
Helium Pressure Cell Mounted on Displex:	
0–5 kbar @ 4–300 K	

Detector distances on locus of constant solid angle in reciprocal space.



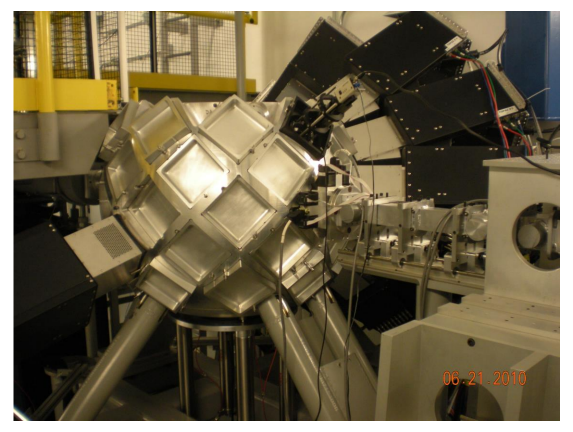
# **ISAW hkl plot**



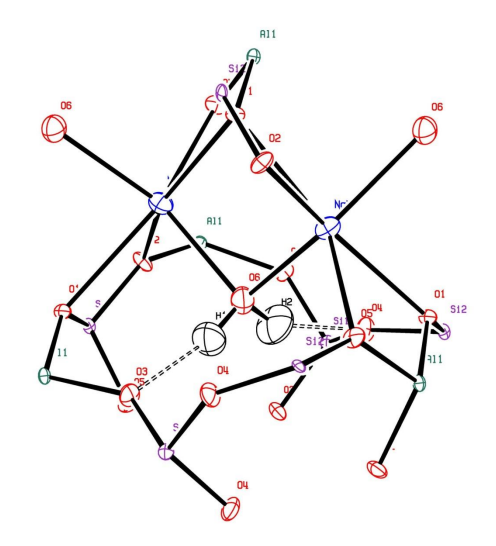


# Topaz

- Project Execution Plan requires a minimum of 2 steradian (approx. 23 detectors) coverage.
- Each detector active area is 150 mm x 150 mm.
- Secondary flight path varies from 400 mm to 450 mm radius and thus cover from 0.148 to 0.111 steradian each.



# **Natrolite structure from TOPAZ data**



# **Outline of single crystal structure analysis**

- Collect some initial data to determine the unit cell and the space group.
  - Auto-index peaks to determine unit cell and orientation
  - Examine symmetry of intensities and systematic absences
- Measure a full data set of observed intensities.
- Reduce the raw integrated intensities,  $I_{hkl}$ , to structure factor amplitudes,  $|F_{obs}|^2$ .
- Solve the structure.
- Refine the structure.

#### Data reduction - single crystal TOF Laue

<u>Data reduction</u>: convert raw integrated intensities,  $I_{hkl}$ , into relative structure factor amplitudes,  $|F_{hkl}|^2$ .

 $I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) (V_s/V_c^2) |F_{hkl}|^2 \lambda^4 / \sin^2 \theta$ 

k = scale factor

 $\phi(\lambda)$  = incident flux spectrum

 $\varepsilon(\lambda)$  = detector efficiency as a function of wavelength  $\lambda$ 

 $A(\lambda)$  = sample absorption

 $y(\lambda)$  = secondary extinction correction

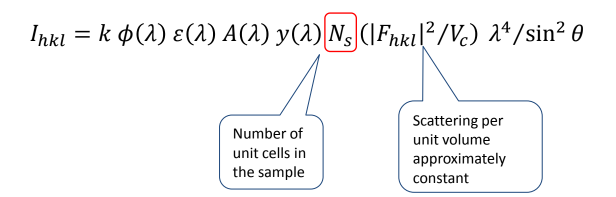
 $V_{\rm s}$  = sample volume

 $V_{\rm c}$  = unit cell volume

#### Intensity vs. sample volume and unit cell volume

 $I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) (V_s/V_c^2) |F_{hkl}|^2 \lambda^4 / \sin^2 \theta$ 

 $I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) (V_s/V_c) (|F_{hkl}|^2/V_c) \lambda^4 / \sin^2 \theta$ 



#### Wilson plot

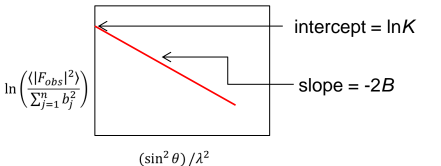
$$\langle |F_{hkl}|^2 \rangle = \sum_{j=1}^n b_j^2 = n \langle b_j^2 \rangle$$

$$\langle |F_{obs}|^2 \rangle = K \sum_{j=1}^n b_j^2 e^{-2B \, (\sin^2 \theta)/\lambda^2}$$

n = number of atoms in the unit cell  $b_j$  = neutron scattering length, or  $f_i$  = x-ray form factor

*K* = scale factor *B* = temperature or thermal parameter

$$\ln\left(\frac{\langle |F_{obs}|^2\rangle}{\sum_{j=1}^n b_j^2}\right) = \ln K - 2B\left(\sin^2\theta\right)/\lambda^2$$



$$V_c = \sum_{j=1}^n v_j = n \langle v_j \rangle$$

 $\langle |F_{hkl}|^2 \rangle / V_c = \langle b_j^2 \rangle / \langle v_j \rangle$ 

 $V_c$  = unit cell volume  $v_j$  = volume of atom j

For crystals containing similar types of atoms in similar ratios, this is a constant.

#### Lorentz factor

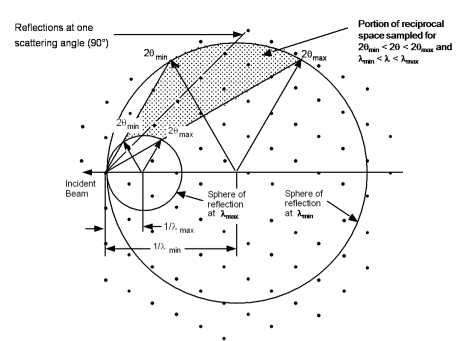
The Lorentz factor is geometric integration factor related to the time or angular range during which a peak is reflecting.

Laue integration:

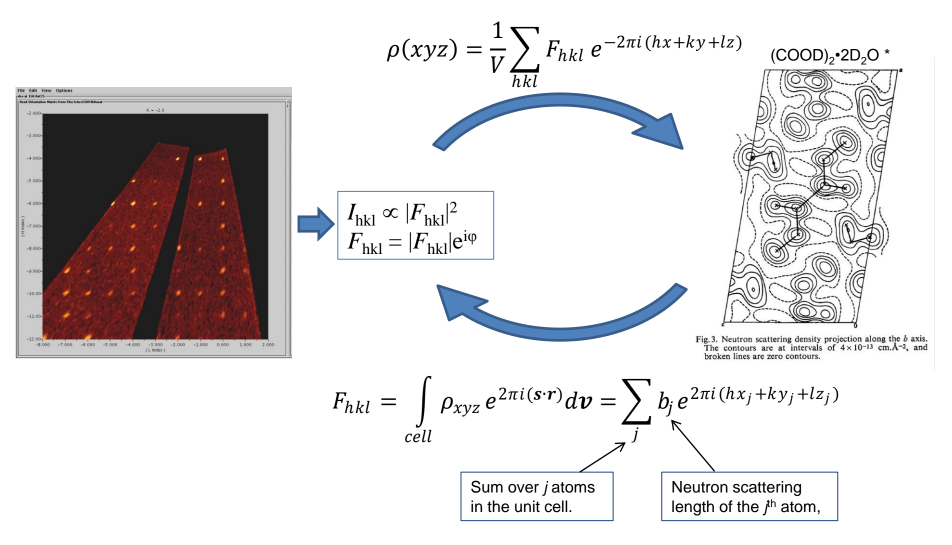
$$I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) N_s (|F_{hkl}|^2 / V_c) \lambda^4 / \sin^2 \theta$$
$$I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) N_s (|F_{hkl}|^2 / V_c) \lambda^2 d^2 / 4$$

Constant wavelength integration:

 $I_{hkl} = k \phi(\lambda) \varepsilon(\lambda) A(\lambda) y(\lambda) N_s (|F_{hkl}|^2 / V_c) \lambda^3 / \sin 2\theta$ 



#### Fourier transforms

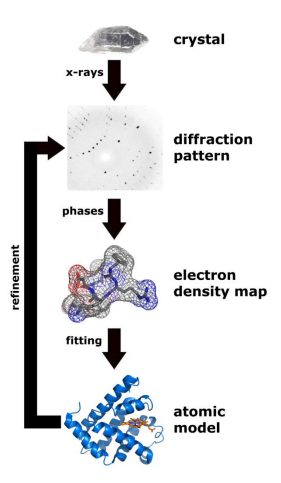


\* Iwasaki, Iwasaki and Saito, Acta Cryst. 23, 1967, 64.

# Solutions to the phase problem

- Patterson synthesis using the  $|F_{obs}|^2$  values as Fourier coefficients
  - Map of inter-atom vectors
  - Also called the heavy atom method
- Direct methods
  - Based on probability that the phase of a third peak is equal to the sum of the phases of two other related peaks.
  - J. Karle and H. Hauptman received the 1985 Nobel Prize in Chemistry
- Shake-and-bake
  - Alternate between modifying a starting model and phase refinement
- Charge flipping
  - Start out with random phases.
  - Peaks below a threshold in a Fourier map are flipped up.
  - Repeat until a solution is obtained
- MAD
  - Multiple-wavelength anomalous dispersion phasing
- Molecular replacement
  - Based on the existence of a previously solved structure with of a similar protein
  - Rotate the molecular to fit the two Patterson maps
  - Translate the molecule

# **Structure Refinement**



$$\chi^{2} = \sum_{hkl} w (|F_{0}| - |F_{c}|)^{2}$$

$$F_{hkl} = \sum_{i} b_{i} \exp\left[2\pi i (hx_{i} + ky_{i} + lz_{i})\right] \exp\left[-8\pi^{2} U_{i} \sin^{2} \theta / \lambda^{2}\right]$$

#### GSAS, SHELX, CRYSTALS, OLEX2, WinGX...

Nonlinear least squares programs. Vary atomic fractional coordinates x, y, z and temperature factors U (isotropic) or  $u_{ij}$  (anisotropic) to obtain best fit between observed and calculated structure factors.

Workflow for solving the structure of a molecule by X-ray crystallography (from http://en.wikipedia.org/wiki/Xray\_crystallography).

## Neutron single crystal instruments in the US

- SNAP @ SNS: high pressure sample environment (<u>http://neutrons.ornl.gov/instruments/SNS/SNAP/</u>)
- TOPAZ @ SNS: small molecule to small protein, magnetism, future polarized neutron capabilities (<u>http://neutrons.ornl.gov/instruments/SNS/TOPAZ/</u>)
- Four-Circle Diffractometer (HB-3A) @ HFIR: small molecule, high precision, magnetism (<u>http://neutrons.ornl.gov/instruments/HFIR/HB3A/</u>)
- MaNDi (Macromolecular Neutron Diffractometer) @ SNS: neutron protein crystallography, commissioning in 2012 (<u>http://neutrons.ornl.gov/instruments/SNS/MaNDi/</u>)
- IMAGINE (Image-Plate Single-Crystal Diffractometer) @ HFIR: small molecule to macromolecule crystallography, commissioning in 2012 (<u>http://neutrons.ornl.gov/instruments/HFIR/imagine/</u>)
- SCD @ Lujan Center, Los Alamos: general purpose instrument, currently not available due to budget constraints (<u>http://lansce.lanl.gov/lujan/instruments/SCD/index.html</u>)
- PCS (Protein Crystallography Station) @ Lujan Center, Los Alamos: neutron protein crystallography (http://lansce.lanl.gov/lujan/instruments/PCS/index.html)

# Books and on-line tutorials

- M. F. C. Ladd and R. A. Palmer, Structure Determination by X-ray Crystallography, Third Edition, Plenum Press, 1994.
- J. P. Glusker and K. N. Trueblood, Crystal Structure Analysis: A Primer, 2<sup>nd</sup> ed., Oxford University Press, 1985.
- M. J. Buerger, *Crystal-structure analsysis*, Robert E. Krieger Publishing, 1980.
- George E. Bacon, *Neutron Diffraction*, 3<sup>rd</sup> ed., Clarendon Press, 1975.
- Chick C. Wilson, Single Crystal Neutron Diffraction From Molecular Crystals, World Scientific, 2000.
- Interactive Tutorial about Diffraction: www.totalscattering.org/teaching/
- An Introductory Course by Bernhard Rupp: http://www.ruppweb.org/Xray/101index.html

