

MSE-238  
Structure of Materials

Week 8 - Diffraction  
Spring 2025

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# Overview

Diffraction:

- The scattering vector
- Ewald sphere
- Interaction with different radiation
- sample types and measurement methods
- influence of imperfect microstructure

mixed with repetition of crystallography on examples

- hard sphere model
- interstitial sites
- crystal symmetry
- crystal planes, family of planes
- quasi-crystal

→(Hammond Chapter 8-10)

→ Phil Willmott, “Introduction to Synchrotron Radiation” Chapter 6

# Scattering/Diffraction: the scattering vector

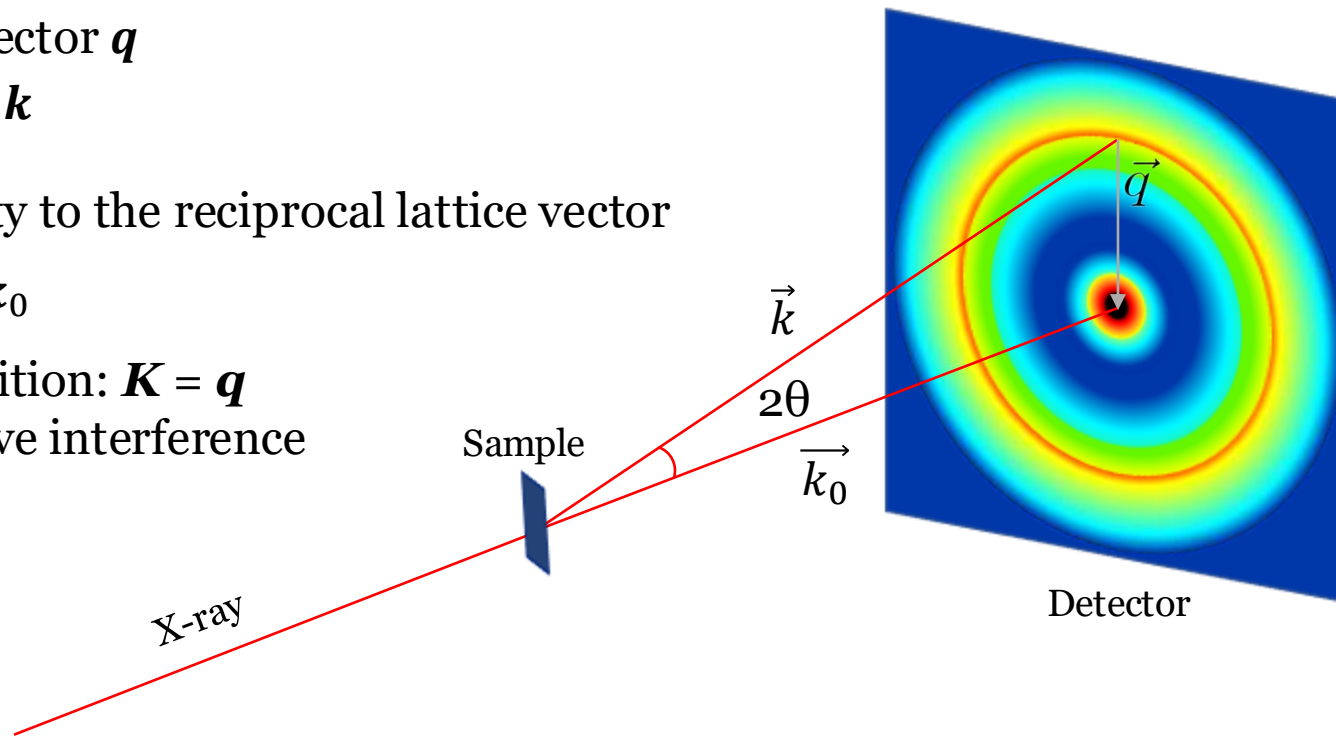
scattering vector  $\mathbf{q}$

$$\mathbf{q} = \mathbf{k}_0 - \mathbf{k}$$

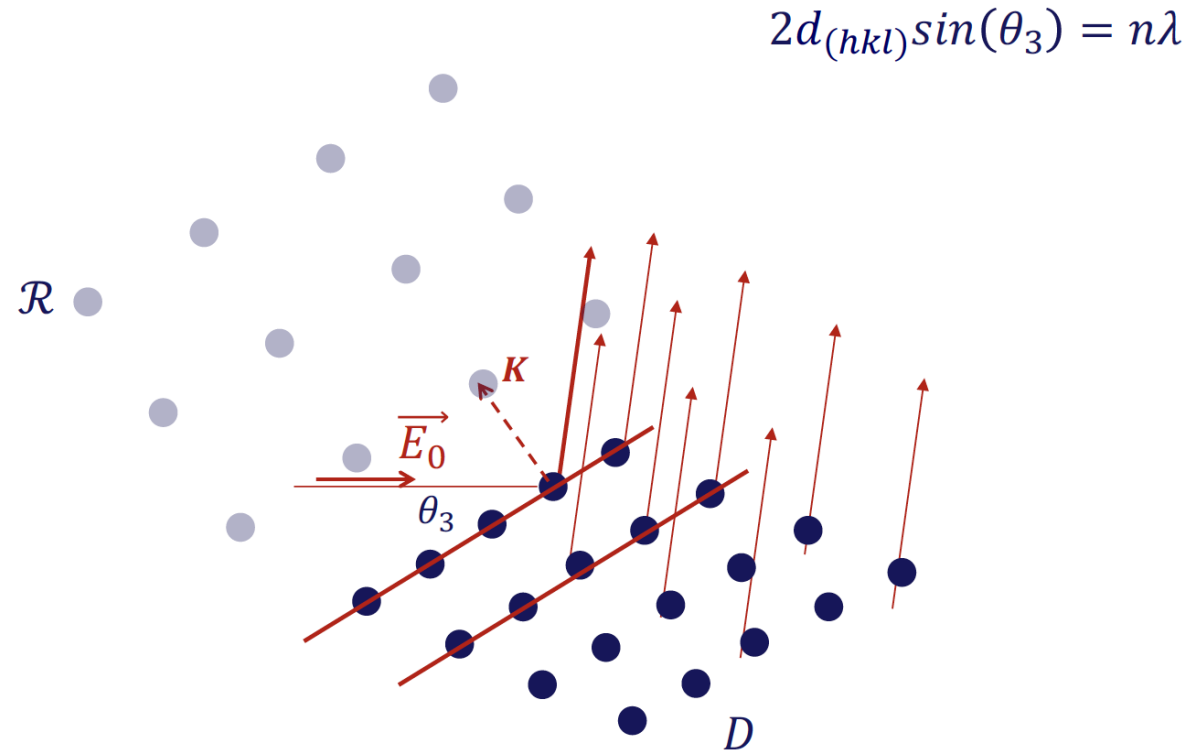
see similarity to the reciprocal lattice vector

$$\mathbf{K} = \mathbf{k} - \mathbf{k}_0$$

→ Laue condition:  $\mathbf{K} = \mathbf{q}$   
constructive interference

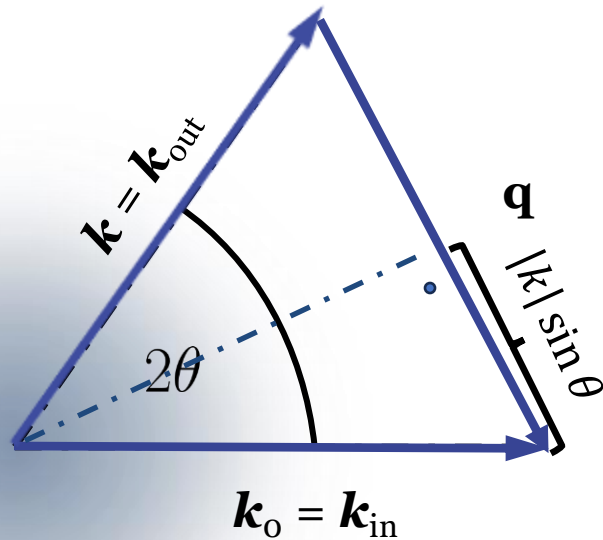


# Repetition: The reciprocal space lattice



reciprocal lattice vector  $K = k - k_0$

# Elastic scattering



wave vector  $k = \frac{2\pi}{\lambda}$

elastic scattering: no loss in photon energy but direction of the photon can change with a scattering angle  $2\theta$

$$|k_{in}| = |k_{out}|$$

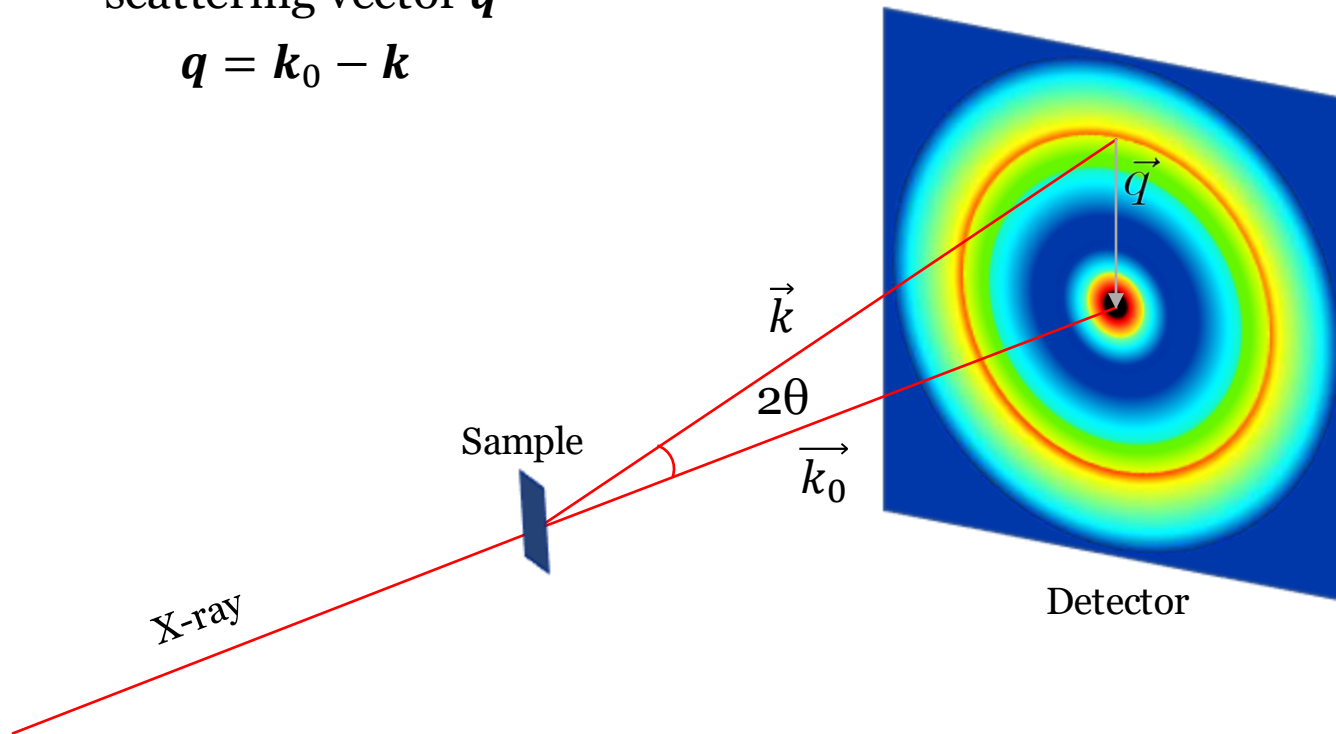
scattering vector  $q = k_0 - k$

$$q = 2|k| \sin \theta = \frac{4\pi \sin \theta}{\lambda}$$

# Scattering/Diffraction

scattering vector  $q$

$$\mathbf{q} = \mathbf{k}_0 - \mathbf{k}$$



$$|\vec{q}| = q = \frac{4\pi \sin(\theta)}{\lambda}$$

light  $\lambda = 400$  to  $600$  nm

X-ray tube  $\lambda = 1$  to  $2$  Å

Cu Kα =  $1.5406$  Å

synchrotron  $\lambda = 0.1$  to  $5$  Å

thermal neutrons  $\lambda = 1$  to  $10$  Å

electrons  $\lambda = 0.025$  Å

X-ray energy mostly given in keV

Electronvolt = eV

Energy of an electron after being accelerated from rest in a potential of 1 V

$$1 \text{ eV} = 1.6022 \times 10^{-19} \text{ J}$$

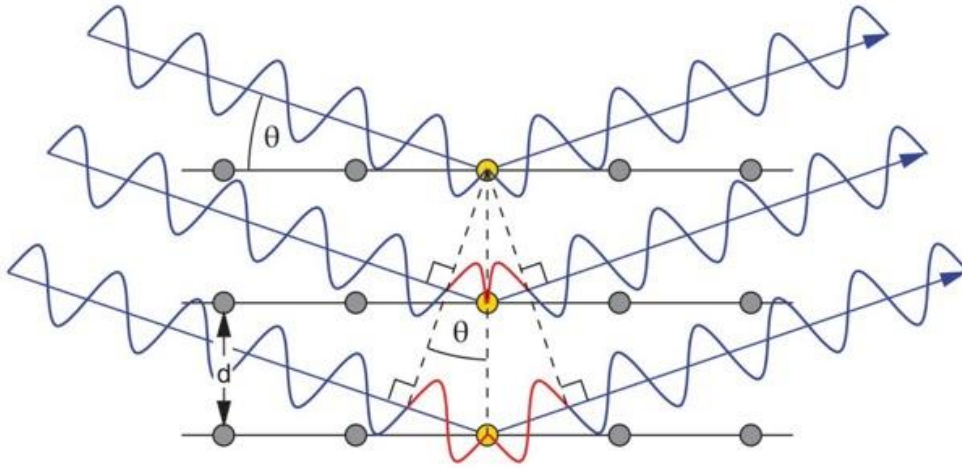
$$E = hc / \lambda$$

$h$  is Planck's constant ( $6.6261 \times 10^{-34}$  Js)

$c$  is the speed of light ( $2.9979 \times 10^8$  m/s).

$$\lambda [\text{Å}] = 12.3984 / E [\text{keV}]$$

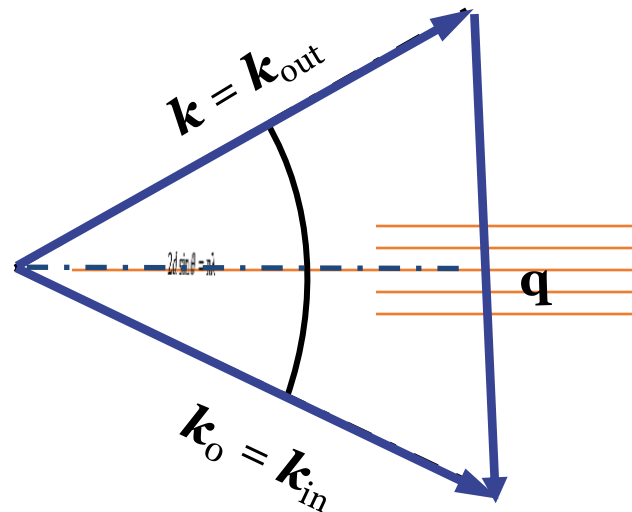
# Bragg law



The scattering vector  $\mathbf{q}$  always lies perpendicular to the scattering planes if the Bragg condition is fulfilled

the angle subtended by  $\mathbf{k}_{\text{in}} = 2\pi/\lambda$  (or  $\mathbf{k}_{\text{out}}$ ) and the scattering planes is  $\theta$ .

The scattering angle is  $2\theta$



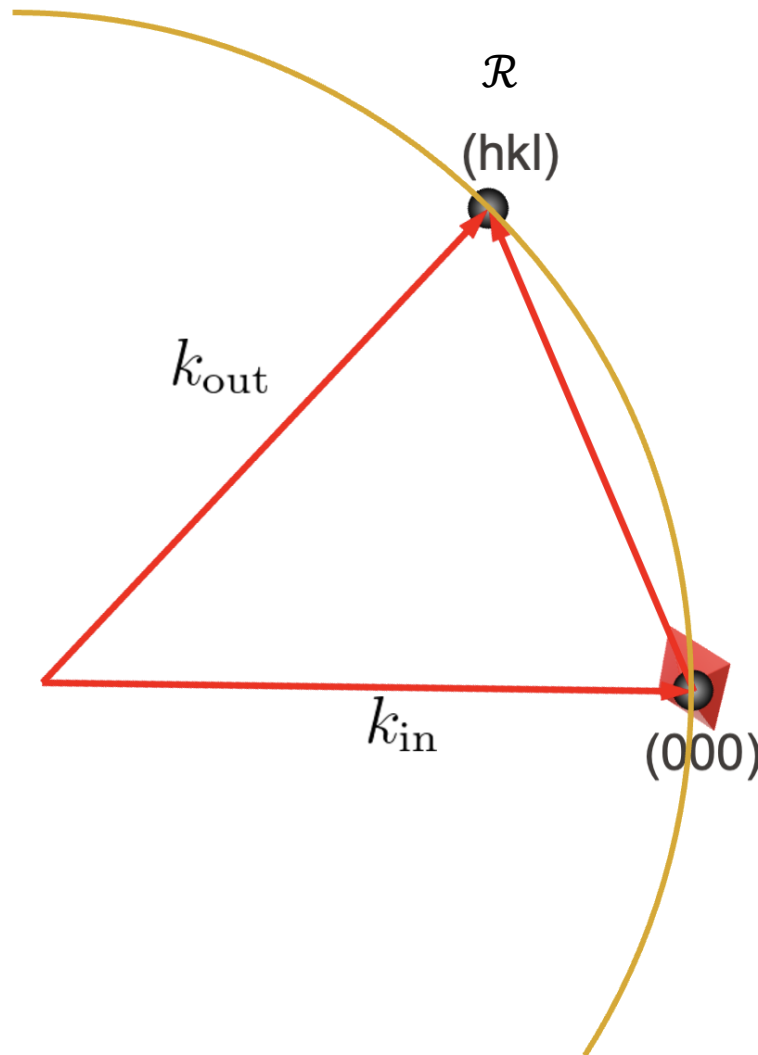
Bragg law:  $2d \sin \theta = n\lambda$

$$|\mathbf{q}| = q = \frac{4\pi \sin(\theta)}{\lambda}$$

$$d = n \frac{2\pi}{q}$$

independent of wave length  
(experimental condition)

# Bragg condition and the Ewald sphere



elastic scattering:  $|k_{in}| = |k_{out}| = 2\pi/\lambda$

To see a diffraction peak @ (hkl) :

the Bragg points (000), which is at the position of the direct incoming beam

and the Bragg point (hkl) from the reciprocal space lattice

must lie on a sphere of radius equal  $|k|$  (the wave vector of the experiment) in reciprocal space  $\mathcal{R}$ , the so called Ewald sphere

Join at [menti.com](https://menti.com/join/38486882) | use code 3848 6882

Radius of Ewald sphere: X-rays vs. electrons?



# Which radiation?

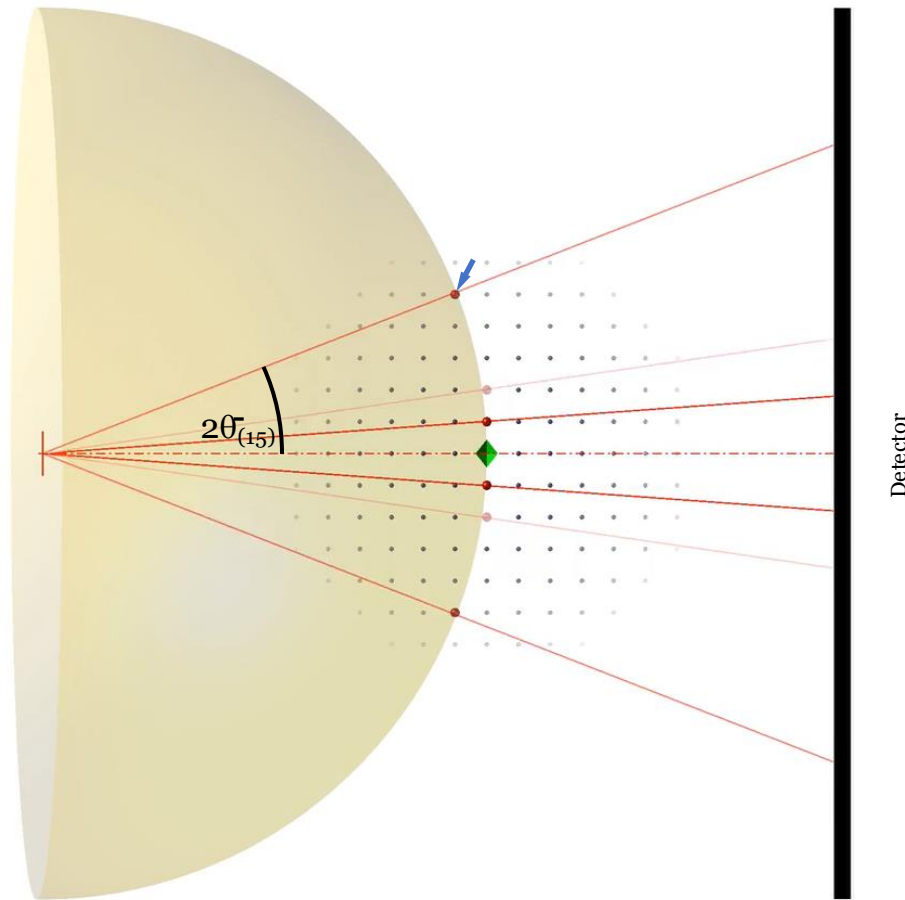
visible light will not be a good probe since the wavelength is 400nm-700nm, which is much bigger than the distance between the atoms, typically of the order of a few  $10^{-10}$ m. For instance,

- the lattice parameter of the element Ni is  $3.6 \cdot 10^{-10}$ m,
- the size of an atom is about  $1.4 \cdot 10^{-10}$ m.

	Energy	wavelength
Neutrons	1 – 5 meV (cold)	9 - 4 Å
	25 – 50 meV (thermal)	1.8 – 1.3 Å
Xrays	100keV	0.12 Å ( hard Xrays)
	40 keV	0.31 Å
	5 keV	2.48 Å (soft Xrays)
Electrons	200 keV	0.025 Å

# The Ewald sphere

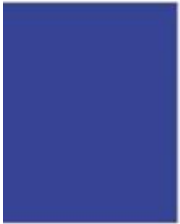
Bragg law defines on a purely geometrical basis for which angles constructive interference **can** occur



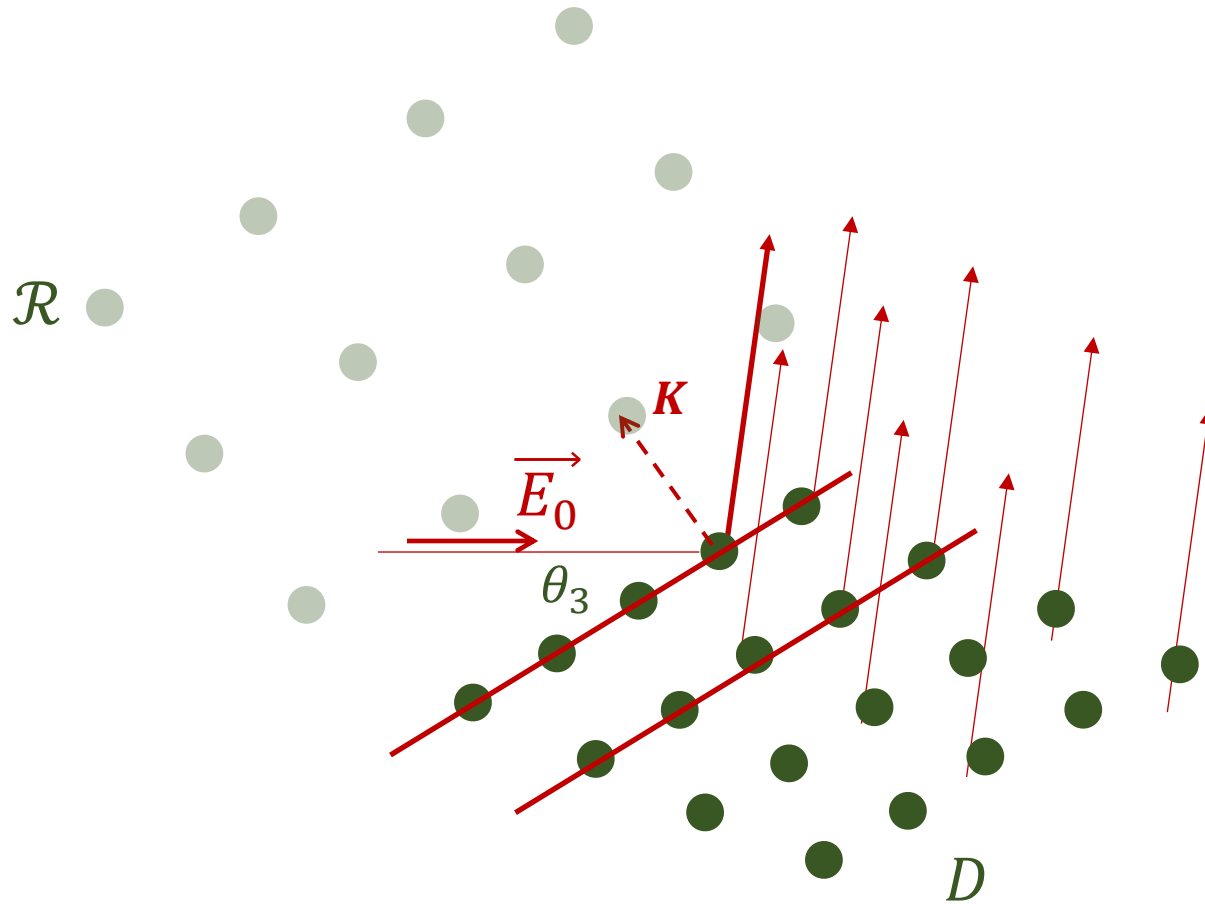
Philip Willmott: Synchrotron and X-ray Free Electron Laser (Part 2)

# Sample types

single  
crystal

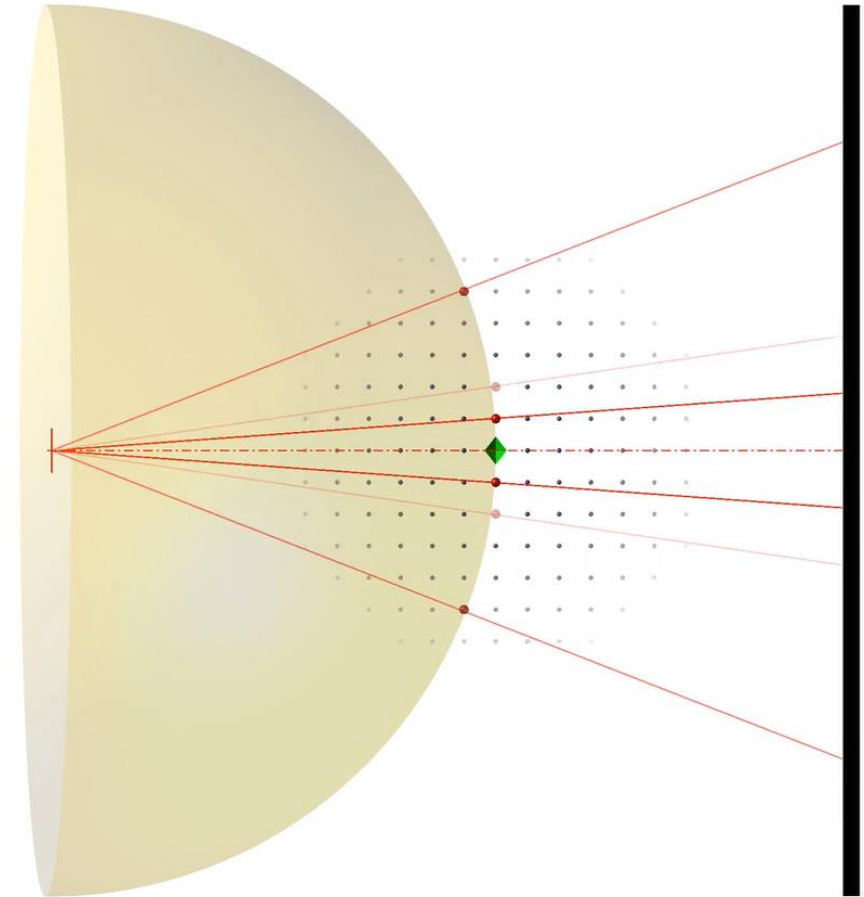


$$2d_{(hkl)}\sin(\theta_3) = n\lambda$$



# Single crystal diffraction – rotation method

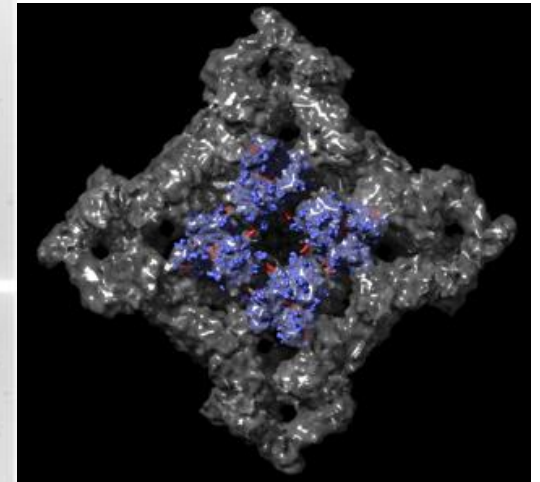
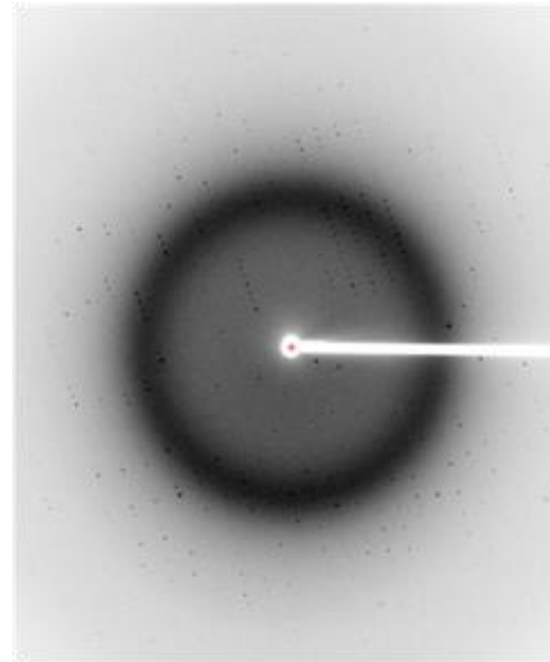
- monochromatic radiation is used
- By rotating the crystal around an axis perpendicular to the incident beam ( $\phi$ ), diffraction maxima pass through the surface of the Ewald sphere and are registered on a 2D x-ray detector



# Application: protein crystallography

study the three-dimensional structure of biological macromolecules.

Data collection: wave length = 0.097nm, Canadian Light Source - 1 second per frame, total of ~360 frames, step size 0.5 degrees



Courtesy: F. Van Petegem, UBC, Canada

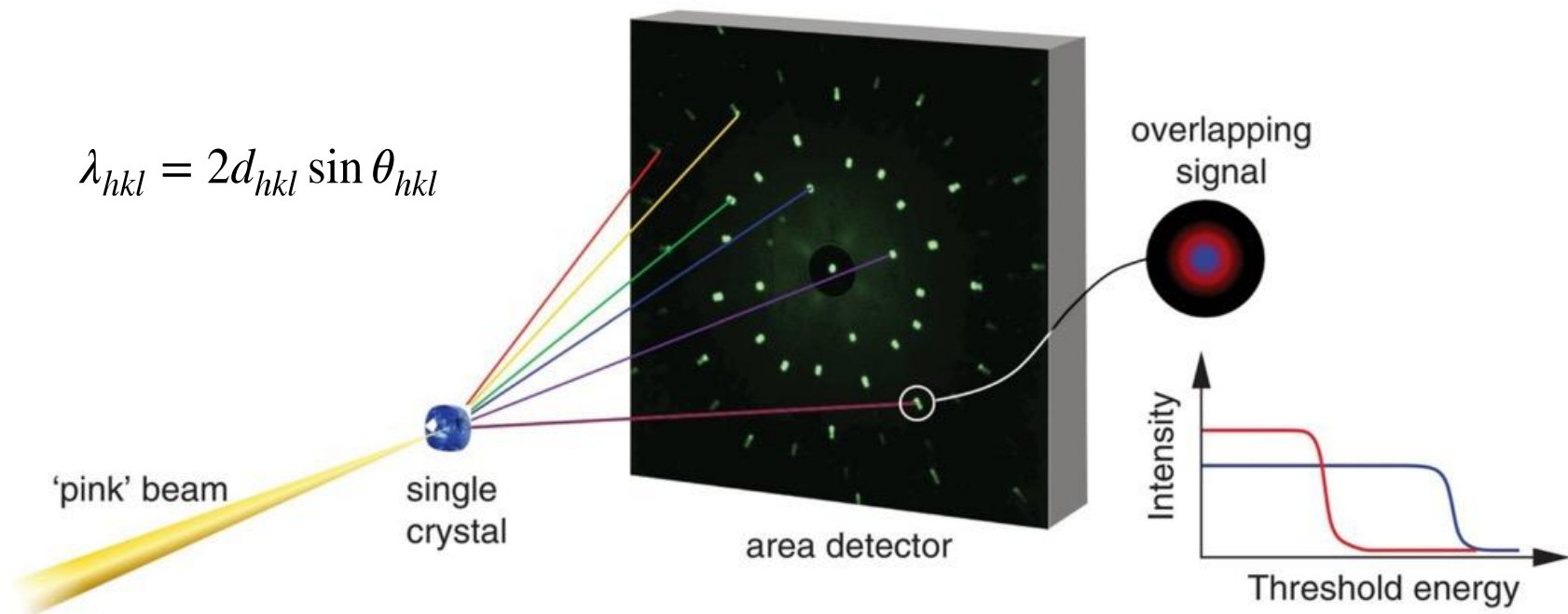
For structure determination from single crystal diffraction one needs to retrieve...

in order to solve this problem, the space group will restrict the model **fewer atoms** because symmetry will automatically place equivalent atoms.

→ Refine the structure more efficiently using fewer parameters.

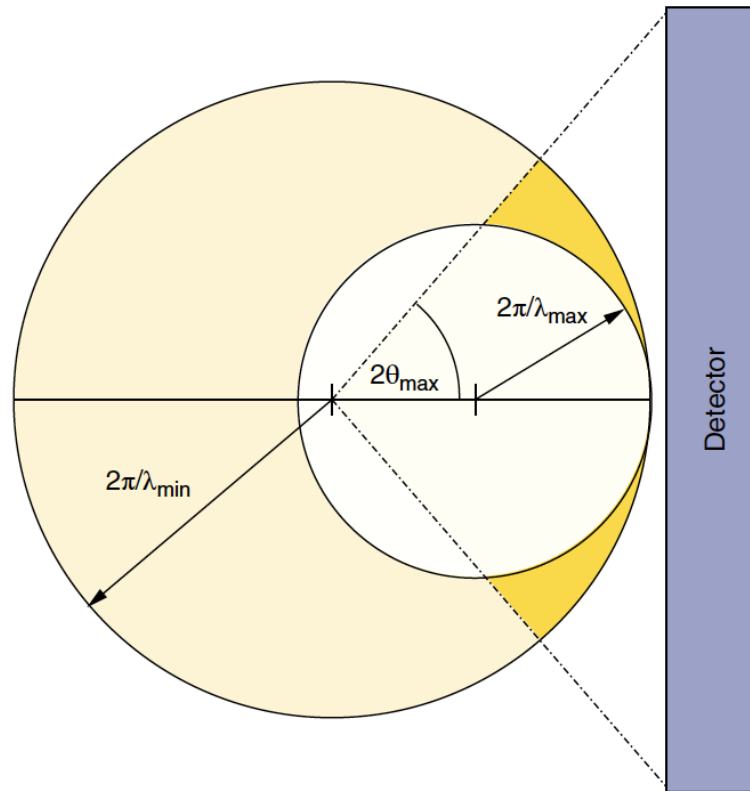
to reach higher resolution in the structure determination one needs....

# Single crystal diffraction – Laue diffraction



- Pink beam: many wavelengths
- Bragg equation fulfilled for multiple combinations of  $d$  and  $\theta$

# Single crystal diffraction – Laue diffraction



**Figure 6.31** The volume of reciprocal space that can be simultaneously accessed in Laue diffraction, shown here in bright yellow, depends on the range of photon energies of the polychromatic beam and the maximum angle that can be subtended by the area detector.



# Sample types

single  
crystal



twinned  
crystal



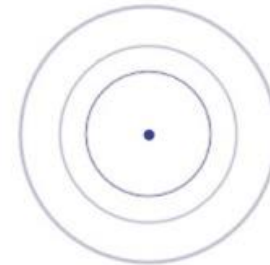
crystal with  
mosaic spread



textured  
sample



powder  
sample



# Material structure?

- Solid materials: Structures at different length scales

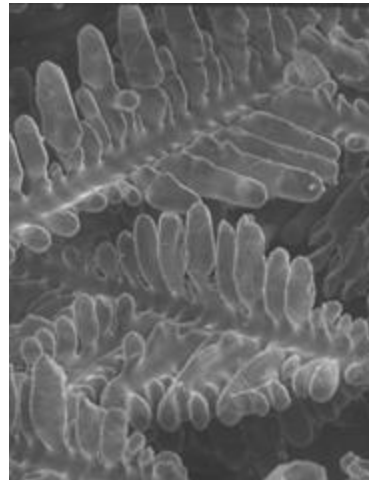
## Example: metal



Part of a turbine - Ni  
(10 cm)



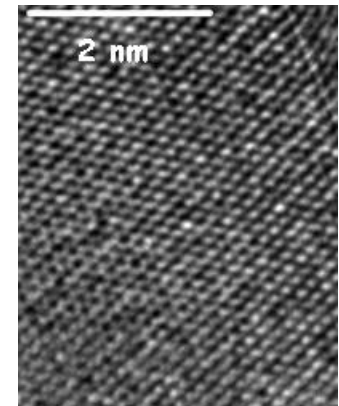
Grains  
(mm)



Dendrites  
(10-100 mm)



Alloys  $\text{Ni}_3\text{Al}$   
(10-100 nm)



Atoms  
(0.1 nm)

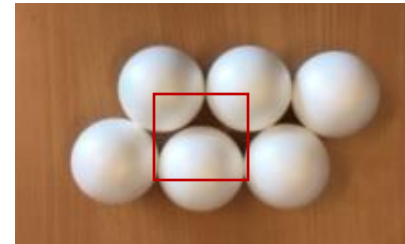
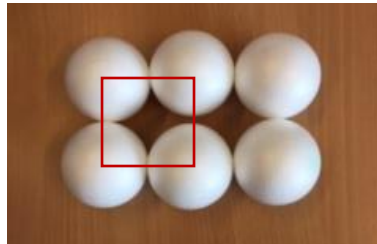
# XRD

## Imperfect microstructure

- large crystal with perfect atomic arrangement give rise to perfectly sharp peak (except of instrumental broadening)
- imperfections such as grain boundaries, defects at dislocations, stacking faults, stresses → peak broadening, as well as possibly peak position shifts

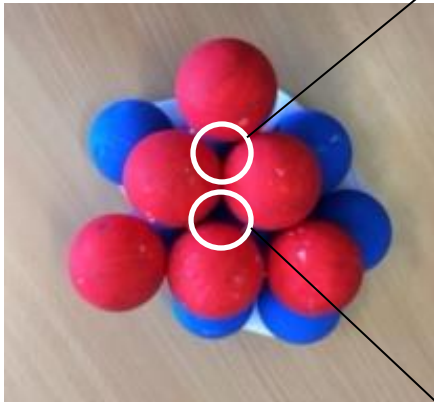
# The hard sphere model

- 2D configuration:

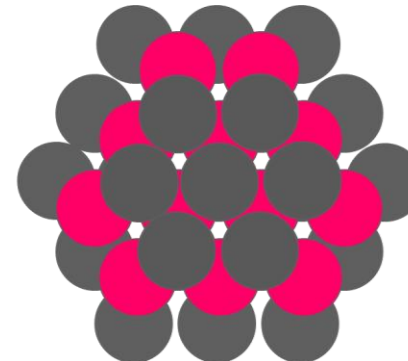


- 3D configuration:

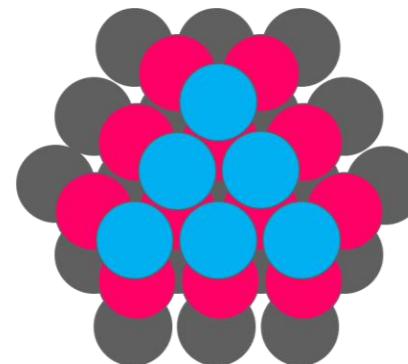
2 possibilities:



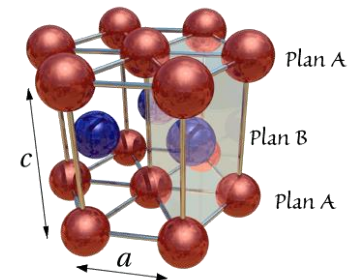
MSE-238



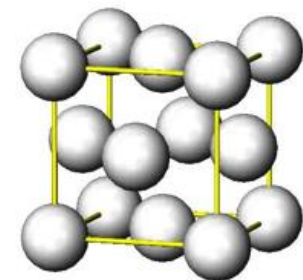
A-B-A



A-B-C

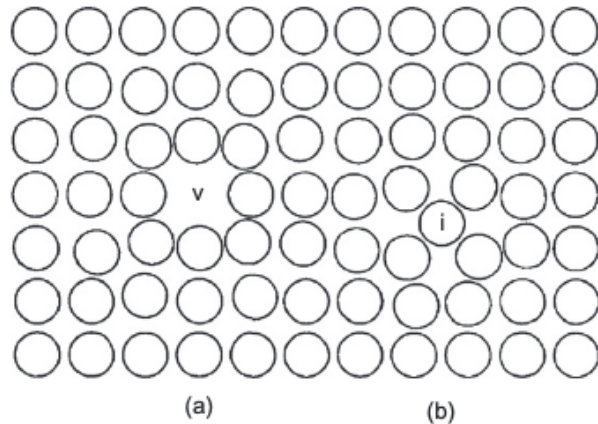


Hexagonal Compact

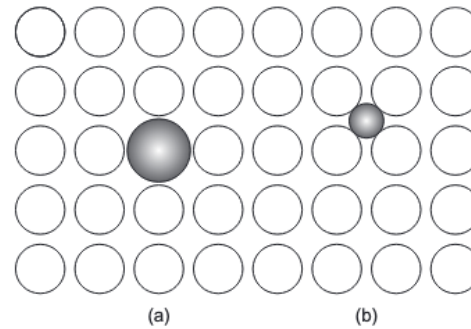


Face-centered Cubic  
(= cubic-close packed)

# Single crystals: defects!



point defects in a simple cubic crystal



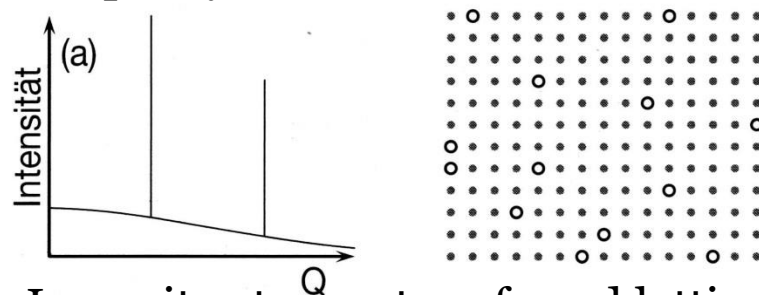
impurities

Hull, D. & Bacon, D. J. Defects in Crystals. *Introd. to Dislocations* (2011)

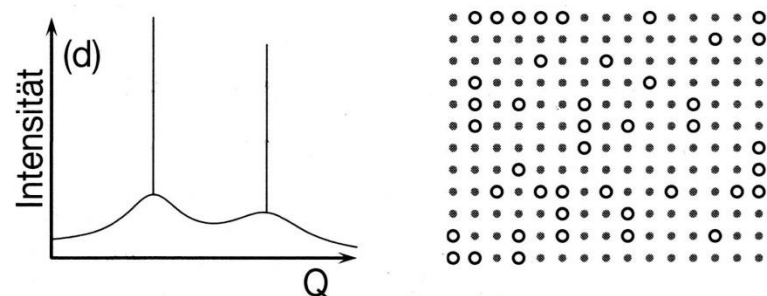
# XRD: Impurity and disorder

Consider crystal with original crystal atoms • and impurity species ○ (or holes)

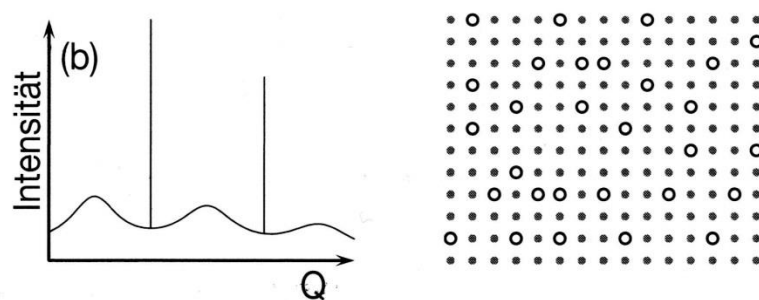
Impurity atoms at random lattice sites



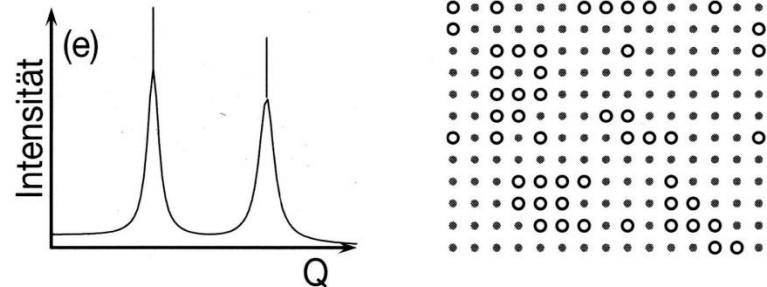
Impurity atoms tending to cluster



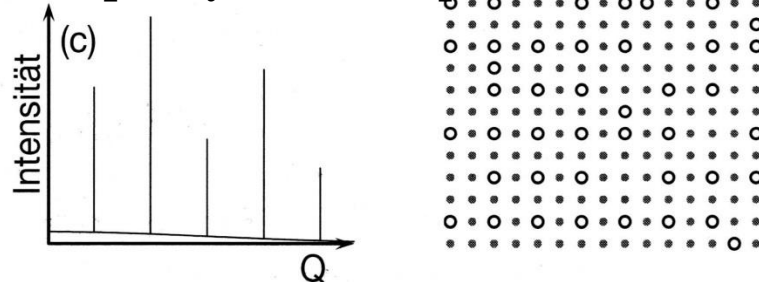
Impurity atoms at preferred lattice sites



Impurity atoms residing in clusters

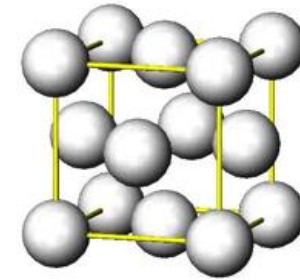
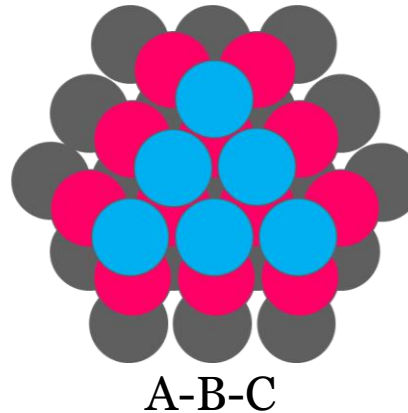


Impurity atoms at preferred lattice sites, long range order

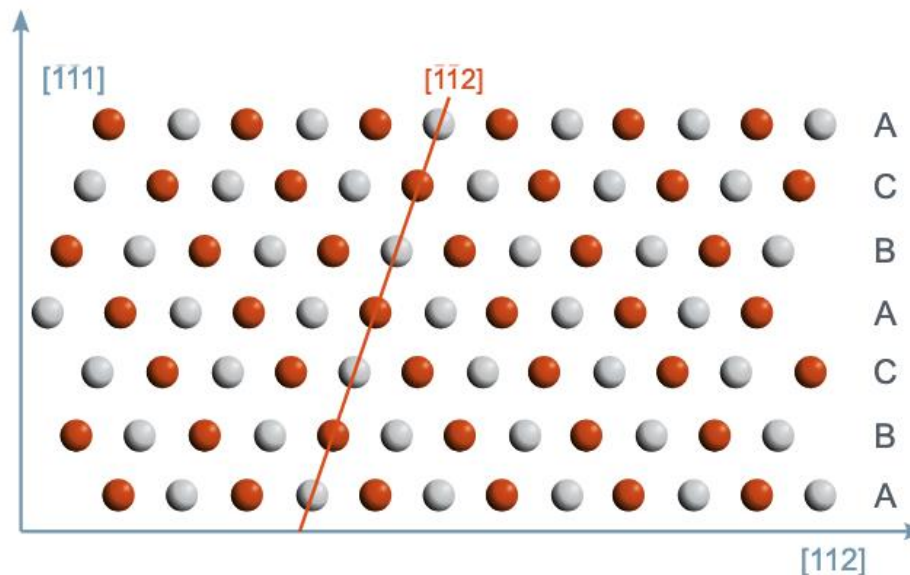




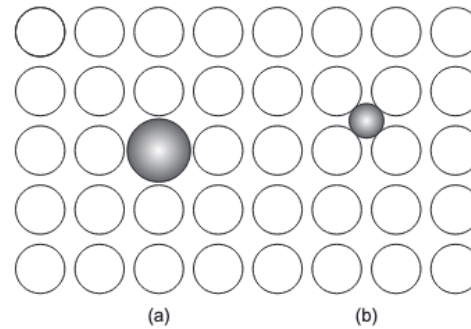
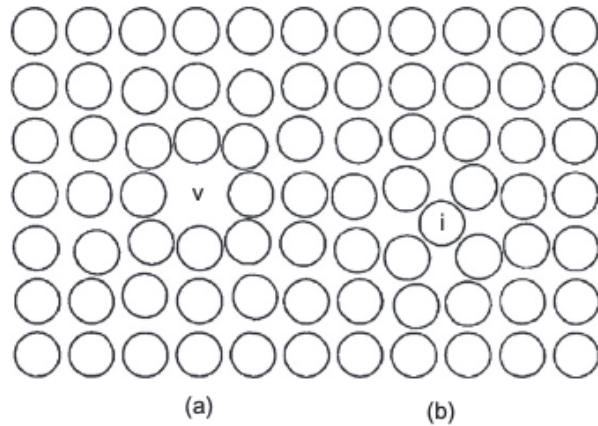
# FCC structure as hard sphere model



Face-centered Cubic  
(= cubic-closed packed)



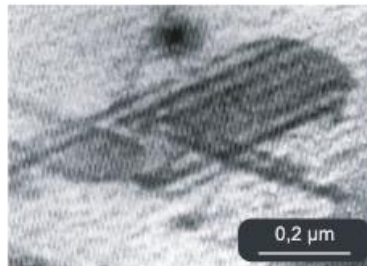
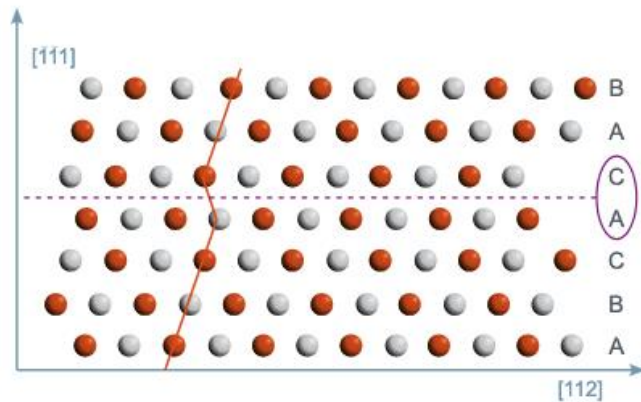
# Single crystals: defects!



Hull, D. & Bacon, D. J. Defects in Crystals. *Introd. to Dislocations* (2011)

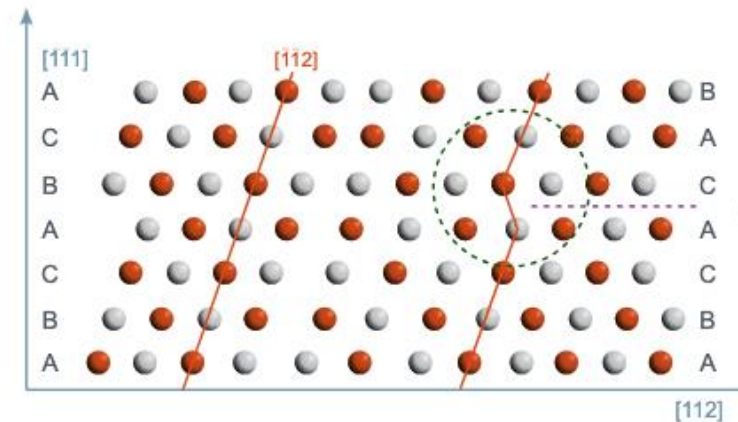
point defects in a simple cubic crystal

impurities



planar defect: stacking fault

<https://nte.mines-albi.fr/>

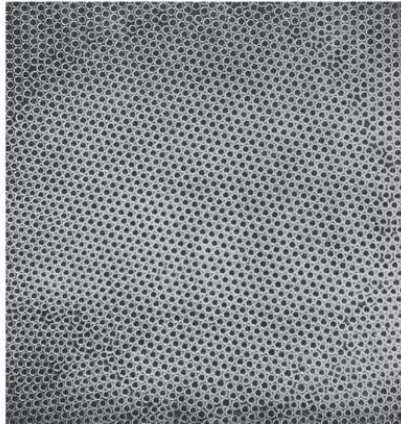


partial dislocation



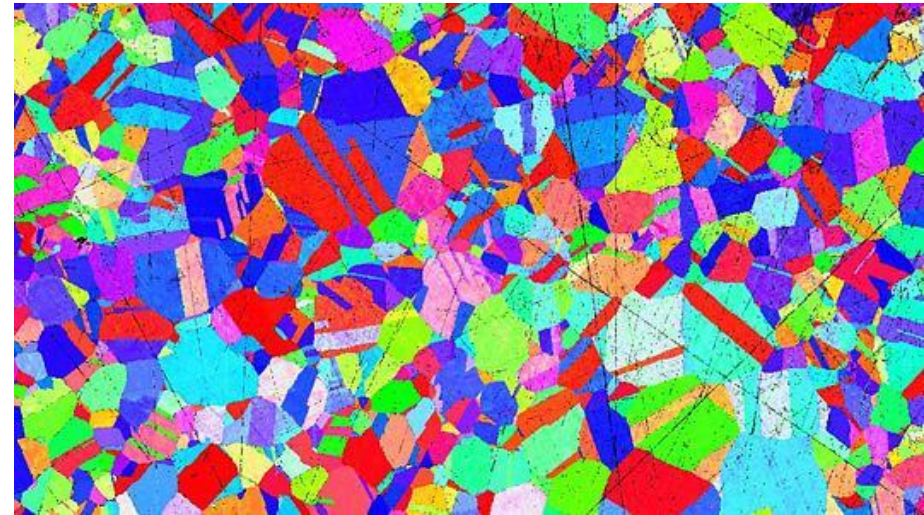
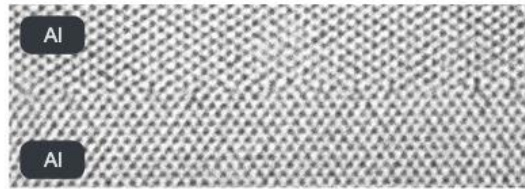
# Polycrystalline material grain boundaries


grain boundaries



grain boundaries are both sources and traps for point defects and dislocations. Grain boundaries also play an important role in plastic deformation as they can induce dislocations under the action of a stress and constitute as well obstacles to the movement of dislocations.

Hull, D. & Bacon, D. J. Defects in Crystals. *Introd. to Dislocations* (2011)



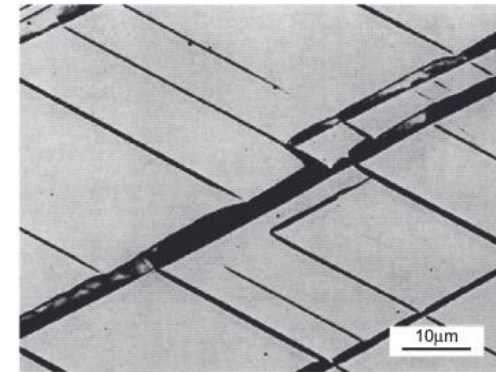
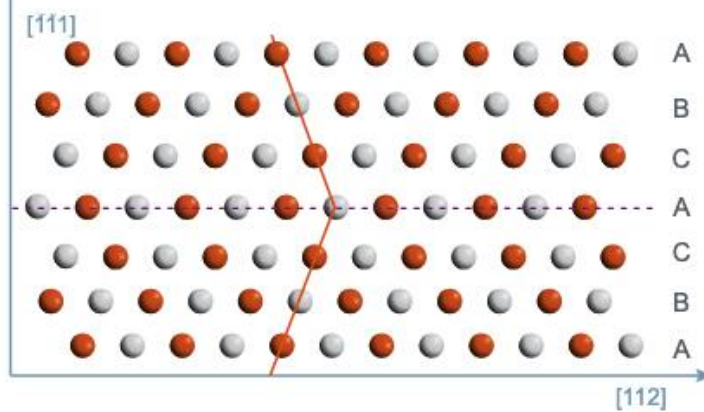
Optical micrographic views of a stainless steel showing grains and grain boundaries constituting (left) and grain boundaries (right) viewed using a TEM (interface between aluminium and -aluminium at top; and aluminium and-germanium at bottom). 

<https://nte.mines-albi.fr/>

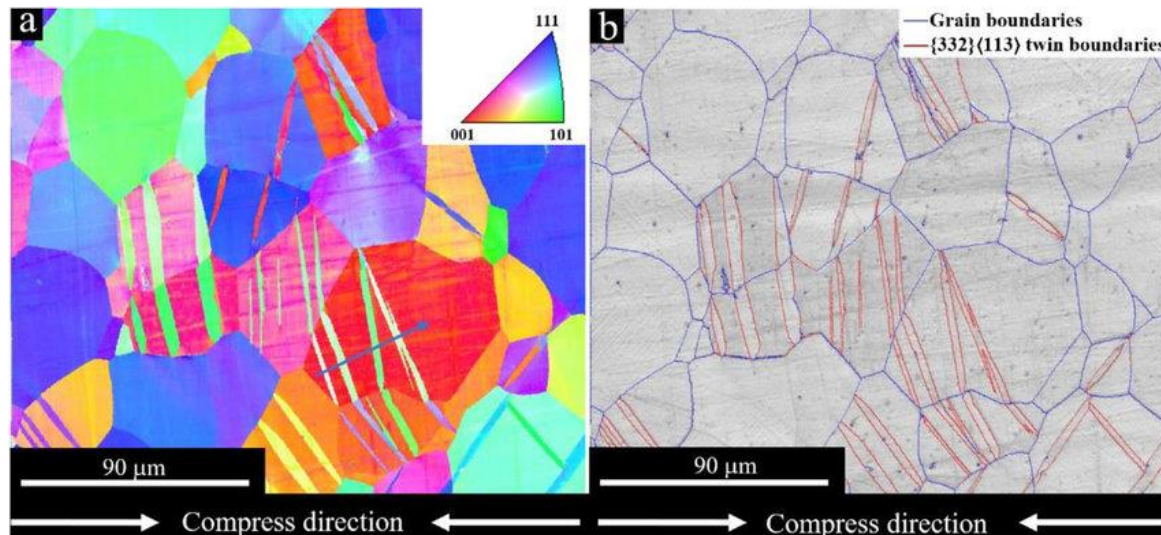
electron back scatter diffraction (EBSD) shows different orientations of grains

# Twins

a twin refers to a crystal that is composed of two or more domains (individual crystal parts) that are **related by a specific symmetry operation**, but not part of the crystal's internal symmetry (space group). These domains **share some lattice points**.



deformation twins in silicon iron



Yang et al. J. of  
Materials Science &  
Technology 73 (2021)  
52-60

# Sample types

single  
crystal



twinned  
crystal



crystal with  
mosaic spread



textured  
sample





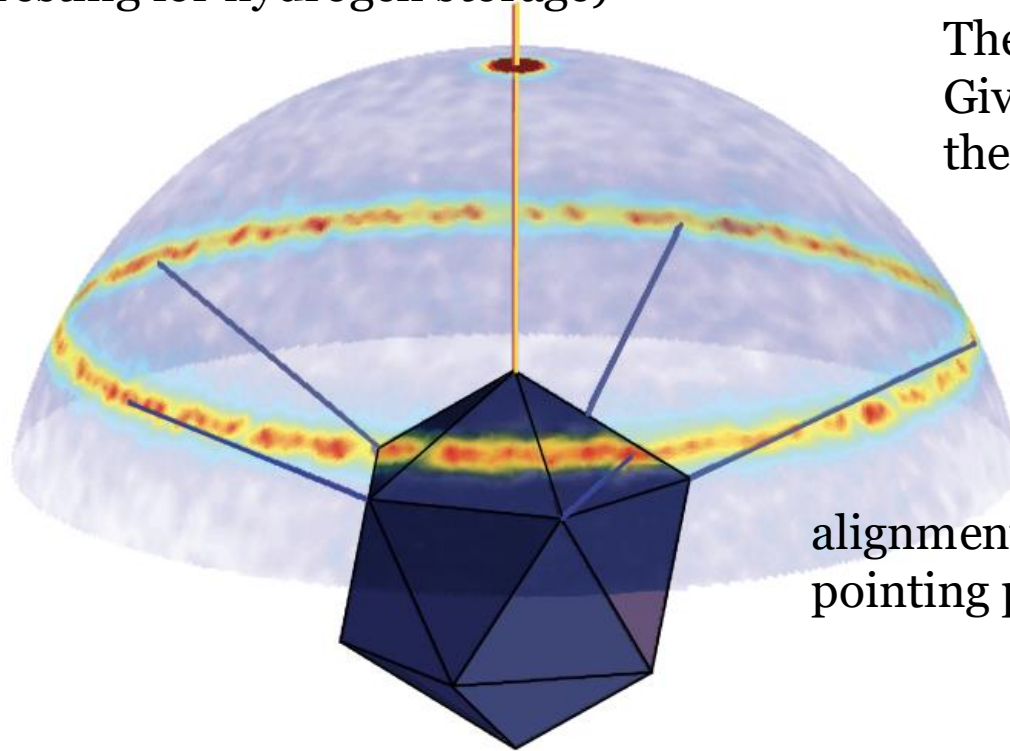
# Textured sample and pole figure

Ti-Ni-Zr alloy thin films  
(interesting for hydrogen storage)

Given a specific set of reciprocal lattice vectors,  $\{hkl\}$

The pole figure  $P_{hkl}(q)$

Gives the probability of finding that plane in the direction,  $q$



alignment of one of the symmetry axis  
pointing perpendicular out of the thin film

illustrated here is the structure of a icosahedron which has a five-fold symmetry

→ is this a crystal system?

# Symmetry operations in 2D

For discrete objects, rotational symmetries can only be discrete:  $\frac{2\pi}{n}$

and they rotational symmetry must be compatible with a translational symmetry!

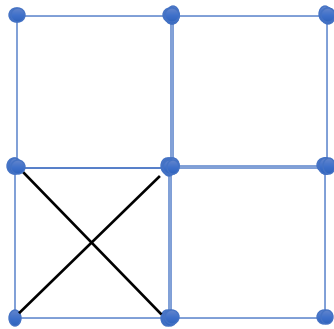
$n=1 \rightarrow$  1-fold, no symmetry

$n=2 \rightarrow$  2-fold,  $180^\circ$  rotation

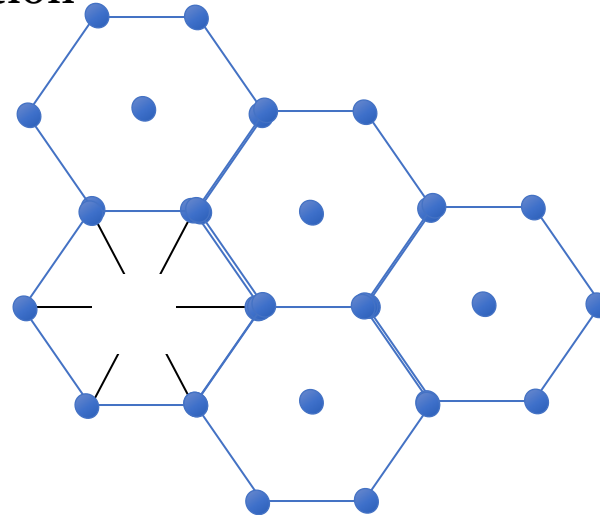
$n=3 \rightarrow$  3 fold,  $120^\circ$  rotation

$n=4 \rightarrow$  4 fold,  $90^\circ$  rotation

$n=6 \rightarrow$  6 fold,  $60^\circ$  rotation

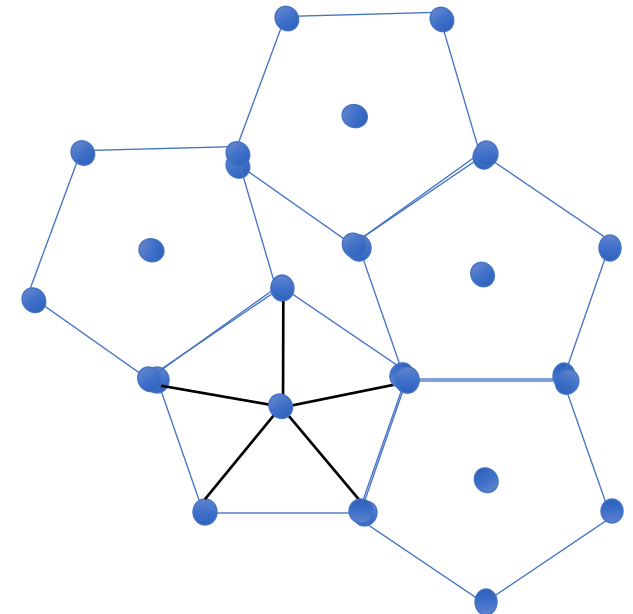


4-fold



6-fold

what about 5-fold?

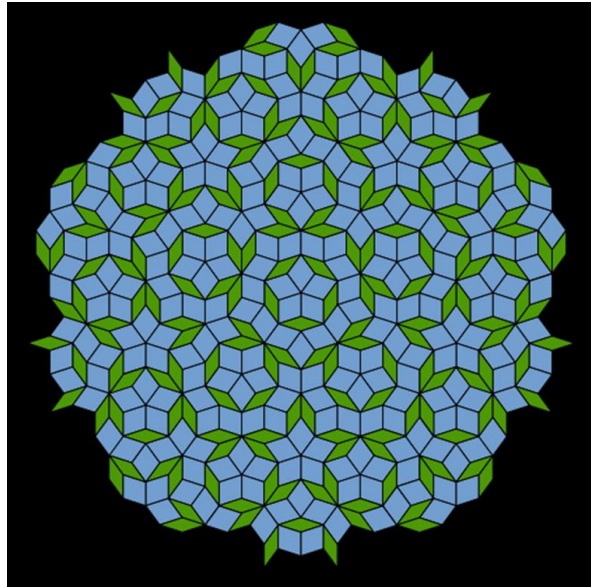


and mirror symmetry (m)

# Patterns with 5-fold symmetry $\rightarrow$ Quasi-crystals

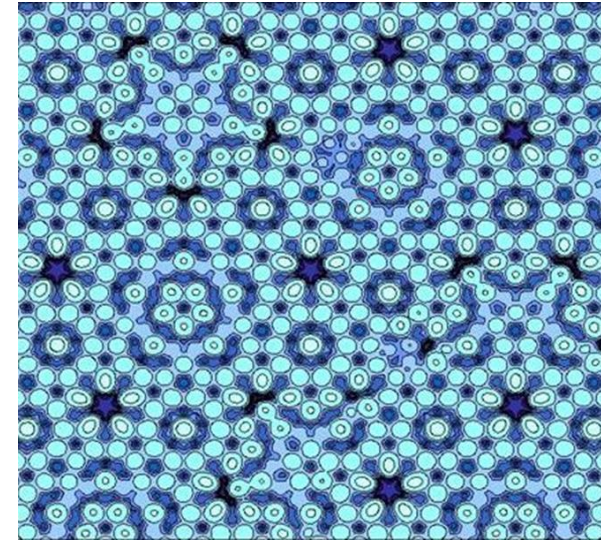
A quasiperiodic crystal (quasicrystal) is a structure that is ordered but not periodic.

A quasi-crystalline pattern can continuously fill all available space, but it lacks translational symmetry



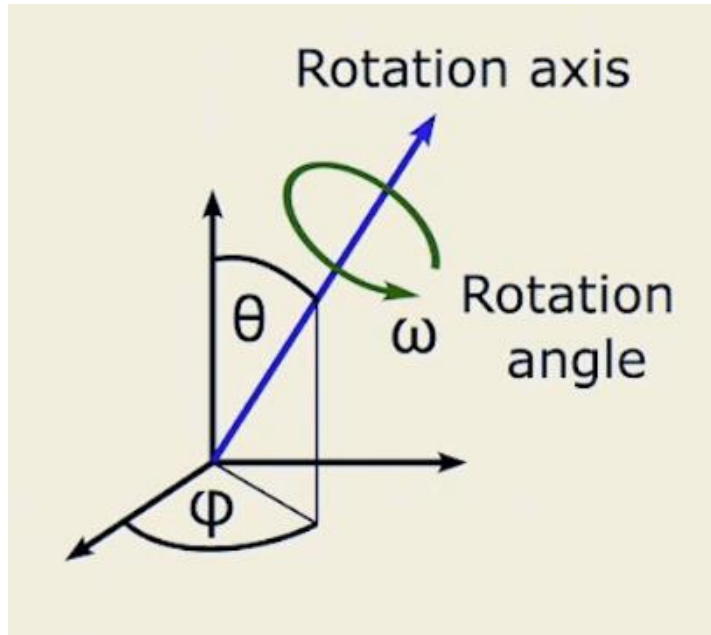
Penrose tiling gives a quasicrystal

[http://en.wikipedia.org/wiki/Penrose\\_tiling](http://en.wikipedia.org/wiki/Penrose_tiling)

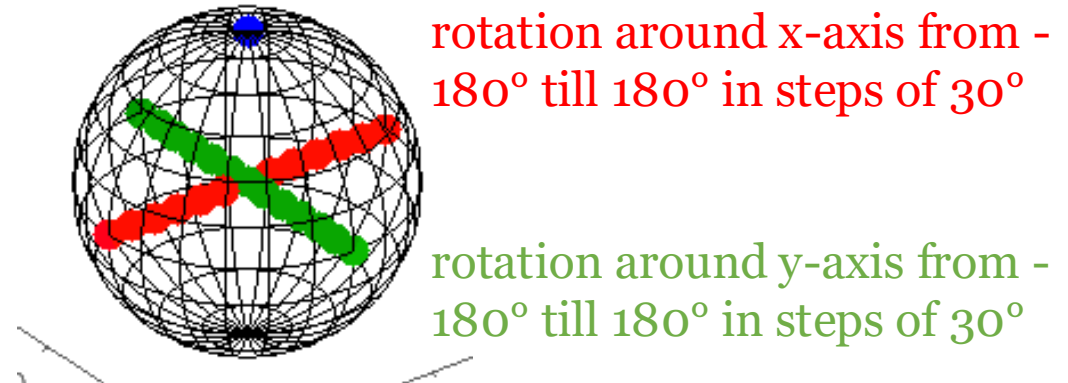
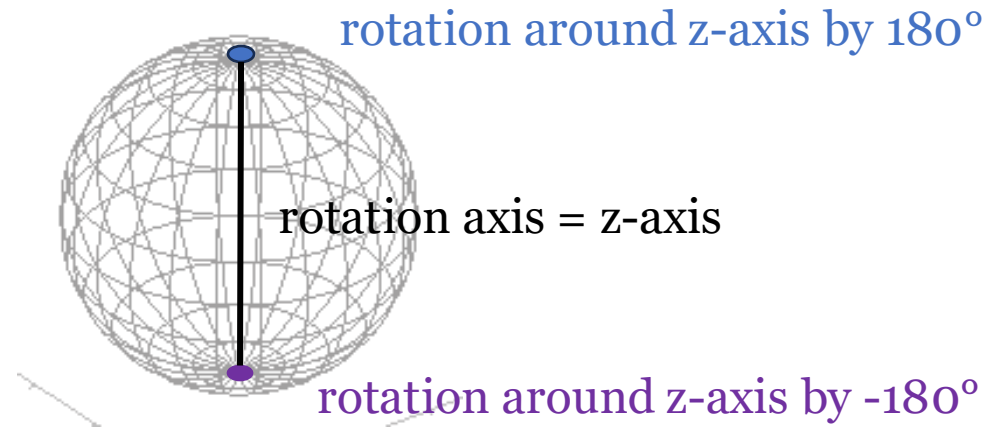


Atomic model of an aluminium-palladium-manganese (Al-Pd-Mn) quasicrystal surface.

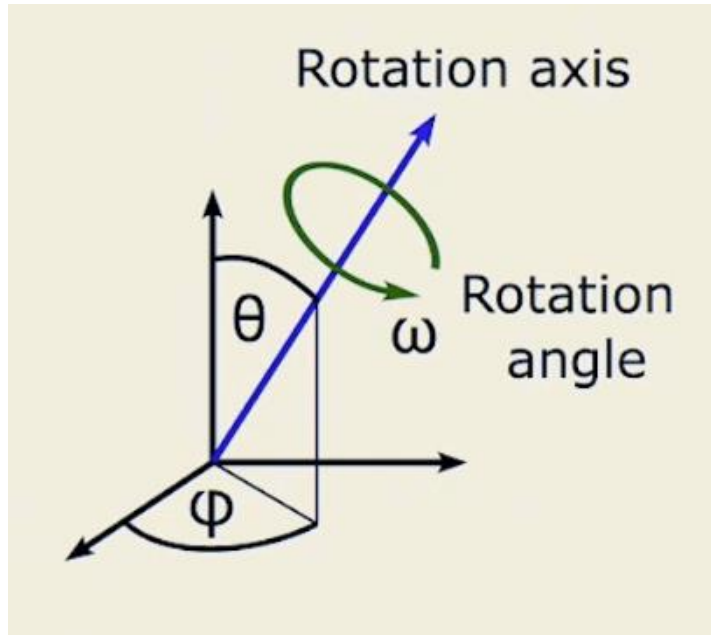
# Textured samples: orientation information



axis-angle representation of 3D orientation  
one vector defining the axis, and an angle defining the rotation around it

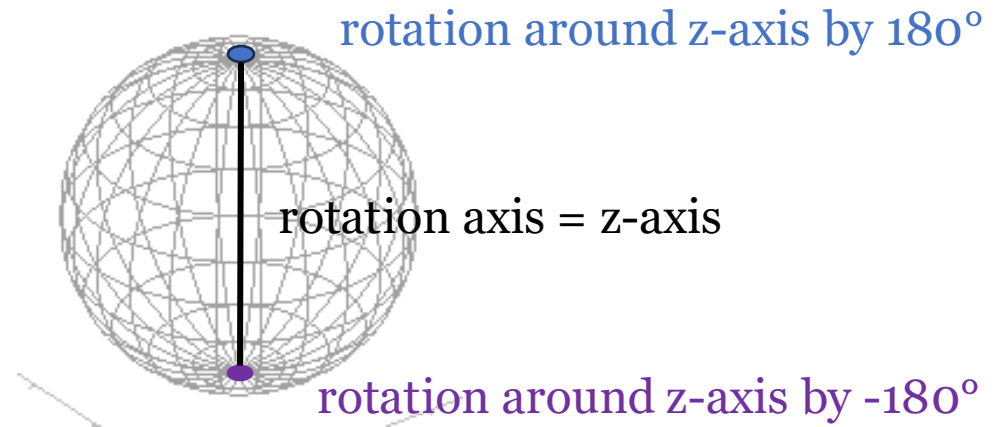


# Textured samples: orientation information



axis-angle representation of 3D orientation

one vector defining the axis, and an angle defining the rotation around it



if there is a two-fold **symmetry** around that axis, rotation of  $0^\circ$  and  $180^\circ$  and  $-180^\circ$  are equivalent!

what defines if certain orientations are equivalent?



# crystal system

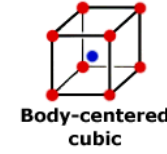
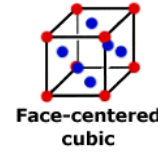
## Bravais lattices

## defining symmetry

Cubic

$$a = b = c$$

$$\alpha = \beta = \gamma = 90^\circ$$

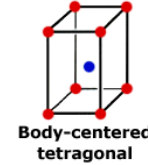
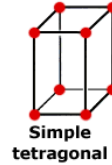


4x 3-fold axis  
3x 4-fold axis

Tetragonal

$$a = b \neq c$$

$$\alpha = \beta = \gamma = 90^\circ$$

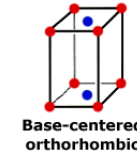
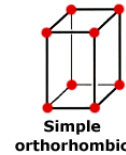


4-fold axis

Orthorhombic

$$a \neq b \neq c$$

$$\alpha = \beta = \gamma = 90^\circ$$

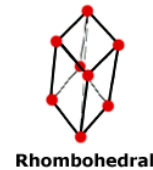


3x 2-fold axis

Trigonal or rhombohedral

$$a = b = c$$

$$\alpha = \beta = \gamma \neq 90^\circ$$

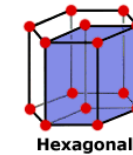


3-fold axis

Hexagonal

$$a = b \neq c$$

$$\alpha = \beta = 90^\circ; \gamma = 120^\circ$$

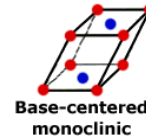


6-fold axis

Monoclinic

$$a \neq b \neq c$$

$$\alpha = \gamma = 90^\circ \neq \beta$$



2-fold axis

Triclinic


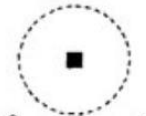

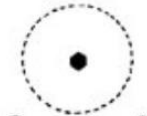


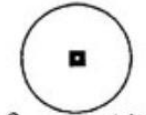


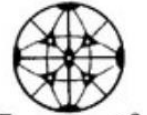
















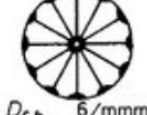




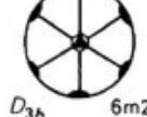
$$a \neq b \neq c$$

$$\alpha \neq \beta \neq \gamma$$



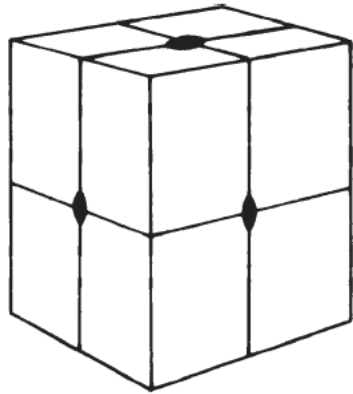
1-fold axis

# 32 Point groups in 3D

Triclinic and Monoclinic	Orthorhombic	Tetragonal	Hexagonal		Cubic (Isometric)
 $C_1$ 1		 $C_4$ 4	 $C_3$ 3	 $C_6$ 6	 $T$ 23
 $C_i$ 1		 $C_{4h}$ 4/m	 $S_6$ 3	 $C_{6h}$ 6/m	 $T_h$ m3
 $C_2$ 2	 $D_2$ 222	 $D_4$ 422	 $D_3$ 32	 $D_6$ 622	 $O$ 432
 $C_s$ m	 $C_{2v}$ 2mm	 $C_{4v}$ 4mm	 $C_{3v}$ 3m	 $C_{6v}$ 6mm	 $T_d$ $\bar{4}3m$
 $C_{2h}$ 2/m	 $D_{2h}$ mmm	 $D_{4h}$ 4/mmm	 $D_{3d}$ $\bar{3}m$	 $D_{6h}$ 6/mmm	 $O_h$ $m\bar{3}m$
		 $S_4$ $\bar{4}$		 $C_{3h}$ 6	
		 $D_{2d}$ 42m		 $D_{3h}$ 6m2	

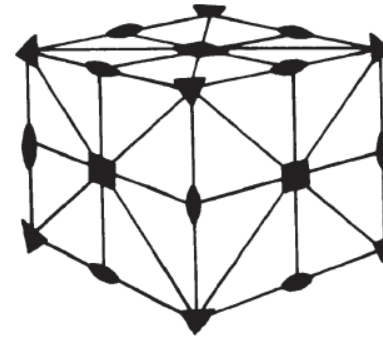
# Point symmetry elements examples

orthorhombic



- 3 times 2-fold axis,  
perpendicular to the faces
- three mirror planes  
parallel to faces planes

cubic

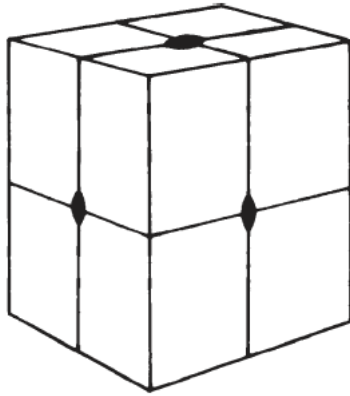


→ highest symmetry,  
makes it hard to see!

- 3 times 4-fold axis  
perpendicular to the faces
- 4 times 3-fold axis between  
opposite cube corners
- 6 times 2-fold axis between  
opposite center of edges
- 9 mirror planes
  - 3 parallel to faces planes
  - 6 parallel to the face diagonals
- plus center of inversion and  
rotoinversions!

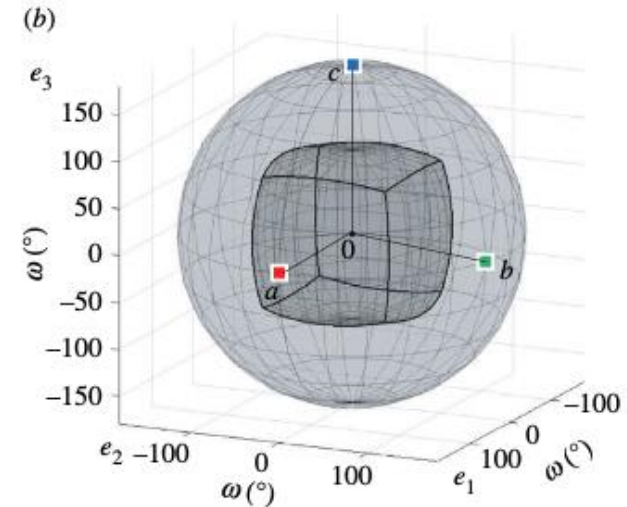
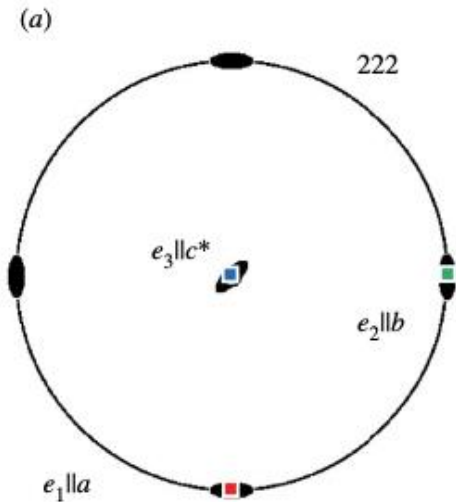
# Orientation of a crystal and crystal symmetry

orthorhombic



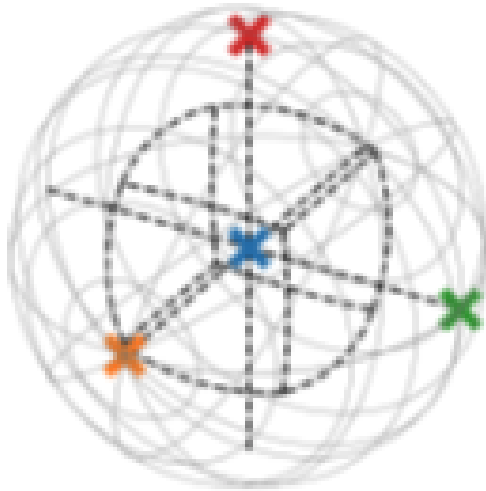
→ 3 times 2-fold axis,  
perpendicular to the faces  
222

→ three mirror planes  
parallel to faces planes



for plotting all possible orientation, only a subset of the  
sphere is needed → called the fundamental zone

# Orientation of a crystal and crystal symmetry

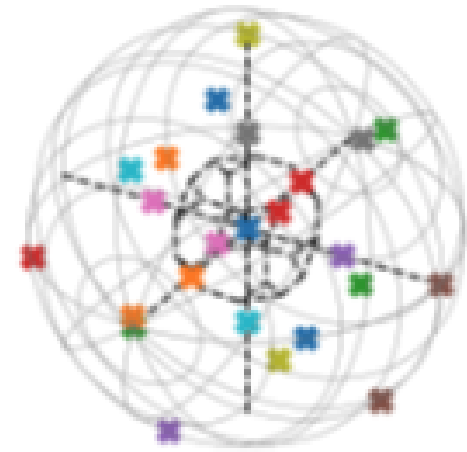


point group 222

crystal system



622



432

the more symmetry elements a point group has, the smaller is the fundamental zone since more and more orientation become equivalent

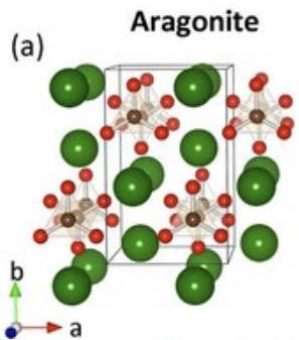
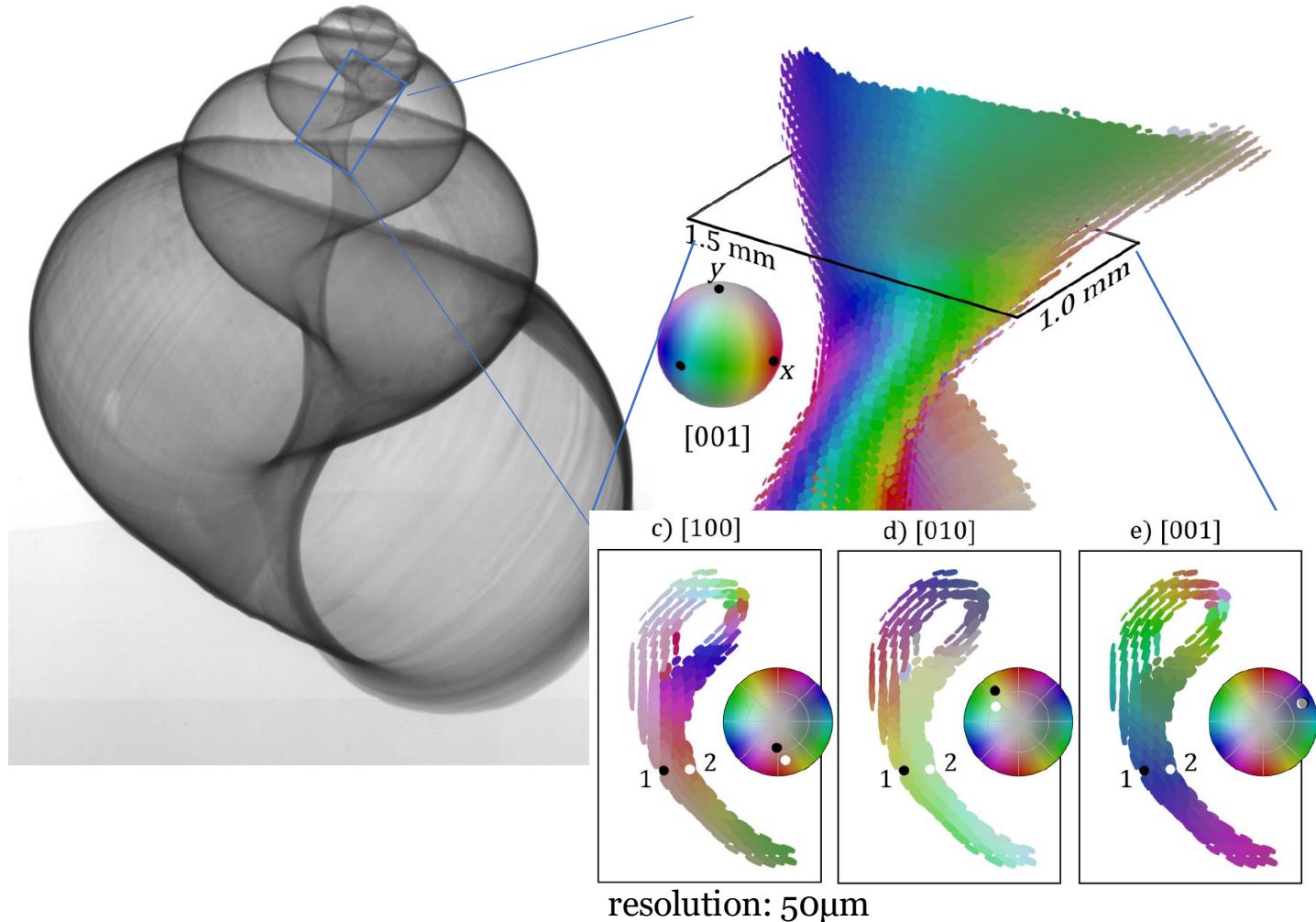
→ this also means for a full dataset in single crystal diffraction fewer orientations of a crystal need to be measured if there is a higher symmetry

# Outer shape and symmetry: Biomineral

Roman snail shell



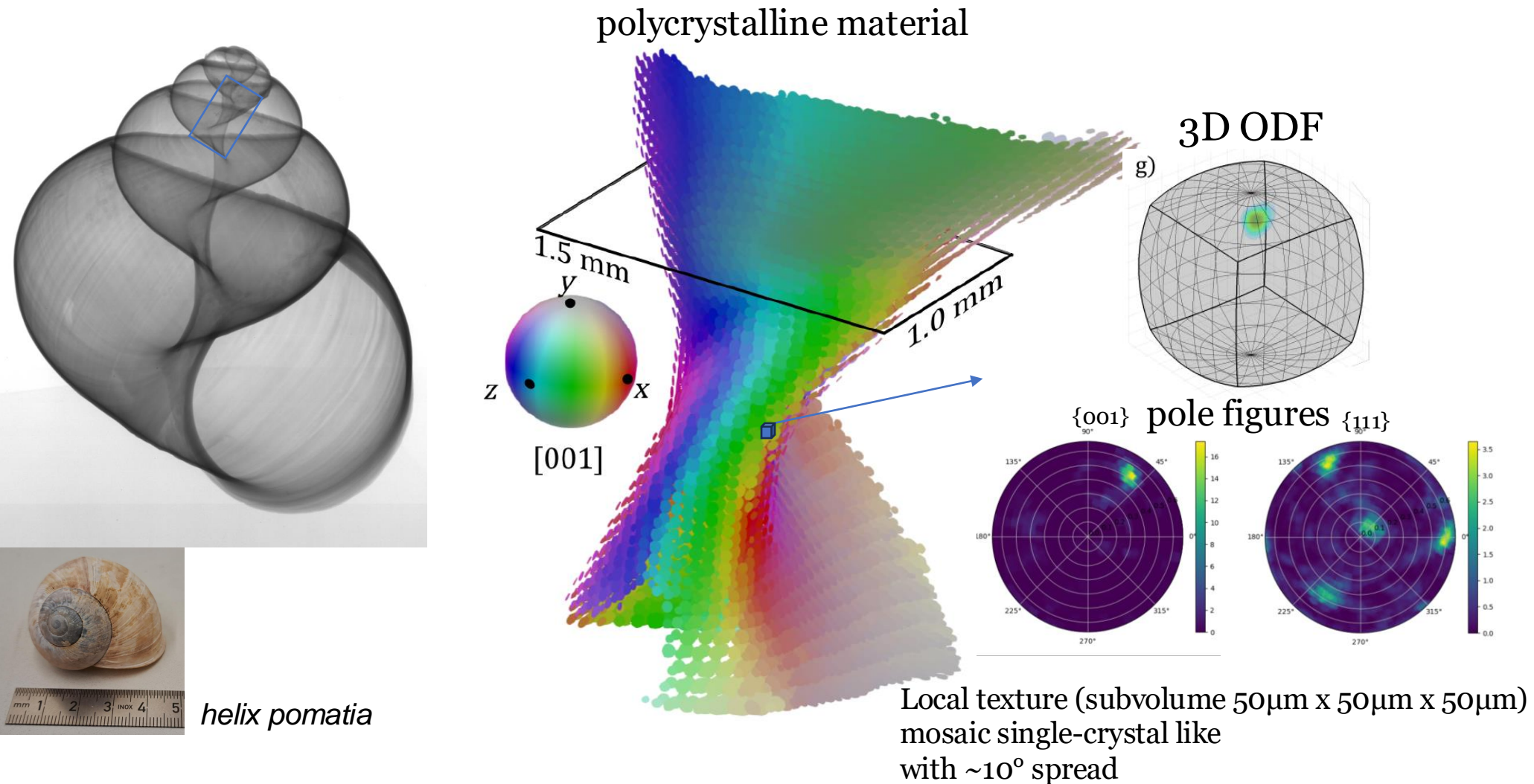
polycrystalline material



aragonite orthorhombic



# Roman snail shell: local texture as mosaic single crystal



M. Carlsen *et al.* Applied Crystallography 2025

# Sample types

single  
crystal



twinned  
crystal



crystal with  
mosaic spread



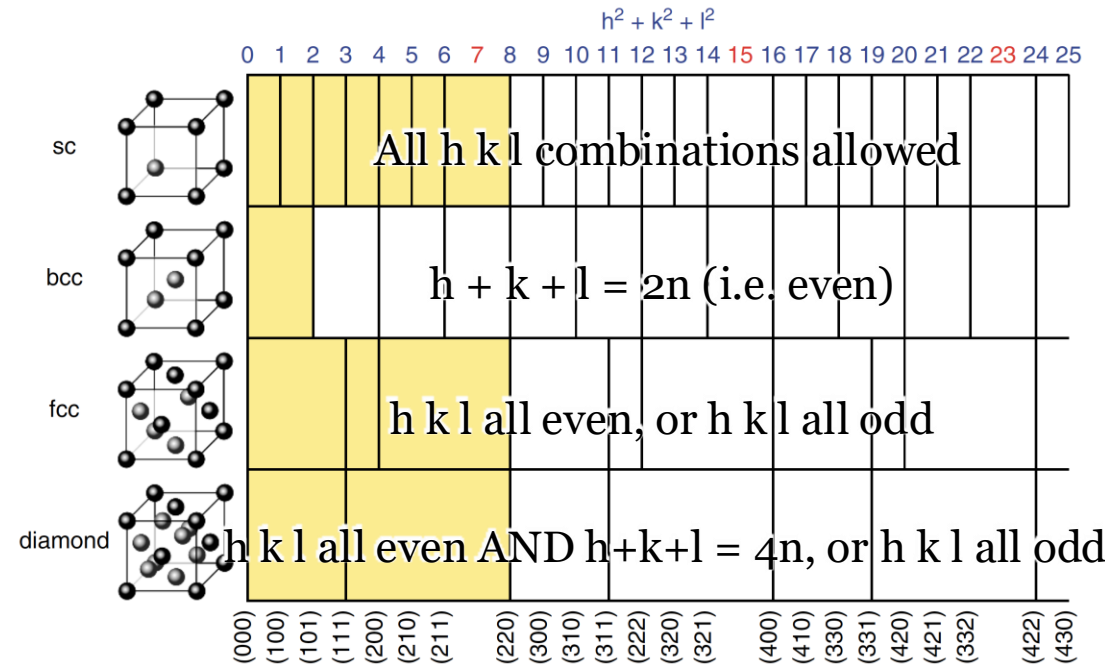
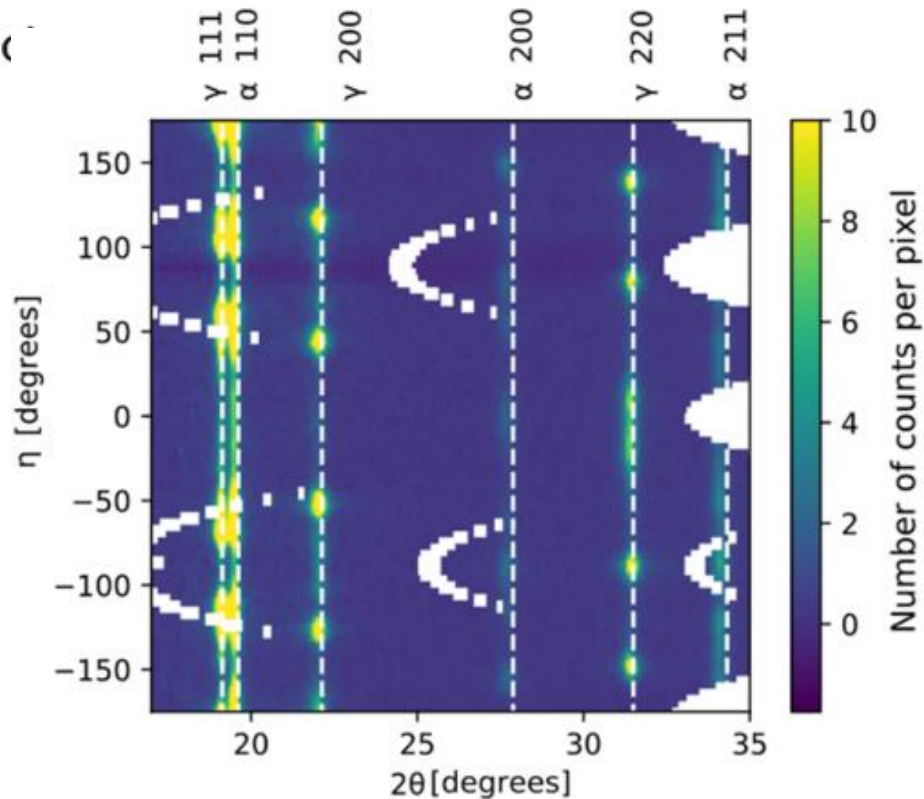
textured  
sample





# Fibre texture: drawn metal wire

drawn metal wire made of stainless steel

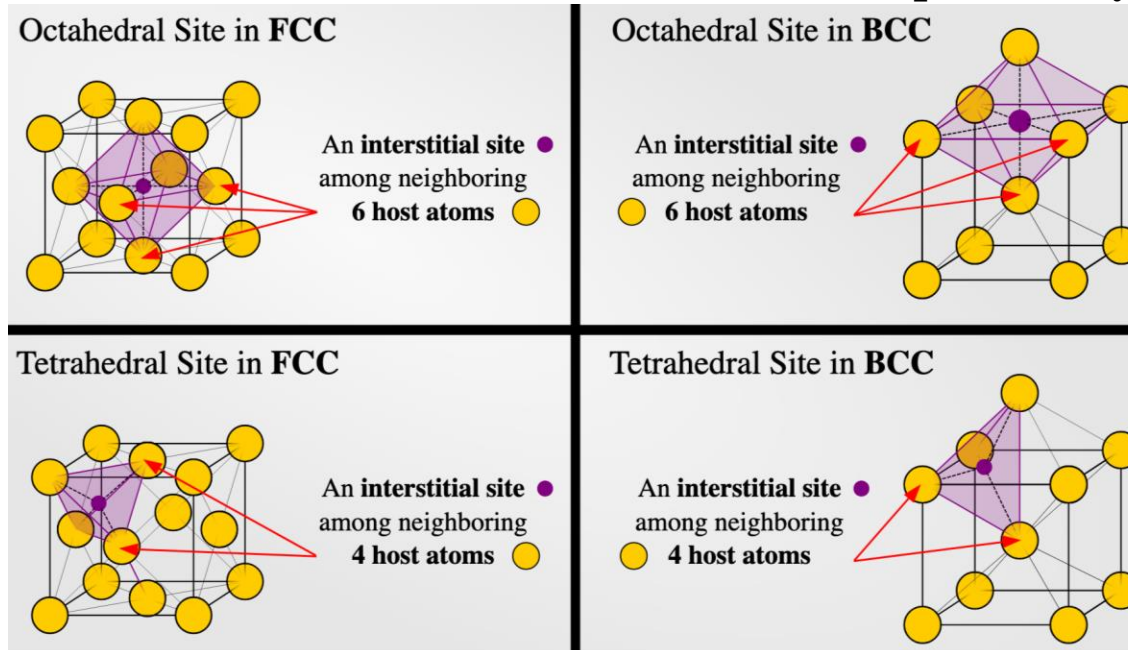


what is the crystal system of the  $\alpha$  phase?  
what is the crystal system of the  $\gamma$  phase?

# Fibre texture: drawn metal wire

drawn metal wire made of stainless steel

there is 0.12% carbon inside that material, in which phase do you expect it to be?

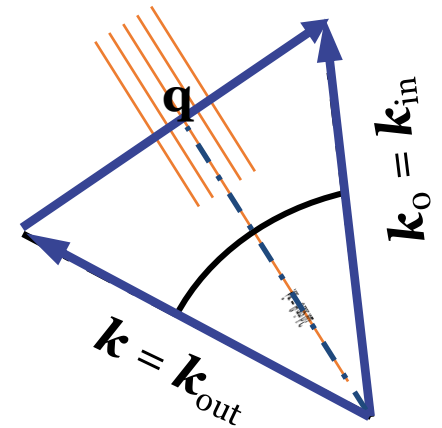
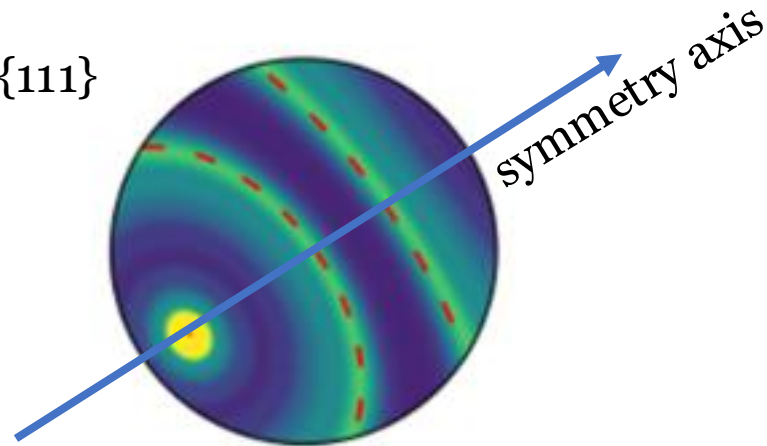


Crystal Structure	FCC	BCC
Number and Size of Octahedral Voids	4 voids, $r = 0.414 R$	6 voids, $r = 0.155 R$
Number and Size of Tetrahedral Voids	8 voids, $r = 0.225 R$	12 voids, $r = 0.291 R$

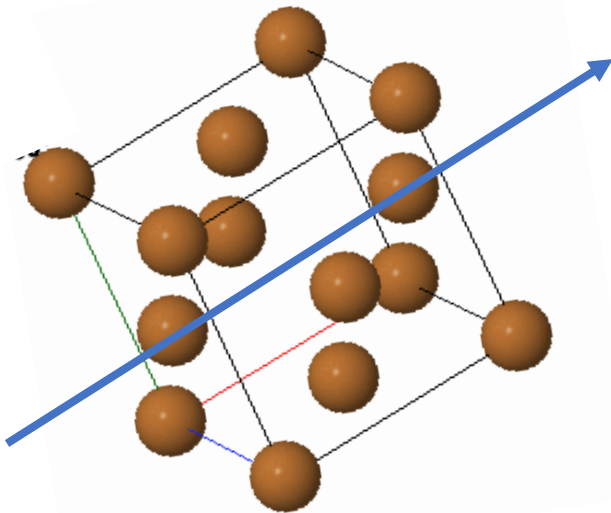
**Carbon: 0.12% maximum**  
 Manganese: 2.00% maximum  
 Phosphorus: 0.045% maximum  
 Sulfur: 0.030% maximum  
 Silicon: 1.00% maximum  
 Chromium: 17.0–19.0%  
 Nickel: 8.0–10.0%  
 Nitrogen: 0.10% maximum  
 Iron: Balance

# Fibre texture: drawn metal wire

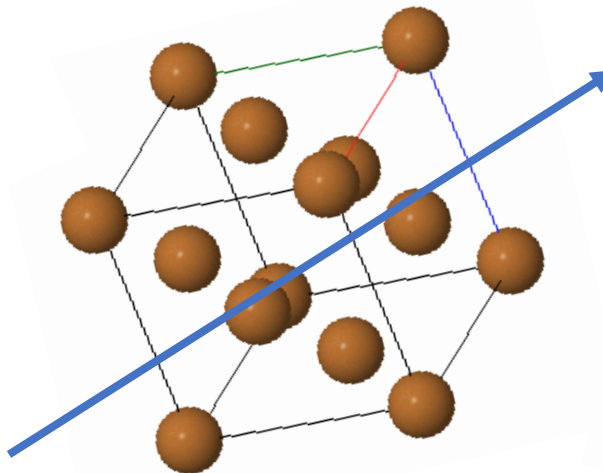
pole figure for  $\{111\}$



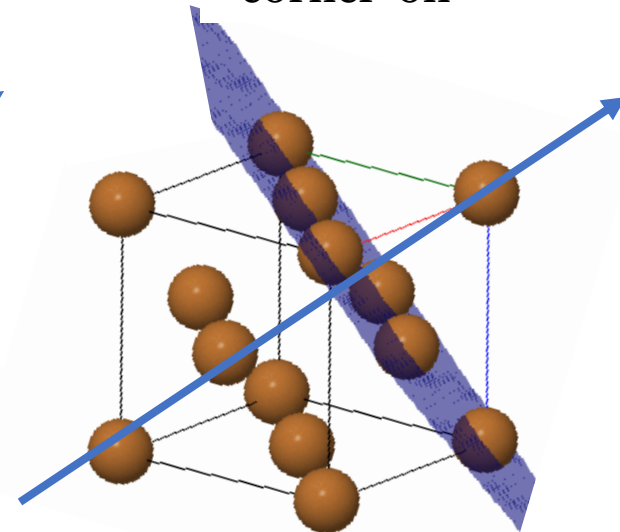
"face-on"



"edge-on"

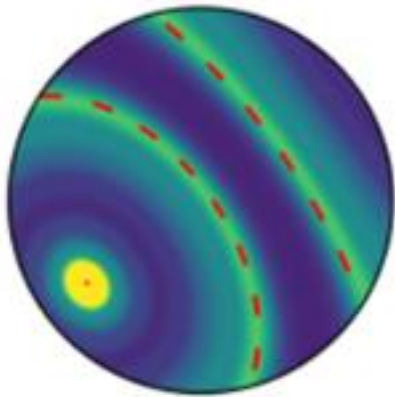


"corner-on"

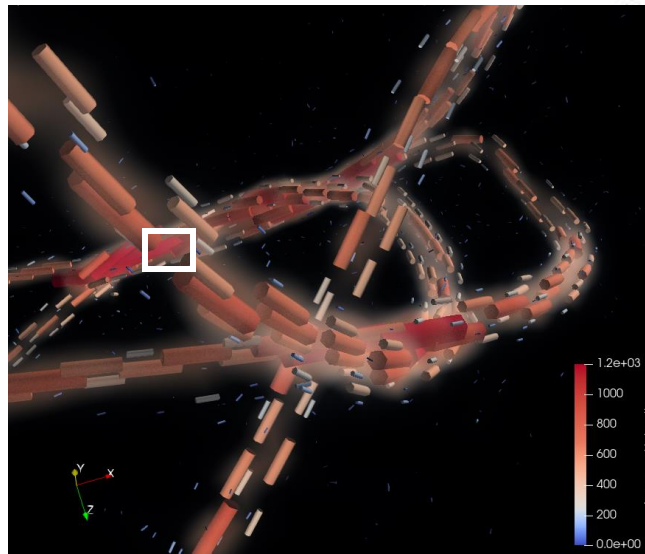


# Texture analysis

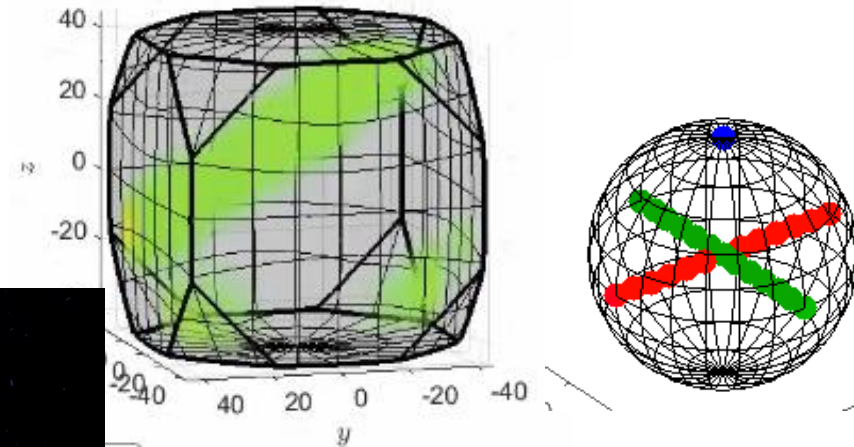
- The pole figure
  - Given a specific set of reciprocal lattice vectors,  $\{hkl\}$
  - The pole figure gives the probability of finding that plane in the direction,  $q$



fiber texture: peaks smear into a ring



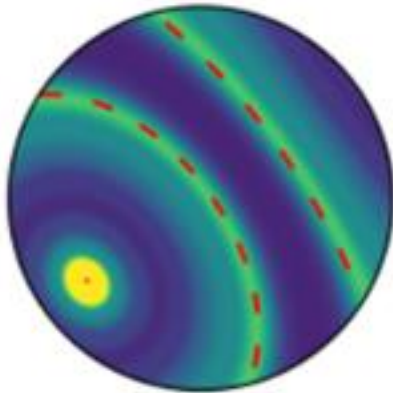
3D orientation distribution function (ODF)  
how is the crystal lattice oriented with respect to the actual sample



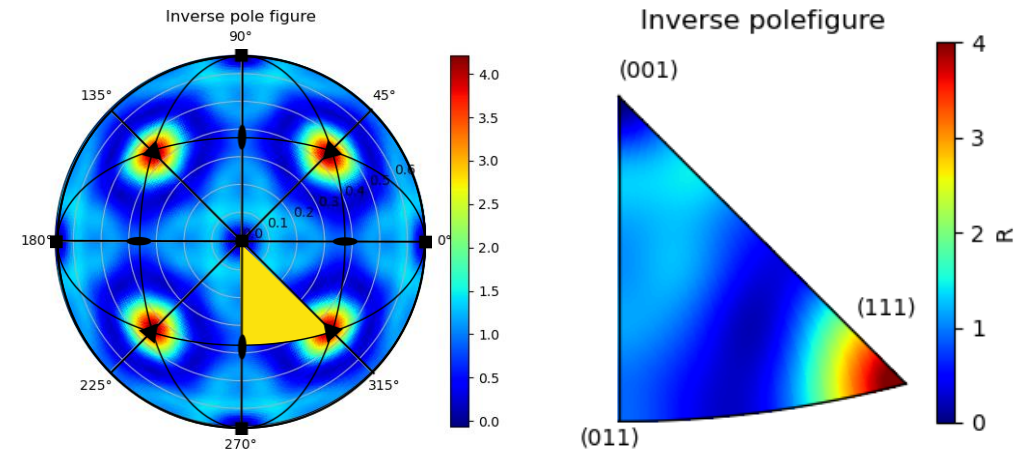
fiber texture: fixed symmetry axis, but free rotation around  
→ a line

# Texture analysis

- The pole figure
  - Given a specific set of reciprocal lattice vectors,  $\{hkl\}$
  - The pole figure gives the probability of finding that plane in the direction,  $q$



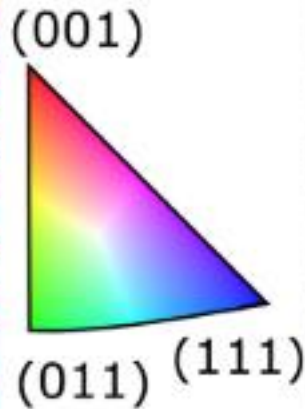
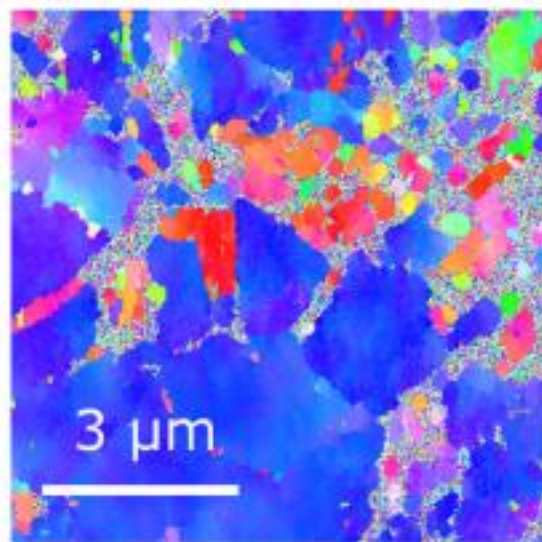
- The inverse pole figure
  - Given a specific direction  $y$  in the sample (here: the wire direction = draw direction)
  - The inverse pole figure gives the probability that  $y$  falls in a certain lattice orientation.



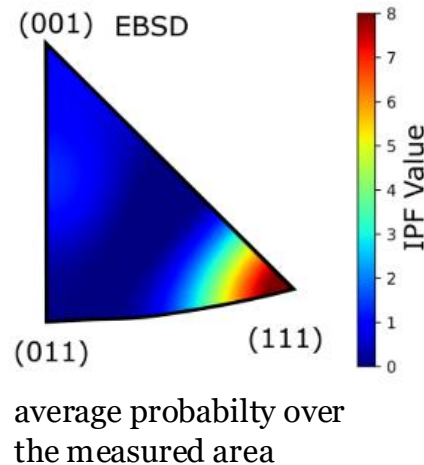


# Inverse pole figures

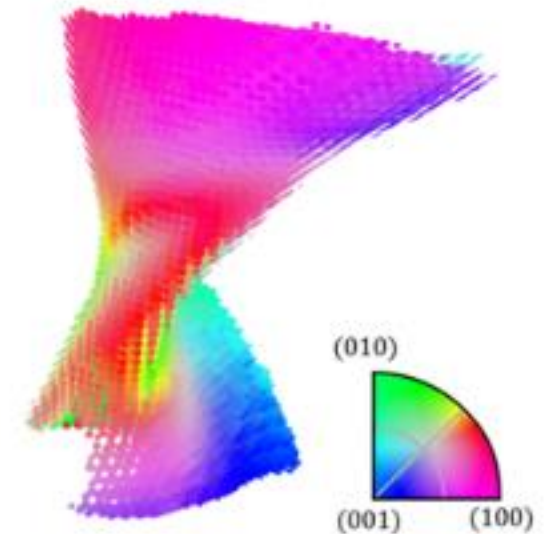
electron back-scattered diffraction (EBSD)  
on a cut with the surface normal to the wire direction  
inverse pole figure map



fcc



inverse pole figure map of the snail shell

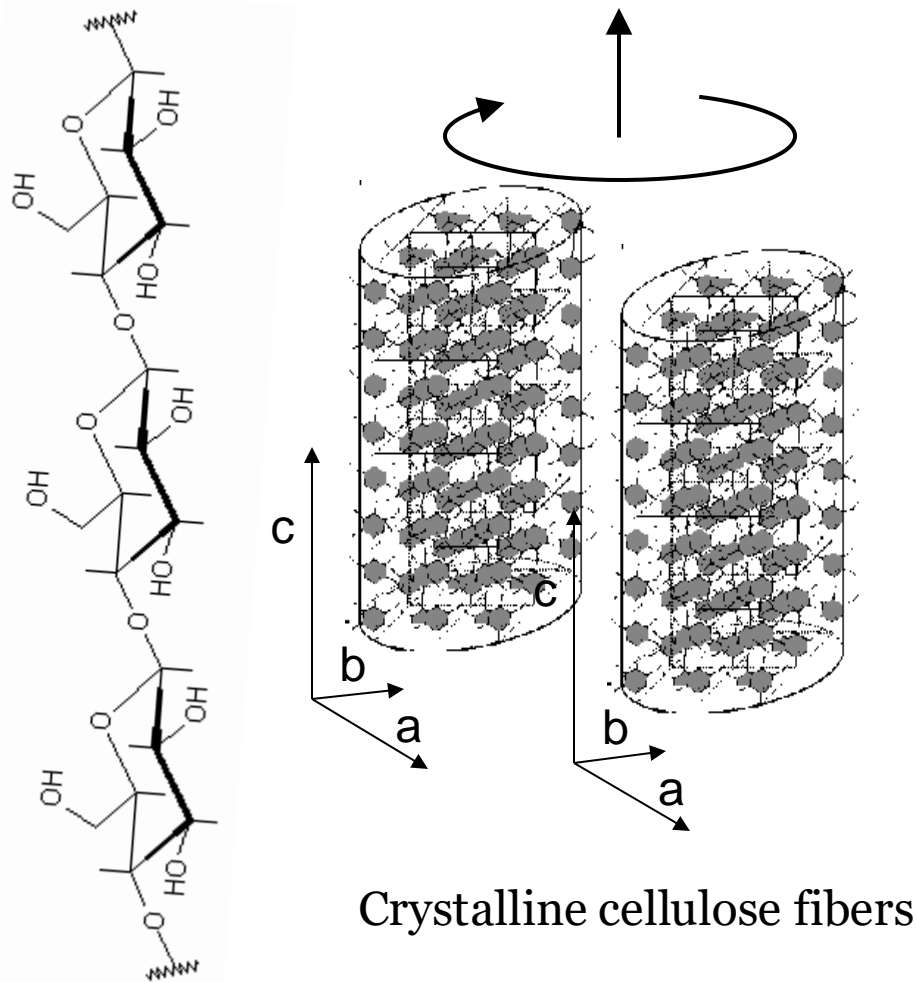


orthorhombic

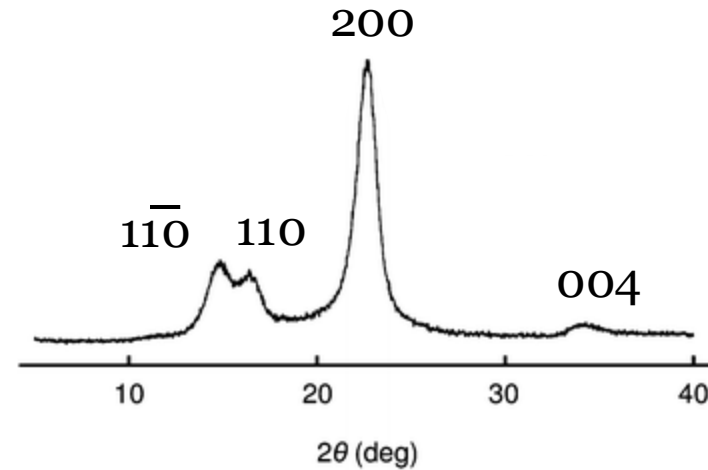
note that the portion from the circle shown depends on crystal symmetry!

# Fibre texture are very common

cellulose fiber

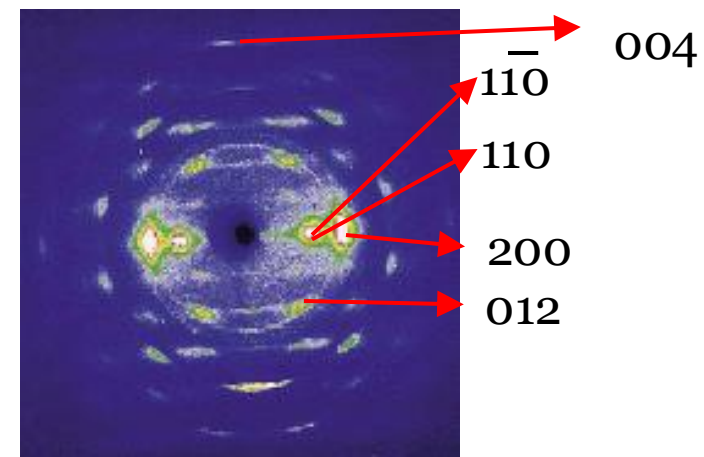


Crystalline cellulose fibers



2D diffraction pattern

→ information of crystalline orientation



# Sample types

single  
crystal



twinned  
crystal



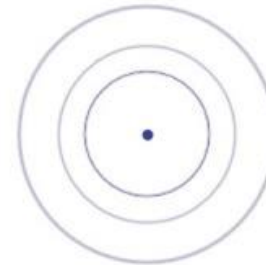
crystal with  
mosaic spread



textured  
sample

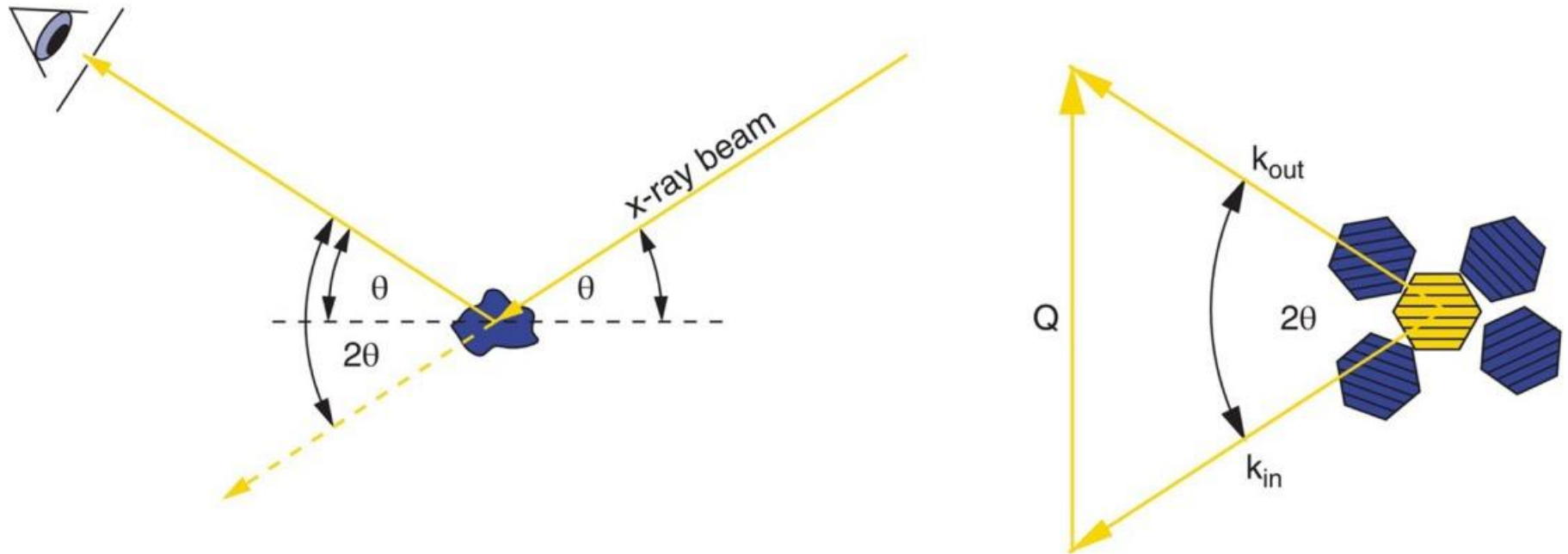


powder  
sample



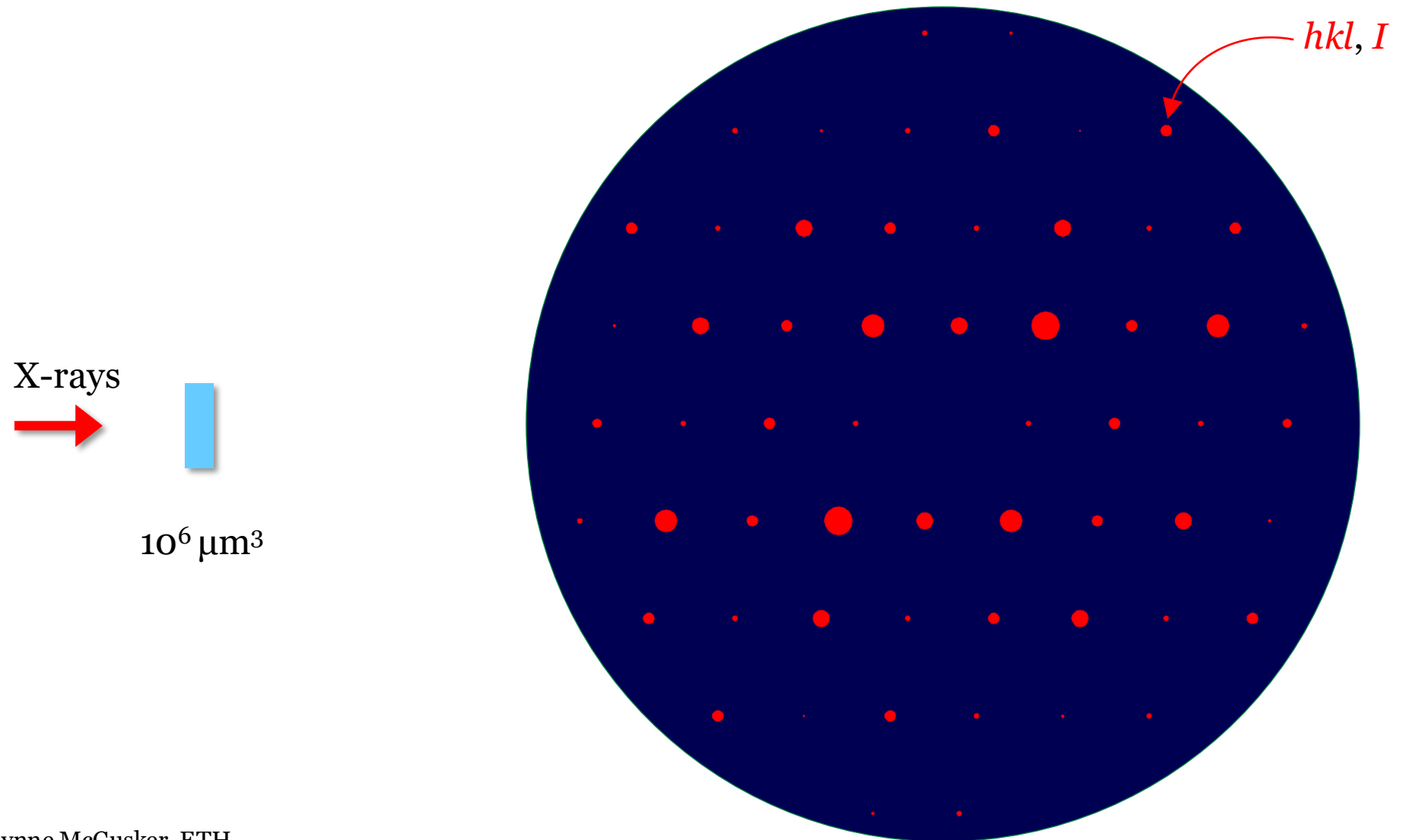


# Powder diffraction



Conditions for diffraction in a powder sample. A detector will only see a diffracted signal if the  $d_{hkl}$  spacing, the orientation of the crystallite, and the angle of the detector  $2\theta$  to the incident x-ray beam lead to the diffraction condition being satisfied. This is fulfilled by the yellow-highlighted crystallite.

# Single Crystal Diffraction

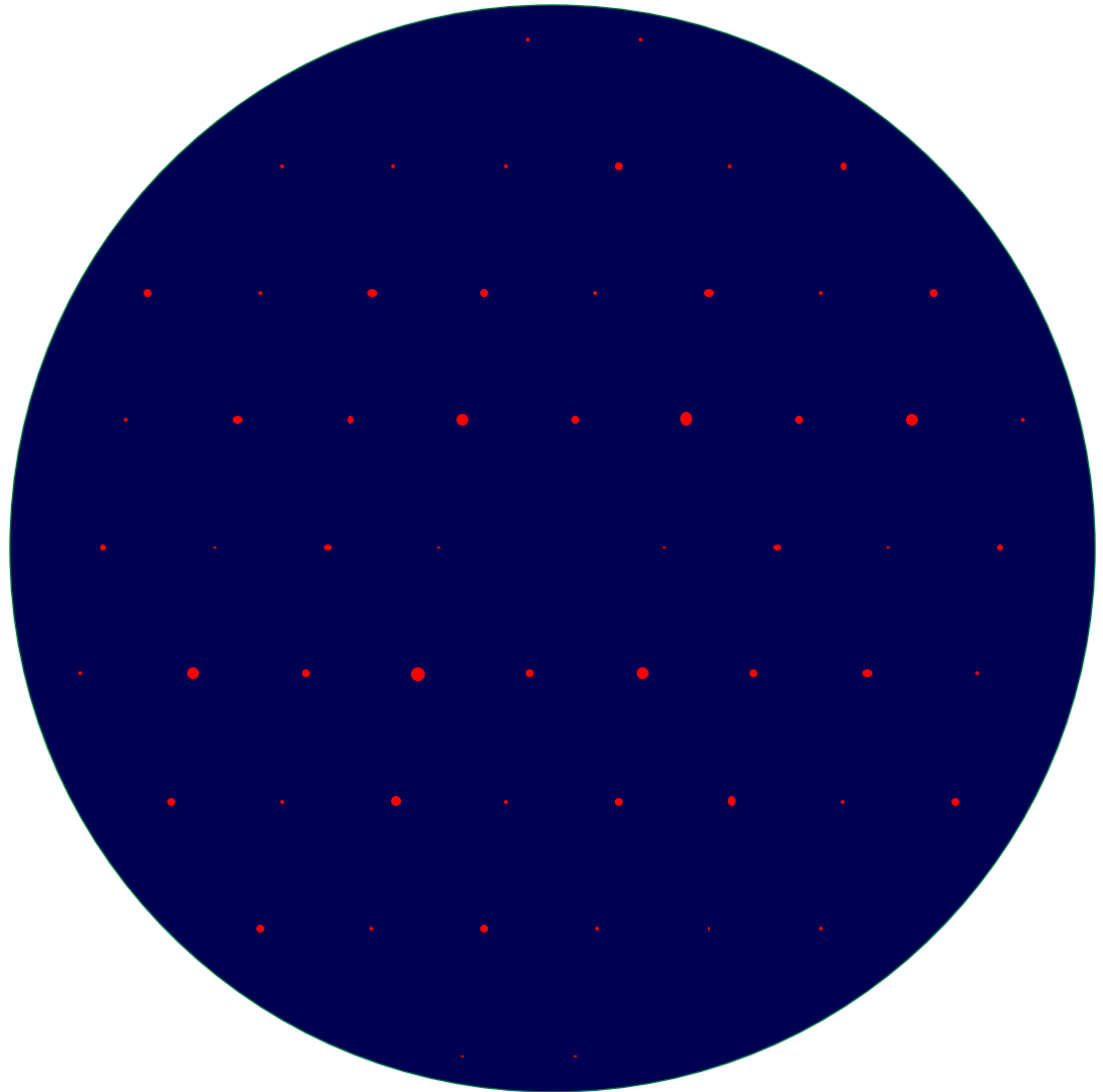


# Powder Diffraction

X-rays

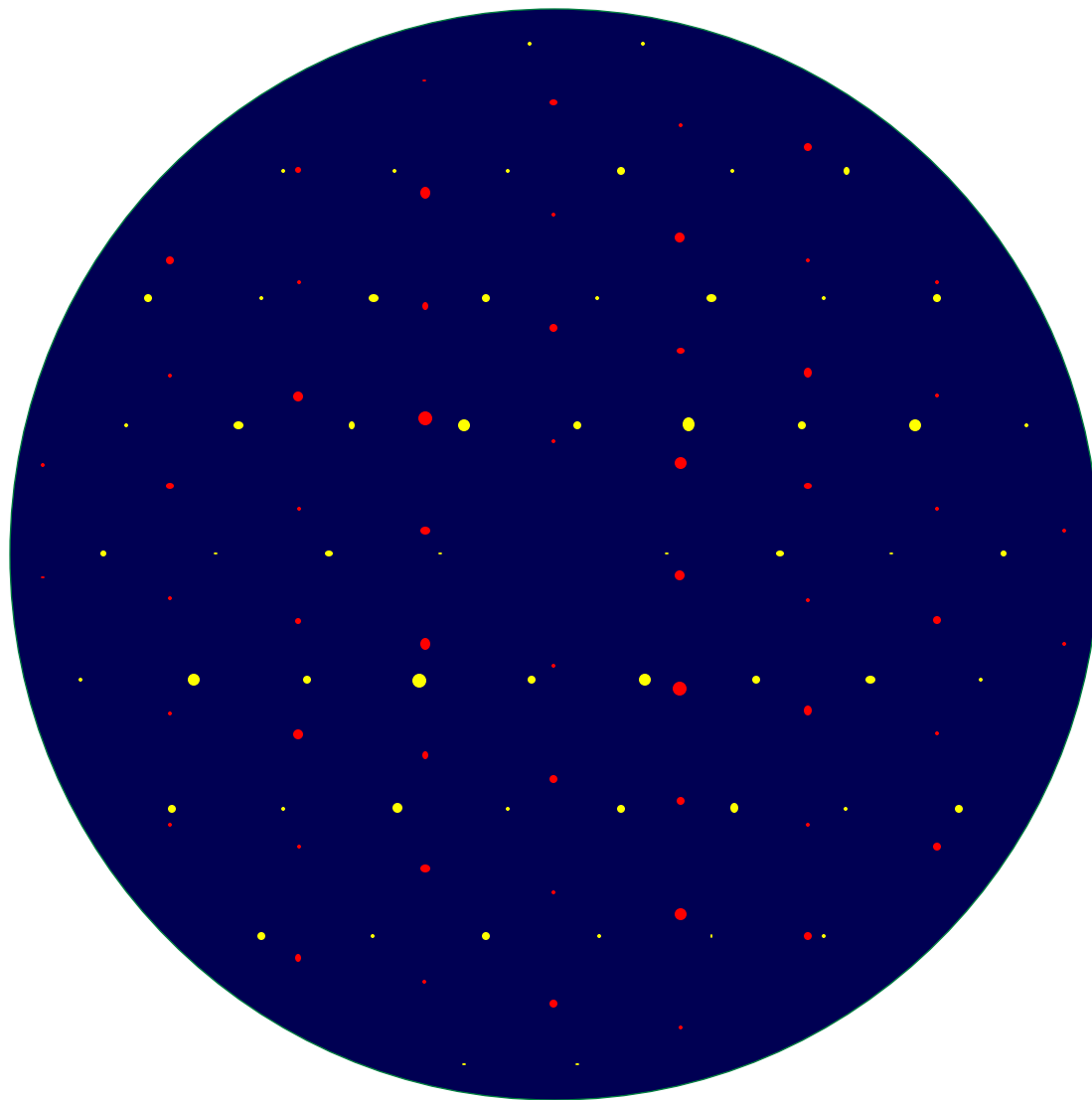


$1\ \mu\text{m}^3$



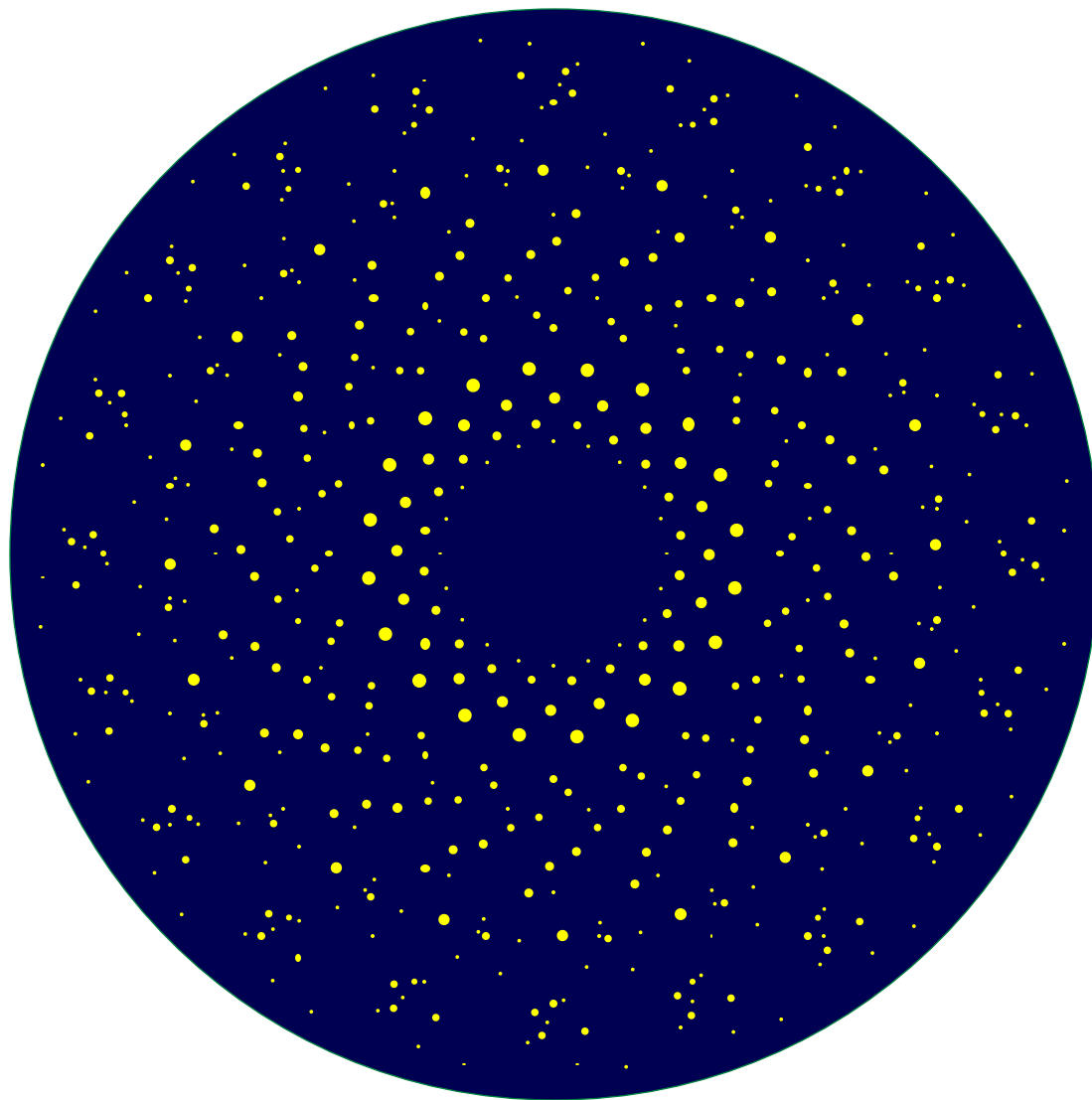
# Powder Diffraction

X-rays

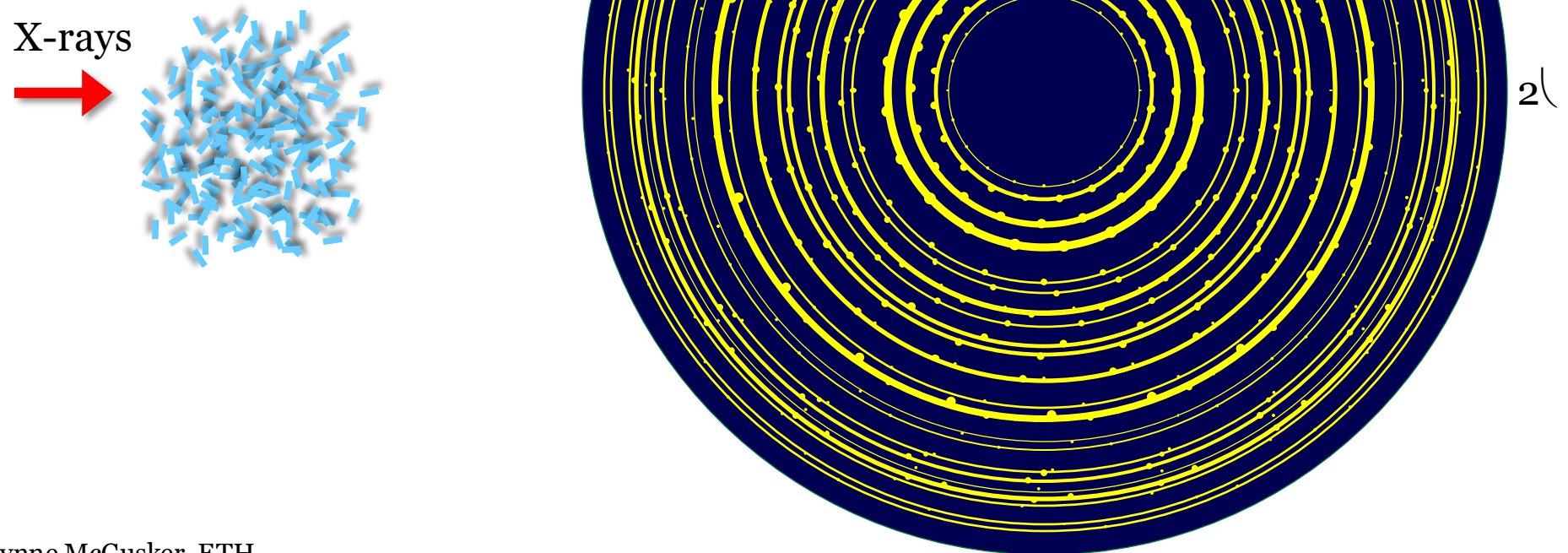


# Powder Diffraction

X-rays

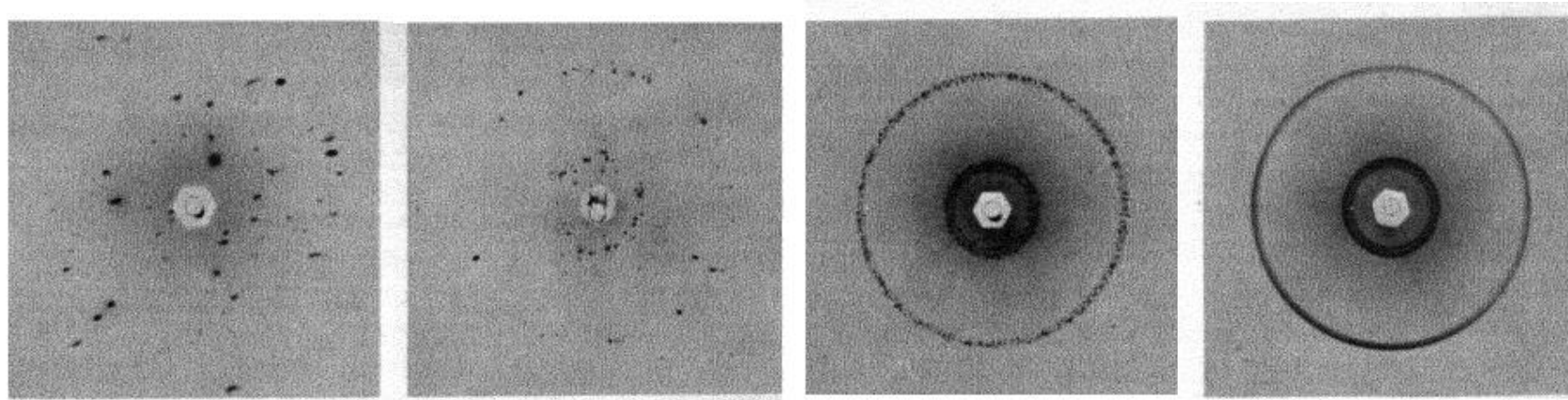


# Powder Diffraction



# Polycrystallinity

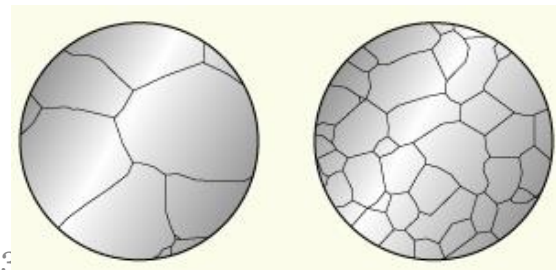
Example: XRD of recrystallized aluminium



Decreasing grain size

Grain size be determined by comparing XRD-patterns recorded under **identical conditions** (in particular beam size!)

Schematic: number of grains in beam:  
(smaller beams „see“ fewer crystallites)





# Sample types

single  
crystal



twinned  
crystal



crystal with  
mosaic spread



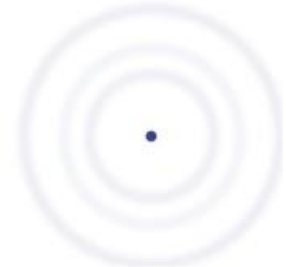
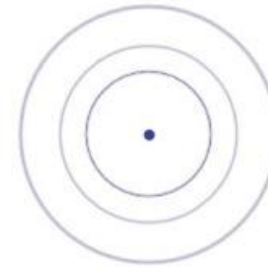
textured  
sample



powder  
sample

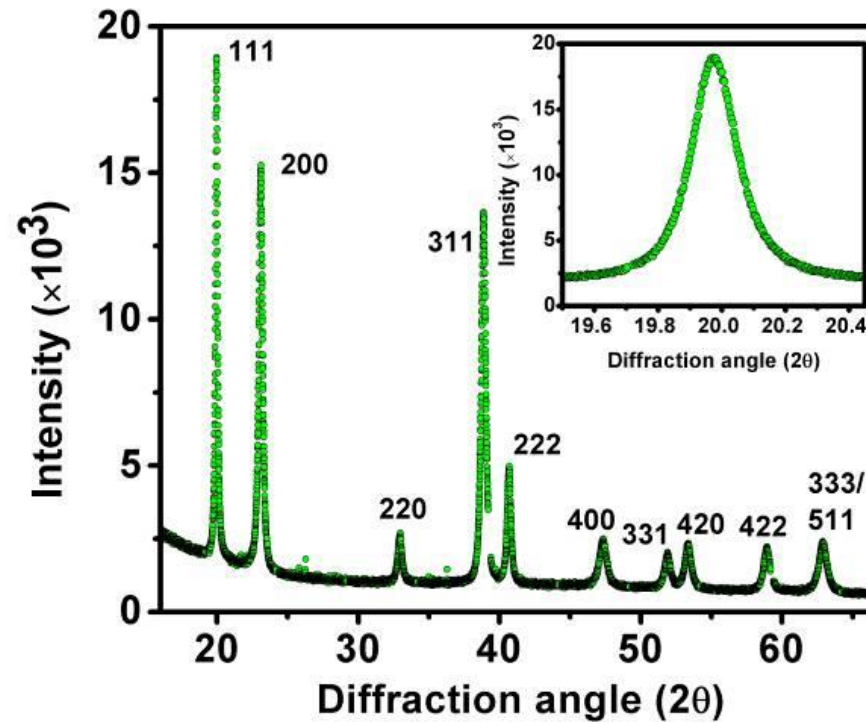


nanocrystalline  
powder



# Nanocrystalline powder

- Example: nanocrystalline Ni



# XRD

## Imperfect microstructure

- large crystal with perfect atomic arrangement give rise to perfectly sharp peak (except of instrumental broadening)
- imperfections such as grain boundaries, defects at dislocations, stacking faults, stresses → peak broadening, as well as possibly peak position shifts
- small crystal size: “defect” as the long-range atomic arrangement is disrupted at the interface → peak broadening

# XRD: Crystal size

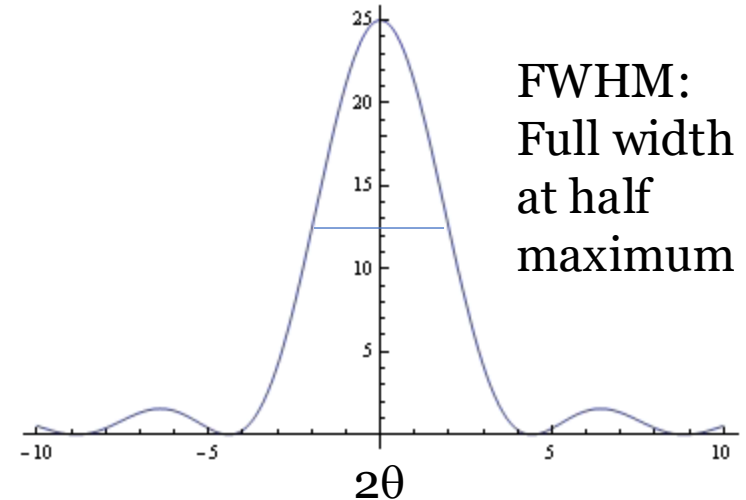
Scherrer width  $B = \frac{K \cdot \lambda}{D \cdot \cos \theta}$

B: broadening of diffraction line at half maximum, in  $2\theta$  measured in radians

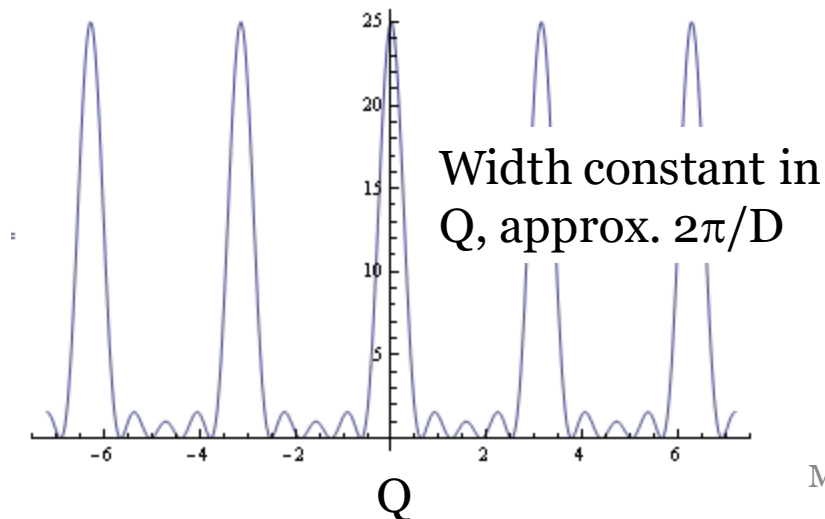
D: diameter of crystallite

K: constant,  $\approx 1$ , depends on crystal structure

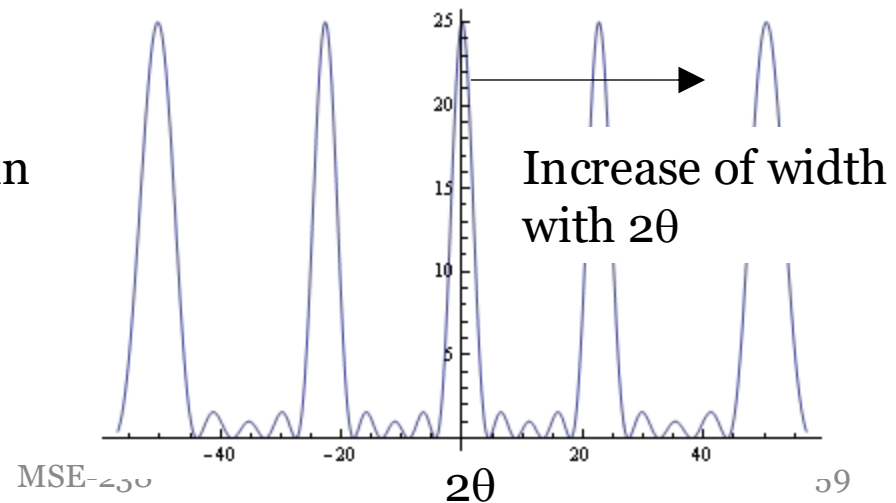
**small crystal  $\rightarrow$  broad peak**



Plot versus  $Q$



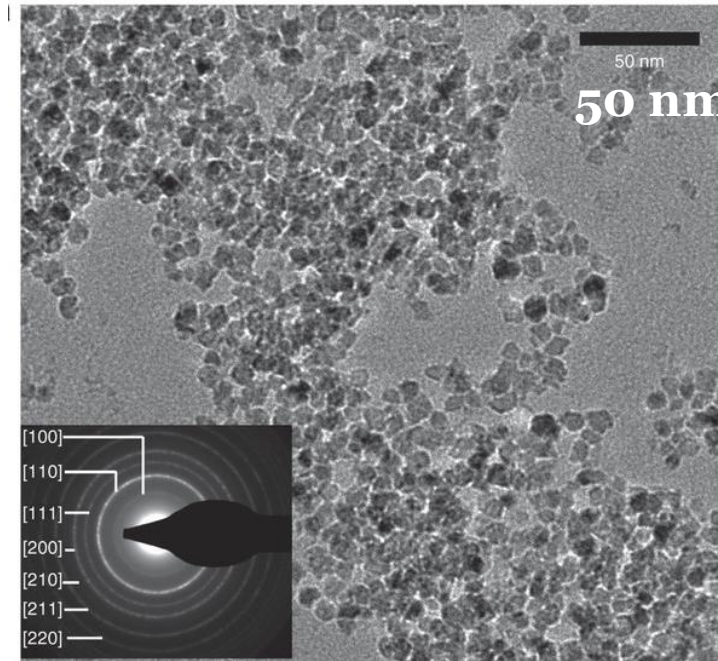
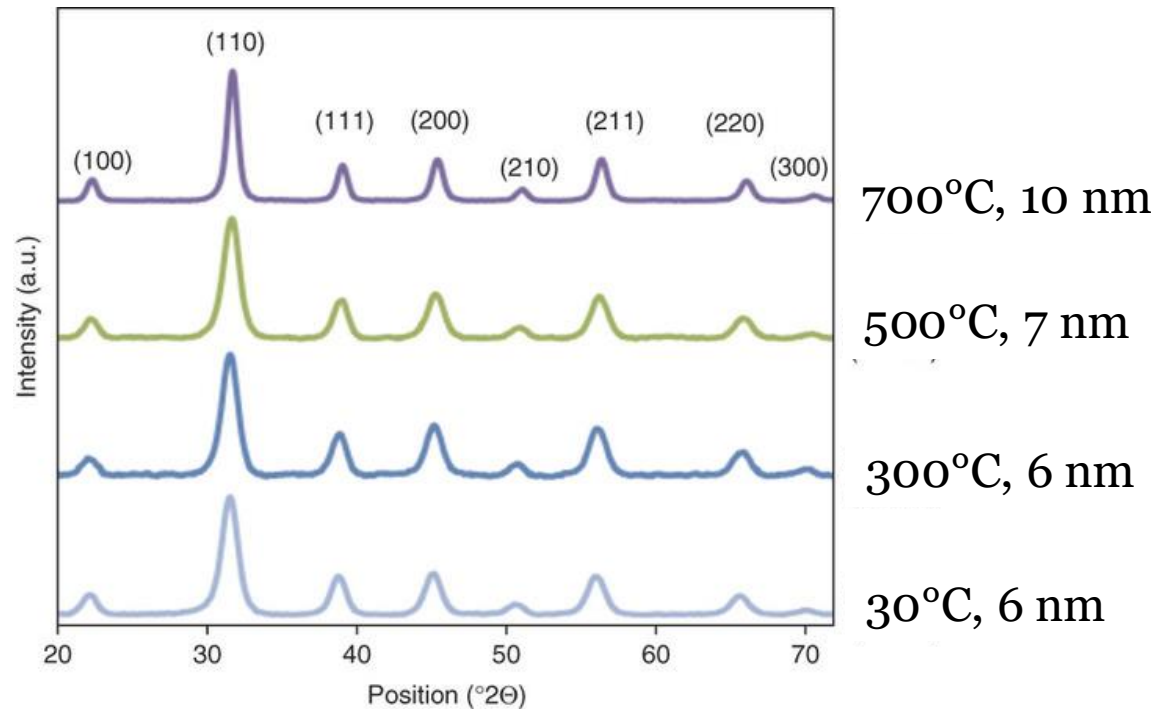
Plot versus  $2\theta$



# XRD: Crystal size

## BaTiO<sub>3</sub> nanoparticles

Large-scale synthesis of BaTiO<sub>3</sub> nanopowders using a bioinspired process at nearly room temperature (25 °C). Size changes during sintering:



# XRD: Strain broadening

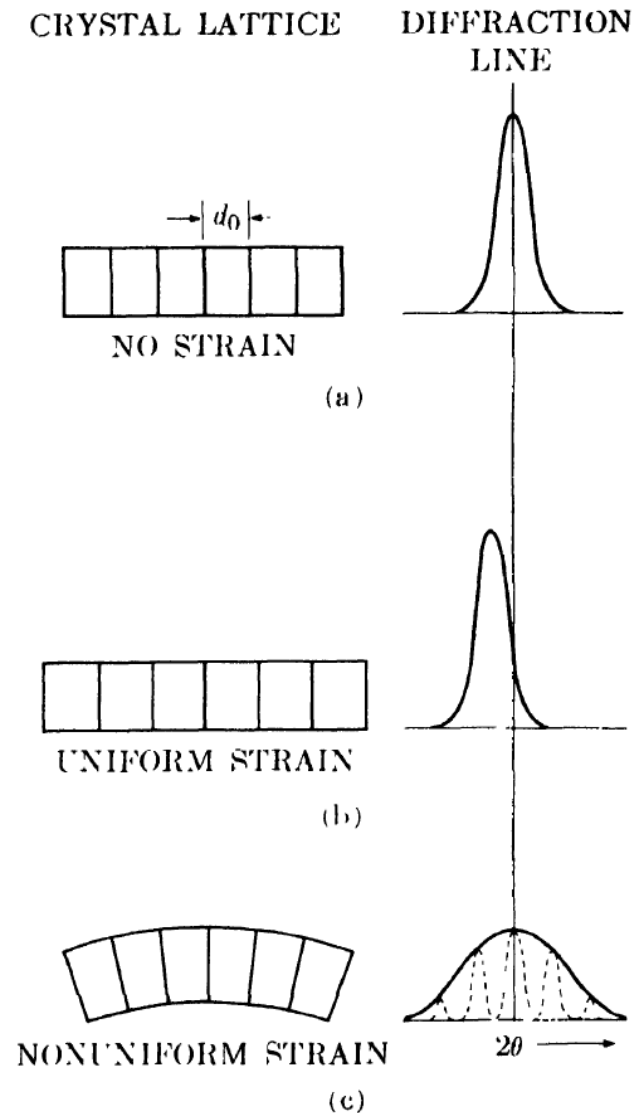
Line broadening due to non-uniform lattice distortions.

## Broadening related to strain:

$$b = \Delta 2\theta = -2 \frac{\Delta d}{d} \tan \theta = -2\varepsilon \tan \theta$$

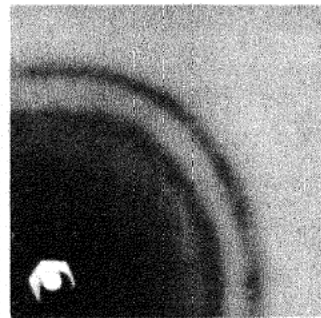
Often occurs with size broadening, difficult to separate.

Stronger dependence on  $\theta$  (width increases for higher order reflections). Also different line shape.

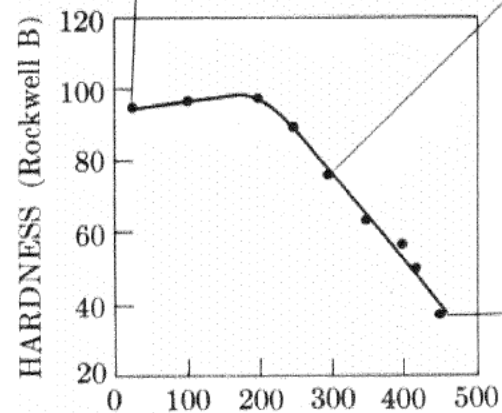


# XRD: Strain broadening

Strain broadening  
after cold rolling



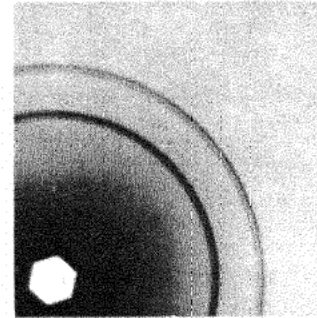
As rolled



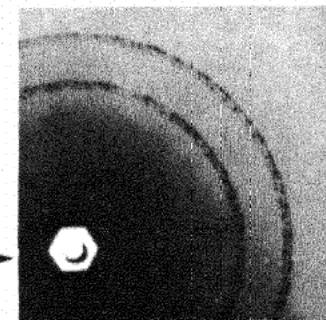
ANNEALING TEMPERATURE (°C)

(a) Hardness curve

MSE-238



1 h at 300°C



1 h at 450°C

Recovery

Recrystallization



# XRD

## Imperfect microstructure

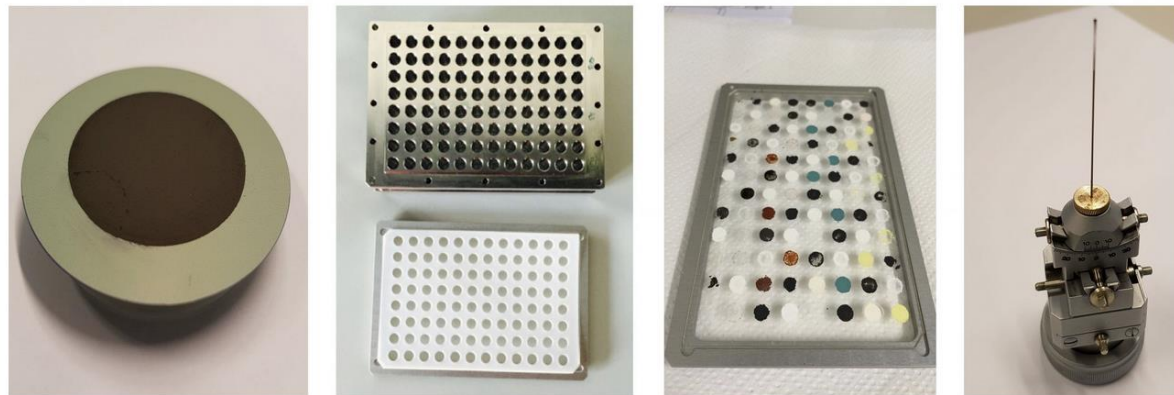
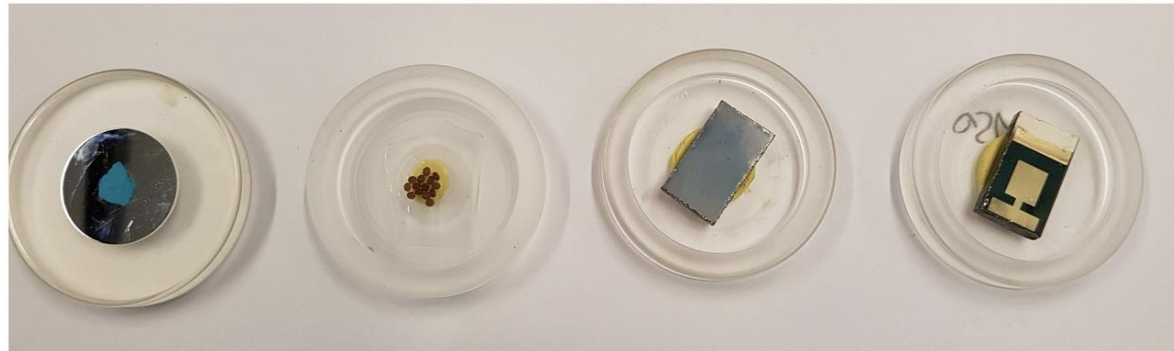
- large crystal with perfect atomic arrangement give rise to perfectly sharp peak (except of instrumental broadening)
- imperfections such as grain boundaries, defects at dislocations, stacking faults, stresses → peak broadening, as well as possibly peak position shifts
- small crystal size: “defect” as the long-range atomic arrangement is disrupted at the interface → peak broadening
- when looking at more than one order of a reflection, the effect of “size” and “strain” can be separated
- Rietfeld refinement to get all information out of powder diffraction data

# Peak profiles

- Peak profiles are determined by many factors. The most important ones include:
  - Resolution function
  - Coherent scattering length
  - Microstrain
    - Inhomogeneous elastic strain
    - Anti-phase boundaries
    - Faulting
    - Dislocations
    - Grain surface relaxation
  - Solid solution inhomogeneity
  - Temperature factors
- Peak profile is a convolution of the profiles from all of these contributions

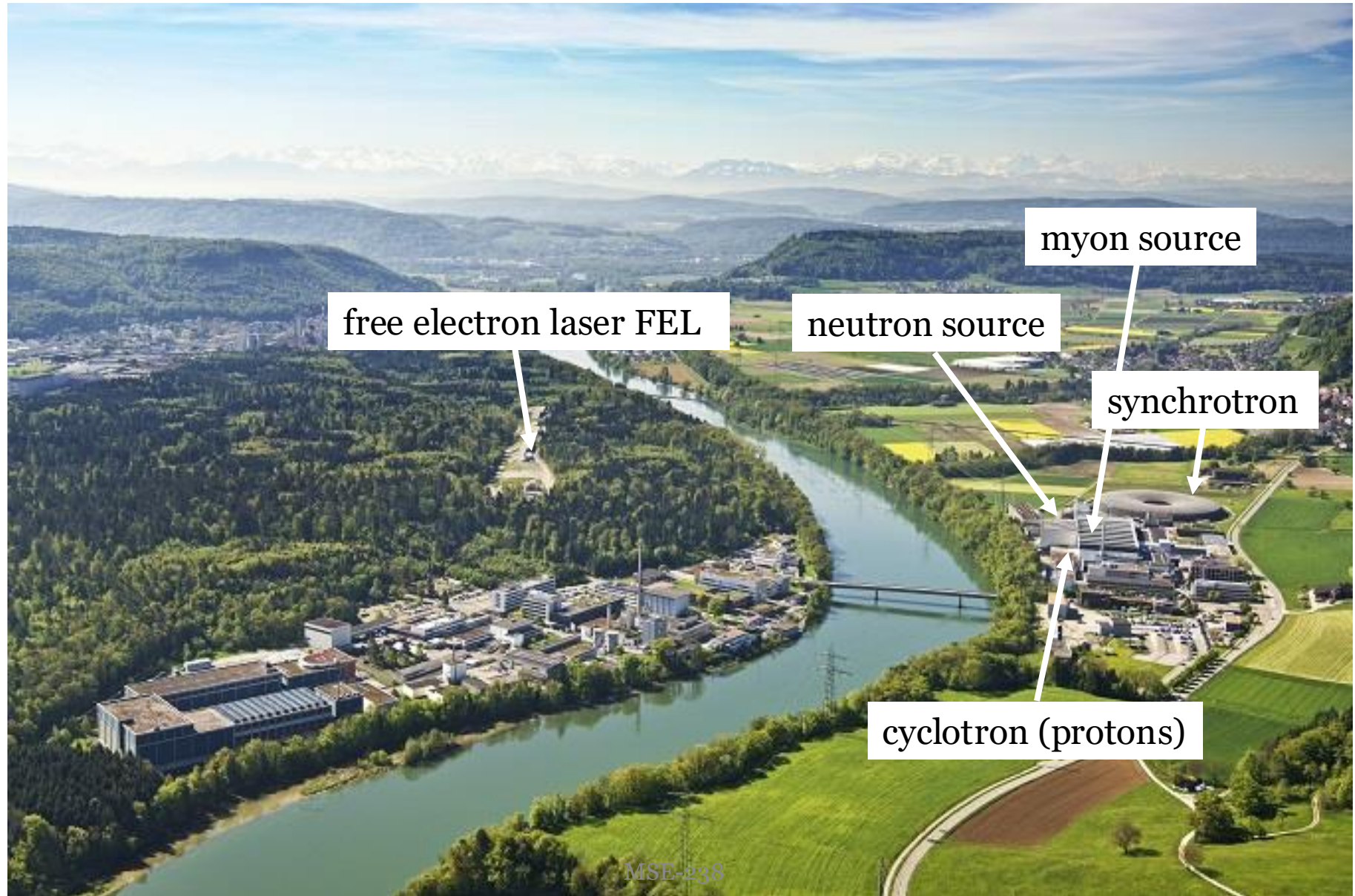
# Lab-source diffractometers for powder and single-crystal diffraction

<https://www.epfl.ch/schools/sb/research/isic/platforms/x-ray-diffraction-and-surface-analytics/x-ray-instrumentation/x-ray-scattering-instrumentation/>

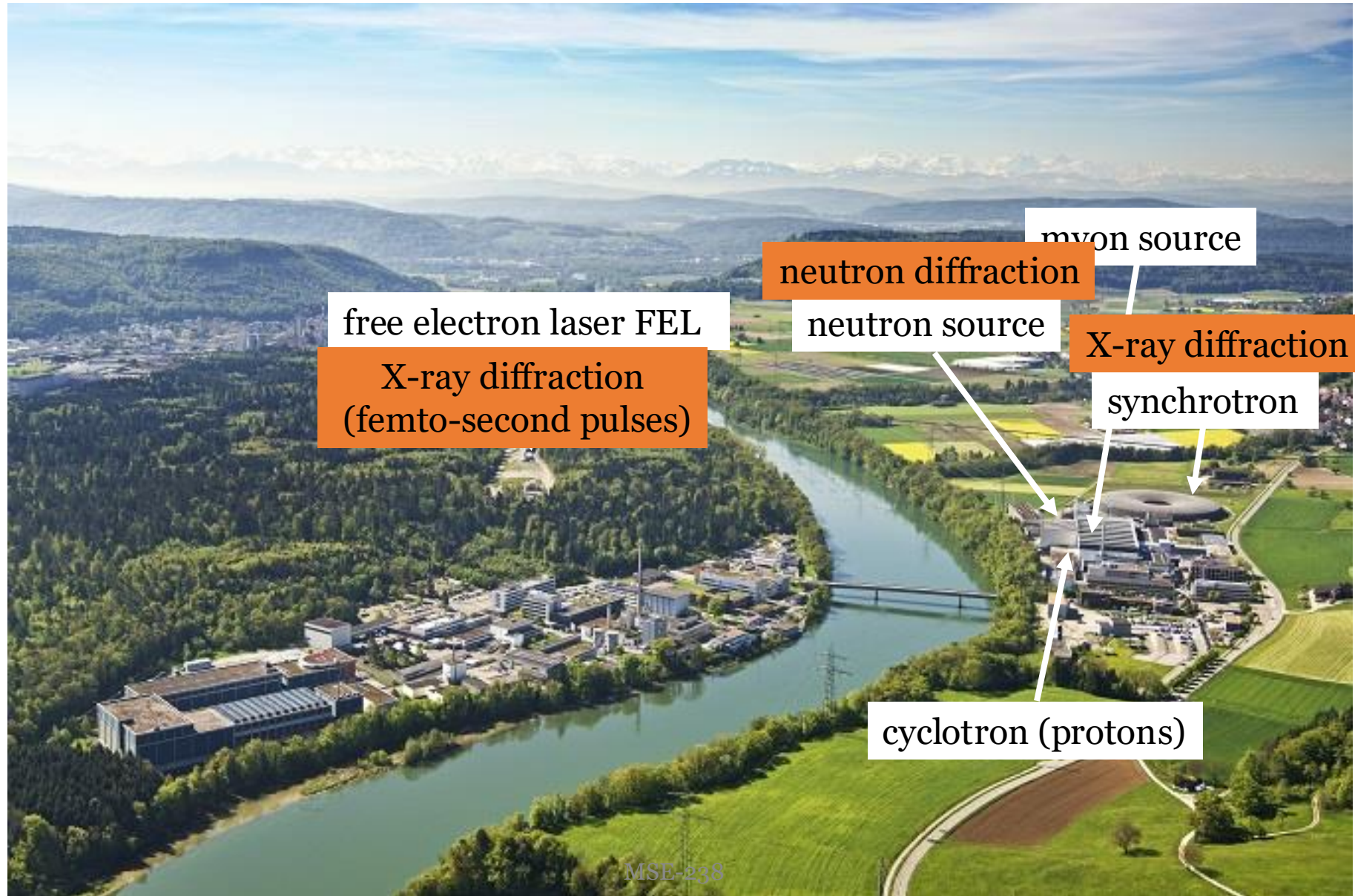


**Samples:** some examples of commonly measured samples are shown on **top**, from left to right: Metalorganic framework (MOF) powder sample, polymer beads, perovskite thin film, perovskite film with electrode. **bottom**, left to right: Solid Oxide Fuel Cell (SOFC), high throughput sample changer for synthesis robot (before and after loading), powder under Argon atmosphere in glass capillary.



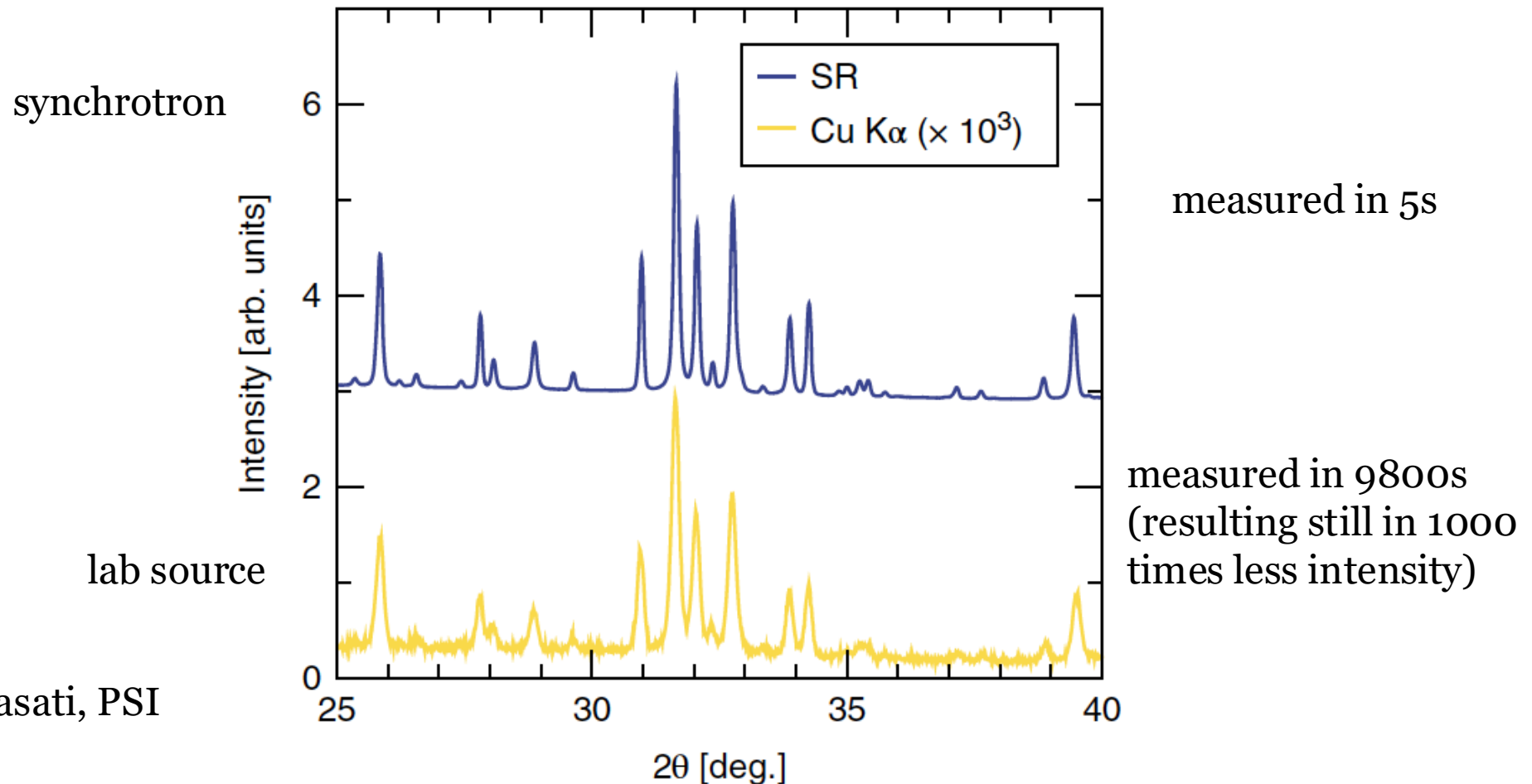






# Lab source vs. Synchrotron

powder diffraction of Calcium Phosphate



N. Casati, PSI

# Summary

## Diffraction:

- The scattering vector  $q$
- Ewald sphere to determine which reciprocal lattice points are in Bragg condition in a certain geometry
- single crystal diffraction: rotation method or Laue diffraction
- imperfect crystals: defects, grainboundaries, strain: change peak width and/or position
- preferred direction of certain crystallographic directions: textured sample influence of symmetry
- polycrystalline material without preferred direction: “powder sample”
- X-ray diffraction Lab sources vs. synchrotrons