

# Biology I: Structural Biology and Macromolecular Crystallography (MX)

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18/6/2013  
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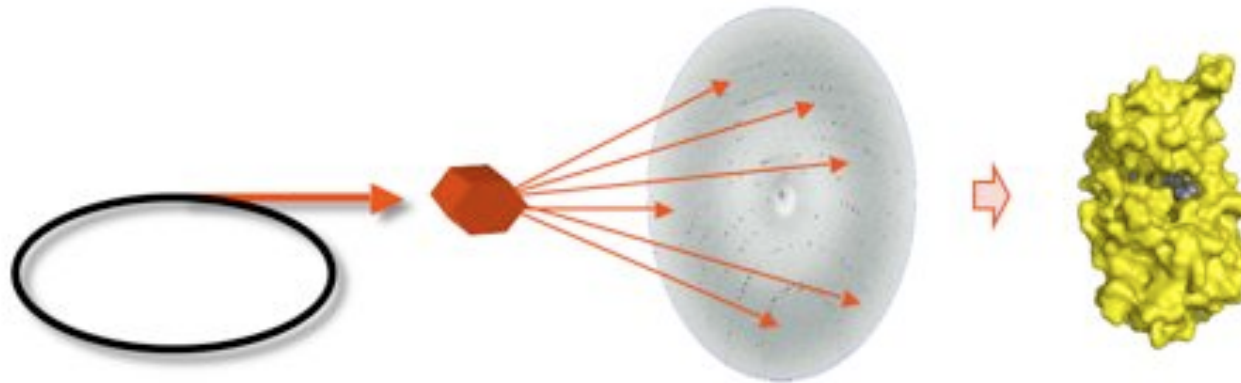
# EMBL

- Funded 1974
- Inter-governmental organization
- 20 member states
- One associate state
- Missions
  - Basic Research in Molecular Biology
  - Technology and Instrumentation
  - Facilities and Services
  - Teaching and Training
  - Technology Transfer
- 5 sites
- 1400 employees from 60 nations
- Annual Budget ca. 85 M€



# Biology I to IV

- Structural Biology and Macromolecular Crystallography (MX)
- MX – The method
- MX – Collection and processing of diffraction data
- MX – Building models and the future



# Today

- What is a protein?
- The central dogma of molecular biology
- Milestones in macromolecular crystallography
  - The first structures
  - Recombinant production of proteins
  - Use of synchrotron radiation
  - Detector technologies
  - The protein data bank
  - Cryogenic sample cooling
- Crystallographic Workflow

# Look at Abl kinase

- 2148 atoms per molecule, 275 amino acids
- A protein consists of chained amino acids (20 types)
- Backbone vs. Sidechain
- alpha-helices
- beta-sheets
- Schematic 'cartoon' representations of proteins
- Surface representation of proteins (with properties mapped)
- Interactions with ligands can be studied
- Gleevec / Imantinib
  
- Folding (Stretch out 275 amino acids  $\rightarrow$   $3.8 \text{ cm} \times 275 = 10.41 \text{ m}$ , compare to  $53 \text{ \AA} / \text{cm}$ )

Downloads:

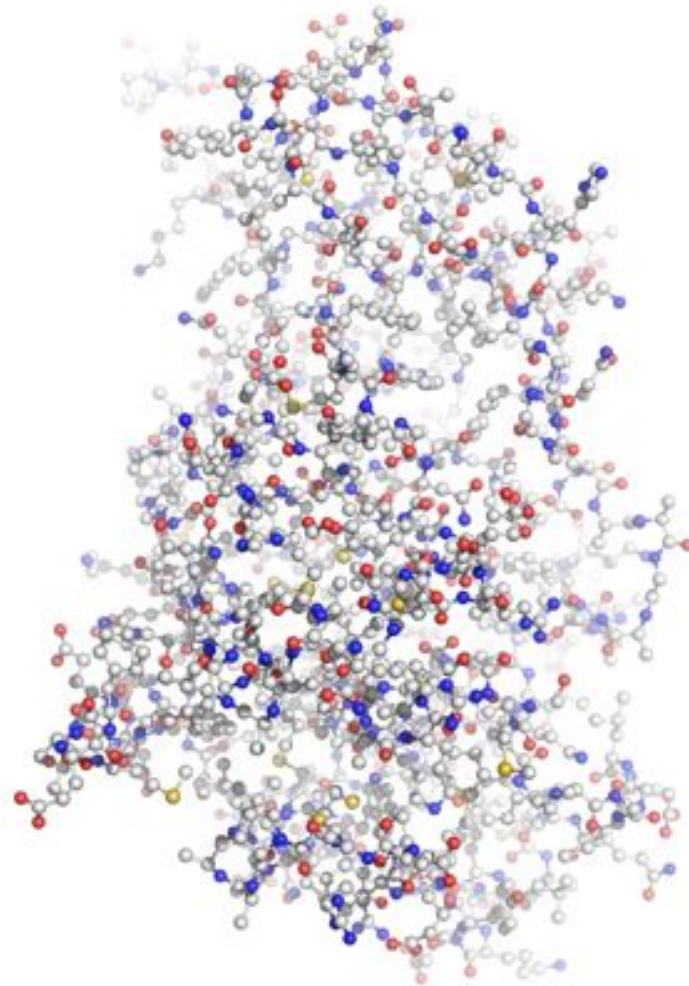
molecule viewer: <http://pymol.org/educational/>

pdb-model: <http://www.rcsb.org/pdb/explore/explore.do?structureId=1FPU>

viewer script: see course website

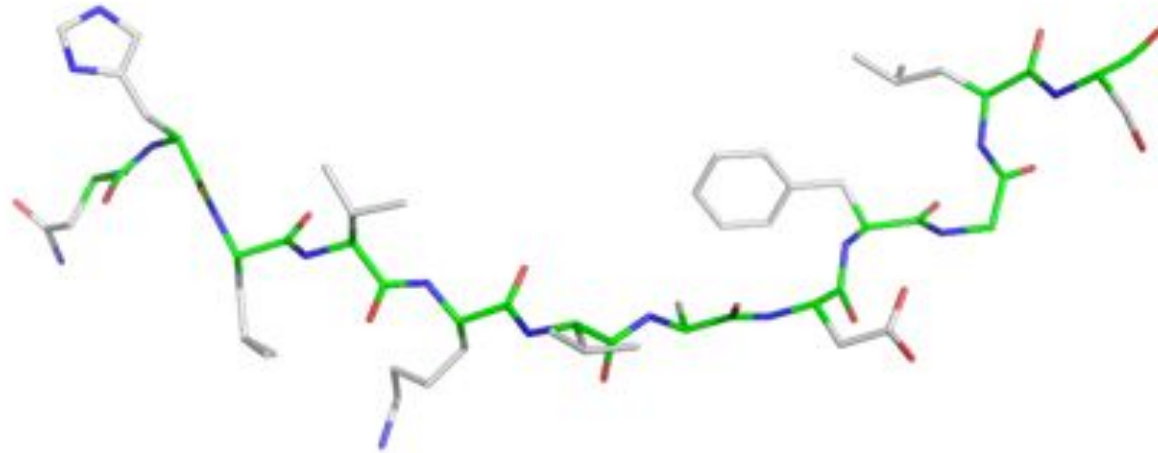
# A protein molecule

- Looking at a collection of >2000 atoms can be rather confusing:



# Polypeptide chain

- Proteins are heteropolymers of amino-acids:



# Protein Backbone

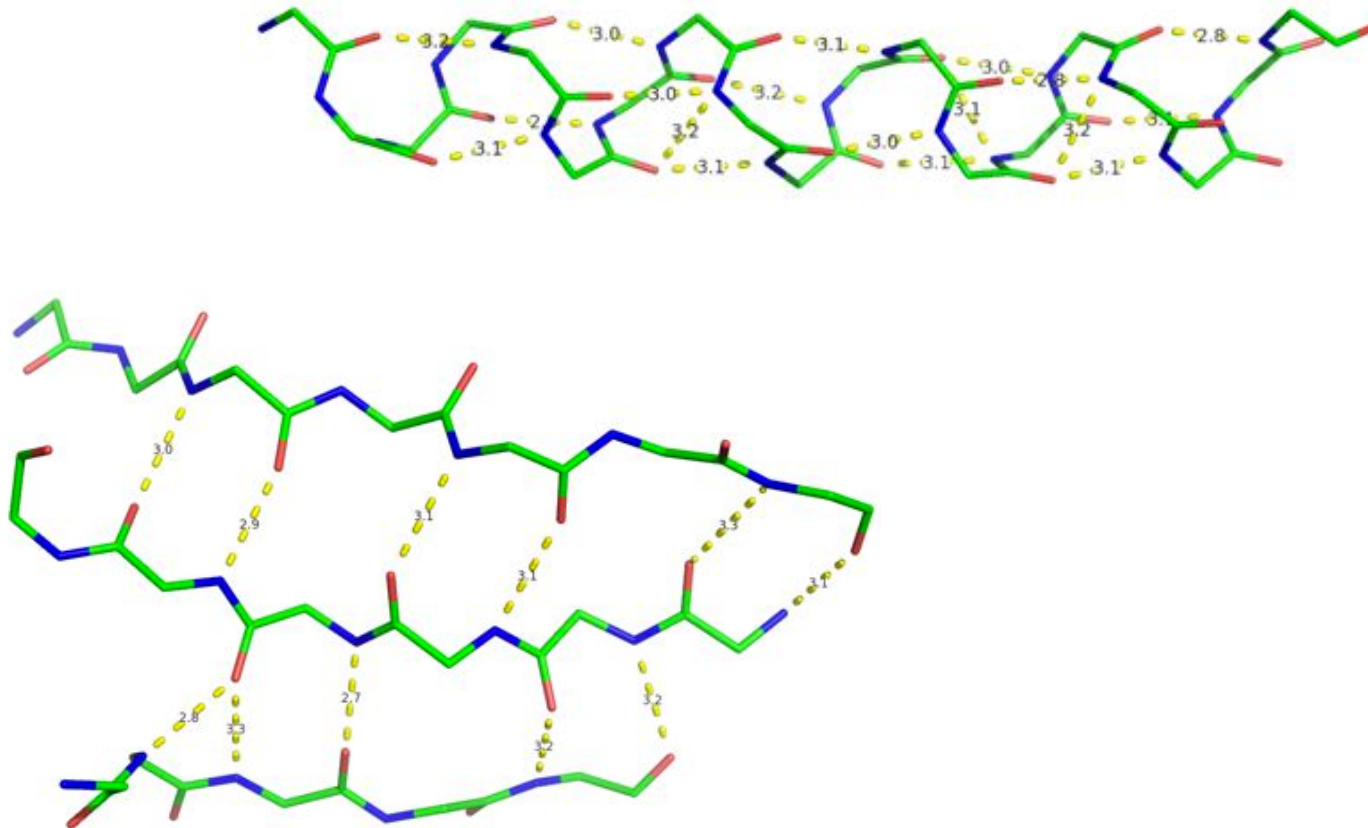
- Looking at the backbone only allows to follow the polypeptide chain across the protein





# Protein conformation

- alpha helices and beta sheets contain repeating pattern / hydrogen bonds:



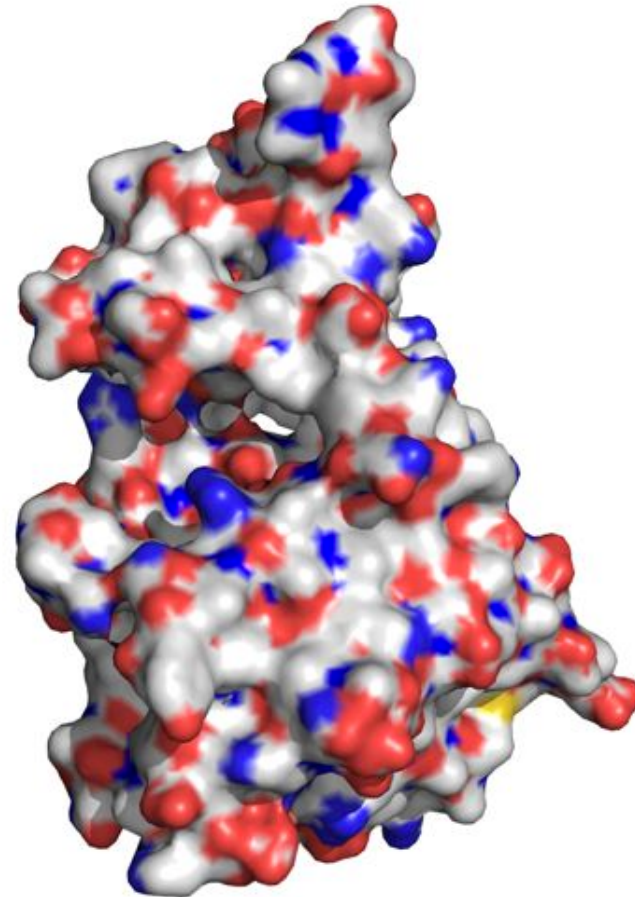
# Protein conformation

- Schematic representation of helices and sheets illustrates the 'fold'



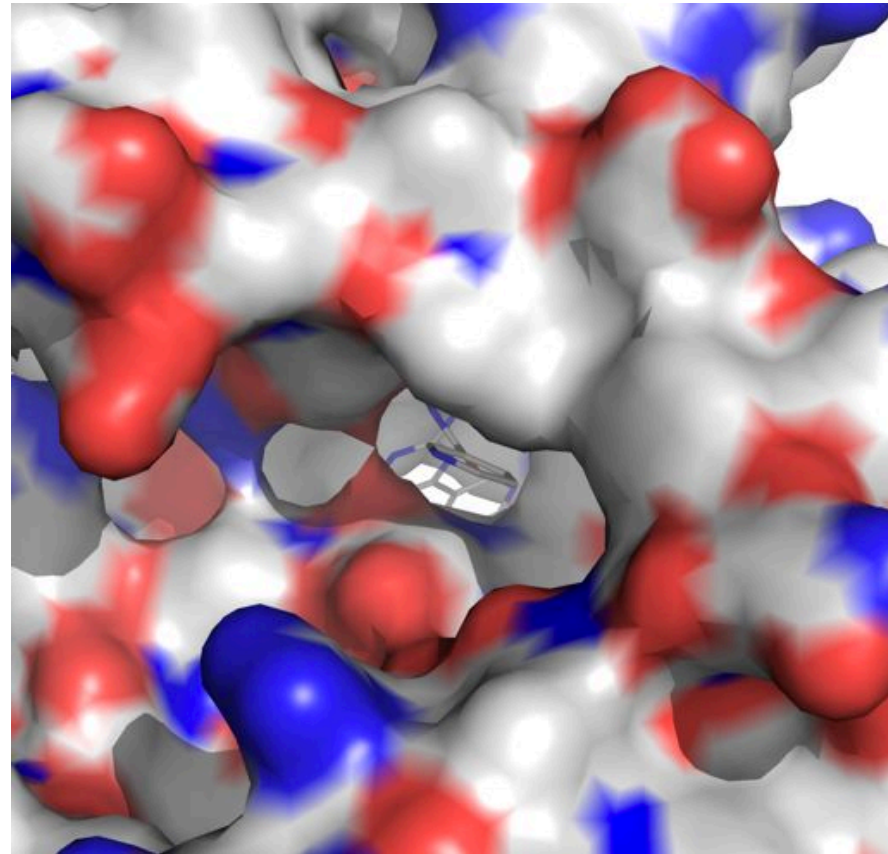
# Protein surfaces

- What other molecules 'see' is a molecular surface.
- Active sites are often found in cavities



# Active sites

- Knowledge of the active site geometry allows to design ligands binding to the active sites thus acting as drugs.
- Here: Gleevec binding Abl-Kinase

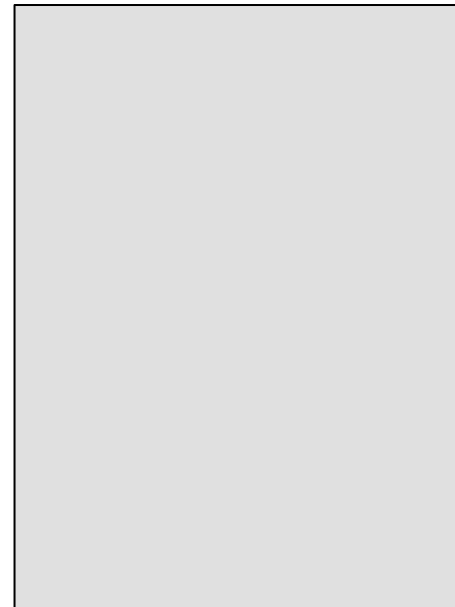


# Crystal structures can be used to understand the action of drugs



## The drug

Mortality rate reduced by 80% for patients where interferon did not work

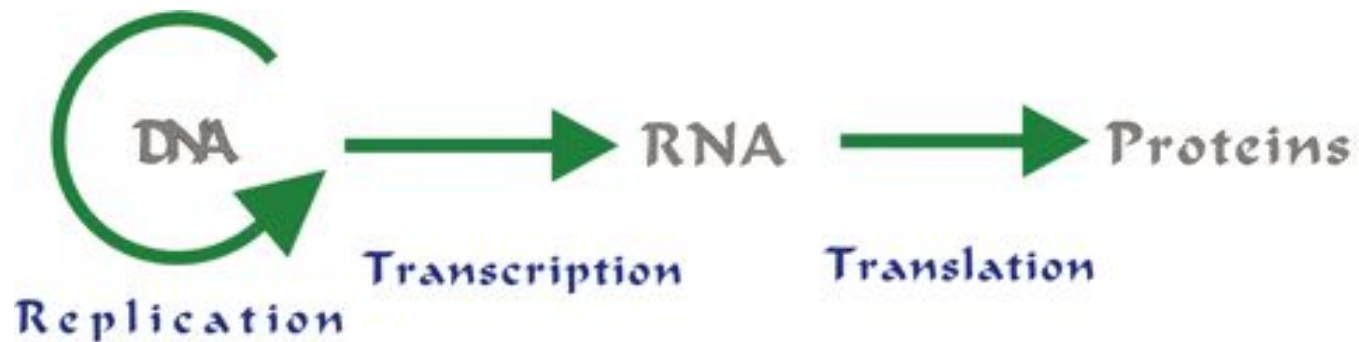


## Crystal structure of the drug bound to its target Abl-Kinase

red shows resistant mutation

# The Central Dogma of Molecular Biology

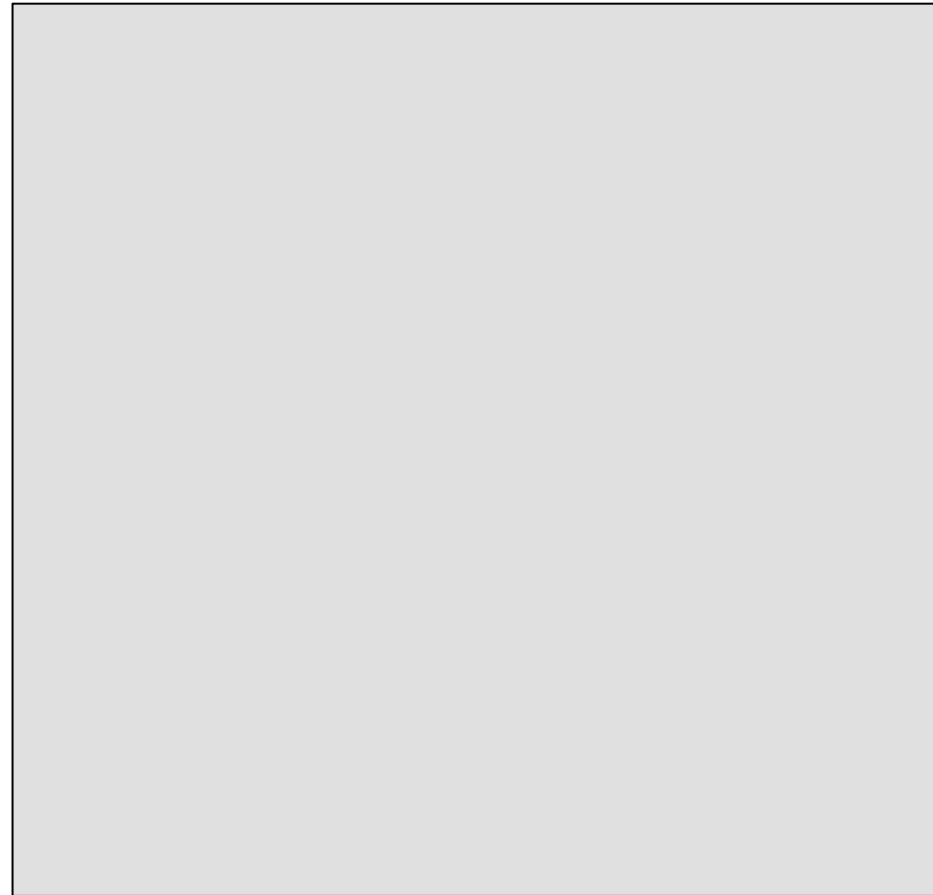
- 'DNA makes RNA makes protein' (Nirenberg 2010)



<http://library.thinkquest.org/C0122429/intro/genetics.htm>

# Crystal Structures support Basic Biology

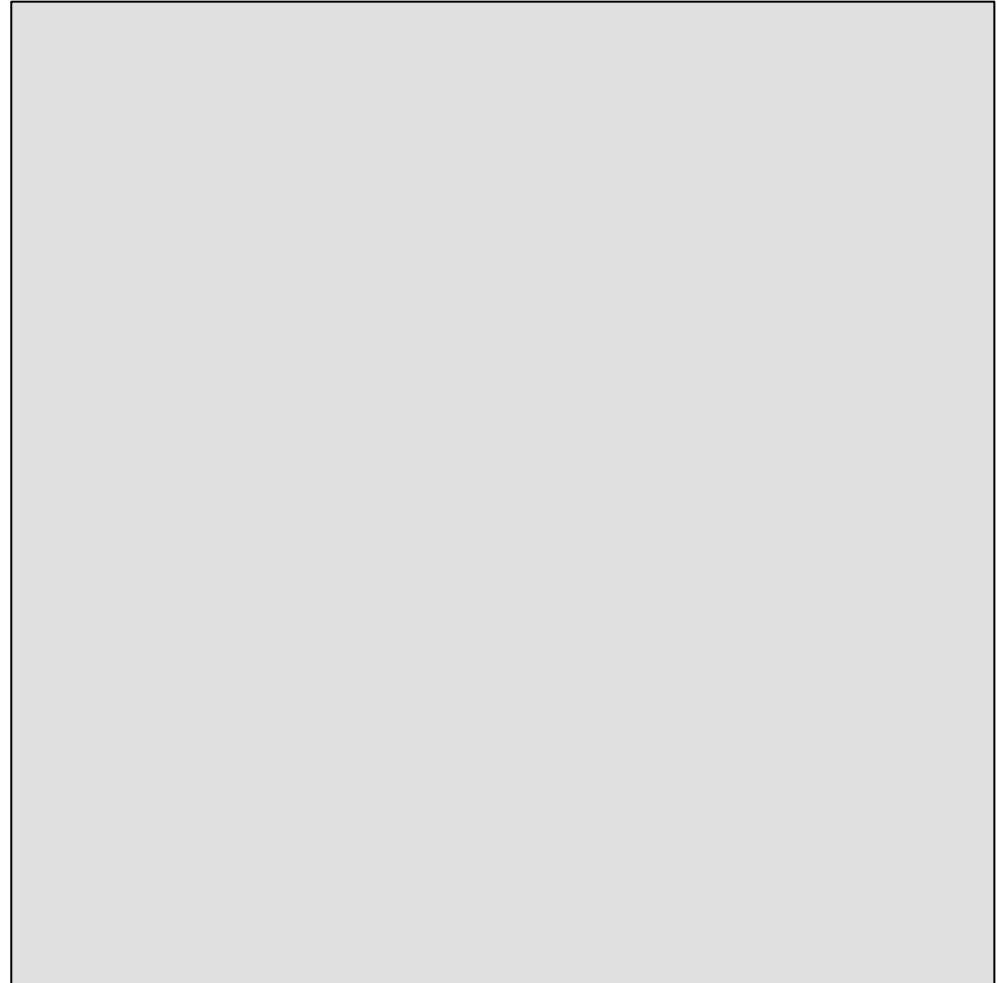
- RNA Polymerase II – Structural Basis of Transcription
- 12 proteins
- ~30000 atoms
- Nobel Prize 2006 to Roger Kornberg



<http://www.youtube.com/watch?v=6QMPU9nuQso>  
<http://www.lmb.uni-muenchen.de/cramer/pr-materials/index.htm>

# Crystal Structures support Basic Biology

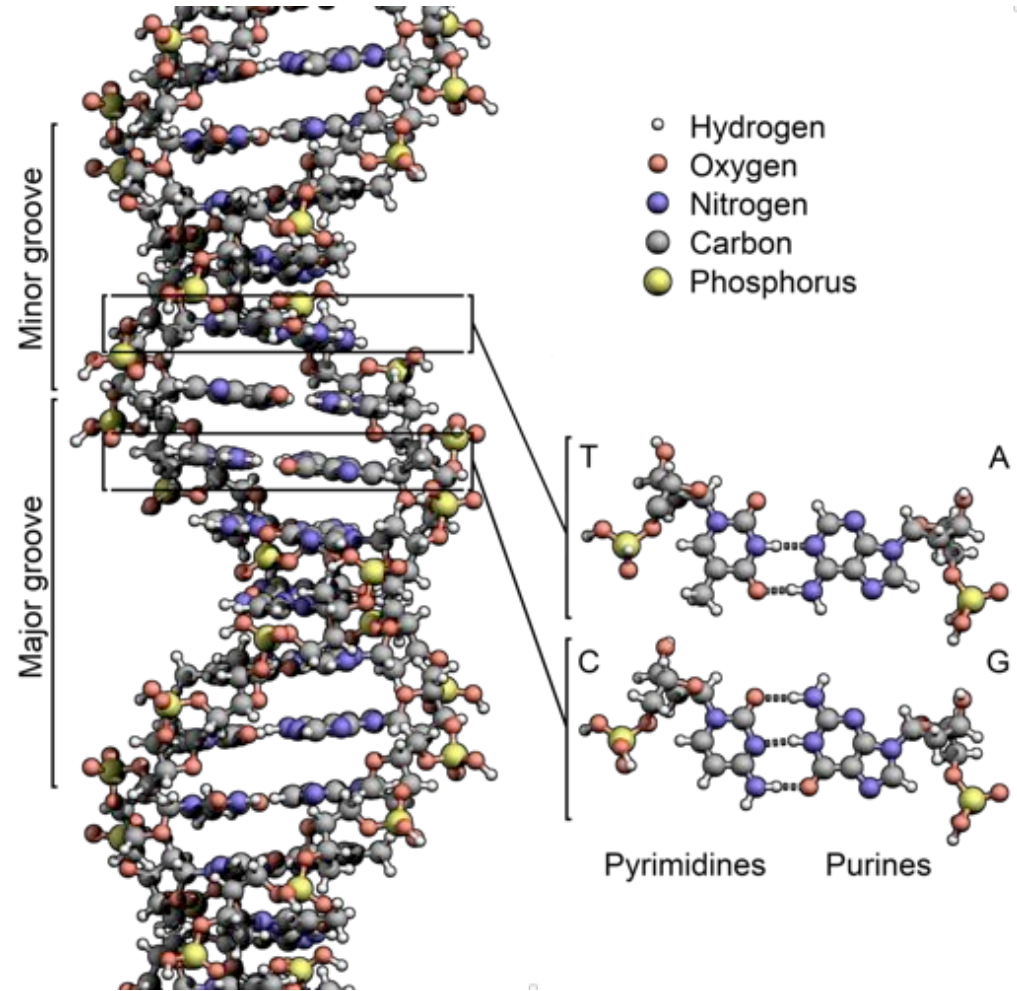
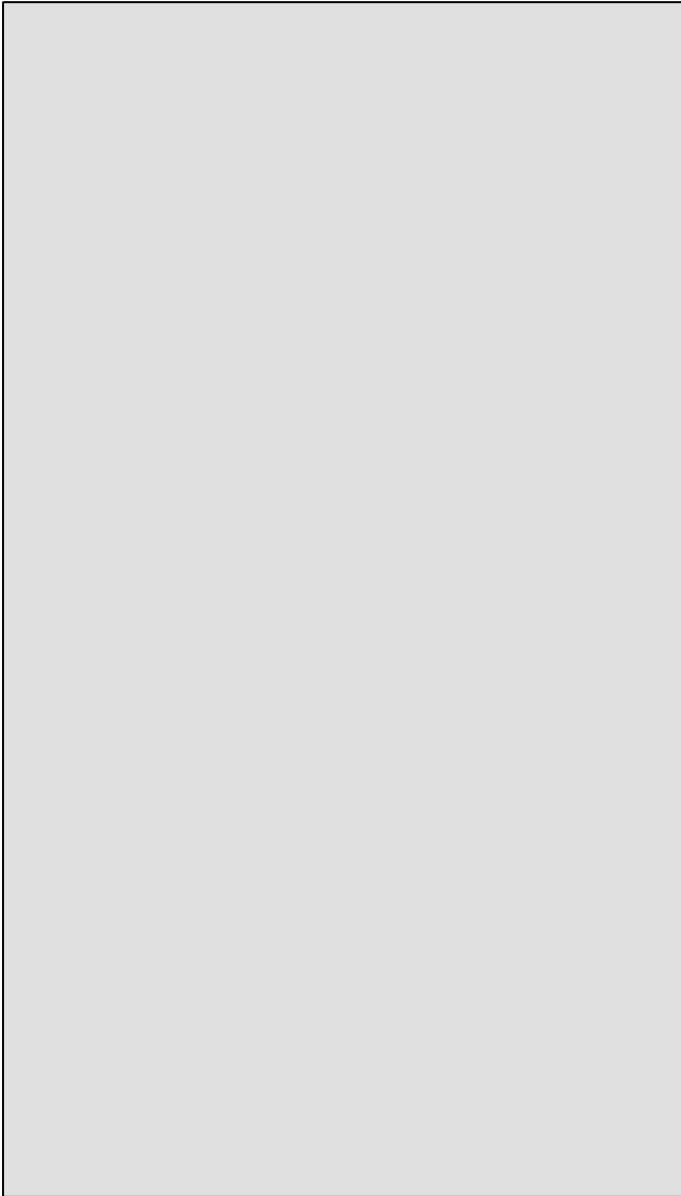
- Ribosome – Structural Basis of Translation
- 3 stretches of RNA, 52 proteins
- ~37000 RNA atoms, ~22000 protein atoms
- Nobel Price 2009 to Steitz, Yonath, Ramakrishnan





# Some milestones

# DNA (1953)



Watson & Crick (1953) Nature 172:137

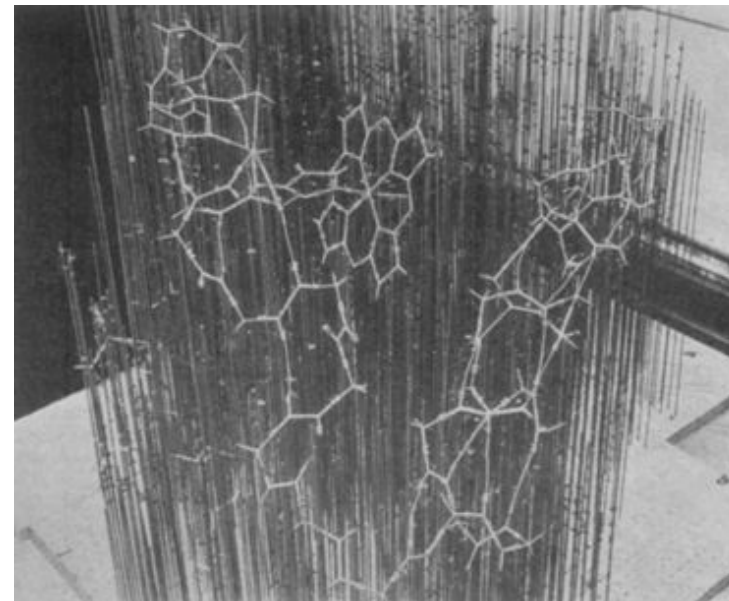
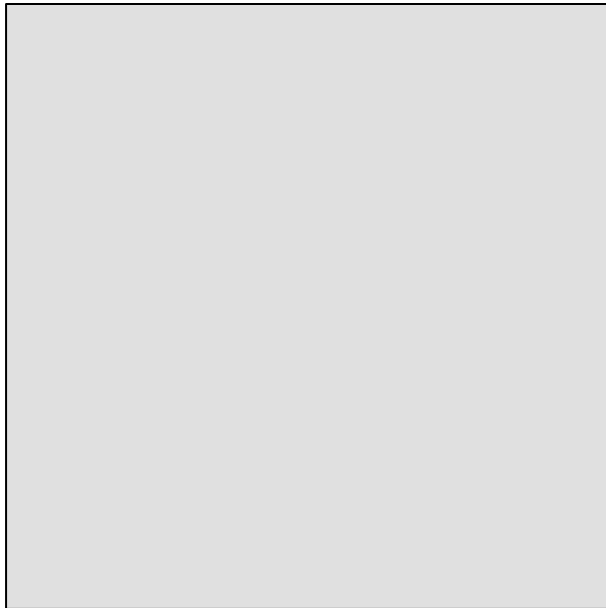
[http://www.nobelprize.org/nobel\\_prizes/medicine/laureates/1962/](http://www.nobelprize.org/nobel_prizes/medicine/laureates/1962/)

[http://commons.wikimedia.org/wiki/](http://commons.wikimedia.org/wiki/File:DNA_Structure%2BKey%2BLabelled.pn_NoBB.png)

[File:DNA\\_Structure%2BKey%2BLabelled.pn\\_NoBB.png](http://commons.wikimedia.org/wiki/File:DNA_Structure%2BKey%2BLabelled.pn_NoBB.png)

# Myoglobin (1960)

- Started in 1954
- Structure at 2 Å resolution published in 1960
- Key-step: choose sperm-whale as source



KENDREW, J. C. et al. (1960) *Nature*, 185, 422–427.

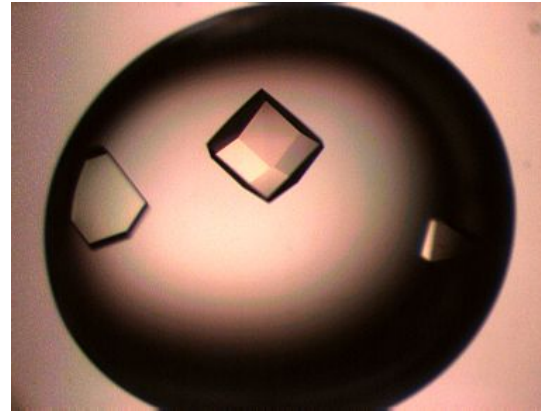
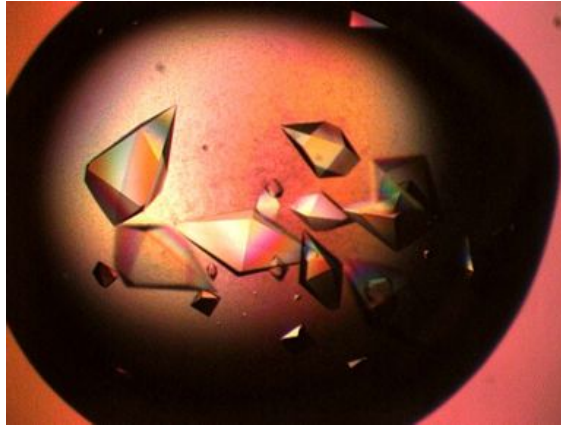
[http://www.nobelprize.org/nobel\\_prizes/chemistry/laureates/1962/kendrew-lecture.html](http://www.nobelprize.org/nobel_prizes/chemistry/laureates/1962/kendrew-lecture.html)

pdb: 1MBN

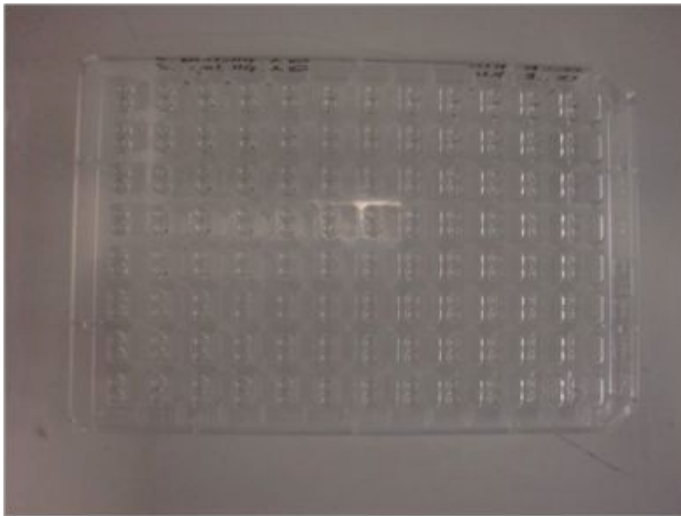
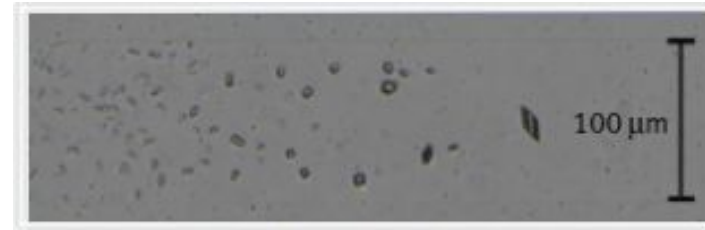
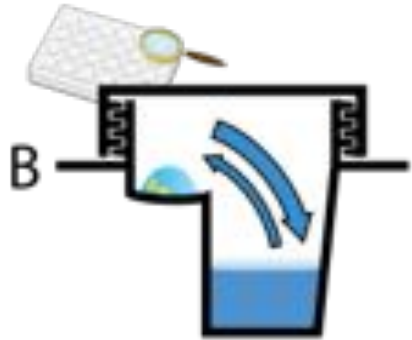
# Crystals

- Are difficult to grow
- Are difficult to reproduce
- Are often very small (microns)
- Are often inhomogeneous
- Are mechanically fragile
- Are radiation sensitive

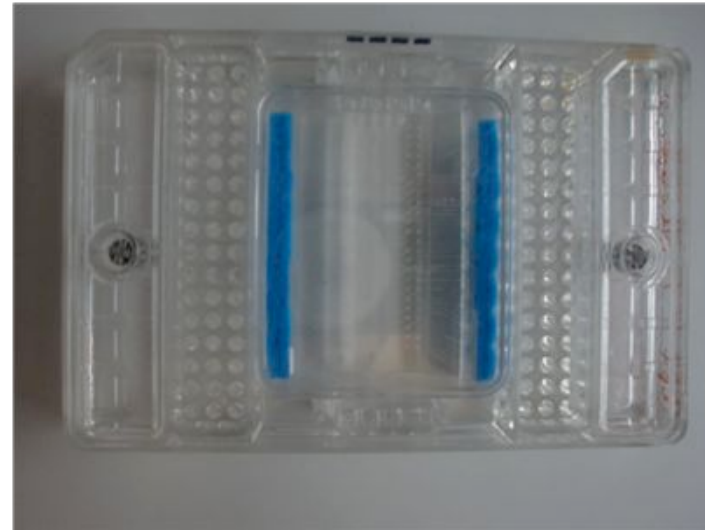
# Finding Conditions



# Small Volume HTP Crystallization



96 x 50-200 nl



96 x 10 nl

# High-Throughput Crystallization at EMBL Hamburg

- Miniaturization
  - 200nl/exp for vapour diffusion
  - 10nl/exp for counter diffusion
- Reference:
  - J. Müller-Dieckmann (2006) Acta Cryst. D64:1146-1152.

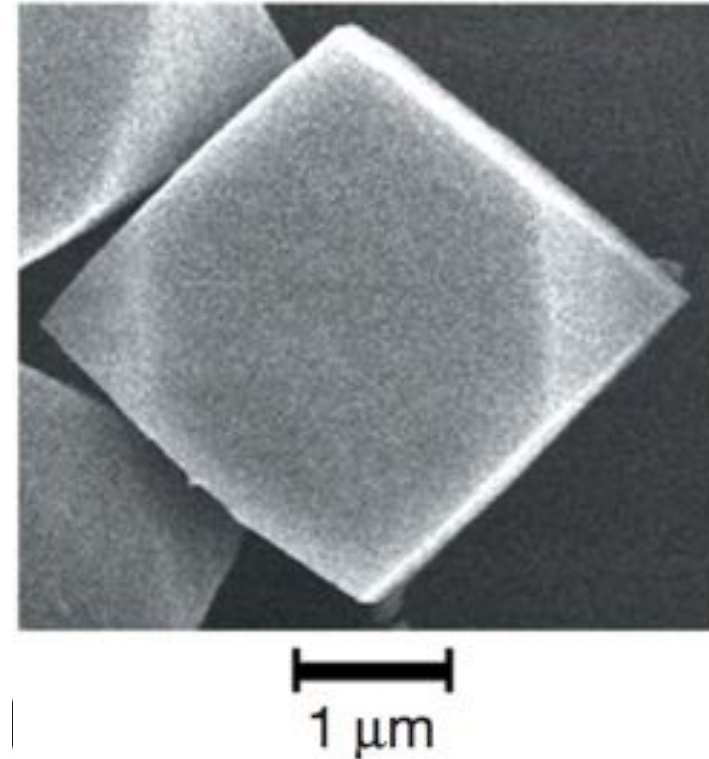


Jochen Müller-Dieckmann  
Xandra Kreplin



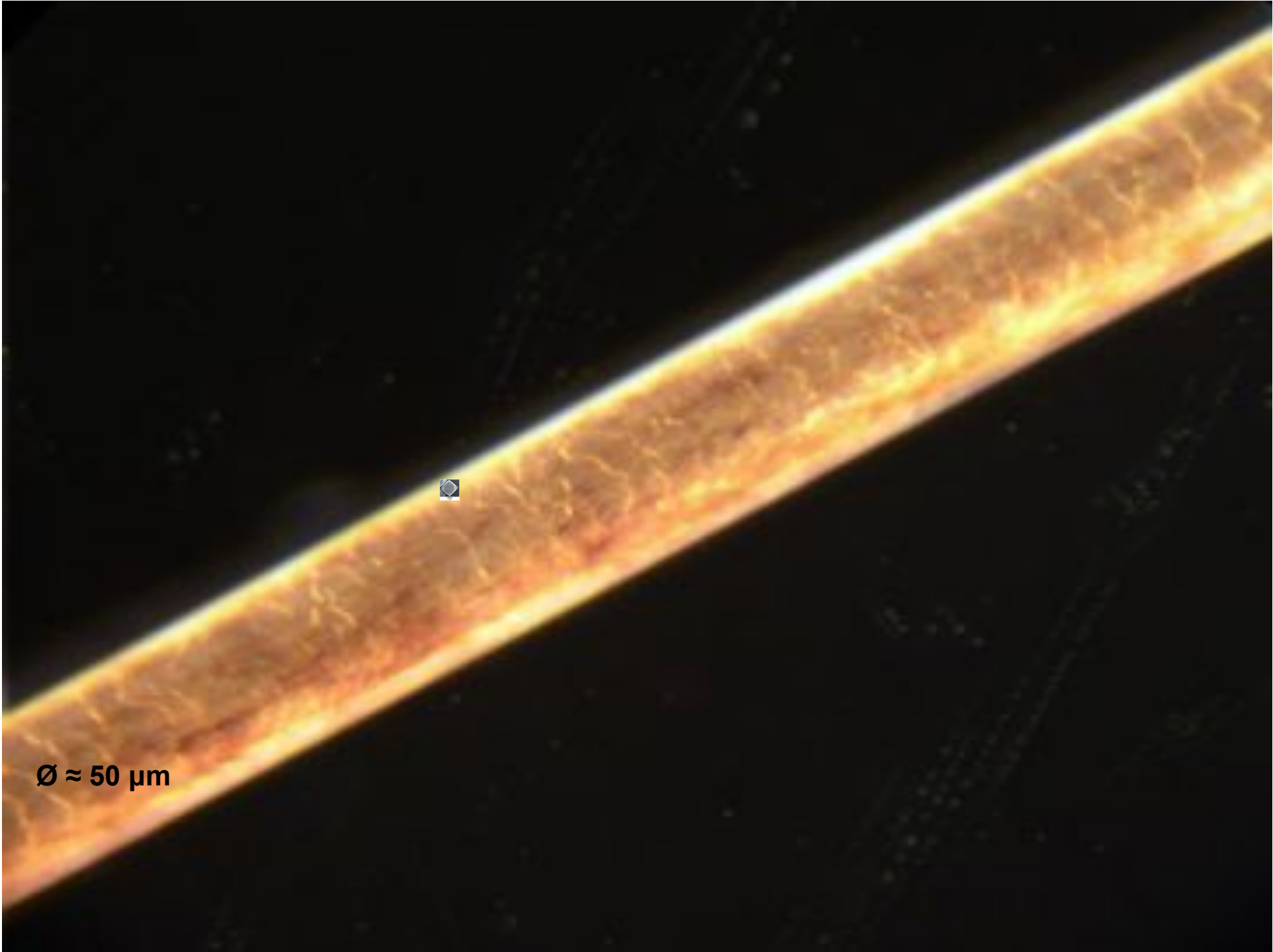
# Small Crystals

- Crystal dimensions of 1-10 micron are not rare.
- Often these crystals are of high quality
- Small and parallel beams needed.

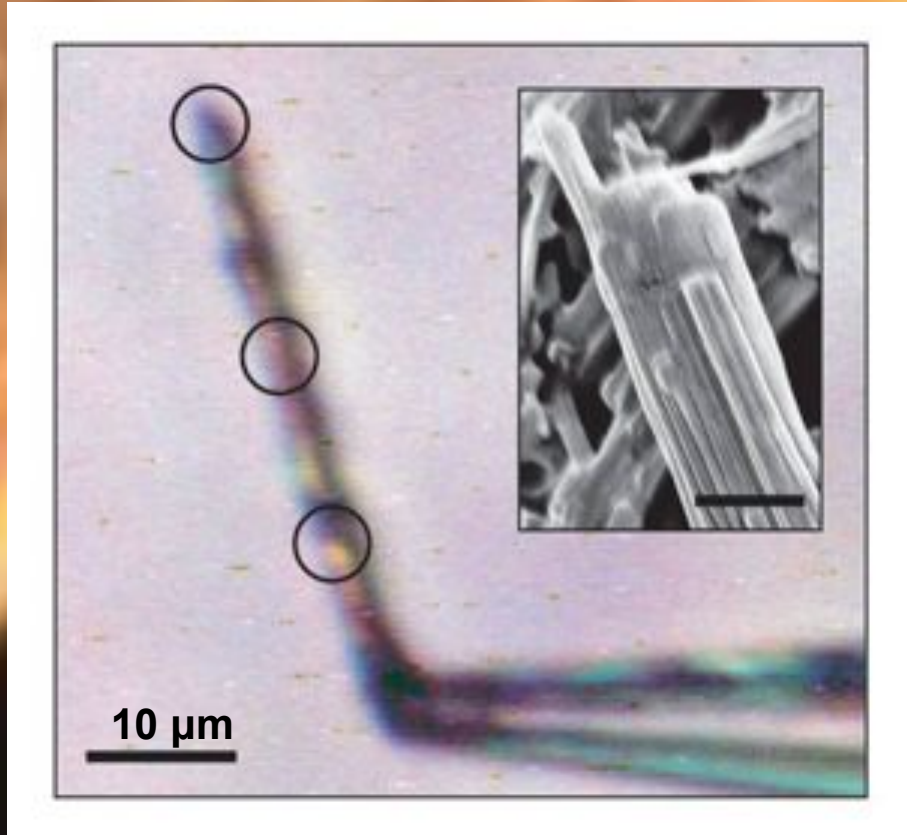
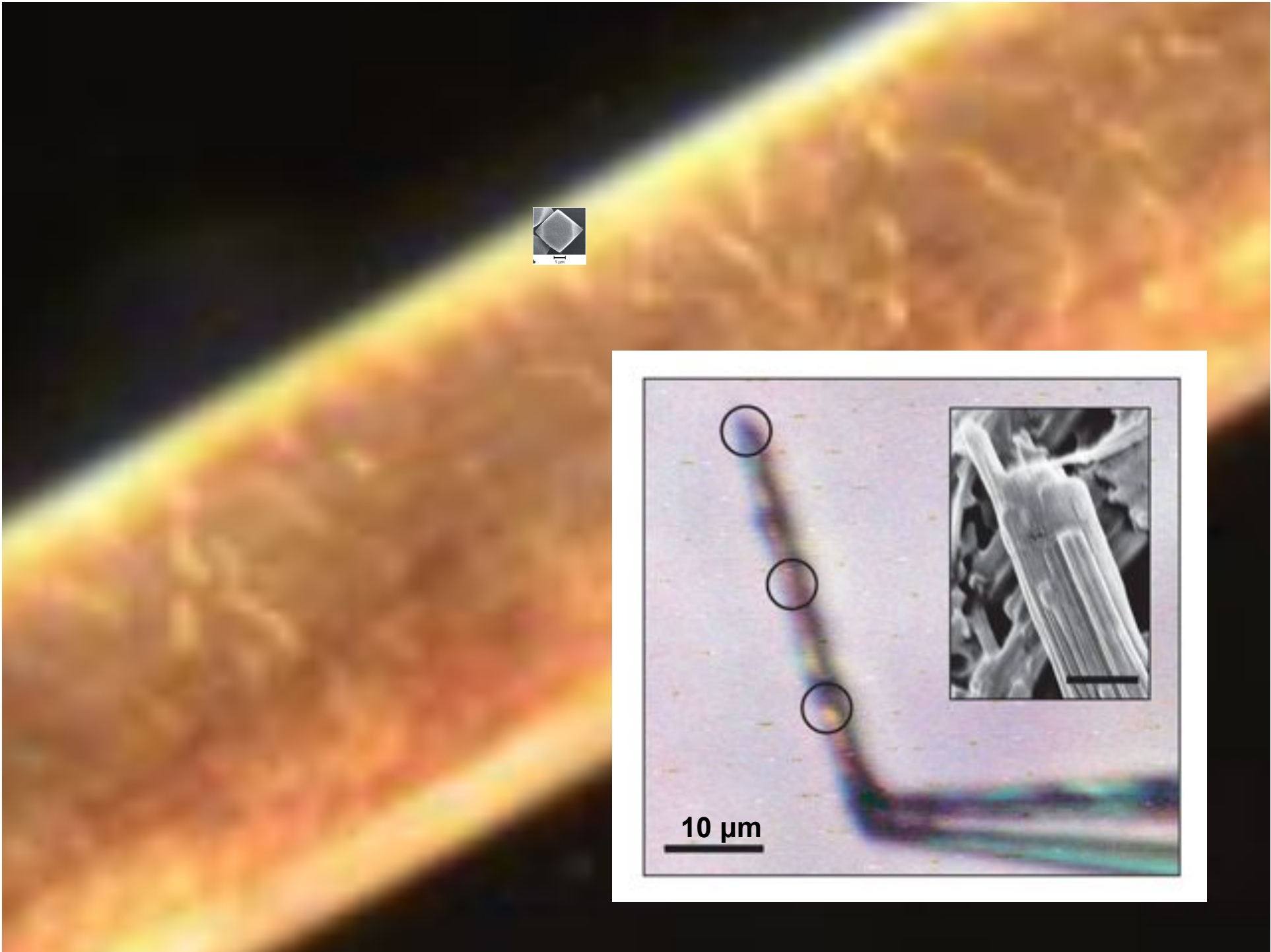


Coulibaly et al. The molecular organization of cypovirus polyhedra. *Nature* (2007) 446: 97-101



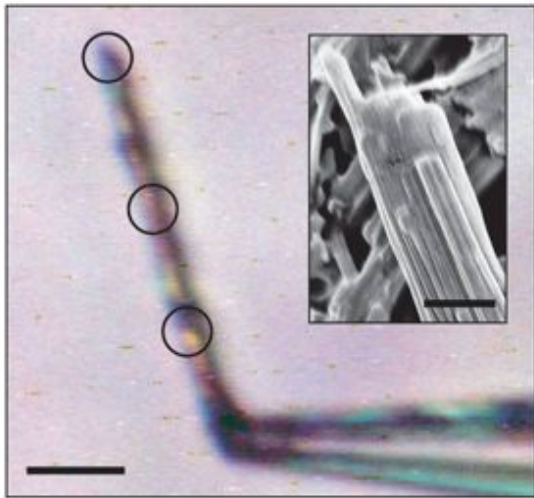


$\text{Ø} \approx 50 \mu\text{m}$

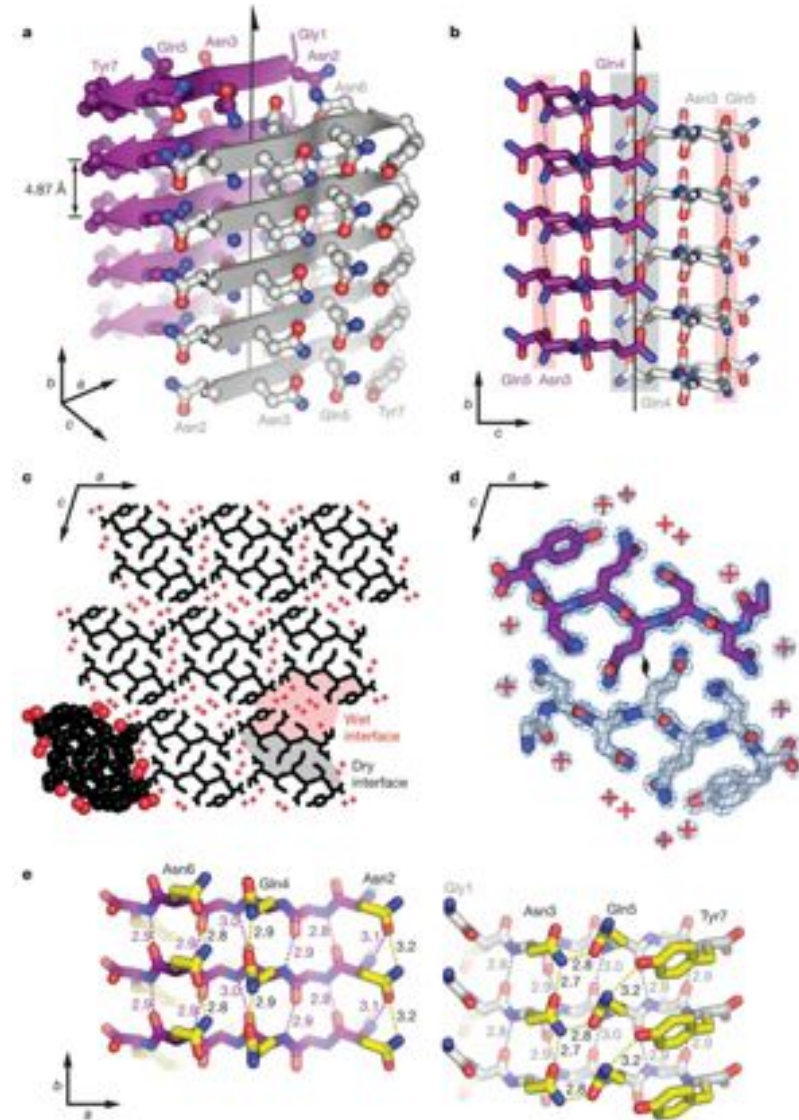


10 μm

# Amyloid fibrils

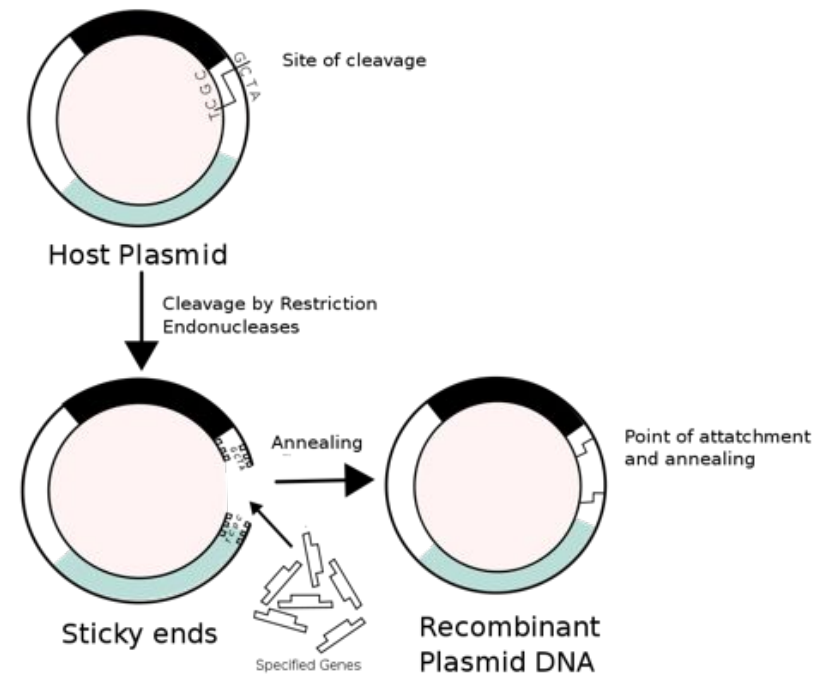
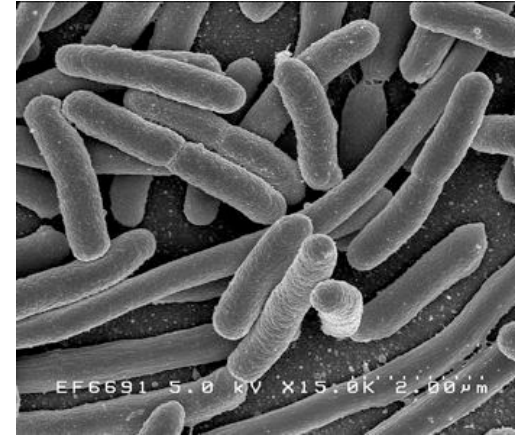


Nelson et al. Structure of the cross-beta spine of amyloid-like fibrils. Nature (2005) 435: 773-8



# Recombinant production of proteins

- Insert DNA from other organism into bacteria (Lobban?, 1972) using enzymes as tools for the manipulation of DNA
- 1982 synthetic human insulin (Genentech & Eli Lilly) entered the market for diabetes therapy.
- Recombinant production (and 'overexpression') of protein molecules is crucial for macromolecular crystallography as large amounts of material are needed to produce crystals.





# Use of synchrotron radiation in Biology

## Synchrotron Radiation as a Source for X-ray Diffraction

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J. WITZ

Laboratoire des Virus des Plantes, Institut de Botanique de la Faculté des Sciences de Strasbourg, Strasbourg

Some preliminary results have been obtained with synchrotron radiation from the 7.5 GeV electron synchrotron Deutsches Elektronen-Synchrotron (DESY) in Hamburg as a source for X-ray diffraction.

When an electron is accelerated it emits radiation. At the very high energies used in DESY, the emitted radiation is confined to a narrow cone about the instantaneous direction of motion of the electron. Thus the synchrotron radiates tangentially. Synchrotron radiation is polychromatic, with a peak in the X-ray region for an electron energy of 7.5 GeV (see ref. 1 for the original theoretical description and refs. 2-4 for experimental details).

The DESY synchrotron uses bursts of 50 pulses and each 10 ns pulse contains  $6 \times 10^{11}$  electrons (10 mA average beam current). The injection energy is relatively low and the electrons are accelerated up to 7.5 GeV in the HRS.

Most of the X-radiation is emitted during the last 3 ns of each pulse; little radiation is produced at the lower electron energies, and so the time averaged intensity at 1.5 Å is about 30% of the peak value.

Table 1 Data for Quartz Monochromator in Synchrotron Radiation Beam

Synchrotron	7.5 GeV, 10 mA beam current
Electron beam diameter	approximately 4 mm (= effective X-ray source diameter)
Distance	37 m from synchrotron to monochromator
Close-fit of the incident beam	approximately $10^{-4}$ rad
Polarization	85% at 1.5 Å in the eighth ns of the cycle, polarized in the plane of the quadrupoles
Be-window	0.5 mm (96 mg cm <sup>-2</sup> )
Crystal	quartz cut at $\alpha = 8^\circ 30'$ to the 1011 plane, dimensions 45 × 13 × 6.3 mm <sup>3</sup>
Bender	pins: outer pair 40.3 mm inner pair 39.3 mm radius of curvature of crystal, 9 m
Wavelength	1.51 Å ( $\theta = 11^\circ 12'$ )
Wavelength spread	$\Delta\lambda = 3 \times 10^{-4}$ Å (due to deviation from Johann focusing and to finite source size)
Focus	1.7 m from crystal line focus 180 μm wide
Angular aperture of reflected beam	horizontal: 2 mrad (convergence) vertical: 3-4 mrad (divergence)
Measured flux in line focus	$1.3 \times 10^{12}$ photons s <sup>-1</sup> mm <sup>-2</sup> (at focal length) (at the eighth ns of the cycle)

Table 2 Biological Applications

Specimen	Elliott fine-focus X-ray tube*	DESY synchrotron with Siemens post-focusing monochromator †
Single crystal	Standard collimator 0.5 mm diameter	
$a = 0.5$ mm	$A = 12.3$ cm	$D = 1$ m
$b = 0.5$ mm	$d = 0.3$ mm	$d = 120$ μm
$L = 1.5$ cm	$F = 10^7$ photons s <sup>-1</sup>	$F = 4 \times 10^7$ photons s <sup>-1</sup>
	$I = 2 \times 10^7$ photons s <sup>-1</sup> mm <sup>-2</sup>	$I = 2.5 \times 10^{11}$ photons s <sup>-1</sup> mm <sup>-2</sup>
Tobacco mosaic virus gel	Double-crystal focusing monochromator ‡	
$a = 0.6$ mm	$d = 80$ μm	$D = 0.8$ m
$b = 1$ mm	$F = 10^7$ photons s <sup>-1</sup>	$d = 100$ μm
$L = 12$ cm	$F = 2 \times 10^7$ photons s <sup>-1</sup> mm <sup>-2</sup>	$F = 3 \times 10^7$ photons s <sup>-1</sup>
		$I = 3 \times 10^{11}$ photons s <sup>-1</sup> mm <sup>-2</sup>
Insect muscle	Double-crystal focusing monochromator ‡	
$a = 3$ mm	$d = 100$ μm	$D = 1.5$ (10) m
$b = 0.3$ mm	$F = 3 \times 10^7$ photons s <sup>-1</sup>	$d = 180$ (150) μm
$L = 40$ cm	$F = 3 \times 10^7$ photons s <sup>-1</sup> mm <sup>-2</sup>	$F = 2 \times 10^7$ (2 × 10 <sup>6</sup> ) photons s <sup>-1</sup>
		$I = 1.5 \times 10^{10}$ photons s <sup>-1</sup> mm <sup>-2</sup>

$a$ , Width of specimen;  $b$ , height of specimen;  $L$ , specimen film distance;  $A$ , anode specimen distance;  $D$ , focal length, that is, monochromator film distance;  $d$ , spot or focus diameter on film;  $F$ , X-ray power reaching the specimen; and  $I$ , flux density at the focus.

\* Loaded with 40 kV, 50 mA into a  $0.2 \times 2$  mm<sup>2</sup> electron focus at the anode in the first case, and 40 kV, 15 mA into a  $0.14 \times 0.7$  mm<sup>2</sup> focus in the other two cases. This set is the most powerful fine-focus X-ray tube currently available.

† The setting of this Johann-type\* monochromator is optimized for each type of specimen.

‡ Conditions of the synchrotron are as in Table 1, computed for 1.5 Å radiation.

We have evaluated the spectral luminance (that is, the power in photons per second radiated per unit area, solid angle, and wavelength interval) of both the synchrotron and a fine-focus rotating anode X-ray tube (see Table 2). The values are  $2 \times 10^{11}$  (time averaged) and  $3 \times 10^{11}$  photons s<sup>-1</sup> sterad<sup>-1</sup> cm<sup>-2</sup> Å<sup>-1</sup> respectively at 1.54 Å, showing clearly that the synchrotron is, relative to present X-ray tubes, a very bright source. The actual advantage to be gained in a diffraction experiment depends critically on the optical system necessary to focus and monochromate the radiation. Three types of focusing mono-

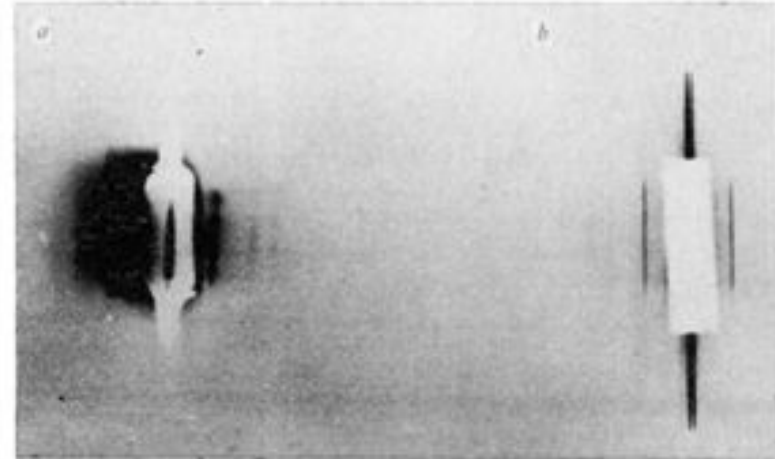


Fig. 3 Equatorial reflexions from dorsolongitudinal flight muscle of *Lethocerus maximus* recorded with: *a*, monochromated synchrotron radiation; electron energy 5 GeV, beam current 8 mA, exposure time 15 min, specimen film distance 40 cm; note the parasitic scattering on the left of the backstop arising from fluorescence from the monochromator holder; *b*, Elliott fine-focus rotating anode tube at 40 kV, 15 mA, exposure time 1 h, specimen film distance 36 cm. The strong line is the 20 reflexion ( $d = 231$  Å); the weak lines are the 21, 31 and 32 reflexions.

# Use of synchrotron radiation in MX

## Applications of synchrotron radiation to protein crystallography: Preliminary results

(x-ray diffraction/anomalous dispersion/rubredoxin/azurin/nerve growth factor/glutaminase-asparaginase)

JAMES C. PHILLIPS, ALEXANDER WLODAWER, MARGUERITE M. YEVITZ, AND KEITH O. HODGSON\*

Department of Chemistry and Stanford Synchrotron Radiation Project, Stanford University, Stanford, California 94305

Communicated by Richard H. Holm, October 23, 1975

**ABSTRACT** X-ray diffraction photographs of protein single crystals have been obtained using synchrotron radiation produced by an electron-positron storage ring. The diffracted intensities observed with this unconventional source are a factor of at least 60 greater than those obtained with a sealed x-ray tube using the same crystal and instrumental parameters. Diffraction data have been collected by the precession method to higher resolution and using smaller protein crystals than would have been possible with a conventional source. The crystal decay rate in the synchrotron beam for several proteins appears to be substantially less than that observed with Ni-filtered Cu radiation. The tunable nature of the source (which allows selective optimization of anomalous contributions to the scattering factors) and the low angular divergence of the beam make the source very useful for single crystal protein diffraction studies.

The use of synchrotron radiation as a source for single crystal x-ray diffraction studies has recently been the subject of considerable discussion and controversy. In contrast to con-

tation of synchrotron radiation produced by the SPEAR electron-positron storage ring at the Stanford Linear Accelerator Center, Stanford, Calif.

Synchrotron radiation is emitted tangentially to the instantaneous path of the charged particles as they are main-

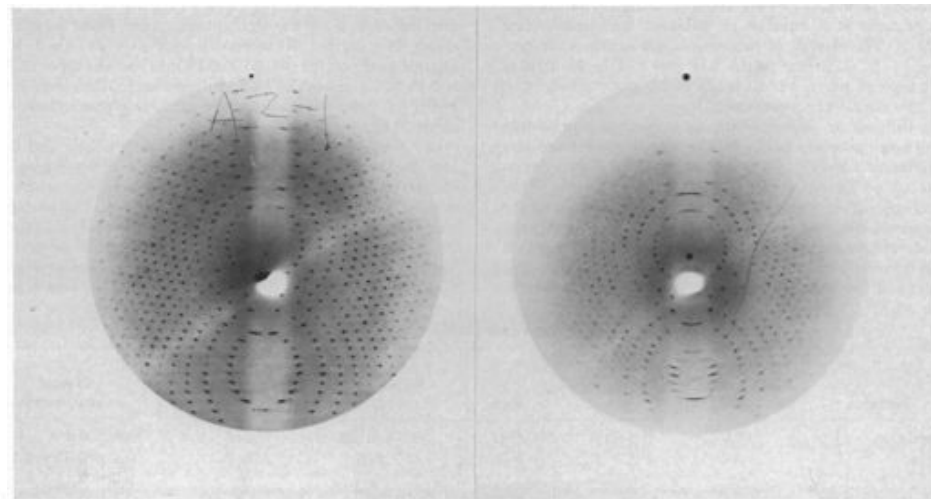


FIG. 3. Cone-axis oscillation photographs of the same azurin crystal. Precession angle  $6.5^\circ$ , oscillation angle  $20^\circ$ . (left) Synchrotron source,  $E = 3.7$  GeV,  $I = 40$  mA, only electrons present,  $\lambda = 1.740$  Å, exposure time 10 min. (right) Philips fine-focus sealed Cu anode tube, operated at 40 kV, 30 mA, exposure time 6 hr, Ni filter.

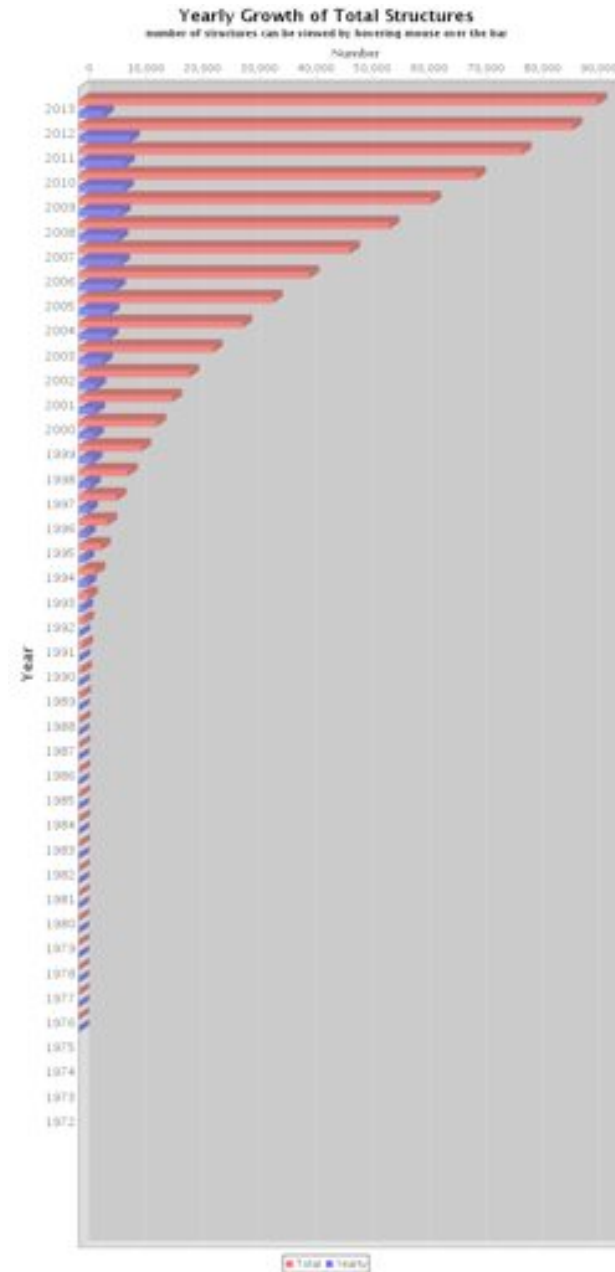
# Detectors

Technology	When	Readout	Remark
X-ray film	1940	30 min	grain size < 10 $\mu\text{m}$
Image Plates	1990	1.5 min	pixel: 150 x 150 $\mu\text{m}^2$ + PSF
CCD detectors	1997	1 sec	pixel: 80 x 80 $\mu\text{m}^2$ + PSF
Pixel-Array Detectors (2010)	2010	3 msec	pixel: 173 x 173 $\mu\text{m}^2$ sharp
Pixel Array Detectors (2014)	2104	??	pixel: 50 x 75 $\mu\text{m}$

- PSF = Point spread function

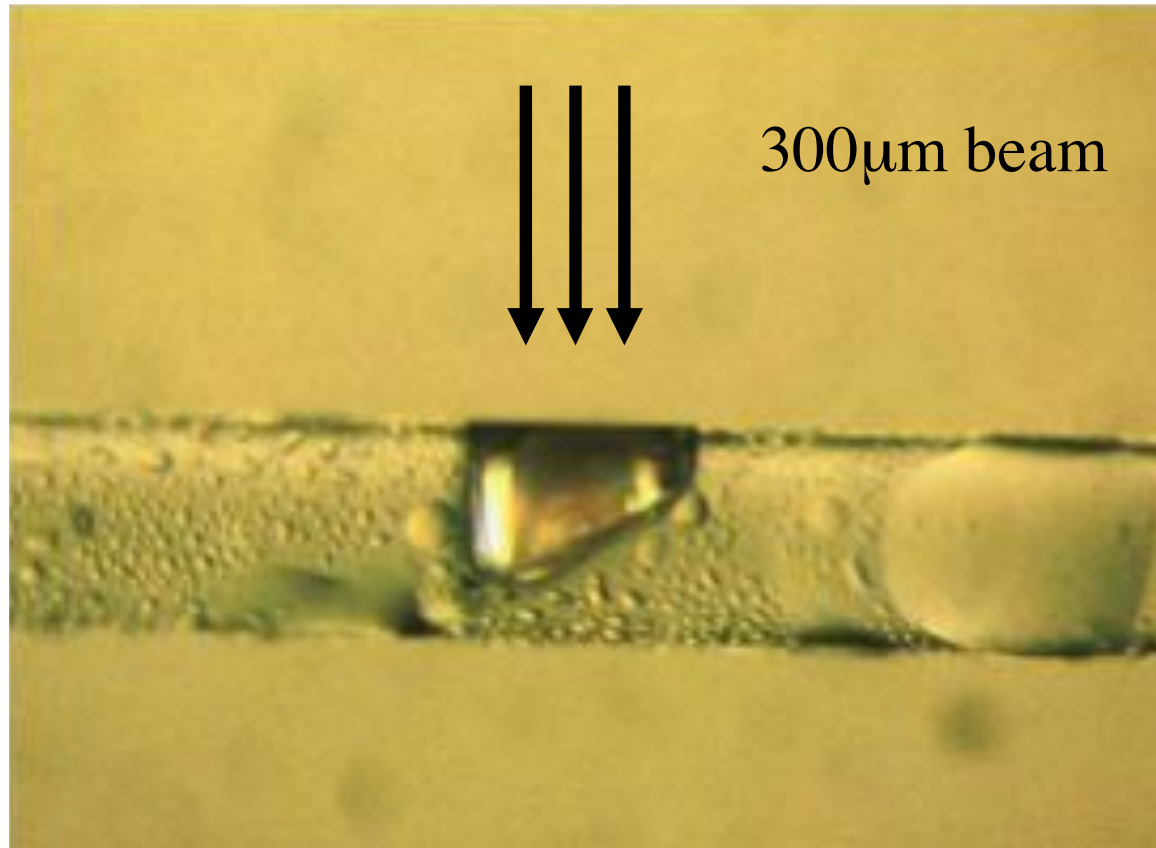
# The Protein Data Bank

- Started as a grass-root movement in the 70's
- As of Tuesday Jun 11, 2013 17:00 PDT, **91359 structures** are in the protein data bank.
- Out of these 80569 were determined by X-ray crystallography
- 1976:13
- 1980:69
- 1990: 507
- 2000: 13596
- 2010: 70013
- A large fraction of structures determined today are determined by 'Molecular Replacement'





# Radiation Damage



Garman & Schneider (1997) J. Appl. Cryst. 30:211

# Data collection at 100 K

- Mounting protein-crystals in a free-standing film revolutionized the field.

Teng, T. Y. (1990)  
J. Appl. Cryst 23: 387-391

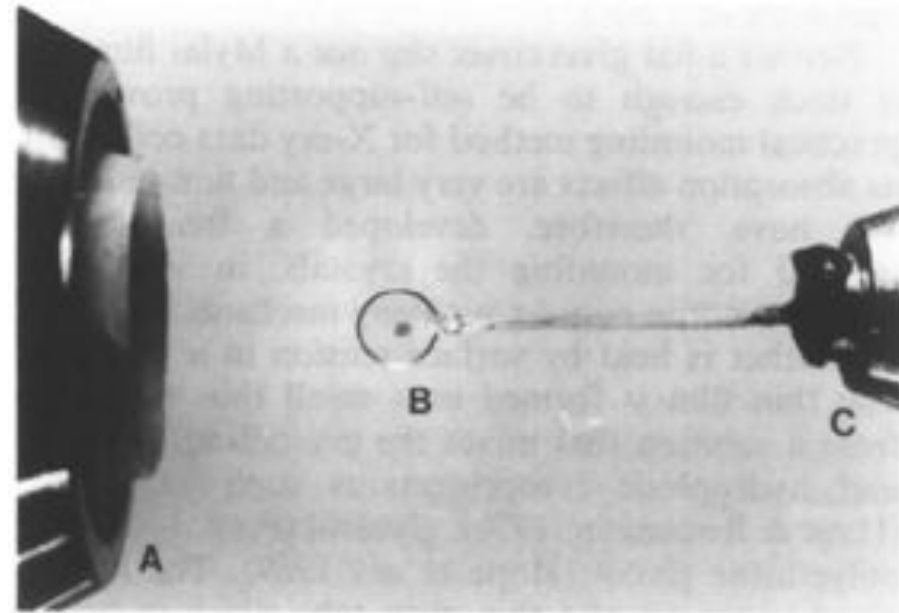
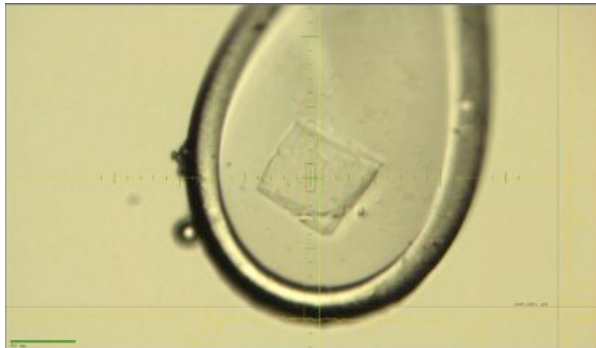
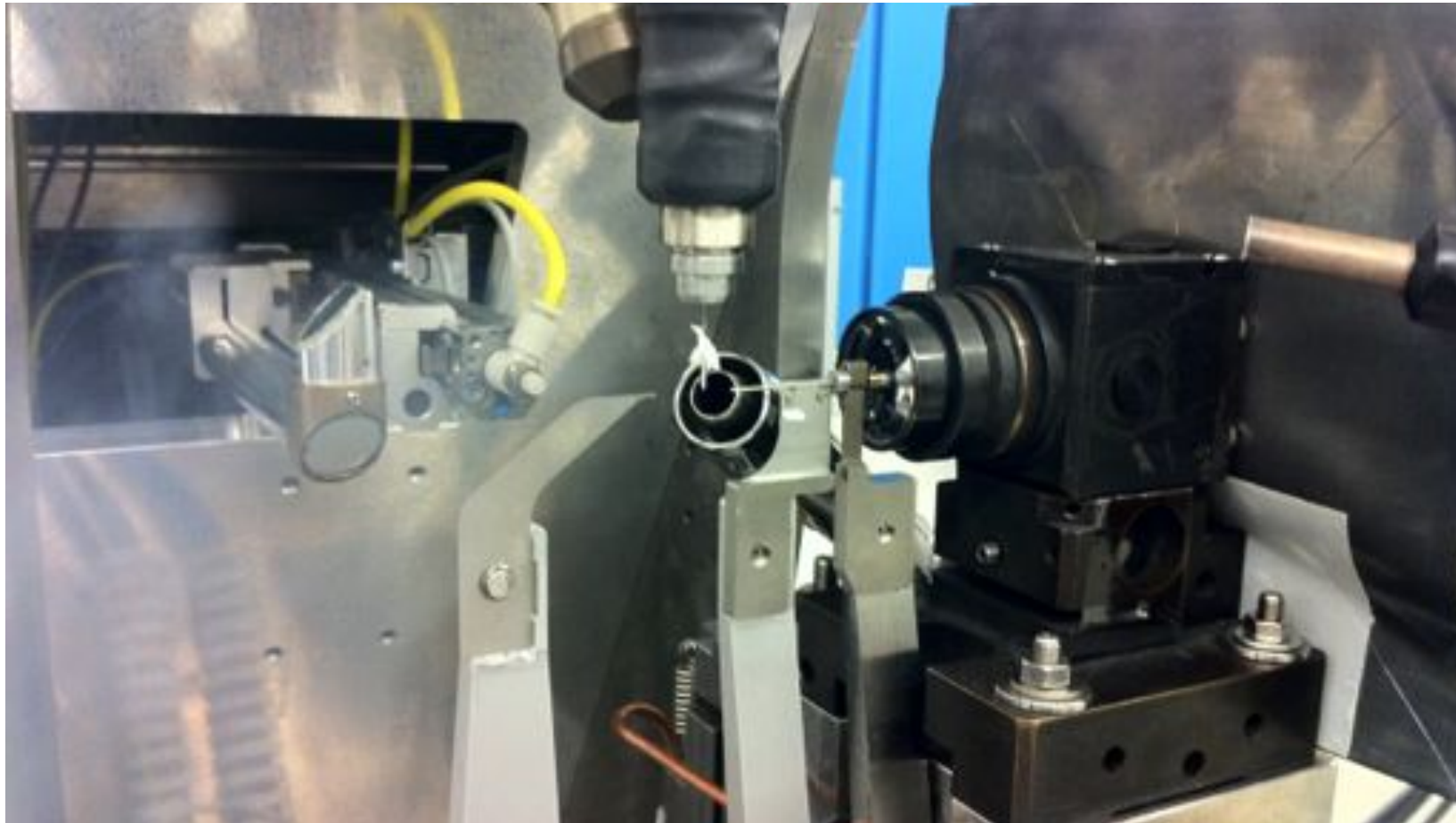


Fig. 2. The loop mounted on an oscillation camera used at CHESS: *A* cold nitrogen nozzle; *B* the loop with a frozen crystal; *C* cold-gas-stream reflector mounted on the goniometer head. The cold nitrogen nozzle (*A*) of a transfer line is 1/4 in in diameter which delivers a gas stream at from 80 to 230 K. Within a cone-shaped working volume of 65 mm<sup>3</sup>, the temperature gradient is less than 4 K, at an initial exit temperature of 85 K. Fog or ice formation around the nozzle, crystal and goniometer head is avoided by a coaxial warm and dry nitrogen stream that surrounds the cold stream, and by a built-in heater on the base of the cold-gas-stream reflector (*C*).

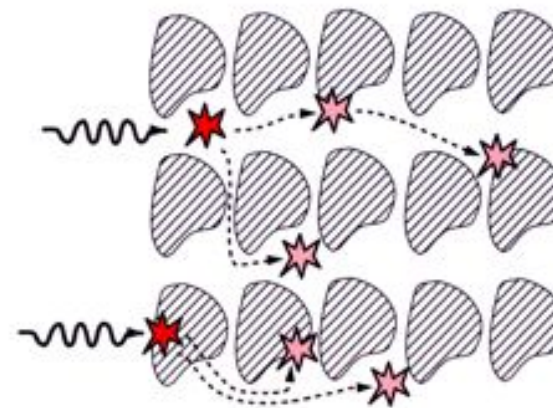
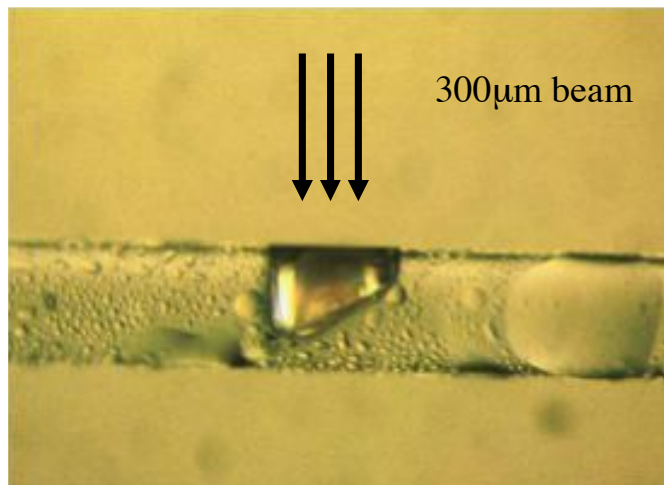
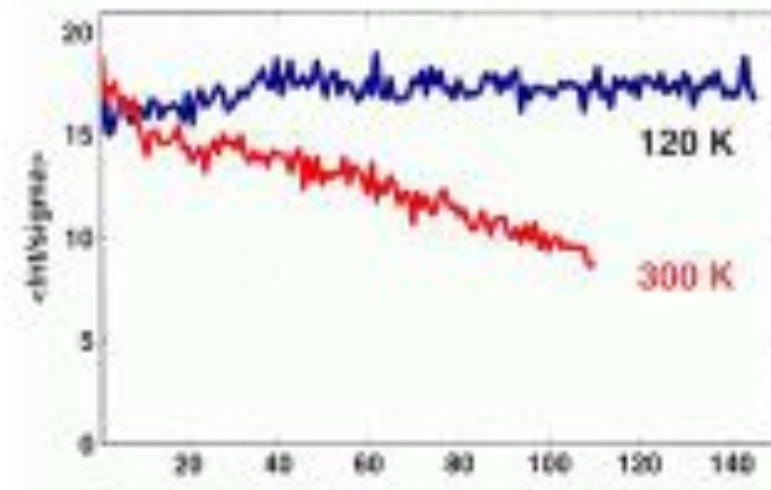
## HPGonioV: Sample environment



- 400 micron thick blade.
- Motorized adjustable distance to sample: 7-36 mm.

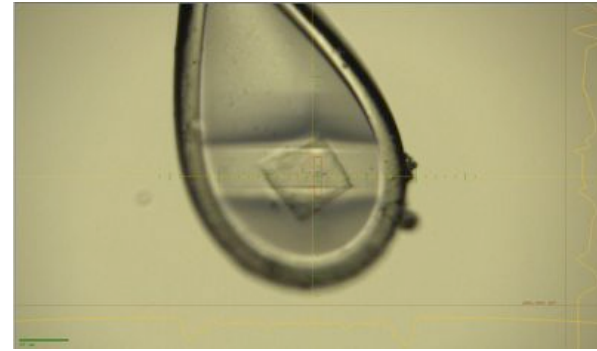
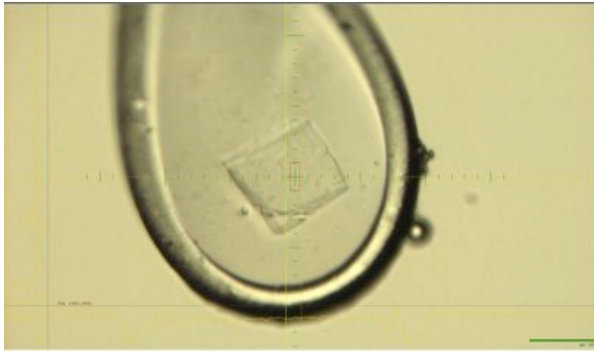
# Radiation Damage

- Data collection at 100 K significantly extends crystal lifetime

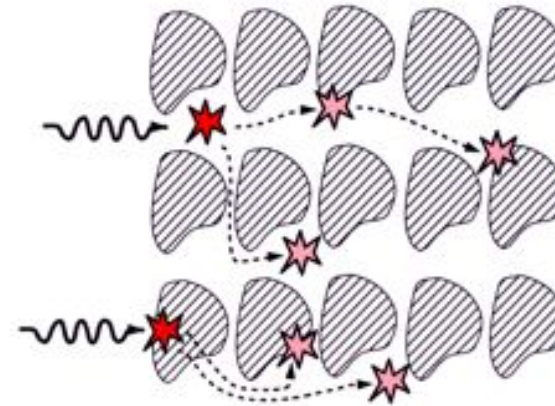


Garman & Schneider (1997) J. Appl. Cryst. 30:211

# Radiation damage at 100 K



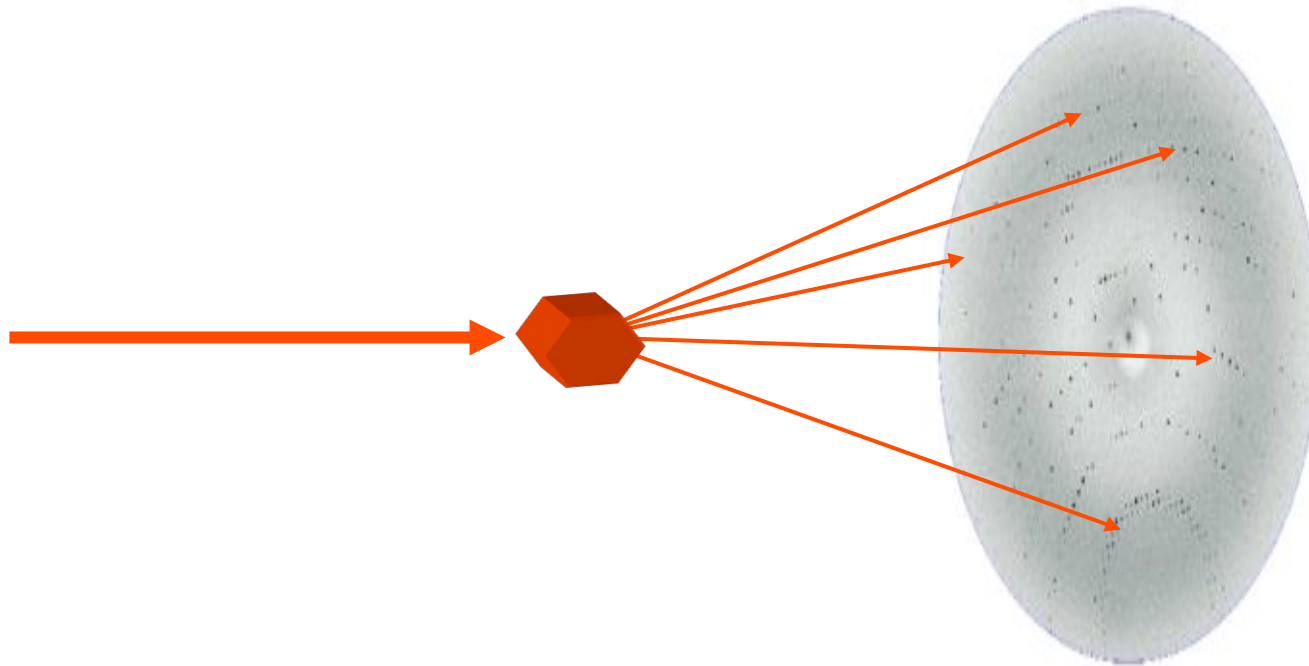
- On modern synchrotron beamlines the lifetime of protein crystals at 100k is on the order of seconds to minutes.
- Going back to Room temperature, Owen et al. (2012) Acta Cryst. D68:810 have shown that 'in the first 200 msec of a room temperature experiment, one can outrun hydroxyl radicals.



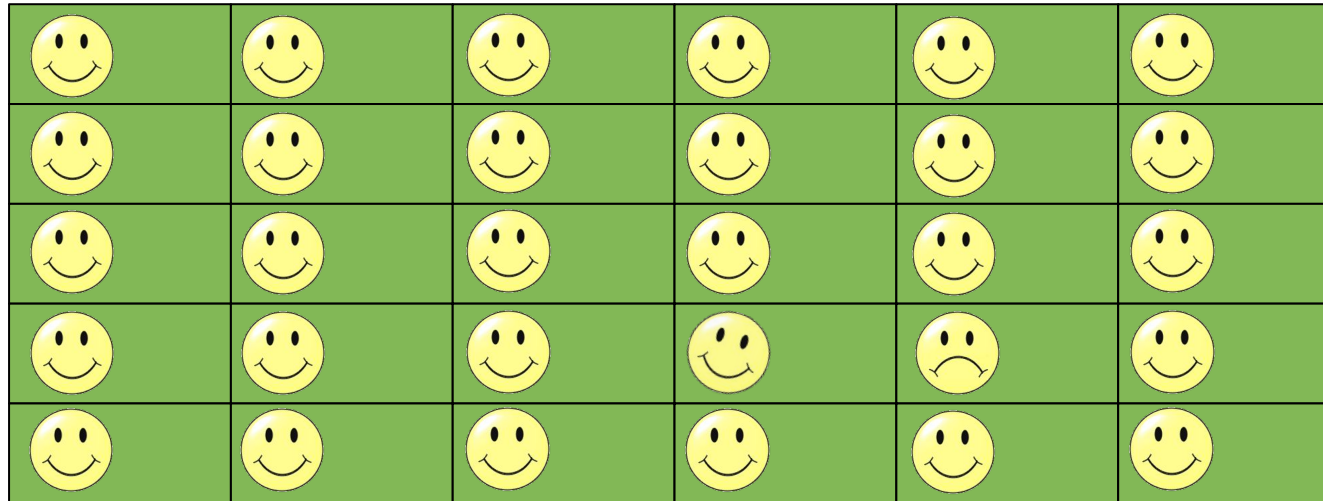
# Crystal Structure Determination



# Diffraction from a Crystal



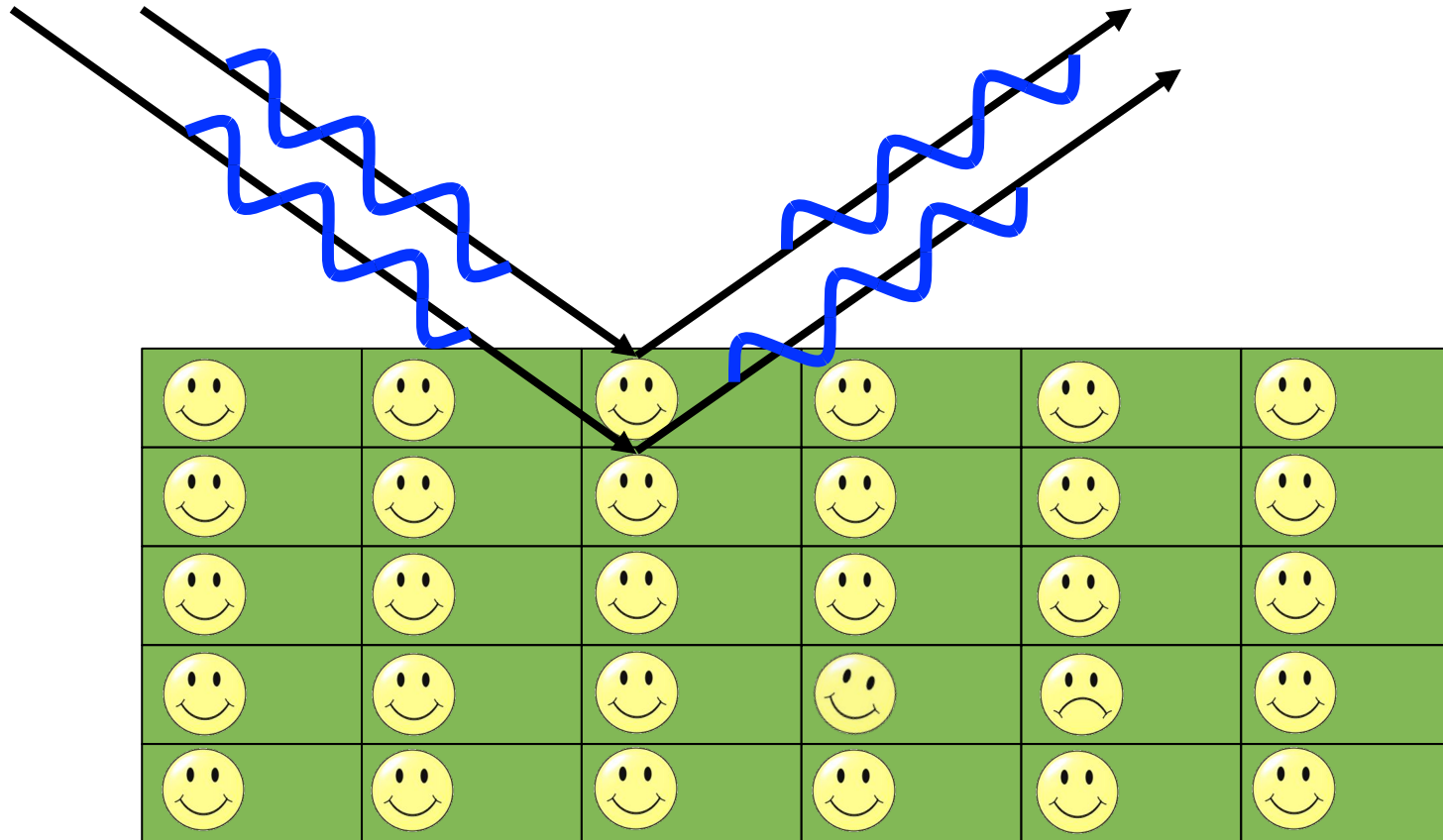
# Inside a crystal



- A crystal consists of repeating units, the crystallographic unit cells.
- Each unit cells has 'the same' content, i.e. the same molecules in the same conformation and in the same orientation
- In real crystals, there is always some amount of 'disorder'



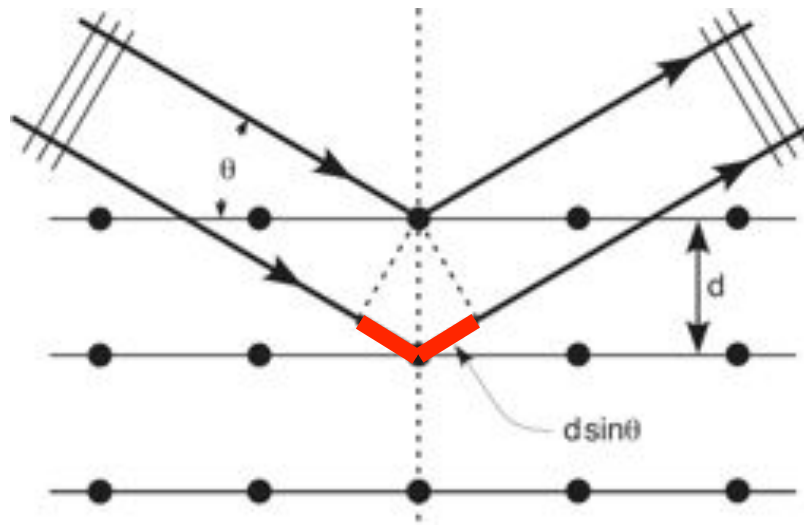
# Diffraction from a crystal



- When electromagnetic waves are interacting with a periodic structure, interference effects will occur.

# Diffracting planes

[http://en.wikipedia.org/wiki/X-ray\\_crystallography](http://en.wikipedia.org/wiki/X-ray_crystallography)



Constructive interference occurs when Bragg's law is fulfilled:

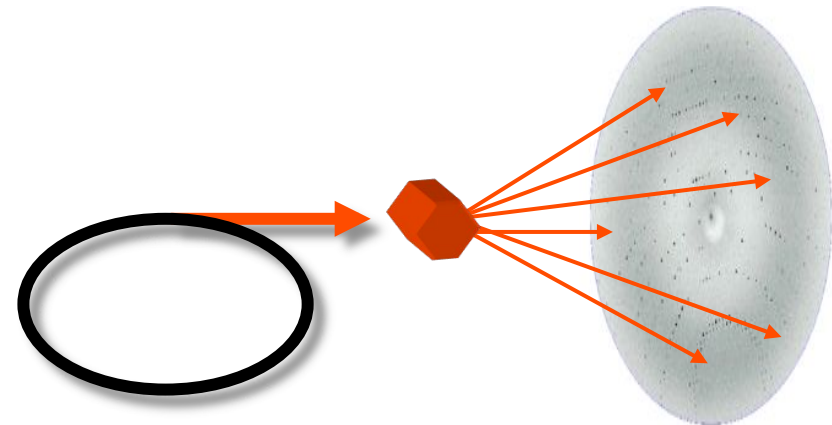
$$2 d \sin\theta = n \lambda$$



Nobel prize for physics 1914 to Max von Laue

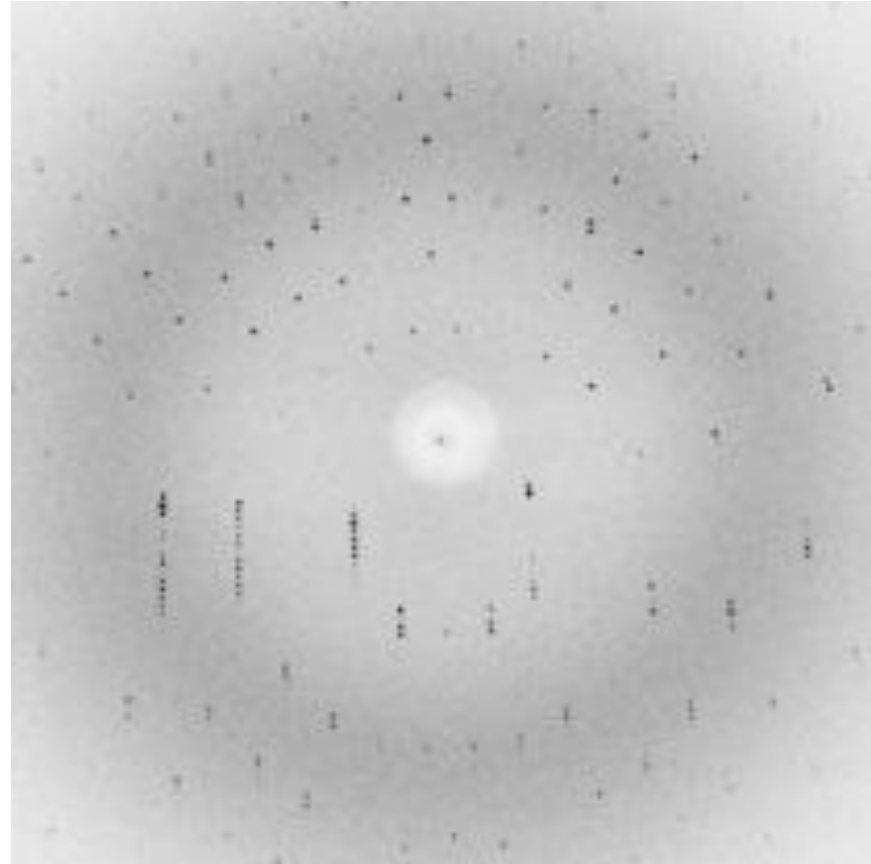


Nobel prize for physics 1915 to William and Lawrence Bragg



# Diffraction Data

- The diffraction pattern changes when the crystal is rotated.
- By rotating the crystal, different Bragg-planes are brought into their diffracting position



[http://www-structmed.cimr.cam.ac.uk/Course/Basic\\_diffraction/data\\_animation.html](http://www-structmed.cimr.cam.ac.uk/Course/Basic_diffraction/data_animation.html)

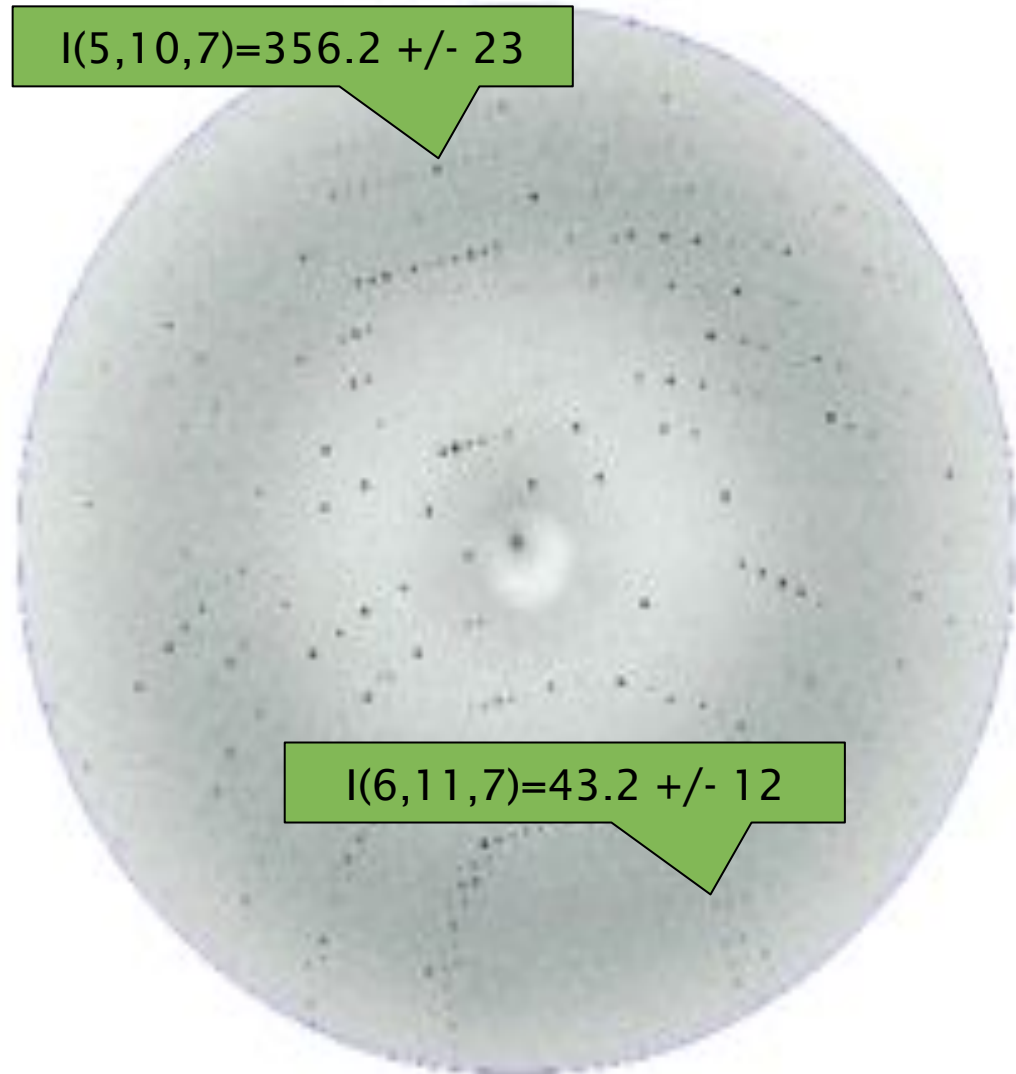
# Diffraction and forests ...



<http://flickr.com/photos/rossogiallobianco/2486114038>

# Indices and Structure factor amplitudes

- Every diffraction spot is marked by an index  $hkl$
- For every diffraction spot an Intensity  $I$  is measured.
- The result of the experiment is an indexed set of  $I$ 's
- Diffraction at low  $2\theta$  is stronger than at high  $2\theta$



# Diffraction data

h	k	l	I	sig(I)
0	20	35	4980.5	122.6
0	20	36	6906.6	216.6
0	20	37	8302.3	231.7
0	20	38	3209.5	89.3
0	20	39	459.6	22.1
0	20	40	1017.4	33.8
0	20	41	-5.6	18.3
0	20	42	33.8	15.6
0	20	43	4545.7	133.4
0	20	44	210.5	19.2
0	20	45	808.8	29.1

# Structure Factor Amplitudes

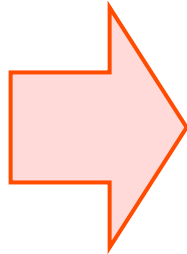
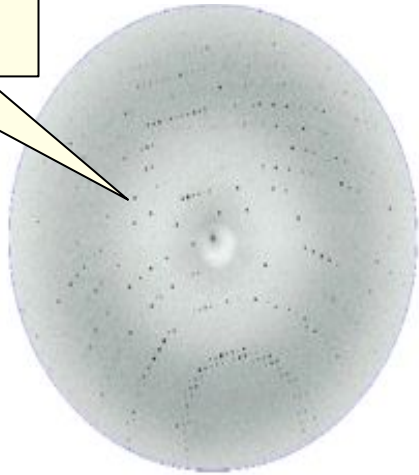
- For formal reasons, the measured I's are usually converted to 'Structure Factor Amplitudes' F by:

$$F = \sqrt{I}.$$

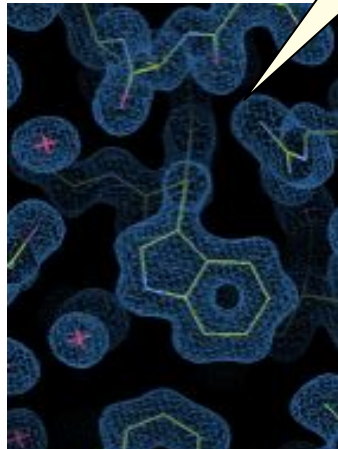


# Calculating Electron density

**Intensity  $I_{hkl}$**  for some set of planes with Miller indices  $hkl$



**Electron density** for all points in the unit cell



$$\rho_{xyz} = \sum_{hkl} |F_{hkl}| e^{-i\varphi_{hkl}} e^{-2\pi i(hx+ky+lz)}$$

**Electron density** at some point  $xyz$  in space

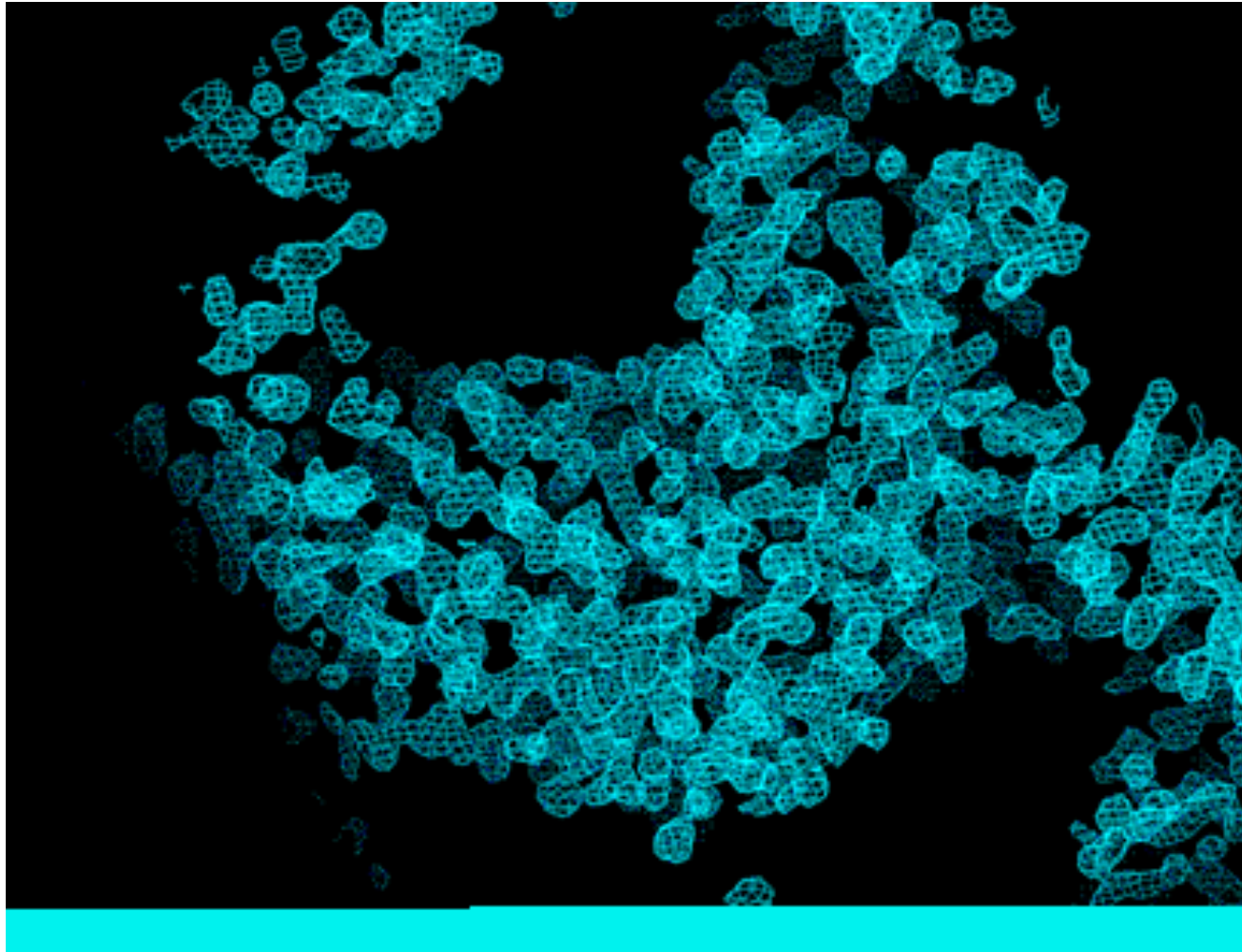
Sum over all Bragg reflections

**Structure Factor Amplitude** for each reflection  $hkl$ .  $F_{hkl} \approx \sqrt{I_{hkl}}$

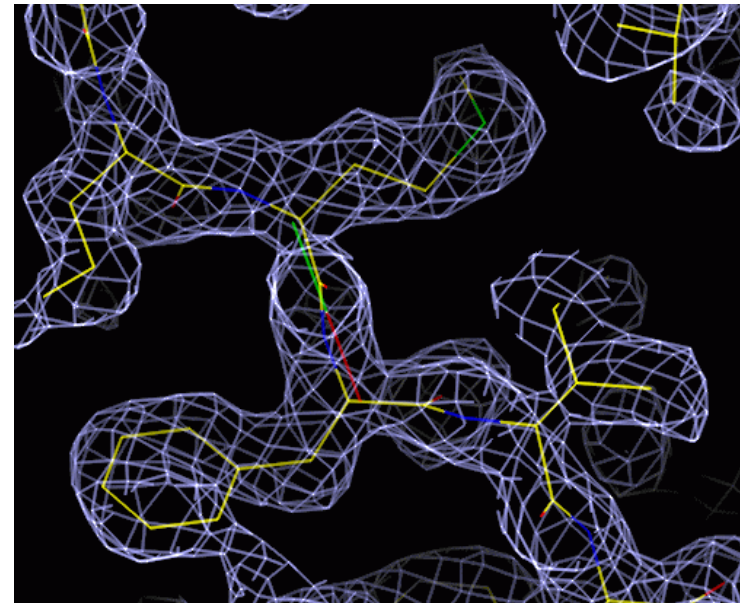
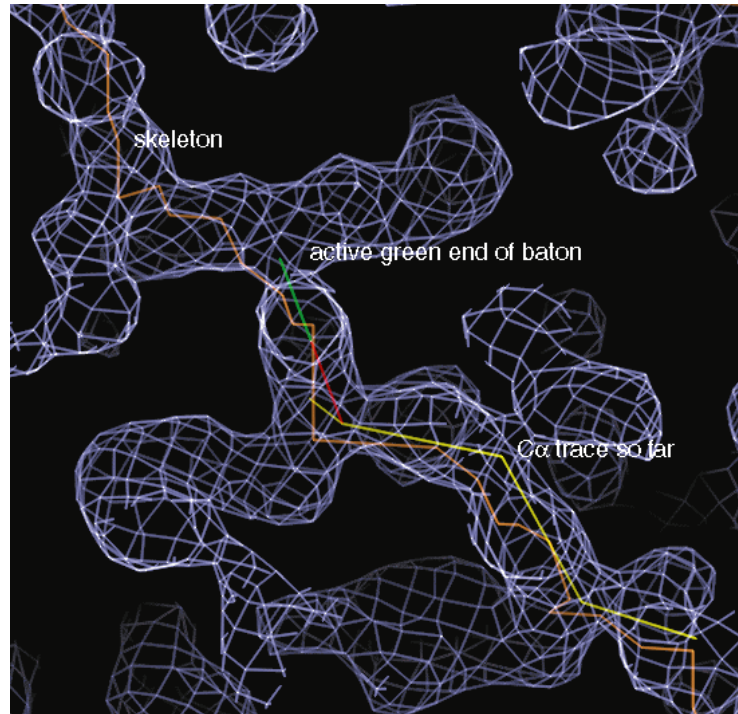
**Structure Factor Phase** for each reflection  $hkl$ .

**Phase shift** depending on  $hkl$  and position in space

# Typical initial electron density map



# Interpretation of the electron density map



# A 'pdb'-file (www.rcsb.org)

```
HEADER      TRANSFERASE                      06-MAR-03   10PJ
TITLE      STRUCTURAL BASIS FOR THE AUTO-INHIBITION OF C-ABL TYROSINE
TITLE      2 KINASE
.
.
ATOM       1  N   ALA A 243      20.064  -2.529  43.315  1.00  51.64      N
ATOM       2  CA  ALA A 243      19.658  -1.370  42.459  1.00  51.36      C
ATOM       3  C   ALA A 243      20.832  -0.838  41.643  1.00  50.91      C
ATOM       4  O   ALA A 243      20.650   0.010  40.776  1.00  50.77      O
ATOM       5  CB  ALA A 243      19.092  -0.263  43.322  1.00  50.20      C
ATOM       6  N   MET A 244      22.030  -1.350  41.906  1.00  50.62      N
ATOM       7  CA  MET A 244      23.218  -0.888  41.195  1.00  50.73      C
ATOM       8  C   MET A 244      23.537  -1.709  39.950  1.00  50.43      C
ATOM       9  O   MET A 244      24.554  -1.488  39.298  1.00  50.33      O
ATOM      10  CB  MET A 244      24.420  -0.882  42.146  1.00  51.32      C
ATOM      11  CG  MET A 244      24.215   0.020  43.361  1.00  52.32      C
ATOM      12  SD  MET A 244      25.597   0.070  44.515  1.00  54.49      S
ATOM      13  CE  MET A 244      26.730   1.130  43.627  1.00  53.37      C
ATOM      14  N   ASP A 245      22.662  -2.653  39.620  1.00  49.87      N
ATOM      15  CA  ASP A 245      22.857  -3.505  38.446  1.00  49.28      C
ATOM      16  C   ASP A 245      22.115  -2.895  37.257  1.00  47.55      C
```